Design and implementation of the TRACIA: intracoronary autologous transplant of bone marrow-derived stem cells for acute ST elevation myocardial infarction

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Abstract

Objective: To describe the design of a protocol of intracoronary autologous transplant of bone marrow-derived stem cells for acute ST-elevation myocardial infarction (STEMI) and to report the safety of the procedure in the first patients included.

Methods: The TRACIA study was implemented following predetermined inclusion and exclusion criteria. The protocol includes procedures such as randomization, bone marrow retrieval, stem cells processing, intracoronary infusion of stem cells in the infarct-related artery, pre- and post-MRI, pre- and post SPECT with radioisotope ventriculography, and clinical follow-up at 6 months.

Results: Eight patients with a diagnosis of acute STEMI and duration of symptoms of ≤24 hours that were perfused successfully through primary percutaneous coronary intervention (PPCI) with a LVEF of ≤45% were assigned randomly to two groups (n = 4 each). One group treated with stem cells and the other corresponded to the control group. Neither death, re-infarction, no need for revascularization or thrombosis of the stent were observed at follow-up.

Conclusions: The initial experience at the Instituto Nacional de Cardiología Ignacio Chávez in the treatment of acute STEMI by means of autologous transplantation of bone marrow-derived stem cells is encouraging. Implementation was possible in the first eight patients with no complications.
Introduction

Treatment of acute ST elevation myocardial infarction (STEMI) has changed radically in the last two decades, placing as treatment of choice reperfusion therapy with thrombolysis and, preferably, through primary percutaneous transluminal coronary angioplasty (PTCA). Early restoring of the flow in the artery responsible for the infarct has decreased mortality significantly. However, the myocardial necrosis process starts rapidly after coronary occlusion and an irreversible loss of muscle is produced in many patients despite successful reperfusion. Myocardial necrosis activates a series of repair changes that include formation of scar tissue and ventricular dilation, and this alteration of the ventricular geometry can continue for weeks or months fostering chronic cardiac failure. One way of reverting or ameliorating this process would be the regeneration of cardiomyocytes and by stimulating neangiogenesis within the infarcted tissue. Experimental studies have suggested that the use of bone-marrow-derived adult stem cells can contribute to regenerate the infarcted myocardium and foster neangiogenesis. The first clinical experience with acute infarctation and injecting intracoronary hematopoietic stem cells derived from the bone marrow of the same patient was reported by Strauer in 2002. Several subsequent studies have demonstrated an improvement in the left ventricular ejection fraction (LVEF) of patients receiving stem cells, such as the BOOST, REPAIR-AMI studies and that of Fernández-Avilés. However, information reported until now is still not conclusive and, therefore, it is justified to continue performing prospective and randomized studies.

Objective

To describe the design and implementation of a protocol of intracoronary autologous transplant of bone marrow-derived stem cells for acute STEMI, and to report the safety of the procedure in the first patients included.

Methods

Design of the study and randomization. This study is a controlled, randomized, single-blind, two-arms clinical trial performed at the Hemodynamics Department of the National Institute of Cardiology (Instituto Nacional de Cardiología Ignacio Chávez), in collaboration with the National Center for Blood Transfusion. The study was performed following the principles of good practices of the Helsinki Declaration and the Health Regulations of Mexico, and with the approval of the institutional Research and Ethics Committees. Each patient signed the corresponding informed consent form.

The design of the study is depicted in Figure 1. After the primary PTCA, and between days 3 and 5, patients were assigned randomly at a 1:1 proportion to the stem cells group or the control group. For this, closed envelopes with consecutive numbers were used. In the stem cells group, bone marrow was obtained, performing an intracoronary injection of the tissue, as well as a ventriculography between days 6 and 7. In both groups, a nuclear magnetic resonance imaging (MRI) study, a two-dimensional echocardiogram and simple photon emission computed tomography (SPECT) with isotope ventriculography were performed between days 5 and 7. Clinical follow-up was performed on the first, third, and sixth months in both groups, and cardiac catheterism with ventriculography, SPECT, and MRI were performed.

Inclusion criteria. The study included adult patients of either gender, between 20 and 75 years of age, with a diagnosis of acute STEMI and a duration of symptoms of ≤ 24 hours, who had been perfused successfully with primary PTCA (with or without previous administration of thrombolytics) and with an implanted stent, and who, after the reperfusion, had an LVEF of ≤ 45% measured by post-catheterism ventriculography and/or echocardiogram.
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Figure 1. TRACIA study design flowchart.

Exclusion criteria. Excluded from the study were patients with previous Q-wave infarction, cardiogenic shock, suspicion or evidence of mechanical complications due to the infarct, a history of sustained ventricular tachycardia or ventricular fibrillation after the first 24 hours of the infarct, treatment during the 4 previous weeks with any drug under research, current use of antineoplastic and/or immunosuppressor drugs, history of cancer in the last 5 years, women in fertile age unless they had a negative pregnancy test, previous malignant hemopathy, known renal failure with creatinine >2.6 mg/dL, vascular cerebral accident of any type in the last year, and any disease that might affect the survival of the patient during the time of the study.

Technique to obtain the bone marrow. Extraction of stem cells was performed in the hemodynamics study room, with the patient in ventral decubitus, under conscious sedation and analgesia. Along the whole procedure, the patient was kept under surveillance by means of non-invasive arterial pressure monitoring, electrocardiogram, and O₂ saturation measurements. Extraction was achieved by repeated punctures in the posterior iliac crest with a puncturing needle connected to a 20 mL syringe, after skin asepsis with an iodine-base solution (Isodine®). In each puncture, an average of 5 to 10 mL was aspirated. After each puncture-aspiration, the needle was washed with saline solution and heparin, repeating the process until 80 to 100 mL of bone marrow was obtained. The sample was placed in a special 100 mL bag for bone-marrow collection with a double filtration system, and containing 10 mL of saline solution and 1 mL of 1% non-fractionated heparin. The bag was transported in fresh to the National Center of Blood Transfusion.

Processing and collecting of stem cells or hematopoietic progenitor cells (HPC). The obtained sample was processed immediately at its arrival to the cell therapy area following the protocol to separate bone marrow-derived mononuclear cells, using a Ficoll® density gradient with the CS-900 kit in the automated Sepax® S-100 system (Biosafe America Inc, USA). The initial documents included request for the processing of hematopoietic progenitor cells (HPC), clinical summary of the patients, demographic data, and informed consent. The total volume of the obtained bone marrow was calculated, and captured in the TESI® Hemodata database, to be assigned a specific bar code; and samples were obtained for the initial blood count. During collection of mononuclear cells with the Sepax® equipment, 100 mL of pyrogen-free sterile Ficoll® and a washing buffer solution (2.5% albumin) were used. The process lasted approximately 1 hour and 15 minutes. Once the process ended, a bag with a fixed volume of 50 mL of mononuclear cells (MNC) was obtained. From this bag, samples were taken to perform a final blood count, clonogenic cultures (STEMCELL Technologies, USA), to determine CD34+, CD35+, and cell viability through flow cytometry (FACScalibur, Bektion Dickinson, USA). From the residual washing bag, 20 mL were taken to perform aerobic and anaerobic microbiological cultures (BacTALERT, Biomeriux, USA). Cells recovered with the automated process corresponded in average to: MNC, 56%; CD34+, 99.7%; and platelet depletion, 88%; erythrocytes depletion, 99.7%; and granulocytes depletion, 99.7%; and platelet depletion, 66%. The final dose was adjusted from 1.0 to 2.0 × 10⁶ CD34+ in 20 mL of buffer. HPC were transported in fresh for their immediate infusion.

Intracoronary infusion of stem cells or HPC. This procedure was performed between the 6th and 7th day after the primary PTCA according to conventional angioplasty techniques. That is, the femoral artery was punctured with a 6 Fr system and the artery responsible for the infarct was cannulated using the corresponding guiding catheter. Before performing the basal shot with contrast medium, 200 µg of nitroglycerine were administered. Once the responsible artery had been identified, a hydrophilic 0.014” angioplasty guide was advanced from 300 cm until the distal portion of the vessel and at that moment a dose of non-fractionated heparin was administered according to the weight of the patient. Afterwards, an over-the-wire balloon catheter was introduced until the site of the implanted stent (relation balloon-stent, 1:1), insufflating the ba-
loon at 2 or 3 atmospheres (atm). At this moment, the angioplasty guide was removed, infusion of stem cells was started at a rate of 1 mL per min during 2 minutes, and then the balloon was deflated for 1 min. This procedure was repeated until completing the total dose of stem cells (approx. 10 mL to 13 mL).

**Magnetic resonance protocol.** The used scanner was a 1.5-Tesla Sonata optimized for the heart. The study is initiated with localizing images, followed by orthogonal sections in the three planes in the HASTE (half-acquisition turbo spin echo) sequence as reference for the acquisition of T2-weighted high resolution images, obtaining 16 images of a single pulse of activity by means of the semi-Fourier reconstruction. The functional morphological analysis included 4- and 5-chamber images in gradient plane echo movie, showing the outflow tract of the left and right ventricles. In each projection, at least one cardiac cycle is captured (R-R interval of the electrocardiogram) with a fast Turbo-FLASH sequence. To evaluate myocardial viability, intravenous gadolinium was administered, acquiring immediate images in two cameras and short axis, named “first pass”. Fifteen to twenty minutes thereafter, images were acquired in four cameras, short axis, and two cameras in an inversion-recovery sequence in search of delayed reinforcement.

**SPECT protocol.** A Siemens Orbiter 2000 gamma camera provided with an ICON A/P processor and the quantitative Gated-SPECT software Cedars-Sinai was used. We used the one-day technetium-99m sestamibi (99mTc) protocol. The SPECT images were obtained 45 min to 60 min after the administration of 10 to 15 mCi at rest and 20 to 30 mCi after effort. To perform the radioisotope ventriculography in equilibrium, we used the in vitro labeling of erythrocytes technique with 99mTc (ultra Tag-99mTc Mallinckrodt) at a 30 mCi dose, in a General Electric ELGEMS-Millenium MPS gamma camera.

**Statistical analysis.** Continuous variables are presented as means and standard deviation (SD), and categorical variables are expressed as frequencies and percentages.
Discussion

The TRACIA (for its initials in Spanish; autologous transplant of bone marrow-derived stem cells in the acute myocardial infarct with ST segment elevation) study represents the first clinical institutional experience in the setting of acute STEMI. In this randomized, two-institutional study, we report the first eight cases with an angiographic and clinical follow-up by means of SPECT and MRI at 6 months, and illustrate, essentially, the feasibility and safety of performing a stem cells study at our institution.

Regarding the ideal time for the intracoronary injection of stem cells, we chose days 5 and 7, as there is evidence that during the first 5 days of the acute STEMI the inflammatory process is more intense and stem cells could easily be destroyed. It has been recommended that the ideal time for the transplant is between days 5 and 14 after presentation of the infarct.15-17 Regarding safety of the procedure, there were no complications during either bone marrow extraction or the intracoronary injection of stem cells, as has been reported previously.

The mechanism of action of stem cells in the treatment of infarcts is still uncertain and can be multifactorial. Most published data derive from experimental studies, which have suggested that the beneficial effect is exerted through the activation of the stem cells residing in the myocardium. This activation is accomplished through paracrine mechanisms. In fact, it has been demonstrated in mice that bone marrow-derived stem cells transplanted into an infarcted myocardium, after a certain time, lose their hematopoietic characteristics and acquire cardiogenic and endothelial lineages to form functional cardiomyocytes and vascular structures.18

Although the evidence published until now suggests that transplantation of bone marrow-derived stem cells can be clinically relevant, studies with large number of patients are required, as well as new experimental models able to elucidate the biology of these cells and their action mechanism in the infarcted myocardium.

Conclusions

The initial experience at the Instituto Nacional de Cardiología Ignacio Chávez in the treatment of acute STEMI by means of autologous transplantation of bone marrow-derived stem cells is encouraging. Implementation was possible in the first eight patients with no complications.

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References