The Pharmacology of Regenerative Medicine

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I. Introduction to Regenerative Pharmacology

Historically, small molecule (i.e., compounds of <500–800 mol. wt.) pharmaceutical research and development has focused on compounds with increasingly selective mechanisms of action. This makes sense from a symptom-based approach to the treatment of disease, wherein one wishes to focus on the primary mechanism of action required for drug efficacy while simultaneously limiting off-target effects and minimizing adverse events/side effects. The development requirements for regenerative pharmacology will be much more demanding. In fact, the challenges associated with regenerative pharmacology, that is, curative therapeutics, will in many instances require complex mixtures of compounds [i.e., growth factors such as 3D, three dimensional; 6-OHDA, 6-hydroxydopamine; AADC, L-amino acid decarboxylase; AAV, adeno-associated virus; AFs, amniotic fluid cells; Ale, alendronate; AR, adrenoreceptor; BAM, bladder acellular matrix; BMPs, bone morphogenic proteins; BrdU, bromodeoxyuridine; CHIR99021, 6-(2-(4-(2,4-dichlorophenyl)-5-(4-methyl-1H-imidazol-2-yl)-pyrimidin-2-ylaminoethyl)-aminonitritile; Col IV, collagen type IV; DA, dopamine; ECM, extracellular matrix; EGF, epidermal growth factor; EHNA, erythro-9-(2-hydroxy-3-nonyl)adenine; EPO, erythropoietin; ES, embryonic stem; FDA, Food and Drug Administration; GCSF, granulocyte colony-stimulating factor; GDNF, glial cell line-derived neurotrophic factor; GSK, glycogen synthase kinase 3; HSC, hematopoietic stem cells; IF, insulin-like growth factor; iPS, induced pluripotent stem; I-Q-1, 2-[4-(acetylphenyl)diazeyn]-2-(3,3-dimethyl-2,4-dihydro-1H-isooquinol-1-yl)acetamide; molecular formula C21H21N4O2; L-DOPA, L-3,4-dihydroxyphenylalanine; LP, lamina propria; MP, muscularis propria; MPTP, 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine; MSC, mesenchymal stromal cells; NE, a protein strongly expressed in neural tissue encoding epidermal growth factor-like domain; NELL1, NELL-like molecule-1; NGF, nerve growth factor; NSC, neural stem cells; NTN, neurturin; PD, Parkinson’s disease; PLAGa, poly(lactic-co-glycolic acid); RM, regenerative medicine; RRT, Rett syndrome; SCNT, somatic cell nuclear transfer; STC, subtotal cystectomy; TE, tissue engineering; TGF, transforming growth factor; VEGF, vascular endothelial growth factor; XAV939, C14H11F3N2OS.]
fibroblast growth factor (FGF), epidermal growth factor (EGF), platelet-derived growth factor, nerve growth factor (NGF), vascular endothelial growth factor (VEGF), insulin-like growth factor (IGF), bone morphogenetic proteins (BMPs), etc.] for restoration of tissue/organ function. These latter compounds have significantly higher molecular weights (generally ~10,000 to >100,000 mol. wt.) than those traditionally developed by the pharmaceutical industry.

In this article, we attempt to pull together a rather vast amount of scientific and technical information from increasingly intersecting interdisciplinary fields of research to emphasize the significant role that pharmacologists can play in developing curative therapeutics. So, what are the potential implications of regenerative pharmacology? Imagine the day when:

1. Drugs can be targeted to specific nuclei in the brain (e.g., the center affected in Parkinson’s Disease) or any desired region(s) of organs/tissues to exert local therapeutic or healing effects without untoward side effects;
2. Multiple bioactive compounds can be loaded into a sophisticated drug delivery system(s) that is locally placed to orchestrate a complete functional regenerative response;
3. One can sufficiently recapitulate the complexity of the internal milieu to permit new functional tissue and organ formation in vitro for subsequent implantation in vivo.

In his recent State of the Union address President Obama alluded to the crucial impact of such efforts on scientific innovation:

“If we want to make the best products, we also have to invest in the best ideas. Every dollar we invested to map the human genome returned $140 to our economy. Today, our scientists are mapping the human brain to unlock the answers to Alzheimer’s; developing drugs to regenerate damaged organs; devising new material to make batteries ten times more powerful. Now is not the time to gut these job-creating investments in science and innovation.” (Read more: http://www.whitehouse.gov/state-of-the-union-2013.)

A major goal of this report is to emphasize that the success of such an effort will be accelerated by the rigorous application of the pharmacological sciences. We currently lack a broad knowledge of the complex pharmacology of mammalian wound healing and functional regeneration. Correction of this knowledge gap demands a global multidisciplinary, collaborative research and effort to stimulate the conversations that must occur at the intersections of pharmacology, biomaterials, biomedical/tissue engineering, nanotechnology, stem cell and developmental biology, etc. We believe that the conceptual framework and scientific foundations, as well as many of the technologies required for success, are already in place or being developed, but the effort is not organized and the necessary conversations are not happening. We hope that the readers of this report will grasp the considerable value of this effort and form the sustained alliances and collaborations required to begin the journey.

However, before launching into a comprehensive discussion of regenerative pharmacology and its central role in the continued development of regenerative medicine technologies, as outlined in Fig. 1, it is important to provide some fundamental background information about the nature of tissue/organ regeneration and the current status of regenerative medicine technologies.

A. Regeneration and Regenerative Medicine.

Tissue and organ regeneration occurs throughout the animal kingdom, and this phenomenon has understandably captured the scientific imagination for hundreds of years (Nachtrab and Poss; 2012). There are large disparities in regenerative capacity both between species (e.g., amphibian versus mammalian) and among organs (e.g., liver versus kidney). Exploration of these differences has offered insights regarding the mechanistic basis of regeneration and the diminished or apparently absent regenerative potential in certain systems, including many human tissues (Stocum, 2002; Taub, 2004; Sanchez Alvarado and Tsonis, 2006; Stocum and Cameron, 2011; Baddour et al., 2012). In this scenario, the extensive attention focused on regenerative medicine is understandable given the potential for repair or replacement of old, missing, damaged or diseased cells, tissues, and organs. In fact, regenerative medicine technologies are specifically developed for this purpose. The complexity of endogenous regeneration, the relatively limited mammalian capacity for regeneration, and the vast shortages of donor organs coupled with the seemingly ever-increasing life span of humans have combined to create a huge demand for regenerative medicine.

The goal of regenerative medicine can be concisely codified as the repair and/or replacement of damaged cells, tissues, and organs for functional restoration. It is a global, interdisciplinary effort with a translational research focus on development of therapies for patients afflicted with a variety of age- and disease-related disorders/dysfunction. Regenerative medicine (RM) and its companion field tissue engineering (TE) have provided a variety of current technologies for functional tissue/organ restoration, and these approaches have been described in detail in numerous publications (Freed et al., 2006; Mikos et al., 2006; Grayson et al., 2009; Corona et al., 2010; Atala et al., 2011; Badylak et al., 2012), and thus, only the most salient aspects are discussed herein. Figure 2 provides a general conceptual framework for many aspects of the TE/RM process.
B. Overview of Current Regenerative Strategies. Regardless of the precise strategy used for reconstruction, restoration, or repair of the tissue/organ of interest, cells and biomaterials (i.e., scaffolds) provide the basic constituents required for creating new tissue; they represent the “raw materials” from which tissues and organs are built. However, the exact TE/RM approach taken will necessarily depend on the degree of tissue/organ dysfunction. For example, if sufficient tissue/organ viability remains in vivo, then either cells alone (i.e., cell therapy) or scaffolds alone (biomaterial therapy) may be adequate to provide the required regenerative response. Such an approach is feasible at this point in the disease process, as it may still be possible to leverage existing endogenous mechanisms for relatively complete tissue repair and/or restoration of organ function. In contrast, when there is a dearth of viable tissue remaining, as in many cases of traumatic injury and in many congenital and acquired conditions, the degree of end organ dysfunction may be so great that it exceeds the endogenous regenerative capacity of the organ or tissue. In this scenario, any remaining endogenous repair mechanisms will require much greater augmentation via the implementation of TE strategies that produce more fully developed native-like tissue/organ biomimetics (i.e., biologic substitutes), up to and including whole organ replacement—including biomaterial strategies such as whole knee or hip replacements. One method commonly contemplated for creation and maturation of engineered tissues/organisms in vitro involves utilization of bioreactor technologies. Bioreactors are laboratory devices that...
recapitulate relevant aspects of the in vivo physiologic environment such as stretch, flow, compression, etc. By use of this approach, cells may be seeded on a biomaterial/scaffold, placed in a bioreactor, and subjected to appropriate environmental cues that are critical to tissue formation and function. In this fashion, bioreactors may be used to create more advanced three-dimensional (3D) tissue constructs in vitro prior implantation in vivo (see Freed et al., 2006; Goldstein and Christ, 2009; Grayson et al., 2009; Corona et al., 2010; Badylak et al., 2012). Alternatively, bioprinting, which simultaneously deposits cells and materials, can be in complex geometries reminiscent of native tissue architectures and may provide another feasible approach to the creation and assembly of 3D tissues and organs (Boland et al., 2006; Mironov et al., 2009; Jakab et al., 2010; Chang et al., 2011; Marga et al., 2012). The key point here, with respect to the focused aim of this report, is that pharmacology can play an obvious role in all currently contemplated approaches to TE/RM. This point is highlighted in Table 1.

C. Status of the Regenerative Medicine Enterprise. One index of the growing prominence, popularity, and expectations of regenerative medicine is the observation that a Google search for this phrase reveals nearly 6.1 million results (October 15, 2012). A number of substantive national efforts were recently launched to promote a sustained commitment to regenerative medicine. For example, the 2012 Annual Industry Report of the Alliance of Regenerative Medicine (http://alliancerm.org/sites/default/files/ARM-Annual-Industry-Report-2012.pdf) clearly indicates that pharmacology is poised to make a major contribution to the advancement of all major sectors of the regenerative medicine industry. In fact, the top 15 regenerative medicine products are already estimated to have treated 500,000 patients between 1998 and the end of 2011. As described in more detail below, contributions for pharmacology to the development of translational regenerative medicine technologies and therapies can be envisioned for cell-based therapies, as well as the small molecules, biologics, synthetic materials, biomaterials, and scaffolds—all of which are the subject of the Alliance of Regenerative Medicine 2012 Annual Report. Moreover, this same Washington, DC-based nonprofit organization has outlined a national strategy for regenerative medicine (http://www.alliancerm.org/). The overall mission of this organization is to educate key policymakers about the potential of regenerative medicine and, furthermore, to advocate for public policies that establish advantageous environments for funding, regulatory approval, and reimbursement strategies for regenerative medicine technologies/therapies. Such efforts have been aided by the introduction in the United States House of Representatives of the Regenerative Medicine Promotion Act of 2011 (HR 1862).

Another example of the increasing national commitment to regenerative medicine is the Armed Forces Institute of Regenerative Medicine (www.afirm.mil), which was officially formed in March 2008. The Armed Forces Institute of Regenerative Medicine consists of two civilian research consortia working with the U.S. Army Institute of Surgical Research in Fort Sam Houston, TX. Each consortium is a multi-institutional network with a combined total of more than 30 academic and 15 for-profit members. The recent establishment of an National Institutes of Health Center for Regenerative Medicine (www.crm.nih.gov) further bolsters the national effort in this emerging field. The National Institutes of Health also recently published a fact sheet on the past, present, and future of regenerative medicine research and clinical translation (http://report.nih.gov/NIHfactsheets/Pdfs/RegenerativeMedicine(NIBIB).pdf). In short, the present environment provides an excellent opportunity to bring

<table>
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<th>TE/RM Process/Need</th>
<th>Pharmacological Application</th>
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<tr>
<td>Functional evaluation of engineered and regenerating tissues</td>
<td>Preclinical assessment and pharmacological characterization of tissue/organ phenotype in vitro and in vivo*</td>
</tr>
<tr>
<td>Modulation of stem/progenitor cell expansion and differentiation</td>
<td>Screening of growth factor and small molecule libraries; development of improved culture systems (overlap of pharmacology and engineering) **</td>
</tr>
<tr>
<td>Targeted cellular delivery of drugs/chemicals to modulate regeneration in vivo</td>
<td>Development of novel drug delivery systems including biomaterials, nanomaterials, and bifunctional compounds that target active agents to specific tissue locations**</td>
</tr>
<tr>
<td>Biomaterials as reservoirs for bioactive agents and cell delivery vehicles for accelerated tissue formation and function in vitro and in vivo</td>
<td>Development of functionalized “smart” biomaterials**</td>
</tr>
<tr>
<td>Real-time modulation of tissue formation/regeneration/ morphogenesis*</td>
<td>Pharmacological modulation of the entire regenerative process: may incorporate all of the above elements, with the added complexity of replicating the exquisite spatiotemporal regulation characteristic of morphogen gradients in normal development**</td>
</tr>
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* denotes a “passive or dissecting” contribution of regenerative pharmacology, **denotes an “active or directing” role.
pharmacology to bear in the realm of regenerative medicine and tissue engineering.

With respect to the continued development of regenerative medicine therapies/technologies, we recently noted that: “… the broader clinical use of these groundbreaking technologies awaits improved understanding of endogenous regenerative mechanisms, more detailed knowledge of the boundary conditions that define the current limits for tissue repair and replacement in vivo, and the parallel development of critical enabling technologies (i.e., improved cell source, biomaterials, bioreactors)” (Corona et al., 2010). In fact, as outlined by Parenteau et al. (2012), the opportunity and need for regenerative medicine therapies to drive medical advances is tremendous, and moreover, the interdisciplinary effort that would be required to make this theoretical possibility a reality would be a significant driver of innovation and productivity per se. In addition, investigators have already begun to recognize the importance of the union of traditional pharmacology and regenerative medicine (Stayton et al., 2005; Mooney and Vandenburgh, 2008; Pucéat, 2008; Sakurada et al., 2008; Palatinius et al., 2010; Lee et al., 2011; Jadczyk et al., 2013), and others have begun to use similar terminology to describe this interface (Mozzetta et al., 2009). There is an entire volume devoted to regenerative pharmacology (Christ and Andersson, 2013). In this scenario, regenerative pharmacology is clearly poised to make major contributions to the development of novel therapeutics, and as outlined herein, there are numerous scientific tracks by which pharmacologists can become fully engaged and further accelerate the development of these next-generation clinical therapies. Some representative examples of the spectrum of potential therapeutic possibilities for regenerative pharmacology are presented throughout this document, and for the convenience of the reader, these examples are summarized in Table 2. Nonetheless, the effort remains at a very callow stage at this point.

Regenerative medicine can leverage important insights not only from studies of regeneration, as noted above and below, but significant advances can also be derived via improved understanding and application of mechanisms known to be responsible for tissue formation in the first place, that is, from the field of developmental biology. Below we provide a short overview of how understanding the pharmacology of morphogenesis can make important contributions to regenerative medicine.

D. Regenerative Pharmacology and Morphogenesis. Perhaps the importance of developmental biology (and endogenous regeneration of course) to regenerative pharmacology was intuitively obvious from the outset. That is, chemical processes guide the most fundamental aspects of tissue and organ formation and growth (i.e., morphogenesis) as well as regeneration. The implications of this for regenerative pharmacology are clear, because extracellular signaling molecules known as morphogens modulate the fate, movement, and organization of cells during morphogenesis in both embryos and adults (Wilson et al., 1997; Gurdon et al., 1998, 1999; Gurdon and Bourillot, 2001; Brookes and Kumar, 2008; Wolpert, 2011; Rogers and Schier, 2011; Bentzinger et al., 2012). Commonly studied growth factors, cytokines, and hormones such as the transforming growth factor (TGFβ) superfamily (i.e., TGFβ and BMPs), the fibroblast growth factor (FGF) family, Wnt/β-catenin signaling, retinoic acid, Wnt family members, hedgehog family members, and many others, are known to contribute to morphogenesis through a carefully orchestrated series of events. The activities of these factors are influenced by their respective diffusion profiles, effective concentration gradients, and concentration response relationships, as well as their potential modulation/quenching by the extracellular matrix and other components of the extracellular environment. Without doubt there are many unresolved questions regarding the precise mechanisms by which morphogen gradients guide tissue formation and development. Nonetheless, their impact on gene regulatory networks (Davidson, 2010) is increasingly being appreciated. These considerations form the basis for excellent recent reviews (Rogers and Schier, 2011; Kicheva et al., 2012) and entire volumes (Briscoe et al., 2010). In short, the large size and apparently exquisite

**TABLE 2**

List of diseases and disorders for which regenerative pharmacology approaches are currently being investigated/developed

<table>
<thead>
<tr>
<th>Disease, Injury, or Disorder</th>
<th>Section in Document</th>
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<tr>
<td>Heart and cardiovascular disease</td>
<td>II.D.1, IV.D.2, VI.A, VI.B</td>
</tr>
<tr>
<td>Diabetes</td>
<td>II.D.2, IV.E.</td>
</tr>
<tr>
<td>Genetic diseases</td>
<td>II.D.4</td>
</tr>
<tr>
<td>Scar reduction and wound healing</td>
<td>II.D.5</td>
</tr>
<tr>
<td>Bladder disease</td>
<td>III.A.</td>
</tr>
<tr>
<td>Parkinson’s disease</td>
<td>III.B., IV.E.3</td>
</tr>
<tr>
<td>Osteoporosis, bone fractures, bone grafting, spinal fusion</td>
<td>IV.B, IV.C.</td>
</tr>
<tr>
<td>Spinal cord injuries, amyotrophic lateral sclerosis, Alzheimer’s disease, cerebral palsy, macular degeneration</td>
<td>IV.A</td>
</tr>
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distribution requirements of morphogens for normal tissue formation and development indicate that novel drug delivery technologies will be required to ensure that morphogen gradients can be efficiently modulated for curative therapeutics. Certainly, this provides yet another important link to regenerative pharmacology (biomaterials development and drug delivery systems). A more detailed discussion of this point is beyond the focused aim of this report, and the interested reader is referred to the aforementioned references for additional information.

Having reviewed the general characteristics of, and requirements for, tissue and organ regeneration and engineering, the next key question is how exactly can regenerative pharmacology contribute to the development of novel therapeutics?

E. The Relationship of Regenerative Pharmacology to the Disease Process and Development of Novel Therapeutics. The explicit goal of regenerative pharmacology is to modulate cell, tissue, and organ physiology to accelerate, improve, or enhance functional outcomes (Andersson and Christ, 2007). However, this approach requires a radical change in thinking about the therapeutic development paradigm. Figure 3 outlines the progressive nature of the disease process and contrasts regenerative pharmacology with traditional pharmacotherapy. Most importantly, regenerative pharmacology can be used throughout the life cycle of the disease process, a major distinction from the more traditional pharmacological approaches. That is, the symptomatic treatment of age- or disease-related decrements in tissue or organ function are defined by a therapeutic window in which a sufficient amount of viable tissue must still be present to ensure efficacy. In stark contrast, the uses of regenerative pharmacology range from prophylactic applications through mitigation of reduced function to complete tissue/organ replacement in the advent of end organ failure (Fig. 3). However, as noted above, rigorous application of the pharmacological sciences toward creation of cures for disease requires a major paradigm shift in the discovery and development process for novel therapeutic modalities.

How might this dramatic shift in pharmacotherapy be achieved? As depicted in Fig. 1, we argue that pharmacology provides a critical lynchpin for the continued advancement of regenerative medicine and the discovery and development of novel curative pharmacotherapeutics. Ultimately, the seamless integration of the pharmacological sciences into regenerative medicine will require the concerted application of both passive and active processes (see Fig. 4). The active approach refers to the use of growth factors and other pharmacological agents to alter cell growth, differentiation, and function. (i.e., "direct,"); enhance, or repress as required; both in vitro and in vivo). The complementary passive approach relies on the use of established pharmacological methods to characterize endogenously regenerated or bioengineered cells and tissues and to dissect the regenerative process. Both of these approaches are currently used in regenerative medicine. However, more systematic application will be required to fully understand regeneration at the levels of molecules, cells, tissues, and organs, and thereby accelerate translational applications.

In this regard, expression of cell- and tissue-specific molecular markers and the presence of characteristic tissue and organ structure and architecture are necessary, but not sufficient, metrics for assessing the potential utility of engineered or regenerating tissues.

Fig. 3. Regenerative pharmacology and the disease process. Schematic diagram shows the initiation, development, and progression of tissue and organ dysfunction, leading ultimately to end organ failure. The potential utility of regenerative pharmacology approaches to the maintenance of normal tissue and organ function or the prophylaxis of continued decline is noted. However, the long-term goal is to develop curative pharmacological approaches that address the entire spectrum of tissue and organ function and dysfunction, so that regardless of the particular circumstance, a potentially curative therapy can be developed and applied. As described in detail in the text, regenerative pharmacology represents a significant departure from more traditional approaches that have necessarily focused on palliation and symptomatic relief of pathologic alterations in tissue and organ function.
Clearly, the most important barometer of success for tissue/organ engineering or regeneration technologies is their capacity for functional restoration (i.e., normal physiology). Thus, it is of critical importance that comprehensive physiologic evaluation of engineered and regenerating tissues/organs is embedded in the translational research paradigm.

A key aspect to the development of curative therapeutics will be effective delivery of potentially complex mixtures of high molecular weight compounds in a controllable spatiotemporal fashion. This fact points toward the absolute requirement for vastly improved biomaterials and drug delivery technologies and systems. As such, we devote the next portion of this review to a relatively comprehensive description of biomaterials and how they impact regenerative pharmacology.

**F. Drug Delivery Systems/Technologies and Biomaterials.** Advances in research at the intersection of biology, chemistry, and materials science have led to the development of increasingly sophisticated functionalized biomaterials, as well as novel drug delivery systems, as shown in Fig. 5. A comprehensive review of the latest functionalized biomaterials and modern drug delivery systems alone would require a lengthy dedicated report. Moreover, it should be emphasized that extant drug delivery systems and technologies comprise a wide array of mostly application-specific technologies. However, the potential uses of the existing technologies reviewed herein point toward future possibilities.

Most relevant to the focused aims of this review is the utilization of these technologies to 1) overcome the common set of barriers limiting the effectiveness of traditional pharmacotherapy and 2) extend the domain of deliverable therapeutic agents to a wider array of compounds (e.g., large molecular weight growth factors, gene therapies, etc.; see Fig 5). The first major barrier to a systemically delivered therapeutic is directing the agent to its tissue-level site of action. This involves achieving vascular extravasation or creating technologies...
that more efficiently deliver the “payload” (e.g., a drug, compound, or gene) from the systemic circulation to within the tissue(s) of interest. Approaches to achieve tissue-level localization include the use of long-circulating nanoparticles that home to sites of high vascular permeability (so-called “leaky vasculature”), delivery via transdermal delivery systems, or (most commonly in regenerative medicine approaches) direct injection or implantation routes. Once the desired agent has been delivered to its tissue target, a second major barrier exists with respect to local diffusion barriers within that tissue. Finally, any requirement for cellular and subcellular targeting specificity (e.g., gene therapy) provides a third major barrier to therapeutic success, that is, the chemical and structural barriers of the cell itself. We address all three of these key issues below.

II. Biomaterials in Regenerative Pharmacology

The field of biomaterials has undergone a transformation from the use of inert substances to the development of materials that are bioactive and can integrate into host tissues. The use of functionalized biomaterials can range from modifications of biomaterials to promote highly selective cell targeting—as in the case of nanoparticulate delivery systems—to the surface modification of implantable materials that promote cell attachment and tissue integration. In both instances, two classes of functionalized biomaterials often used in regenerative medicine applications are 1) particulate (micro- and nanoparticles) for cell and drug delivery and 2) scaffolding systems for tissue engineering approaches that carry or support cellular growth and tissue formation and/or regeneration. This report seeks to emphasize the former, but these two classes of biomaterials are highly interrelated in terms of their potential applications to regenerative medicine. For example, as shown in Fig. 5, I–K, scaffolding systems are often further functionalized by the incorporation of drug delivery systems into the materials. Alternatively, the functionalized biomaterial systems themselves can be drug delivery systems, either through release of exogenous therapeutic agents or through cell-based therapeutic release (Fig. 5, A–H).

A. Particulate Systems for Cell and Drug Delivery

Micro- and nanoparticulate delivery systems owe much of their development to the field of cancer therapeutics. The intent of many of these particulate delivery systems was to provide enhanced systemic delivery of therapeutic agents through improved pharmacokinetics (e.g., longer blood circulation) and pharmacodynamics (e.g., site-directed specificity). Systemic delivery systems offer the advantage of multiple dose administration at well-defined time points. The short half-lives of growth factors and nucleic acids commonly employed in regenerative medicine and tissue engineering suggests that such particulate delivery systems would also be advantageous for these applications because they provide protection from enzymatic degradation and hydrolysis. The ability of these delivery systems to protect therapeutics also makes them useful for inclusion in scaffolding systems.

Understanding each of these systems is important to understanding the potential breadth of their application(s) to tissue regeneration, repair, or replacement using pharmacological approaches. In fact, there are numerous examples of both particulate and implantable biomaterial systems being used for drug delivery applications. The nanoscale particulate systems are mostly based on self-assembly processes. Salient aspects of several of these technologies, which are specifically relevant to regenerative medicine and tissue engineering, are illustrated in Fig. 5.

1. Quantum Dots and Imaging Nanoparticles. Quantum dots are a crystalline lattice of atoms that act as semiconductors. These materials are gaining increasing usage in cancer studies and regenerative medicine (Fig. 5A). Their popularity as an imaging tool is largely related to their tunability, and applications to medical imaging include fluorescence and near infrared imaging technologies. Quantum dots are fabricated by dissolving an inorganic precursor (e.g., CdO may be used to serve as the Cd component of a CdSe crystal quantum dot) in organic surfactant (e.g., stearic acid) and solvent (e.g., octadecene) at relatively high temperature (e.g., 200°C). After cooling and addition of, e.g., an organophosphorous compound, the second component of the crystal (e.g., Se) may be added at elevated temperature to generate, in the examples above, CdSe nanocrystal quantum dots that are colloidal in nature (Li et al., 2003). Such technologies are critical to nondestructive imaging of engineered and regenerating tissues (theragnostics)—a key aspect to improved regenerative pharmacology approaches (see Fig. 5, E–H). Recently, the use of quantum dots to provide information on pharmacokinetic aspects of nanoparticles (see section II.A.3) was reviewed and indicates the potential for applying these technologies for nondestructive imaging in both pharmacological and tissue engineering realms (Probst et al., 2012).

2. Liposomes, Polymersomes, Micelles, and Cation/Anion Complexes for Encapsulation of Small Molecules, Peptides, Nucleic Acids, or Proteins. These are also self-assembling systems that are widely used for drug delivery because of the versatility of their payload. The self-assembly processes may be dictated by the materials themselves (liposomes, polymersomes, micelles) or through interaction of the materials with biologic molecules or drugs. An example of the latter is the self-assembly as polymer-DNA complexes (polyplexes) formed through electrostatic interactions of cationic polymers and negatively charged DNA (Fig. 5B). These technologies are critical components of regenerative
Fig. 5. Methods to generate functionalized biomaterials for regenerative medicine. Micro- and nanoparticle systems for cell and drug delivery (center): several micro- and nanoparticle systems are highlighted schematically. (A) Nanoparticles used for imaging modalities include quantum dots (fluorescence) and iron oxide nanoparticles (magnetic resonance imaging). Nanoparticles with hollow centers can also be loaded with iodine or other image contrast agents. A schematic of the structure of a quantum dot nanoparticle is shown. (B) In addition to contrast agents, small molecule drugs, nucleic acids, peptides, and protein drugs can be loaded into a variety of self-assembling nanoparticle systems that typically range from 10 to 200 nm. Schematics of DNA-polymer complexes, liposomes, and micelles are shown. (C) These nanoparticles can be surface modified with polyethylene glycol (left) to improve pharmacokinetics or can be modified with targeting motifs to improve cellular uptake (right). (D) Larger microscale constructs can also be formed from natural and synthetic polymers for release of therapeutic agents (right) or the delivery of cells (left) to provide cell-based delivery of, for example, insulin in the treatment of diabetes (Opara et al., 2010). Injectable delivery materials (left): the delivery systems described in the center panel have multiple applications to regenerative medicine when delivered either systemically or locally. (E) Shown is the in vivo tracking of implanted scaffolds containing cells loaded with ultrasmall superparamagnetic iron oxide nanoparticles (Reprinted with permission from Harrington et al., 2011). (F) The use of cationic liposomes to deliver DNA encoding for IGF-1 (and Lac-Z for imaging purposes) is shown at left (Reproduced with permission of BENTHAM SCIENCE PUBLISHERS LTD; Jeschke MG, Herndon DN, Baer W, Barrow RE, and Jauch KW (2001) Possibilities of non-viral gene transfer to improve cutaneous wound healing. Curr Gene Ther 1:267–278), whereas the delivery and protection of Wnt proteins for control of hair follicle stem cells to promote dermal thickening and follicle neogenesis in mice is shown at right (Morrell et al., 2008). (G) The ability to not only localize drugs but have the release of their payload triggered by internal (e.g., pH, temperature change, enzymes) or external (temperature, ultrasound, or as shown, light sources (Reprinted with permission from Azagarsamy MA, Alge DL, Radhakrishnan SJ, Tibbitt MW, and Anseth KS (2012) Photocontrolled nanoparticles for on-demand release of proteins. Biomacromolecules 13:2219–2224. Copyright © 2012, American Chemical Society)). (H) Incorporation of antigens into microparticles or nanoparticles for improved vaccine delivery is shown at left (Reprinted with permission from Demento SL, Cui W, Criscione JM, Stern E, Tulipan J, Kaech SM, and Fahmy TM (2012) Role of sustained antigen release from nanoparticle vaccines in shaping the T cell memory phenotype. Biomaterials 33:4957–4964) while the use of biomaterial implants is aiding in elucidating and ultimately minimizing inflammatory responses to implanted materials (Reprinted with permission from Norton LW, Park J, and Babensee JE (2010) Biomaterial...
pharmacology approaches when dealing with labile compounds/agents such as naked DNA and growth factors.

3. Functionalized Delivery Systems for Long Circulation Time, Receptor Targeting, or Vaccine Delivery. A key aspect to the development of these materials is surface modification. Such modifications may include, for example, grafting of poly(ethylene glycol) to decrease opsonization (Fig. 5C) and improve circulation time. Alternatively, one can include surface coupling of targeting ligands to the carrier (or polyethylene glycol chain as shown at the right) to improve selectivity of cell targeting and enhance cellular uptake. Such systems have been applied to target vascular sites of injury with applications to delivery therapeutics that can promote healing (Shi et al., 2012) or can be used to improve imaging modalities (Yang et al., 2011). It is also important to note that nanoparticle systems are gaining increased emphasis related to vaccine delivery (Fig. 5H) (Reddy et al., 2007; Foster et al., 2010). Given the role of implanted cells in regenerative medicine applications (including some involving autoimmune aspects such as diabetes), the role of nanoparticulate delivery systems for vaccine may be useful for several applications.

4. Synthetic or Natural Polymer Microparticles for Encapsulation of Cells or Therapeutic Agents. The use of microparticles is also common attributable to the ability to achieve sustained release of individual compounds/agents (Fig. 5D) or, alternatively, differential release profiles for distinct compounds/agents when used in conjunction with different materials or material scaffolds. Such technologies would be absolutely critical, for example, with respect to any attempt to recapitulate the exquisite features of development (morphogenesis) in vivo, as described above. Microcarriers are also an important aspect of regenerative medicine technologies with respect to cell encapsulation for various conditions including the delivery of insulin from islet cells as an approach to treat diabetes (Opara et al., 2010) and other diseases/disorders, and this application is considered in more details in section VI.

The methods for fabrication of nano- and microscale particulates are widely varied and too numerous to describe here. Nonetheless, it is worth noting some important applications in which these carrier systems are finding utility, which serve as a foundation for their inclusion in regenerative pharmacology approaches (Fig. 6). Most U.S. Food and Drug Administration (FDA) approved synthetic and natural polymers have been or can be formed into microscale carriers (~1–50 μm diameter). When delivered systemically, their dimensions restrict these systems to the vascular compartment, but approaches have been developed to localize delivery of the therapeutic to a specific region of the vasculature. Furthermore, microbubbles carrying therapeutics or imaging contrast agents can be disrupted by external application of ultrasound (Villanueva et al., 2007; Gao et al., 2008) and thereby achieve intracellular delivery of their payload (Barbarese et al., 1995). Delivery to specific sites of vascular injury can be accomplished by the coupling of targeting ligands (e.g., ligands or antibodies that target selectins or cell adhesion molecules upregulated at sites of vascular injury) to the surface of the microcarriers (Omolola Eniola and Hammer, 2005; Banquy et al., 2008), which essentially mimics the behavior of leukocyte rolling and adhesion. Such techniques are also finding applications in improving imaging modalities important to regenerative medicine (see above). Ultrasmall paramagnetic iron oxide nanoparticles (Harrington et al., 2011) or other imaging contrast agents encapsulated in nanocarriers can be used to improve nondestructive imaging modalities (i.e., theragnostics). Specifically, iron oxide is useful for magnetic resonance imaging (Xu et al., 2012a), iodine, or gold nanoparticles for computed tomography (Kao et al., 2003), and as noted above, quantum dots (de Mel et al., 2012) are also being used for imaging modalities in regenerative medicine.

As described above for vascular targeting, nanoscale systems (~10–200 nm) provide the ability to achieve a greater degree of tissue-specific targeting by
Methods to fabricate biomaterial scaffolds for regenerative medicine applications. There are many approaches to fabricating materials. These approaches range from inexpensive and relatively simple to expensive and quite complex. Several commonly used techniques are shown in this schematic. (A) Solvent evaporation/particulate leaching. A particulate (e.g., sodium chloride) that is insoluble in a particular solvent (e.g., chloroform) is cast with a polymer (e.g., PLGA) in solvent. After the solvent is evaporated, the material can be placed into an alternative solvent in which the particulate is soluble but the polymer is not to form the pores. (B) Sintering—particulate leaching that allows formation of interconnected pores of well-defined architecture. In this approach, leachable polymers are packed together and heated (to above their glass transition temperature) to allow partial fusion of the beads and provide a template. After cooling, a second polymer is cast around the sintered bead template to back-fill the empty regions. The polymer used to fabricate the bead template must be selectively soluble in a solvent. As described above, the bead template is then selectively dissolved in an appropriate solvent to yield a highly porous scaffold with interconnected pores (Fukano et al., 2010; Underwood et al., 2011). (C) Phase separation to introduce porosity (Nam and Park, 1999). This approach involves dissolution of a polymer into a solvent. The temperature is raised to one such that the polymer is fully solubilized. By cooling, the solution can phase separate depending on the concentrations of the solvent and the polymer. This phase separation can achieve solvent-rich regions or polymer-rich regions. Removal of the solvent (e.g., by evaporation) can achieve desirable pore architecture within scaffolds. These can be liquid-liquid phase separations, but it is also possible to introduce gaseous materials to achieve “gas foaming” of the desired pore architecture of the material (Riddle and Mooney, 2004). (D) Electrospinning—polymer dissolved in solvent is ejected through a small orifice (typically a needle). An electrical drop is applied between the orifice and collection device and fine nano-fibers are produced. It is also possible to incorporate nano or microparticles into these electrospun scaffolds (Guo et al., 2012). (E) Microfabrication techniques to introduce very high resolution into materials. Typically, such approaches are not used to produce large three-dimensional scaffolds for implantation. However, the techniques allow for very high levels of control over drug delivery or surface topography, allowing investigation of these effects at the individual cell level. (F) Three-dimensional printing/solid free-form fabrication techniques. These methods achieve high levels of dimensional precision for material applications.
In the application of nanoscale materials to the treatment of cancer, it is recognized that the vasculature of tumors has increased gaps in the endothelium and a reduction in the presence of lymphatic drainage, leading to the so-called enhanced permeation and retention effect. Although reliant on poorly defined vasculature and minimal lymphatic drainage, these principles may have the potential for application to certain aspects of regenerative medicine as well. Targeting the endothelium by conjugating ligands as noted above has also been accomplished with nanoscale carriers (Haun and Hammer, 2008), although not with the same success as with microscale systems (Charoenphol et al., 2010). For nanocarrier systems, the use of appropriate targeting motifs can achieve transcytosis in the case of endothelial or epithelial targeting (Ke et al., 2009) or endocytosis if the carrier is able to extravasate. Taken together, such technologies provide the potential to more selectively deliver therapeutics to target cells while limiting off-target sites (Saul et al., 2006).

To date, these methods have been largely emphasized in the cancer literature. However, the application of tissue- and cell-targeted carrier systems to the regenerative medicine space is increasing. For example, nonviral delivery of plasmid DNA encoding for IGF-1 (and Lac-Z for imaging purposes) when delivered via direct injection has been effective in improving wound healing responses (Fig. 5F) (Jeschke et al., 2001). Conceptually, nonviral DNA delivery may be advantageous to viral methods by reducing inflammatory response and achieving transient expression attributable to lack of genomic incorporation (Li and Huang, 2007). Liposomal or other nanocarriers may also be effective for the delivery of payloads that are sensitive to degradation such as through proteolysis. For example, Wnt proteins are important in various aspects of stem cell renewal and proliferation processes, but lack known agonists and stability. Wnt proteins, however, have been packaged into liposomes (see Fig. 5F). Delivery of Wnt via liposomes maintains their bioactivity and leads to dermal thickening and hair follicle neogenesis in mice, indicating the importance of their actions on stem cells in the follicle niche (Morrell et al., 2008).

Unlike cancer therapeutics where it is generally desirable to deliver a large payload of chemotherapy drugs to a tumor cell, regenerative medicine approaches typically are considered to benefit from spatiotemporally controlled delivery of therapeutic agents, thereby recapitulating relevant aspects of the carefully orchestrated tissue and organ development process (i.e., morphogenesis; again, see description above). Much effort in the development of such delivery systems involves methods that are crude relative to the exquisite morphogen gradients that guide tissue formation and development but nonetheless quite elegant in terms of polymer chemistry. Numerous release triggers exist, with temperature (Bessa et al., 2010), ultrasound (Borden et al., 2008), and light among the most commonly used methods to allow control over the timing of release. In one example shown (Fig. 5G), photocleavable cross-linkers are used for assembly of the nanoparticle encapsulating a therapeutic protein, which is released upon presentation of the triggering light source (Azagarsamy et al., 2012).

As noted above, systemic biomaterial drug delivery technologies, such as those currently under development for vaccines, are being increasingly applied to regenerative medicine. The use of nanoparticle-based biomaterials for vaccine delivery offers the potential to protect antigens, prolong release, or overcome biologic barriers attributable to the small size of the carrier technologies. Functionalized materials for these approaches take numerous forms. As shown in Fig. 5H, antigens may be encapsulated within FDA-approved polymers such as PLGA formed at diameters of several hundred nanometers up to several micrometers (Demento et al., 2012). In addition, the use of biomaterials as scaffolds for regenerative medicine applications emphasizes the importance of understanding the immunologic response to implanted materials per se. In fact, the selection of a preferred formulation or composition of a biomaterial or drug delivery system may even require the use of sustained release of anti-inflammatory agents at the site of implantation (Norton et al., 2010) to improve the biologic response to the material and thus facilitate the regenerative process.

As mentioned, overall, there are limited examples of systemically delivered materials for regenerative medicine applications. In general, drawbacks to the systemic delivery of therapeutic agents via functionalized biomaterial drug delivery systems suffer mainly from low accumulation at their site of action. However, progress in nonviral gene delivery and chemotherapy targeting are moving toward more effective compound delivery to target sites. Chemical constituents of the system, diameter and shape of the carrier (Gratton et al., 2008), surface charge (Georgieva et al., 2011), and the presence of targeting motifs (Ng et al., 2009) have been identified as key parameters for more effective systemic delivery of biomaterials and their cargo.

In summary, there are numerous combinations and permutations of functionalized biomaterials and drug
delivery systems that can be contemplated for use in regenerative pharmacology to promote tissue and organ regeneration and repair. Below we consider biomaterials that provide novel pharmacological approaches suitable for “building” tissue, that is, tissue engineering.

B. Biomaterials as Scaffolding Systems

Biomaterials are a key component of the tissue engineering paradigm, serving as a provisional matrix for cell infiltration and as depots for the delivery of therapeutic agents (see below). The point has been made that we need to think very differently about biomaterials (Williams, 2009), and without question, scaffolds for future generations of TE/RM technologies are expected to differ considerably from present-day implantable materials. Regardless, important design criteria for these scaffolds include 1) architecture and porosity; 2) mechanical properties and their role in directing cellular response; 3) physical and chemical cues for the promotion of cell attachment, migration, and differentiation; 4) compatibility with cell seeding or infiltration; and 5) degradation profiles suitable for tissue-specific regeneration.

Fabrication techniques play a significant role in defining these parameters for more effective scaffolds for TE/RM applications, and numerous approaches have been used to fabricate scaffolds from a variety of biomaterials. Several of the most promising or highly used are shown in Fig. 6. The techniques in use for regenerative pharmacology-based scaffolds differ significantly from the classic biomaterials fabrication techniques. Each technique has advantages and disadvantages, but generally speaking, higher levels of architectural organization are sacrificed for ease and speed of fabrication (Dalton et al., 2009). Porosity is an important aspect of biomaterial scaffolds for tissue engineering to allow cellular infiltration and ultimately optimized tissue regeneration.

1. Controlling Porosity. Several techniques for customizing the porosity of biomaterials are illustrated in Fig. 6, A–C. One widely used and inexpensive approach to introduce porosity into cells is particulate leaching (Fig. 6A). In this method, a particulate (e.g., sodium chloride) that is insoluble in a particular solvent (e.g., chloroform) is cast with a polymer (e.g., PLGA) in solvent. After the solvent is evaporated, the material can be placed into an alternative solvent in which the particulate is soluble but the polymer is not (e.g., in the case of sodium chloride for PLGA scaffolds, the material can be placed in water; the sodium chloride dissolves quickly, whereas PLGA is insoluble and even with hydrolysis degrades slowly during the leaching process). With particulate leaching, one challenge is to achieve interconnected pores. Pore interconnectivity is important to ensure that cells are able to navigate the scaffolds to repopulate it and ultimately promote optimal tissue formation. An approach not only to ensure pore connectivity but ultimately define it with considerable precision is the concept of sintering (Fig. 6B) (Murphy et al., 2002; Linnes et al., 2007). In this approach, leachable polymers (e.g., polystyrene) are packed together and heated (to above their glass transition temperature) to allow partial fusion of the beads and provide a template. After cooling, a second polymer is cast around the sintered bead template to backfill the empty regions. The polymer used to fabricate the bead template must selectively be soluble in a solvent (that is, soluble in a solvent in which the cast polymer is not). As described above, the bead template is then selectively dissolved in an appropriate solvent to yield a highly porous scaffold with interconnected pores. These physical dimensions have been observed to play a role in regenerative environments, including dermal healing processes (Fukano et al., 2010; Underwood et al., 2011). Another technique to introduce porosity is phase separation (Fig. 6C) (Nam and Park, 1999). This approach involves dissolution of a polymer into a solvent. The temperature is raised to one such that the polymer is fully solubilized (note that this may not require true heating as many polymers are soluble in certain solvents at room temperature or lower). By cooling (or reducing pressure) the solution can phase separate depending on the concentrations of the solvent and the polymer. This phase separation can achieve solvent-rich regions or polymer-rich regions. Removal of the solvent (e.g., by evaporation) can achieve desirable pore architecture within scaffolds. These can be liquid-liquid phase separations, but it is also possible to introduce gaseous materials to achieve “gas foaming” of the desired pore architecture of the material (Riddle and Mooney, 2004).

2. Electrospinning. The architecture of scaffolds formed by many of the particulate leaching techniques is rudimentary compared with native tissue structure. Electrospinning (Fig. 6D) is a textile fabrication technique that has recently been revived for TE applications. For reviews on this topic see Greiner and Wendorff (2007) or Sill and von Recum (2008). In short, a polymer dissolved in solvent is ejected through a small orifice (typically a needle). An electrical drop is applied between the orifice and collection device (e.g., a flat sheet or spinning mandrel) and fine nanofibers are produced. These electrospun fibers are of critical importance because they can provide topographical cues to cells. This is particularly useful for cells in which cell alignment is important to function such as when creating scaffolds for neural (Wang et al., 2008) or skeletal muscle regeneration (Choi et al., 2008). It is also possible to incorporate nano- or microparticles into these electrospun scaffolds either by using a particle not soluble in the polymer solvent (and spinning the particles with the polymer in solvent) or to add the particles during the spinning process (Guo et al., 2012). Because achieving suitable porosity into electrospun
materials is a challenge, this approach has also been combined with the particulate leaching approach described above (Wright et al., 2010). One key advantage of electrospinning techniques is the ability to incorporate cellular cues at the micro- and nanolevel (e.g., topographical cues).

3. Microfabrication. Microfabrication techniques (Fig. 6E) also allow for incorporation of topographical as well as other key design parameters. Although microfabrication technology is not typically used to produce large three-dimensional scaffolds for implantation, it has numerous applications in allowing better understanding of processes used to direct tissue regeneration. For example, such approaches are also useful for looking at microfluidic effects on cells—an approach difficult to study in vivo or under traditional in vitro cell culture systems. These techniques are used to create so-called “lab/organ-on-a-chip” technologies that allow for the high-throughput testing and screening of pharmacological agents on individual cells (Huh et al., 2010; Ingber and Whitesides, 2012; Neuzil et al., 2012). One approach to microfabrication is to print a photomask, which can be placed over a material surface. During subsequent etching processes, the mask allows control over which areas are, for example, photocross-linked.

4. Three-dimensional Printing. More recently, three-dimensional (3D) printing or solid free form fabrication (Fig. 6F) has been used for scaffold fabrication, with high levels of spatial resolution for applications in bone, nerve, and cardiovascular tissue engineering as a means to determine optimal scaffold parameters (Boland et al., 2006; Mironov et al., 2009; Jakab et al., 2010; Chang et al., 2011; Marga et al., 2012). With this technique, a polymer (in solvent or melt form) is ejected through a small orifice with high precision on a stage with x-y control. A single “layer” is printed and is akin to printing on a piece of paper with a laser printer. However, the “ink” (polymer) itself is three-dimensional and the “paper” is a stage with z-direction control as well. So, individual layers are printed one on top of the other (so-called layer-by-layer approach). By controlling x-, y-, and z-direction resolution, it is possible to fabricate scaffolds with very precise architecture. It is important to note that certain types of materials are more compatible with this approach. Those that are not compatible may be modified with “fillers” to allow printing, but this may have undesirable effects on resultant material properties or biologic responses. The major drawback to all of these fabrication approaches is that the equipment is highly specialized.

C. Functionalizing Biomaterials

Regardless of the method of scaffold fabrication, there are several properties that are important for biomaterials used for regenerative applications. Two general classes of materials used in regenerative medicine applications are natural and synthetic polymers. One advantage often provided by protein-based natural polymer scaffolds is their ability to promote cell attachment and proliferation through their inherent cell-binding motifs. Collagen (RGD), fibrin (RGD), laminin (YIGSR), and keratin (LDV) all contain three to five amino acid sequences that promote cell binding through integrin or other interactions (Fig. 7Ai). Because polysaccharide natural polymer scaffolds and synthetic scaffolds lack these integrin-binding sites, it is not uncommon to covalently graft binding motifs into/onto the material or to mix the material with naturally based polymers that contain binding motifs (Connelly et al., 2011; Šapir et al., 2011; Rafat et al., 2012).

As shown in Fig. 7, it is desirable to have controlled rates of degradation (material aspect) to promote tissue healing (biologic aspect). In the case of natural polymers (Fig. 7Ai), proteolytic sequences are often inherently present. However, such sequences can also be built in to the polymer backbone through a type of synthetic-natural polymer hybrid (David et al., 2012). Traditionally, the more common approach for synthetic materials is to build in hydrolytically cleavable sequences such as ester groups (Fig. 7Aii). There are reports of other synthetic-natural polymer hybrids as well (Xu et al., 2012b) designed to allow functionalization of materials that otherwise lack biologic function.

For both proteolytic and hydrolytically cleavable sequences, the main concept, from a tissue engineering perspective, is to achieve controlled rates of scaffold degradation. However, this property may be useful in not only allowing the scaffold matrix to remain in place for various lengths of time (as desired and designed) but may also be useful in terms of drug delivery. In fact, a number of different materials are known to achieve release of therapeutic agents (small molecule drugs or growth factors) not through diffusion but through degradation of the scaffold material (Saul et al., 2011).

In addition to or instead of hydrolytic or proteolytic “internal triggers,” it may be advantageous to have internal or external triggers such as pH change, temperature change, enzyme, ultrasound or other energy input, or light-triggered degradation (Balmayor et al., 2008; Narayanan et al., 2012; Nelson et al., 2012) (Fig. 7Bi). It is important to note that the opposite of photodegradable linkages (photocross-linkable gels) are an area of active investigation because this allows an in situ solution to gel (sol-gel) transition, potentially allowing more minimally invasive “implantation” or delivery of soft hydrogels to their site of action. Again, these triggering events may promote gel degradation and/or the triggered release of therapeutic agents such as growth factors.

For all of these approaches, a last consideration is the fashion in which the material degrades. Specifically, one
can use materials to achieve bulk degradation of the material, or, for example, it is also possible for degradation to occur only at the surface (Fig. 7B, iii and iv). Clearly, the type of degradation has important implications for drug delivery, cell ingrowth, and the regenerative process.

In summary, the scaffold types described above are commonly used with local delivery of therapeutic agents, certain growth factors, and nucleic acids. Biodegradable polymers and hydrogels (many of which are biodegradable) are the most commonly used scaffolding materials for therapeutic delivery. These systems typically elicit minimal and temporary inflammatory responses, can be tailored for favorable degradation profiles, and can achieve sustained release of therapeutics. Drug release profiles can vary from minutes or hours to years and can therefore be used to “jump start” regenerative processes or provide a sustained impact.

**D. Examples of Biomaterials Applications to Tissue Engineering/Regenerative Medicine Technologies**

In the preceding sections, we mainly described the general desired characteristics of biomaterials for TE/RM applications. Below we provide a few examples of their implementation.

1. **Cardiovascular Disease.** Biomaterials are one of the foundations for preventative and curative approaches to cardiovascular disease. Drug-eluting stents are perhaps the best and most well-known example of drug delivery systems within the context of a biomaterial (although not necessarily in a regenerative sense) (Mani et al., 2007; Wessely, 2010). Other examples of biomaterials in cardiovascular applications include pacemakers (in particular, pacing lead wires and their insulators) (Crossley, 2000; Santerre et al., 2005) and tissue engineered blood vessels (Peck et al., 2012). Biomaterials within a regenerative medicine approach are also one of the most promising technologies in achieving functional recovery of heart tissue after myocardial infarction. For example, derivatives of polyurethanes have garnered attention because they have mechanical properties that mimic those of heart tissue—namely, elasticity and strength. These materials can be synthesized to allow biodegradation as heart tissue regenerates (Fujimoto et al., 2007a,b), including controlled rates of degradation (Hong et al., 2010). Advances to date have focused primarily on the mechanical aspects of these materials. However, it is becoming recognized that these systems are compatible with controlled release of important protective or stimulatory molecules such as IGF-1 and...
hepatocyte growth factor, which may aid in the regenerative process (Nelson et al., 2011), and suggests the next generation of drug delivery in cardiovascular applications beyond drug-eluting stents.

2. Modulation of Stem and Progenitor Cells. Biomaterials are being used increasingly to direct cell differentiation and behavior. By using poly(acrylamide) gels of varying rigidity, it has been shown that mesenchymal stromal cells (MSC; sometimes referred to as mesenchymal stem cells) can be directed to different lineages ranging from neurons to myoblasts to osteoblasts (Norton et al., 2010). The use of topographical cues can also guide cell behavior. For example, the diameter of electrospun pol(ethersulfone) fibers impacts the attachment, spreading, and differentiation fate of neural stem cells (Gratton et al., 2008). Lastly, and of most importance to regenerative pharmacology, chemical signals released from scaffolds can help to direct cell fate. For example, a growth factor cocktail released from fibrin scaffolds promoted the differentiation of neural progenitor cells toward neuronal and oligodendrocyte phenotypes via heparin-binding methods (Willerth et al., 2008). Moreover, nanoparticulate delivery systems that can achieve endocytosis have recently been used for delivery of proteins involved in the Wnt signaling cascades and affecting cellular proliferation and differentiation (or lack thereof) (Shah et al., 2011).

3. Diabetes. Hydrogels based on alginate and other polymers have been in use for nearly 30 years to encapsulate insulin-producing pancreatic islet cells (Lim and Sun, 1980). This approach could circumvent the need (or at least serve as a bridge) for the development of an engineered pancreas. However, traditional biomaterial challenges of protein deposition, foreign body response, and fibrous encapsulation have been barriers to achieving the long-term delivery of insulin required for type I diabetes.

Transdermal delivery systems using microneedle technology are an alternative, noncellular approach to insulin delivery. As shown in Fig. 5L (Davis et al., 2005), these systems present an array of needles on the microscale that can be attached to a reservoir of drug. The purpose of the microneedles is to allow the drug to bypass the stratum corneum layer of the epidermis, thus overcoming a significant diffusion barrier to drug delivery through the dermal route. These technologies have now reached human trials (Gupta et al., 2009). Reduced pain and inflammation have been reported for these types of delivery devices in which insulin is delivered from a reservoir device. Therefore, these systems may provide an alternative to the current standard of subcutaneous delivery within the context of a reservoir-type material that requires less frequent dosing/application.

4. Treatment of Genetic Diseases. The primary application of biomaterials for treatment of genetic diseases is in the development of nonviral gene delivery systems for nucleic acids, primarily DNA. Systemically deliverable nanoscale carriers are the primary focus, and many of the barriers described above (extravasation, cellular uptake, subcellular localization) must be overcome to treat genetic diseases. Poly(ethylenimine) has been considered a benchmark for biomaterial-based nonviral gene delivery because it enables high levels of transfection by promoting endosomal escape of DNA. Systems with reduced levels of toxicity and improved transgene expression are being developed through increased understanding of the role of chemical constituents of the delivery vehicle (Li and Huang, 2007). Peptidic sequences have been conjugated to various nonviral gene delivery systems to improve cellular uptake (Huang et al., 2010), subcellular transport (Kwon et al., 2008; Moseley et al., 2010), and nuclear localization (Jeon et al., 2007; Moore et al., 2009). Based on safety and toxicity profiles, it is conceivable that these approaches may supplant viral technologies for gene delivery (Li and Huang, 2007). Gene therapy has been of interest in the treatment of cancer for many years, for example to restore mutated tumor suppressor genes such as P53 (Fukushima et al., 2007; Gaspar et al., 2011). Muscular dystrophy also has a complex genetic picture (Kornegay et al., 2012), but canine models (Kornegay et al., 2012) are providing important insights into the disease and potential opportunities for gene therapy through both viral and nonviral methods (Poster et al., 2006; Markert et al., 2008; Wang et al., 2012).

5. Scar Reduction and Wound Healing. The use of biologically based products to treat burns and skin conditions has reached the clinic, and products such as AlloDerm, Apligraf, and Dermagraft are approved for marketing. The ability to provide improved threedimensional architecture within the context of a biodegradable system that is readily implanted or sutured would augment existing technologies. Several materials, including hyaluronic acid (Scuderi et al., 2008), chitosan (Boucard et al., 2007; Yang et al., 2010), and alginate (Lee et al., 2009) hydrogels as well as calcium hydroxyapatite (Goldberg et al., 2006), have been used to treat skin conditions, including acne, burns, and melanocytic nevi. In addition, several types of therapeuticsuch as antibiotics for prophylaxis (Kim et al., 2008), growth factors to promote healing (Fujihara et al., 2008), and other compounds (Queen et al., 2007) have been incorporated into biologically active materials (Luo et al., 2010) to promote regeneration. Novel technologies, such as synthetic peptides derived from gap junction proteins, have also been introduced to potentially promote healing after biomaterial implantation (Soder et al., 2009). Clearly, methods to properly control the spatiotemporal presentation of molecules that promote improved wound healing may have significant implications for the treatment of skin diseases.
healing are part of the next generation of treatments made possible by delivery systems incorporated into biomaterials. In the above sections, we attempted to highlight several aspects of biomaterials that are or will play a role in regenerative pharmacology and regenerative medicine. This list is by no means exhaustive, and many of these technologies transcend any one application. Below we describe existing multidisciplinary efforts to establish new experimental models and paradigms for further exploring the potential utility of regenerative pharmacology.

III. Broad Applications of Regenerative Pharmacology

Moving from palliative to curative approaches in complex organs will be a significant challenge. Bladder regeneration and Parkinson’s disease are described below as representative of the biologic complexities that will need to be considered and overcome to achieve curative approaches. First, we will discuss the bladder, because it provides an excellent model system for exploring both the passive (dissecting) and active (directing) components of regenerative pharmacology to future therapeutics (see Table 1). Parkinson’s disease may be amenable to one of several distinct regenerative pharmacology strategies, including drug-, cell-, or gene-based approaches.

A. Bladder Disease

The aim of regenerative medicine/pharmacology is, ideally, to restore normal organ function either by replacing a nonfunctioning organ (end stage disease) or, in the instance when significant viable tissue still remains, improving organ function when it is severely impaired but not yet irreparably damaged/diseased. In the case of the bladder, the physiologic prerequisite is the capacity to store urine at increasing volumes (without increasing intravesical pressure or spontaneous bladder contractions) until complete emptying can be achieved when socially acceptable. Diverse disease etiologies (e.g., neurogenic, congenital, trauma, infections, etc.) compromise the low-pressure, high-volume function (decreased compliance) of the bladder, leading to a number of lower urinary tract symptoms such as urgency, urgency incontinence, frequency, and nocturia. With such diverse etiologies for bladder dysfunctions and a large demand (over 50 million people are estimated to have some type of urinary incontinence), many different classes of drugs have been investigated for symptomatic relief. Antimuscarinic drugs (e.g., oxybutynin, tolterodine, solifenacin, darifenacin) are now the first-line therapy for treatment of detrusor overactivity and the overactive bladder syndrome, but lower urinary tract symptoms can also be treated with, e.g., α-adrenoreceptor (AR) blockers alone or in combination with antimuscarinics or with onobotulinum toxin A (Andersson et al., 2009; Mangera et al., 2011).

In severe cases refractory to pharmacological treatment, high bladder pressures may develop and lead to upper urinary tract deterioration. Patients that display poorly compliant bladders attributable to structural or neurogenic etiologies are at risk for end stage renal disease and are thus candidates for surgical intervention (Reyblat and Ginsberg, 2008). Augmentation cystoplasty has been performed in bladder diseases arising from many different etiologies, including spinal cord injury, myelomeningocele, interstitial cystitis, idiopathic detrusor overactivity, radiation cystitis, multiple sclerosis, and schistosomiasis. Because transplantation of donor bladders is not an available option, attention turned to regenerative medicine/tissue engineering technologies for this organ. In pioneering studies by Atala et al. (2006), bladder neo-organs were constructed by seeding synthetic scaffolds (collagen or collagen/polyglycolic acid composites) with urothelial cells on the inside and smooth muscle cells on the outside and subsequently implanted into subjects with myelomeningocele. Numerous reports suggested that both animal (Liang and Goss, 1963; Oberpenning et al., 1999; Frederiksen et al., 2004; Burmeister et al., 2010) and human (Sisk and Neu, 1939; Liang, 1962; Tucci and Haralambidis, 1963) bladders have significant regenerative potential after removal of a large portion of the organ (subtotal cystectomy; STC).

Novel pharmacological strategies aimed at harnessing the intrinsic regenerative capacity of the bladder will undoubtedly benefit from an improved basic understanding of de novo bladder regeneration. In this regard, animal models can be used to characterize this compelling regenerative phenomenon, opening up new approaches to regenerative pharmacology. In a multidisciplinary effort to characterize bladder regeneration morphologically, physiologically, and pharmacologically, Burmeister et al. (2010) used a trigone-sparing cystectomy (STC) performed in 12-week-old female rats. By 8 weeks post-STC, the bladder had regrown to a normal size via both computed tomography imaging and in vivo cystometric analysis. Moreover, the bladder displayed urothelial, lamina propria, and detrusor muscle layers, and regained normal thickness upon histologic evaluation. However, there was a decrease in bladder smooth muscle contractility when subjected to cholinergic and electrical stimulation. Specifically, 2 weeks post-STC, cholinergic activation resulted in contractile responses that were ~20% of normal, noncystectomized controls. There was some functional recovery of detrusor muscle contraction by 8 weeks post-STC, although maximal steady-state contractions remained low (~37% of normal values). In addition, the presence of a response to electrical field stimulation indicated innervation of newly formed tissue. This
agrees with an earlier study by Frederiksen et al. (2004) in which whole mount staining of acetylcholinesterase revealed the pattern of nerves in newly formed detrusor.

To put this in proper perspective, despite the observed reduction in smooth muscle contractility, a complete functional rodent bladder regeneration response occurs after surgical removal of 70–80% of the bladder (STC). This is a very different phenomenon than the bladder augmentations that are commonly used to study the impact of various stem cells and biomaterials (i.e., tissue engineering) on bladder regrowth after implantation (see below for more details). In summary, within 8 weeks STC rodents have a regenerated bladder that is both structurally and functionally (with respect to micturition and continence) identical to the native bladder that it replaced (Burmeister et al., 2010). This is true with respect to bladder capacity and bladder wall thickness, as well as the presence of all three bladder wall layers: urothelium, muscularis propria, and lamina propria. To our knowledge, bladder regeneration therefore holds a unique position with respect to its regenerative potential, because there is no other mammalian organ capable of this type of regeneration; this includes the liver, which is the most well studied model of regeneration, but which is more accurately referred to as compensatory hyperplasia (Columbano and Shinozuka, 1996).

A more recent follow up study (Peyton et al., 2012) used fluorescent bromodeoxyuridine (BrdU) labeling to quantify the spatiotemporal characteristics of the proliferative response that mediates the functional regeneration observed by Burmeister et al. (2010), as occurs during the critical first week post-STC. In this study, less than 1% of cells in the bladder wall were labeled with BrdU in control bladders under resting conditions (i.e., no damage), but this percentage increased significantly, by 5- to 8-fold, at all time points post-STC. The spatiotemporal characteristics of the proliferative response were characterized by a significantly higher percentage of BrdU-labeled cells within the urothelium at 1 day than in the muscularis propria (MP) and lamina propria (LP). However, a time-dependent shift at 3 and 5 days post-STC revealed significantly fewer BrdU-labeled cells in the MP than in the LP or urothelium. By 7 days, the percentage of BrdU-labeled cells was similar among urothelium, LP, and MP. STC also caused an apparent increase in immunostaining for Shh, Gli-1, and BMP-4. These studies clearly documented that the early stages of functional bladder regeneration are characterized by time-dependent changes in the location of the proliferating cell population in distinct bladder wall layers and, furthermore, demonstrated time-dependent expression of several evolutionarily conserved developmental signaling proteins during this same 1-week period. This report extends our previous observations (Burmeister et al., 2010) and provides further evidence for the rodent bladder as an excellent model for studying novel aspects of mammalian organ regeneration.

The idea that the bladder tissue formed spontaneously after cystectomy is similar to that which remains was proposed by Frederiksen et al. (2004) who described the pharmacology of regenerating bladder. Fifteen weeks after STC in female rats, transverse strips were excised from the bladder body and were exposed to contractile stimuli, taking into account the proximity of the strip to the trigone. The authors used antagonists of muscarinic receptors (scopolamine) and $\alpha_1$ARs (prazosin) as well as a desensitizing agent of P2X1 receptors ($\alpha$, $\beta$-methylene ATP) to examine the contribution of each receptor type to contractions evoked by electrical field stimulations. Additionally, they used agonists of muscarinic receptors (carbachol), $\alpha_1$-ARs (phenylephrine), and purinergic receptors ($\alpha$, $\beta$-methylene ATP) on separate strips. These authors concluded that although the newly formed bladder smooth muscle is well innervated, it has pharmacological properties similar to the supratrigonal tissue from which it had developed.

Clear species-dependent variations in regenerative capacity of the bladder are to be anticipated (Lin et al., 1989). Furthermore, complete, functional, de novo bladder regeneration may not occur in all species and under all experimental conditions, and thus bladder regeneration has also been studied using implantable tissue engineered grafts in the absence and presence of seeded cells. The rationale for this approach is that provision of a scaffold (biomaterial) and the appropriate cells/stem cells to the site of injury or surgical resection would provide for improved/enhanced bladder regeneration and integration with remaining host tissue. Although this is an exciting strategy, unfortunately there is so little still known about “normal” bladder regeneration that there is no a priori rationale for selecting the most appropriate cell and/or biomaterial for maximizing bladder regeneration. In fact, clinical experience with this technology is limited and not yet ready for wide dissemination, pointing to the need for further investigations into the potential applications of tissue engineering/regenerative medicine applications to bladder reconstruction (Atala et al., 2011). As such, examining the pharmacological performance of regenerated bladders after incorporation of a tissue engineered graft is important to evaluate the proximity to normal bladder function and to develop future approaches based on more detailed mechanistic information. Clearly, more work needs to be done to fully characterize the many different aspects of bladder pharmacology during and after regeneration. Determination of the time course and nature of any changes in, for example, receptor...
showed that using this glycosaminoglycan and growth factor in combination produced the best epithelialization, neovascularization, and smooth muscle regeneration 10 weeks after surgery. It is reasonable to assume that delivering many different growth factors, small molecules, or other compounds may aid in regeneration of the urinary bladder and, furthermore, that active regenerative pharmacology will fill the need to explore these possibilities. The use of cells may facilitate delivery of some factors and also may contribute directly to regeneration. Thus, by further exploring the potential applications of novel biomaterials and drug delivery technologies (as illustrated in Figs. 5–7 and described in the text above), it should be feasible to provide more precise spatiotemporal control over growth factor delivery during bladder regeneration, perhaps decreasing the time course and increasing the efficacy of functional recovery.

B. Parkinson’s Disease

One of the main changes in Parkinson’s disease (PD) is a progressive loss of nigrostriatal dopaminergic neurons. The degeneration of the dopaminergic neurons in the substantia nigra pars compacta, which project to the striatum, leads to a reduction of the dopamine (DA) input to this target structure of the nigrostriatal pathway (Dauer and Przedborski, 2003: Lees et al., 2009; Shulman et al., 2011). These modifications in the functional organization of the basal ganglia circuitry lead to the typical motor features of PD (Moore et al., 2005). The precise etiology of PD remains unclear, but pathologic processes such as inflammation, mitochondrial dysfunction, oxidative stress, proapoptotic mechanisms, and accumulation of toxic proteins may play a role (Moore et al., 2005; Shulman et al., 2011).

Treatments of PD focus on symptom relief, neuroprotection, and neurorestoration (Fig. 8). Symptomatic relief can be provided by dopamine substitution therapy and deep-brain stimulation; however, there is an unmet need for the identification of neuroprotective/neurorestorative agents that can modify the progression of the underlying disease processes. Because current treatments aimed at symptomatic relief have numerous limitations (Lees et al., 2009; Hickey and Stacy, 2011), there has been an intense focus on novel therapies, especially those that might provide a definitive cure for the disease. In this regard, stem cell-based therapy offers promise for future treatment of neurodegenerative diseases, including PD. There are two major approaches to stem cell-based therapy for PD, both of which use the cell as the drug delivery vehicle and, therefore, fall under the auspices of regenerative pharmacology. One strategy is aimed at simply replacing the lost cells by transplanting exogenous stem cells, and in this instance clearly the stem cell becomes the direct source of the missing pharmacological agent, that is, dopamine. A second
approach is to use the implanted cells as vectors that contain and secrete neuroprotective agents to preserve the surviving neurons or to induce renewal of axonal sprouting; in this latter case we are using pharmacology more indirectly to maintain, restore, or regenerate the endogenous cellular source of the dopamine.

The effect of various cell sources have been investigated in animal models of PD, as well as in humans, and these include, embryonic stem (ES) cells (Freed et al., 2001; Kim et al., 2002, see Lindvall and Kokaia, 2009), induced pluripotent stem (iPS) cells (Wernig et al., 2008), fetal and adult brain-derived neural stem cells (NSC; Hermann et al., 2006; Schwarz et al., 2006), mesenchymal stem cells (MSC; Hellmann et al., 2006; Cova et al., 2010), and amniotic fluid stem (AFS) cells (Donaldson et al., 2009). In an interesting application of regenerative pharmacology, neural progenitors generated from pluripotent stem cells in culture were induced to give rise to dopaminergic neurons, which hold therapeutic potential for PD. Studer and colleagues recently reported that the application of CHIR99021, a potent GSK-3β inhibitor known to strongly activate canonical WNT signaling, to certain midbrain precursors cells derived from human pluripotent stem cells strongly promotes differentiation to midbrain dopaminergic neurons, precisely the class that degenerates in PD. These neurons survived well and functioned appropriately when grafted into the brain in several animal models, most notably Parkinsonian nonhuman primates (Kriks et al., 2011). Clinical trials using human mesencephalic tissue provided the proof-of-principle for cell replacement in PD patients but also showed clinical limitations (Koch et al., 2009; Lindvall and Kokaia, 2009, 2010; Meyer et al., 2010).

Gene therapy provides another strategy to restore the ability of the brain to deliver dopamine and...
other agents to the striatum to provide both symptomatic benefit and possibly neuroprotection/neuroregeneration. For this application, genes have thus far been packaged into viral vectors and injected into the brain with the goal of either delivering genes for DA-synthesizing enzymes to the striatum or providing neuroprotection to block or slow ongoing degenerative processes by providing genes for growth factors, antioxidant molecules, or antiapoptotic substances (Muramatsu et al., 2003). Adeno-associated virus (AAV) has been the most commonly used vector because of its ease of use and safety profile (Ozawa et al., 2000; Bankiewicz et al., 2006). This approach is currently being tested in a number of Phase I and II clinical trials (Hickey and Stacy, 2011). Details of some relevant examples are provided below, and furthermore, these strategies and their corresponding pharmacological details are summarized in Fig. 9.

1. Enhancing Striatal Dopamine Effects. Local production of dopamine in the striatum can be achieved by inducing the expression of enzymes involved in the biosynthetic pathway for dopamine. As pointed out by Hickey and Stacy (2011), the potential benefits are compelling: the ability for selective basal ganglia stimulation by bypassing the need for systemic medications, the avoidance of undesirable side effects induced by indiscriminate dopamine activation, and even the possibility for individualized treatment regimens. For example, one strategy was indicated by in vitro experiments showing that triple transduction with separate AAV vectors expressing tyrosine hydroxylase, L-amino acid decarboxylase (AADC), and GTP cyclohydrolase 1, respectively, increased dopamine production (Shen et al., 2000; see Fig. 8). This study demonstrated that stereotaxic intrastratial injection of these factors in 6-hydroxydopamine (6-OHDA)-lesioned rats produced sustained behavioral improvement for up to 12 months. Consistent with the rodent study, the same group confirmed and extended their original observations to document that the same triple AAV transduction of striatal cells with dopamine-synthesizing enzymes also produced behavior recovery in a primate model of PD (i.e., 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine; MPTP lesion) for up to 10 years who had progressive levodopa-responsive Parkinson’s disease and showed a significant improvement from baseline in Unified Parkinson’s Disease Rating Scale scores compared with the sham-treated group over the 6-month course of the study. Certainly, some of the newer biomaterial strategies for nonviral gene delivery, as described above, may ultimately be of benefit to the gene-based treatment of Parkinson’s disease.
PD in the future, as may some of the stem cell-based regenerative pharmacology approaches described below.

**IV. Regenerative Pharmacology and Stem Cells**

Modulation of the behavior of stem and progenitor cells plays a central role in TE/RM technologies, and is a major target of regenerative pharmacology (Table 3). These cells are able to undergo mitotic division, sometimes very extensively, and also to give rise to more specialized progeny. The defining characteristic of stem cells is the continued maintenance of a pool of unaltered daughter cells that retain this dual potential for proliferation and differentiation, a property known as self-renewal. Stem cells from certain sources, such as the inner cell mass of early stage embryos (ES cells),...
can give rise to any specialized cell type found in normal development and are designated pluripotent. Other stem cells and progenitors, notably those present throughout life in many adult organs, appear restricted to limited sets of cell lineages. Such cells are termed multipotent or, if constrained to a single fate, unipotent. The restriction of the developmental fate of cells to a particular lineage is called commitment and precedes the acquisition of overt specialized features during cell differentiation (Smith, 2006; Sheridan and Harris, 2009). Whether by recapitulation of normal development or through novel bioengineered pathways for tissue formation and organogenesis (Ingber and Levin, 2007), therapies based on regeneration almost inevitably must engage stem/progenitor cells (Nirmalanandhan and Sittampalam, 2009). Although still in its earliest phase, an era of clinical testing of stem cell therapies has begun (Trounson et al., 2011).

Pharmacology potentially can enhance stem and progenitor cell-mediated regenerative therapies through at least five distinct strategies: A) stem/progenitor cell expansion in culture (Fig. 9A); mobilization of endogenous stem/progenitor cells in situ (Fig. 9B); lineage-specific differentiation of stem/progenitor cells (Fig. 9C); differentiation from pluripotent stem cells (Fig. 9D); and reprogramming to generate pluripotent stem cells and lineage-restricted cells (Fig. 9E). We discuss selected examples of each of these approaches below. In addition, patient-derived or genetically engineered stem cells serve as novel sources for cell-based systems to mimic features of various human illnesses. Such “disease in a dish” models are emerging as powerful tools for drug discovery in regenerative medicine and, more broadly, in essentially any therapeutic arena (Gage, 2010; Walker, 2010). We also introduce this promising area of basic research, which opens up numerous new opportunities for pharmacology.

Growth factors, hormones, and other biologic signaling molecules may exert pharmacological effects to control the expansion, commitment, or differentiation of stem/progenitor cells and progenitors. These molecules can modulate the proliferation, commitment, and differentiation of stem/progenitor cells and may be classified into five distinct strategies: A) stem/progenitor cell expansion in culture (Fig. 9A); mobilization of endogenous stem/progenitor cells in situ (Fig. 9B); lineage-specific differentiation of stem/progenitor cells (Fig. 9C); differentiation from pluripotent stem cells (Fig. 9D); and reprogramming to generate pluripotent stem cells and lineage-restricted cells (Fig. 9E). We discuss selected examples of each of these approaches below. In addition, patient-derived or genetically engineered stem cells serve as novel sources for cell-based systems to mimic features of various human illnesses. Such “disease in a dish” models are emerging as powerful tools for drug discovery in regenerative medicine and, more broadly, in essentially any therapeutic arena (Gage, 2010; Walker, 2010). We also introduce this promising area of basic research, which opens up numerous new opportunities for pharmacology.

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of stem and progenitor cells. As noted earlier in this article, polymeric biomaterials also can exert pharmacological actions on these cells, both as vehicles for delivery of regulatory molecules and as physical substrates that provide functional cues through their mechanical properties (Furth et al., 2007; Keatch et al., 2012). Increasingly, stem cell biology also intersects with small molecule chemistry to expand the tool kit for regenerative pharmacology (Xu et al., 2008).

A. Stem/Progenitor Cell Expansion in Culture

A clinical experiment at Boston’s Peter Bent Brigham Hospital in 1980 marked a pivotal step toward the development of the first cell-based regenerative medicine products. Expansion in vitro of autologous keratinocytes from small remaining areas of skin enabled the rescue of two children who had suffered extensive third-degree burns (Green, 2008). In time this led to a commercialized product (Epicel: cultured epidermal autograft) (De Bie, 2007). Although fibroblasts had long been cultured successfully in the laboratory, the successful expansion of human keratinocytes, the essential covering cells of the skin, was new. Green and colleagues used epidermal growth factor (EGF) along with factors provided by “helper” murine fibroblasts (irradiated to prevent their own growth) to maintain long-term keratinocyte cultures. On the basis of his pioneering studies of transplantation of such expanded epithelial cell populations from skin or cornea, Green concluded that an essential feature for regenerative cell therapy was the presence in the graft of an adequate pool of stem cells (De Luca et al., 2006).

Various additional cell-based therapies now depend on culture conditions that enable substantial multiplication of donor cells. For example, expanded populations of autologous chondrocytes have received regulatory approval for use in cartilage repair (Manfredini et al., 2007). In some cases, animal or human serum may be used as a crude source of the necessary hormones and growth factors, without pharmacological analysis. However, for many cell types, as exemplified by keratinocytes, the use of specific growth factors in a defined, serum-free medium is essential to achieve proliferation of the stem/progenitor cells without premature terminal differentiation. In addition, the elimination of serum from expansion media mitigates regulatory concerns about animal-derived components and potential pathogens.

The best-characterized stem cells used to date in medicine are those of the blood-forming system. These hematopoietic stem cells (HSC) serve as precursors to all the specialized types of blood cells, including erythrocytes, megakaryocytes, and the components of the innate and adaptive immune systems (Bryder et al., 2006; Seita and Weissman, 2010). The presence of HSC and downstream progenitors derived from them in bone marrow and umbilical cord blood provides the basis for stem cell transplantation in the treatment of disorders such as leukemia and aplastic anemia. The cells actually capable of long-term self-renewal and full reconstitution of the hematopoietic system are rare, comprising less than 0.02% of nucleated cells in the bone marrow (Kent et al., 2009; Bonnefoix and Callanan, 2010). Although the HSC displays a remarkable capacity for proliferation in vivo (Cao et al., 2004; Gazit et al., 2008; Notta et al., 2011), only limited success has been achieved in culturing these cells without loss of “stemness.” Recent reviews survey efforts to employ a wide variety of growth factors and cytokines, as well as small molecules, to enable extensive expansion without excessive commitment to differentiation and concomitant loss of self-renewal capacity (Watts et al., 2011; Walasek et al., 2012; Furth et al., 2013). Some agents have been reported to induce up to 1,000-fold expansion of cells expressing CD34, a surface marker characteristic of many hematopoietic stem and progenitor cells. However, the most immature HSC, i.e., those capable of long-term reconstitution of the blood system, still become relatively depleted. A defined culture medium containing a cocktail of protein factors, including angiopoietin-like 5 does enable ~ 20-fold expansion of long-term HSC from human cord blood (Drake et al., 2011). A small molecule purine derivative designated StemRegenin 1 (SR1), which inhibits signaling through a ligand-dependent transcription factor associated with responses to toxic xenobiotics such as dioxin, likewise drives HSC proliferation to a comparable extent (Boitano et al., 2010). Sophisticated automated culturing systems that reduce the exposure of stem cells to inhibitory paracrine signals from more differentiated progeny cells also facilitate expansion of long-term human HSC (Csaszar et al., 2012). Taken together, the studies of the hematopoietic system reveal the challenges faced by regenerative pharmacology in seeking to regulate the complex balance between maintenance of stem cell identity versus commitment to progenitors that lose self-renewal capacity and give rise to large numbers of postmitotic, terminally differentiated cells.

By contrast to HSC, some classes of stem cells present in fetal and adult tissues can proliferate readily in vitro and retain their essential characteristics. For example, in the mid-1990s, Weiss et al. (1996) made the remarkable discovery that stem cells could be isolated from the mammalian central nervous system, not only during fetal development, but also throughout adult life. These neural stem cells (NSC) expand in aggregated spheres when cultured in serum-free, hormonally defined media containing EGF, sometimes further supplemented with fibroblast growth factor 2 (FGF-2). [Note that the name given to a growth factor often is based on the first cell type
shown to be responsive, but does not necessarily convey the full range of the factor’s biologic activities.\] Multipotent NSC can give rise to neurons, glia, and oligodendrocytes (Gage, 2000; Bergstrom and Forsberg-Nilsson, 2012). The neural stem cells enable life-long neurogenesis in some parts of the brain. Eventually, they may provide a basis to develop curative therapies for devastating neurodegenerative diseases. The use of defined media and bioreactors facilitates the uniform production of clinical grade cell populations (Baghbaderani et al., 2011).

First-in-human clinical studies of neural stem/progenitor cells have begun. For example, StemCells Inc. has initiated four trials of its HuCNS-SC product, although its first program in neuronal ceroid lipofuscinosis (Batten’s disease, a lysosomal storage disorder) was terminated because of insufficient patient enrollment. Possible areas of application for neural stem/progenitors include spinal cord injury, amyotrophic lateral sclerosis, Parkinson’s disease, Alzheimer’s disease, age-related macular degeneration, and cerebral palsy (Feng and Gao, 2012; Glass et al., 2012; Luan et al., 2012; Politis and Lindvall, 2012; Riley et al., 2012; Sandner et al., 2012).

**B. Mobilization of Endogenous Stem Cells**

In principle, it would be advantageous to find ways to effect regenerative responses of stem and progenitor cells in situ, without the need to isolate and transplant cells. However, pharmacological manipulation of stem cell proliferation in the body seems a tall order. Many of the growth factors or cytokines that have been found to promote stem cell proliferation and self-renewal act on multiple cell lineages, both during development and in normal tissue turnover. These include stem cell factor, which is the ligand for the c-Kit receptor, and members of the Wnt, Notch, and Hedgehog families of developmentally important signaling factors. Nonetheless, factors that have multiple cellular targets sometimes can exert relatively specific pharmacological effects. For example, pleiotrophin, a heparin-binding cytokine that acts on the receptor protein tyrosine phosphatase-\(\beta^c\) has a wide array of cellular targets: it enhances outgrowth of nerve fibers and can regulate cellular proliferation and/or differentiation of ES cells and in representative cell types derived from each of the three embryonic germ layers (Deuel et al., 2002). Even so, administration of recombinant pleiotrophin selectively stimulates the recovery of bone marrow in irradiated mice, accompanied by a 20-fold increase in long-term HSC (Himburg et al., 2010). Thus, it is possible that a pool of specific stem cells (e.g., the HSC) depleted by an insult such as radiation or chemotherapy could be restored by administration of a broadly acting drug without perturbing stem/progenitor cells in unaffected compartments of the body, which may be held in check by other homeostatic mechanisms.

Targeted delivery of a pleiotrophic factor may accomplish a similar end. The Wnt pathway plays a major role in regulating proliferation and differentiation of multiple categories of stem/progenitor cells, including those involved in bone formation. By packaging purified recombinant Wnt3a protein in liposomal vesicles that could be administered locally to skeletal defects in mice, investigators were able to promote more rapid bone regeneration through enhanced proliferation of skeletal progenitors and accelerated differentiation into osteoblasts (Minehar et al., 2010).

Mobilization of stem cells from niches and their recruitment to damaged tissue sites also may support regeneration. Several cytokines, including granulocyte colony-stimulating factor (G-CSF; approved by the FDA under the names filgrastim and lenograstim) (Frampton et al., 1994), stimulate HSC to migrate from quiescent niches in the bone marrow and enter into the circulation. This has practical value in the collection of HSC from transplant donors and probably also contributes to recovery of the hematopoietic system from various insults. A small molecule named plerixafor (Mozobil; AMD3100) mobilizes HSC through another mechanism (Brave et al., 2010). It interferes with the action of a chemokine, stromal cell-derived factor-1, at its receptor CXCR4. Although the stromal cell-derived factor-1/CXCR4 axis is widely used to control cell migration in development, inflammation, and other contexts, the pharmacological activity of Plerixafor is sufficiently specific to displace HSC from bone marrow niches without causing unacceptable toxicities. The drug has received marketing approval from both the FDA and the European Committee for Medicinal Products for Human Use.

Regenerative pharmacology approaches can be used to selectively concentrate stem and progenitor cells at sites in need of repair. For stem cells that, unlike HSC, rarely enter into the circulation, transplantation may be best accomplished through grafting of cells embedded in an appropriate biomaterial. The scaffold could be augmented with growth factors and additional extracellular matrix components to help drive proliferation and/or differentiation (Turner et al., 2010). Validation of this approach came from a demonstration that delivery in hyaluronan hydrogels greatly improves retention of HpSC (hepatocyte stem cell) in the liver, promoting more efficient engraftment, proliferation, and vascularization (Turner et al., 2013) (Figs. 5–7). Grafting cells in a hydrogel matrix thus enhances tissue repair and decreases the risk of cells being carried through the bloodstream to become trapped and possibly to survive and even proliferate at distant sites.

A recent report showcases an elegant approach to promote bone regeneration by targeting stem/progenitors to an injured area. MSC are well established as precursors to bone-forming osteoblasts. The
number of mesenchymal stem/progenitors in the bone marrow goes down with aging, leading to decreased osteogenesis and potentially contributing to the development of osteoporosis. However, transplantation of exogenous MSC neither leads to long-term engraftment of the marrow nor restores the cellularity of the correct part of the bone structure. To concentrate potentially regenerative cells at the appropriate location, Lane, Lam, and colleagues designed a compound to bridge between a membrane protein present on MSC and the bone surface (Guan et al., 2012). From a peptidomimetic library they selected a high-affinity ligand against integrin α4β1, designated LLP2A, and coupled it to alendronate (Ale), a bone-binding bisphosphonate. The resulting bifunctional reagent (LLP2A-Ale) was assessed for activity in immunodeficient mice grafted with human MSC. The compound stimulated homing and retention of the MSC at the bone surface, where they contributed to increased trabecular bone formation and bone mass over many weeks. Although Ale is used to treat osteoporosis, the bisphosphonate alone, without the targeting peptide, was not effective at the low doses used in these studies. The conjugated compound LLP2A-Ale also could direct endogenous MSC to the bone surface, as judged by stimulation of osteoblast activities. The drug prevented osteopenia (bone loss) with aging after peak bone mineral density had been achieved. Moreover, it curtailed the drastic loss of bone that occurs because of estrogen deprivation after ovariecotomy of female mice. The data thus support a novel paradigm for pharmacological direction of stem cells to a target tissue where they can engage in repair and regeneration.

C. Lineage-Specific Differentiation of Stem and Progenitor Cells

Pharmacological administration of individual cytokines or growth factors can drive the production of specific differentiated cell types from multipotent or lineage-restricted stem and progenitor cell populations. The accelerated production of red blood cells after administration of recombinant human erythropoietin (EPO; epoetin) (Faulds and Sorkin, 1989) and of infection-fighting granulocytes after injection of recombinant G-CSF (Frampton et al., 1994) can be considered the first commercially significant examples of regenerative pharmacology. EPO primarily targets lineage-specific erythrocyte progenitors to drive rapid increases in the number of circulating red blood cells to combat anemia or, notoriously, as a performance-enhancing drug for endurance athletes. G-CSF targets committed granulocyte progenitors and speeds the recovery from neutropenia, for example after treatment of cancer patients with cytotoxic chemotherapeutic agents. In both cases the mature blood cells have a limited life span; human erythrocytes survive approximately four months and nonactivated granulocytes less than 1 week. EPO and G-CSF do not permanently alter the balance of stem and progenitor cell compartments in the bone marrow, so their effects are transient.

Pharmacological use of growth factors and cytokines on stem/progenitor cell populations also can have more long-lasting outcomes. United States and European regulatory agencies approved two members of a large family of bone morphogenetic proteins (BMPs) for several orthopedic applications. These proteins induce the proliferation and differentiation of bone precursor cells, likely including MSC. Recombinant human BMP-7, also known as osteogenic protein-1 or eptotermin alfa, delivered via a sponge made of type I collagen, received FDA approval under a Humanitarian Device Exemption for treatment of long-bone fractures that did not heal spontaneously (Friedlaender et al., 2001). The same cytokine, formulated as a putty with collagen and carboxymethylcellulose, is approved for revision spinal fusions. Similarly, a collagen sponge carrying recombinant human BMP-2 (INFUSE Bone Graft, Medtronic Spinal and Biologics, Memphis, TN) has been approved under an Humanitarian Device Exemption for certain fracture repair, bone grafting, and spinal fusion procedures (McKay et al., 2007).

In the United States alone, the market for BMP devices is in the range of $1 billion annually (Burks and Nair, 2010). However, recent critical reviews point out risks of significant and even catastrophic complications that can result from ectopic bone production attendant to BMP therapy (Carragee et al., 2011; Mirza, 2011). Therefore, finding improved agents to promote bone repair remains an important goal for regenerative pharmacology. Simply improving the delivery of BMPs might have significant benefit, as leakage of excess cytokine can result from overloading of collagen sponges. Genetically modified cells represent one possible system to provide BMP locally (Kimelman-Bleich et al., 2011). Osteoconductive materials, permissive for bone development, might synergize with BMPs and potentially reduce side effects. A promising new osteoconductive scaffold material was identified from a combinatorial library of biodegradable poly(β-amino ester)s; screening of a small molecule library revealed a number of "hits" that either promoted or inhibited osteogenic differentiation of MSC (Brey et al., 2010; Brey et al., 2011).

Other protein factors may synergize with or eventually replace BMP-7 or BMP-2 for many orthopedic applications. For example, the growth factor NEL-like molecule-1 (NELL1) acts on osteochondral lineage progenitor cells to promote bone regeneration, while suppressing the competing differentiation pathway to fat cells (adipocytes) (Zhang et al., 2010; Zou et al., 2011; James et al., 2012; Shen et al., 2012b). NELL1, alone or in combination with a BMP, enhances bone formation in a number of in vivo repair models (Li et al., 2011; Siu et al., 2011; Zhang et al., 2011; Zhu et al., 2011).
Although it may appear that there is significant redundancy among the sets of factors that can induce bone regeneration, the optimal choice of pharmacologic agent for a given application may depend on detailed understanding of the physiology of the target tissue and its repair. BMP-2 rapidly and irreversibly induces formation of both cartilage and new bone, but it also can cause unwanted ectopic bone formation (Noel et al., 2004). However, the broadly acting factor Wnt3a, when packaged in liposomes for local delivery, induces a bone-specific pattern of regeneration without ectopic bone formation (Minear et al., 2010). The effect appears consistent with enhancement of normal repair activities carried out by stem/progenitor cells migrating to areas of injury in the periosteum and to the bone marrow cavity. Thus, agents that promote Wnt signaling may have superior characteristics for regenerative pharmacology in orthopedics. Candidate drugs include not only recombinant Wnt3a protein, but also small molecule agonists of Wnt signaling (Gwak et al., 2012) and antagonists of endogenous inhibitors of the Wnt pathway (Agholme and Aspenberg, 2011).

D. Differentiation from Pluripotent Stem Cells

Despite the presence of lineage-restricted stem/progenitor cells in most or all of the body's tissues, in many circumstances they are not readily accessible and/or may be difficult to propagate in culture. Since the first isolation of human ES cells, 17 years after the initial description of mouse ES cells, these have been touted as a possible source for replacement therapy of any specialized cell type (Odorico et al., 2001; Thomson et al., 1998). The generation of induced pluripotent stem (iPS) cells by genetic reprogramming of somatic cells (Takahashi and Yamanaka, 2006), for which Yamanaka shared the 2012 Nobel Prize in Physiology or Medicine, raised the ante by potentially enabling production of autologous specialized cells of any lineage for any human being (Yu et al., 2007; Nakagawa et al., 2008; Park et al., 2008; Okita and Yamanaka, 2011). A long-term goal would be to develop curative therapies for conditions in which crucial cell populations have been lost through injury or disease, without a need for immunosuppression. Potential clinical targets include such major disease areas as heart failure, liver failure, kidney failure, neurodegeneration, osteoporosis, and diabetes.

There is abundant evidence that ES and iPS cells can be induced to give rise to a wide variety of specialized cells types, and already a significant body of pharmacological data helps to define molecular mechanisms that contribute to differentiation toward various lineages (Atkinson et al., 2013). Nevertheless, achieving efficient, highly specific differentiation of pluripotent stem cells and successfully grafting the resulting stem cell-derived progeny cells to treat human disease pose formidable obstacles of scale, economics, and safety. Detailed discussion of these problems lies beyond the scope of this review. However, at least two other general points appear particularly salient to pharmacological approaches.

First, differentiation from an initial pluripotent stem cell to a desired differentiated cell type in almost all instances will require a series of steps corresponding to distinct milestones in the complex normal morphogenesis of mature tissues and organs. Current strategies generally are based on known developmental sequences. They begin by inducing the pluripotent cells to restrict to one of the three germ layers (ectoderm, mesoderm, and endoderm) that segregate at the gastrulation phase of embryogenesis (Murry and Keller, 2008). In most cases, investigators have next focused on the empirical identification, guided by knowledge of embryonic development, of growth factor/cytokine mixtures required to promote each of the sequential stages toward the desired mature cell type. However, for many cell types only inefficient and/or incomplete differentiation has been achieved to date. Processes that occur over many weeks in the tightly regulated environment of the developing embryo and fetus often cannot be perfectly replicated with cell lines in laboratory culture. 3D methodologies and chemical biology offer powerful new tools that should enhance our ability to derive therapeutic cell types and useful in vitro disease models from stem cells.

The application of bioengineering principles, in particular the development of suspension culture methodologies for the expansion and differentiation of pluripotent stem cells and their derivatives, offers significant opportunities to optimize the production of specialized cell types. For example, a recent report documents an 18-fold increase in definitive endoderm yield using optimized growth factor cocktails and a suspension bioreactor system (Ungrin et al., 2012). However, additional hurdles still must be overcome to generate differentiated cells at high purity and yield.

A second major concern lies in the relative immaturity of differentiated progeny derived from ES or iPS cells. For example, to generate β-like cells able to maintain normal blood sugar levels and respond to metabolic challenges in vivo, the Baetge team found it necessary to allow human ES cell-derived pancreatic progenitors to mature for more than 3 months after implantation into immune-deficient mice (Kroon et al., 2008). Similar observations come from critical assessment of hepatocytes derived from human pluripotent stem cells (Snykers et al., 2009; Delaforest et al., 2011). Careful comparison of hepatocyte-like cells obtained from human ES cells with authentic adult liver progenitor cells revealed significant differences (Funakoshi et al., 2011). After 3 weeks of differentiation in culture, the ES-derived cells lacked certain key adult features and more nearly resembled human fetal hepatocytes at least 20 weeks of gestation.
Failure to achieve a normal, adult phenotype by differentiation of pluripotent stem cells in culture appears to be a general problem for many cell fates. A recent study used global transcriptional profiling to assess differentiation to multiple lineages, including representatives of each of the three germ layers (Patterson et al., 2012). In no case was the progeny from ES cells or iPS cells identical to mature tissue-derived cells. Of special concern, the differentiated cells continued to express a subset of genes associated with early embryonic development (e.g., LIN28A, LIN28B, and DPPA4). Overall, they showed characteristics of cells present within the first 6 weeks of human prenatal development. Although the results reinforce the value of ES and iPS cells in understanding human embryology, the immaturity of differentiated progeny derived from pluripotent stem cells may limit their utility in modeling adult human diseases and producing safe cells for regenerative medicine applications.

1. Pharmacological Tools for Differentiation from Pluripotent Stem Cells. Further application of 3D culture technologies and tissue-specific matrix components, in addition to soluble growth factors and cytokines, have the potential to improve in vitro differentiation from both pluripotent stem cells and lineage-restricted stem/progenitor cells present in fetal and adult tissues (Ott et al., 2008; Baptista et al., 2011; Bratt-Leal et al., 2011; Kraehenbuehl et al., 2011; Leipzig et al., 2011; Mari-Buye and Semino, 2011; Spence et al., 2011; Wang et al., 2011c; Azarin et al., 2012; Cardinale et al., 2012; Purpura et al., 2012). In this arena there exists great potential for regenerative pharmacology to explore synergies among peptidic signaling molecules (growth factors, cytokines), ECM components and synthetic biomaterials, and small molecule pharmaceutical compounds.

Modulation of stem cell survival, growth, and differentiation by small molecules has been validated in concept and represents an area of intense research activity (Ding and Schultz, 2004; Barbaric et al., 2010; Li and Ding, 2010; Zhu et al., 2010; Yuan et al., 2011a; Choi and Nam, 2012; Li et al., 2012; Atkinson et al., 2013). The use of phenotypic screens to identify compounds that can serve as probes for the identification of specific cellular functions associated with differentiation can be viewed as an important application of “chemical genetics” (Sachinidis et al., 2008). Brief mention was already made of applications to Parkinson’s disease. Although there are examples of differentiation modulators for cell types from each of the three embryonic germ layers, for the focused aim of this review, below we provide a more detailed example highlighting potential cardiac applications.

2. Cardiac Lineage as an Example of Regenerative Pharmacology for Guiding Phenotypic Differentiation. In vertebrates the first functional organ to develop is the heart. Turnover of adult cardiac muscle is extremely limited. In light of the high prevalence of heart disease, replacement of damaged or dead cardiomyocytes stands as an enormous challenge and opportunity for regenerative medicine (Gersh et al., 2009; Bartunek et al., 2010). Human cardiomyocytes also can serve as tools for toxicity testing and drug discovery. Generation of beating cardiomyocytes from pluripotent stem cells has been achieved through several approaches that have been reviewed recently (Rajala et al., 2011; Bernstein, 2012; Zwi-Dantsis and Gepstein, 2012). As with other cell types, the best results for directed differentiation have been achieved using a sequential approach patterned after normal development (Kattman et al., 2011). In this case the first step is induction of mesoderm, followed by progression to cardiac mesoderm, cardiac/cardiovascular progenitors, and cardiomyocytes. The differentiated cells can beat but have a relatively immature phenotype. Key mechanistic steps in the formation of cardiogenic mesoderm and the differentiation of committed progenitors to cardiomyocytes are still poorly understood. Moreover, authentic cardiovascular progenitors must actually give rise to a number of subtypes of cells corresponding to the first heart field, yielding left ventricular cardiomyocytes, and the second heart field, yielding smooth muscle and endothelial cells, sinoatrial nodal and atrioventral nodal cells (involved in pacemaker activity and controlled beating), atrial cardiomyocytes, and right ventricular cardiomyocytes. Recapitulating the full cardiac lineage tree in a controlled, directed manner remains a goal for future studies. Isolation of iPS cell-derived cardiovascular progenitor cells with potential to differentiate to endothelial, smooth muscle, and cardiomyocyte lineages represents an encouraging step in this direction (Nsair et al., 2012). Key steps in lineage specification appear to be determined by surprisingly subtle quantitative changes in signaling by TGF-β family members (Activin/Nodal and BMPs), which poses a significant challenge for effective pharmacology (Kattman et al., 2011). Assessment of temporal changes in chromatin structure provides a powerful new method to identify major regulators of cardiac development and may facilitate the identification of protein factors and small molecules to modulate differentiation (Paige et al., 2012).

Several practical methods have been devised to purify cardiomyocytes generated from pluripotent stem cells. For example, the high level of mitochondrial staining with the dye tetramethylrhodamine methyl ester perchlorate, facilitates enrichment of cardiomyocytes to >99% purity by fluorescence-activated cell sorting (Hattori et al., 2010). A different strategy rests on a genetic trick—linking expression of a drug resistance gene to a promoter expressed exclusively in the cardiac lineage to enable elimination of all undifferentiated stem cells and any cells
that have committed to noncardiac fates (Zandstra et al., 2003).

Pharmacological studies have revealed both protein factors and small molecules capable of modulating cardiomyogenesis from pluripotent stem cells. Among members of known families, fibroblast growth factor 10 ([FGF-10—a misleading name because, like FGF-7, also known as keratinocyte growth factor, this factor is mitogenic for keratinocytes but not for fibroblasts) appears crucial for normal cardiomyocyte differentiation from ES cells, based on inhibition studies with a neutralizing monoclonal antibody against FGF-10 and the use of inhibitors of the ligand’s cognate receptor protein-tyrosine kinase, FGF receptor-2 (Chan et al., 2010). Screening of a combinatorial compound library led to the discovery of small molecules that could induce cardiomyocyte differentiation from pluripotent stem cells (Wu et al., 2004). Four related diaminopyrimidine compounds, designated Cardiogenol A–D, showed this activity on a mouse embryonal carcinoma cell line, with the most potent having an EC₅₀ of 0.1 µM. Cardiogenol C also induces a cardiomyocyte-like phenotype in multipotent stem cells derived from murine hair follicles (Yau et al., 2011). Studies in that system suggested that the compound enhances Wnt/β-catenin signaling, possibly through downregulation of Kremen1, a receptor for Dickkopf protein, which is known to negatively modulate the Wnt pathway. By contrast, in the ES cell system the administration of an inhibitor of Wnt signaling, XAV939, immediately after the generation of mesoderm progenitor cells strongly enhanced the production of cardiomyocytes (Wang et al., 2011a). XAV939 functions as an inhibitor of Tankyrase, thereby stabilizing Axin and inhibiting Wnt signaling. The apparently contradictory observations on the effects of Cardiogenol C and XAV939 may be reconciled by the precise timing of their administration. Murry and colleagues reported that Wnt/β-catenin signaling has a biphasic role in cardiac lineage differentiation from human pluripotent stem cells; at an early stage it promotes mesoderm induction, while at a later point it limits the production of cardiomyocytes from committed progenitors (Paige et al., 2010).

Yet another library screen led to the discovery of a distinct small molecule that enhanced cardiomyogenesis from ES cells by approximately 3-fold (Shen et al., 2012a). The “hit,” designated compound 62 (a 2,6-disubstituted 4-anilinoquinazoline derivative), potently inhibits the tyrosine kinase activity of the EGF receptor (IC₅₀ = 101 nM). However, other EGF receptor inhibitors do not show the same enhancement of cardiac differentiation, suggesting that compound 62 may have another as yet unidentified molecular target that is more specifically associated with the development of the heart.

As noted at several points in this article, regenerative pharmacology can encompass ECM components and other elements of the 3D microenvironment that may complement small molecule signaling modulators. Characterization of a niche for cardiovascular progenitor cells in developing mammalian hearts led investigators to formulate 3D cell culture inserts by electrospinning (see Fig. 6D) and in some experiments to coat them with collagen type IV (Col IV). They observed that Col IV indeed enhanced the expansion of the cardiac progenitor population, most notably when used in the 3D context (Schenke-Layland et al., 2011). In this case the addition of an inhibitor of Wnt/β-catenin signaling (IQ-1) (Miyabayashi et al., 2007) further facilitated expansion of the cardiac progenitors, with the effect additive to that of Col IV (Schenke-Layland et al., 2011). In a related approach, culturing of pluripotent stem cells in a sandwich configuration between layers of Matrigel together with sequential addition of known growth factors (activin A, BMP-4, FGF-2), promoted robust differentiation to cardiomyocytes (Zhang et al., 2012). Matrigel is a commercially available ECM, extracted from a transplantable mouse tumor line, that contains high levels of Col IV and laminin but also a number of less well-defined components including some growth factors (Vukicevic et al., 1992; Kleinman and Martin, 2005). Although much work remains to be done, there is clear evidence that such an approach may eventually yield a variety of novel therapeutics for cardiac disease/dysfunction.

E. Reprogramming to Generate Pluripotent Stem Cells and Lineage-Restricted Cells

Cell differentiation to a great extent reflects epigenetic control—that is, heritable changes in gene expression without alterations in DNA sequence (Ng and Gurdon, 2008). Although the underlying mechanisms remain incompletely understood, packaging of DNA into chromatin and modifications such as DNA methylation clearly play central roles. The understanding that the nucleus of a differentiated vertebrate cell could be reprogrammed to a ground state similar to that of cells in the earliest stages of embryonic development came from Gurdon’s remarkable experiments on nuclear transfer in Xenopus published 50 years ago (Gurdon, 1962; Gurdon et al., 2003) and recognized in 2012 with the Nobel Prize shared with Yamanaka. Gurdon showed that when a nucleus was transplanted from an intestinal cell into the cytoplasm of an enucleated egg, a viable tadpole clone could develop, demonstrating that the somatic cell contained an intact genome. Several decades later Wilmut accomplished the same feat of somatic cell nuclear transfer (SCNT) in a mammal to generate cloned sheep such as the famous “Dolly” (Wilmut et al., 1997).

Yamanaka and Blau (2010) discovered, astonishingly, that the forced expression of four transcription factors that are normally found in early embryonic cells suffices to reprogram differentiated somatic cells to an

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ES-like state, in much the same way as could be accomplished by SCNT or fusion with pluripotent cells. His group and that of Thomson confirmed that reprogramming of human cells, as with mouse cells, can reset pluripotency (Takahashi et al., 2007; Yu et al., 2007). The implications of these findings for regenerative medicine are enormous. Human iPS cell technology in principle enables the creation of patient-specific cellular therapeutics for any specialized lineage. Perfect histocompatibility matching would presumably obviate any need for immunosuppression (Fairchild, 2009). This assumption has been supported by studies with cloned bovine tissues produced by SCNT (Lanza et al., 2002), but challenged by recent experimental data with murine iPS cells (Zhao et al., 2011). In the case of genetic disorders, it should be possible to correct mutations prior to returning cells to the patient (Xu et al., 2009; Kazuki et al., 2010; Khan et al., 2010; Howden et al., 2011; Wong and Chiu, 2011). For patients with chronic infections such as a hepatitis or immunodeficiency virus, it may be possible to arm cells with protective antiviral genetic elements (Kamata et al., 2010; Rahman et al., 2011).

Taking diabetes as an example, a patient could donate a small sample of skin, hair, or blood for reprogramming to iPS cells. These would be expanded in culture, induced to differentiate to the pancreatic β-cell lineage, and implanted to replace the insulin-producing cells lost because of the disease. In type 1 diabetes, measures might still be required to counter the autoimmune attack that led to destruction of the patient’s original β-cells. For example, triggering antigens might be eliminated or masked by genetic modification. Great basic and practical hurdles remain to be overcome before the iPS cell technology and adjuncts such as the correction of genetic defects can be implemented at a meaningful scale in human medicine. Nevertheless, the overall promise of reprogramming cells to enable autologous therapies for regenerative medicine is spectacular (Nishikawa et al., 2008; Csete, 2010; Okita and Yamanaka, 2011; Ji et al., 2012; Robinton and Daley, 2012).

1. Genetic Reprogramming. In the first reports of iPS technology, integrating recombinant vectors were used to deliver four genes (OCT4/ SOX2/ KLF4/ c-MYC or OCT4/ SOX2/ NANOG/ LIN28) to fibroblasts or other differentiated cells. The risks of insertional mutagenesis or reactivation of an oncogene (e.g., c-MYC) were perceived as significant safety issues for clinical applications of reprogrammed cells. It quickly became apparent that the expression of the inserted transgenes shuts off relatively early in reprogrammed cells, and endogenous pluripotency genes turn on. Therefore, transient expression of the reprogramming factors should suffice, allowing the use of self-inactivating or deletable viral vectors or nonintegrating viral or plasmid vectors (Okita et al., 2008; Stadtfeld et al., 2008; Chang et al., 2009; Yu et al., 2009). The introduction of the reprogramming transcription factors can also be accomplished in the form of purified recombinant proteins, sometimes with peptide tags to facilitate entry into the cell (Kim et al., 2009; Zhou et al., 2009; Cho et al., 2010; Pan et al., 2010; Zhou and Ding, 2010). Another emerging approach uses synthetic modified mRNA encoding the reprogramming factors to avoid the risk of permanent genetic modification of cells by DNA transduction (Warren et al., 2010).

2. Pharmacological (Chemical) Reprogramming. Pharmacology already is being applied to the reprogramming process. In fact, much recent attention has focused on the potential of small molecules to make reprogramming more efficient, complete, accurate, and safe and possibly to replace some or even all of the transcription factors now used to reset cells to the pluripotent state (Shi et al., 2008; Ichida et al., 2009; Lyssiotis et al., 2009; Desponts and Ding, 2010; Wang et al., 2011b). Recent review articles from Ding’s laboratory, which has been a major contributor to this field, highlight the rapid progress that has been made (Li and Ding, 2010; Li et al., 2012). In some cases the choice of a target cell type that already expresses some of the four “pluripotency factors” can simplify the reprogramming cocktail. Under favorable circumstances iPS cells can be obtained using either NANOG or OCT4 in combination with small molecules (Theunissen et al., 2011; Yuan et al., 2011b). Studies in ES cells have shown that these two transcription factors are closely associated with the pluripotent phenotype (Chambers, 2004; Kashyap et al., 2009). An important goal will be to produce pluripotent cells that most nearly resemble normal cells of the early embryo, especially in features relevant to the safety and efficacy of future regenerative medicines.

Classes of small molecules that can contribute to reprogramming have been reviewed recently (Sidhu, 2011; Yuan et al., 2011a; Choi and Nam, 2012). Examples include compounds that influence obvious targets for epigenetic regulation such as inhibitors of histone deacetylase and histone methyltransferase (e.g., HMTase G9a) inhibitors, histone demethylase (e.g., LSD1), and inhibitors of DNA methyltransferase (e.g., 5-Aza). Inhibition by a compound designated AMI-5 of a previously unexpected target, methylation of proteins on arginine residues catalyzed by protein arginine methyltransferase, together with inhibition of TGF-β signaling, enable reprogramming to pluripotency by Oct4 as a single genetic factor (Yuan et al., 2011b). Additional compounds that influence cell signaling also can contribute to reprogramming. These include inhibitors of GSK-3, already discussed in other contexts; inhibitors of the Src protein-tyrosine kinases and of the MEK dual-specificity kinase that phosphorylates the tyrosine and threonine residues of ERK kinases.
required for activation in cellular signal transduction; and an L-type calcium channel agonist.

Attention also has focused on the use of small molecules to facilitate maintenance of pluripotent cells, once lines are established. For example, inhibitors of the Rho-associated protein kinase, a regulator of cell shape and motility acting via modulation of the cytoskeleton, prevents apoptotic death of dissociated ES and iPS cells, which makes their large-scale cultivation significantly easier (Watanabe et al., 2007; Harb et al., 2008; Koyanagi et al., 2008). A more direct effect on pluripotency per se is exerted by erythro-9-(2-hydroxy-3-nonyl)adenine (EHNA) and various analogs. These enable long-term growth of ES cells without differentiation, while allowing the rapid reacquisition of differentiation potential once the compound is removed (Burton et al., 2010a). EHNA, therefore, can replace FGF-2, a costly growth factor that is often used to maintain pluripotent human stem cells. EHNA inhibits both cyclic nucleotide phosphodiesterase 2 and adenosine deaminase, but other inhibitors of these enzymes do not have the same biologic effect on pluripotent cells, suggesting that a critical molecular target remains to be determined (Burton et al., 2010b).

3. Transdifferentiation. Concerns still remain that reprogramming of somatic cells to a pluripotent state by any combination of genetic factors and small molecules may intrinsically be a disruptive process that is likely to induce unpredictable mutations, genetic instability, and epigenetic alterations. This consideration, together with recognition of practical translational advantages, has led a number of groups to explore the possibility of concerted switching of cells from one specialized lineage to another—a process sometimes called transdifferentiation. Viewed broadly, the directed differentiation of pluripotent cells and the direct reprogramming of mature cells to specific lineages may be viewed as complementary technologies for “turning straw into gold,” that is, to produce useful cells for regenerative medicine (Cohen and Melton, 2011).

Initial reports of success with directed transdifferentiation have catalyzed intensive investigation of both genetic and chemical methods for direct reprogramming to a range of specialized cell fates, including candidates for cellular therapeutics. A key paper from Melton’s laboratory reported the conversion of pancreatic enzyme-secreting exocrine cells to insulin-producing endocrine cells by transduction of three transcription factors (Ngn3, Pdx1, and MafA) in the mouse pancreas in vivo (Zhou et al., 2008). Other switches within cell types of related lineage have been reported. For example, embryonic mouse mesodermal cells can be induced to a cardiomyocyte fate in culture or to develop rapidly into cardiomyocytes if injected into the heart, after introduction of three factors, one encoding a cardiac-specific subunit of a chromatin remodeling complex (Baf60c) and two encoding cardiac lineage transcription factors, Gata4 and Tbx 5 (Takeuchi and Bruneau, 2009). Similarly, introduction of Gata4, Tbx5, and a third cardiac transcription factor, Mef2c, converted fibroblasts to functional cardiomyocytes (Ieda et al., 2010).

In experiments focused on neuronal lineages, mouse fibroblasts have been reprogrammed by introduction of three transcription factors (Ascl1, Brn2/Pou3f2, and Myt1l) into neurons that can generate action potentials and form synapses (Vierbuchen et al., 2010). Similarly, genes for five transcription factors, Mash1, Ngn2, Sox2, Nurr1, and Pitx3, induced conversion of fibroblasts into cells resembling dopaminergic (DA) neurons that exhibited characteristic dopamine uptake and electrophysiological profiles (Liu et al., 2012). These DA-like neurons relieved Parkinson disease-like symptoms in a rat model. Thus, induced transdifferentiation may provide yet another regenerative pharmacology strategy to treat PD (see above). Finally, microRNAs involved in terminal differentiation of neural progenitors, augmented by neural transcription factors, promoted transdifferentiation of human fibroblasts to neurons (Yoo et al., 2011). It is noteworthy that fibroblasts, which originate in mesoderm, derive from a different embryonic germ layer than neurons, which normally derive from ectoderm.

In another switch involving cell types representing different germ layers, several groups have reported the generation of induced hepatocyte-like cells from adipose-derived stromal cells or fibroblasts (Lue et al., 2010; Huang et al., 2011; Sekiya and Suzuki, 2011). For example, a combination of the transcription factors Gata4, Hnf1alpha, and Foxa3, together with inactivation of the cell cycle regulator p19(Arf), gave rise to induced hepatocyte-like cells that were able to restore liver function in a mouse genetic model in vivo (Huang et al., 2011).

A related strategy entails selection of lineage-specific stem/progenitor cell populations that can be expanded extensively. In some cases this apparently has been accomplished by relatively minimal changes that appear to reset differentiated cells back to a progenitor stage within the same lineage. For example, mature, pigmented melanocytes were reverted to neural crest stem cells in response to expression of the active intracellular form of Notch1 (Zabierowski et al., 2011). Similarly, astrocytes were converted to neural stem/progenitors (Corti et al., 2012).

A powerful, general means to obtain lineage-committed cells, both mature and expandable stem/progenitor populations, follows from observations that transient exposure to subsets of the four “conventional” pluripotency factors also can mediate cell fate switching across unrelated lineages without the need to isolate stable iPS cells (Nie et al., 2012). Deng and
colleagues (Efe et al., 2011) tested this idea in the context of conversion of fibroblasts to cardiomyocytes. Exposure of starting cells to Oct4, Klf4, and Sox2 in the presence of a Janus kinase-signal transducer and activator of transcription inhibitor to limit iPS cell formation was followed by exposure to the cardioinductive growth factor BMP-4 in chemically defined medium. This protocol indeed led to the rapid, efficient production of cardiomyocytes (Efe et al., 2011). The same strategy of transient exposure of a population to the pluripotency factors followed by additional lineage-specific genetic factors and/or inductive signals enabled reprogramming of fibroblasts to neural stem/progenitors capable of both extensive self-renewal and differentiation to neurons, astrocytes, and oligodendrocytes in culture (Kim et al., 2011; Han et al., 2012; Thier et al., 2012). Remarkably, Oct4 alone can induce lineage conversion in cells capable of proliferation, without acquisition of a pluripotent phenotype. Bhatia and colleagues observed that colonies of fibroblasts transduced with Oct4 sometimes express a surface marker found on virtually all blood cells (CD45) (Szabo et al., 2010). They demonstrated that these cells have activated hematopoietic transcriptional programs. Furthermore, the fibroblast-derived cells behaved as multipotent progenitors of the granulocytic, monocytic, megakaryocytic, and erythroid lineages and were able to engraft mice and generate the corresponding mature blood cells.

Rigorous testing of cells obtained by various reprogramming approaches will be required to ascertain the fidelity relative to normal lineage programs, the presence of embryonic and fetal versus adult molecular/genetic signatures, as well as for genomic instability and epigenetic idiosyncrasies. Nonetheless, it seems clear that combinations of genetic and pharmacological manipulation increasingly will enable the directed production of lineage-restricted stem/progenitors and mature specialized cells. As these technologies become robust and cost-effective, they will likely drive the creation of individualized cell-based therapies for multiple genetic disorders and degenerative diseases.

V. Disease Models from Patient-specific Reprogrammed Cells

Going beyond regenerative medicine, the ability to produce cells of essentially any lineage with the genotype of any person, coupled with advances in human genetics and genomics, holds promise to literally transform pharmacology through the creation of new, readily accessible models of human disease. The notion is captured in the phrase “disease-in-a-dish” (Saha and Jaenisch, 2009; Gage, 2010; Walker, 2010). The fundamental assumption is that disease phenotypes can be replicated in specialized cells, produced from iPS cells or by direct reprogramming, that carry the genetic constitution of affected individuals. This appears most straightforward for diseases that result from single mutations with high penetrance, i.e., those that demonstrate simple Mendelian inheritance. A first example came from the identification of motor neuron abnormalities in neural-lineage cells differentiated in vitro from patient-specific iPS cells for human spinal muscular atrophy, an autosomal recessive genetic disorder resulting from mutations that decrease levels of the protein survival motor neuron 1 (Ebert et al., 2009; Ebert and Svendsen, 2010).

The concept of in vitro replication of human disease phenotypes in specialized cells derived from patient-specific iPS cells already has gained at least partial validation for a number of neurologic, hematopoietic, cardiovascular, and metabolic disorders (Unternaehrer and Daley, 2011; Ebert et al., 2012; Maury et al., 2012; Rajamohan et al., 2013). Examples of specific conditions for which patient-specific iPS cells have been obtained and differentiated into disease-relevant cell types include Parkinson’s disease (Byers et al., 2012; Jang et al., 2012) and other neurodegenerative disorders (Ito et al., 2012; Jung et al., 2012), Wilson’s disease (Yi et al., 2012), lysosomal storage disorders (Huang et al., 2012b), and diabetes (Fujikura et al., 2012).

A comprehensive review of specific abnormal phenotypes modeled via the disease-in-a-dish approach lies beyond the scope of this article. However, one example can be presented briefly to highlight the potential to generate predictive in vitro models for disorders in which the underlying genetics and biology are relatively complex. Rett syndrome (RTT) exemplifies autism spectrum disorders, a set of neurodevelopmental diseases characterized by behavioral phenotypes such as impaired social interaction and repetitive behaviors. RTT results from mutations in the MECP2 gene, encoding methyl-CpG binding protein 2. This X-linked gene is inactivated randomly in females, so that heterozygous individuals display mosaic expression of an abnormal protein (or complete absence of the protein in about 50% of cells, in the rarer case of null alleles). RTT patient-specific iPS cells have been isolated and shown to undergo X-chromosome inactivation in the course of differentiation to yield functional neurons (Marchetto et al., 2010a; Ananiev et al., 2011; Cheung et al., 2011). Of great importance, the RTT iPS cell-derived neurons showed a number of abnormalities compared with normal controls, such as reduced numbers of synapses and spine density and functional deficits in calcium signaling and electrophysiology (Marchetto et al., 2010a). The studies pointed to a developmental window during which deficits might be corrected. Similar studies with glutaminergic neurons derived from iPS cells obtained from a mouse model of RTT also showed electrophysiological abnormalities, possibly resulting from...
Making use of cells to produce therapeutic agents per se is certainly not a new concept. Before the era of high-throughput drug screening and molecular modeling of drug-receptor interactions, many medications were initially isolated from plant sources (e.g., paclitaxel)—indicating their production by cells (Cragg and Newman, 2013). Similarly, the biochemical industry produces important therapeutics from penicillin (Ligon, 2004) to insulin (Ladisch and Kohlmann, 1992) through the use of microbial, plant, and mammalian cultures both with and without recombinant DNA technology. Although these technologies lead to the production of therapeutic agents through more traditional pharmaceutical manufacturing methods (reactors, separation trains, and related quality control measures), the concept of using cells in situ to produce pharmaceutical agents is a more recent development. Examples of this approach range from the production of growth factors from cells within biomaterial scaffolds that take up viral or nonviral particles (Saul et al., 2007; De Laporte et al., 2010) to injecting cells with viral gene delivery particles in situ (Barton-Davis et al., 1998) to genetically engineering cells before implantation (Edwards et al., 2005). Each of these approaches ultimately leads to cell-based production of growth factors. Typically nonviral methods and non-genome DNA delivery from viruses leads to transient expression of the therapeutic, whereas nucleic acid delivery from viruses that incorporates into the genome can lead to long-term expression. The temporal need for the therapeutic agent must therefore be considered when choosing the construct used to deliver the nucleic acids. Conceptually, cellular biochemical pathways might be used to produce drugs ranging from small molecules to proteins, although the methods to optimize delivery in situ pose as a future challenge.

The use of cells to produce a therapeutic agent, though, is clearly not dependent on genetic manipulation of the cells; certain cell types naturally produce a needed therapeutic. The prototype for materials-based approaches to using cells as factories for production of therapeutic agents is the encapsulation of Islet cells for the production of insulin. Alginate encapsulation of pancreatic Islet cells provides a means to achieve a glucose-responsive system that can respond to the physiologic state in a natural way. Furthermore, such approaches have used components that prevent immunologic response, allowing the use of allogeneic or xenogeneic cell sources to be considered (Opara et al., 2010).

One challenge associated with such approaches to using cells to produce pharmacological agents is that this approach implies the use of these cells in their nonnative state. For example, the encapsulation of pancreatic Islets in alginate (a material derived from algae) is not reflective of the native state of the Islet...
cells in the pancreas where these cells rest on a specific protein-based extracellular matrix. Specifically, it might be expected that the three-dimensional conformation of Islets or any other cell type interacting with a material would differ from its native state, potentially affecting biochemical synthesis of the therapeutic agents. Approaches to better replicate the native state to improve materials for cell-mediated delivery of “native therapeutics” are increasingly investigated, that is, methods for improved recapitulation of native architecture for fabrication of de novo tissues. This has been recognized for some time with the development of small diameter vascular grafts in which proper flow conditions and cell-cell interactions promote proper endothelial cell phenotype for the production of vaso-dilators such as nitric oxide and proper regulation of cell surface markers of endothelial cell health such as selectins. Recently, we demonstrated the importance of the architecture in a tissue engineered ovary through more native-like cell-cell interactions. Through the proper orientation of granulosa and theca cells in an alginate material, more physiologically relevant levels of estrogen and progesterone were produced (Sittadjody et al., 2013).

B. Mesenchymal Stem Cells as a Therapeutic Delivery Vehicle

Stem cells provide the cellular building blocks of new tissue formation, and moreover, pharmacology can be used to guide this process. However, some stem cell populations, notably mesenchymal stem cells (MSC), also have other unique characteristics that make them useful for direct provision of chemicals or other therapeutic agents for pharmacological modulation of tissue and organ regeneration (Porada and Alemeida-Porada, 2010), as they are well known to elaborate a host of bioactive molecules and factors (Caplan, 2013). Related to these properties, there are two applications we will consider herein: 1) the use of MSC as "factories" to deliver trophic factors—immunosuppressive and anti-inflammatory factors that leverage the cells’ intrinsic ability to home to damaged tissues and tumors; and 2) their use as cellular delivery machines for focal regenerative pharmacology, by local production of factors that modulate regeneration, repair, and restoration of tissue and organ function (Mooney and Vandenburgh, 2008). These are considered below.

1. MSCs as Factories for Trophic Factors. There is good evidence that MSCs are able to distribute and then lodge/engraft within multiple tissues in the body and release trophic factors that trigger the tissue’s own endogenous repair pathways, and in some cases (e.g., bone, fat, cartilage) provide a source of tissue-specific cells (Pretheeban et al., 2012; Caplan, 2013). However, the potential of MSCs to mediate repair is often observed in the absence of any evidence of sustained engraftment of the transplanted cell in the damaged organ. Rather, the injected MSCs home to the injured area, in particular to hypoxic, apoptotic, or inflamed regions, and release trophic factors that hasten endogenous repair by producing tissue protection, enhancing angiogenesis, inhibiting fibrosis and apoptosis, and stimulating recruitment, retention, proliferation, and differentiation of tissue-residing stem cells (Joyce et al., 2010). In short, a significant body of evidence now indicates that MSCs can stimulate regeneration and repair (Caplan, 2013), and thus are likely to play important roles in promoting tissue recovery of, for example, the myocardium (Brink and Cohen; 2013; Shim et al., 2013; Williams et al., 2013; Zhao and Huang, 2013), the central nervous system (Joyce et al., 2010; Huang et al., 2012b; Kramer et al., 2012), and the liver (Krishna et al., 2011). Studies on the injured heart, in particular, have provided evidence that many of the beneficial effects of MSCs in the repair/regeneration of the damaged myocardium, may be caused by promotion of angiogenesis (Huang et al., 2009). MSCs appear to secrete vascular VEGF and bFGF upon contacting the injured myocardium, which stimulates the formation of new vessels and increases capillary density to increase/restore blood flow to an infarcted region (Li et al., 2009). In addition to these paracrine and trophic activities, it would seem that MSCs have properties that help not only to reduce existing damage but promote the healing process (Porada and Almeida Porada, 2010). Thus in the liver, it has been shown that MSCs can enhance fibrous matrix degradation, likely through the induction of matrix metalloproteinases. Also in the heart, MSCs may release paracrine factors that attenuate fibroblast proliferation and inhibit collagen synthesis/deposition, apparently by stimulating cardiac fibroblasts to secrete matrix metalloproteinases. Taken together, these studies clearly emphasize the intrinsic pharmacological properties of MSCs to modulate tissue/organ regeneration and repair.

2. Mesenchymal Stem Cells as Delivery Machines. In addition to MSCs, several other cell systems have recently emerged as biologic drug carriers, such as carrier erythrocytes, bacterial ghosts, and genetically engineered stem and dendritic cells (Gutierrez Millan et al., 2012). However, adult MSCs have been widely studied because they are easy to isolate from different tissues (not only from bone marrow) and to differentiate into cells of various organs (Chiu and Rao, 2011; Dhara et al., 2011; Caplan, 2013). These properties, together with their hypoimmunogenicity, make them good candidates either for tissue regeneration or as vehicles in gene therapy.

It is possible either to augment the natural MSC production of specific proteins and to enable the cells to express proteins outside of their native repertoire, which greatly broadens the spectrum of diseases for...
which these cells may provide therapeutic benefit. For example, with respect to novel treatments for arrhythmias, human bone marrow MSCs, which express connexins and can form functional gap junctions in the heart (Brink and Cohen, 2013), can be gene modified to express a desired ion channel (the “funny current” or HCN channel responsible for pacing), and then can be focally implanted to provide a “biological” pacemaker.

In addition, MSCs can be readily transduced with all of the major clinically prevalent viral vector systems, including those based upon adenovirus, the murine retroviruses, lentiviruses, and AAV (Porada and Almeida-Porada, 2010), to efficiently produce a wide range of cytoplasmic, membrane-bound, and secreted protein products. This ease of transduction coupled with the ability to subsequently select and expand only the gene-modified cells in vitro to generate adequate cell numbers for transplantation, combine to make MSCs one of the most promising stem cell populations for use in gene therapy studies and trials. The ability of MSCs to migrate to a target tissue in vivo suggests the potential use of hMSCs as a cellular delivery system for a variety of bioactive molecules, and recent studies indicate that small interference RNA and microRNA are also among these, as they are able to cross gap junction channels (Brink et al., 2012). Use of genetically recombinant stem cells and biomimetic nanostructured scaffolds for the development of novel biomimetic drug delivery systems has received widespread attention as a promising strategy for wound treatment, in which multipotent stem cells, encoded with plasmid DNA coding for polypeptides, are used both as the cellular therapeutic medium as well as the vehicle for the delivery of functional genes to the wound site (Peng et al., 2012).

The majority of studies using gene-modified stem cells have been undertaken with the purpose of enhancing the natural abilities of stem cells to mediate repair within various tissues; however, MSCs have the ability to accumulate at the site of not only tissue/organ damage/inflammation but may also locate to cancer tissue when administered in vivo (Studeny et al., 2002, 2004; Hall et al., 2007). The MSCs seem to have the ability to “sense” the forming tumor, migrate to the tumor, and contribute to the newly forming tumor stroma (Hall et al., 2007). This property is now recognized as a powerful and unique means of selectively delivering anticancer gene products to tumor cells in vivo (Studeny et al., 2002, 2004; Hall et al., 2007; Hu et al., 2012). Analogous strategies may be applicable to tissue and organ regeneration as well.

VII. Summary and Future Directions

There has been an explosion of information and a parallel increase in technology development since we first attempted to bring the new field of regenerative pharmacology to the attention of pharmacologists (Andersson and Christ). However, our overall goal for doing so remains unaltered, that is, to get pharmacologists more involved in this growing field of research by exposing them to the tools, opportunities, challenges and expertise that will be required to increase awareness and spur excitement within the pharmacological community. Our overall goal was to provide sufficient detail from each of the critical intersecting fields of research to emphasize the necessity for multidisciplinary collaboration and hope that one outcome of this report is that it will indeed generate the required conversations among all of the stakeholders. In short, we believe that the field of regenerative medicine and its companion field, tissue engineering, will benefit tremendously from the more rigorous application of pharmacological sciences. That is, despite the unequivocal success and enormous potential of regenerative medicine and tissue engineering technologies, a greater focus on evaluation of functional outcomes and endpoints is still required. Even from a macroscopic perspective it is clear that a more extensive characterization of basic pharmacokinetic and pharmacodynamics principles is required. Specifically, this should include, among others, assessment of excitation-contraction coupling mechanisms, more rigorous analysis of concentration-response curve data using standard pharmacological analyses/methods, estimation of receptor affinity, receptor subtypes, intrinsic activity, efficacy, potency, etc. In fact, a greater emphasis on the pharmacology and physiology of various regenerative medicine and tissue engineering approaches is critical to increase understanding of tissue/organ regeneration and repair processes and therefore is a necessary prerequisite to increasing the rate of technology development and eventual clinical translation. To this end we have attempted to unite, in a single report, the salient features of diverse fields of research—ranging from materials chemistry and functionalized biomaterials to stem cells, organ/tissue regeneration, wound healing, and development biology—in the hope that providing all of this information at this time would provide the foundation for future interactions and discussions. This report represents the first leg in a long journey.

Authorship Contributions

Participated in research design: Christ, Saul, Furth, Andersson.
Wrote or contributed to the writing of the manuscript: Christ, Saul, Furth, Andersson.

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