Generation of Induced Pluripotent Stem Cells from Hair Follicle Bulge Neural Crest Stem Cells

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Abstract

Induced pluripotent stem cells (iPSCs) are promising candidates for the study of disease models as well as for tissue engineering purposes. Part of a strategy to develop safe reprogramming technique is reducing the number of exogenous reprogramming factors. Some cell types are more prone to reprogramming than others. iPSC induction with less reprogramming factors has been described in cells with endogenous expression levels of pluripotency genes, such as neural stem cells. Because multipotent neural crest stem cells (NCSCs) from mammalian hair follicle bulges also express pluripotency genes, we argued that this property would facilitate reprogramming of hair follicle bulge NCSCs and could substitute for the use of exogenous reprogramming factors. Although we confirmed the expression of pluripotency genes in hair follicle bulge cells, our results show that these cells do require a full set of reprogramming factors for iPSC induction. Hair follicle bulge–derived iPSCs were created with efficiencies similar to fibroblasts. We conclude that high endogenous levels of pluripotency factors are no guarantee for facilitated induction of pluripotency.

Introduction

Mammalian somatic cells can be reprogrammed toward a pluripotent state by means of transduction with (combinations of) specific transcription factors, such as Oct4, Sox2, Klf4, c-Myc (Yamanaka factors), Nanog, or Lin28 (Liao et al., 2008; Takahashi and Yamanaka, 2006; Takahashi et al., 2007; Yu et al., 2007). These so-called induced pluripotent stem cells (iPSCs) are in many aspects similar to embryonic stem cells (ESCs) and provide a potential autologous source of stem cells for application in tissue engineering and regenerative medicine (Chin et al., 2009; Hanna et al., 2007; Wernig et al., 2008). To develop safe and reproducible iPSC technology for future clinical use, focus over the past few years has been to find strategies to: (1) minimize the number of reprogramming factors needed (Kim et al., 2008), (2) avoid potentially oncogenic factors like c-Myc (Nakagawa et al., 2007), (3) replace integrative viruses with other nonintegrative transgene carriers to avoid potentially mutational insertions (Okita et al., 2008; Zhou and Freed, 2009), (4) simplify and standardize reprogramming using defined small molecules instead of transcription factors (Desponts and Ding, 2010; Jia et al., 2010; Yu et al., 2009), and (5) find cell types that are both easily accessible and more prone to reprogramming (Eminli et al. 2008; Kim et al., 2009; Tsai et al., 2010). In the present study, we focus on the latter strategy.

The neural crest is a population of migratory multipotent progenitor cells in vertebrates that emerges and delaminates from the dorsal neural tube during neurulation. It contributes significantly to the development of mammalian craniofacial tissues (for review, see Santagati and Rijli, 2003). As a result, neural crest stem cells (NCSCs) persist in several craniofacial structures in the adult organism and therefore are a relatively accessible source of stem cells for transplantation studies and tissue engineering (for review, see Kaltschmidt et al. 2012; Shakhova and Sommer, 2010).

We isolated nestin-expressing NCSCs from the adult mouse whisker follicle (Amoh et al. 2005; Amoh et al., 2009; Sieber-Blum et al. 2004; Sieber-Blum and Hu, 2008) and analyzed the expression of pluripotency genes in these adult hair follicle NCSCs (HFNCSCs) both at the mRNA and protein levels. We hypothesized that (high) endogenous expression of one or more pluripotency factors might facilitate pluripotency induction compared to, for instance, skin fibroblasts and/or that less transcription factors would be necessary to induce pluripotency. We confirmed the pluripotent expression profile of hair follicle bulge NCSCs and also showed that this does not...
facilitate their reprogramming into iPSCs and that a full set of Yamanaka factors is still required.

Material and methods

Isolation of HFNCSCs and cell culture

NCSCs were isolated from mouse whisker follicles as previously described (Sieber-Blum et al., 2004). In short, whiskers of C57BL/6 mice and nestin–green fluorescent protein (GFP) transgenic mice were dissected. Hair follicle bulges were cultured on poly-d-lysine (PDL)-laminin–coated culture plates (one per well) in Neuralbasal (Gibco Invitrogen 21103-049) 1% glutamax (Gibco Invitrogen 35050-038), 1% l-glutamine (PAA M11-006), 5 μg/mL heparin (Sigma H4784), 2% B27 serum-free supplement (Gibco Invitrogen 17504-044), 20 μg/mL epidermal growth factor (EGF) (Gibco Invitrogen PHG0311), 20 μg/mL basic fibroblast growth factor (bFGF) (Gibco Invitrogen PHG0021), and 0.2% Primocin (Amaxa VZA-1022). The medium was changed every 2–3 days.

iPSC induction

For reprogramming of the mouse HFNCSCs, we have used both the retroviral and lentiviral transduction approaches. For retroviral transduction, retroviral particle–containing supernatants were produced using Phoenix Eco packaging cells that were transfected using FuGENE, with separate pMX-based vectors encoding murine Oct4, Sox2, Klf4, and cMyc plasmids (Addgene), as described previously (Czepiel et al., 2011). For lentiviral transduction, HEK293T cells were transfected with a lentiviral self-inactivating pRRL.PPT.SF plasmid containing murine Oct4, Klf4, Sox2, c-Myc, and dTomato complementary DNA and co-transfected with pMD2-VSV-G, and pCMV-A8.91 packaging vectors. Lentiviral supernatant was collected 48 h posttransfection and filtered through a 0.45-μm filter (Whatman). Around 20,000 HFNCSCs were transduced with viral supernatant supplemented with 8 μg/mL Polybrene (Sigma-Aldrich), which was replaced by fibroblast medium after 24 h. Transfection efficiency was evaluated on the basis of fluorescent dTomato expression. Four days posttransduction, cells were dissociated using 0.05% trypsin-EDTA and seeded onto irradiated mouse embryonic fibroblast (MEF) feeder layers on a 1% gelatin-coated six-well plate and cultured in ES medium. For in vitro default differentiation, medium was switched to N2 medium consisting of DMEM/F12 supplemented with 1% N2 supplement (R&D Systems), 2 mM l-glutamine, 100 U/mL penicillin/streptomycin, and 1 mM sodium pyruvate for 5 days.

Immunocytochemistry

Immunocytochemistry was performed as previously described (Czepiel et al., 2011). The following antibodies were used: Mouse anti-mouse nestin (1:200; Chemicon, MAB 353), mouse anti-mouse Oct4 (1:100; Santa Cruz, 5279), rabbit anti-mouse Sox2 (1:200; Abcam, 15830), mouse anti-mouse Sox2 (1:200; Cell Signaling, 4900S), rabbit anti-Nanog (1:200; Abcam, 80892), mouse anti-mouse SSEA-1 IgM (1:200; Santa Cruz, 21702), mouse anti-mouse desmin (1:200; Dako M0760), mouse anti-mouse GATA (1:200; Santa Cruz, 25310), and mouse anti-mouse βIII-tubulin (1:400; Abcam 7751). The following fluorescently labeled secondary antibodies were used: Alexa Fluor 488-conjugated donkey anti-rabbit (1:300; Molecular Probes, A21206), Alexa Fluor 488-conjugated goat anti-mouse IgM (1:300; Invitrogen, M31601), DyLight 488-conjugated goat anti-rabbit (1:100; Jackson Immunoresearch), and Cy3-conjugated donkey anti-goat (1:300; Jackson Immunoresearch). For nuclear visualization, cells were counterstained with Hoechst 33342 (1:1000; Jackson Immunoresearch) in phosphate-buffered saline (PBS). The alkaline phosphatase (AP) activity of iPSC colonies was detected using an AP detection kit (Sigma).

qRT-PCR

Total RNA was extracted (Chomczynski and Sacchi, 1987) and transcribed into cDNA using random hexamers and Moloney murine leukemia virus (M-MLV) reverse transcriptase (Fermentas). Purity of the RNA was confirmed by means of gel electrophoresis. Quantitative real-time PCR was performed in 384-well plates using iQ SYBR Green Supermix (Bio-Rad) on an ABI7900HT system (Applied Biosystems) with the following cycling parameters: 95°C for 3 min, 39 cycles with 95°C for 10 sec, and 58°C for 30 sec. qRT-PCR primers were designed with PeriPrimer for the transcripts listed in Table 1. Housekeeping genes hydroxymethylbilane synthase (HMBS) and glyceraldehyde-3-phosphate dehydrogenase (GAPDH) were used as internal controls for normalization. The data were processed with SDS version 2.3 analysis software (Applied Biosystems). The relative expression ratio of target genes was quantified using the 2^{-ΔΔCt} calculation method (Schmittgen and Livak, 2008).

Western blot

Proteins were extracted from cell samples using lysis buffer supplemented with protease inhibitors. Cell lysates were sonicated and quantitated by DC Protein Assay (Bio-Rad, Hercules, CA), and proteins were separated on 15% sodium dodecyl sulfate polyacrylamide gels and then transferred onto polyvinylidene fluoride membranes. Membranes were incubated in Odyssey blocking buffer and incubated overnight in primary antibodies diluted in PBS + 0.1% Tween-20. The following primary antibodies were used: Mouse anti-mouse Oct4 (1:1,000; Santa Cruz, 5279), mouse anti-mouse Sox2 (1:1,000;...
Table 1. Primers Used for Quantitative Real-Time PCR (qRT-PCR)

<table>
<thead>
<tr>
<th>Accession no.</th>
<th>Gene</th>
<th>Forward primer (5’-3’)</th>
<th>Reverse primer (5’-3’)</th>
<th>Product size (bp)</th>
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<tr>
<td>NM_013633</td>
<td>Oct4</td>
<td>CCTGGCGGTCTCTTGGGA</td>
<td>CTCGGAACGCGAGATGGTG</td>
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<tr>
<td>NM_011443</td>
<td>Sox2</td>
<td>AACCAAGAACAGCCGGA</td>
<td>CGGACCATTCCAAGAAGG</td>
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<tr>
<td>NM_010637</td>
<td>Klf4</td>
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<td>CTCACGCAACGTTAATGC</td>
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<tr>
<td>NM_001177352</td>
<td>c-Myc</td>
<td>GAGGGCAAGTGGACAGTGG</td>
<td>GCGTGTGTGTGATCTGCT</td>
<td>51</td>
</tr>
<tr>
<td>NM_028016</td>
<td>Nanog</td>
<td>GATTCGAAAGGGCTACGACC</td>
<td>TAAGCTGCAATTGGATCTG</td>
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<tr>
<td>NM_001110251</td>
<td>HMBS</td>
<td>CCGAGCCAAGGGACAGGA</td>
<td>CTCTCTCCAGTGCTCAGA</td>
<td>107</td>
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</table>

Results

Expression of pluripotency factors in HFNCSCs

We isolated HFNCSCs from the adult mouse whisker follicle of nestin–GFP transgenic mice as well as from C57BL/6 mice (Fig. 1A). Explants were placed in lamin-coated six-well plates, and NCSCs were allowed to migrate out of the follicle bulges (Fig. 1B, C). For assessment of pluripotency gene expression with immunohistochemistry, C57BL/6 cells were replated in small badges of 10–100 cells on laminin and allowed to expand clonally in selective proliferation medium. In this way, pure populations of HFNCSCs could be obtained that proliferated in monolayers as described previously (Sieber-Blum et al., 2004). After 1 week, cells were analyzed by means of immunocytochemistry, and 100% of the cells were found to be nestin-positive, whereas >95% strongly expressed Sox2 (Fig. 1D). Also the pluripotency markers Oct4 and Nanog were expressed, although weaker in comparison to Sox2. Oct4 expression was seen mainly in the cytoplasm but not in the nucleus (in accordance with previous findings; Atlasi 2008). Monolayers of nestin-GFP HFNCSCs (p0) were sorted by fluorescence-activated cell sorting (FACS) for GFP after 1 week and analyzed for mRNA expression of pluripotency genes using qRT-PCR (Fig. 1E). Pluripotency gene expression was compared with that in MEFs, ESCs, and previously generated F-iPSCs. Compared to MEFS and ESCs, Sox2 and Nanog are expressed 100-fold higher in HFNCSCs (ESCs, p > 0.05; Sox2 and Nanog, p < 0.05). HFNCSCs expressed Klf4, Sox2, c-Myc, and Nanog at levels that did not differ significantly from those in ESCs (p > 0.05 for all factors); only Oct4 is expressed 100-fold higher in ESCs (p < 0.05).

Reprogramming of HFNCSCs into iPSCs

In view of the specific endogenous pluripotency expression profiles of the HFNCSCs, we tested whether we could reprogram these cells into iPSCs while omitting the transcription factors already highly expressed by the HFNCSCs (Sox2, Klf4, and/or c-Myc were left out in different combinations as well as altogether; Oct4 was always present). After 4 weeks, no colonies were observed in any of the conditions. Only lentiviral transduction of all four Yamanaka factors in a polycistronic vector resulted in the appearance of iPSC colonies within 5–7 days posttransduction (Fig. 2A). Of 2 × 104 transduced cells, we counted an average of five colonies per field of view at 100× magnification, or a total of 1125 colonies. Efficiency was calculated at 0.056%, which is slightly higher than efficiencies with fibroblasts and other cell types. Colonies displayed a proper ESC-like morphology with a large nuclear/cytoplasmic ratio and prominent nucleoli. HF-iPSCs were characterized as to their reprogramming and pluripotent characteristics.

Discussion

The mammalian hair follicle contains several compartments of stem cells that are responsible for regeneration of these structures during the anagen phase of the hair follicle as well as after injury (for review, see Fuchs and Horsley, 2008; Tiede et al., 2007). A cell population of neural crest origin has been identified in the hair follicle bulge region by the expression of β-galactosidase expression under the Wnt1 promoter in transgenic mouse models (Nagoshi et al., 2008; Sieber-Blum et al., 2004; Wong et al., 2006). These HFNCSCs express neural crest markers Sox10 and neural crest/neural stem cell marker nestin and can be differentiated toward neural crest lineages such as neurons, smooth muscle cells, and melanocytes in vitro (Fernandes et al., 2004;
Hunt et al., 2008; Li et al., 2003; Sieber-Blum et al., 2004; Toma et al., 2001). Amoh and Hoffman (Amoh et al., 2005; Amoh et al., 2009; Hoffman, 2006) isolated nestin+/-keratin 15- multipotent skin-derived cells from the hair follicle bulge of nestin-driven GFP transgenic mice, which could be differentiated into neurons, glia, keratinocytes, smooth muscle cells, and melanocytes in vitro.

Interestingly, some authors have reported the expression of pluripotency-related genes in HFNCSCs. Yu et al. described the presence of nestin-positive cells within the human

FIG. 1. (A) Location of bulge region within the hair follicle (B); other structures depicted are blood sinus (BS), capsula (C), dermal papilla (DP), outer root shaft (ORS), and sebaceous gland (SG). (B) Isolation of the bulge of a mouse whisker follicle. Sequentially—isolated hair follicle; transversal cut is made just below the ring sinus and above the papilla (dotted lines); a longitudinal incision is made through the middle section of the follicle (dotted line); the capsula is opened and the bulge region with the attached hair is isolated (arrow indicates bulge). Scale bar, 200 μm. (C) Adult mouse HFNCSCs migrating out of the bulge after 5 days on laminin (left panel), and 2 weeks after isolation (and after removal of bulge) (middle panel, magnification in right panel). (D) Characterization of HFNCSCs with immunocytochemistry. HFNCSCs strongly express nestin and weakly express Oct4; Sox 2 and Nanog are also expressed. Nuclei are counterstained with Hoechst. Scale bar, 200 μm. (E) qRT-PCR analysis for mRNA expression levels of Oct4, Sox2, Klf4, c-Myc, and Nanog in ESCs and HFNCSCs showing relative values normalized to HMBS and compared to expression levels of MEFs (logarithmic scale). Error bars represent standard deviation (SD) (n = 3); (*) p < 0.05, one-way ANOVA, Bonferroni post hoc multiple comparisons test.
hair follicle bulge, which in addition were positive for pluripotency factors Nanog and Oct4 (Yu et al., 2006). Sieber-Blum and Hu also showed the expression of (different) pluripotency factors in bulge NCSCs; c-Myc, Klf4, and Sox2 were expressed at similar levels compared to ESCs, whereas Oct4 and Nanog were expressed significantly lower than in ESCs (Sieber-Blum and Hu, 2008). Altogether these studies show that NCSCs persist in the mammalian hair follicle bulge and display several pluripotency-related expression profiles. By means of qRT-PCR, we confirmed that the nestin-expressing NCSCs from the bulge region of the adult mouse whisker follicle (HFNCSCs) co-express high mRNA levels of pluripotency genes Oct4, Sox2, Klf4, and Nanog. Expression of Oct4 was lower than in ESCs, but in contrast to Sieber-
Blum and Hu, we showed Nanog expression levels similar to ESCs. Like Yu et al., we were able to detect expression of both Oct4 and Nanog at the protein level.

We hypothesized that because of their endogenous expression of several pluripotency genes, HFNCSCs might be more prone for pluripotency induction, eventually with fewer factors. However, our results show that HFNCSCs can only be reprogrammed using a full set of Yamanaka factors, with an efficiency comparable or only slightly higher than generally found for fibroblasts. Others have shown that endogenous expression levels of pluripotency genes should be able to substitute for ectopic expression, and thus no transgenes would be necessary for iPSC induction in case of high-level expression. The group of Jaenisch and Hochedlinger showed expression levels of pluripotency genes should be able to substitute for ectopic expression, and thus no transgenes would be necessary for iPSC induction in case of high-level expression.

Characterization of four-factor derived HF-iPSCs did show ESC-like colony morphology, EB formation and differentiation patterns, and pluripotency marker expression, confirming proper pluripotency induction. The expression of transgene mRNAs was higher than in fibroblast-derived iPSCs, especially for Oct4 and Nanog; this might reflect the high expression levels in the original cells. Assessment of the methylation profiles of the promoter regions of key pluripotency genes would be insightful for determining whether these high expression levels in HF-iPSCs in comparison to F-iPSCs are due to epigenetic differences.

Altogether we have confirmed the pluripotent expression profile of hair follicle bulge NCSCs; however, HFNCSCs could be reprogrammed into fully characterized iPSCs only with a full set of exogenous pluripotency transcription factors (Oct4, Sox2, Klf4, and c-Myc), and without higher efficiency. We conclude that high endogenous levels of Yamanaka factors are no guarantee for facilitated induction of pluripotency.

**Author Disclosure Statement**

The authors declare that no conflicting financial interests exist.

**References**


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