Durable Benefits of Cellular Postconditioning: Long-Term Effects of Allogeneic Cardiosphere-Derived Cells Infused After Reperfusion in Pigs with Acute Myocardial Infarction

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Background—Infusion of allogeneic cardiosphere-derived cells (allo-CDCs) postreperfusion elicits cardioprotective cellular postconditioning in pigs with acute myocardial infarction. However, the long-term effects of allo-CDCs have not been assessed. We performed a placebo-controlled pivotal study for long-term evaluation, as well as shorter-term mechanistic studies.

Methods and Results—Minipigs underwent 1.5-hour mid-left anterior descending balloon occlusion followed by reperfusion and were randomized to receive intracoronary allo-CDCs or vehicle 30 minutes postreperfusion. Left ventriculography (LVG) demonstrated preserved ejection fraction (EF) and attenuation of LV remodeling in CDC-treated pigs. Pigs underwent cardiac magnetic resonance imaging (MRI) and LVG 1 hour and 8 weeks after therapy to evaluate efficacy. MRI showed improvement of EF and attenuation of LV remodeling immediately after allo-CDC infusion. In addition, allo-CDCs improved regional function and decreased hypertrophy 2 months post-treatment. Histological analysis revealed increased myocardial salvage index, enhanced vascularity, sustained reductions in infarct size/area at risk and scar transmurality, and attenuation of collagen deposition in the infarct zone of allo-CDC-treated pigs at 2 months. Allo-CDCs did not evoke lymphohistiocytic infiltration or systemic humoral memory response. Short-term experiments designed to probe mechanism revealed antiapoptotic effects of allo-CDCs on cardiomyocytes and increases in cytoprotective macrophages, but no increase in overall inflammatory cell infiltration 2 hours after cell therapy.

Conclusions—Allo-CDC infusion postreperfusion is safe, improves cardiac function, and attenuates scar size and remodeling. The favorable effects persist for at least 2 months after therapy. Thus, cellular postconditioning confers not only acute cardioprotection, but also lasting structural and functional benefits. (J Am Heart Assoc. 2016;5:e002796 doi: 10.1161/JAHA.115.002796)

Key Words: allogeneic transplantation • cardioprotective effect • cardiosphere-derived cells

Decades of work have attempted to limit acute myocardial infarction (AMI)-related heart damage. Though numerous interventions work when applied before or during ischemia or in the immediate reperfusion period, nothing seems to work after a delay of 30 minutes or more. A notable exception is the phenomenon of cellular postconditioning: Administration of allogeneic cardiosphere-derived cells (allo-CDCs) after reperfusion reduces infarct size, prevents microvascular obstruction, and attenuates adverse remodeling of the left ventricle (LV). The use of 48-hour endpoints in the initial description of cellular postconditioning demonstrated that the benefits were attributable to true cardioprotection, rather than to regeneration, which plays out over several weeks. However, a cardioprotective strategy would need to produce lasting benefits in order to be clinically useful, and the long-term consequences of cellular postconditioning have yet to be investigated.

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Accompanying Videos S1 through S4 are available at http://jaha.ahajournals.org/content/5/1/e002796/suppl/DC1

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Here, we sought to test the long-term safety and efficacy of allo-CDC administration postreperfusion in pigs with AMI. We also probed the mechanism of the cardioprotective effect.

**Methods**

All animal studies were performed in an American Association for Accreditation of Laboratory Animal Care–accredited facility with approval from the institutional animal care and use committee of the Cedars-Sinai Health System. Thirty-two Yucatan minipigs were studied, in either a placebo-controlled pivotal trial (n=28; Figure 1A) or in a short-term mechanistic study (n=6; Figure 1B). Allo-CDCs were grown from a freshly explanted Sinclair male minipig heart.

**Culture of Porcine CDCs**

We obtained allogeneic CDCs from a Sinclair minipig heart from Sinclair Bio Resources, LLC (Columbia, MO), for the placebo-controlled pivotal study and the mechanism assessment study to create a master cell bank (MCB). The heart arrived on ice submerged in cardioplegic solution. Hearts were grossly dissected, and biopsy-sized pieces (~25 mg) were seeded to create explant-derived cells (EDCs). After ~14 days, EDCs were harvested to create an MCB. MCB vials were thawed and cultured as cardiospheres (CSPs) in suspension culture. CSPs were grown on Ultra Low Cell STACK vessels (Corning Life Sciences, Tewksbury, MA). allo-CDCs were grown by seeding CSPs on Nunc Triple Flasks (Thermo Fisher Scientific, Waltham, MA) and passaging when confluent. allo-CDCs were resuspended (1.25 M/mL for a total dose of 12.5

![Diagram](https://via.placeholder.com/150)

**Figure 1.** Study protocols. Placebo-controlled pivotal trial (A) and mechanistic study (B). AMI indicates acute myocardial infarction; CDCs, cardiosphere-derived cells; CK, creatine kinase; ECG, electrocardiogram; LVG, left ventriculography; MRI, magnetic resonance imaging.
Durable Benefits of Cellular Postconditioning

Kanazawa et al

millions of CDCs in 10 mL in CryoStorCS10 (BioLife Solutions, Inc., Bothell, WA) in cryobags (PL07 PermaLife Bags; Origen Biomedical, Inc., Austin, TX), placed directly in a CryoMed controlled-rate freezer, and then transferred to liquid nitrogen. CDCs were thawed at the day of the infusion. Upon thawing, 1 mL of heparin (100 USP units/mL) and 0.1 mL of nonglycerin (50 μg/mL) were added as diluents for a total 10-mL dose for administration. Cell-dose preparation was performed by Capricor, Inc. (Beverly Hills, CA).

Myocardial Infarct Creation and CDC Infusion

On day 0, animals were premedicated with ketamine 20 mg/kg intramuscularly (IM), atropine 0.05 mg/kg IM, and acepromazine 0.25 mg/kg IM. Animals were subsequently induced with propofol 2 to 4 mg/kg intravenously (IV) to effect, intubated, and maintained on isoflurane 2% to 3%. Amiodarone 10 mg/kg IV loading dose, then 0.5 mg/kg IV, as needed, and lidocaine 0.03 μg/kg/min were given for ventricular arrhythmias. Heparin 100 IU/kg IV was given for anticoagulation. AMLs were created in adult Yucatan minipigs by injection of an angioplasty balloon in the mid-left anterior descending artery (LAD) for 1.5 hours. For the placebo-controlled pivotal study, a total of 28 minipigs received placebo or allo-CDCs. allo-CDCs were formulated, frozen, and count-verified just before intra-coronary infusion, which was performed 30 minutes postreperfusion by an over-the-wire balloon catheter, placed in the mid-LAD. Cells were administered in 3 equally divided cycles of balloon inflation separated by 3 minutes of deflation (Figure 1A). For the mechanism assessment study, intracoronary infusion of allo-CDCs in thawed bags (exact cell number was 7.5 million in 10 mL) was performed 30 minutes post-MI (myocardial infarction) using a standard clinical over-the-wire angioplasty balloon (TREK, Abbott Vascular, Abbott Park, IL) in 3 boluses of 3.3 mL with intermediate washes of 2 mL between each bolus to ensure complete cell delivery.

Left Ventriculography

To measure global LV function and volumes, left ventriculography (LVG) was performed before infusion and before sacrifice on day 56 using a pigtail catheter inserted retrogradely into the LV, with imaging in the 40-degree left anterior oblique projection (30 frames per second) using nonionic contrast (27≈30 mL/3 sec). The value of LV volume should be matched for indexed to body surface area (BSA), BSA can be calculated as per the formula (≈0.121 × BW kg0.575).4

Cardiac Magnetic Resonance Imaging

All magnetic resonance imaging (MRI) studies were performed on a 3.0T clinical scanner (Siemens MAGNETOM Verio; Siemens Healthcare GmbH, Erlangen, Germany). An early post-treatment cardiac MRI was performed ≈60 minutes postinfusion of either allo-CDCs or placebo. Typical in-plane resolution was 1.3×1.3 mm, and slice thickness was 6 mm, with no gaps. Global LV function, regional systolic thickening, and regional end systolic thickness were assessed using retrospectively (ECG)-gated and breath-held cine steady-state free precession acquisitions.3 Follow-up MRIs were performed 2 months post-AMI. LV ejection fraction (EF), volumes, global function and regional function were assessed. Contrast-enhanced MRI was performed to assess area at risk (AAR; acutely)5 and scar mass (at 2 months).3

Minipigs undergoing baseline MRI subsequently received an IV injection of gadolinium (Gd)-based contrast agent gadoversetamide (0.2 mmol/kg body weight; OptiMARK; Covidien Imaging Solutions, Hazelwood, MO), and, 8 minutes later, delayed contrast-enhanced images were acquired with an prospectively (ECG) gated, breath-hold, interleaved, two-dimensional Turbo FLASH sequence. Inversion time was adjusted by the technician performing the scan to null signal from noninfarcted remote myocardium.

Image analyses related to tissue viability, that is, scar mass, viable myocardial mass (total mass minus scar mass), relative scar size (scar mass divided by total mass), and regional function, were performed using a validated imaging processing software (QMass MR; Medis Medical Imaging Systems BV, Leiden, The Netherlands).

For transmurality calculation, delayed enhancement images were used to quantify the infarct’s endocardial length (length of the endocardium that is occupied by scar), scar thickness, and wall thickness. The center of the scar on the endothelial border was noted, and scar thickness and wall thickness were measured at this point. Then, the same measurements were performed at half the distance between the center of the scar and the lateral edges of the scar, as previously described.6 Scar transmurality was calculated as the percentage of scar thickness over wall thickness. All values (3 values per each MRI slice containing scar) were averaged for each animal.

Regional function was evaluated by dividing each of the short-axis slices at end-systole into 6 segments. The posterior right ventricular insertion was used as a reference point. Scar size from late Gd-enhanced cardiac MR images was defined based on full width at half maximum to delineate scarred myocardium. Global function was reevaluated using ARGUS software (Syngo Multiple Modality Workplace, version 40A; ARGUS Software, Houston, TX).

Safety Evaluation (Coronary Flow and Arrhythmia)

Coronary flow of the LAD was evaluated utilizing the classical Thrombolysis In Myocardial Infarction (TIMI) grading system.
pre- and postcell infusion according to 4 grades of flow, as described. ECGs were recorded before induction of AMI, before CDC/vehicle infusion, immediately after CDC/vehicle infusion, day 2, 7, and 28 after CDC/vehicle infusion, and before sacrifice on day 56. In addition, ECGs were continuously monitored during intracoronary infusion for arrhythmias.

**Histopathological Evaluation**

Minipigs were euthanized and hearts explanted and sectioned into 1-cm-thick short-axis slices to measure infarct size (IS). Then, transverse ventricular slices were incubated with 2% TTC (2,3,5-triphenyl tetrazolium chloride) for 20 minutes at 37°C to stain viable myocardium. TTC stains viable myocardium brick red, whereas the infarct appears as non-TTC-stained tissue (white/yellow). Each slice was imaged digitally; infarcted areas were determined by manual tracing by a researcher blinded to treatment allocation (Adobe Photoshop CS2; Adobe Inc, San Jose, CA). Myocardial salvage weight percent of AAR or LV (ImageJ, version 1.45s; National Institutes of Health [NIH], Bethesda, MA). Myocardial salvage index was calculated by formula with AAR-IS/AAR, as described previously. Then, 8-μm sections from myocardial samples (fixed in 10% formalin, paraffin-embedded) were obtained from the infarct, border, and remote zones for immunostaining.

To measure myocyte cross-sectional area, sections underwent immunostaining for α-sarcomeric actinin (ab9465; Abcam, Cambridge, MA) and wheat-germ agglutinin (Molecular Probes, Grand Island, NY) (to visualize cell borders). Vascular density was quantified in pigs sacrificed 48 hours postreperfusion. A total of 3 to 5 sections obtained from the border zones were evaluated per heart with immunostaining for Isolectin (Invitrogen I21411; Invitrogen, Carlsbad, CA) and α-smooth muscle actin (α-SMA; ab5694; Abcam). Masson's trichrome staining (Sigma-Aldrich, St. Louis, MO) was performed on serial sections for tissue collagen quantification.

To measure apoptotic cardiomyocytes, sections from myocardial samples underwent immunostaining for terminal deoxynucleotidyl transferase dUTP nick end labeling (TUNEL; Roche 12156792910; Roche, Indianapolis, IN) and α-sarcomeric actinin (ab9465; Abcam). For inflammatory infiltration, CD3 (ab16669; Abcam), CD45 (A1222GA; AbD Serotec, Oxford, UK), CD68 (MAB1715; Abnova, Taipei City, Taiwan), CD206 (M2 marker; sc-48758; Santa Cruz Biotechnology, Santa Cruz, CA), and inducible nitric oxide synthase (iNos; M1 marker; NBP1-50606; Novus, Saint Charles, MO), and rhodamine phalloidin (R415; Invitrogen) were used for immunostaining. Alexa Fluor–conjugated secondary antibodies (Molecular Probes, Eugene, OR) were used, and counterstaining with 4′,6-diamidino-2-phenylindole (DAPI; Molecular Probes) was performed. Sections were imaged using a confocal laser scan microscope (Leica Microsystems, Wetzlar, Germany) and images were processed by Leica Application Suite software. Border (peri-infarct) zone was defined as the region at the edges of the scar (comprising areas of both viable and scarred myocardium).

To evaluate possible immune responses in the heart, hematoxylin and eosin (H&E) staining was performed in sections obtained from the infarct, the border, and the remote LV myocardium. Sections were analyzed by independent pathologists blinded to treatment allocation. Analysis was graded according to the International Society for Heart and Lung Transplantation (ISHLT) guidelines, a system used in clinical practice to grade cardiac allograft rejection.

**Histocompatibility**

To assess histocompatibility, low-resolution swine leukocyte antigen typing was performed on the donor Sinclair minipig and all recipient Yucatan minipigs, as described. Divergent pig strains were used to stimulate the typical allogeneic conditions in human transplantation where donor and recipient are frequently mismatched with regard to human leukocyte antigen (HLA). Complete mismatch of HLA was confirmed if there was at least one Class I or one Class II antigen mismatch at the allele group level between donor and recipient.

**Circulating Donor-Specific Antibodies**

In the placebo-controlled pivotal study, to evaluate humoral immune response, recipient pig serum was obtained at baseline and day 2, 7, 28, and 56 postinfusion. Serum was screened for circulating antidonor immunoglobulin G (IgG) antibodies utilizing flow cytometry at each point.

**Blood Examination**

In the placebo-controlled study, complete blood count, chemistry panel, C-reactive protein, and erythrocyte sedimentation rate were analyzed before induction of AMI, day 2, 7, and 28 after CDC/vehicle infusion, and before sacrifice on day 56. In the mechanism assessment study, blood samples were taken at baseline, before infusion, and 30, 60, 90, and 120 minutes postinfusion to measure creatinine kinase (CK)-MB as a cardiac injury.

**Statistical Analysis**

Results both in text and in figures are presented as mean±SEM, unless specified otherwise. Comparisons between any 2 groups were performed using the 2-tailed unpaired Student t test for comparison between placebo and
treatment groups, the paired t test for comparison between baseline and endpoint, and repeated-measures ANOVA for comparison of data measured serially in time in the 2 groups. Differences were considered statistically significant when $P<0.05$.

Results

Adverse Events and Mortality

The placebo-controlled pivotal trial (Figure 2) and the mechanistic study were performed contemporaneously. Two of 34 pigs (5.8%) died because of ventricular arrhythmia during myocardial infarction (MI) creation. Three pigs were excluded because of technical failure or incomplete MI creation. There was no further mortality (0%), nor were there any adverse events during the infusion procedure in either study. In the pivotal trial, 4 and 3 animals died in the perioperative period in placebo- and allo-CDC-infused groups, respectively. One pig was excluded because of death during anesthesia for a blood draw on day 28. One pig in placebo and 3 pigs in allo-CDC-treated groups were excluded with EF $\geq 55\%$ in the 1-hour MRI, per prespecified exclusion criteria. Thus, a total of 11 pigs completed the placebo-controlled pivotal trial. There was no further mortality, nor were there any other adverse events in the mechanistic study.

Placebo-Controlled Pivotal Trial

We performed the pivotal trial with animals randomized to placebo or allo-CDC groups. The allo-CDC group mimicked the clinical situation in which a preformulated bag is acutely thawed for infusion at the site of reperfusion. From cell counting before infusion, actual delivered cell dosage was estimated at 7.58 million, with postthaw cell recoveries of 75.9±5.9% and cell viability of 89.8±2.7% ($n=6$; Table 1).

Long-Term Functional Efficacy of Allogeneic CDCs Assessed by LVG

To assess the functional efficacy of allo-CDCs, we measured cardiac function and volumes by LVG. Unlike MRI, LVG could logistically be used post-AMI, but before treatment, to establish the baseline value of function and thus to confirm the fidelity of randomization. Indeed, EF is comparable at baseline (43–44%) in the 2 treatment groups (Figure 3B). Representative ventriculograms at 2 months post-MI show amelioration of LV anterior wall motion in the CDC-treated group.
heart compared to placebo (yellow arrows indicate akinetic region; Figure 3A), but there was no difference in absolute EF between 2 groups at 2 months. Although the EF from baseline post-MI to endpoint (2 months) in the placebo group was decreased \( (P < 0.05) \), EF in the CDC-treated group was not significantly changed (Figure 3B). Left ventricular end-diastolic volume index (LVEDVI) at endpoint was higher in placebo than in CDC-treated pigs \( (P < 0.05; \text{Figure 3C}) \). In addition, paired left ventricular end-systolic volume index (LVESVI) shows that placebo \( (P < 0.01) \) underwent more LV dilatation \( (P < 0.05) \), with significantly greater LV end-systolic volumes at end-point \( (P < 0.05; \text{Figure 3D}) \). Thus, infusion of allo-CDCs preserved global cardiac function and minimized adverse remodeling.

### Functional Benefit After allo-CDC Infusion Assessed by Cardiac MRI

To assess detailed cardiac function and tissue characteristics, we performed cardiac MRI \( \approx 60 \) minutes postintervention and at endpoint. By the time of the first MRI, EF was already significantly better in allo-CDC-treated pigs than in placebo (Figure 4A and 4B; Videos S1 through S4). Likewise, LV dilatation was significantly attenuated in allo-CDC pigs both at end-diastole (Figure 4C) and end-systole (Figure 4D) \( 1 \) hour post-treatment. These differences between placebo- and allo-CDC-infused groups cannot be attributed to baseline differences pretreatment (Figure 3). Thus, the functional and structural benefits of cellular postconditioning are apparent with little delay. By 2 months of follow-up, all the differences of indices are less than in the short term and there were no significant differences in the 2 groups. Nevertheless, regional wall motion and percent thickening are greater in CDC-treated pigs compared to placebo at 2 months (Figure 4F and 4G). Taken together, these data indicate that allo-CDCs preserve cardiac function and prevent acute LV remodeling promptly after CDC infusion. Furthermore, allo-CDC infusion improves regional wall motion in the chronic phase.

### Assessment of Cardiac MRI With Structural Effect After allo-CDC Infusion

Gd contrast enhancement in the acute phase of MI is not a reliable measure of scar,\(^1\)\(^5\) but it does correlate with AAR (the borders of AAR appear bright, but regions of microvascular occlusion in the center exclude Gd and remain dark; Figure 5A, left panels, yellow arrows). The volume of myocardium bounded by early Gd-enhancement was equal in placebo- and CDC-treated pigs at 1 hour (pooled data in Figure 5B), consistent with the notion that AARs are identical in the 2 groups. Two months later, MRI analysis showed that the zone of late Gd enhancement, a reporter of scar size (Figure 5C), was not significantly different in the 2 groups at 2 months. However, total LV mass was lower in allo-CDC-treated animals relative to placebo (Figure 5D), presumably reflective of a decrease in compensatory hypertrophy; likewise, scar transmurality was lower in CDC-treated hearts (Figure 5E). All of the conclusions regarding tissue

### Table 1. Summary of Coronary Flow and Dosing Data in Placebo- and CDC-Treated Pigs

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<th>Pre</th>
<th>Post</th>
<th>Cell Dose (M)</th>
<th>Cell/kg</th>
<th>Recovery (%)</th>
<th>Viability (%)</th>
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BW indicates body weight; CDCs, cardiosphere-derived cells; N/A, not applicable; TIMI, Thrombolysis in Myocardial Infarction.
characteristics at 2 months were subjected to histological validation, as discussed next.

**Histological Assessment of Scarred and Viable Myocardium**

Immediately after the 2-month MRIs, hearts were removed and processed for histology. Figure 6A shows representative slices stained with TTC to quantify infarcted area. The placebo heart has a thin infarcted wall in which the scar is mostly transmural, whereas the heart treated with CDCs has viable myocardium surrounding the scar with a visibly thicker infarcted wall. Pooled data show that scar transmurality was decreased in CDC-treated hearts (Figure 6B), confirming the MRI results (Figure 5E). Myocardial salvage index was also higher in the CDC group (Figure 6C). Absolute infarct size showed a strong trend toward reduction in CDC-treated pigs (Figure 6D), which was significant when normalized to AAR (53.4% in CDC-treated vs 69.3% in placebo; Figure 6E). In addition, the remote/LV area ratio is higher in placebo- than in CDC-treated hearts (Figure 6F), which suggests that placebo hearts developed hypertrophy in the remote myocardium (consistent with the MRI analysis; Figure 5D). To verify this conjecture, we evaluated cardiomyocyte morphology by microscopic analysis. Myocyte cross-sectional area was smaller in CDC-treated pigs than in placebo (Figure 6G and 6H). Thus, one of the key long-term benefits of cellular postconditioning is a decrease in compensatory myocardial hypertrophy.

**Histopathological and Serum Evaluation of Intracoronary Infusion of allo-CDCs**

In order to investigate the effect of CDCs on vessels, we evaluated immunohistochemistry with isolectin and α-SMA.
Vessel density in the IZ was higher in the CDC-treated group (Figure 7A and 7B); meanwhile, fibrosis was attenuated, as is evident from representative Masson’s trichrome–stained sections from the infarcted wall (Figure 7C and 7D). In addition, quantification of apoptosis revealed fewer TUNEL-positive myocytes in the border zone in CDC-treated hearts than in placebo (Figure 7E and 7F), consistent with our previously reported findings at an earlier time point.18 Taken together, these results indicate that CDCs promote vessel formation and attenuate collagen deposition and cardiomyocyte apoptosis.

Figure 4. Allogeneic CDCs exhibit improved global function and remodeling 1 hour after cell treatment compared to placebo, as assessed by MRI. Matched cine short-axis images (at end-diastole [ED] and end-systole [ES]) at 1 hour and 2 months post-treatment for pigs treated with placebo and allogeneic CDCs (A). Comparison of ejection fraction (B), LVEDVI (C), and LVESVI (D) at 1 hour and 2 months in placebo- and CDC-treated pigs. Endocardial (red) and epicardial (green) contours of the LV are shown. Cardiac slices were divided into 6 segments to calculate regional wall motion and thickening in infarct and remote myocardium at 2 months (E). Regional wall motion (F) and percent thickening (G) in infarct and remote myocardium in placebo and CDC groups at 2 months. CDCs indicates cardiosphere-derived cells; EF, ejection fraction; LVEDVI, left ventricular end-diastolic volume index; LVESVI, left ventricular end-systolic volume index; MRI, magnetic resonance imaging. Values are means±SEM. *P<0.05 and **P<0.01 between 2 groups.

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Immune Reaction to allo-CDCs in Pivotal Study

To evaluate immune and inflammatory reactions to cellular postconditioning, we first checked the peripheral leukocyte count and differential at 48 hours. Leukocyte and lymphocyte counts were equal before ischemia, but were attenuated in CDC-infused animals compared to placebo 48 hours post-MI (Figure 8A and 8B). Serum neutrophil and monocyte fractions exhibited a similar trend (Figure 8C and 8D). This apparent attenuation of inflammation occurred despite the fact that there was complete immunological mismatch for swine leukocyte antigens between donor and recipient pigs (Table 2). No circulating alloreactive IgG antidonor antibodies were detected in any recipients of allo-CDCs at any time point (Figure 8E and 8F). Histopathology demonstrated focal minimal and mild lymphohistiocytic infiltration in the border zone in placebo- and CDC-treated pigs, respectively (Figure 8G). The infiltrating cells were localized within interstitial and perivascular spaces (Grade 1R according to the ISHLT grading system; Figure 8H). Remote myocardium was consistently clear of inflammatory infiltrates.

Probing Mechanism

The results in Figure 4 indicate that functional and structural benefits of cellular postconditioning are evident as early as 1 hour after CDC infusion. We further investigated the time course of cardioprotection by measuring CK-MB level at various time points (Figure 9A). Whereas CK-MB levels were comparable preischemia and at baseline (pretreatment) post-MI, a decrease in the CDC group is evident by 30 minutes post-treatment and persists for at least 2 hours. We therefore selected 2 hours post-treatment as the endpoint for the mechanistic study. Once again, TUNEL-positive myocytes were decreased in the border zone of CDC-treated animals versus placebo (Figure 9C), signifying a decrease in apoptosis. In contrast to the anti-inflammatory response observed systemically (Figure 8A through 8D), the myocardium exhibited no overall changes in CD3- or CD45-positive cell content between placebo- and CDC-treated hearts. Nevertheless, there was a significant increase in the number of CD68+ cells (macrophages; Mφ) in hearts of CDC-treated pigs (Figure 9D and 9E). Mφ have the intrinsic capacity to polarize between M1 (proinflammatory) and M2 (reparative) phenotypes; interestingly, we observed a strong shift toward an M2-like Mφ phenotype (Figure 9F and 9G). Taken together, these data suggest that salutary changes in Mφ polarization from a “proinflammatory” toward a “healing” phenotype may contribute to cellular postconditioning, consistent with more-extensive mechanistic studies in rats and in isolated macrophages.19,20
Discussion

Here, we report on the long-term benefit of allo-CDC therapy immediately postreperfusion in a clinically relevant pig model of AMI. The main findings are: (1) allo-CDC infusion after AMI is clinically feasible and safe without immunosuppression. (2) CDCs prevent LV dilatation acutely (=60 minutes) and attenuate LV remodeling post-AMI chronically (2 months).

Figure 6. Allogeneic CDCs prevent hypertrophy, decrease infarct size, and salvage myocardium, as assessed by postmortem histology. Representative short-axis heart slices of a placebo- and a CDC-treated pig after incubation with TTC. Viable myocardium stains brick red, whereas scarred myocardium appears white (A). Comparison of scar transmurality (B), myocardial salvage index (C), scar mass (D), and percent IS/AAR (E) at 2 months in placebo and CDC groups. Comparison of remote area (as percent of LV) at 2 months in placebo and CDC groups (F). Cardiomyocyte cross-sectional area in the remote zone (G and H). AAR indicates area at risk; AU, astronomical unit; CDCs, cardiosphere-derived cells; IS, infarct size; LV, left ventricle; TTC, 2,3,5-triphenyl tetrazolium chloride. Values are means±SEM. *P<0.05 and **P<0.01 between 2 groups. Scale bar in (A) is 1 cm.
(3) CDCs increase LV wall thickness and improve regional LV wall motion chronically. (4) CDCs preserve myocardium and prevent hypertrophy. (5) CDCs decrease inflammatory response and reduce reperfusion injury. (6) Underlying mechanisms include enhanced angiogenesis, decreased apoptosis, and polarization of Mφ toward a reparative phenotype.

**Optimal Timing of Cell Administration and Advantage of Allogeneic Products**

Here, we have followed our originally described cellular postconditioning protocol, wherein cells are administered 30 minutes postreperfusion. Though this protocol is effective, few data are available regarding the timing window of...
therapeutic opportunity post-AMI. In the rat model, allo-CDC administration 20 minutes, but not 2 hours, post-AMI is cardioprotective. Administration of cells immediately postreperfusion is known to be more effective than delivery 7 days post-AMI in increasing neovascularization and expression of vascular endothelial growth factor in rats and pigs, respectively. Within the early time window, an important consideration is the ability of cells to reach their targets when delivered by the coronary route. Microvascular obstruction (MVO) progressively evolves starting several minutes postreperfusion, reaching a peak 48 hours afterward. Early cell administration not only has the potential to...
mitigate MVO, but also should logically be timed to precede significant MVO so as to maximize cell access to injured tissue. In terms of cardioprotection, myocardium at risk is irreversibly damaged after 3 hours of occlusion, so earlier administration is required for meaningful salvage.\textsuperscript{24} The notion that prompt intervention is optimal is supported by our observations that LV EF is augmented, and ventricular volumes are preserved, very quickly after allo-CDC therapy (Figure 4B through 4D).

The use of allogeneic cells as “off-the-shelf” products is mandatory for prompt administration postreperfusion. Allogeneic cell therapy has the additional advantage that it avoids variations of cell efficacy attributable to patient age or comorbidities, plus the risks and costs of patient-specific tissue harvesting and cell processing. Although immune rejection is a possible disadvantage, allogeneic mesenchymal stem cells\textsuperscript{25} and CDCs (http://clinicaltrials.gov/ct2/show/NCT01458405) have been used in various early-phase clinical trials of heart disease without safety concerns,\textsuperscript{25} consistent with preclinical studies indicating that allogeneic cell therapy without immunosuppression is safe and effective.\textsuperscript{3,14,20,26} Indeed, our study demonstrates only nonsignificant local inflammatory response and the absence of circulating alloactive antibodies with long-term follow-up after the administration of allo-CDCs.

Long-Term Beneficial Effects of Cellular Postconditioning

The present work demonstrates that prompt infusion of CDCs postreperfusion preserves regional wall motion and wall thickening functionally with long-term follow-up. This reduction of wall stress leads to attenuated LV remodeling, as well as reduced hypertrophy. The attenuation of hypertrophy is important for 2 reasons: First, reduction of hypertrophy is itself a salutary principle, given that post-MI hypertrophy has been associated with poor long-term outcomes\textsuperscript{27}; second, the reduction of hypertrophy in CDC-treated animals confounds the assessment of scar-size reductions, especially when IS is conventionally expressed as a percentage of LV mass.\textsuperscript{28} Thus, correction for AAR, as in Figure 6E, is particularly important.

Mechanisms of Benefit

We now widely recognize that most of the beneficial effects of transplanted CDCs are indirect, and not attributable to canonical stem cell mechanisms. CDCs have been argued to be cardiac progenitor cells,\textsuperscript{29–31} but that feature of CDCs is an epiphenomenon when it comes to mechanism of action. In the extreme, allo-CDCs are cleared completely within several weeks, but their functional and structural benefits persist at least 6 months.\textsuperscript{14} Thus, long-term transplanted cell survival is not required for sustained benefit.\textsuperscript{32} Consistent with the mechanistic data in Figure 9F and 9G, our recent publication in a rat AMI model implicates CDC-secreted exosomes in the mechanism of action, through effects on macrophage polarization.\textsuperscript{20}

The implication of macrophages breathes new life into the long-standing, but much-maligned, concept that inflammation may be a useful target to mitigate reperfusion injury.\textsuperscript{1,33} Inflammation post-AMI is important for healing and influences short- and long-term outcomes.\textsuperscript{34,35} Leukocytosis is a strong predictor of mortality and heart failure in patients

Table 2. Summary of SLA Haplotype

<table>
<thead>
<tr>
<th>Donor Pig</th>
<th>Breed</th>
<th>Sex</th>
<th>Blood Type</th>
<th>Inferred Low-Resolution (Lr) SLA Haplotype</th>
<th>Predicted No. of Allele Mismatch</th>
<th>No. of Antigen Mismatch*</th>
</tr>
</thead>
<tbody>
<tr>
<td>0111</td>
<td>Sinclair</td>
<td>M</td>
<td>Non-A</td>
<td>Lr-2.2/11.12</td>
<td>SLA-1: 0  SLA-3: 2  SLA-2: 2  DRB1: 1  DQB1: 2  DQA: 6</td>
<td>3</td>
</tr>
</tbody>
</table>

Low-resolution SLA genotyping revealed complete Class I (SLA-1, SLA-2, and SLA-3) and Class II (DRB1, DQB1, and DQA) mismatches between the donor Sinclair pig and CDC recipients (Yucatan minipig). CDCs indicates cardiosphere-derived cells; SLA, .

*Maximum number of antigen mismatch at SLA-1, SLA-2, SLA-3, DR, and DQ is typically 10 to 12 (2 alleles at each locus with some haplotypes express duplicated SLA-1 antigen); with 0111 as the donor, the maximum number of antigen mismatch is predicted to be 11 given that both haplotypes express 2 SLA-1 antigens each, whereas Lr-2.2 does not express SLA-3.

†Based on mismatch at DRB1 and DQB1.

‡One allele appears to match at the low-resolution allele-group level, but allelic-level mismatch may be identified by high-resolution typing.
Moreover, a subset of leukocytes (viz., neutrophils and monocytes) are associated with microvascular injury, LV remodeling, and function. In addition, monocyte/macrophage heterogeneity post-AMI is associated with the impairment of myocardial salvage and adverse LV remodeling. Here, we found that peripheral leukocyte counts are lower in CDC-treated animals compared to placebo in the first 2 days postreperfusion. In addition, CDC infusion reduces myocardial injury and apoptosis as soon as 2 hours after cell treatment, at which time changes in macrophage polarization markers iNos (M1) and CD206 (M2) were utilized to differentiate the proportion of infiltrating M1 and M2 macrophages within the border zone tissue 2 hours postintervention. Data are presented as the number of CD206+ vs iNos+ macrophage per field of view (G). Values are means±SEM. *P<0.05 and **P<0.01 between 2 groups. Scale bars are 50 µm in (B, D, and F). CDCs indicates cardiosphere-derived cells; CK, creatine kinase; iNOS, inducible nitric oxide synthase; TUNEL, terminal deoxynucleotidyl transferase dUTP nick end labeling.
polarization are already evident. As tantalizing as it may be, the limited mechanistic information in the present study serves more to generate hypotheses than to test them. The data are therefore large-animal correlates to our recent detailed characterization in a rat AMI model of the mechanistic roles of inflammation in general, and macrophages specifically, in cellular postconditioning.20

Limitations
This study has several limitations. First, infarct size and AAR could not be measured at baseline (after reperfusion, but before therapy) in this protocol. For this reason, the only paired pre-/postanalyses are based on LVG, which is accurate for overall ventricular performance, but provides no tissue-level insights. Second, to produce AMI, we used an iatrogenic balloon occlusion model without any thrombus or atherosclerotic plaque in the absence of comorbidities, such as diabetes, and pigs were pretreated with amiodarone to prevent loss of animals attributable to ventricular arrhythmia. These do not reflect the real clinical situation; indeed, the preadministration of amiodarone has the potential to mask a possible effect of CDCs on arrhythmogenesis,39 and diabetes is known to interfere with some classical forms of cardioprotection.40 In addition, pigs were not receiving medical therapy, such as beta-blockers or renin-angiotensin system inhibitors, which are established as optimal medical therapy for preventing adverse remodeling and progression to heart failure. So, it is unclear whether CDCs would have any incremental benefits above and beyond optimal medical therapy. Finally, our phenotyping ended 2 months after cell treatment in this study. Although our previous work indicates that the benefits of CDCs can last even longer,10,14 more-extended evaluation would be valuable, particularly in terms of hard clinical endpoints, such as progression to heart failure.

Conclusions
Here, we demonstrate that intracoronary infusion of allogeneic CDCs postreperfusion, utilizing strategies consistent with standard clinical practice, reduced infarct size, preserved regional function, and elicited cytoprotection in pigs with AMI. The durable benefits indicate that cellular postconditioning has the potential to improve long-term outcomes post-AMI.

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Disclosures
E. Marbán and L. Marbán own equity in Capricor, Inc. L. Marbán and Kreke are employed by Capricor. Malliaras is a consultant for Capricor, Inc. Other authors have no relationships with industry.

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