Introduction

Coronary artery disease remains one of the main causes of morbidity and mortality in the world. Despite advances in medical treatment and improvements in revascularisation techniques, thousands of patients are still not amenable to these treatments thus denying them a symptom-free life. For the past few decades researchers have been attempting to develop and utilise cellular therapy to rebuild or revascularise the heart [1-3]. Early clinical trials have shown that implantation of autologous bone marrow cells was feasible with improvement in ischaemic status and left ventricular function [4-11]. Indeed this technique has the potential to become one of the emerging new modalities for treatment of ischemic heart disease. However, most of the previous studies with autologous bone marrow stem cells were performed with concomitant percutaneous coronary intervention (PCI) or coronary artery bypass graft surgery (CABG). In this report we describe our preliminary clinical trials and report our result on the safety and feasibility of direct intramyocardial injections using high cell count of autologous bone marrow stem cells as a sole therapy for severe ischaemic heart disease.

Methods

This was a prospective non-randomized clinical trial. All patients who had severe ischemic heart disease with stable angina that was refractory to maximum medical therapy and not amenable for percutaneous or surgical
revascularization methods either because of diffuse disease or the target vessels were too small, were eligible for recruitment. This has to be validated by an independent interventional cardiologist and cardiothoracic surgeon. Additional inclusion criteria include the presence of myocardial reversibility as confirmed by preoperative technetium scan. This study was approved by the institutional ethical committee and informed consent was obtained from all the patients. Exclusion criteria included: life expectancy of less than a year, active malignant neoplasm, alcohol or drug dependency, severe diabetic retinopathy, poorly controlled diabetes and those requiring concomitant open-heart procedures. Since September 2003, four patients were enrolled.

Preoperative screening

All patients had a repeat coronary angiogram within a month of the procedure to ensure that the coronary arteries were not suitable for percutaneous coronary intervention or surgical revascularization and this was assessed by an independent interventional cardiologist and cardiothoracic surgeon. During cardiac catheterization, left ventriculogram was performed to measure the left ventricular ejection fraction (LVEF), left ventricular end-diastolic pressure (LVEDP), left ventricular end-systolic volume (LVESV) and left ventricular end-diastolic volume (LVEDV) were measured. A SPECT (single photon emission computed tomography) myocardial perfusion-imaging scan using Gamma camera (GE model Optima nx dual detector) was used to detect areas of defect or reversibility. A one day stress-rest or rest-stress protocol was performed. Exercise treadmill (Bruce’s protocol) or intravenous dobutamine infusion was aimed at achieving a target heart rate of more than 85%. Then 10-15 mCi of technetium ($^{99m}$Tc-tetrofosmin) was injected. After about half-hour later, images were acquired over 15 minutes (stress scan). About 3 to 4 hours later another 25 to 30 mCi of technetium scan was injected again. Patient was rested for 45 minutes to one hour before a second scan was acquired (rest scan). Data was collected in optical disc with Dicom compatible format with HP (Hewlett Packard) Kayak as workstation computer. The areas of defect and reversibility were calculated by Cedar Sinai software. Transthoracic two-dimensional echocardiography was also used to assess the global LVEF, regional wall motions and absence of valvular abnormalities. A preoperative 24-hour holter was also used to exclude presence of any arrhythmia. A detailed computed tomography of the abdomen complimented with abdominal ultrasound, chest roentgram, faecal occult blood test, urinary analysis and serum tumor markers such as carcinoembryonic antigen (CEA) CA19-9 and alpha feto protein (AFP) were done to exclude the presence of malignancy. Eye assessment was done by independent ophthalmologist to exclude presence of severe diabetic retinopathy. Blood sugar monitoring in particular glycosylated hemoglobin and fasting blood sugar were optimized prior to the procedure.

Bone Marrow Aspiration

This was done on the same day of the intramyocardial injection. Patients were anaesthetized with full intubations and placed in prone position. The patient was heparinised with monoparin (heparin sodium bovine, CP Pharmaceuticals) 70 iu/kg body weight just prior to the commencement of the bone marrow harvest. The marrow was aspirated using Islamic needle from both the posterior superior iliac crests. Aliquots of 3-5 mls were aspirated each time. The marrow was then deposited into ACD (acid-citrate-dextrose) and heparinised collection bag. At the start of the harvest, 20mls of ACD and 20mls of hepsal (hepsal = 15,000 iu of preservative-free heparin in 500mls of normal saline) were added into the empty collection bag (Fenwal Bone marrow Collection Kit, Baxter Healthcare Corporation). This combination was sufficient to anticoagulate 100mls of bone marrow. After each 100mls of bone marrow collection, an additional of 20mls of ACD and 15mls of hepsal were added. A total of 500mls of marrow was collected from each patient.

Isolation of Autologous Bone Marrow Stem Cells

Following harvest of about 500mls of the marrow, the collection was sent to the pathology laboratory. A buffy coat was prepared using semi-automated cell
processor Cobe 2991. At the end of the process, the volume is reduced to about 250ml. This procedure depleted the red cells and also reduced the volume. This is later centrifuged at 4000rpm for 20 minutes at 18 degrees celsius without braking. From this about 15ml of concentrated buffy coat was then collected using an automated collection system called ‘Optipress’ into a transfer bag and ready for injection. Tests on the final harvest included flow cytometry (CD34, B and T-cells subset enumeration) and culture and sensitivity to check for sterility.

Cell implantation

The patients either had a median sternotomy or lateral thoracotomy depending on areas to be injected. Direct intramyocardial injection of high cell count of autologous bone marrow stem cell was done under general anaesthesia and with the heart beating without the aid of cardiopulmonary bypass. A total of 30-40 injections with a mean number of \(2.4 \times 10^8\) cells (ranges from \(1.1 \times 10^8\) to \(4.2 \times 10^8\)) injected directly under vision into the myocardium as determined preoperatively from the preoperative technetium scan (Table 1). The injections were spaced at about 0.5cm from each other using 27 gauge needle. There were no complications during the injections and in particular no significant arrhythmias or bleeding were noted. No patients required conversion to cardiopulmonary bypass support or intra aortic balloon pump postoperatively. The chest was closed in the routine manner. There were no special requirements postoperatively apart from low dose of intravenous nitroglycerine for the first 24 hours. All preoperative medications were restarted on the following postoperative day. The patients had repeated invasive and non-invasive assessments or investigations at 1 month, 4 month and 12-month postoperative period. This included coronary angiogram with left ventriculography, SPECT scan, 2-Dimensional echocardiography, 24-hour holter and tumour markers.

Statistics

Results are reported as the mean value ± standard deviation. Comparison of the changes from baseline to 12 months in these patients was assessed with Friedman test. A probability value <0.05 was considered statistically significant. Statistical analysis was done with SPSS version 12.0 for windows.

Results

All four patients enrolled were male. The median age was 59.3 years (range from 58 to 60 years old). The bone marrow aspiration went uneventfully and the intramyocardial implantation was completed successfully without any complication. Two patients had resternotomies for previous coronary artery bypass graft surgery, one had lateral thoracotomy and one other median sternotomy. Transesophageal echocardiography was used routinely during the intramyocardial injections to exclude accidental systemic embolisation, in which there was none. The patients were restarted on their preoperative medication but antianginal medication was withdrawn at 1 month follow-up. All except one patient completed the 12-month assessment postoperatively. There was no death, myocardial infarctions or cerebrovascular accidents during the procedure or at our follow up median of 14.3 months (ranges from 10 to 22 months). Periodic holter study showed no arrhythmias. Patients’ symptoms improved as early as 1 month postoperatively and continued to do so at their last follow up. The New York Heart Association (NYHA) functional class improved from a mean of 3.2 preoperatively to a mean of 1.5 and 1.0 at 1 and

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Dm=Diabetes mellitus, HPT=Hypertensive, HLP=Hyperlipidemia, CABG=coronary artery bypass garft

Table 1. Patient clinical characteristics and intraoperative data
4 month follow up (Figure 1). This showed statistical significance with p=0.041. The Canadian Classification Score (CCS) for angina also showed a statistically significant change (p=0.049) with improvement from 2.3 preoperatively to 0 at 4 month postoperatively (Figure 2). The two dimensional echocardiography revealed an upward trend in left ventricular ejection fraction from 50±9% preoperatively to 58±14% at 12 months (p=0.117). Results from LV angiography (LVEF, LVESV and LVEDV) at baseline and at 1, 4 and 12-month follow up showed improvement but statistical significance was not reached (Figure 3). The areas of defect and reversibility from the technetium scan illustrated improvement but again did not show statistical significance (Figure 4). Most importantly all the patients managed to be independent and able to go back to work.
The present report describes the first clinical trial of direct intramyocardial injections using high cell count of autologous bone marrow stem cell with a mean of $2.4 \times 10^8$ cells as a sole therapy for patients who have severe ischemic heart disease whom are not suitable for normal revascularization methods. The number of patients are small because of the strict criteria used in patients selection. This is also compounded by the aggressiveness of most interventional cardiologist in treating triple vesse disease with percutaneous intervention and the improvement of surgical techniques in coronary bypass surgery. Nevertheless our results...
showed that the intramyocardial injections using high cell count of autologous bone marrow stem cell procedure is relatively safe. There was no death or cerebrovascular events during the implantation or during the follow up period. All the mononuclear cells from the bone marrow aspirate were used as a whole, rather than subpopulation as each fraction of the bone marrow cells might contribute to the regeneration of ischemic myocardium. There were no recorded arrhythmias with the 24 hour holter during the follow-up period. There was significant improvement in symptoms of angina and effort tolerance as early as 1 month as shown in Figures 1 and 2. Although this highly subjective, this is supported by the transthoracic echocardiography which showed improvement in left ventricular ejection fraction from 50±9% preoperatively to 57±10% at 4 month and complimented with improvement in left ventriculogram ejection fraction from 61±6% preoperatively to 69±17% at 4 months. The analysis of the areas of defect and reversibility demonstrated some very interesting findings (Figures 3, 4 and 5). We had expected the direct trauma caused by the intramyocardial injections and thus predicted a time interval taken for angiogenesis and regeneration to occur, might make the areas of defect and reversibility to increase before any improvement was to be expected. This is clearly shown from the Figures 4 and 5 where the areas of defect and reversibility were increased at I month but subsequently decreased during follow-up period. These results demonstrate that the injection of high cell count of autologous bone marrow stem cells may lead to angiogenesis and repair of the ischemic myocardium although the objective measurements do not reached values of statistical significance due to the small number of patients. It also showed that the simple approach of direct intramyocardial injection into areas of interest as determined by the preoperative SPECT scan is feasible.

The evidence that cardiac cells may regenerate and the presence of cardiac stem cells stimulated an exciting journey and opportunities for myocardial repair [12-15]. This lead to the idea that the heart may be a terminally differentiated organ but does not composed wholly of terminally differentiated cells. Recent clinical trials using autologous bone marrow stem cells in infarcted hearts have shown functional improvements [4-11]. Other clinical trials with the use of skeletal myoblast transplantation were equally encouraging [16,17]. The postmortem findings in some of these patients proved that engraftment and transdifferentiation may have occurred in both methods [18,19] However both methods of cellular therapy trials were done concomitantly with coronary artery bypass graft surgery or following angioplasties. Our study records two new approaches that differ from previous trials. Primarily we used a high cell count of autologous bone marrow stem cells, and secondly we performed this procedure as a sole therapeutic modality. Our preliminary findings showed that it is relatively safe and very encouraging early results were obtained. It is however must be reiterated that this therapy is not intended to be a first- or second-line therapy. This study aimed to find an alternative therapy that is reserved for those who are symptomatic despite on maximum medical therapy and are not suitable for any means of revascularisation methods. Therefore larger prospective randomized trials are required before such novel therapies could routinely be implemented. Hopefully we would know then which cells to use, how much to use and the best method of delivery for such cells.

Acknowledgement

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References


