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Prolonged cultivation of hippocampal neural precursor cells shifts their differentiation potential and selects for aneuploid cells

Abstract: Neural precursor cells (NPCs) are lineage-restricted neural stem cells with limited self-renewal, giving rise to a broad range of neural cell types such as neurons, astrocytes, and oligodendrocytes. Despite this developmental potential, the differentiation capacity of NPCs has been controversially discussed concerning the trespassing lineage boundaries, for instance resulting in hematopoietic competence. Assessing their in vitro plasticity, we isolated nestin+/Sox2+, NPCs from the adult murine hippocampus. In vitro-expanded adult NPCs were able to form neurospheres, self-renew, and differentiate into neuronal, astrocytic, and oligodendrocytic cells. Although NPCs cultivated in early passage efficiently gave rise to neuronal cells in a directed differentiation assay, extensively cultivated NPCs revealed reduced potential for ectodermal differentiation. We further observed successful differentiation of long-term cultured NPCs into osteogenic and adipogenic cell types, suggesting that NPCs underwent a fate switch during culture. NPCs cultivated for more than 12 passages were aneuploid (abnormal chromosome numbers such as 70 chromosomes). Furthermore, they showed growth factor-independent proliferation, a hallmark of tumorigenic transformation. In conclusion, our findings substantiate the lineage restriction of NPCs from adult mammalian hippocampus. Prolonged cultivation results, however, in enhanced differentiation potential, which may be attributed to transformation events leading to aneuploid cells.

Keywords: cellular transformation; hippocampus; neural precursor cells; neural stem cells; stem cell plasticity.

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Introduction

In the last decades, neural stem cells (NSCs) gained increasing interest as cells being responsible for adult neurogenesis and participating in tissue homeostasis in the mammalian brain. The importance of understanding identity and function of NSCs today has been elevated by their prospective application for treatment of neurological diseases in the human system.

Defined as self-renewing, multipotent cells that give rise to most cell types of the central nervous system (CNS), NSCs were first successfully isolated by Reynolds and Weiss (1992) in the early 1990s from adult mouse brain tissue. Nowadays, NSCs are commonly known to primarily reside within the subventricular zone (SVZ) of the lateral ventricle (Palmer et al., 1997). As a second prominent region, the subgranular zone of the hippocampal dentate gyrus was also shown to contain NSCs (Palmer et al., 1997), although more recent studies show evidence of hippocampal NSCs being neuronal precursors (NPCs) due to their lineage-restricted differentiation capability and limited capacity to self-renew (Seaberg and van der Kooy, 2002; Bull and Bartlett, 2005). Despite these two well-known regions, NPCs have also been detected in other regions of the mammalian CNS, namely the adult spinal cord and the subcallosal zone of the brain (Weiss et al., 1996; Seri et al., 2006). Concerning their in vivo function, NPCs have been proven to modify CNS tissue homeostasis by giving rise to neurons that can integrate into stable neural circuits (Cameron and McKay, 2001; Imlieski et al., 2012). In the adult mammalian brain, such generation of functional neurons was shown under normal conditions upon physiological stimuli or after injury (reviewed by Ma et al., 2009). In the adult
hippocampus, NPCs participate in adult neuronal plasticity by generation of newborn neurons (Imielski et al., 2012).

Besides the broad range of studies focusing on in vivo characterization, NSCs have been well studied in vitro as cell clusters called ‘neurospheres’ (Reynolds and Weiss, 1992; Gritti et al., 1996, 1999). Notably, a single NSC was also shown to exhibit the ability to form new neurospheres under appropriate medium conditions and give rise to both neurons and glial cells, suggesting the self-renewal capability and multipotency of NSCs (Morshead et al., 1994; Bonaguidi et al., 2008). Next to differentiation into their developmentally expected derivatives, NSCs were reported to trespass lineage boundaries, e.g., resulting in hematopoietic cell progeny or skeletal myotubes (Bjornson et al., 1999; Galli et al., 2000).

From the viewpoint of a cell simply changing its phenotype due to extracellular signals, the concept of plasticity has been altered over the last years. Today’s understanding of plasticity can be summarized as the shift of an undifferentiated but developmentally restricted cell toward another type of terminal differentiation (Wagers and Weissman, 2004; reviewed by Bonfanti et al., 2012). Adult stem cells from different sources than the CNS have also been shown to exhibit an extraordinary high developmental plasticity (Krause et al., 2001; Wagers et al., 2002), although this phenomenon is suggested to be associated with either spontaneous cell fusion or genomic abnormalities (Terada et al., 1994; Bonaguidi et al., 2008). In general accordance with the findings by (Seaberg and van der Kooy, 2002), neural precursor cells (NPCs) began to grow adherently after 2 weeks. Proliferating adherent NPCs homogeneously exhibited characteristic elongated bipolar morphology with oral nuclei (Figure 1B) (Conti et al., 2005). After reaching about 80% confluency, cells could be harvested and passaged, followed by further cultivation in an adherent manner. Proving their NPC characteristics in the first place (Babu et al., 2007), NPCs were shown to form neurospheres under appropriate culture conditions (Figure 1C).

**NPC cultures homogeneously express neural precursor markers while lacking expression of neural crest and mesenchymal stem cell markers**

Isolated NPCs in early passage (<7) were characterized using immunocytochemical staining revealing high expression of neural stem cell marker nestin (98.0±1.3%, Figure 2A) and NPC marker Olig2 (80.4±1.6%, 100%, Figure 2B). In accordance to these observations, only few cells were immunoreactive for differentiation markers β-III-tubulin (neuronal cells, 1.35±0.08%, data not shown), glial fibrillary acidic protein (GFAP) (glial cells, 3.0±0.5, Figure 2C), and O4 (oligodendrocytes, 7.6±0.9%, data not shown), suggesting the presence of a small subpopulation of spontaneously differentiated NPCs. Moreover, immunocytochemical staining demonstrated no expression of alpha smooth muscle actin (αSMA) (data not shown). Reverse transcription PCR (RT-PCR) analysis substantiated nestin expression in NPCs, further demonstrating the message of typical neural precursor and NSC markers Sox2, Sox9, Sox10, brain lipid-binding protein (BLBP), sonic hedgehog (SHH), vimentin, and Pax6 (Figure 2D). Because Sox10 is also known to be expressed in neural crest-derived stem cells (NCSCs) (reviewed by Kaltschmidt et al., 2012), we tested the potential expression of a further NCSC-specific transcription factor Snail and found no expression in cultivated NPCs (Figure 2D).

**Results**

**Adult hippocampal neural precursor cells can be efficiently cultivated as monolayer and show the ability to form neurospheres**

Isolated cells from the mouse hippocampus were cultured on poly-β-lysine and laminin-coated surface in NPC medium containing the growth factors EGF and FGF-2 (see Figure 1A). In general accordance with the findings by (Seaberg and van der Kooy, 2002), neural precursor cells (NPCs) began to grow adherently after 2 weeks. Proliferating adherent NPCs homogeneously exhibited characteristic elongated bipolar morphology with oral nuclei (Figure 1B) (Conti et al., 2005). After reaching about 80% confluency, cells could be harvested and passaged, followed by further cultivation in an adherent manner. Proving their NPC characteristics in the first place (Babu et al., 2007), NPCs were shown to form neurospheres under appropriate culture conditions (Figure 1C).
Figure 1 (A) Schematic diagram of the cultivation procedure. Hippocampi were carefully isolated and mechanically and enzymatically dissociated followed by cultivation of the resulting suspension on poly-d-lysine/laminin-coated cell culture dishes. NPCs cultivated adherently reach 80% confluency after at least 30 days. (B) Phase contrast image of NPCs cultivated adherently on poly-d-lysine/laminin in presence of EGF and FGF-2 showing typical neural progenitor cell morphology. (C) Phase contrast image of adult hippocampus-derived neurospheres using uncoated, low adhesion flasks.

Figure 2 Characterization of cultivated NPCs. (A, B) NPs cultures express neural precursor markers. Adherently grown NPCs were fixed and stained against the neural stem cell-specific intermediate filament nestin and neural precursor marker Olig2. DNA was counterstained using SYTOX green. Nestin was expressed in 98.0±1.3% of NPCs, whereas Olig2 was expressed in 80.4±1.6% of NPCs. (C) Small subpopulations of NPCs showed expression of GFAP (3.0±0.5). (D) RT-PCR analysis demonstrated expression of characteristic neural precursor markers nestin, Sox2, Sox9, Sox10, BLBP, SHH, vimentin, and Pax6, whereas neural crest stem cell-specific transcription factor Snail as well as MSC marker Thy1 were not detectable. Non-template approaches served as controls.
perivascular MSCs (Paul et al., 2012), we assured the absence of MSC-marker Thy1 (Figure 2D) in NPC cultures.

**Early-passage NPCs give rise to neuronal, glial, and oligodendrocytic cell types**

Determining the differentiation capability of NPCs, a spontaneous differentiation assay was used by applying DMEM/F-12 medium containing 10% fetal calf serum (FCS). Immunocytochemical stainings revealed expression of differentiation markers β-III-tubulin (neuronal cells, 10.63±1.59%), GFAP (glial cell, 25.70±2.23%), and O4 (oligodendrocytes, 27.28±0.47%) in spontaneously differentiated NPCs (Figure 3A), which was significantly increased compared with undifferentiated control approach. Differentiated NPCs further revealed significantly decreased expression of the NSC marker nestin (17.04±0.55%) and

![Figure 3](image_url)

**Figure 3** Early-passage NPCs give rise to neuronal, glial, and oligodendrocytic cells. (A) NPCs undergo differentiation into neuronal, glial, and oligodendrocytic cells. After cultivation for 7 days in the presence of FCS, NPCs were fixed and immunocytochemically stained against neuronal, glial, and oligodendrocyte markers. Confocal microscopy reveals change in morphology in numerous cells after differentiation and the presence of β-III-tubulin-positive neuronal cells, GFAP-expressing glia, and O4-positive oligodendrocyte-like cells. (B) Statistical evaluation reveals significantly decreased percentage of nestin-positive cells and increase of differentiation markers, namely 10.63±1.59% of β-III-tubulin positive cells, 25.70±2.23% of GFAP-expressing cells, and 27.28±0.47% of O4 immunoreactive cells. (C) NPCs reveal pronounced neuronal differentiation potential. NPCs were subjected to directed neuronal differentiation in the absence of growth factors and presence of 5 μM retinoic acid. Immunocytochemical staining revealed a highly increased percentage of β-III-tubulin-expressing cells (73±0.35%), whereas the amount of cells expressing nestin was decreased (5.18±3.50%) and the percentage of GFAP-positive cells only slightly changed (23.16±7.64%) compared with spontaneously differentiated NPCs.
the NPC marker Olig2 (55.24±1.90%), as shown by statistical evaluation of ICC analysis (Figure 3B).

Although NPCs showed only a low capability in spontaneously increasing neurons (10.63±1.59%), a directed neuronal differentiation assay resulted in 73% of β-III-tubulin immunoreactive neuronal cells, indicating a high neuronal differentiation potential. We also observed a further decrease in the amount of nestin-expressing cells (5.18±3.50%), whereas the percentage of GFAP-positive cells was found to be only slightly changed (23.16±7.64%) in comparison to spontaneously differentiated NPCs (Figure 3C).

Extensively cultivated NPCs reveal reduced potential for ectodermal differentiation

As the differentiation capability of a precursor cell is suggested to be changeable during culture (Morshead et al., 2002), we assessed the presence of respective differentiation markers in high-passage NPCs (>25) in a spontaneous differentiation assay. In comparison to early-passage NPCs (passage 4), significantly decreased differentiation into β-III-tubulin-positive neuronal cells and O4-expressing oligodendrocytes was observed in extensively cultivated NPCs. On the contrary, the percentage of undifferentiated nestin- and Olig2-positive NPCs was found to be increased within the passage 26 culture, whereas the amount of GFAP-expressing glial cells seemed unaffected (Figure 4A).

Early- and late-passage NPCs were able to give rise to early osteogenic cells under directed differentiation conditions

Further investigating the differentiation potential of late-passage NPCs, we assessed a potential crossing of germ layer boundaries by applying a directed differentiation assay into the osteogenic lineage. Broadly described to commonly induce osteogenic differentiation of MSCs (Jaiswal et al., 1997), the respective differentiation medium comprised dexamethasone, β-glycerophosphate, and L-ascorbic acid-2-phosphate. Here, early- and late-passage NPCs were found to give rise to early osteogenic cell types in a directed differentiation assay, indicated by changes in morphology from bipolar to cuboidal after 10 days of induction as well as by alkaline phosphatase (ALP) activity (data not shown, Figure 4B). These findings suggest a slightly enhanced plasticity of NPCs even in early passages. Although NPCs of all passages showed ALP activity, the amount of ALP-active cells was found to be significantly lower in early-passage NPCs in comparison to late-passage NPCs (p26) (see Figure 4C).

Long-term cultured NPCs spontaneously gave rise to αSMA-positive mesodermal as well as β-catenin-expressing pre-endodermal cells

Contrary to directed osteogenic differentiation conditions, we observed the presence of mesodermal αSMA-expressing cells in a spontaneous differentiation assay (in the presence of 10% FCS) in late-passage NPCs (Figure 4D), whereas early-passage NPCs showed no differentiation into αSMA-positive progeny (data not shown). In accordance to these findings, a small subset of spontaneously differentiated late-passage NPCs showed expression of β-catenin, an early endodermal marker (Sinner et al., 2004). We observed the presence of such β-catenin-positive cells again exclusively in differentiated late-passage NPCs (Figure 4D), in contrast to early-passage NPCs lacking β-catenin expression under the same differentiation conditions (data not shown).

Prolonged culture of NPCs results in enhanced osteogenic and adipogenic differentiation potential under specific differentiation conditions

To further determine potential effects of long-term culture on the plasticity of NPCs, we applied the directed osteogenic differentiation assay over a longer period. After exposure of high-passage NPCs to osteogenic differentiation medium for 10 days, we again observed changed cellular morphology from bipolar to cuboidal as well as ALP activity, suggesting a phenotype switch into immature osteoblast-like cells. NPCs osteogenically differentiated for 20 days further displayed calcium deposition as shown by intense Alizarin Red and von Kossa signal, whereas untreated controls showed no positive signals (Figure 5A). Importantly, no Alizarin Red and von Kossa-positive cells were founds in early-passage NPCs after 20 days of osteogenic differentiation (data not shown). RT-PCR analysis further substantiated osteogenic differentiation of long-term cultured NPCs by revealing expression of two specific bone matrix proteins osteopontin and osteonectin (Figure 5B).

Investigating the differentiation capacity of long-term cultured NPCs into mesenchymal cell types in more detail, we applied a directed adipogenic differentiation methods
developed for differentiation of MSCs (Janderova et al., 2003). Although a high degree of cell death was observed in NPC cultures during the differentiation assay (data not shown), few NPCs were shown to form Oil Red O-positive lipid droplets, suggesting their conversion into pre-adipogenic cells (Figure 5C). On the contrary, no Oil Red O-positive lipid droplets were observed in early-passage NPCs applied to the same differentiation conditions (data not shown). In accordance to these findings, we observed the expression of adipocyte marker lipoprotein lipase using RT-PCR in differentiated NPCs (Figure 5D).

Long- and intermediate-term cultured NPCs show aberrant chromosome numbers, whereas no abnormal karyotype was observed in early passages

Because fate switches crossing germ-layer boundaries are suggested to be associated with transformation events during in vitro culture (Morshead et al., 2002), we further assessed karyotypes of cultivated NPCs in early and late passages. As shown in Figure 6A, control
mouse embryonic fibroblasts (MEFs) and passage 6 NPCs showed normal karyotypes comprising 40 chromosomes. In contrast, NPCs cultivated up to passages 8, 10, 12, 19, and 27 revealed highly abnormal karyotypes, for instance, 68.5±1.8 chromosomes in passage 27 NPCs (Figure 6B).

### Extensively cultivated NPCs grow independently of EGF and FGF-2 without changes in clonal efficiency

In addition to the aneuploidy of long-term cultured NPCs during prolonged culture, we observed the formation of abnormal worm-like spherical structures in NPCs cultivated up to passage 27 under neurosphere conditions. In contrast, passage 6 NPCs revealed normal capability of sphere formation (Figure 6C) with round, densely packed neurospheres. Because such structures may be associated with enhanced growth capacities due to transformation events, we investigated potential differences in clonal growth between passage 6 and passage 27 NPCs. Long-term cultivated NPCs revealed no significantly changed clonal efficiency (Figure 6D) and proliferation capability, as depicted by no significant changes in total cell numbers between NPCs of passage 7 (2.11E+05±4.76E+04 cells), passage 13 (2.40E+05±4.09E+04 cells) and passage 22 (2.12E+05±4.15E+04 cells) after 96 h of cultivation. Extensively cultured NPCs were shown to grow independently to growth factors FGF-2 and EGF (Figure 6E).
Discussion

This study demonstrates for the first time a change of _in vitro_ plasticity in adult murine NPCs, which is associated closely with transformation events occurring during long-term culture. In particular, we demonstrate a dramatic reduction in neuronal and oligodendrocytic differentiation capacity and acquisition of mesodermal differentiation in late-passage NPCs.

With respect to their origin within the CNS, neural stem cells and precursor cells have been broadly described to possess the ability of differentiating into neurons, astrocytes, and oligodendrocytes (Reynolds and Weiss, 1992; Morshead et al., 1994; Gritti et al., 1996). Surprisingly,
adult NSCs and NPCs were described to harbor an extraordinary high degree of plasticity, suggested by trespassing of lineage restrictions. NSCs were shown to give rise to skeletal muscle cells by direct exposure to myoblasts (Galli et al., 2000) as well as to melanocytes, chondrocytes, and smooth muscle cells in co-culture with quail neural crest cells (Alexanian and Sieber-Blum, 2003). Wurmser et al. (2004) extended these reports by demonstrating the differentiation of NSCs into endothelial cells in vitro, evidenced by stable expression of respective markers and the capacity to form capillary networks.

Moreover, in vitro-cultivated NSCs were described to undergo differentiation into cells of all germ layers after injection into the chicken and mouse blastocyst, thereby contributing to the formation of chimeric embryos (Clarke et al., 2000). Transplantation of cultivated, genetically labeled NSCs into irradiated hosts was further suggested to result in differentiation into a broad range of blood cell types (Bjornson et al., 1999). On the contrary, the said differentiation capacity was questioned only 3 years later by Morshead et al. (2002), demonstrating no sign of NSC-derived hematopoietic cell types in 128 host animals after intravenous injection of NSCs. The authors also observed changes in growth properties, growth factor dependence, and cell cycle kinetics in cultured neurosphere cells, thereby assuming that the fate switch reported by Bjornson et al. (1999) may be due to rare transformation events (Morshead et al., 2002).

In case of adult stem cells, such transformation events are commonly known to be associated with prolonged in vitro culture (Izadpanah et al., 2008; reviewed by Widera et al., 2008; Rosland et al., 2009). With respect to neural stem cells, Wu et al. (2011) reported spontaneous malignant transformation of human NSCs cultured over an extended period (passage 17). Subsequent transplantation of transformed human NSCs into nude mice led to the formation of xenografts expressing neuroendocrine tumor markers, whereas a xenograft-derived cell line showed abnormal karyotype and enhanced proliferation. Stem cells from the adult rat SVZ were likewise shown to transform into tumorigenic cell lines after their expansion in vitro, as demonstrated by multiple acquired chromosomal aberrations and malignant tumor formation after syngeneic transplantation into the brain of adult rats (Siebzehrubl et al., 2009). As reported by Kaus et al. (2010), long-term culture of adult rat NSCs not only results in aberrant, aneuploid DNA contents, and malignant tumor formation after in vivo transplantation but also in cell growth independent to exogenous growth factors. Substantiating these findings, we observed abnormal chromosome numbers as well as growth factor-independent proliferation in NPCs from murine adult hippocampus after long- and intermediate-term culture.

Long-term culture of NPCs did not only result in cellular transformation but also in enhanced plasticity, particularly including successful differentiation of long-term cultivated NPCs into adipogenic and late osteogenic cell types. Besides this enhanced potential for mesodermal differentiation, we also observed a reduced ectodermal differentiation capacity in long-term cultivated NPCs. These findings are in accordance to several studies demonstrating a reduced potential of long-term cultured NPCs to undergo ectodermal differentiation (Bull and Bartlett, 2005; Louis et al., 2008). Our findings further indicate a slightly enhanced plasticity of NPCs even in early passages, a phenomenon revealed by the extraordinary high neuronal differentiation potential as well as the capability to give rise to early osteogenic cell types. However, in our hands, early-passage NPCs never gave rise to late osteogenic cell types and showed a lower potential to differentiate into early ones compared with late-passage NPCs. In addition, early-passage NPCs did not differentiate spontaneously into mesodermal and endodermal cell types. Given these data, we suggest that although fate switches may occur in early passages, their severity is highly increased during prolonged cultivation. In accordance to our further findings, fate switches are broadly associated with transformation events (Morshead et al., 2002; Terada et al., 2002; Ying et al., 2002; Lotem and Sachs, 2006), although a direct link between the osteogenic and adipogenic differentiation potential of NPCs and the transformation events associated with in vitro culture has not been described so far.

In vivo, a chromosome-specific accumulation of aneuploidy has been recently demonstrated with a frequency of 50% in aged mouse brain (Faggioli et al., 2012). However, because the NPCs described here showed normal karyotype in early passage, the aneuploidy shown in our study seems to be selected by prolonged culture.

In conclusion, we demonstrate here for the first time that cultivation of adult murine hippocampal NPCs is closely associated with cell fate switches from neuronal and oligodendrocytic toward the mesodermal lineage as well as aberrant karyotypes and growth factor-independent proliferation.

Our findings particularly emphasize the potential relevance of transformation events concerning the investigation of cellular plasticity of NPCs and NSCs. Moreover, this study underlines the importance of rigid quality control of the stem cells for basic research as well as for potential clinical use.
Materials and methods

Isolation of NPCs from adult mouse hippocampus

All tissue was extracted according to local (Bezirksregierung Düsseldorf) and international guidelines. In particular, adult 129/BL6 or C57/BL6 mice (between 2 and 4 months age) were killed by cervical dislocation. Complete brains were immediately removed. Hippocampi were carefully separated from the adjacent tissue, dissected, and chopped into small fragments using surgical disposable scalpels (Aesculap, Malsungen, Germany) and placed in ice-cold 1× HBSS (Life Technologies, Darmstadt, Germany) supplemented with 15 mm HEPES (pH 7.5; PAA, Pasching, Austria), d-glucose (5.4 mg/ml; Sigma-Aldrich, Munich, Germany), amphotericin B (1:100; PAA), and penicillin/streptomycin (1:100; PAA). Further stem cell-isolation procedure from the dissected tissue was performed as described by Widera et al. (2006).

Cultivation of NPCs

The cell suspension was plated on poly-d-lysine/laminin-coated dishes (for the coating protocol, see Martin et al., 2012) in serum-free DMEM/F12 (Biochrom AG, Berlin, Germany) supplemented with EGF (20 ng/ml; R&D Systems, Wiesbaden, Germany), FGF2 (60 ng/ml; laboratory made), 3× NS2/B27 supplement (Chen et al., 2008), and heparin (0.5 U/ml; Sigma-Aldrich) and subsequently cultivated at 37°C, 5% CO₂, and 5% O₂ until confluency of ~80% (at least for 30 days) with medium change every 2 days. Passaging was performed using Accutase (PAA) at 37°C for 15 min with subsequent cultivation as monolayer or as neurosphere cultures. For neurosphere cultivation, uncoated low-adherence 25-cm² Tissue Culture Flasks (Greiner Bio One, Frickenhausen, Germany) were used. Spheres were dissociated using Accutase at 37°C for 15 min.

Limited dilution assay

Limited dilution assay was used to examine the clonal efficiency of NPCs as described by Widera et al. (2009). In particular, NPCs were collected and dissociated as described above. Afterward, the dissociated cells were diluted to 1 cell/100 μl and placed into 96-well, round-bottom microwell plates in NPC medium. Four hours after plating, 384 wells were analyzed for the presence of single cells. Newly forming neurospheres were examined regularly using inverse microscopy (AMG EVOS xl; PeqLab, Erlangen, Germany) over a period of 1 month.

Immunocytochemistry

Cells were fixed with 4% paraformaldehyde (PFA; Sigma-Aldrich) for 20 min at room temperature (RT), followed by three wash steps with 1× PBS and permeabilization with 0.02% Triton X-100 (Sigma-Aldrich). Blocking was achieved using 5% of appropriate normal serum (Dianova, Hamburg, Germany) for 30 min at RT followed by incubation with primary antibodies (see Table 1). After three wash steps using 1× PBS, secondary antibodies were applied for 1 h at RT in the dark. The following secondary antibodies were used including Alexa Fluor 555 goat anti-mouse (A21422; 1:300; Life Technologies), Alexa Fluor 555 goat anti-rabbit (A21428; 1:300; Life Technologies), and Alexa Fluor 555 goat anti-mouse IgM (μ chain, A21426, 1:300; Molecular Probes/Life Technologies). Subsequently, the cells were washed three times with 1× PBS and the DNA was counterstained with SYTOX green (1:10,000; Life Technologies) supplemented with RNase A (500 U/ml; Thermo/Fermentas, St. Leon-Rot, Germany) followed by mounting using in Mowiol (Carl Roth). For O4 surface antigen staining, cells were washed in PBS supplemented with 0.1% bovine serum albumin followed by 20-min incubation with primary antibody. Afterward, cells were washed with 1× PBS and fixed for 15 min in 4% PFA. Secondary antibodies were applied as described above. Fluorescence imaging was performed using confocal laser scanning microscopy (LSM 510; Carl Zeiss).

Reverse transcription PCR

Total RNA was purified using RNeasy Mini Kit or RNeasy Micro Kit (Qiagen, Hilden, Germany) according to the manufacturer’s guidelines. The RNA concentration was measured and qualified with Nanodrop UV spectrophotometry (Peqlab). cDNA was generated using First Strand cDNA Synthesis Kit (Thermo/Fermentas) according to the manufacturer’s guidelines. PCR was performed using KAPA2G Robust PCR Kit (Peqlab) according to the manufacturer’s guidelines. For primer sequences, see Table 2.

Spontaneous differentiation

For spontaneous differentiation assay, cells were seeded at a density of 1×10⁶ cells/well onto 18-mm coverslips, placed in a 12-well plate, and cultured in DMEM/F-12 supplemented with P/S, amphotericin B, and 10% FCS (Sigma-Aldrich). Cultivation was performed for at least 10 days at 37°C, 5% CO₂, and atmospheric O₂ in a humidified incubator. The quantification was performed after immunocytochemical

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Table 1  List of primary antibodies used for ICC.

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<th>Antibody</th>
<th>Species of origin</th>
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</table>
staining for the respective marker using ImageJ software (NIH, Bethesda, MD, USA) and statistically evaluated using GraphPad Prism (GraphPad Software, La Jolla, CA, USA) (Student’s t-test).

**Directed neuronal differentiation**

For directed neuronal differentiation, cells were seeded at a density of $2 \times 10^5$ cells/well on PDL/laminin-coated 18-mm coverslips placed in a 12-well plate and pre-cultured for 48 h in NPC medium supplemented with 5 $\mu$m retinoic acid (Sigma-Aldrich). Afterward, the medium was switched to growth factor-free Neurobasal medium (Life Technologies) supplemented with 5 $\mu$m retinoic acid. Further cultivation was performed for at least 7 days at 37°C, 5% CO$_2$, and atmospheric O$_2$ in a humidified incubator.

**Osteogenic differentiation**

For osteogenic differentiation, cells were seeded at a density of $1 \times 10^5$ in DMEM high glucose (PAA) supplemented with P/S, amphotericin B, 10% FCS, insulin (2 $\mu$m; Sigma-Aldrich), 3-isobutyl-1-methylxanthine (500 $\mu$m; Sigma-Aldrich), dexamethasone (1 $\mu$m; Sigma-Aldrich), and indomethacin (200 $\mu$m; Sigma-Aldrich). After 8 days, adipogenic induction medium was completely removed and changed to regeneration medium composed of DMEM high glucose, 10% FCS and 2 $\mu$m insulin for 2 days. Cells were treated in this manner for 21 days. Adipogenic differentiation was assessed via Oil Red O staining and RT-PCR analysis.

**Adipogenic differentiation**

For adipogenic differentiation, cells were seeded at a density of $1 \times 10^5$ in DMEM high glucose (PAA) supplemented with P/S, amphotericin B, 10% FCS, insulin (2 $\mu$m; Sigma-Aldrich), 3-isobutyl-1-methylxanthine (500 $\mu$m; Sigma-Aldrich), dexamethasone (1 $\mu$m; Sigma-Aldrich), and indomethacin (200 $\mu$m; Sigma-Aldrich). After 8 days, adipogenic induction medium was completely removed and changed to regeneration medium composed of DMEM high glucose, 10% FCS and 2 $\mu$m insulin for 2 days. Cells were treated in this manner for 21 days. Adipogenic differentiation was assessed via Oil Red O staining and RT-PCR analysis.

**Detection of ALP activity**

After 10 days of differentiation, ALP activity in osteogenically induced NPCs was detected by determination of ALP activity after 10 days and histological stainings as well as PCR analysis after 20 days of differentiation.

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**Table 2** List of primers used for RT-PCR.

<table>
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<th>Product size (bp)</th>
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Alizarin Red S staining

After 20 days of differentiation, cells were fixed with 4% PFA for 15 min. After removal of fixation solution, cells were washed two times with ddH2O. Alizarin Red S staining solution (Waldeck GmbH) was incubated for 5 min at RT followed by rinsing with ddH2O. Staining was visualized using AMG EVOS xl inverse microscope (Peqlab).

Von Kossa staining

After 20 days of differentiation, cells were fixed with 4% PFA for 15 min and washed two times with ddH2O followed by incubation with 5% silver nitrate solution (Fluka Chemie AG) under ultraviolet light for 60 min followed by two wash steps with ddH2O. Subsequently, 5% sodium thiosulfate solution (resolved in ddH2O; Fluka Chemie AG) was applied for 3 min. Unreacted silver was removed followed by two wash steps with ddH2O and counterstaining for 5 min with Kernechtrot solution (100 ml ddH2O, 5 g aluminum sulfate, and 0.1% Kernechtrot; Merck, Darmstadt, Germany). Cells were washed once with 100% ethanol then twice with 96% ethanol. Staining was visualized with AMG EVOS xl inverse microscope (Peqlab).

Oil Red O staining

The medium was completely removed followed by two wash steps with ddH2O. Afterward, cells were fixed with 10% PFA for 1 h at RT. Oil Red O staining solution was prepared with three parts from Oil Red O Stock solution (0.5% Oil Red O resolved in isopropanol; Sigma-Aldrich) and 2 parts ddH2O. The mixture was filtered two times using Whatman paper to remove precipitates. Oil Red O staining solution was applied to the cells and incubated for 2 h at RT. The solution was subsequently removed and cells were washed two times with ddH2O. Staining was visualized using inverse microscopy AMG EVOS xl inverse microscope (Peqlab).

Preparation of NPCs metaphase chromosomes

Metaphase chromosomes of NPCs were prepared according as described by Kaus et al. (2010). In particular, 1×10⁶ NPCs of respective passages (p6, p12, p19, and p27) as well as MEFs were metaphase-arrested, treated with pre-warmed hypotonic solution, and progressively fixed in methanol/acetate. For chromosome spreading, the suspension was dropped onto coverslips and air dried. Staining of chromosomes was done using DAPI (50 ng/ml).

Proliferation assay of cultivated NPCs

Defined numbers of passage 7, passage 13, and passage 22 NPCs were cultivated in NPC medium for 4 days. Total cell numbers were determined after dissociation using Accutase at 37°C for 15 min. GraphPad Prism was applied for statistical analysis of cell proliferation (GraphPad Software, La Jolla, CA, USA).

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References


