Cardiomyopathy and Cell Therapy: Ejection Fraction Improvement and Cardiac Muscle Mass Increasing, after a Year of Bone Marrow Stem Cells Transplantation, by Magnetic Resonance Image

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Abstract

The idiopathic dilated cardiomyopathy (IDC) is one of the major public health problems in the western world. Patients with IDC in functional class IV (New York Health Association - NYHA), even after therapeutic optimization, have high mortality. Stem cell therapy has emerged as a potential therapeutic option for cell death-related heart diseases and several positive effects were assigned to cell therapy in cardiomyopathy. The aim of this study was identify short-term result of cell transplantation in idiopathic dilated cardiomyopathy patients (IDC) who were treated by transplantation of autologous bone marrow mononuclear cells (BMMC). Intracoronary injections of autologous BMMC were performed in eight patients with severe ventricle dysfunction (mean of left ventricle ejection fraction – LEVF=20.03%), cardiac mass muscle around 156.2 g and NYHA between III and IV grades, other 8 IDC patients received placebo. The IDCs were followed - up for one and two years, by magnetic resonance imaging (MRI). The results after one year showed significant improvement in LVEF (mean=181.4) and muscle mass increasing (mean=181.4 g), after two years the LVEF continued improving, reaching a mean of 32.00% and the cardiac muscle mass kept stable (mean=179.4 g). Excepted for one patient, all the other had improvement in the NYHA functional class. The placebo group did not show any improvement. We believe that BMMC implant may be a beneficial therapeutic option for IDC patients.

Keywords: Cardiomyopathy; Bone marrow mononuclear cells transplantation; Left ventricle ejection fraction; Cardiac muscle mass; Magnetic resonance image

Introduction

Cardiovascular disease is one of the major public health problems in the western world. Costs of hospitalization amount to more than 10 billion U.S. dollars annually, quality of life is poor, and thousands of lives are lost every year [1,2]. In Brazil, it is estimated that 2% of the population are affected by dilated cardiomyopathy [3] and the cardiovascular mortality rate, in 2004, was 286 people per 100,000 inhabitants [2,3-8].

Dilated cardiomyopathy is a disease without effective therapeutic options in addition to cardiac transplantation [8-14]. Unfortunately, organ rates are low and most of the patients on transplant waiting list died before receiving a new heart [15-20]. Therefore, a therapy capable of improving cardiac function and more accessible to the population is of great interest [21].

In order to cure heart failure, some revolutionary treatment approaches were proposed. Tremendous excitement and controversy have seen in the fields of stem cell biology and cardiac regeneration [22]. Studies in ischemic heart disease demonstrated that stem cells obtained from various sources, including bone marrow, can contribute indirectly to cardiac regeneration [23-26], and indicated the potential use of stem cell therapy in other cardiomyopathies [27,28].

Several positive effects were assigned to cell therapy in cardiomyopathy, for instance: improvement of left volume ejection fraction (LVEF), decreasing end-diastolic and systolic volumes, myogenesis and angiogenesis, after treatment [28-30]. Stem cell therapy has emerged as a potential therapeutic option for cell death-related heart diseases [31,32]. Preclinical and a number of early phase human studies suggested that cell therapy may augment perfusion and increase myocardial contractility, furthermore, the clinical efficacy of cell therapy remains to be proven [33].

Imaging allows for in vivo tracking of cells and can provide a better understanding in the evaluation of the functional effects of cell-based therapies [33,34]. Among these are direct labeling of cells with super-paramagnetic agents, radionuclide, and the use of reporter genes for imaging of transplanted cells. Magnetic resonance imaging (MRI) is the best suited to meet such broad objective thanks to its resolution and clinical applicability [35]. The use of non-invasive imaging modalities in pre clinical cell therapy studies revealed key aspects of cell biology that will not be observed by other approaches excepted for histological analysis [36,37].

The aim of this study is to identify short-term result of cell transplantation in idiopathic dilated cardiomyopathy patients (IDC)

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Received December 03, 2013; Accepted December 21, 2013; Published December 23, 2013

Citation: Greco OT, Filho IJZ, Bellini MF, Bilaqui A, Souza AS Junior, et al. (2013) Cardiomyopathy and Cell Therapy: Ejection Fraction Improvement and Cardiac Muscle Mass Increasing, after a Year of Bone Marrow Stem Cells Transplantation, by Magnetic Resonance Image. J Stem Cell Res Ther S6: 008. doi:10.4172/2157-7633.S6-008

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who were treated by intracoronary transplantation of autologous bone marrow mononuclear cells (BMMC) and compare these results with the placebo (PL) group, by magnetic resonance imaging [33-38].

Materials and Methods

Materials

The materials used were trypan blue (Sigma Aldrich, St. Louis, MO, USA), Kit monoclonal antibody CD34 (+), CD34(-), CD 133 (+) and CD 133(-) (BD Biosciences, South America, USA), Kit SEPAX (Biosafe America, Inc.1225 North Loop West, Suite 120, Houston, TX 77008, USA), Flow Cytometry (BD Accuri C6™, BD Biosciences, South America, USA). As the density gradient Ficoll-Hypaque was used (Amersham Biosciences, Piscataway, NJ, USA).

Methods

Experimental design – patients: The study was blinded, randomized and performed at 16 patients diagnosed with idiopathic dilated cardiomyopathy (IDC) at Instituto de Moléstias Cardiovasculares (IMC). Bone marrow mononuclear cell (BMMC) injection was performed in 8 and 8 patients received placebo, at the period between 2005 and 2008. From 16 IDC patients, 12 were male, 4 were female and the mean age was 51.8 years (range: 44-62 years). Following the functional classification of New York Health Association (NYHA), 10 IDC patients was grade III and 6 was grade IV. The study was approved by Brazilian National Research Ethics Committee (CONEP) (Registration #: 15342).

The Randomized Assessment of Digoxin on Inhibitors of Angiotensin-Converting Enzyme (RADIANCE) trial found that discontinuing digoxin in patients with low ejection fraction and HF resulted in worsening HF, Current American Heart Association/ American College of Cardiology guidelines classify digoxin use as IIa in patients with current or prior symptoms of HF and reduced left ventricular ejection to decrease hospitalizations for HF. B-Blockers should be maintained in this hemodynamically stable patient (a Class I indication) because there is no evidence that routinely discontinuing β-blockers in this setting is beneficial. Patients such as this usually respond well to intravenous loop diuretics (avoiding the absorption issues of oral diuretics in the setting of intestinal edema), with improved renal function as preload is optimized. In patients who are true refractory to diuretics, intravenous nitroglycerine or ultrafiltration can be helpful (Class IIa) (1). Bone marrow aspiration, isolation of mononuclear cells and cell injection: Before the arrival and manipulating bone marrow sample, it was "check list" materials and reagents required for handling Kit (SEPAX) and bone marrow, such as syringes, needles, 70% ethanol, gas, human albumin, ficoll and serum. Called up the equipment SEPAX, at least thirty minutes prior to stabilization of electric current. Sterilized room and laminar flow hood with UV light for fifteen minutes. The Kit was manipulated inside the laminar flow hood. The first step to manipulation of the kit was to examine whether all three "taps" were in the position "T". The second step was to close all clamps. The third step was to inject 100 mL of Ficoll puriscence washing. The fourth step was to discard 62.5 mL serum bottle and inject the same volume of albumin and connect the bottle right output Kit. The fifth step is to connect the empty bag (for storing the mononuclear cell layer) and the sixth step in the Kit was connected the bag with the bone marrow. For handling the SEPAX was necessary to follow the instructions of the standard protocol for operating the machine in the category of "separation by ficoll." After mounting, the verification was made of three settings: sample volume (60 mL), number of washes (two washes for 500 mL serum). After this, the "enter" checked the Kit after checking Kit, opened up all the clamps. Soon after, the software asked to do the pumping purse bone marrow and then tightened "up" twice to adjust the trajectory of the spinal cord to close the entrance. Thereafter, if pressed "enter" to start the process. First, the Ficoll was aspirated into the tube. Then, the marrow was slowly aspirated into the tube on top of the Ficoll. It started spinning and then the fraction of erythrocytes and granulocytes was discarded and mononuclear ficoll layer was aspirated and washed with saline. After one hour the process ends and the mononuclear cell layer was available for quantifying CD 34 (+) and CD 133 (+), after obtaining the pellet by centrifugation [28]. The final suspension of bone marrow mononuclear cells contained 3.0 × 10^7 totals cells (being 5.0 × 10^6 CD 34+ cells, 2.5 × 10^6 CD 133+ cells and others cells as monocytes) was injected intracoronary in the affected area. The placebo group received intracoronary injection of saline plus autologous serum of the patient solution, to ensure that the study was double-blind, keeping the color of the injected solutions. Before every injection, the catheter was positioned perpendicular to the endocardium with excellent loop stability and the extension of the needle had to induce a premature ventricle contraction.

Magnetic Resonance Imaging (MRI): Magnetic resonance imaging has become a key surrogate end point to demonstrate efficacy in early phases, small-sized studied. MRI can provide detailed morphologic and functional information and therefore, seems ideally suited to integrate efficacy assessments with the capability for cell tracking [16]. MRI studies were performed using a 1.5-Tesla (GE Medical System, Milwaukee, WI, USA) with 8 channels cardiac array and vector electrocardiographic gating. Two experienced observers, blinded to all clinical data, analyzed the images. Previously validated software was used to determine parameters of global systolic function (Advantage Workstation – AW Volume Share 2 (AW 4.4), Report Card 4.0, GE Medical Imaging System, and Milwaukee, WI, USA).

Statistical Analysis

The preliminary evaluation of data was performed based on Descriptive statistics and univariate analysis. The nonparametric Friedman test was applied to assess the time of assessment effect (T0, T1 and T2) was significant. Nonparametric comparisons post hoc Friedman test was also used. The level of significance was set as p<0.05. The statistical analysis was carried out with the statistical software MINITAB® 15.1.

Results

The left ventricle ejection fraction (LVEF) descriptive statistics can be observed in Table 1 and these results were calculated based on measurements made both before cell therapy (T0), after one year of therapy (T1) and after two years (T2). By the statistical univariate analysis, BMMC treated patients reacted positively to the infusion, considering that the average values of LVEF increased from 22.64% to 29.75% at the T0 to T1, reaching 32.69% at the T2.

The analysis of extreme values (minimum and maximum) shown that there was a positive evolution on LVEF, which is most significant in the first year. We can also observe important data variability, after a year and two years of therapy. While at the T0 the LVEF varied between 7.4 to 44.1%, after one year (T1) the percentage increased to
18.8% of the minimum and 58.9% of the maximum LVEF value (Figure 1). In the second year, patients were maintained between 20 and 56.5%. By the data dispersion and the number of patients, LVEF median estimates were considered more reliable than the means.

Some patients had remarkable improvement in the first year of treatment, increasing LVEF in 16.1%. For others, the increase was in second year and also those who had remained stable or showed a slight reduction of 4.2% in the LVEF within the period. Overall the trend is growth in the first year and a lower increase from first to second year (Figure 2). The placebo group did not show any improvement, neither on LVEF (p=0.429) nor for cardiac mass (p=0.438). The data of this group is also on Table 1.

Table 1 also presents the results of univariate analysis performed for the cardiac mass (CM), where we can verify that most patients remained at levels near the values unchanged at both times, the first year and the second year of treatment. Baseline the mean weight was 177.2 g with a standard deviation of 87.8 g. The values ranged from 57.1 to 387.3 g for 8 analyzed IDC patients, half of them had initial CM at least 161.8 g. One year after the infusion of stem cells, the average rose to 189.6 g, standard deviation fell to 53.1 g and the values of CM were included a narrower band (121.6 to 290.7 g). The median increased in proportion of the average, and reached 174.2 g. After 163.1 g, both estimates similar to those initials. The standard deviation (SD) has remained in the first year (SD=51.9 g), as well as range of variation, which was between 130 and 290.7 g. The results in Table 1 indicate differences obtained by the Friedman test (P=0.031) and the significance found by post hoc test (P=0.021), which indicates changes in cardiac mass after one year, suggesting a CM increase (Figure 1). For the period of two years there is no significant difference (P=0.141). There is an increasing trend of CM after 1 year, but with the passing of another year, there is a reduction to initial levels (Figure 3).

The patients had a good evolution in the post-implant with improvement of NYHA functional class. Only one patient kept his NYHA grade (III), 50% of IDC patient’s improved one class in NYHA grade (IV to III, and III to II) and 37.5% improved two classes, changing from IV to II, or III to I grades.

The comparisons between placebo and BMMC groups no showed statistically significant difference for LVEF (p-value=0.1307) but an increasing on cardiac mass was observed (p-value=0.0005) between BMMC and PL therapy on the both times (p-value<0.001).

Discussion

The reparative effects of cardiac tissue may either be due to paracrine effect caused by stem cells in the bone marrow, stimulating the process of "homing" of macrophages, neutrophils, monocytes, stem cells and telocytes, and by differentiation processes of infused cells. In the case of a paracrine effect, the first highlighted are macrophages can be induced by stem cells to produce nitrogen oxide (NO), it is important for tissue regeneration. The second highlight is the telocytes - interstitial cells of Cajal - which play an important role in intercellular communication [18,19]. In the case of cell differentiation, the small fraction of mesenchymal stem cells may have differentiated into cardiomyocytes, and hematopoietic stem cells may have contributed to the cardiac tissue angiogenesis process and may also have transdifferentiated in cardiomyocytes. Furthermore, in adult heart, stem cells there are at rest, but can rapidly proliferate when stimulated with growth factors released by the infused stem cells and macrophages. This is solid evidence that the heart is in a continuous process of growth, cell death and renewal [3].

In preclinical studies, the effects of cell therapy were analyzed in experimental models of infection with Trypanosoma cruzi in mice with chronic Chagas cardiomyopathy. It was observed that infusion of bone marrow stem cells in mice have reduced heart...
Using a combination of electrophysiological and imaging techniques it is concluded that electrically active cardiomyocytes derived from human embryonic stem cells are able to actively pacing at rest, the recipient ventricular cardiomyocytes in vitro and in vivo ventricular myocardium, suggesting an alternative or complementary method for correction of defects in cardiac impulse generation, such as pacemakers based on cells [7]. In experimental models, the infarct size after - myocardium and remained constant in the placebo showed a trend towards reduction in low dose, high dose therapy MSC reduced infarct size from 18.2 (± 0.9%) to 14.4 (± 1.0%) with (p-value=0.02), left ventricular mass. In addition, both treatments of low and high dose increased regional contractility and myocardial blood flow in both infarct and border areas [19]. Saw an increase in muscle mass 156.2 to 181.4 during the first year but decreased to 179.4 in the second year after cell therapy, indicating that the ideal for the regeneration of muscle mass period is the first year post treatment and these results are statistically significant when compared with placebo. Another group showed a significant improvement in cardiac function and volume, resolution of healing and increased wall thickness for severe ischemic dilated cardiomyopathy patients with cardiac magnetic resonance imaging at 6 months of infusion of bone marrow stem cells in relation to baseline [23].

The analysis of the ejection fraction of the right ventricle indicates an improvement of heart function six months after the stem cell transplant. Similarly, we observed an increase in walking time and race test [14]. Patients undergoing cell therapy in this study also showed an improvement in functional class, except for one patient who remained NYHA class III, the other patients showed good recovery post- implant, where 50% of IDC patients reduced one NYHA class grade (IV to III and III to II) and 37.5% showed an improvement of two classes, from IV to II or III to I.

Conclusion
Considering the results obtained by magnetic resonance imaging, we can consider that trends of improvement in ejection fraction and increased cardiac muscle mass in the first year after cell therapy, but these clinical parameters are less pronounced between the first and second year. However the idiopathic dilated cardiomyopathy patients had a good clinical response to the implant of autologous stem-cells. Thus, we conclude that the infusion of bone marrow mononuclear cells has greater therapeutic efficiency in dilated cardiomyopathies.

Acknowledgements
The work was financially supported by the HMC Hospital / BMI of São José do Rio Preto - Brazil. We appreciate the support of Duke University (Durham, NC, USA) in the field of research and statistical studies and also appreciate the support of the Life Group (Brazil), Biosafe (Switzerland, EC) and Celartia (Ohio, USA). The authors are grateful to Dra. Adriana Barbosa Santos for statistical help.

Disclosure of Potential Conflicts of Interest
The authors declare that they have no conflicts of interests.

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