The cell biology of regeneration

Ryan S. King and Phillip A. Newmark

Howard Hughes Medical Institute, Department of Cell and Developmental Biology, University of Illinois at Urbana-Champaign, Urbana, IL 61801

Regeneration of complex structures after injury requires dramatic changes in cellular behavior. Regenerating tissues initiate a program that includes diverse processes such as wound healing, cell death, dedifferentiation, and stem (or progenitor) cell proliferation; furthermore, newly regenerated tissues must integrate polarity and positional identity cues with preexisting body structures. Gene knockdown approaches and transgenesis-based lineage and functional analyses have been instrumental in deciphering various aspects of regenerative processes in diverse animal models for studying regeneration.

Regeneration: Sometimes truth is stranger than (science) fiction

Fans of Doctor Who, the quintessential British science fiction television series, have been able to enjoy the adventures of its main character (the Doctor) for over three decades. As a Time Lord from the planet Gallifrey, the Doctor can regenerate his body completely as he nears death. The Doctor’s regenerative abilities, however, are not without flaw: after he regenerates, he takes on different features and characteristics. This imperfect process enables the series to continue with a new actor portraying the Doctor upon the departure of his predecessor. The Doctor’s slightly defective regenerative power is not only a boon for the show’s fans (and producers), but it is also a rare example of science fiction being outdone by the real world. On Earth, a wide range of actual organisms can regenerate missing parts after injury. Unlike the fictional Doctor, these animals can rebuild new structures that are indistinguishable from those they are replacing. With the goal of understanding the factors that enable some organisms, but not others, to restore missing structures, biologists have been studying regeneration for well over two centuries (Morgan, 1901; Lenhoff and Lenhoff, 1986).

Although the Doctor’s regenerative abilities remain a mystery, recent research on various earthlings has started to reveal the mechanisms leading to regeneration of complex structures. During the past decade the application of molecular genetic techniques, including double-stranded RNA-mediated genetic interference (RNAi) and transgenesis, has revitalized studies of classical animal models of regeneration (Poss, 2010). Here we highlight a sample of recent work addressing some of the critical questions facing regeneration researchers: What signals initiate regeneration? What is the nature of the cells that produce the regenerate? What role does the nervous system play in this process? The answers to these and related questions should help developmental biologists, tissue engineers, and clinicians to understand and one day overcome the limits on human regeneration.

Back from the dead: Apoptosis and regeneration

After amputation, local responses at the site of the wound play important roles in the initiation of regenerative processes. Wound healing (Gurtner et al., 2008), ion flux (Levin, 2009), and interactions between the wound epidermis and the underlying tissue appear critical for regenerative outgrowth. Recent work suggests that programmed cell death may play a role in triggering regenerative responses in many different organisms, including Hydra (Bergmann and Steller, 2010).

Hydra, a freshwater polyp belonging to the Phylum Cnidaria (which includes jellyfish, sea anemones, and corals), was the subject of the first scientific investigations of regeneration in animals (Lenhoff and Lenhoff, 1986). The Hydra body consists of ectodermal and endodermal cell layers separated by an extracellular matrix; neurons and interstitial cells (stem cells that produce neurons, gland cells, and germ cells, among other cell types) reside between these two layers. The animal is radially symmetric, and is polarized along the oral (head and tentacles)/aboral (foot) axis (Fig. 1 A). Small fragments of Hydra tissue can regenerate a complete organism; even dissociated single cells can reaggregate, reestablish polarity, and form a new animal (Noda, 1971; Gierer et al., 1972).
An apoptotic program is initiated asymmetrically after transection midway through the body column. The fragment that will regenerate a new head displays a robust apoptotic response near the wound site, whereas the fragment regenerating a new foot does not (Fig. 1 A; Chera et al., 2009). Inhibition of apoptosis blocks head regeneration, and activation of apoptosis at a foot-regenerating wound leads to ectopic head formation (Chera et al., 2009). Thus, induction of apoptosis is both necessary and sufficient for head regeneration. The apoptotic response stimulates a synchronous burst of proliferative activity in neighboring cells and leads to the establishment of a head organizer via secretion of Wnt3 ligand by apoptotic interstitial cells (Chera et al., 2009). RNAi knockdown of Wnt3 prevents the proliferative burst induced by apoptosis, whereas treatment with exogenous Wnt3 rescues proliferation when apoptosis is inhibited (Chera et al., 2009). The induction of apoptosis after injury appears to require the mitogen-activated protein kinase (MAPK) pathway, including ribosomal S6 kinase (RSK), and cAMP response element–binding protein (CREB). RNAi knockdown of RSK and CREB blocks apoptosis at the head-regenerating tip, as does treatment with U0126, a pharmacological inhibitor of MEK, the kinase that phosphorylates MAPK (Chera et al., 2011). In fragments that will regenerate a new head, the transcription factor CREB is phosphorylated by RSK within minutes of amputation (Fig. 1 A; Chera et al., 2011). Intriguingly, MAPK/CREB-induced apoptosis is not required in fragments
regenerating a foot, and cell proliferation is not up-regulated in foot regeneration. Rather, foot regeneration proceeds through cellular rearrangement and transdifferentiation (morphallactic regeneration). The exact nature of the signal that leads to asymmetric MAPK activation (and, thus, apoptosis) at the site of injury in head-regenerating fragments remains to be determined.

The apoptosis-induced compensatory proliferation (Fan and Bergmann, 2008) observed in Hydra may be a conserved mechanism for stimulating proper wound healing and regeneration. Apoptotic cells have been observed during early phases of regeneration in several animals that can regenerate missing tissues: planarians (Fig. 1 B), Xenopus (Fig. 1 C), and newts (Hwang et al., 2004; Vlaskalin et al., 2004; Tseng et al., 2007; Chera et al., 2009; Pellettieri et al., 2010). Although the requirement for apoptosis during regeneration has not been addressed in planarians and newts, treatment of Xenopus larvae with caspase inhibitors during the first 24 h after amputation blocks tail regeneration (Tseng et al., 2007). As observed in Hydra, inhibition of apoptosis impedes proliferation; in addition, nerves fail to extend appropriately toward the site of amputation. Because nerves release important regenerative signals (see penultimate section), it is unclear whether reduced proliferation results directly from lack of signals derived from apoptotic cells, or indirectly from defective innervation. Furthermore, because these assays rely on caspase inhibition it is possible that nonapoptotic roles for caspases (Kuranaga and Miura, 2007) may be involved rather than apoptosis by itself.

Apoptotic cells have been shown to provide a number of signals that can regulate wound healing and regeneration. In Drosophila, the larval imaginal discs are capable of regenerating: wing discs can produce appropriately sized adult wings after radiation-induced killing of over 50% of the cells (Haynie and Bryant, 1977). MAPK signaling through Jun kinase is important both for initiating apoptosis and production of Wnt (Wg) and BMP (Dpp) mitogens (Pérez-Garito et al., 2009; Bergantiños et al., 2010; Morata et al., 2011). The extent to which these factors are required for stimulating compensatory proliferation remains unclear. After radiation-induced apoptosis, neither dpp nor wg were required (Pérez-Garito et al., 2009). However, after ectopic expression of a pro-apoptotic gene, wg was required for the proliferative response (Smith-Bolton et al., 2009). Furthermore, wg is also required for disc regeneration after surgical transection (Schubiger et al., 2010). Additional roles for apoptosis have been reported in epidermal wound healing and liver regeneration in mouse. Caspase 3 and caspase 7 mutant mice have defects in both processes, and these mutants show reduced cell proliferation in these contexts (Li et al., 2010). Caspases 3 and 7 can activate Ca$^{2+}$-independent phospholipase A$_2$, leading to production of arachidonic acid and prostaglandin in apoptotic cells, the latter of which can stimulate proliferation (Li et al., 2010).

Injury-induced apoptotic signals are also required to maintain tissue homeostasis. When cells of the adult Drosophila midgut are injured by toxins or induced to undergo apoptosis, intestinal enterocytes secrete the cytokine Unpaired, which stimulates proliferation of intestinal stem cells through activation of the Jak/Stat pathway (Jiang et al., 2009). Similarly, in the mouse intestine massive induction of apoptosis (via intestine-specific knockout of the p53 inhibitor Mdm2) is eventually compensated in adults by increased proliferation and expansion of the stem cell pool (Valentín-Vega et al., 2008). Apoptotic cells also contribute to homeostasis in epithelia by lipid-based signaling (sphingosine-1-phosphate) that triggers actomyosin contraction in the surrounding cells, leading to the extrusion of the dying cells (Gu et al., 2011). These observations suggest many potential roles for dead and dying cells to alter cell behavior at sites of injury.

Conjuring up spare parts: Cellular sources of regeneration

In most regenerating organisms, replacing an amputated structure requires the production of new cells. Therefore, one of the main functions of early signaling events after injury is to stimulate the production of additional cells that are capable of rebuilding lost structures. New cells coalesce near the site of injury, giving rise to a mass of undifferentiated cells called the regeneration blastema. Subsequent signals then regulate outgrowth and patterning of the newly formed tissue. To understand how early signaling events initiate regeneration and stimulate blastema formation, it is crucial to identify the cells upon which these signals act. New cells can be generated in a variety of ways, including proliferation of a resident stem cell population, division of terminally differentiated cells, or dedifferentiation/transdifferentiation of mature cells to a stem cell–like precursor or another cell type (Fig. 2 A). The extent to which each mode is used varies between species and even across tissues within the same species.

Stem cells and transdifferentiation both contribute to the regenerative abilities of Hydra. These animals possess three distinct stem cell populations: ectodermal, interstitial, and endodermal stem cells (Galliot et al., 2006). Development of transgenesis in Hydra (Wittlieb et al., 2006) has enabled in vivo tracking of the stem cell lineages: for example, investigating the differentiation of interstitial stem cells and migration of their progeny in intact animals (Khalturin et al., 2007). Transgenesis has also aided the analysis of transdifferentiation. By expressing eGFP in the zymogen gland cells (a derivative of the interstitial stem cells), Siebert et al. (2008) showed that these cells move up the body column and down-regulate the expression of a zymogen gland cell–specific marker. By using histology, electron microscopy, and in situ hybridization with cell type–specific markers, they identified cells with characteristics of both zymogen gland cells and another cell type (granular mucous cells), suggesting that zymogen gland cells could convert to granular mucous cells as they were displaced up the body column and entered the head region (Siebert et al., 2008). The ability of Hydra cells to transdifferentiate permits these animals to regenerate even in the absence of cell proliferation (Cummings and Bode, 1984).

Freshwater planarians (another classical model for studying regeneration) owe their amazing regenerative abilities to stem cells. In these animals, a population of mesenchymal stem cells, called neoblasts, is the source of cells for regeneration (Baguña et al., 1989). Neoblasts are the only dividing somatic cells and have been defined by their high nuclear/cytoplasmic
Figure 2. Cellular sources of regeneration. (A) The ability to regenerate amputated structures often requires the production of new cells. These new cells can be derived from amplification and differentiation of resident stem cells, proliferation of differentiated cells, dedifferentiation of cells to a more primitive state, or transdifferentiation of one cell type to another cell type. (B) Transmission electron micrograph of a neoblast from the planarian *Schmidtea mediterranea*. Planarians owe their impressive regenerative abilities to these adult stem cells. Neoblasts are the only mitotic somatic cells and are defined by their high nuclear (blue) to cytoplasm (green) ratio and the presence of cytoplasmic ribonucleoprotein complexes called chromatoid bodies (arrows). Image courtesy of Ana Vieira (University of Illinois at Urbana-Champaign, Urbana, IL). (C) Rescue of a lethally irradiated planarian by introduction of a single neoblast. *S. mediterranea* exists as two genetically distinct strains: an asexual strain (brown) that reproduces by transverse fission; and a sexual, hermaphroditic strain (gray) that reproduces by cross-fertilization. Wagner et al. (2011) have shown that lethally irradiated sexual animals can be rescued by injection of a single asexual neoblast. Because the asexual donor neoblast is the only source of new cells, the sexual host is eventually converted into an asexual animal after repeated rounds of amputation, regeneration, and tissue turnover.
ratio and sensitivity to γ-radiation, as well as the expression of several post-transcriptional regulators and markers of proliferation (Fig. 2 B; Baguñà et al., 1989; Newmark and Sánchez Alvarado, 2000; Orii et al., 2005; Reddien et al., 2005; Salvetti et al., 2005; Guo et al., 2006; Yoshida-Kashikawa et al., 2007; Rouhana et al., 2010). Classical experiments showed that injection of neoblasts could restore regenerative abilities and long-term viability to lethally irradiated planarians; furthermore, injection of neoblasts derived from a sexual planarian strain could “transform” lethally irradiated asexual individuals into sexuals (Baguñà et al., 1989). Because these injections used thousands of cells, neoblasts as a population have long been considered pluripotent. However, whether individual neoblasts are pluripotent or if there are subsets of lineage-committed neoblasts has remained an open question. Recent work by Wagner et al. (2011) provides convincing evidence that a subset of neoblasts is pluripotent. They injected single asexual donor neoblasts into lethally irradiated sexual hosts (the irradiated sexual animals survive longer after irradiation than the asexuals do, permitting sufficient time for clonal expansion of the injected cells). Single neoblast injections were able to restore viability to the sexual animals and convert them to an asexual mode of reproduction. Restriction fragment length polymorphism and haplotype sequencing confirmed the genotypic conversion of the host to that of the donor strain (Fig. 2 C; Wagner et al., 2011). Characterizing this pluripotent subset of the neoblast population will be an important avenue for future research.

The source of regenerative cells in vertebrates varies between tissues and organisms, and in some cases remains a matter of continued debate. Many vertebrate tissues contain adult stem cells that play important roles in tissue turnover and homeostasis. Nonetheless, division, dedifferentiation, and transdifferentiation of differentiated cells contribute to regeneration in several different contexts. For example, whereas liver progenitor cells appear to be major sources of new hepatocytes under conditions of extreme damage or chronic disease, restoration of liver mass after partial heptectomy or mild liver injury is largely accomplished through proliferation of remaining hepatocytes (Riehle et al., 2011). Similarly, recent lineage-tracing experiments indicate that after damage to the zebrafish heart, existing cardiomyocytes undergo dedifferentiation and proliferate to generate new cardiomyocytes for replacing lost heart mass (Jopling et al., 2010; Kikuchi et al., 2010). Pigmented epithelial cells in the newt dorsal iris can regenerate a new lens via transdifferentiation: these cells from the dorsal iris can dedifferentiate, reenter the cell cycle, and differentiate to produce new lens cells (Henry and Tsonis, 2010).

Dedifferentiation also contributes new cells during appendage regeneration in Urodele amphibians (news and axolotls). Near the site of amputation, syncytial skeletal myotubes fragment and produce mononucleate cells that reenter the cell cycle (Lo et al., 1993; Echeverri et al., 2001; Kumar et al., 2004). Skeletal muscle from adult newts also contains Pax7-positive muscle stem cells (satellite cells) that become activated and contribute to new tissues during regeneration (Morrison et al., 2006, 2010). The respective contributions of muscle dedifferentiation and satellite cell activation during limb regeneration in the axolotl and newt remain open questions. In contrast, larval tail regeneration in *Xenopus laevis* appears to be driven largely by progenitor cells; in lineage-tracing experiments the extent of labeling of new muscle correlates with the amount of satellite cell labeling before amputation (Slack et al., 2004). Lineage tracing experiments in both axolotl and *Xenopus* indicate that multiple cell types contribute to formation of the blastema (Slack et al., 2004; Kragl et al., 2009). However, it is still unclear whether these other tissues contribute cells through proliferation of progenitor cells or dedifferentiation of mature cells. Such an understanding will be crucial for deciphering the mechanisms by which early regenerative signals trigger blastema formation.

**To thine own self be true: Cellular memory during regeneration**

In addition to identifying the cellular sources of regeneration in different systems, researchers have been examining whether blastemal cells are pluripotent, multipotent, or have more limited potential. As discussed already, pluripotent neoblasts are the source of new cellular material that drives regeneration in planarians. However, cell proliferation is restricted largely to regions outside the blastema (Reddien and Sánchez Alvarado, 2004; Wenemoser and Reddien, 2010); thus, post-mitotic neoblast progeny are the predominant cells within the blastema that rebuild lost tissues. Eisenhoffer et al. (2008) have exploited the sensitivity of neoblasts to γ-radiation to identify numerous neoblast as well as early and late neoblast progeny markers. Interestingly, not all early neoblast progeny express the same panel of markers (Fig. 3 A). Although this difference could reflect transient temporal alterations in gene expression in early neoblast progeny, it more likely indicates that these cells are in various early stages of lineage commitment and/or en route to different terminal cell types. This heterogeneity has also been suggested by gene expression profiling on individual neoblasts (and their progeny) isolated using fluorescence-activated cell sorting (Hayashi et al., 2010). A subset of radiation-sensitive cells with G2 DNA content and expressing various neoblast markers also expressed a marker of muscle differentiation, suggesting that not all lineage-committed neoblasts are post-mitotic (Hayashi et al., 2010). Consistent with this result, S phase or G2 neoblasts have been observed to express markers of differentiated cell types, including excretory and neuronal markers (Nishimura et al., 2011; Scimone et al., 2011).

In planarians, specification of blastema positional identity (i.e., as anterior or posterior) likely occurs early during regeneration. Sengel (1960) removed anterior blastemas after 3 d of regeneration and cultured them as explants in vitro; these blastema explants produced only anterior tissues, including photoreceptors and cephalic ganglia (Fig. 3 B; Sengel, 1960). By contrast, explanted posterior blastemas did not produce anterior tissues. Therefore, these early experiments suggested that specification of blastemas to produce heads or tails occurs during early stages of regeneration. This idea has been confirmed by recent molecular studies showing rapid up-regulation of polarity signals (Petersen and Reddien, 2009, 2011; Gurley et al., 2010). Surprisingly, when anterior blastemas were co-cultured together with posterior blastemas, they were now able to produce small planarians
containing head, central body, and tail structures (Fig. 3 B). These results suggest that, although neoblast progeny are specified early during regeneration, they may retain some developmental plasticity; in response to altered position cues (e.g., juxtaposition of anterior and posterior tissues) they may be able to change their fates and generate different cell types and tissues than they would have formed otherwise. Alternatively, the blastema explants may have contained some neoblasts that responded to the juxtaposition of anterior and posterior tissues to generate distinct body structures.

Recent experiments using transgene-based lineage tracing suggest that vertebrate blastema cells can “remember” the tissue from which they are derived, and that their fates are largely restricted to forming similar tissues in the regenerate. For example, transgenesis in axolotl and *Xenopus* has been combined with embryonic grafting to specifically label various tissues, including muscle, Schwann cells, spinal cord, and dermis; this tissue-specific labeling enables the contribution of various tissues to be analyzed during regeneration (Fig. 4 A; Gargioli and Slack, 2004; Kragl et al., 2009). These experiments revealed that regenerated muscle tissue is derived solely from muscles present in the limb before amputation. However, in the axolotl, interconversion of dermis and cartilage was observed. Both of these tissues are derivatives of lateral plate mesoderm, suggesting that these tissues may dedifferentiate to produce progenitors restricted to lateral plate fates or that the dermis may contain an uncommitted stem cell population (Kragl et al., 2009).

Lineage restriction has also been observed during fin regeneration in zebrafish. Tu and Johnson (2011) have used transposon-based clonal analysis to examine the potency of various cell lineages during development, growth, and regeneration (Tu and Johnson, 2010, 2011). This method stably labels one to a few cells in the developing fin bud, enabling the progeny of single fin bud cells to be monitored. Examination of hundreds of animals indicated that fin bud cells are greatly restricted in developmental potential, only generating one to a few cell types in the adult fin (Tu and Johnson, 2011). As in axolotl and *Xenopus*, zebrafish caudal fin cells remain lineage committed during regeneration, and do not contribute to lineages other than that from which they are derived (Tu and Johnson, 2011). Similar results were obtained in studies of bone regeneration in the zebrafish fin. Osteoblasts near the amputation site downregulate osteoblast differentiation markers, lose their differentiated morphology, proliferate, and give rise to new bone in the regenerate (Knopf et al., 2011). These results reveal that regeneration in these contexts does not require dedifferentiation to a pluripotent state, and that the cells that make up the blastema are lineage restricted. These lineage-restricted progenitors even occupy distinct spatial domains in the blastema, indicating that the initial sorting and positioning of these cells may be crucial for producing a properly patterned appendage (Kragl et al., 2009; Tu and Johnson, 2011).

Although the lineage restrictions observed during limb regeneration (Kragl et al., 2009) are similar to those observed during limb development (Pearse et al., 2007), there are dramatic differences between development and regeneration. In addition to nerve dependence (see next section) and differences in scale of the structures being formed, limb regeneration differs significantly from limb development in that some components of the proximo-distal axis of the limb are retained after amputation. To prevent duplication or deletion of limb structures, cells at the site of amputation must be able to determine their position along the proximo-distal axis and use this information to
Based on its differential expression along the proximo-distal axis, da Silva et al. (2002) identified a cell surface protein, referred to as Prod1, that is expressed at high levels proximally and low levels distally. Treatment with anti-Prod1 antibodies blocked the encapsulation of distal blastema by proximal blastema tissues, suggesting that Prod1 plays a role in cell–cell interactions that mediate identity along the proximo-distal axis. Furthermore, when distal blastema cells overexpressing Prod1 were transplanted into proximal blastemas, they contributed to proximal rather than distal structures (Echeverri and Tanaka, 2005). The mechanisms by which differences in Prod1 expression along the proximo-distal axis translate into faithful regeneration of the limb are not completely clear; however, Prod1 has been found to interact with a secreted protein called newt Anterior Gradient (nAG; see next section), providing a link between proximo-distal patterning and the role of innervation in regeneration (Kumar et al., 2007a).
"Anything's possible if you've got enough nerve": Nerve dependence in regeneration

The above, out-of-context quote (Rowling, 2003) introduces the important role that innervation plays during regeneration in many organisms. For example, earthworms in which a portion of the ventral nerve cord was removed near the amputation plane failed to regenerate new heads, whereas manipulations resulting in two anteriorly exposed nerve cords produced two heads (Morgan, 1901). In another annelid, the polychaete *Spirographis spallanzani*, deviation of the nerve cord toward the lateral body wall induced the formation of supernumerary structures: the anterior-directed cut nerve cord led to the formation of an ectopic head, whereas the posterior-directed cut nerve cord led to the production of an ectopic tail (Kiortsis and Moraitou, 1965). The nervous system also appears to be required for arm regeneration in Echinodermes because removal of the radial nerve in the starfish prevents arm regeneration (Huet, 1975).

In planarians, the nervous system is required for proper regeneration of the gonads and accessory reproductive organs. Collins et al. (2010) identified a single neuropeptide that mediates the effects of the nervous system on the postembryonic development and maintenance of the reproductive organs. Furthermore, the formation of ectopic cephalic outgrowths in planarians with improperly connected cephalic ganglia and ventral nerve cords suggests that discontinuities between the brain and nerve cords result in the production of signals that trigger proliferative outgrowth (Cebrià and Newmark, 2007).

The role of innervation is best understood in the context of amphibian limb regeneration. Classical experiments by Singer showed that denervated limbs failed to regenerate, and that the amount of innervation was critical for proper regeneration (Carlson, 2007). Kumar et al. (2007b) identified the nAG protein as a ligand for Prod1, the cell surface protein involved in proximo-distal patterning. During regeneration nAG is expressed in the Schwann cells of the nerve sheath and, later, in gland cells of the wound epidermis. Expression in both cell types is dependent on proper innervation. Remarkably, introduction of a vector encoding nAG into denervated limbs was able to rescue regeneration to the digit stage, thus restoring proximo-distal patterning. Experiments in which nAG was added to blastemal cells in culture revealed that nAG exerts its effects upon regeneration by stimulating blastemal cell proliferation. The mechanism by which nAG stimulates blastema cell proliferation remains an open question. However, when Prod1 is expressed in cultured newt blastema cells, it leads to the activation of MMP9 expression and ERK signaling; it also interacts with the epidermal growth factor receptor (Blassberg et al., 2011). It is important to determine whether nAG plays a role in mediating these activities of Prod1.

Given the wide distribution of regenerative abilities throughout the animal kingdom, it has been postulated that some regenerative mechanisms have been conserved throughout evolution (Sánchez Alvarado and Tsonis, 2006; Brockes and Kumar, 2008; Bely, 2010; Poss, 2010). According to this view, lack of regenerative abilities reflects evolutionary loss. For good reasons, most investigators have focused their studies on broadly conserved developmental regulators. However, Prod1 provides an intriguing example of a molecule that has apparently evolved within a specific evolutionary lineage, the Urodele amphibians (Garza-García et al., 2010). Thus, it will also be important for investigators to consider the roles of such taxon-specific genes in regeneration in diverse organisms. Ultimately, we need to know a great deal more about the cellular and molecular mechanisms operating during various regenerative processes to better understand the distribution of regenerative abilities throughout the metazoa.

Concluding thoughts

As discussed in this Review, investigations of model organisms have begun to characterize the cellular sources of regeneration, define their potency, and identify molecules required for restorative events. Thus, there is now a great opportunity for cell biological studies to link gene function to cellular behavior. For example, the cell biology of dedifferentiation is poorly understood: How is the complex cytoskeleton of cardiomyocytes or skeletal muscles reconfigured during the course of dedifferentiation to enable assembly of a mitotic spindle? How are mononucleate cells produced by syncytial myotubes? In addition, we need to understand how individual cell behaviors are integrated across tissues to permit a coordinated response to tissue loss. For example, the observation that tissues from proximal limb blastema encapsulate those from distal limb blastema suggests that cell–cell interactions are critical for proper sorting and organization within the blastema (Nardi and Stocum, 1984; da Silva et al., 2002). The nature of such differential adhesion remains to be determined in regenerating tissues. Similarly, it will be important to characterize how the extracellular environment is altered during regeneration, and how different extracellular matrices shape cell behaviors in the blastema (Calve et al., 2010). One of the long-term goals of regeneration research is to understand why humans have such limited regenerative potential and what, if anything, can be done to improve it. The knowledge gained from studying cell biological questions in model organisms should help drive such future efforts of regenerative medicine.

We thank Jim Collins, David Forsthoefel, Rachel Roberts-Galbraith, and the anonymous reviewers for helpful comments on the manuscript, as well as Ana Vieira for providing the electron micrograph for Figure 2. We extend our sincere apologies to the many colleagues whose work we were unable to cite due to space limitations.

Research in P.A. Newmark's laboratory is supported by the National Institutes of Health (RO1-HD043403) and the National Science Foundation (IOS-0744689). P.A. Newmark is an investigator of the Howard Hughes Medical Institute.

Submitted: 19 May 2011
Accepted: 24 January 2012

References


