Adult neural stem cell behavior underlying constitutive and restorative neurogenesis in zebrafish

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**ABSTRACT**

Adult Neural Stem Cells (aNSCs) generate new neurons that integrate into the pre-existing networks in specific locations of the Vertebrate brain. Moreover, aNSCs contribute with new neurons to brain regeneration in some non-mammalian Vertebrates. The similarities and the differences in the cellular and molecular processes governing neurogenesis in the intact and regenerating brain are still to be assessed. Toward this end, we recently established a protocol for non-invasive imaging of aNSC behavior in their niche \textit{in vivo} in the adult intact and regenerating zebrafish telencephalon. We observed different modes of aNSC division in the intact brain and a novel mode of neurogenesis by direct conversion, which contributes to stem cell depletion with age. After injury, the generation of neurons is increased both by the activation of additional aNSCs and a shift in the division mode of aNSCs, thereby contributing to the successful neuronal regeneration. The cellular behavior we observed opens new questions regarding long-term aNSC maintenance in homeostasis and in regeneration. In this commentary we discuss our data and new questions arising in the context of aNSC behavior, not only in zebrafish but also in other species, including mammals.

In the adult vertebrate brain, new neurons are produced\textsuperscript{1,4,23,39,59} and turned over\textsuperscript{38,44} throughout the animal’s life, with an age-dependent decline\textsuperscript{14,15,21,33,56}. These neurons differentiate from neural stem cells (NSCs) that reside in specialized niches within the brain\textsuperscript{11,17,29,54}. The existence of adult/neonatal NSCs, also in the human brain\textsuperscript{22,23,59}, raised hope for their use in regenerative therapies for neurodegenerative diseases or brain injuries. Indeed, adult NSCs (aNSCs) provide new neurons engaged in the repair of the injured brain in regeneration-competent species, such as zebrafish\textsuperscript{8,10,11,36,37,58,65}. Although the aNSCs contribute with new, mature neurons in some regeneration-competent species, the first attempts to utilize the endogenous aNSCs for repair in the mammalian brain largely failed\textsuperscript{7,58,61,62}, probably due to the lack of understanding of basic aNSC biology. Moreover, it also remains unclear to which extent the regeneration of the injured brain requires changes in the behavior of aNSCs compared to the intact brain in order to complete the regeneration process. For example, repair of the injured cerebral cortex would require the generation of pyramidal neurons, a cell type never produced by the aNSCs in the intact brain\textsuperscript{46}. Therefore, it is crucial to compare the cellular behavior at the single stem cell level in the intact and injured brain of regeneration-competent and regeneration-incompetent species. The first approaches to understand the cellular behavior of single aNSCs in the neurogenic zones of the intact mammalian brain based on clonal analysis\textsuperscript{13,15} revealed the fast consumption of a single aNSC that produces a heterogeneous neuronal output. However, different cellular processes such as cell death, selective proliferation and terminal differentiation could yield into the described features of adult mammalian neurogenesis. The methods used in these studies could not provide complete information on the continuous behavior/dynamics of aNSCs in their intact niche, highlighting the need for complementary methods that allow repeated observation of the same aNSCs in their biological environment.

In the mammalian brain the NSC niches are rare (Sub-ependymal zone (SEZ), dentate gyrus (DG) and...
hypothalamus) and are located several hundreds of μm away from the brain surface rendering them inaccessible for direct observations. In contrast, the neurogenic niches in the adult zebrafish are widespread throughout the whole brain and accessible for live imaging, particularly in the dorsal telencephalon, due to its location at the outer surface of the brain (Fig. 1a). The privileged location of aNSCs in the dorsal zebrafish telencephalon, and the exceptional regenerative potential of this brain region, makes the zebrafish pallium an attractive system to pursue in vivo imaging experiments.

Live in vivo imaging allows for the integrative view on the changes in behavior of the single aNSCs and their progeny in the regenerating and intact zebrafish brain. This commentary will discuss the behavior of the aNSCs in the zebrafish telencephalon in both conditions and its possible implications for the processes of aging and regeneration.

Output of the adult neural stem cells in the intact and injured brain

The neurogenic niche in the adult zebrafish pallium, accessible for live imaging, contains radial glia-like aNSCs (Fig. 1b) with their cell bodies lining the ventricular wall. Radial processes of aNSCs span the brain parenchyma to contact the basement

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**Figure 1.** Neurogenic niches in the adult zebrafish telencephalon and behavior of aNSCs in the pallium. (a) Representation of a coronal section through the adult zebrafish telencephalon illustrating the ventricular zone (green), containing aNSCs and progenitors, and the parenchyma (brown), mostly composed of neurons. The two hemispheres are linked by a dorsal ependymal lining (DEL) that closes the ventricle (v). (b) and (c) Scheme of the dorsal (b) and ventral (c) ventricular zones, composed of different progenitor types: RG-like aNSCs, non-glial intermediate progenitors and neuro-epithelial-like cells. (d) and (e) Modes of cell division and neurogenesis in the intact (d) and injured (e) zebrafish telencephalon, assessed by live imaging. In the intact brain (d) the newborn neurons are deposited immediately adjacent to the progenitor cells. After injury (e) there is an increase in proliferating aNSCs at the VZ, a change in their mode of cell division and the migration of progeny to the injured parenchyma, where the newborn neurons contribute to tissue regeneration. The dashed circle marks the region where the lesion was. Abbreviations: DEL-dorsal ependymal lining; NE-neurepithelial; RG-radial glia; VZ-ventricular zone.
membrane. The morphology and the antigen profile of aNSCs in the zebrafish pallium are reminiscent of radial glial (RG) cells in the developing mouse telencephalon. The aNSCs in the zebrafish pallium do not only share the morphological and immunohistochemical characteristics with the mammalian RG cells in the developing brain, but also have the capacity to generate new neurons. However, in contrast to the new neurons produced in the developing mammalian cerebral cortex, the new neurons produced by the pallial aNSCs in the intact brain of zebrafish do not migrate away from the stem cell zone and are instead deposited directly below the progenitor zone (Fig. 1d). These new neurons are intermingled with fast dividing progenitors (Fig. 1b) that do not have stem cell characteristics (intermediate progenitors (IPs) or non-glial progenitors). Traumatic brain injury induces a specific program in the aNSCs and intermediate progenitors resulting in the production of new neurons necessary for regeneration. In contrast to the intact brain, these newborn neurons migrate larger distances to repopulate the damaged brain areas (Fig. 1e). Importantly, the small stab wound injury induces the restorative neurogenesis without an impact on the ongoing neurogenesis present in the intact brain. This indeed raises the question of the origin of the new neurons engaged in the repair process. Moreover, it remains unclear to which extent the cellular processes underlying restorative neurogenesis recapitulate those sustaining the normal generation of adult-born neurons.

To address these questions, we repetitively imaged aNSCs in the Tg(gfap::GFP)mir2001 transgenic fish line that expresses GFP in all aNSCs. To reliably re-identify aNSCs in different imaging sessions, we sparsely labeled a small number of them by electroporation of a reporter (TdTomatomem) encoding plasmid. As the ubiquitous cytomegalovirus (CMV) promoter drives the expression of the reporter, we could follow not only aNSCs, but also their progeny that lost the radial morphology and glial marker expression. Our results confirmed previous observations that in the intact brain aNSCs are mostly quiescent and only a small proportion of aNSCs divide or change their identity to generate progeny (aNSCs activation) at any discrete time point. Both live imaging and clonal analysis suggest that aNSCs predominantly generate neurons upon their activation. In addition, we could observe the generation of gfp::GFP-positive glial cells indistinguishable from the mother cell indicative of self-renewal. Indeed, all aNSC divisions in the intact brain generated at least one cell with radial morphology and also expressing gfap::GFP. Whether some of these glial cells are already a differentiated cell type (possibly homologous to the ependymal cells in the mammalian brain) or a quiescent stem cell is not clear yet. However, virtually all GFAP-positive glial cells in the zebrafish pallium can be recruited into the cell cycle upon Notch inhibition, strongly suggesting that at least some of the newly generated gfp::GFP-positive cells with radial morphology are quiescent aNSCs. Longer imaging periods would be required to tackle this question in a comprehensive manner.

The generation of oligodendrocytes from aNSCs in the zebrafish pallium has not been directly assessed. However, aNSC-derived cells mostly express either GFAP or neuronal lineage markers (Sox2 and HuCD) (unpublished data), accentuating the idea that only neurons and aNSCs/ependyma are generated from NSCs in this brain region. This is indeed very similar to the adult mouse DG, where aNSCs do not generate oligodendrocytes. Oligodendrocytes are generated in this region exclusively by neural/glial antigen2 (NG2)-positive progenitors. In the adult mouse SEZ oligodendrocytes are also generated, but the neuronal and oligodendroglial lineages are separated in two independent populations of progenitors.

Although the predominantly neuronal output appears to be a hallmark of aNSCs in the Vertebrate brain, the cohort size produced by a single aNSC differs greatly between species and analyzed areas. Despite the difference in clone size, the output of a single aNSC seems to be defined by the number of divisions of intermediate progenitors. In the zebrafish pallium, these progenitors divide once or twice before terminally differentiating into neurons. Interestingly, this low degree of lineage amplification in the zebrafish pallium is reminiscent of the behavior of progenitors in the adult mouse DG, supporting the suggested homology between the two regions. Also in the developing rodent dorsal telencephalon, basal/intermediate progenitors divide few times before generating neurons. However, this situation contrasts with the mouse ventral telencephalon, in which multiple rounds of division of progenitors greatly amplify the neuronal output. Similarly, in the SEZ niche a high degree of
amplification occurs at the transit amplifying progenitor (3–4 divisions) and neuroblast (1–2 divisions) level.\textsuperscript{15,50} Moreover, aNSCs also divide asymmetrically several times each 2–3 weeks to produce a larger neuronal output containing several cohorts (Fig. 2b).\textsuperscript{15} Unlike in the pallium, the neurogenic niche in the zebrafish subpallium appears more similar to the mouse SEZ (Fig. 1c), as there are proportionally more IPs\textsuperscript{34,42} and some of these also migrate to the olfactory bulb.\textsuperscript{34} Interestingly, in the zebrafish subpallium the aNSCs exhibit low levels or no expression of glial markers and, because they express nestin and the tight junction component zona occludens 1 (ZO-1),\textsuperscript{24} they resemble neuroepithelial cells (Fig. 1c). The mixture of radial and tangential migration of the NSC progeny in this region makes a clonal analysis challenging. Thus, the behavior of subpallial aNSC in the zebrafish at the single cell level remains to be assessed.

The broad spectrum of the single aNSC behaviors leading to the production of differently sized neuronal cohorts prompted us to search also for the mechanisms enlarging the neuronal output during regeneration, that allow the replacement of the lost neurons without an obvious impact on the ongoing
Several cellular mechanisms such as increased aNSC proliferation, proliferation of the non-glial progenitors, decreased cell death etc. could account for this net increase in neuronal production. Indeed, both aNSCs and non-glial progenitors increase their proliferation after brain injury. However, we could not observe multiple divisions of a single aNSC using live imaging in vivo, but rather increased recruitment of quiescent aNSCs. Interestingly, we observed that some of the aNSCs dividing after injury did not self-renew, but rather exhausted themselves by a symmetric division generating two non-glial progenitors. This mode of division, not present in the intact brain, might constitute an advantage after injury, since it generates a larger neuronal output from a single aNSC compared to an asymmetric division. On the other hand, it also leads to the depletion of stem cells. Curiously, enhanced aNSC depletion has also been observed in the DG of a mouse model of neuronal hyperactivity, suggesting some conservation of aNSC reaction to challenge in different Vertebrate species. This NSC exhaustion phenomenon would predict a decreased capacity of the zebrafish brain to regenerate damages induced by the repetition of the insult. Alternatively, aNSCs in the zebrafish might be heterogeneous and could contain more primitive, undifferentiated cells with the capacity to repopulate aNSCs depleted after injury and enable proper regeneration even after multiple injuries. Indeed, in the zebrafish caudal fin amputation paradigm, progenitor cells completely reconstitute the entire damaged tissue without losing efficiency even after repeated amputations. Therefore, it would be important to address if aNSCs have the capacity to repopulate the neurogenic zone and allow tissue restoration even after multiple insults or if the different organs have different regeneration capacity depending on the characteristics of the somatic stem cells residing within the given organ.

Direct conversion

An important novel feature of adult neurogenesis in the zebrafish telencephalon uncovered by our in vivo imaging was the direct conversion of a considerable proportion of aNSCs (50% of all aNSCs generating neurons) into a neuron without any cell division. Interestingly, this type of neurogenesis is also described in developing brains. In fact, single progenitors labeled at the neural rod stage directly convert into neurons without cell division in the developing zebrafish hindbrain. Also in Xenopus laevis, time-lapse imaging demonstrated that the majority of RG cells in the developing optic tectum directly differentiate into neurons. As direct conversion had never been observed in the adult brain, this could either mean that aNSCs in the zebrafish telencephalon possess this unique capacity, or that direct conversion has not yet been detected in the adult brain of other species due to the lack of suitable technical approaches such as live imaging.

At present, we cannot elucidate if the population of aNSCs that directly convert to neurons is a specific population or if these cells also have the capacity to divide. In fact, both in the zebrafish and in the mouse embryo, dividing RG cells generate neurons directly without going through an intermediate progenitor state (Fig. 2a, direct neurogenesis). In our study however, we followed aNSCs for at least 2 weeks before they directly converted into neurons, suggesting that these cells either have an extremely long cell cycle or do not need to divide at all to generate neurons.

Notably, after injury aNSCs divide more and change their division mode to produce large cohorts of neuronal progeny needed in these hostile conditions. In contrast, direct conversion would allow slow, but constant addition of new neurons to the slowly growing adult. It is still to be addressed if the aNSCs undergoing direct conversion in the intact brain would become activated after injury and divide in a symmetric neurogenic manner (a mode of aNSCs division that we observe only in the injured brain) to produce a larger neuronal output and deplete themselves after injury. Alternatively, direct conversion might be actively blocked but these aNSCs would be kept at the VZ for the slow neuronal production after the brain is regenerated.

Stem cell depletion—a common feature of the vertebrate neurogenesis

In the adult zebrafish pallium, new neurons are added either by asymmetric division of aNSCs, thus maintaining the stem cell pool, or by direct conversion depleting the stem cell pool. Our live imaging showed that the proportion of aNSCs that directly convert to neurons and deplete themselves (17%) is considerably higher than the proportion of aNSCs dividing
symmetrically to amplify the stem cell population (1%) in the intact brain. This finding implies a gradual consumption of aNSCs with age. Indeed the number of proliferating aNSCs was decreasing in 6 and 10 months old animals compared to 3 months old animals, which correlates with the previously described decrease in neurogenesis in the aged zebrafish pallium. These data therefore would support the hypothesis that aNSC depletion is the cellular basis for the age-dependent decline of neurogenesis in zebrafish. However, Edelmann et al observed that the total number of glap::GFP-positive glia (regardless of their proliferative status) does not decrease with aging. The cellular mechanisms that maintain the ependymoglia in the adult brain remain to be investigated, but one possible explanation would be that the modes of stem cell division/behavior could be changed toward the gliogenic/ependymoglial fate in aging animals. In our study we used 2–3 months-old animals and followed single cells for one month but it is possible that in older animals there is a prevalence of symmetric aNSCs divisions that would replenish the pool of ependymoglia. More in vivo imaging of single aNSCs at these later stages would be needed to clarify these issues. Importantly, in the mouse neurogenic niches, SEZ and dentate gyrus, there is a limited number of self-renewing divisions of aNSCs followed by terminal differentiation into the neuronal or astrocytic lineage. Therefore the exhaustion of aNSCs in Vertebrates is a common trait, possibly including the glial fate as the final differentiation step.

Future directions

In summary, the in vivo imaging we established allowed for the first time the visualization of aNSCs in their natural niche in a vertebrate model, not only in physiological conditions but also after brain damage. Consequently, several features of the behavior of individual aNSCs were revealed at the single cell level, at an extent that could not be assessed by previous population-based studies. The knowledge acquired with the observation of NSC behavior in the zebrafish brain may serve as a basis to conduct studies on modulating NSC activity in the diseased brain. For the application of these findings in mammalian disease models it will be important to compare the NSC behavior in zebrafish and mammals.

Disclosure of potential conflicts of interest

No potential conflicts of interest were disclosed.

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