Neural crest–derived cells with stem cell features can be traced back to multiple lineages in the adult skin

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Given their accessibility, multipotent skin-derived cells might be useful for future cell replacement therapies. We describe the isolation of multipotent stem cell–like cells from the adult trunk skin of mice and humans that express the neural crest stem cell markers p75 and Sox10 and display extensive self-renewal capacity in sphere cultures. To determine the origin of these cells, we genetically mapped the fate of neural crest cells in face and trunk skin of mouse. In whisker follicles of the face, many mesenchymal structures are neural crest derived and appear to contain cells with sphere-forming potential. In the trunk skin, however, sphere-forming neural crest–derived cells are restricted to the glial and melanocyte lineages. Thus, self-renewing cells in the adult skin can be obtained from several neural crest derivatives, and these are of distinct nature in face and trunk skin. These findings are relevant for the design of therapeutic strategies because the potential of stem and progenitor cells in vivo likely depends on their nature and origin.

Introduction

Embryonic, fetal, and adult tissues are used as sources to investigate the developmental and therapeutic potential of stem cells. Because of their accessibility and the possibility that the patient could act as a stem cell donor, adult stem cells from the skin have received particular attention (Slack, 2001). Apart from multipotent epithelial stem cells that form hair follicles, sebaceous glands, and epidermis (Taylor et al., 2000; Oshima et al., 2001; Blanpain et al., 2004; Claudinot et al., 2005) and so-called melanocyte stem cells that generate pigmented cells (Nishimura et al., 2002), a multipotent cell dubbed skin-derived precursor cell (SKP) has been isolated from both the murine and human skin (Toma et al., 2001). SKPs have the potential to produce in vitro cell types normally not found in the skin, such as neuronal cells. Subsequently, several laboratories reported the existence of self-renewing cells present in the skin of mice, pigs, and humans and able to differentiate in vitro into cells expressing neuronal, glial, osteoblast, chondrocyte, smooth muscle, melanocyte, and adipocyte lineage markers (Belicchi et al., 2004; Dyce et al., 2004; Joannides et al., 2004; Sieber-Blum et al., 2004; Amoh et al., 2005; Shih et al., 2005; Toma et al., 2005).

The formation of cells normally not present in skin might be due to transdifferentiation, which describes the conversion of a cell type of a specific tissue lineage into a cell type of another lineage (Wagers and Weissman, 2004). Alternatively, cells from a given lineage might dedifferentiate into a more naive state that allows the cell to redifferentiate along new lineages. Finally, multipotent cells with stem cell features might persist until adulthood, able to generate a broad variety of cells, depending on their environment. To distinguish among these possibilities, the origin and nature of the cell in question has to be determined and its developmental potential has to be analyzed at the single cell level (Wagers and Weissman, 2004).
The developmental origin and exact localization of skin cells giving rise to neural and nonneural progeny is unclear in many of the reported cases. Multipotent skin-derived cells have been enriched by means of markers found on hematopoietic stem cells (Belicchi et al., 2004) or have been isolated from transgenic animals expressing GFP from promoter elements of Nestin (Amoh et al., 2005), a gene also expressed in neural progenitor cells. One source that has been associated with sphere-forming SKPs is the dermal papilla from whisker follicles (Fernandes et al., 2004). Whisker follicles are large hair follicles of the face that serve as sensory organs for a wide range of mammals, excluding humans. Genetic in vivo cell fate mapping revealed that the dermal papilla of these follicles is of neural crest origin (Fernandes et al., 2004). Similarly, culturing explants of bulge and dermal sheath of whisker follicles allowed the identification of neural crest–derived multipotent cells in the upper part of the whisker follicle (Sieber-Blum and Grim, 2004; Sieber-Blum et al., 2004).

A neural crest origin might explain the multipotency of at least some stem and progenitor cells in the skin. Indeed, the neural crest contributes during vertebrate development to a variety of tissues, including the peripheral nervous system and nonneural cell types such as melanocytes in the skin (Le Douarin and Dupin, 2003). Clonal analysis revealed that multipotent, self-renewing neural crest stem cells (NCSCs) cannot only be isolated from migratory neural crest but also from different tissues at later stages and even from the adult organism (Stemple and Anderson, 1992; Bixby et al., 2002; Kruger et al., 2002). Thus, it is conceivable that apart from the whisker follicle, other neural crest–derived compartments in the skin might contain multipotent neural crest–derived cells.

Results

p75/Sox10-positive neural crest–derived cells with stem cell properties can be isolated from the adult murine and human skin

Floating sphere cultures have previously been used to identify self-renewing cells in both murine and human skin (Toma et al., 2001, 2005; Belicchi et al., 2004; Fernandes et al., 2004; Ioannides et al., 2004). To further characterize sphere-forming cells derived from the trunk skin of adult mice, dorsal and ventral skin biopsies comprising both dermis and epidermis were dissected, dissociated, and cultured, and formation of spheres was observed within 4–7 d of culture. These spheres could be passaged for several months without overt morphological changes (Fig. 1 A), pointing to the self-renewing capacity of cells present in the spheres. Intriguingly, unlike SKPs enriched by marker selection (Belicchi et al., 2004) or cultured in slightly different conditions than used here (Fernandes et al., 2004), 100% of all primary, secondary, and later passage spheres generated from mouse trunk skin (n > 50 spheres) contained cells expressing the low-affinity neurotrophin receptor p75 and the transcription factor Sox10, both markers for NCSCs (Fig. 1, C and D; Stemple and Anderson, 1992; Paratore et al., 2001). In spheres passaged >20 times, 67.0 ± 10.5% of all cells expressed p75, 76.6 ± 4.5% of all cells expressed Sox10, and 58.6 ± 10.5% of all cells were double positive for p75 and Sox10. 15.0 ± 6.2% of all cells were negative for these markers, pointing to a cellular heterogeneity within skin-derived spheres, as also observed in sphere cultures from other tissues (Reynolds and Rietze, 2005). Thus, skin-derived cells expressing NCSC markers can be propagated in culture for prolonged time periods.

Similarly, spheres readily formed from dissociated surgical samples of adult human thigh and face skin (Fig. 1 B). These spheres could be expanded by passaging, such that after 3 mo >106 cells had been generated from a 16-cm2 skin sample used as starting material. Similar to mouse cultures, all spheres contained p75/Sox10-positive cells, which accounted for >60% of all cells (Fig. 1, D and F). However, other markers for pre-migratory or migratory NCSCs, such as Sox9 and HNK-1, were not expressed.

As p75 and Sox10 are markers for NCSCs (Stemple and Anderson, 1992; Paratore et al., 2001), we next examined whether the mouse trunk skin–derived spheres originate from the neural crest. The fate of neural crest cells was mapped in vivo by mating ROSA26 Cre reporter (R26R) mice, which express β-galactosidase upon Cre-mediated recombination, with mice expressing Cre recombinase under the control of the Wnt1 promoter (Jiang et al., 2000; Lee et al., 2004). In Wnt1-Cre/R26R double-transgenic mice, virtually all NCSCs express β-galactosidase (Brault et al., 2001; Lee et al., 2004). Importantly, despite the transient expression of Cre recombinase, the progeny...
of neural crest cells continue to express β-galactosidase because of the genomic recombination event. Anti–β-galactosidase antibody staining revealed that all primary and late passage spheres generated from the back skin of adult Wnt1-Cre/R26R double-transgenic mice were composed of neural crest–derived cells (Fig. 2 and not depicted). In particular, 100% of all p75-positive cells coexpressed β-galactosidase, as revealed by a typical punctuated staining pattern (Lutolf et al., 2002). Because 87.3 ± 6.0% of all p75-positive cells also expressed Sox10 (three independent experiments with spheres obtained after 20–35 passages), the data demonstrate that sphere-forming, Sox10-expressing cells from the adult mouse skin are neural crest derivatives.

To test the developmental potential of sphere cells derived from murine and human skin, spheres containing p75/Sox10-positive neural crest cells were allowed to differentiate at high cellular density. The formation of glia expressing glial fibrillary acidic protein (GFAP), βIII tubulin (TuJ1)–positive neuronal cells, and smooth muscle actin (SMA)–expressing nonneural cells was readily detectable in both mouse and human cell cultures (Fig. 3, A–C and G–I), although the number of neuronal cells generated was highly variable and low in comparison to that of glia and smooth muscle cells. Upon addition of ascorbic acid and bone morphogenetic protein (BMP) 2, the generation of chondrocytes was observed (Fig. 3, D and J), whereas treatment with stem cell factor and endothelin-3 resulted in formation of a few melanocytes (Fig. 3, E and K). Finally, occasional adipocytes were detected (Fig. 3, F and L). However, we never observed the generation of keratinocytes as assessed by staining with a pan-keratin antibody (unpublished data), demonstrating that neural crest–derived sphere-forming cells are distinct from epithelial stem cells of the skin.

The aforementioned data are consistent with the idea that skin-derived spheres contain multipotent cells capable of generating neural and nonneural cell types. In analogy to NCSCs isolated from other stages and locations, it is likely that this broad potential is inherent to the p75/Sox10-expressing neural crest–derived cells found in the spheres. To address this hypothesis, we plated cells from mouse trunk skin–derived spheres at clonal density and prospectively identified and mapped single undifferentiated, unpigmented p75-positive clone founder cells (Stemple and Anderson, 1992; Hagedorn et al., 1999; Lee et al., 2004; Kleber et al., 2005). The clone founder cells were then incubated in culture conditions permissive for neurogenesis, gliogenesis, and nonneural cell formation (Stemple and Anderson, 1992). 57.9% of all p75-positive founder cells were at least tripotent, giving rise to clones consisting of neural and nonneural cell types (Fig. 4 A). Virtually no p75-positive cell was restricted to a single cell lineage. Thus, p75/Sox10-positive neural crest–derived cells prepared from the adult trunk skin are multipotent and can be expanded in culture. Upon isolation, these cells therefore exhibit properties of NCSCs.

Several instructive growth factors, including Wnt, BMP, neuregulin (NRG), and TGFβ, have been shown to promote specific fate decisions in NCSCs at the expense of other possible fates. Although Wnt responsiveness is lost at later developmental stages (Kleber et al., 2005), postmigratory NCSCs isolated from various structures maintain their responsiveness to BMP2, NRG1, and TGFβ, although the biological activity of these factors changes with time and location (Bixby et al., 2002;
Multiple sources of sphere-forming neural crest-derived cells in the whisker follicle

Apart from back skin–derived p75/Sox10-positive multipotent cells (Figs. 1 and 2), the neural crest origin of sphere-forming cells in the adult skin has been demonstrated for whisker follicle–derived SKPs, which, however, are negative for the NCSC markers p75 and Sox10 (Fernandes et al., 2004). This could either reflect differential regulation of NCSC markers in the same cell type because of varying culture conditions or indicate sphere-forming capacity of skin cells from different neural crest derivatives. To address this issue, we first mapped neural crest derivatives in the adult skin and investigated which of these neural crest derivatives express the NCSC marker Sox10 in vivo. We initially focused on the whisker follicle because this structure has been identified before as a source of multipotent neural crest–derived cells (Fernandes et al., 2004; Sieber-Blum et al., 2004). In the head, the neural crest contributes to many mesenchymal structures (Santagati and Rijli, 2003). Thus, many mesenchymal structures in whisker follicles isolated from Wnt1-Cre/R26R double-transgenic mice expressed β-galactosidase (Fig. 5 A). In particular, the capsule, the ringwulst, the dermal sheath, and, as previously published (Fernandes et al., 2004; Sieber-Blum et al., 2004), the dermal papilla turned out to be neural crest derived. The neural crest origin of all these structures was confirmed by fate mapping experiments performed in human tissue plasminogen activator (Ht-PA)-Cre/R26R mice, in which Cre recombinase is expressed in neural crest cells independently from Wnt1 promoter activity (Pietri et al., 2003; Fig. 5 B). As revealed by X-gal staining of whisker follicles isolated from Sox10\(^{lacZ}\) mice (that express β-galactosidase from the Sox10 locus; Britsch et al., 2001), capsula, ringwulst, and dermal papilla did not express Sox10 in vivo, whereas the dermal sheath, glial cells in nerve endings, and melanocytes were Sox10 positive (Fig. 5 C). Thus, the whisker follicle comprises various Sox10-positive and -negative tissues of neural crest origin.

To investigate which of these neural crest derivatives contain cells with sphere-forming potential, dermal papilla, capsula, the upper part of the dermal sheath (without the bulge), and the lower part of the dermal sheath were isolated from whiskers of adult Wnt1-Cre/R26R double-transgenic mice by microdissection, dissociated, and cultured in the same conditions as used before for trunk skin–derived multipotent neural crest cells. In addition, rat whiskers were used to dissect the ringwulst, which in mice was too small to be isolated without contamination from other tissues. Strikingly, all these whisker follicle structures appear to harbor cells with the capacity to generate spheres (Fig. 5 D). X-gal staining of Wnt1-Cre/R26R mouse cell cultures confirmed that the spheres were neural crest derived. Therefore,
neural crest cells with sphere-forming potential are not confined to a particular niche in the whisker follicle.

Glial cells as well as the melanocyte lineage are associated with sphere-forming p75/Sox10-positive cells in the adult skin

Unlike in the head, the mesenchyme in the trunk is not derived from the neural crest (Santagati and Rijli, 2003), and β-galactosidase expression in back skin of Wnt1-Cre/R26R mice was thus restricted to a few locations (Fig. 6, A and C). The same structures were also labeled in the back skin of Ht-PA-Cre/R26R mice (Fig. 6, B and D). In particular, both in the anagen and telogen stage, X-gal staining was found in the permanent part of the pelage follicle, including the bulge region below the sebaceous gland (Fig. 6, A and B; and Fig. S1, available at http://www.jcb.org/cgi/content/full/jcb.200606062/DC1). This area comprises the location of melanocyte stem cells (Nishimura et al., 2002) and glial cells in nerve endings (Botchkarev et al., 1997). In addition, pigmented melanocytes in the bulb region (the lower part of the haır follicles; Fig. 6, C and D) and nerves expressed β-galactosidase. In contrast, other hair follicle structures such as the dermal papilla, dermal sheath, and the outer and inner root sheaths were X-gal negative and, in the trunk skin, do not give rise to melanocyte stem cells and therefore not stained with X-gal. Six independent experiments were performed for each structure. Bar, 50 μm.

To elucidate whether p75/Sox10 expression and the capacity to form spheres are associated with glial cells from skin, we made use of desert hedgehog (Dhh)-Cre mice that express Cre recombinase in the peripheral glial lineage from early stages onward, but not in migrating neural crest cells or in neural crest–derived cells of other than glial lineages (Jaegle et al., 2003). β-Galactosidase activity was detectable in nerves and nerve endings in the back skin of adult Dhh-Cre/R26R mice (Fig. 7 A). As predicted from the proposed location of glial cells associated with nerve endings in the hair follicle (Botchkarev et al., 1997), X-gal staining in pelage follicles of Dhh-Cre/R26R mice was confined to a region around the bulge (Fig. 7 A), corresponding to the area that also contains β-galactosidase–expressing cells in Wnt1-Cre/R26R, Ht-PA-Cre/R26R, and Sox10lacZ mice (Fig. 6, A and B; and Fig. S1). In Dhh-Cre/R26R mice, X-gal–labeled cells of the bulge region were also labeled with anti-Sox10 antibody (Fig. 7 C) and anti-p75 antibody (Fig. 7 E). Pigmented melanocytes in the hair follicle bulge were X-gal negative, however, indicating that cells labeled in Dhh-Cre/R26R mice do not give rise to melanocyte s and thus are not related to the melanocyte lineage (Fig. 7 G).
To directly demonstrate that cells from the glial lineage tracked by Dhh-Cre promoter activity possess sphere-forming potential, these cells have to be prospectively identified and freshly isolated. One possibility to achieve this would be using specific surface antigen markers. However, such markers for the early glial lineage are currently unavailable. Furthermore, nerves present in the skin cannot be isolated by microdissection. Therefore, we used a genetic strategy to prospectively identify and directly isolate cells associated with the glial lineage. Dhh-Cre mice were mated with R26R-EYFP mice that express EYFP upon Cre-mediated recombination (Srinivas et al., 2001). Cells expressing EYFP in the trunk skin of Dhh-Cre/R26R-EYFP double-transgenic mice were isolated by FACS and transferred into medium permissive for sphere formation (Fig. 8, A and B). Although from unselected skin samples >10^6 cells were used to generate ~50 spheres (Fig. 1; see Materials and methods), <10,000 cells from both the EYFP-positive and -negative cell fraction were seeded in these experiments, to assess a possible enrichment in the spherogenic potential of FACS-selected cells. In two independent experiments, the EYFP-positive (Fig. 8 A, green frame), but not the EYFP-negative (Fig. 8 A, blue frame), cell population gave rise to spheres. Moreover, acutely fixed primary spheres of EYFP-positive cells were composed of cells expressing both p75 and Sox10 (Fig. 8 C). Thus, p75/Sox10-positive cells related to the glial lineage can be isolated from the skin and form spheres.

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We next asked whether sphere-forming potential is a common feature of peripheral glia. Therefore, we investigated whether sphere cultures can also be established from adult peripheral nerves. In agreement with others (Toma et al., 2001), we were unable to obtain spheres from cultures of...
dissociated sciatic and trigeminal nerves from adult mice (unpublished data). Thus, nerves or nerve endings in skin, but not peripheral nerves in general, contain cells with sphere-forming potential.

In cell preparations from the trunk skin of Dhh-Cre/R26R mice, only a fraction of all p75/Sox10-positive cells also expressed β-galactosidase (unpublished data). This could point to inefficient Cre-mediated recombination in Dhh-Cre/R26R mice. Alternatively, sources in the skin other than the glial lineage might yield sphere-forming neural crest–related cells. To address whether spherogenic neural crest–derived cells might be connected to the melanocyte lineage, we traced the fate of trunk skin cells in Dct-Cre/R26R mice (Guyonneau et al., 2002). Dct codes for the enzyme dopachrome tautomerase (also called Trp-2), which is required for melanin synthesis and already expressed in melanocyte stem cells (Nishimura et al., 2002). As expected, β-galactosidase activity in the back skin of Dct-Cre/R26R mice was detectable in melanocytes (Fig. 7 H) and in the hair follicle bulge region corresponding to the location of melanocyte stem cells (Fig. 7 B; Nishimura et al., 2002). Moreover, some X-gal–positive cells in the bulge region also expressed Sox10 (Fig. 7 D) and p75 (Fig. 7 F).

To investigate whether, in addition to cells of the glial lineage, the early melanocyte lineage also comprises undifferentiated neural crest–derived cells with the capacity to generate spheres, we isolated EYFP-expressing cells prospectively identified in the skin of Dct-Cre/R26R-EYFP mice. Intriguingly, in two independent experiments, FACS isolation and culturing of <10,000 cells revealed that only EYFP-expressing (Fig. 8 E, green frame), but not EYFP-negative, cells (Fig. 8 E, blue frame) were able to form spheres (Fig. 8 H). Analysis of acutely fixed primary spheres revealed many cells coexpressing p75 and Sox10, whereas pigmented differentiated melanocytes were absent (Fig. 8 G). These data indicate that the early melanocyte lineage comprises p75/Sox10-positive cells that can be propagated as spheres. Thus, as in whisker follicles of the face, the trunk skin contains more than one source of sphere-forming neural crest–derived cells, namely, cells of the glial and melanocyte lineages.

**Discussion**

In the present study, we show that cells with NCSC features can be isolated from the adult trunk skin of both mouse and human. Like NCSCs from other embryonic and postnatal sources, these neural crest–derived cells in the skin express p75 and Sox10 and are multipotent, able to generate several neural and nonneural lineages. Moreover, multipotent neural crest–derived cells from the adult skin display a self-renewing capacity, in that mouse and human skin–derived cells can be grown and expanded for months in floating sphere cultures. Intriguingly, in whisker follicles of facial skin, several structures of neural crest origin appear to comprise cells with sphere-forming capacity. In the trunk skin, however, genetic cell fate mapping, p75/Sox10 expression analysis in vivo, and, importantly, prospective identification and direct isolation demonstrate that cells displaying NCSC properties do not reside in mesenchymal structures of hair follicles but, rather, are associated with the melanocyte and glial lineages. Thus, self-renewing neural crest–derived cells from the skin are not confined to a particular niche but can be attributed to distinct locations in face and trunk skin.

**Sphere-forming neural crest-derived cells reside in distinct structures of the adult skin**

Several reports have described the isolation of multipotent cells from murine and human skin (Belicchi et al., 2004; Dyce et al., 2004; Joannides et al., 2004; Sieber-Blum et al., 2004; Amoh et al., 2005; Shih et al., 2005; Toma et al., 2005), raising the question about the origin of the endogenous cells able to self-renew and to generate multiple cell lineages, including cell types normally not found in the skin. Multipotent cells expressing GFP under the control of a Nestin regulatory element have been isolated from the hair follicle bulge area of transgenic mice and reported to undergo neurogenesis in vivo upon transplantation into the murine subcutis (Amoh et al., 2005). In these mice, Nestin-GFP–expressing cells are associated with the outer root...
sheath (Li et al., 2003), which, however, does not originate from the neural crest (Figs. 5 and 6). The nature of Nestin-expressing cells in the skin remains to be determined, though, as Nestin-GFP in another transgenic mouse line (Kawaguchi et al., 2001) marks the inner but not outer root sheath (unpublished data). Moreover, Nestin may be widely expressed in multiple structures of human skin (Wang et al., 2006).

The dermal papilla from whisker follicles has been reported to be of neural crest origin and to harbor sphere-forming SKPs that are p75/Sox10 negative (Fernandes et al., 2004). However, many structures of the whisker follicle turned out to be neural crest derived, including the dermal sheath, the ringwulst, and the capsule, apart from the dermal papilla (Fig. 5). Intriguingly, upon microdissection, all these tissues appeared to have cells with sphere-forming potential, although we cannot formally exclude possible contamination of the microdissected material by other structures. Thus, the capacity to self-renew appears to be a widespread feature of neural crest-derived cells in the adult skin of the face. Moreover, at least some of these self-renewing cells are also multipotent, as both SKP spheres from whisker pads and explant cultures of the upper part of whisker follicles with dermal sheath and the bulge region contain multipotent neural crest-derived cells (Fernandes et al., 2004; Sieber-Blum et al., 2004).

It has been proposed that, similar to SKPs from face skin, p75-negative cells isolated from the mouse back skin or the human foreskin are also neural crest derived, although this has not been addressed yet (Fernandes et al., 2004; Toma et al., 2005). Most likely, however, these latter sphere-forming cells are not associated with the dermal papilla of hair follicles, because the human foreskin is devoid of hair follicles. Moreover, using in vivo cell fate mapping, we demonstrated that, unlike in whisker follicles of the face, only few structures of pelage follicles in the trunk skin are actually neural crest derived. Dermal papilla, dermal sheath, and other supportive structures do not appear to be neural crest derivatives (Figs. 5 and 6). This is consistent with studies in chicken and mouse, which showed that mesenchymal tissues in the trunk are not neural crest derived (Santagati and Rijli, 2003). These differences in facial versus trunk neural crest contribution might also be relevant for the study of hair follicle development, given that whisker follicles are a widely used model system to investigate mechanisms regulating follicular cell fates (Alonso and Fuchs, 2003; Gambardella and Barrandon, 2003).

Skin structures harboring multipotent neural crest–derived cells are presumably marked by p75 and Sox10 expression in vivo, given that cells displaying NCSC properties after isolation from the adult trunk skin express these markers. Indeed, in Wnt1-Cre/R26R, Ht-PA-Cre/R26R, Dhh-Cre/R26R, and Dct-Cre/R26R mice, β-galactosidase–expressing cells positive for p75 and Sox10 were found in the bulge region encompassing melanocyte stem cells and glial cells in nerve endings (Botchkareva et al., 1999; Nishimura et al., 2002; Figs. 6 and 7). However, the only way to unambiguously demonstrate that these lineages contain resident multipotent cells with the potential to self-renew is by prospectively identifying such cells in the adult skin and testing their potential upon acute isolation. Because of the lack of specific surface markers, we were not able to use antibodies to directly isolate multipotent cells from back skin. In particular, p75 expression does not distinguish between glial and melanocyte lineages (Fig. 7) and, in addition, is found in regions of the skin that are not neural crest derived, such as the outer root sheath of hair follicles, and, at early stages of hair follicle morphogenesis, the dermal papilla (Fig. 6; Botchkareva et al., 1999; Rendl et al., 2005). Similarly, the melanocyte marker c-Kit is not suitable for isolation of prospective multipotent cells from the melanocyte lineage because it is not expressed in the bulge of anagen hair follicles and because in vivo it is also found in epithelial skin cells not originating from the neural crest (Peters et al., 2002, 2003). Moreover, in preliminary experiments, we failed to obtain p75/Sox10-positive spheres from c-Kit–positive skin cells isolated by FACS (unpublished data). Nonetheless, we were able to identify spherogenic neural crest–derived cells in the skin by using a genetic approach. Thereby, the lineage-specific activity of Dhh-Cre and of Dct-Cre, respectively, combined with a Cre reporter allele (Srinivas et al., 2001), led to EYFP expression in cells from either the glial or the melanocyte lineage. Consistent with a dual origin of multipotent neural crest–derived cells in the trunk skin, EYFP-positive cells isolated from both Dhh-Cre/R26R-EYFP and Dct-Cre/R26R-EYFP mice formed spheres of p75/Sox10-positive cells (Fig. 8). Thus, we conclude that in the trunk skin of mice, spherogenic neural crest–derived cells are associated with the glial as well as the melanocyte lineage. Based on their high similarities to mouse cells in terms of marker expression and potential, and based on the fact that humans do not have whisker follicles, we assume that the p75/Sox10-positive multipotent cells obtained from human skin (Figs. 1 and 3) are also related to glial and melanocyte rather than mesenchymal lineages.

That cells of the bulge region marked in Dct-Cre/R26R mice generate melanocytes is in agreement with earlier findings (Nishimura et al., 2002), which identified Dct<sup>−/−</sup>-positive cells of the bulge region as so-called melanocyte stem cells generating differentiated melanocytes in the lower part of the hair follicle. Moreover, a considerable fraction of Dct-positive cells in the bulge also expresses Sox10 (Osawa et al., 2005), in agreement with our results (Fig. 7). Our analysis of neural crest–derived cells isolated from Dct-Cre/R26R mice supports the hypothesis that melanocyte progenitors in the bulge region are not only self-renewing (Nishimura et al., 2002) but indeed represent multipotent cells (Sommer, 2005).

The combined data indicate that the adult skin is host to sphere-forming cells with different identities. In addition to multipotent, undifferentiated neural crest–related cells (Fernandes et al., 2004; this study), the potential to self-renew has also been attributed to pigment cells and possibly other developmentally restricted neural crest–derived cell types (Dupin et al., 2000; Trentin et al., 2004). Similar to NCSCs isolated at different time points and from different peripheral nervous system regions (Kleber and Sommer, 2004), the multipotent neural crest–derived skin cells described in this study display altered responsiveness to instructive growth factors as compared with migratory NCSCs (Fig. 4). Thus, multipotent, self-renewing
SKPs failed to generate CNS neurons upon transplantation. Likewise, these trials suggest that multipotent neural crest–derived cells are a valuable source for cell replacement therapies because sufficient cell material could be obtained for such purposes. However, in our hands, spherogenic neural crest–derived cells from the adult mouse skin displayed a rather restricted potential in vivo (unpublished data). In particular, skin-derived multipotent neural crest–related cells displayed a restricted potential when compared to skin spheres, which are capable of giving rise to neural and nonneural cell types at clonal density (Fig. 4) does apparently not reflect transdifferentiation but, rather, the broad potential inherent to NCSCs. The fact that these cells can be easily expanded in culture, even when isolated from the skin of adult humans, might make them a valuable source for cell replacement therapies because sufficient cell material could be obtained for such purposes. However, in our hands, spherogenic neural crest–derived cells from the adult mouse skin displayed a rather restricted potential in vivo (unpublished data). In particular, skin-derived spheres of neural crest origin, when dissociated and injected into the lateral ventricles of rat and chicken embryos or transplanted onto hippocampal brain slices, remained largely undifferentiated in cell aggregates close to the injection site and failed to integrate into the host central nervous system (CNS) tissue (unpublished data). This is in contrast to neural progenitors obtained from embryonic stem cells or neural stem cells from the CNS assessed in the same experimental paradigms (Brüstle et al., 1997; Benninger et al., 2003). Moreover, we did not observe neural differentiation, tissue integration, or behavioral improvement upon transplantation of skin-derived multipotent neural crest–derived cells into the striatum of a 6-hydroxydopamine–treated mouse (unpublished data). In particular, skin-derived multipotent neural crest–related cells with stem cell properties might be of high value for the generation of Schwann cells, cartilage, or other neural crest–derived tissues potentially useful in clinical applications. Hence, our study underlines the importance of choosing the appropriate stem cell type for a given task.

Materials and methods

Skin sphere cultures

Human thigh skin from an adult man (−45 yr of age) and face skin from an adult woman (−37 yr of age; provided by G. Beer, University Hospital of Zurich, Zurich, Switzerland) were obtained in the frame of cosmetic surgery according to the guidelines of the University Hospital of Zurich. Murine skin was taken from adult C57BL/6 mice of at least 8 wk of age. Skin samples (composed of both dermis and epidermis) were dissected, cut into small pieces, and digested in 0.1% Trypsin-EDTA (Invitrogen) in HBSS without Ca²⁺ and Mg²⁺ (Aniwmed) and digested for 30 min at 37°C. Partially digested skin pieces were dissociated mechanically and filtered through a 40-μm cell strainer (BD Biosciences). The cell suspension was centrifuged and washed with medium, and the cell pellet was resuspended in growth medium (GM) consisting of DME-F12 1:1 containing 1× B-27 supplement (Invitrogen), 20 ng/ml FGF2 (Peprotech), 10 ng/ml EGF (Peprotech), penicillin/streptomycin (P/S), and fungizone. GM for human cells also contained 10 ng/ml leukemia inhibitory factor (Sigma-Aldrich). 2.5–4 million cells were seeded in GM into an uncoated T25 cell culture flask (BD Biosciences). After 4–7 d in culture, sphere formation was observed. For FACS analysis, skin was taken from mice between 10 and 16 d of age. Samples were incubated in 0.5 mg/ml Dispase (Roche) in HBSS for 30 min at 4°C. Fat tissue was removed with forceps, and the rest of the skin was cut into small pieces and digested in 1 mg/ml collagenase ( Worthington) in HBSS for 45 min at 37°C. After a final digestion with 0.1% Trypsin-EDTA in HBSS for 5 min at 37°C, the partially digested skin pieces were dissociated mechanically and treated as described above. FACS was performed with a FACS Aria (Becton Dickinson).

Skin sphere passaging

Once a week, the sphere suspension was transferred into a 15-ml Falcon tube. Cells adhering to the flask bottom were discarded. Spheres were centrifuged, and one third of the supernatant was transferred as conditioned medium into a new T25 flask. Spheres were incubated with 300 μl Trypsin-EDTA solution (0.25%) for 3–5 min at RT. 400 μl of Ovomucoid solution (1 mg/ml Trypsin inhibitor [Sigma-Aldrich] and 10 mg DNase [Roche] in 25 ml medium) were added, and spheres were dissociated mechanically, centrifuged, resuspended in fresh GM, and seeded into a new flask containing one third conditioned medium. After some passages, spheres were cultured in flasks coated with Poly(2-hydroxyethylmethacrylate) (Poly-Hema; Sigma-Aldrich). Coating was performed at RT with a solution of 16 mg/ml Poly-Hema in 95% ethanol.

Microdissection and sphere cultures of whisker follicles

Anagen-phase whisker follicles from mouse and rat were dissected out of the whisker pad and microdissected as described previously (Kobayashi et al., 1993). Follicle structures were incubated in 0.05% Trypsin-EDTA in DME for 1.5–2 h at 37°C. Trypsin activity was stopped with DME containing 10% FCS. After two washing steps with GM, cells of each structure were plated in a well of a 24-well dish. Sphere formation was observed within 1–2 wk. Passaging was performed as described for skin spheres using 0.05% Trypsin-EDTA for 2 min at RT.

Cell differentiation

All differentiation assays were performed using spheres plated on dishes coated with fibronectin (FN) or poly-L-lysine/FN as described previously (Stemple and Anderson, 1992). Neurogenesis, gliogenesis, and smooth muscle formation were observed after 3–7 d in GM. Chondrocyte forma- tion was obtained after 9 d in DME containing 10% FCS, 50 μg/ml ascorbic acid 2-phosphate (Sigma-Aldrich), 10 ng/ml FGF2, and PDGF, followed by DME containing 10% FCS, 50 μg/ml ascorbic acid 2-phosphate, and 10 ng/ml BMP2 (Peprotech) for another 3 d. Adipocytes were occasionally observed when spheres were cultured in DME/F12 containing B-27 supplement and 10 ng/ml BMP2. Melanocytes were observed when cultured in...
MEM containing 10% FCS, 50 ng/ml murine stem cell factor (PeproTech), 100 nM endothelin-3 (Sigma-Aldrich), and P/S for at least 10 d.

Clonal analysis of neural crest-derived stem cells
Murine skin-derived spheres passaged >17 times were dissociated with Trypsin-EDTA as described and plated at clonal density on pDL/FN-coated 35-mm dishes (Corning) in standard medium prepared as reported previously (Stemple and Anderson, 1992). Single p75-positive cells were picked and labeled as described previously (Hagedorn et al., 1999) and incubated in standard medium alone or supplemented with 100 ng/ml BMP2, 1 nM NRG1 (R&D Systems), or 0.1 ng/ml TGFβ (R&D Systems). After 10 d, the cells were fixed and analyzed immunocytochemically.

Immunofluorescence on cells and tissue sections
Anti-p75, anti-Sox10, anti-SMA, and anti-GFAP antibody stainings on cells were done as described by Kleber et al. (2005). Anti-TuJ1 antibody (1:200; Sigma-Aldrich) and anti-Keratin antibody (1:500; Abcam) were used for 2 h at RT, whereas anti-β-galactosidase (1:100; Roche) and anti-NG2 (1:200; Chemicon) antibodies were used with incubation overnight at 4°C. The following secondary antibodies were used for 1 h at RT: Cy3-conjugated goat anti-mouse (1:200; Jackson ImmunoResearch Laboratories), Cy3-conjugated goat anti-rabbit (1:200; Jackson ImmunoResearch Laboratories), Alexa 488-conjugated goat anti-mouse (1:100; Invitrogen), and Alexa 488-conjugated goat anti-rabbit (1:100; Invitrogen). Cell nuclei were stained with DAPI. Paraffin sections of X-gal–treated skin were stained for Sox10 as described previously (Dutt et al., 2006), using a controlled antigen-retrieval device (FSG 120-T/T; Milestone). Heat unmasking for the p75 was done in 10 mM trisodium citrate, pH 6.0, using the same antigen-retrieval device. The antibody (Chemicon) was used at a dilution of 1:5,000. Alexa 594–conjugated goat anti-mouse and Alexa 488–conjugated goat anti-rabbit (1:200; Invitrogen) were used as secondary antibodies. Immunofluorescence of cells was analyzed using a microscope (Axiovert 100; Carl Zeiss Microlmaging, Inc.) and magnifications of 32×. Pictures were made with a camera (AxioCam MRm) and Axiovision 4.2 software (Carl Zeiss MicroImaging, Inc.). Immunofluorescence on cells and tissue sections after 10 d, the cells were fixed and analyzed immunocytochemically. The antibody (Chemicon) was used at a dilution of 1:5,000. Alexa 594–conjugated goat anti-mouse and Alexa 488–conjugated goat anti-rabbit (1:200; Invitrogen) were used as secondary antibodies. Immunofluorescence of cells was analyzed using a microscope (Axiovert 100; Carl Zeiss Microlmaging, Inc.) and magnifications of 32×. Pictures were made with a camera (AxioCam MRm) and Axiovision 4.2 software.

Other stainings
For Alcian blue staining, cells fixed with 4% formaldehyde were incubated with a 3% solution of glacial acetic acid in distilled water for 3 min at RT. For油 red O staining, fixed cells were incubated in a 3% solution of glacial acetic acid in distilled water for 3 min at RT. For Alcian blue staining, cells fixed with 4% formaldehyde were incubated with a 3% solution of glacial acetic acid in distilled water for 3 min at RT. For oil red O staining, fixed cells were incubated in a 3% solution of glacial acetic acid in distilled water for 3 min at RT. For Alcian blue staining, cells fixed with 4% formaldehyde were incubated with a 3% solution of glacial acetic acid in distilled water for 3 min at RT. For oil red O staining, fixed cells were incubated in a 3% solution of glacial acetic acid in distilled water for 3 min at RT.

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