Telencephalon enlargement by the convergent evolution of expanded subventricular zones

Some mammals and birds independently evolved an enlarged telencephalon. They appear to have done so, at least in part, by developing a thick telencephalic subventricular zone (SVZ). We suggest that this correlation between telencephalic enlargement and SVZ expansion is due to a mechanical constraint acting on the proliferative ventricular zone (VZ). Essentially, we argue that rapid proliferation in the VZ after post-mitotic cells in the overlying mantle zone have begun to form limits the VZ's tangential expandability and forces some proliferating cells to emigrate from the VZ and expand the pool of proliferating cells that comprise the SVZ.

**Keywords:** development; neocortex; birds

An SVZ has also been described in the telencephalon of birds (Striedter & Keefer 2000). This avian SVZ is thicker in embryonic parrots than in age-matched quail (Striedter & Charvet 2008), an observation that correlates with a difference in adult telencephalon volume, because adult parrots have a proportionately larger telencephalon than adult quail or other fowl (Boire & Baron 1994). The telencephalic expansion in adult parrots involves the striatum as well as several pallial regions. This correlates with our finding that the parrot's SVZ is thickened in both the striatum and the pallium (Striedter & Charvet 2008). Collectively, these observations suggest that the SVZ expanded convergently in parrots and mammals.

Instances of convergent evolution often indicate the existence of underlying causal principles that can account for the evolved similarities. Of course, some convergences may simply be due to chance. However, the argument for chance becomes less likely if the convergent similarities show up in diverse lineages. It is useful, therefore, to ask whether the SVZ is expanded also in other species that have enlarged their telencephalon.

An obvious taxon to examine in this context are songbirds, as they rival parrots in their degree of telencephalic enlargement (Boire & Baron 1994). Previous studies described only a thin SVZ in the subpallium and lateral pallium of a juvenile songbird, the zebra finch, 30 days after hatching (Dewulf & Bottjer 2002). However, we find that the telencephalon of newly hatched zebra finches contains a large SVZ that is similar in thickness and extent to that of parrots (figure 1). Therefore, songbirds also appear to have enlarged their telencephalon, at least in part, by expanding their SVZ. Whether the SVZ expanded independently in songbirds and parrots remains unclear because songbirds may be closely related to parrots (Hackett et al. 2008). However, the fact that songbird and parrot brains differ in numerous major respects (Stingelin 1958) strongly suggests that these two taxonomic groups enlarged their telencephalons independently of one another.

To discover causal principles underlying the convergent evolution of expanded SVZs in mammals and birds, we begin with Smart’s (1972) insight that the organization of the VZ, with its interkinetic nuclear migration and ventricular mitoses, imposes significant constraints (figure 2). One constraint is that rapid proliferation within the VZ requires, and generally co-occurs with, tangential VZ expansion. This tangential expansion accounts for the ballooning of the telencephalic vesicle early in embryogenesis. However, if tangential expansion cannot occur, then the VZ must slow its proliferation rate or allow some mitoses to occur away from the ventricle in an SVZ (Smart 1972). Thus, the SVZ allows rapid proliferation to occur in regions of the telencephalon that are limited in their degree of tangential expansion. The most obvious such region is the ganglionic eminence, which bulges into the telencephalic ventricle, gives rise to parts of the basal ganglia and contains a thick SVZ (Smart & Sturrock 1979).

Thickening the telencephalic walls by means of an expanded SVZ is one effective strategy for packing many cells into a small space. An alternative strategy is to expand the telencephalon tangentially, causing...
the telencephalic vesicle to balloon more, and then to bend the tissue into complex folds, as happens with the cerebral cortex in large-brained mammals (Prothero & Sundsten 1984). Thus, one can view the expanded SVZ in parrots and songbirds as convergent uses of the same radial expansion strategy for enlarging the telencephalon. Primates, by contrast, employ a hybrid strategy that relies on tangential expansion to enlarge the cortical surface and on an expanded SVZ to thicken the neocortex.

Expanding an SVZ may be a good strategy, but what is its proximate cause? All SVZs initially develop as a thin layer just superficial to the VZ (Kriegstein et al. 2006; Charvet & Striedter 2008), suggesting that the first SVZ cells are descendents of VZ cells that emigrated from the VZ (Fish et al. 2008). The subsequent expansion of the SVZ is probably due to continued emigration from the VZ and to the proliferation of SVZ cells in situ. SVZ cells differ from VZ cells not only in their migratory behaviour but also in their pattern of gene expression. For example, they express \( Tbr-2 \), which is not expressed in VZ cells (Englund et al. 2005). Unfortunately, these molecular differences do not explain why some cells leave the VZ to become SVZ cells, while others remain behind. Perhaps the VZ contains a molecularly distinct subpopulation of cells that is fated to form the SVZ. Some evidence for such molecular heterogeneity has been described (Englund et al. 2005; Pinto et al. 2008).

An alternative possibility is that some VZ cells are mechanically forced out of the VZ. Once they have lost their attachment to the ventricle, these budding SVZ cells may adopt a unique pattern of gene expression and start behaving differently from VZ cells. Analogous mechanical effects on gene expression have been described in other proliferating cells (Huang & Ingber 1999). We hypothesize that the force pushing the cells out of the VZ results when the VZ is proliferating rapidly but its tangential expansion is constrained. This portion of our argument is based on Smart’s (1972) insights. We further suggest that what constrains the VZ is the presence of post-mitotic cells just superficial to the emerging SVZ in the young mantle zone (MZ; figure 2). These post-mitotic cells tend to have long tangential processes that may limit the MZ’s tangential expandability, which then constrains the VZ, to which the MZ is linked by radial glia.

This set of interlinked hypotheses was originally derived from the observation that the SVZ in both mammals and birds consistently begins to form just after the MZ appears. More direct support comes from a recent study of transgenic mice that express a constitutively active form of \( \beta \)-catenin in the neocortex (Wrobel et al. 2007). Because activated \( \beta \)-catenin promotes cell cycle re-entry over cell cycle exit (Chenn & Walsh 2002), these transgenic animals exhibit a significant delay in neurogenesis onset, which means that MZ cells form abnormally late. As our hypothesis predicts, in these transgenic mice, SVZ formation is delayed. The study’s authors did not discuss the possibility of a link between delaying neurogenesis and delaying SVZ formation, but their findings are consistent with our proposal.

The observation that turtles thicken their lateral telencephalic wall but exhibit, at best, a rudimentary SVZ (Molnar et al. 2005; Martínez-Cerdeño et al. 2006) implies that telencephalic walls can be thickened without a proper SVZ. Such SVZ-independent thickening might occur when VZ cells proliferate faster than the MZ can be stretched, but slowly enough to prevent proliferating cells from losing their attachment to the ventricular surface. Slow proliferation might also explain the somewhat surprising absence of an SVZ in opossums. Another complication for our proposal is that brains do expand tangentially long after the SVZ has formed. This late tangential expansion differs from the earlier expansion, however, in being part of a more uniform expansion due to the growth of post-mitotic neurons and the late addition of glial cells.

Figure 1. Sections through the telencephalon of (a, b) parakeets and (c–e) zebra finches labelled with antibodies against (a–c, e) proliferating cells (anti-PCNA) or (d) mitotic cells (anti-phosphorylated histone H3). The thick SVZ in the striatum (St) is marked with an asterisk. Mitotic cells in the hyperpallium, which is probably homologous to at least part of the mammalian neocortex, are indicated with arrowheads. H, hyperpallium; M, mesopallium; N, nidopallium; Hip, hippocampus; Sep, septum. Scale bar, 100 \( \mu \)m.
The most potent means of enlarging a brain region is to prolong its period of precursor proliferation (Smart 1972; Finlay & Darlington 1995). The downside of prolonging proliferation is that it delays maturation, as mature neurons do not divide. One way to minimize this problem is to delay neurogenesis offset without delaying neurogenesis onset (Striedter & Charvet 2008), thereby allowing at least some neurons to differentiate early. According to our hypothesis, this strategy creates a thick SVZ and a thick adult structure, which is what we observe in the telencephalon of parrots and songbirds. Of course, a delay in telencephalic neurogenesis offset causes most telencephalic neurons to mature relatively late. This is not a problem for parrots or songbirds, as these altricial birds feed their babies. However, precocial species might well adopt a different strategy for enlarging their telencephalon. Testing of this hypothesis is ongoing.

We thank Edwin Monuki for his valuable feedback and the NSF for grant support (IOS-0744332).

Georg F. Striedter* and Christine J. Charvet
Department of Neurobiology and Behavior, University of California Irvine, Irvine, CA 92697–4550, USA *(georg.striedter@gmail.com)


Smart, I. 1972 Proliferative characteristics of the ependymal layer during the early development of the spinal cord in the mouse. J. Anat. 111, 365.


