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Revisiting the relationship between regenerative ability and aging

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See research article: http://www.biomedcentral.com/1741-7007/10/103

Abstract
Contrary to the longstanding view that newts (Notophthalmus viridescens), but not axolots (Ambystoma mexicanum), can regenerate a lens, a recent report in BMC Biology by Panagiotis Tsonis and colleagues shows axolots indeed possess this ability during early larval stages. In contrast, they show that zebrafish never possesses this ability, even as embryos. This underscores the importance of comparing regenerative ability across species and reinforces the need to consider organ regeneration in the context of evolution, development, and aging.

New insights in regenerative biology will continue to arise from studies of animal models that present a diversity of regenerative responses. To better leverage these models, it will be important to consider regenerative ability within the context of evolution, life history, physiology, and development [1]. Here we highlight a long-standing problem, the loss of regenerative ability as an organism ages. Considering this fundamental relationship between aging and regeneration in naturally regenerating systems may help translate new discoveries into effective applications in regenerative medicine.

While humans possess varied and diverse mechanisms for physiological regeneration - regeneration to maintain and renew organ functions throughout life - this renewal declines with age; furthermore, we possess a very limited ability for the regeneration of tissues following injury. Enviously, we watch flawless regeneration of limbs, lens, retina, spinal cord, brain, heart, and neuro-sensory cells, seemingly throughout life, by distant vertebrate relatives. There are, however, limits to the ability of any organism to regenerate, and understanding these limits may be the key to understanding why humans cannot regenerate whole organs. Although examples of tissue regeneration are rare among birds and mammals, tissue regeneration occurs in amphioxus, the most basal living chordate [2]. This suggests that humans likely share mechanisms that are used by salamanders and other vertebrates to regenerate organs. Salamanders provide a good stepping-off point for considering factors that limit regenerative ability because they present a basal tetrapod condition that most closely approximates ancestral lineages leading to amniotes.

Regenerative potential is generally higher during early life stages
To understand why and how regenerative ability varies across ontogeny and phylogeny requires a comparative approach that evaluates regeneration of homologous structures throughout development, preferably using species that diverged from a common ancestor. Unfortunately, few rigorous studies have examined regeneration throughout development. Still, ontogenetic changes in regenerative potential have been described for a few species and several lines of evidence support the idea that embryos, larvae, and juveniles of many species have greater potential for tissue repair and regeneration than adults (Figure 1). Frogs and mice, for example, are capable of regenerating skin during early life stages but lose this ability later in development, while axolotls retain the capacity for skin regeneration even after metamorphosis ([3] and references therein). Similarly, the ability of zebrafish to regenerate pectoral fins and frogs to regenerate limbs is reduced in older animals, while heart regeneration appears unaffected even in very old fish ([1,4] and references therein). Adding to these studies, a report by Suetsugu-Maki et al. [5] now shows that axolots can regenerate a lens during early life stages, while most vertebrates cannot regenerate lenses at all. And yet, newts are capable of lens regeneration throughout life and if the progenitor cells that restore lenses in...
non-regenerating species, including humans, are cultured in vitro, cells with lens phenotypes differentiate [6]. Thus, while the potential for lens regeneration is apparently a shared derived trait of vertebrates, that potential is restricted during development in most species. Together, these examples suggest that even in some salamanders with boundless regenerative abilities, constraints imposed by aging, either at the physiological or cellular level, work to limit regenerative capacity in organ- and species-specific manners.

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**Figure 1. Reparative regeneration as a function of developmental stage among vertebrate models of regeneration.** Panels depict how the regenerative ability of homologous structures varies across ontogeny for the primary vertebrate models of regeneration. Regenerative ability is presented on a 0 to 5 scale; five represents perfect regeneration and zero no regeneration. Major ontogenetic stages are represented as embryo, larva, juvenile and adult with metamorphosis, puberty or both indicated for each species. (a) Zebrafish exhibit lifelong regenerative capacity of spinal cord, brain, and heart. They cannot regenerate a lens at any time in development and while pectoral fins (homologous to tetrapod limbs) regenerate in juveniles, their regenerative capacity is reduced following puberty, with females retaining a higher capacity for complete regeneration. (b) Salamanders and newts are the archetypical tetrapod regenerator, retaining near perfect regeneration of most organs and appendages well into adulthood (although almost no studies have tested these abilities in old animals). They do, however, experience a decline in limb regeneration following metamorphosis, which usually manifests as patterning defects and loss of limb elements. Axolotls can only regenerate lenses as early stage larvae. (c) Although most frogs exhibit some degree of regeneration as larvae, with the exception of limited digit and very restricted limb regeneration in some species, they do not exhibit regenerative ability as adults. (d) Mammals exhibit some regenerative capacity as embryos but lose almost all of this ability before or shortly after birth for the structures listed.
What limits regenerative ability during aging?
Understanding how aging changes cells, tissues, and physiological systems is key to identifying mechanisms that limit regenerative ability. During both development and regeneration, relatively undifferentiated cells become specified to form organs that then undergo tremendous growth, but the overall process differs between the two. Regeneration is activated in response to injury, depends upon tissue-specific progenitor cells, and occurs under physiological conditions and within an extracellular environment that differs from the embryonic state. Embryonic cells have great potential for cellular reprogramming, and cellular reprogramming through epigenetic modifications and changes in transcription are associated with regenerative responses [7]. During development, cells exhibit changes in transcription that limit signaling pathways associated with cellular plasticity. For example, cells differentiate and lose the ability to enter the cell cycle, both of which must be reversed for limb regeneration to occur in salamanders. The maintenance of plastic cellular states and cell-cycle re-entry are likely associated with the actions of tumor suppressor proteins like retinoblastoma protein (RB), the levels and activation states of which are known to vary during regeneration in salamanders and across developmental stages in mammals [8,9]. Taken together, studies suggest that the abundances of key regulatory molecules are permissive for cell cycle re-entry from a quiescent state in young life stages, but restrictive as an organism ages, and this limits regenerative ability. Future studies that quantify levels of such molecules in young and old salamanders may shed new light on progenitor cell activation, regenerative ability, and potentially, diseases of aging such as cancer.

How do systemic factors affect regenerative ability?
Early, heightened cellular plasticity in response to injury reflects not only local but also systemic factors that are difficult to disentangle without ontogenetic perspective. The importance of systemic factors is dramatically shown in studies of parabiotic mice that differ in age but share the same circulatory system. Serum from young animals in studies of parabiotic mice that differ in age but share the same circulatory system. Instead, regeneration also depends upon serum factors that have broad access to tissues and regeneration, relatively undifferentiated cells become specified to form organs that then undergo tremendous growth, but the overall process differs between the two. Regeneration is activated in response to injury, depends upon tissue-specific progenitor cells, and occurs under physiological conditions and within an extracellular environment that differs from the embryonic state. Embryonic cells have great potential for cellular reprogramming, and cellular reprogramming through epigenetic modifications and changes in transcription are associated with regenerative responses [7]. During development, cells exhibit changes in transcription that limit signaling pathways associated with cellular plasticity. For example, cells differentiate and lose the ability to enter the cell cycle, both of which must be reversed for limb regeneration to occur in salamanders. The maintenance of plastic cellular states and cell-cycle re-entry are likely associated with the actions of tumor suppressor proteins like retinoblastoma protein (RB), the levels and activation states of which are known to vary during regeneration in salamanders and across developmental stages in mammals [8,9]. Taken together, studies suggest that the abundances of key regulatory molecules are permissive for cell cycle re-entry from a quiescent state in young life stages, but restrictive as an organism ages, and this limits regenerative ability. Future studies that quantify levels of such molecules in young and old salamanders may shed new light on progenitor cell activation, regenerative ability, and potentially, diseases of aging such as cancer.

Many animals undergo post-embryonic growth and developmental phases that commence in response to circulating hormones that are released at relatively specific times during ontogeny. In the case of amphibian metamorphosis, thyroid hormone (TH) reprograms juvenile cells and activates adult progenitor cells, and this brings about the conversion of tadpole aquatic larvae into more terrestrial adults. Interestingly, while newts always undergo metamorphosis, axolotls rarely do unless treated exogenously with TH [12]. Future studies that use hormones to induce metamorphosis at different times during ontogeny may be able to disentangle the effects of aging from intrinsic regenerative ability, as it has already been shown that prolonging the larval state enhances regenerative ability compared to same-aged animals that have undergone metamorphosis [13].

Regeneration insight from comparative approaches
In coming years, we envision a new, golden age in regenerative biology. Animals present a diversity of regenerative responses that vary across organs, developmental stages, and phylogeny. Increasingly, advances in genetic and genomic technologies will make it possible to compare regenerative responses within and among animal models to identify factors that cause regenerative ability to change with aging.

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