Cardiac stem cell therapy: Does a newborn infant’s heart have infinite potential for stem cell therapy?

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Feature Editor’s Introduction—The recent decade has witnessed unprecedented research activity in the field of myocardial regeneration with particular enthusiasm displayed around cardiac stem cell therapy. This enthusiastic research resulted in a number of clinical studies, in which cardiac progenitor cells were delivered to the myocardium during heart surgery in children and adults, demonstrating improvement in heart function. The stem cell therapy appeared to be of great promise in children, particularly neonates, who are likely to have higher regenerative potential. A fascinating and groundbreaking study was recently published in Nature, and is discussed below, demonstrating that injection of live or dead cardiac progenitor cells, live or dead mononuclear cells, or zymosan—a nonspecific inducer of inflammation obtained from baker’s yeast—into the heart resulted in activation of innate immunity and sustained improvement of cardiac function. No new cardiomyocytes were generated. This beneficial effect was abrogated by immune-suppression or macrophage blockade, suggesting that innate immunity is necessary to improve cardiac function. If sustained improvement of cardiac function can equally be obtained by the substance coming from dead cardiac progenitor cells or from baker’s yeast, should we continue to refine stem cell therapy or focus on nonexpensive ways to induce acute immune response? Could injection of tomato juice instead of cardiac progenitor cells provide the same end result? It is now becoming increasingly clear that the immune system plays a key role in regeneration of an injured heart. Where to from here? Below is a feature article produced by Shunji Sano, the world’s leading expert in clinical application of cardiac stem cell therapy, and his team discussing the current state of stem cell therapy for a newborn infant’s heart. The feature article and accompanying commentaries provide stimulating reading for inquisitive surgical minds.

CENTRAL MESSAGE
A newborn infant’s heart has an infinite potential for stem cell therapy and neonates might be the best candidates for stem cell therapy.

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For the past 2 decades, stem cell therapies have emerged as among the most highly investigated treatments for cardiovascular disease. A majority of the preclinical and clinical research in stem cell therapy for heart disease has been focused on the recovery or regeneration of ischemic myocardium in adult patients.1-3 Several stem cell populations from people of different ages (neonate to adult) and from different tissues (noncardiac or cardiac) are under active investigation. Nonembryonic stem cells have been found in the bone marrow, peripheral blood, adipose tissue, skeletal muscle, umbilical cord blood (UCB), and cardiac tissue. As shown in Table 1, various types of stem cells have been used for cell therapy in cases of ischemic heart disease and heart failure (HF).4 Several routes of delivery of stem cells, including intracoronary, intramyocardial, and intravenous, are applied to the heart. Initial systematic reviews and meta-analysis related to stem...
cell therapy for HF and acute myocardial infarction (AMI) have revealed the benefit of cell therapies for cardiac function, exercise tolerance, and quality of life in adult populations.2

The discovery of endogenous cardiac progenitor cells (CPCs) and cardiosphere-derived cells (CDCs) is among the exciting new technologies in the field of cardiac regeneration, which has yielded much interest relating to their proliferative and differentiation potentials. Several investigators have revealed resident populations of CPCs in postnatal human hearts.5,6 The term c-kit+ CPCs refers to a multipotent cell population expressing the tyrosine kinase receptor c-kit. They are believed to be a primary source for the generation of new myocardium in the setting of myocardial injury. CDCs, derived from the mesenchymal component of heart tissue, have been found to possess a multipotent ability. CDCs consist of a heterogeneous cell population and unlike c-kit+ CPCs, CDCs do not require cell sorting. The heart tissue from which CDCs can be isolated and expanded is typically obtained by endomyocardial biopsy, but may also be collected during open cardiac surgery. CDCs have been shown to exert a greater functional benefit in experimental MI with respect to their remarkable potential for myogenic differentiation, angiogenesis, and paracrine factor secretion compared with other cell types.7 Based on these results, the first-in-human phase I Food and Drug Administration-approved clinical trial using autologous c-kit+ CPCs in patients with myocardial ischemia commenced in 2009 (the Cardiac Stem Cell Infusion in Patients with Ischemic Cardiomyopathy trial).8 The Cardiosphere-Derived Autologous Stem Cells to Reverse Ventricular Dysfunction and Autologous Cardiosphere-Derived Cells trial was also commenced using autologous CDCs infusion in patients with MI in 2009.8 Although many clinical trials are being studied in adult patients with HF and AMI using allogeneic CDCs and autologous cells, most meta-analyses seem to agree that the potential beneficial effect of cell therapies is still inconclusive and statistically underpowered. Furthermore, the recently published global position paper on cardiovascular regenerative medicine stated that, even if cell based therapy in HF patients proved to be safe, the results are neither positive nor consistent and true mechanism of stem cell therapy is still unclear.4,9

MECHANISMS OF CARDIOVASCULAR REGENERATIVE RESPONSE

Somatic stem cells, including cells derived from the heart, bone marrow, adipose tissue, and even circulating nonmyocardial progenitors, are the major cell types that might give rise to cardiomyocytes that in theory could be used in clinical trials. Among the cells described above, CDCs have been shown to exert a greater functional benefit in experimental MI with respect to their remarkable potential for myogenic differentiation, angiogenesis, and paracrine factor secretion compared with other cell types.10 Human CDCs could be isolated from right atrial specimens, and resident c-kit-expressing CPC subpopulation has shown to be the most abundant and proliferative when grown from neonatal CDCs, but this subpopulation declined with age.11

A large number of clinical trials have shown stem cell therapy to be a promising therapeutic approach for the treatment of HF; however, the precise underlying mechanisms of action of stem cells are still unclear. Initially, transplanted stem cells were believed to differentiate into cardiomyocytes that integrate with innate myocytes. However, further investigation has revealed that the engraftment rate of transplanted cells was low in the long-term, suggesting a likelihood of insufficient number of differentiated cardiovascular cells to directly replenish the damaged tissue to produce measureable improvement in cardiac function.12

<table>
<thead>
<tr>
<th>Type of stem cells</th>
<th>Origin</th>
<th>Characteristic</th>
<th>Advantage</th>
<th>Disadvantage</th>
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<tbody>
<tr>
<td>Skeletal myoblasts</td>
<td>Muscles</td>
<td>Unipotent</td>
<td>Easy to harvest</td>
<td>Low functional integration between cardiac muscles</td>
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<tr>
<td>Bone marrow mononuclear cells</td>
<td>Bone marrow</td>
<td>Multipotent</td>
<td>Easy to harvest</td>
<td>No specific to cardiac</td>
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<tr>
<td>Mesenchymal stem cells</td>
<td>Bone marrow</td>
<td>Multipotent</td>
<td>Easy to harvest</td>
<td>No specific to cardiac</td>
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<tr>
<td>Mesenchymal stem cells</td>
<td>Umbilical cord</td>
<td>Multipotent</td>
<td>Easy to harvest</td>
<td>No specific to cardiac</td>
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<tr>
<td>Peripheral blood-derived stem cells</td>
<td>Peripheral blood</td>
<td>Multipotent</td>
<td>Easy to harvest</td>
<td>No specific to cardiac Low number of cells</td>
</tr>
<tr>
<td>Adipose-derived stem cells</td>
<td>Adipose</td>
<td>Multipotent</td>
<td>Easy to harvest</td>
<td>No specific to cardiac Low number of cells</td>
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<tr>
<td>Umbilical cord blood-derived stem cells</td>
<td>Umbilical cord blood</td>
<td>Multipotent</td>
<td>Easy to harvest</td>
<td>No specific to cardiac Low number of cells</td>
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<tr>
<td>Resident cardiac stem cells</td>
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<td>Multipotency</td>
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In addition, the lineage tracing experiments have revealed that vasculogenesis rather than cardiomyogenesis might be an alternative mechanism of action that could be critically involved in the processes of myocardial regeneration. Experimentally, vasculogenesis may be important in models of chronic coronary occlusion. Clinically, formation of new vessels may contribute to improved cardiac performance in some patients with ischemic heart disease, but it is difficult to imagine how it could do so in the setting of nonischemic cardiomyopathy or in patients with ischemic heart disease.

Recent studies have revealed that stem cell transplantation stimulates endogenous cardiac repair process via paracrine signaling, direct cell to cell interaction, and/or transfer of microRNAs from exosomes that influence the transcriptional activity of host cells. The important role of exosomes has been highlighted as a mechanism of action after stem cell therapy. Exosomes released from human CDCs were cardioprotective. They inhibited stress-induced cardiomyocyte apoptosis, induced cardiomyocyte proliferation, and stimulated angiogenesis compared with dermal fibroblast-derived exosome in vitro. Exosomes derived from CDCs promote angiogenesis, cardiomyocyte survival, and an antifibrotic effect, which is mediated by microRNA transfer.

Although recent evidence suggests that exosomes may act as a vector of genetic information, and messenger RNAs carried by exosomes can be translated into proteins in the target cell, further studies are needed to enhance our understanding of both cellular and molecular mechanisms of cardiac repair with stem cell therapy.

In the issue of an acute immune response underlines the questions of cardiac stem-cell therapy. Vagnozzi and colleagues tried to disclose the content of acute sterile immune response mainly consisting of the induction of C-C chemokine receptor 2 (CCR2)+ and CX3C chemokine receptor 1 (CX3CR1)+ macrophages in mice treated by cardiac stem cell or sterile antigen exposure after ischemia–reperfusion injury. The authors focused on acute sterile immune response characterized by transient and local tried induction of CCR2+ and CX3CR1+ positive macrophages.

The induction of CCR2+ and CX3CR1+ macrophages at the middle stage of ischemia-reperfusion injury (approximately 3 to 7 days) were induced not only by cell therapy but also by intracardiac injection of cells that have been killed by freeze-thaw or chemical inducers of the innate immune response.

After all, both cell therapy and inducers of the innate immune response resulted in functional rejuvenation—the formation of new cardiomyocytes (ie, pericentral material 1+) in the heart after ischemia-reperfusion injury. In these respects, the functional benefits of cardiac cell therapy have shown the potential for an acute inflammation-based wound healing response that rejuvenates the infarcted area of the heart.

In turn, CX3CR1-null mice did not benefit from bone marrow mononuclear cells (MNCs) therapy and showed a much larger total inflammatory response. CX3CR1 deficiency does not impair the contents of macrophages present in tissues, but affects the function of these macrophages and increases tissue inflammation. However, a significantly higher mortality rate after MI was observed in CX3CR1-null mice. This suggests that CX3CR1 cells play a key role in late infarct maturation and remission. Injecting bone marrow MNCs around the ischemia-reperfusion injury area significantly reduced extracellular matrix content in the border area around the infarction. Certain subtypes of macrophages recruited by MNCs therapy differentially affect the passive mechanical properties of the MI area by influencing cardiac fibroblast activity. These results indicate an important functional distinction between CCR2 and CX3CR1 macrophages in cardiac wound healing.

On the other hand, there is another perspective has been associated with the early stage of wound healing (approximately 3 days) after MI inducing a loss of normal immunosuppressive mechanisms. A key player of immunosuppressive mechanisms are regulatory T lymphocytes (Tregs). Tregs are considered important suppressors of the immune response. In fact, CD4+ Foxp3+ Tregs strongly expanded in the heart, circulation, spleen, and lymph nodes after MI. An inducible Foxp3-null mouse, at age 8 to 10 weeks, with ischemic cardiomyopathy reverses about 10% of left ventricle remodeling and dysfunction 4 to 8 weeks after artificial MI. Left ventricle remodeling and dysfunction while suppressing circulating CD4+ T cells and systemic inflammation, promoting tissue angiogenesis and hypertrophy. Acute Treg dysfunction after ischemia–reperfusion injury was also closely associated with angiogenesis.

These aspects suggest that regulation of the innate immune response is important in practicing stem cell transplantation therapy. In addition, it is also understood that there are various factors to be verified in the process of healing an ischemic lesion.

ADVANTAGE OF PEDIATRIC CARDIAC STEM CELL THERAPY

Human CDCs isolated from neonatal heart tissue have revealed a remarkable regenerative capability compared with those from adults in experimental MI, suggesting a possible contribution of CDCs in congenital cardiac repair. Direct comparison of CDCs between children and adults has shown that there might be a greater abundance of these cells within the heart of patients with congenital heart disease (CHD). Mishra and colleagues investigated the prevalence and proliferation capacity of different stem cell-like cells acquired from cardiac tissue of children
undergoing open heart surgery. They showed that expression of c-kit was highest within the right atrium (5.2%) compared with the ventricle (1.4%). Also, c-kit expression was highest in neonates (8.9%) and declined with advancing age in infants (6.4%) and children >2 years of age (3.2%). In addition, c-kit+ human cardiac progenitor cells (hCPCs) were 3 times more frequently found in neonates than in children over 2 years. The proliferation and differentiation potential of the hCPCs was also greater in neonates, as shown by the higher expression levels of c-kit and Ki67, as well as the expression of NKK2, NOTCH1, and NUMB, the genes responsible for proliferation and differentiation. Furthermore, heart tissue samples of children with CHD contained an increased number of c-kit+ cells, and these cells expressed cardiac lineage and endothelial transcription factors during differentiation under in vitro conditions.19

There is more evidence that CPCs obtained from neonates are superior to those from adults in the recovery of cardiac function.20 The greater expression of heat shock factor-1 in neonatal CPCs may account for this superiority. Similarly, Agarwal and colleagues21 found that when studied in rats with MI the exosomes, lipid bilayer nanovesicles secreted by a variety of the cells when multivesicular endosomes fuse with the plasma membrane, obtained from CPCs of human neonates were superior to those from older children. These age-dependent functional characteristics of CPCs were further investigated by using c-kit– expressing CPCs. Transplantation of neonatal CPCs into rat infarcted heart tissue produced significantly greater functional recovery through increased neovascular formation and paracrine factors secretion when compared with that paracrine factors secretion when compared with that of adult heart-derived CPCs. Neonatal CPC-derived total conditioned media could recapitulate these beneficial effects but not in that of adult CPCs, suggesting neonatal CPC secretome may play an important role in these processes.

CLINICAL TRIAL OF PEDIATRIC CARDIAC STEM CELL THERAPY

The most common cause of late complications in these single ventricle (SV) physiology patients is HF and our aim of regenerative therapy in patients with SV is simply to increase healthy myocardium, particularly in hypoplastic left heart syndrome with right ventricle-dominant anatomy.

The first clinical and comparative study of stem cell therapy in patients with SV physiology were conducted in Okayama University as the phase I Transcoronary Infusion of Cardiac Progenitor Cells in Patients with SV Physiology (TICAP) and as the phase II Cardiac Progenitor Cell Infusion to Treat Univentricular Heart Disease (PERSEUS) trials. The TICAP trial was designed to evaluate the safety and feasibility of stem cell therapy for HF in children with hypoplastic left heart syndrome.22,23 In our protocol, right atrium tissue was obtained during stage II or stage III operation to isolate CDCs. The cells were cultured to reach a cell number of 30,000 per kilogram of body weight and then infused into the coronary artery by cardiac catheterization 4 to 5 weeks after the surgical procedure. The initial results at 18 months after injection of the CDCs reported no adverse events related to the therapy and a mean improvement in ejection fraction from 46.9% to 54.0% (vs a change of 46.7% to 48.7% in the control group). In addition, a significant improvement of growth was observed at 18 months, whereas there was no change in growth in the control group. The phase II study (PERSEUS) has been conducted since April 2013 after the encouraging results of the TICAP trial.24 In PERSEUS, we expanded the indication to patients with SV. A total of 34 patients were randomly assigned to the treatment or control group in a 1:1 ratio. The results of the PERSEUS trial suggest robust evidence that intracoronary infusion of autologous CDCs have a positive effect for cardiac function. At 3 months, the absolute changes in ventricular function were significantly greater in the CDC-treated group than in the controls (+6.4% vs +1.3%). These beneficial effects of CDCs implantation lead to a reduction of HF status; acceleration of somatic growth, which causes improvement of quality of life; and relief of parental stress. A retrospective cohort study to evaluate the therapeutic efficacy of CDCs infusion in patients with SV physiology at 2 years demonstrated that intracoronary delivery of CDCs significantly improved ventricular function and reduced the incidences of late failure, adverse events, and unplanned catheter intervention in patients with SV who underwent stage 2 or 3 palliation at 2 years of follow-up.25 An industry-sponsored, multicenter, randomized phase III clinical trial for SV (Cardiac Stem/Progenitor Cell Infusionin Univentricular Physiology [APOLLON] trial; ClinicalTrials.gov identifier: NCT02781922) started in 2016 at 3 children’s hospitals, including Okayama University, to verify the risk/benefit of intracoronary infusion of autologous CDCs in a large population.

There are additional 8 ongoing cell-based clinical trials for pediatric HF (Table 2). Out of these 8, 3 clinical trials are using autologous UCB-derived stem cells. Recently, Burkhart and colleagues26 reported the result of phase I clinical trial using autologous USB directly into the right ventricle myocardium of patients with hypoplastic left heart syndrome. They found that stem cell therapy using autologous USB is safe and feasible, although they found no improvement in cardiac function.

FUTURE DIRECTIONS

Although the functional benefits of cell therapy might be largely mediated by autocrine or paracrine factor secretion to repair the injured myocardium rather than de novo
myocardial differentiation in situ, the achieved consensus has shed light on proteins and noncoding RNAs transferred by stem cell-secreted exosomes to mediate the protective effects. The clinical efficacy of stem cell therapies in adult heart disease remains elusive, and the mechanisms underlying the regenerative processes in children are still unclear.

The advantage of the biological significance of resident CPCs from CHD should be taken into account for disease modeling and development of novel regenerative therapies. New insights into CPCs and mesenchymal stem cells isolated from pediatric patients, as well as molecular mechanisms controlling the cell proliferation and functional differentiation of these cells during stressed workload, may provide the foundation for exciting regenerative approaches for CHD treatments.

Many groups are currently investigating various avenues of stem cell optimization such as cell preconditioning using drugs or environmental stressors and genetic manipulations to increase angiogenesis, enhance survival pathways, and boost the promoters to differentiate into effective cardiomyocytes. The potential for optimizing stem cell therapy using preconditioning, genetic modification, or other approaches for CHD treatments.

Conflict of Interest Statement
The authors reported no conflicts of interest.

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References


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