Heart failure is a life-threatening disease prevalent worldwide. Cardiac transplantation is the last resort for patients with severe heart failure, but donor shortages represent a critical issue. Cardiac regenerative therapy is beneficial, but it is currently unsuitable as a substitute for cardiac transplantation. Human induced pluripotent stem cells (hiPSCs) are excellent sources for the generation of terminally differentiated cells. The preparation of a large number of pure cardiomyocytes (CMs) is the major premise for translational studies. To control the quality of the generated CMs, an efficient differentiation method, purification strategy, and mass-scale culture must be developed. Metabolic purification and large-scale culture systems have been established, and pure hiPSC-derived CMs of clinical grade are now available for translational research. The most critical challenge in cell therapy is the engraftment of transplanted cells. To overcome the low engraftment ratio of single CMs, aggregations of CMs are developed as cardiac spheroids. A cardiac transplantation device with domed tips and lateral holes has been developed for the transplantation of cardiac spheroids. Large animal models are necessary as the next step in the process toward clinical application. The transplant device has successfully been used to inject cardiac spheroids uniformly into myocardial layers in swine, and this approach is progressing toward clinical use. Remaining issues include immunological rejection and arrhythmia, which will require further investigation to establish safe and effective transplantation. This review summarizes the present status and future challenges of cardiac regenerative therapies. (DOI: 10.2302/kjm.2020-0009-IR)

Keywords: heart failure, regenerative medicine, induced pluripotent stem cell, cardiomyocyte, translational research

Background

Heart failure (HF) is a life-threatening disease and the leading cause of hospitalization in many developed countries. Both the mortality and morbidity resulting from HF are very widespread, and the number of HF cases is increasing annually worldwide, a phenomenon referred to as the “HF pandemic.” Severe HF is treated using device therapies (cardiac resynchronization therapy and implantable cardioverter defibrillator) in addition to optimal medical therapies. However, many patients eventually require ventricular assist devices and cardiac transplantation at stage D heart failure in the American Heart Association and American College of Cardiology’s A-to-D staging system. Ventricular assist devices can cause severe infections and thrombosis, and donor shortages are a persistent challenge for cardiac transplantation. Therefore, cardiac regenerative therapies are currently used to treat patients with severe HF; however, no “standard” cell therapy has so far been established for HF.
Human pluripotent stem cells (human embryonic stem cells and induced pluripotent stem cells) are potent sources of human cardiomyocytes (CMs) in vitro. Specifically, human induced pluripotent stem cells (hiPSCs) have strong potential to avoid immunological rejection by using patient-derived iPS cells. In recent years, the clinical application of hiPSCs has been realized in many fields (e.g., macular degenerative disease and Parkinson’s disease), and hiPSCs also have great potential for the treatment of HF. However, a large number of hiPSC-derived CMs must be prepared in vitro for the effective improvement of cardiac function. CMs developed in vitro are immature and it is difficult to determine whether they will function effectively along with host CMs. Therefore, transitioning from basic research to translational studies is the key to realizing clinical applications (Fig. 1).

In vivo experiments are critical for the translational approach known as “bench to bedside.” Most basic research is performed on rodents (mice and rats) because they are readily available and relatively inexpensive compared with larger animals. In particular, mice are useful for studying molecular mechanisms thanks to the large number of tools available for this type of analysis. Some clinical studies have been approved after rodent experiments; however, confirmatory studies involving larger animals are desirable before clinical trials for regenerative medicine. The physiological characteristics of rodents are different from those of humans. For example, the heart rate is approximately 600 beats per minute (bpm) in mice and more than 300 bpm in rats. The heart rate is particularly important for studies of cardiovascular diseases, because arrhythmogenicity must be assessed for clinical applications. Therefore, larger animals, such as swine, sheep, and dogs, are utilized for translational studies because their sizes are more suitable for the evaluation of medical devices, although the molecular techniques available are very limited. This review focuses on in vitro and in vivo experiments that should lead to clinical applications for cardiac regenerative medicine.

**hiPSC-derived CMs: differentiation, purification, and large-scale culture systems**

Cardiac differentiation from hiPSCs mirrors the developmental steps of the human heart. Pluripotent stem cells maintain pluripotency with core transcription factors (Oct4, Sox2, and Nanog), and their instability leads to the spontaneous differentiation of lineage cells.
Several factors promote lineage differentiation to mesodermal cells. In particular, Wnt/β-catenin signaling potentially induces mesoderm formation. After mesodermal differentiation, CMs are sequentially developed by the inhibition of Wnt signaling. The differentiation ratio varies on a case-by-case basis, so that the differentiation efficiency must be evaluated for each batch. As a result, many non-CMs are also generated after differentiation, including undifferentiated and proliferating cells, leading to the formation of teratomas and other tumors. Therefore, the purification strategy is the most critical issue for exploiting hiPSC-derived CMs for clinical applications. Several purification strategies have been proposed, most of which are based on fluorescent cell sorting. However, large numbers of CMs are difficult to purify using fluorescent cell sorting systems because the process is time consuming and leads to severe cell damage. We developed a purification system that utilizes metabolic differences between pluripotent stem cells and CMs. The depletion of glucose and glutamine promotes cell death in hiPSCs and non-CM derivatives, cell types that potentially cause tumor formation, whereas lactate supports the survival of CMs. After metabolic purification, less than 0.001% of hiPSCs remain. No teratomas developed when the purified CMs were subcutaneously transplanted to immunodeficient mice. Finally, massive cell culture systems are necessary for the clinical and industrial-scale production of CMs. Usually, large cell culture systems use a suspension cell culture technique, such as a bioreactor. However, suspension culture involves the formation of embryoid body-like masses from hiPSCs, and these tend to initiate spontaneous differentiation. Therefore, a two-dimensional (2D) culture system is a fundamental requirement for maintaining the pluripotency of hiPSCs. We developed a large-scale culture system for hiPSCs and hiPSC-derived CMs with multilayer culture plates and active gas ventilation. This 2D culture system can generate \( 1.5 \times 10^9 \) hiPSC-derived CMs efficiently and sequentially. To reduce the cost of large-scale culturing, a new cell culture coating system must be developed. UV/ozone effectively modifies the surfaces of cell culture plastics and drastically reduces the amount of matrix coating. Adherent cells are relatively difficult to collect in multilayer cell culture plates because mechanical stimuli cannot be applied to all layers evenly, and prolonged incubation with detaching solutions causes severe cell damage. To overcome these disadvantages, we also developed an effective cell detachment system that uses resonance vibrations. This approach reduces the amount of detaching solution and yields a high number of hiPSC-derived CMs that are ready for use in translational studies.

Transplantation strategies: cardiac spheroids and a transplant injection device

The transplantation strategy is a key factor in cardiac cell therapy. Single CMs exhibit limited survival in host hearts. Additionally, the engraftment ratio of transplanted single CMs is very low. We found that only approximately 3% of transplanted cells survived in mouse hearts. Most cells were pumped out by the beating hearts. These results led to the development of improved transplantation strategies. Cell sheets are the most popular cell transplantation technique in regenerative medicine. A cell sheet is relatively noninvasive because it is simply placed on the surface of the heart. However, the application of more than three layers can result in ischemia, making it difficult to synchronize the host heart. To improve the engraftment ratio and cell survival, the cardiac spheroid technique was developed at Keio university. The efficient production of hiPSC-derived cardiac spheroids was also established by using specific microwell plates. Purified hiPSC-derived CMs were reaggregated for 7 days in the microwell plates. These spheroids had a diameter of approximately 200 μm and comprised approximately 1000 pure hiPSC-derived CMs. Fluorescent beads were injected into the swine myocardial layer to determine the most effective diameter for the cardiac spheroids. The 175-μm beads showed a better retention rate compared with the 20-μm beads, suggesting the better retention of cardiac spheroids compared with single CMs. In summary, cardiac spheroids have the potential to be an efficient intramyocardial transplantation strategy to improve the engraftment ratio of hiPSC-derived CMs.

Next, we developed a cardiac injection device specifically for the transplantation of hiPSC-derived cardiac spheroids. For ideal cell engraftment, cardiac spheroids must be distributed uniformly in all myocardial layers. Transplanted cells that were injected using a single needle condensed at only one point in the myocardium, and the condensed hiPSC-CMs were exposed to a strong risk of ischemia. Consequently, the cardiac transplant device was designed to have six needles with domed tips and several lateral holes. The domed tips minimize myocardial injuries. hiPSC-derived cardiac spheroids injected from the lateral holes were distributed uniformly in all myocardial layers. Furthermore, to prevent perforation of the myocardial wall, the needles were angled at 45° to the cardiac surface so that they could not be inserted deeper than 7 mm. Finally, the transplant device was tested in swine hearts. Tissue-marking dye and cardiac spheroids were injected using the device and were found to be distributed uniformly in the myocardial layers. The retention rate of transplanted cardiac spheroids was much higher when using this device than when using single needles. These findings show that the combination of cardiac spheroids and a transplant device successfully improves the engraftment rate of hiPSC-derived CMs.

Clinical studies with hiPSC-derived CMs

The above-described successful in vitro and in vivo
studies have provided a sound basis for clinical studies. Because the most critical steps have been overcome, hiPSC-derived cardiac spheroids are currently ready for clinical application (Fig. 3). The final challenge to overcome is immunological rejection. There are several ways to achieve immunological tolerance. The first and most effective strategy is the auto-transplantation of patient-derived hiPSC-derived CMs. This is the theoretically ideal method, but it is practically difficult because the generation of clinical-grade iPSCs is very expensive. Clinical-grade iPSCs must have a high differentiation capacity and no genetic abnormalities. Each hiPSC derivative must be examined for tumorigenicity in vitro and in vivo. The second strategy for acquiring immunological tolerance involves the use of human leukocyte antigen (HLA)-matched iPSCs, which are especially suitable for the Japanese population.26 It was previously reported that 50 iPSC lines would be adequate to cover a three-locus match (HLA-A, B, DR) in 90.7% of the Japanese population. The third strategy involves the use of immunosuppressive agents. However, no data are currently available for the most effective regimen for the transplantation of hiPSC-derived CMs. From these immunological concerns, the idea of universal donor PSCs (HLA-knockout PSCs) has emerged.27 The critical challenge limiting the adoption of this outstanding strategy is how to avoid the natural killer cell-mediated missing-self responses against HLA-knockout PSCs. Retaining HLA-C has been shown to help HLA-knockout PSC derivatives evade natural killer cell-dependent lysis.28

**Discussion**

Since hiPSCs were first reported, several projects have been undertaken to address their clinical application. hiPSCs have been used clinically against macular degenerative disorder.6 Because of the unique nature of the heart, however, a large number of CMs are required for cardiac regenerative therapies. As this review has stated, most of the challenges related to clinical application have already been resolved; however, some still remain. The maturation of CMs is the most critical remaining concern. Current protocols can generate ventricle-type CMs, but they have only the fetal phenotype. Their contraction force and expression of ion channels are very immature. Three-dimensional cardiac tissues that can be stimulated mechanically and electrically may be a key to acquiring
mature cardiomyocytes. Organoid formation can become very advanced and may take the place of spheroids. In terms of present technology, pure CMs are the best form for clinical application. A mix of proliferative cells, such as endothelial cells and fibroblasts, could increase the risk of tumorigenesis. When the growth of vascular and interstitial cells is controlled in the future, high-quality organoids with vascular formations could help improve the engraftment ratio.

Tumor formation and arrhythmia are the most significant concerns in cardiac regenerative medicine involving hiPSC-derived CMs. Metabolic purification systems can successfully purify hiPSC-derived CMs; however, tumorigenesis still raises concerns because hiPSCs are artificially generated, and genetic variance could induce the activation of cancer-driver genes. Even if CMs are purified completely, there is no guarantee that hiPSC-CMs will not cause tumor formation as a result of transformations 10 or 20 years later; however, this possibility appears to be low because fully differentiated CMs should lose their proliferative potential.

Arrhythmia is a common cardiac disorder, and transplanted hiPSC-derived CMs can cause fatal arrhythmias in patients with severe HF. Arrhythmia can occur through automaticity, triggered activity, and reentry. In addition to these fundamental mechanisms, cell transplantation causes inflammation in the host heart, which potentially aggravates arrhythmia. It has been reported that hiPSC-derived CMs can synchronize with a host heart but may induce arrhythmia, including ventricular tachycardia. These results suggest that we must be prepared for arrhythmia in patients with severe HF in a clinical setting. The accumulation of preclinical data with small and large animal models is necessary for the establishment of effective and safe cardiac regenerative medicine.

**Conclusion**

To realize cardiac cell therapies, in addition to in vitro studies, in vivo experiments are critical as translational studies to ensure safety and efficacy. Small to large animals must be used in accordance with the experimental purposes. Conclusive results from animal studies will accelerate clinical applications and promote further progress of regenerative medicine.
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Conflicts of Interest

The author owns equity in Heartseed Inc. The author is one of inventors named in patent 6333378.

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