Satellite glia of the adult dorsal root ganglia harbor stem cells that yield glia under physiological conditions and neurons in response to injury

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SUMMARY

The presence of putative stem/progenitor cells has been suggested in adult peripheral nervous system (PNS) tissue, including the dorsal root ganglion (DRG). To date, their identification and fate in pathophysiological conditions have not been addressed. Combining multiple in vitro and in vivo approaches, we identified the presence of stem cells in the adult DRG satellite glial population, and progenitors were present in the DRGs and sciatic nerve. Cell-specific transgenic mouse lines highlighted the proliferative potential of DRG stem cells and progenitors in vitro. DRG stem cells had gliogenic and neurogenic potentials, whereas progenitors were essentially gliogenic. Lineage tracing showed that, under physiological conditions, adult DRG stem cells maintained DRG homeostasis by supplying satellite glia. Under pathological conditions, adult DRG stem cells replaced DRG neurons lost to injury in addition of renewing the satellite glial pool. These novel findings open new avenues for development of therapeutic strategies targeting DRG stem cells for PNS disorders.

INTRODUCTION

The adult dorsal root ganglion (DRG) harbors cell bodies of sensory neurons that discriminate a large number of sensory stimuli because of a high degree of morphological and functional diversity (Usoskin et al., 2015). Adult DRG neurons are metabolically and physically supported by two types of glia. First, satellite glial cells (SGCs) extend thin processes around each neuronal body to form an envelope and provide a unique environment of neuroglial communication that regulates neuronal activity (Hanani, 2005; Pannese, 1981). Second, the well-known myelinating and non-myelinating Schwann cells support axons along the peripheral nerve. Macrophages, fibroblasts, and perivascular structures formed by endothelial cells and pericytes also contribute to DRG homeostasis.

Most adult DRG cells are neural crest cell derivatives that migrate extensively to their destination during development. However, whether undifferentiated stem or progenitor cells persist in the adult DRG is still a matter of controversy. Few studies support the possibility that the number of DRG sensory neurons increases progressively during life (Devor and Govrin-Lipmann, 1985), and others refute this hypothesis (La Forte et al., 1991; Pover et al., 1994). An average of 20%–50% of sensory DRG neurons die in response to axotomy (Groves et al., 2003) or capsaicin injection (Czaja et al., 2008). The total number of neurons recovers to normal levels over time in injured DRG of juvenile rats. Injection of bromodeoxyuridine (BrdU) combined with stereological quantification in injured rats revealed newly born cells comprising glial cells and a few neurons (Gallaher et al., 2010, 2014), suggesting, but not proving, replacement of adult DRG cells lost to disease by newly born ones.

The stemness potential (Snippert and Clevers, 2011) of adult DRG was also assessed in vitro based on the sphere forming capacity of explants (Li et al., 2007; Namaka et al., 2001). Cells emerging from explanted tissues have been shown to proliferate and form multipotent spheres giving rise to neurons, glial cells, and myofibroblasts. Adult DRG multipotency also occurs in vivo and is strongly modulated by environmental cues, with DRG sphere-forming cells giving rise to perivascular pericytes after transplantation in the murine developing brain but myelin-forming Schwann cells when grafted in demyelinated adult spinal cord (Vidal et al., 2015). Fate mapping studies using PO-Cre and Wnt1-Cre/floxed EGFP have demonstrated that the in vitro DRG stem cell (SC) properties belong essentially to the PNS neural crest-derived fraction (Nagoshi et al., 2008).

Although these studies provided some hints about the presence of adult DRG neurogenesis, the experiments were not conclusive because they were performed with mixed or global adult DRG populations. Late progenitors derived from multiple neural crest cells (Sabourin et al., 2009) or SCs (Vidal et al., 2015) that persist in the DRG through adulthood could have been the source of these newly formed neurons (Farel, 2003). Because of technical limitations, clear identification of the putative DRG SCs and their stemness behavior under physiological and pathological conditions was never established. This fundamental biological question is crucial at a clinical level.
because these presumed DRG SCs could be ideal targets to modulate in PNS disorders such as peripheral neuropathies. To resolve these unanswered questions, we combined multiple in vivo and in vitro approaches and SC-specific transgenic tools to identify, purify, and characterize the most putative DRG SCs in vitro and study their fate under physiological and pathological conditions in vivo. Taking the adult spinal cord as reference to validate our transgenic tools, we report that the adult DRG contains SCs and progenitor cells. Adult DRG SCs are a subpopulation of SGCs, which have the capacity to generate neurons and glia in vitro and in vivo in response to injury.

RESULTS

The adult mouse DRG contains slowly proliferating cells

Because the adult DRG is composed of multiple cell populations (Figure 1A), we first considered that each DRG cell type could constitute the putative DRG SCs. To eliminate some candidates, we used the EdU long-label retention assay to detect the presence of quiescent or slowly proliferating SCs in the adult DRG (Taupin, 2007) and discriminate them from the more proliferative progenitor cells. Spinal cords of the same animals were used as controls to validate our tools because they harbor neural SCs localized in the central canal wall (Meletis et al., 2008) and glial progenitors in the parenchyma (Horner et al., 2000) and referred to CNS SC.

The 5-Ethynyl-2’-deoxyuridine (EdU) incorporation assay without (Figures 1B–1D) or with 1-month chase (Figures 1E–1G) identified the presence of proliferating cells in the adult DRG (Figures 1B–1E). Their low number with chase (<1%) in 4-month-old mice was consistent with the relative quiescence of SCs. This assay validated the presence of EdU+ progenitor cells in the spinal cord parenchyma without chase only (Figures 1C and 1F) and EdU+ SCs in the central canal wall (Figures 1D and 1G).

We assumed that the more cells divide rapidly (i.e., progenitors), the more their EdU content is diluted. The EdU dilution effect was assessed by comparing EdU levels without and with chase for each tissue (DRG and spinal cord parenchyma or central canal) based on the percentage of BrdU staining overlap with Hoechst (EdU+low < 30%, EdU+medium < 66%, EdU+high > 66%). Without chase, the 3 regions contained a high percentage of EdU+low cells and a small percentage of EdU+medium and EdU+high cells (Figure 1J, light versus dark gray columns). Although, after the chase, the amount of EdU+low cells in the DRG and the parenchymal spinal cord increased slightly, EdU+high cells were detected in the DRG and the spinal cord central canal wall only (Figures 1J and 1K). Values of the EdU dilution pattern of DRG cells were comprised between the spinal cord parenchymal progenitors and the spinal cord central canal SCs, suggesting that DRG SCs proliferate at a different rate than spinal cord SCs or that the DRG contains multiple populations of proliferating cells.

Without excluding the first possibility, we validated the second one, characterizing the EdU+ populations in these different tissues. The marker cell specificity according to tissues type is listed in Table S1. In the spinal cord central canal, EdU+ cells were IBA1+ microglia/macrophages and Glial Fibrillary Acidic Protein+ (GFAP+) SCs (Figures S1A–S1C), and in the spinal cord parenchyma, EdU+ cells were IBA1+ microglia and NG2+ oligodendrocyte progenitors but not GFAP+ astrocytes (Figures S1D–S1F). In the DRG, EdU+ cells were subdivided into at least 3 distinct populations comprising IBA1+ microglia/macrophages (45% ± 4%; Figures 1H and 1L), GFAP+ perineuronal SGCs (30% ± 3%; Figures 1I and 1L), and an unidentified population (25% ± 3%; Figure 1L). This population included possibly pericytes, fibroblasts, endothelial cells, non-myelinating Schwann cells, or SGCs that did not express GFAP, given that there is no specific and selective marker for the entire SGC population. Therefore, we define SGCs as DRG cells that are labeled with GFAP/SOX10 and display morphological features consistent with perineuronal ensheathment.

Even though, at this point, several DRG cell types remained SC candidates, some could be excluded de facto from the list. These comprised myelinating Schwann cells and neurons that do not proliferate and epineurial fibroblasts localized at the DRG periphery that proliferate too rapidly to retain EdU during the chase period (compare

Figure 1. In Situ characterization of slowly proliferating cells in the adult DRG

(A) Schematic of the different adult DRG cell types.
(B–I) Immunodetection of EdU+ cells (arrows) without or with chase in the DRG (B, E, H, and I), spinal cord parenchyma (C and F), and spinal cord central canal (D and G).
(H and I) DRG cells immunolabeled for EdU, IBA1, or GFAP.
(J and K) Comparative EdU dilution pattern before and after chase (J) and between the three different regions after chase (K).
(L) Proportion of the different EdU+ cell types after chase.
Scale bars, 50 μm. Boxed areas are enlarged in the right panels. Data are shown as mean ± SEM (n = 3–4). Two-way ANOVA and multiple comparisons: Dunnett’s test (J and K), Tukey’s test (L); **p < 0.01, *p < 0.05. See also Figure S1.
Because of the low proportion of EdU$^{\text{high}}$ cells and the lack of tools specifically discriminating each DRG cell type from potential immature cells, identification and characterization of the putative DRG SCs remained difficult using solely the EdU incorporation assay.

**Figure 2.** Characterization of transgene expression in Nestin-GFPcyt and Nestin-CFPnuc transgenic spinal cords

(A–L) Double immunohistochemistry for GFP (A–F) and CFP (G–L) and cell-specific markers in the spinal cord parenchyma (A–C and G–I) or central canal (D–F and J–L): GFAP$^+$ astrocytes (A, D, G, and J), IBA1$^+$ microglia (B, E, H, and K), and NG2$^+$ oligodendrocyte progenitors (C, inset a, and I, inset a) or pericytes (C, inset b; I, inset b; and L). Boxed areas are enlarged in the right panels.

(M and N) Visualization by EM of immunogold-labeled ependymal cell nuclei in the Nestin-CFPnuc spinal cord. E, ependymal cell.

Scale bars: 50 μm (A–L), 5 μm (M and N), and 1 μm (E2).
Figure 3. Characterization of GFPcyt+ and CFPnuc+ cells in the adult DRGs of Nestin-GFPcyt and Nestin-CFPnuc reporter mice
(A–L) Immunodetection and quantification of EdU+ cells in Nestin-GFPcyt (A–F) or Nestin-CFPnuc (G–L) mice.
(C–F and I–L) Double immunohistochemistry to reveal GFP (C–F) or CFP (I–L) and IBA1+ macrophages (C and I), PV25+ proprioceptive
neurons (D and J), GFAP+ glial cells (E and K), and NG2+ SGCs (F, inset a, and L, inset a) or pericytes (F, inset b, and L, inset b).

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Validation of the Nestin-GFPcyt and Nestin-CFPnuc transgenic lines to identify SCs and progenitor cells in the adult spinal cord

Nestin is a marker of CNS SCs. To clearly identify the putative DRG SCs in situ and isolate them in vitro, we explored use of Nestin-Green Fluorescent Protein (GFP) cytoplasmic (GFPcyt) (Mignone et al., 2004) and Nestin-Constitutively Fluorescent Protein (CFP) nuclear (CFPnuc) (Encinas et al., 2006) transgenic mice. The Nestin-GFP line encompasses stem/early progenitors and late progenitor populations in the brain (Mignone et al., 2004), and the Nestin-CFP line labels stem/early progenitors only (Encinas et al., 2006).

First we validated transgene expression in the adult spinal cord as a positive control in the same tissue section of connected DRG. Three populations of GFP+ cells were detected in the Nestin-GFPcyt spinal cord (Figure 2). These included parenchymal GFAP+ astrocytes (Figure 2A) and parenchymal NG2+ oligodendrocyte progenitors and pericytes (Figure 2C) in addition to ependyma-associated GFAP+ SC population (Figure 2J) but excluded from the parenchymal astrocyte, microglia/macrophage, and oligodendrocyte progenitor populations (Figures 2G and 2I). These results were confirmed by immunogold electron microscopy for CFPnuc based on the presence of immunogold particles restricted to the nucleus of ependymal ciliated cells of the central canal (Figures 2M and 2N). These immunofluorescent and immunogold electron microscopy observations in the adult spinal cord validate those in the adult brain for late progenitors in GFP mice (Mignone et al., 2004) but SCs/early progenitors in CFP mice (Encinas et al., 2006).

The Nestin-GFPcyt mouse line labels three distinct cell types in the DRG

Nestin-GFPcyt+ and Nestin-CFPnuc+ cells were also found in the DRG (Figure 3), raising the possibility that the adult DRG contains SCs and progenitor cells. Using the EdU chase assay as above, we defined that some DRG GFPcyt+ (Figure 3A) and CFPnuc+ cells (Figure 3G) were also EdU+. The percentage of EdU+/GFPcyt+ cells over the entire DRG EdU+ population was superior to that of the EdU+/CFPnuc+ population after chase (37% ± 2% versus 21% ± 4%, p = 0.03) (compare Figures 3B and 3H), suggesting that a difference between the two transgenic lines existed in the DRG as well. To validate this hypothesis and identify the putative DRG stem and progenitor cells, we characterized the DRG GFPcyt+ and GFPcyt+ cells by immunohistochemistry.

DRG GFPcyt+ cells were localized around DRG neurons (Figure 3D), a location typical for the previously described neuroglial unit (Hanani, 2005; Milosavljević et al., 2021; Pannese, 1981). The great majority of these perineuronal GFPcyt+ cells expressed the SGC markers GFAP (78% ± 9%; Figure 3E) and SOX10 (47% ± 17%; Figure S2), which label neural crest cells and their derived progeny including SGCs (Britsch et al., 2001), and the oligodendrocyte progenitor marker NG2, also known to be expressed in SGCs (Rezajooi et al., 2004; Figure 3F, inset a). Perineuronal GFPcyt+ cells never expressed IBA1 (Figure 3C) nor the neuronal marker PV25 (Figure 3D). However, not all of the Nestin-GFPcyt+ cells were associated with neurons because some of them were located in the endoneurium. These endoneurial Nestin-GFPcyt+ cells expressed NG2 (Figure 3F, inset b), the endoneurial fibroblast/endothelial marker CD34 (Hirose et al., 2003; Richard et al., 2012; Figure S3B), or the pericyte marker CD13 (Figure S3C), but none expressed the myelinating Schwann cell marker P0 (Figure S3A).

These data indicate that the GFPcyt+ population of the Nestin-GFPcyt adult PNS comprised pericytes and endoneurial fibroblasts in addition to SGCs.

The Nestin-CFPnuc mouse line labels a subpopulation of DRG satellite cells

In contrast, CFPnuc expression in the DRG of Nestin-CFPnuc+ transgenic mouse line was restricted to the ependymal-associated GFAP+ SC population (Figure 2J) but excluded from the parenchymal astrocyte, microglia/macrophage, and oligodendrocyte progenitor populations (Figures 2G and 2I). These results were confirmed by immunogold electron microscopy of CFPnuc based on the presence of immunogold particles restricted to the nucleus of ependymal ciliated cells of the central canal (Figures 2M and 2N). These immunofluorescent and immunogold electron microscopy observations in the adult spinal cord validate those in the adult brain for late progenitors in GFP mice (Mignone et al., 2004) but SCs/early progenitors in CFP mice (Encinas et al., 2006).

(M and N) EM analysis of Nestin-CFPnuc DRGs after CFP immunogold labeling. Gold particles (arrowheads) were detected only in SGC nuclei (M, inset a) and were absent from neuron (M, inset b), pericyte (N, inset a), and endothelial cell (N, inset b) nuclei. Blue and white dotted lines delineate SGCs and neurons, respectively. Boxed areas are enlarged in the right panels. N, nucleus. Scale bars: 50 μm (A–L) and 2 μm (M and N). Data are shown as mean ± SEM (n = 3), two-way ANOVA. See also Figures S2 and S3.
encasing individual neuronal cell bodies, corresponding, by definition, to SGCs (Figure 3M, inset a). The nuclei of the other DRG cell types, including neurons (Figure 3M, inset b), pericytes, endothelial cells (Figure 3N, insets a and b), myelinating Schwann cells (Figures S3G and S3H), macrophages (Figures S3I and S3J), or fibroblasts (Figures S3K and S3L), were never immunolabeled. When comparing the two lines, the fraction of perineuronal GFAP⁺ SGCs expressing CFPnuc (50% ± 7%) was reduced compared with those expressing GFPcyt (89% ± 6%).

The present data indicate that putative progenitors and SCs are present in the adult DRG. Although Nestin-GFPcyt mice had identified parenchymal progenitors and perineuronal SCs in the DRGs, the Nestin-CFPnuc mouse line traced only the presumptive DRG SCs with clear features of SGCs.

Figure 4. In vitro analysis of SC properties of purified Nestin-CFPnuc⁺ and Nestin-CFPnuc⁻ cells
(A–G) Ten days after in vitro expansion under subclonal conditions (<5 000 cells/mL) in EGF/FGF, FACS-purified adult CFPnuc⁺ spinal cord cells (SpCs) (A and B) or CFPnuc⁺ DRG cells (F and G) gave rise to CFP⁺ spheres.
(C–I) After growth factor removal and adhesion, spinal cord primary spheres differentiated into typical CNPase⁺ multi-branched oligodendrocytes (C), GFAP⁺ ramified astrocytes (D), and MAP2–MAP5⁺ process-extending neurons (E), whereas DRG primary spheres differentiated into GFAP⁺ bipolar spindle-shaped Schwann cells (H) and MAP2–MAP5⁺ process-extending neurons (I).
(J and K) Phase illumination shows that, after FACS purification, CFPnuc⁻ SpCs died over time (J), whereas CFPnuc⁺ DRG cells generated adhering bipolar cells (K).
(L and M) After growth factor removal, CFPnuc⁺ DRG primary spheres differentiated into αSMA⁺ flat fibroblasts, IBA1⁺ pancake-shaped microglia/macrophages (L), and GFAP⁺ spindle-shaped PNS glia but not into MAP2–MAP5⁺ neurons (M).

Scale bars: 20 μm (A–G, J, and K) and 100 μm (C–E, H, I, L, and M). See also Figure S4.
Figure 5. Fate mapping of adult DRG SCs
(A) Schematic representation of tamoxifen induction and temporal analysis of Nestin-CreER\textsuperscript{T2}/Rosa26-YFP mice.
(B and C) Detection of adult DRG-YFP\textsuperscript{+} cells 2 and 90 days after tamoxifen induction (dpi).
(D) Number of DRG transgenic positive cells per square millimeter at 2 dpi in CFPnuc mice and ipsi/contra injured or non-injured mice.
(E) Temporal evolution of the number of DRG-YFP\textsuperscript{+} cells per square millimeter.

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DRG SCs give rise to neurons and glia in vitro

Taking advantage of the high specificity of the Nestin-CFPnuc transgenic line, we purified, by fluorescence-activated cell sorting (FACS), the putative DRG SCs to establish their stemness potential in vitro, taking the spinal cord as a positive control. After FACS and seeding, all spinal cord and DRG CFPnuc⁺ viable cells expressed the transgene, validating their successful purification (Figure S4).

First, we obtained proof that the FACS-purified spinal cord CFPnuc⁺ fraction was the only fraction capable to form free-floating spheres (Figures 4A and 4B), whereas, under the same growth conditions, the spinal cord CFPnuc⁻ fraction did not generate spheres and died (Figure 4J). After adhesion and growth factor retrieval to induce differentiation, spinal cord CFPnuc⁺ spheres were tripotent, as described originally (Meletis et al., 2008; Reynolds and Weiss, 1992), differentiating into oligodendrocytes 2', 3' Cyclic Nucleotide 3' Phosphodiesterase + (CNPase⁺), astrocytes (GFAP⁺), and neurons (MAP2–MAP5⁺; Figures 4C–4E), validating that the Nestin-CFPnuc mouse line identifies CNS SCs.

Next we isolated, by FACS, the Nestin-CFPnuc⁺ and Nestin-CFPnuc⁻ fractions from the DRGs. After 10 days under epidermal growth factor (EGF)/fibroblast growth factor (FGF) conditions, cells of the positive fraction were able to proliferate and formed free-floating spheres that maintained Nestin-CFPnuc expression (Figures 4F and 4G). In the absence of EGF/FGF, these CFPnuc⁺ spheres differentiated, in great majority, into the major DRG constituents, including bipolar Schwann cells (GFAP⁺), but also into typical neurons (MAP2–MAP5⁺) with round cell bodies and long processes but never into Smooth Muscle Actin⁺ (SMA⁺) flat myofibroblasts (Figures 4H and 4I). The observation of de novo neurogenesis and in vitro bipotency of the FACS-sorted DRG CFPnuc⁺ cells reinforced the stemness of the putative DRG SCs. Under proliferation conditions, the DRG CFPnuc⁻ fraction, unlike the spinal cord CFPnuc⁺ fraction, generated adhering bipolar cells (Figure 4K) that eventually formed aggregates at later times. Under differentiation conditions, CFPnuc⁻ cells gave rise to flat fibroblasts or pericytes, microglia/macrophages, and spindle-shaped Schwann cells but never to neurons (Figures 4L and 4M).

Fate mapping of adult DRG SCs highlights their gliogenic potential in physiological condition

We showed that, under healthy conditions, DRG SCs proliferate slowly in situ without elucidating the phenotype of their newly born cells. In vitro, adult DRG SCs formed bipotent spheres giving rise to neurons and glial cells. Because the “stemness potential” of the DRG SC could have been overestimated in vitro (Singec et al., 2006; Snippert and Clevers, 2011), we questioned whether these newly identified cells behaved as bona fide SCs in vivo and whether they participated in physiological and/or pathological tissue homeostasis. To fate map the progeny of the adult DRG SCs, we crossed the Nestin-CreERT² line with the Rosa26-Yellow Fluorescent Protein (YFP) reporter line (Imayoshi et al., 2006). Nestin-CreERT² activity was induced by daily tamoxifen gavage during 5 days in 3- to 4-month-old Nestin-CreERT²:Rosa26-YFP mice (Figure 5A). To evaluate the percentage of recombined DRG SCs in Nestin-CreERT²:Rosa26-YFP mice, we compared the density of DRG transgene-expressing cells (number of YFP⁺ cells per square millimeter) with that of CFPnuc mice. At 2 days post induction (dpi), their number was significantly lower in non-injured (control, 663 ± 60) Nestin-CreERT²:Rosa26-YFP DRGs compared with CFPnuc DRGs (1,340 ± 180), suggesting a YFP recombination efficiency of the DRG SC population of around 44% (Figure 5D).

After a resting period of 10 days to eliminate systemic tamoxifen, the permanently recombined YFP⁺ cells were analyzed in non-injured animals at different time points after induction under physiological conditions (Figures 5B and SC). YFP⁺ cells were never detected in sham-induced animals (oil administration only; data not shown). We next examined the evolution of this YFP⁺ population by quantifying their number at 2, 15, 60, and 90 dpi (Figure 5E). The DRG-YFP⁺ cell density (total amount per square millimeter) remained stable over time, but a progressive but not significant increase occurred with time (664 ± 65, 749 ± 87, 755 ± 144, and 821 ± 195 YFP⁺ cells/mm² at 2, 15, 60, and 90 dpi, respectively), suggesting slow proliferation of DRG SCs or frequent turnover of their progeny under healthy endogenous conditions.

YFP⁺ SGC cells, identified by their perineuronal, small crescent moon-like phenotype and strongly Hoechst-stained nuclei, were easily distinguished from YFP⁺ DRG neurons, which had large round cell bodies and large faintly Hoechst-stained nuclei. Based on these criteria, all recombined YFP⁺ cells at 2 dpi were SGCs and not neurons (Figures 5B, 5F, 7A, and 7D). Immunohistocharacterization at 2 dpi demonstrated that YFP expression was restricted to typical perineuronal SGCs (GFAP⁺: 92% ± 6%, Figure S5A; SOX10⁺: 69% ± 11%, Figures 5F and 5G). Recombined

(F and G) Immunodetection of adult DRG-YFP⁺ cells with SOX10 (glial cells) and NEU-N (neurons) at 2 (F) and 90 dpi (G) in non-injured Nestin-CreERT²/Rosa26-YFP mice. Boxed areas are enlarged in the right panels.

Data are shown as mean ± SEM (n = 3). One-way ANOVA and multiple comparisons: Dunnett’s test (D) and Tukey’s test (E); **p < 0.01, *p < 0.05. See also Figure S5.
Figure 6. Effect of sciatic nerve axotomy on DRG neurons in Nestin-Cre-ERT2/Rosa26-YFP DRGs

(A–D) Immunohistochemistry for YFP, activated caspase-3, and NEU-N on the DRG contra side or ipsi side at 15 dpi (A and C) and 60 dpi (B and D). Boxed areas are enlarged in the right panels.

(E) Temporal evolution of the density of apoptotic caspase-3+ cells (contra and ipsi sides) after axotomy.

(F) Proportion of apoptotic neurons (caspase-3+) over the total neuronal (NEU-N+) population.

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YFP+ cells never expressed markers of neurons (Neuronal Nuclear Antigen (NEU-N); Figures 5F and 5G) or of other DRG cell types (Figures S5C–S5H), such as macrophages (IBA1), Schwann cells (P0), pericytes (CD13), or endothelial cells/fibroblasts (CD34). Analysis of the YFP population phenotype at 90 dpi indicated that, under physiological conditions, the DRG-YFP+ progeny had the unique fate to become SGCs (GFAP+, 89% ± 6%; SOX10+, 67% ± 0.7%) because no cell type other than SGCs expressed YFP (Figures 5G and S5B). These results validate that, under physiological conditions, YFP expression was restricted to a subpopulation of SGCs, DRG SCs, that expressed Nestin, as described for Nestin-CFPnuc mice.

**Long-term lineage tracing of adult DRG SCs reveals their glio- and neurogenic potential after PNS injury**

To examine the role of DRG SCs under pathological conditions, we challenged their response to nerve injury and questioned their participation in the regenerative process, using the same tracing paradigm as under physiological conditions (Figure 5A). Previous studies indicated that a variety of cellular changes occur in DRGs after sciatic nerve axotomy, including death of neurons and SGCs, followed by proliferation of microglia/macrophages, Schwann cells, and SGCs (Cherkas et al., 2004; Hu and McLachlan, 2002; Pierucci and de Oliveira, 2006). We therefore performed unilateral sciatic nerve axotomy after tamoxifen induction of Nestin-CreERT2/Rosa26-YFP (Nestin-YFP) mice. Tamoxifen induction followed by a 10-day chase prior to axotomy ensured that all recombined DRG cells were DRG SCs and that some did not dedifferentiate or become reactivated cells, such as Schwann cells re-expressing Nestin after injury (Jessen and Mirsky, 2008). Evaluation of the YFP population at 2 dpi showed no significant difference in the density of YFP+ (number per square millimeter) before (663 ± 60) and after injury (ipsilateral side, 515 ± 180; contralateral side, 493 ± 134) (Figure 5D).

Taking the contralateral side of the axotomy as a control, we next temporally assessed the effect of axotomy on cell death in the connected DRGs (L3, L4, and L5) from 2–90 dpi. Caspase-3 immunodetection revealed that cell death was induced in the ipsilateral DRG (Figures 6A and 6D) but not on the contralateral side (Figures 6A and 6B). DRG apoptosis was regulated temporally with a first wave of activated caspase-3+ apoptotic cells appearing at 15 dpi after axotomy in the ipsilateral DRG (significantly higher than at 2 dpi, p = 0.006) (Figure 6E) and remaining stable up to 30 dpi, followed by a second wave of apoptosis at 60 dpi in the ipsilateral DRG (significantly higher than at 2 and 15 dpi, p ≤ 0.0001) and compared with the contralateral side. Although SGC death occurred, in response to sciatic nerve axotomy, as reported previously (McKay Hart et al., 2002; Pierucci and de Oliveira, 2006), we focused our analysis on neuronal death because neurons are the primary target of injury (Pierucci and de Oliveira, 2006). Immunodetection of cleaved caspase-3 together with NEU-N revealed that some of the dying cells were neurons, with a peak of apoptotic neurons at 15 dpi. Although the number of dying neurons was low (0.63% ± 0.3%), neuronal death persisted up to 90 dpi, the latest time point analyzed (Figure 6F). When the numbers of ipsi- versus contralateral neurons per square millimeter were compared (Figure 6G), a slow and progressive loss of neurons was observed in the ipsilateral DRGs up to 30 dpi, with an ipsi-contralateral ratio of 0.85 ± 0.01/mm2 between 30 and 90 dpi. The total number of neurons per unit area in the ipsilateral DRG was significantly lower than in the control non-injured DRG at 15 (p = 0.033) and 60 dpi (p = 0.018) (Figure 6H).

Thus, unilateral sciatic nerve axotomy had an effect on its connected DRGs by inducing cell death over time, including that of neurons.

**SGCs are known to be activated under pathological conditions** (Hanani et al., 2002). We therefore tested whether these “activated SGCs” corresponded to DRG SCs. Taking the contralateral side as a control to normalize the recombination bias between animals (Figures 7A–7C versus 7D–7F), we observed a first significant peak of the YFP+ cell ratio at 15 dpi (p = 0.042), followed by a second peak at 60 dpi (p = 0.011) (Figure 7G). Thus, the ipsilateral YFP+ DRG SCs reacted to PNS injury by increasing cell numbers compared with the contralateral DRG SCs. The YFP+ DRG SC pattern of activation correlated with the waves of DRG cell death (compare Figures 6E and 7G), suggesting their contribution to the DRG regenerative process elicited after injury.

Immunocolorization of YFP+ cells and their progeny at 2 and 90 dpi in the ipsilateral DRGs of injured animals provided evidence showing that the great majority of the newly generated cells were GFAP+ SGCs (ipsilateral [ipsi] 78% ± 5% and contralateral [contra] 87% ± 6% cells at 2 dpi ipsi 78% ± 6% and contra 72% ± 7% at 90 dpi) (Figure S5) and, to a minor extent, SOX10+ SGCs (ipsi 30% ± 10% and contra 55% ± 15% at 2 dpi and ipsi 44% ± 11% and contra 34% ± 12% at 90 dpi) (Figures 7H and 7J). Large YFP+ cells expressing NEU-N were detected only in the

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**Additional annotation:**

- **(G) Ratio of ipsi over contra DRG neuronal density over time.**
- **(H) Histogram of DRG neuron density over time in control non-injured and contra and ipsi injured mice.**
- **Scale bars:** 50 μm (A–D). Data are shown as mean ± SEM (n = 3). Two-way ANOVA and multiple comparisons: Tukey’s test (E and F), Tukey’s test (F), and Dunnett’s test (G); ****p < 0.0001, ***p < 0.001, **p < 0.01, *p < 0.05.
ipsilateral DRG of axotomized Nestin-CreER\textsuperscript{T2}/Rosa26-YFP mice (Figures 7 I, 7K, and S6G–S6I) mice. Quantification of the percentage of NEU-N+/YFP+ cells over YFP+ cells (ipsi, 0% and 0.24% ± 0.11% at 2 and 90 dpi, respectively; contra, 0% at 2 and 90 dpi) indicated that in vivo DRG neurogenesis remains a rare event. The remaining DRG-YFP+ cells tested at 2 and 90 dpi were never microglia/macrophages (IBA1\textsuperscript{+}), myelinating SCs (P0\textsuperscript{+}), pericytes (CD13\textsuperscript{+}), or fibroblasts/endothelial cells (CD34\textsuperscript{+}) (Figures S6A–S6F). Thus, neurogenesis, in addition to gliogenesis, occurred in the adult injured DRG with newly born neurons generated from slowly proliferating Nestin-YFP+ DRG SCs.

DISCUSSION

The observation that dissociated DRG cells can proliferate and form spheres that generate neurons, glia, and myofibroblasts in vitro suggests the presence of SCs in adult DRGs (Li et al., 2007; Nagoshi et al., 2008; Namaka et al., 2001; Vidal...
et al., 2015). However, their stemness characteristics remained difficult to prove because of technical limitations related to stereological quantification and the lack of specific markers to unambiguously trace the neurogenic population. Therefore, the identity of adult DRG SCs as well as their role under physiological and pathological conditions remained elusive. We used multiple approaches to unambiguously demonstrate the presence of a novel Nestin-expressing cell in the adult DRG SGC compartment. These cells have characteristics of adult SCs in situ and in vivo, generating new SGCs under physiological conditions but neurons in addition to SGCs, when required, to replace neurons lost to injury.

**Progenitors and SCs are present in the adult mouse DRG**

We took advantage of two Nestin reporter mice to uncover the identity of the putative DRG SC candidate. Because of the more restricted expression of the Nestin-CFPnuc line, which labels CNS SCs/early progenitors in the spinal cord and a subpopulation of SGC in the DRG, we were able to isolate and study the stemness potential of the DRG CFPnuc+ fraction. Their unique in situ perineuronal location and co-expression of CFP with cell markers such as GFAP, SOX10, and NG2 and EdU incorporation allowed their unambiguous identification as SGCs. In culture, these DRG CFPnuc+ cells were able to form bipotent spheres. In the absence of growth factors and adhesive conditions, CFPnuc+ cells differentiated into neurons and spindle-shaped glial cells. However, in contrast to previous in vitro studies using whole DRG preparations (Li et al., 2007; Nagoshi et al., 2008; Vidal et al., 2015), the DRG CFPnuc+ fraction never generated myofibroblasts. Thus, we formulated the hypothesis that, in addition to SCs, the adult DRG contains progenitors, whose identity remains to be unraveled.

The EdU incorporation assay in non-injured wild-type mice allowed us to identify the presence of multiple proliferating cell types in the DRG. We found EdU+ cells with a typical SGC morphology in addition to other EdU+ PNS cell types, such as the slowly proliferating resident macrophages (Müller et al., 2010), pericytes, and a few fibroblasts. This is in contrast to a previous report stating that SGCs are the only slowly proliferating cells of the DRG under physiological conditions (Gallaher et al., 2014). It also highlights that the EdU incorporation assay alone is not sufficient to identify a putative SC without avoiding detection of other misleading candidates. However, use of this assay reduced the number of potential candidates under physiological conditions because myelinating Schwann cells and endothelial cells did not incorporate EdU.

We also found that not all PNS and CNS EdU+ were GFPcyt+ or CFPnuc+. These cells could have been the differentiated progeny of GFPcyt+ or CFPnuc+ cells that lost Nestin expression during the differentiation process. Alternatively, these EdU+/Nestin− cells could correspond to a cell type that has a physiological turnover (EdU+) without stem or progenitor properties (GFP+ or CFP+), as observed for the PNS/CNS-resident immune cells (Müller et al., 2010). These observations stress the necessity to use multiple transgenic tools combined with in situ and in vitro characterization to carefully and correctly separate progenitors from SC populations. Because of this meticulous analysis, the present study is the first to clearly identify and characterize SCs and progenitors in the adult DRG. Adult sciatic nerves were analyzed in parallel to the DRG and spinal cord assays. Although rare Edu+ cells were present in this tissue, CFPnuc+ cells were never detected in adult sciatic nerves of Nestin-CFPnuc mice when analyzed in situ. Although sciatic nerves generated spheres with time in culture, as demonstrated previously (Takagi et al., 2011), they never generated neurons in vitro. These combined in situ and in vitro observations infer that adult sciatic nerves do not harbor SCs but growth-responsive cells, including macrophages, progenitors, and cells like Schwann cells, that have the capacity to dedifferentiate and proliferate (Mirsyky et al., 2008).

**DRG SCs have gliogenic and neurogenic fates, depending on the environmental context**

Using genetic fate mapping, we demonstrate that adult DRG SC are responsible for DRG gliogenesis and sustain an important turnover of the mature SGC population under physiological conditions. The proliferation capacity attributed previously to SGCs reflects, in fact, a cell-specific characteristic restricted to a subpopulation of SGCs, the Nestin-YFP+ subpopulation. Based on the Edu-CFPnuc assay, only a low percentage of CFPnuc+ DRG SCs (20%) over the total EdU+ population (1.77%) proliferated at a slow rate under physiological conditions (Figures 1D and 3H). The results obtained with the Nestin-CreER line under physiological conditions indicated that, as in the CFPnuc line, the recombined YFP+ cells correspond strictly to a subpopulation of SGCs, with numbers remaining stable over time (Figure 5E). This suggests a homeostatic equilibrium between slow YFP+ SC proliferation and the slow death of their mature SGC-YFP+ progeny to ensure maintenance of the SGC population throughout adulthood.

Proliferation of Nestin-Cre-YFP+ cells increased up to 2 months after injury without reaching a normal plateau, suggesting concomitant DRG SC reactivation and neuronal loss. Characterization of the activated DRG SC progeny in this injury context confirmed the SGC gliogenic potential and neurogenic potential under pathological conditions. The function of mature SGCs under...
pathophysiological conditions and their major involvement in neuroprotection of DRG neurons has been highlighted previously (Hanani and Spray, 2020). SGC accomplish re-uptake of neuroactive substances or peripheral toxins, making them more susceptible to death than other DRG cell types (Scheffler et al., 2010). Thus under pathophysiological conditions, adult DRG SCs may have a dual role. First, they play a crucial role in maintaining the level of mature SGCs, ensuring neuronal survival throughout life. Second, they generate neurons in addition to their glial progeny to refuel the pool of neurons (and glia) lost to axotomy. Although significant, neurogenesis remained a minor event compared with gliogenesis. The cell fate plasticity of DRG SCs is similar in many aspects to the glial SOX10+ SCs of the enteric nervous system, which have gliogenic potential under physiological conditions but give rise to neurons after enteric neuronal loss (Laranjeira et al., 2011).

In contrast to CNS SCs, which are restricted to specific brain and spinal cord compartments, DRG SCs seem to be dispersed in their entire tissue to quickly sense environmental changes and efficiently renew the SGC pool of the DRG neuron-glia unit (one neuron and its SGCs). Similar SCs could exist in other PNS tissues, where they mainly generate glia under physiological conditions without being confined to specific SC niches. This sensitivity to environmental modifications may endow these PNS SCs with an improved capacity to respond and adapt to injury and, therefore, contribute to the more efficient regenerative properties of the PNS compared with the CNS.

Conclusion
Although several studies have proposed the presence of DRG SCs, none considered them a distinct SGC population. Their existence, activation, and involvement in glial/neuronal renewal under pathophysiological conditions open a new field of research, especially in neuro-pathic studies. The “activated” SGC seems to be an entity shared by various peripheral neuropathies caused by chemotherapy (Warwick and Hanani, 2013), injury (Donegan et al., 2013), viruses, and heredity (Sghirlanzoni et al., 2005) and could contribute to the genesis and maintenance of other SGCs (Hanani, 2012) and renewal of neurons. New transgenic models combined with molecular/transcriptomics analysis should provide the ability to further study the contribution and intracellular regulation of DRG SCs under physiological and pathophysiological conditions. These DRG SCs could be pharmacological targets to treat peripheral neuropathies and possibly CNS disorders providing a source of PNS glia known to invade the CNS in response to demyelinating or traumatic conditions (Garcia-Diaz et al., 2019).

EXPERIMENTAL PROCEDURES

Animals
Nestin-CFPnuc (Encinas et al., 2006), Nestin-GFPcyt (Mignone et al., 2004), Nestin Cre ERT2 (Imayoshi et al., 2006), and Rosa26-YFP lines were bred and housed under standard conditions, and experiments were performed according to European Community guidelines (see supplemental information).

EdU pulse-chase assay
EdU (catalog number C35002, Invitrogen) dissolved in PBS (8.33 mg/mL) was administered by intraperitoneal injection at a dose of 75 mg/kg every 12 h during 8 days in 3-month-old wild-type or transgenic (Nestin-CFPnuc or Nestin-GFPcyt) mice (n = 12). Half of the injected mice were sacrificed 12 h after the last injection, and the others were killed after 1 month of chase. DRGs, sciatic nerves, and spinal cords were prepared as described below.

Cre induction
Three to four-month-old double-transgenic mice, Nestin-CreERT2 × Rosa-YFP, received a dose of tamoxifen dissolved in corn oil (30 mg/mL, Sigma, catalog number T5648) by oral administration (210 mg/kg) during 5 days.

Sciatic nerve axotomy
After a resting period of 10 days, tamoxifen-induced animals were anesthetized. In half of the group, the right sciatic nerve was exposed mid-thigh and cut. Injured or non-injured induced mice (n = 3–5 per time point and per group) were sacrificed at different dpi. For detailed information, see supplemental experimental procedures.

Tissue processing for fluorescence immunohistochemistry
DRGs, sciatic nerves, and spinal cords from 4-month-old Nestin-CFPnuc and Nestin-GFPcyt mice (n = 3–4 per line and experimental group) were sacrificed by overanesthesia (Ketamin-Rompan) and perfused trans-cardially with a solution of 4% paraformaldehyde (PFA) in phosphate-buffered saline (PBS) 1x before tissue collection and freezing. For detailed information, see supplemental experimental procedures.

Tissue processing for immunogold electron microscopy
Four-month-old Nestin-CFP (n = 5) mice were sacrificed by overanesthesia (Ketamin-Rompan) and trans-cardially perfused with 4% PFA in PBS (0.1 M). DRGs and spinal cords were post fixed 1 h in 4% PFA and 0.3% glutaraldehyde (Sigma-Aldrich) and processed for electron microscopy (EM). After sectioning and immunogold staining, ultra-thin sections were examined by transmission electron microscope. For detailed information, see supplemental experimental procedures.
Adult DRG and spinal cord cultures

Isolation

Transgenic 3- to 4-month-old mice were euthanized to collect spinal cords and DRGs, and processed for dissociation. Cells from the two tissues were purified by FACS based on their endogenous expression of CFPnuc and GFPcyt (n = 6 independent replicates/transgenic line). For expansion, cells sorted by FACS were seeded at semi-clonal concentration and cultured in suspension as spheres in vitro. For differentiation, 15-20 day primary spheres were induced to adhere and cultured without growth factor for 7–10 days of culture. Cells were fixed and immunolabeled as described above. For detailed information, see supplemental experimental procedures.

Fluorescence imaging, quantification, and statistics

Imaging was carried out with an Apotome system (Carl Zeiss) equipped with the Zen software. Images were processed using Adobe Photoshop (Adobe Systems). All quantifications were conducted using the Fiji software and statistics with GraphPad Prism 6 software. In vitro data were deduced from 3–4 experiments performed in duplicates, as detailed in the supplemental information.

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.stemcr.2022.10.002.

AUTHOR CONTRIBUTIONS

Conception and design, collection and assembly of data, data analysis and interpretation, and manuscript writing, M.M.; conception and design, collection and assembly of data, and data analysis and interpretation, M.V.; collection and assembly of electron microscopy data and data analysis and interpretation, C.B.; collection and assembly of in vitro data and data analysis, C.D.; collection and assembly of immunohistology data, J.C.; data assembly and analysis and manuscript editing and writing, B.G.-D.; conception and design, financial support, administrative support, data analysis and interpretation, manuscript writing, and final approval of the manuscript, A.B.-V.E.

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CONFLICT OF INTERESTS

The authors declare no competing interests.

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