Intravascular Stem Cell Bioreactor for Prevention of Adverse Remodeling After Myocardial Infarction

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Background—Prevention of adverse remodeling after myocardial infarction (MI) is an important goal of stem cell therapy. Clinical trial results vary, however, and poor cell retention and survival after delivery likely limit the opportunity to exert beneficial effects. To overcome these limitations, we built an implantable intravascular bioreactor (IBR) designed to protect contained cells from washout, dilution, and immune attack while allowing sustained release of beneficial paracrine factors.

Methods and Results—IBRs were constructed using semipermeable membrane adhered to a clinical-grade catheter shaft. Mesenchymal stem cell (MSC) viability in and paracrine factor release from IBRs were assessed in vitro and IBR biocompatibility and immune protection confirmed in vivo. In a porcine anterior MI model, IBRs containing 25 million allogeneic MSCs (IBR-MSCs) were compared with IBRs containing media alone (IBR-Placebo; n=8 per group) with adverse remodeling assessed by magnetic resonance imaging. Four weeks after MI, IBR-MSCs had no significant change in end-diastolic volume (+0.33±4.32 mL; P=0.89), end-systolic volume (+2.14±4.13 mL; P=0.21), and left ventricular ejection fraction (−2.27±2.94; P=0.33) while IBR-Placebo had significant increases in end-diastolic volume (+10.37±3.84 mL; P=0.01) and ESV (+11.35±2.88 mL; P=0.01), and a significant decrease in left ventricular ejection fraction (−5.78±1.70; P=0.025). Eight weeks after MI, adherent pericarditis was present in 0 of 8 IBR-MSCs versus 4 of 8 IBR-Placebo (P=0.02), suggesting an anti-inflammatory effect. In a separate study, 25 million allogeneic pig MSCs directly injected in the peri-infarct zone 3 days after MI (n=6) showed no significant benefit in adverse remodeling at 4 weeks compared with IBR-MSCs.

Conclusions—MSCs deployed inside an implantable, removable, and potentially rechargeable bioreactor in a large animal model remain viable, are immunoprotected, and attenuate adverse remodeling 4 weeks after MI. (J Am Heart Assoc. 2019;8:e012351. DOI: 10.1161/JAHA.119.012351.)

Key Words: cytokines • growth factors • myocardial infarction • remodeling heart failure • stem cell

Administration of stem cells following myocardial infarction (MI) promotes regeneration of cardiac tissue, decreases inflammation-associated cell loss, and thereby limits adverse remodeling, which is associated with the subsequent development of heart failure and associated morbidity and mortality.1,2 The effects of stem cell administration on adverse remodeling in many clinical trials, however, are modest.3–6 One reason may be that only a small percentage of cells delivered to the heart using current methods remain there, and of those that do, few survive for any significant time.7,8 There is growing evidence from preclinical studies that many of the positive effects of stem cell therapy result from their release of growth factors and cytokines, collectively referred to as “paracrine factors” (PFs), which enhance endogenous repair mechanisms and reduce maladaptive inflammation and apoptosis after tissue

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Clinical Perspective

What Is New?

- This study tested a novel intravascular bioreactor that was designed to house a large number of stem cells in a protected space within the body and allow free release of soluble paracrine factors (growth factors, cytokines, etc), but not of cells, and in doing so to determine whether this method could limit adverse remodeling after myocardial infarction in a large animal model.

What Are the Clinical Implications?

- The results of this study indicate that it is not necessary to directly inject stem cells in the heart to elicit a beneficial effect and suggest a new method by which adverse remodeling after myocardial infarction, which is known to be an important pathologic mechanism in the development of heart failure, may be prevented or reduced.

Methods

Data and study materials will be made available to other researchers for purposes of reproducing the results or replicating the procedures. Interested researchers may contact the corresponding author.

MSC Isolation and Culture

The in vitro and the preliminary in vivo IBR experiments were conducted using human bone marrow–derived MSCs (hMSCs; Lonza, Walkersville, MD). The in vivo study of IBR therapy in MI was conducted using bone marrow–derived porcine MSCs (pMSCs) isolated from bone marrow aspirates of Yorkshire swine (15–20 kg) using established techniques. The pMSCs were analyzed for expression of MSC-specific surface markers including CD90, CD44, CD34, and CD45 using a BD LSRII flow cytometer (BD Biosciences, San Jose, CA). Data were gated and analyzed using FlowJo software (Tree Star, Ashland, OR). Osteogenic and adipogenic differentiation assays were performed as described previously. Briefly, pMSCs were plated in 6-well plates at 30 000 cells/well (osteogenic assay) and 200 000 cells/well (adipogenic assay). On day 1, cells were treated with specific induction media. For the osteogenesis assay, the media were changed every 3 days, and calcium deposition was quantified at 2 weeks. For the adipogenesis assay, lipid accumulation was quantified following 3 cycles of induction and maintenance as described previously.

Media

In culture, both hMSCs and pMSCs were grown in media composed of: Alpha-MEM (Mediatech, Manassas, VA) supplemented with 20% fetal bovine serum (HyClone, Logan, UT), L-glutamine (350 μg/mL; Mediatech), and penicillin/streptomycin (50 IU/mL/50 μg/mL; Corning Mediatech, Manassas, VA). These media were used for all experiments, including in vitro assessment of PF release and in vivo testing of IBR biocompatibility and postinfarction efficacy.

Bioreactor Construction

The IBR (Figure 1) was constructed from a semipermeable cellulose ester membrane with a molecular weight cutoff of 100 kDa (Spectrum Labs, Rancho Dominguez, CA). A 10-cm membrane segment was attached to a 20-cm modified 6-French vascular catheter (Boston Scientific, Marlborough, MA) with a heat-sealed distal end and the shaft fenestrated at 1-cm intervals using an 18-gauge needle. The membrane was secured to the midportion of the modified catheter using cyanoacrylate medical device adhesive (Henkel, Rocky Hill,
CT), 2-0 silk surgical suture, and 0.125-inch medical grade heat shrink tubing (InsulTab, Woburn, MA). IBRs were sterilized in 70% ethanol under ultraviolet light for 12 hours and washed in phosphate-buffered saline (Corning Mediatech) and media before cell culture experiments and in vivo implantation.

In Vitro IBR PF Release
To assess in vitro PF release, IBRs were loaded with increasing numbers of hMSCs (10⁵, 10⁶, 5×10⁶) and submerged in media in separate flasks for 7 days under standard mammalian cell culture conditions (37°C, 5% CO₂). Samples of media outside the IBRs were collected on days 1, 3, and 7; centrifuged to pellet any debris; and then flash frozen at −80°C. Concentrations of vascular endothelial growth factor (VEGF), basic fibroblast growth factor, hepatocyte growth factor, and interleukin-8 in IBR-conditioned media (CM) were assessed using multiplex ELISA (Quansys, Logan, UT). Bioreactors loaded with media alone (ie, no cells) were used as controls. At 7 days, hMSCs recovered from IBRs underwent viability assessment by Trypan Blue exclusion. To assess PF-release dose response, 6 additional CM experiments were conducted with IBRs loaded with 10⁷, 2.5×10⁷, and 4.5×10⁷ hMSCs. Samples of CM collected from outside the IBRs on days 1, 3, and 7 were assessed for VEGF production by ELISA (R&D Systems, Minneapolis, MN).

In Vivo Biocompatibility and Immune Protection
To assess IBR biocompatibility and immune protection in vivo, IBRs containing 10⁶ hMSCs (ie, xenogeneic) were inserted in the right internal jugular vein in Yorkshire swine (20–30 kg) via surgical cutdown and advanced to the junction of the superior vena cava and right atrium under fluoroscopic guidance (n=2). Animals were assessed daily for 1 week for fever, hemodynamic compromise, and dermatologic reaction that would suggest acute rejection. After 7 days, IBRs were removed, washed with sterile phosphate-buffered saline, and immediately submerged in media and placed in culture. As previously done, samples of CM from outside the IBRs were collected at days 1, 3, and 7, and the concentrations of PFs measured via multiplex ELISA. After 7 days in culture, hMSCs recovered from the IBRs were plated on standard tissue culture flasks to assess morphology and growth.

Preclinical Model of MI
Anterior MI was produced in female Yorkshire swine (20–30 kg) using 90-minute catheter-based balloon occlusion of the mid–left anterior descending artery as previously described (n=22).18,19

Baseline MRI
Three days after MI, baseline cardiac MRI with LGE was performed using a whole-body 3.0 T magnetic resonance scanner to assess left ventricular function, end-diastolic volume (EDV), end-systolic volume (ESV), and LGE before implantation of an IBR. Cine images were acquired using a radiofrequency spoiled gradient recalled echo pulse sequence (repetition time, 4.3 ms; echo time, 2.1 ms; flip α=12°; in-plane resolution 1.3×1.3 mm; 30 phases/cardiac cycle). Images for LGE were acquired after peripheral injection of Magnevist 0.2 mmol/kg (Bayer, Wayne, NJ) with an inversion recovery–prepared T1-weighted gradient echo sequence (repetition time, 5.3 msec; echo time, 2.6 msec; 1 R-R interval; flip α=20°; in-plane resolution 1.4×1.5 mm). Short-axis images from base to apex were acquired with 6-mm slice thickness and no gap for cine and LGE images.

Bioreactor Implantation
After baseline MRI, surgical cutdown of the right neck was performed to isolate the internal jugular vein, and an IBR was inserted as described above. Animals were randomly assigned to an IBR loaded with (1) 2.5×10⁷ allogeneic pMSCs suspended in media (IBR-MSC), or (2) media alone (IBR-Placebo; n=8 both groups). For the IBR-MSC group, allogeneic pMSCs at passages 3 to 6 were harvested, counted, and assessed for viability by Trypan Blue exclusion immediately before loading. A total of 2.5×10⁷ viable pMSCs were

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**Figure 1.** Implantable bioreactor (IBR) prototype. The cell suspension is injected into the delivery port, which is connected to the polymeric semipermeable stem cell chamber via a clinical-grade vascular catheter shaft.
suspended in media to a volume of 1.5 mL for infusion into the IBR, followed by 0.5 mL of media to flush cells into the cell chamber. Placebo IBRs were loaded with 2 mL of media alone. Once loaded, the IBR infusion port was closed with a sterile cap, the proximal shaft secured in place with suture, and the wound closed. Animals were then followed for any sign of immune reaction, rejection, or wound dehiscence.

Four-Week MRI and IBR Removal

Four weeks later, repeat cardiac MRI was performed using the same parameters, followed by surgical cutdown to expose and remove the IBR. Explanted IBRs were washed in sterile phosphate-buffered saline, placed in culture flasks submerged in culture medium (175 mL), and maintained in culture for 7 days. CM collected from outside the IBRs on days 1, 3, and 7 were assessed for VEGF release by ELISA for human VEGF (R&D Systems), which is known to cross react with porcine VEGF.

Eight-Week MRI, Hemodynamic Assessment, and Necropsy

A final MRI was performed 4 weeks after IBR removal, that is, 8 weeks after MI. After MRI, invasive hemodynamic assessment was performed using a solid-state transducer (Millar, Houston, TX) followed by coronary angiography. Animals were then euthanized and necropsy performed with removal and inspection of the heart and pericardium, as well as other vital organs (lungs, liver, spleen, and kidneys). Sections of heart tissue were stained with hematoxylin and eosin or Masson’s trichrome to assess scar and vasculogenesis at the infarct border zone.

MRI Image Analysis

MRI images were analyzed using Cinetool (GE Healthcare, Milwaukee, WI) by 2 observers blinded to the group assignment. When present, LGE was quantified as dense “core” and heterogeneous peri-infarct “gray zone” extents using previously published methods. The core zone was composed of all pixels with signal intensity (SI) >50% of maximal SI within the region of LGE. Gray zone extent comprised all pixels with SI greater than peak SI in normal myocardium, but <50% of maximal SI within the hyperenhanced region. Total infarct size was defined as the sum of the “core” and “gray zone” extents.

Comparison Study of Intramyocardial DI of MSCs

To assess the impact of delivery of the same number of pMSCs (2.5 × 10⁷) to the heart via intramyocardial injection, a separate group of animals (n=6) underwent anterior MI induced using the same protocol. Three days later, cardiac MRI with LGE was performed, followed by DI of allogeneic pMSCs in the MI border zone using the NOGA Myostar injection-catheter system as previously described. Briefly, the mapping catheter was advanced through a 9-French introducer sheath and retrograde across the aortic valve. A complete map of left ventricular geometry and function was generated by collecting local position and electrocardiographic data at >50 points in the endocardium. A viable infarct border zone was determined as a unipolar voltage from 6 to 12 mV. A total of 25 million viable pMSCs, divided into 10 injections, each 0.5 mL in volume, were delivered to the endocardium of the border zone. Injection sites were recorded both in the electroanatomic NOGA map and in 2 orthogonal radiographic projections and marked on a tracing of the endocardial silhouette. Animals receiving DI MSCs were followed for 8 weeks using the same MRI LGE imaging protocol as outlined above with results interpreted by readers blinded to the experimental procedure.

All animal experiments were performed under a protocol approved by either the Johns Hopkins University Animal Care and Use Committee (#SW10M468; IBR experiments) or the University of Miami (#14-193; DI experiment).

Statistical Analysis

All data are reported as mean±SEM. Analyses of paired data were performed using the Wilcoxon signed-rank test and unpaired data using the Wilcoxon rank-sum (Mann–Whitney) test. For ratios, outcomes were compared using the Fisher exact test. For comparisons across the 3 groups, analyses were performed with the Kruskal–Wallis rank test with Dunn’s test for post hoc analyses. A P value of ≤0.05 was deemed statistically significant.

Results

In Vitro PF Release

Human MSCs grown in IBRs in vitro showed high viability after 7 days in culture (mean, 75.4±11.6%; n=6). The majority of MSCs adhered to the inner wall of the IBR cell chamber. When 10⁶ hMSCs were housed in an IBR, there was release of relevant PFs, including VEGF, hepatocyte growth factor, basic fibroblast growth factor, and interleukin-8 over 7 days in culture (n=6; Figure 2A). To determine the dose of MSCs used in the in vivo studies, a dose-response experiment performed using escalating doses of hMSCs over 7 days in culture showed VEGF production increased to a dose of 2.5 × 10⁷ cells (Figure 2B). There was no further increase in VEGF production when the IBR was loaded with a higher
number of cells; hence, the dose of $2.5 \times 10^7$ was chosen for the preclinical MI experiment.

### In Vivo Biocompatibility and Immune Protection

To determine whether the IBR was biocompatible and afforded immune protection for contained cells, devices were implanted in adult pigs and loaded with $10^6$ hMSCs (Figure 3A). Despite use of xenogeneic hMSCs, there was no evidence of immune or transfusion reaction. After 1 week in vivo, the IBRs were removed and placed in culture. Analysis of CM collected over the subsequent 7 days showed continued production and release of relevant PFs (Figure 3B).

Cells recovered from the IBRs were plated and showed continued growth and division with normal MSC morphology (Figure 3C).

### Porcine MSC Isolation and Characterization

For the preclinical MI experiments, pMSCs were isolated from bone marrow biopsies obtained from Yorkshire swine. Assessment of surface marker expression confirmed MSC phenotype with markers typical of MSCs: CD90+ and CD44+; CD34− and CD45− (Figure 4A). Isolated pMSCs also showed typical osteogenic (Figure 4B) and adipogenic differentiation (Figure 4C). The pMSCs used for the study were low passage.

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**Figure 2.** Paracrine factor release from implantable bioreactor (IBR) in vitro. Assessment of conditioned media collected from outside IBRs containing $10^6$ human mesenchymal stem cells (MSCs) over 7 days in culture showed production and release of vascular endothelial growth factor (VEGF; green), interleukin 8 (IL-8; red), basic fibroblast growth factor (bFGF; blue), and hepatocyte growth factor (HGF; orange) as assessed by multiplex ELISA (A; log scale; n=6). A subsequent dose-finding experiment using IBRs loaded with increasing doses of hMSCs showed dose-dependent VEGF production up to a dose of $2.5 \times 10^7$ MSCs (B; log scale; n=6).

**Figure 3.** Biocompatibility testing and human mesenchymal stem cell (MSC) survival in implantable bioreactor (IBR) implanted in pigs. IBR implanted via neck with catheter shaft in right internal jugular vein and stem cell chamber at the level of the right atrium (A). Human MSCs recovered from IBR after 1 week in vivo (n=2) show continued production of basic fibroblast growth factor (bFGF; blue), vascular endothelial growth factor (VEGF; green), and interleukin 8 (IL-8; red) by multiplex ELISA (B), as well as normal MSC morphology (C).
Placebo-Controlled Preclinical IBR MI Study

For the randomized IBR study (Figure 5), MI was induced in 22 pigs, 16 (74%) of which survived to the baseline MRI performed 3 days later. Baseline MRI on these 16 animals showed large anteroapical infarcts involving 28.9 ± 9.7% of the left ventricle by mass and significant systolic dysfunction with mean left ventricular ejection fraction (LVEF) of 44.6 ± 9.1%. There were no significant differences on baseline MRI between the 2 groups in terms of infarct size, LVEF, EDV, or ESV (Table). After baseline MRI, an IBR was implanted and loaded with 2.5 × 10^7 pMSCs (IBR-MSC group) or media alone (IBR-Placebo). Bioreactor implantation was well tolerated and without complication. No animals showed evidence of immune rejection or transfusion reaction over the 4 weeks the IBRs remained implanted.

Week 4 Cardiac MRI

Repeat cardiac MRI at 4 weeks after implantation in the IBR-MSC group showed no significant changes in EDV (+0.33 ± 4.32 mL; P=0.89), ESV (+2.14 ± 4.13 mL; P=0.21), or LVEF (−2.27 ± 2.94%; P=0.33) compared with baseline. In contrast, there were significant increases in both EDV (+10.37 ± 3.84 mL; P=0.01) and ESV (+11.35 ± 2.88 mL; P=0.01), as well as a significant decrease in LVEF (−5.78 ± 1.70%; P=0.025) in the IBR-Placebo group (Figure 6). These changes in the IBR-Placebo group are consistent with pathologic left ventricular remodeling typical after large infarcts. On comparison between the groups, the change in ESV was significantly lower in the IBR-MSC group than in the IBR-Placebo group (P=0.036), although not the change in EDV (P=0.059). There was no significant difference between the groups for the change in LVEF (P=0.40).

The total infarct extent, as assessed as the sum of the core and gray zone masses by LGE decreased significantly 4 weeks after MI in both the IBR-MSC group (−3.99 ± 1.62 g; P=0.049)
and the IBR-Placebo group (-2.85±1.27 g; P=0.01); there was no significant difference between the groups (P=0.75; Figure 7A). When the components of LGE were analyzed separately, there was a significant decrease in core zone at 4 weeks (-6.38±1.45 g; P=0.01) and a significant increase in the gray zone extent (+2.39±0.46 g; P=0.01) in the IBR-MSC group. The IBR-Placebo group had a significant decrease in the core zone at 4 weeks (-3.46±1.10 g; P=0.01) but no significant change in the gray zone extent (+0.61±0.74 g; P=0.67). There was no significant difference between the groups for the change in core zone (P=0.25; Figure 7B), but there was for the gray zone (P=0.036; Figure 7C).

Bioreactor Removal and In Vitro Assessment of VEGF Release
After the 4-week MRI, IBRs were removed, washed, and placed in culture. Analysis of CM collected from outside recovered IBRs showed continued PF production by IBRs from the IBR-MSC group with a mean VEGF concentration of 786.9±256.6 pg/mL after 7 days in culture, indicating survival and continued PF production and release by allogeneic pMSCs after 4 weeks in vivo.

Washout Phase and 8-Week Cardiac MRI
Following IBR removal, animals remained in the study for an additional 4 week “washout” phase. The final cardiac MRI at 8 weeks after MI revealed that following the washout phase there were no significant differences for the changes in EDV (P=0.40), ESV (P=0.12), or LVEF (P=0.14; Figure 6) between the IBR-MSC and IBR-Placebo groups. Similarly, LGE MRI showed no significant differences between the groups in terms of the changes in the total scar size (P=0.42) or the separate core zone (P=0.82) and gray zone extent (P=0.91; Figure 7) following the washout phase.

Hemodynamic Assessment and Coronary Angiography
After the 8-week, final MRI, animals underwent invasive hemodynamics and coronary angiography. The mean dP/dT max was 1182.2±185.8 mm Hg/s in the IBR-MSC group and 852.9±63.6 mm Hg/s in the IBR-Placebo group; this difference did not reach statistical significance (P=0.09). There were also no significant differences in measures of diastolic function, including dP/dT min (-1706.5±185.8 mm Hg/s versus 1493.1±194.2 mm Hg/s; P=0.29) and tau (0.053±0.021 seconds versus 0.050±0.006 seconds; P=0.29). Coronary angiography revealed no significant difference in vessel patency of the left anterior descending artery (infarct vessel) between the groups, with Thrombolysis in Myocardial Infarction-3 (normal) flow in 6 of 8 (75%) of the IBR-MSC and 5 of 7 (71%) of the IBR-Placebo animals (P=NS).
Necropsy

On necropsy, 4 of 8 animals in the IBR-Placebo group had adherent pericardium, consistent with pericarditis, as compared with 0 of 8 animals in the IBR-MSC group (P=0.02). Gross inspection of the lungs, liver, kidneys, and spleen revealed no evidence of neoplasm in either group. All animals (16/16) who underwent baseline MRI and IBR implantation survived the full duration of the 8-week study.

Histology

Histologic comparison of the infarct border zone from IBR-MSC and IBR-Placebo animals at 8 weeks using hematoxylin and eosin and Masson’s trichrome revealed the MRI gray zone correlated with areas of heterogenous tissue composed of myocytes intermixed with fibrotic scar at the infarct border zone (see Figure 7E).

Comparison With DI MSCs

Of the 6 animals that underwent DI of 25 million MSCs, all survived the injection procedure performed 3 days after MI, although 1 was euthanized in the week after DI because of failure to thrive. The remaining 5 animals survived the duration of the study. Cardiac MRI at 4 weeks following DI revealed large-magnitude, although statistically nonsignificant, changes compared with baseline for EDV (+27.07±12.17 mL; P=0.08), ESV (+22.89±9.39 mL; P=0.08), and LVEF (−2.97±1.33; P=0.08) on paired analyses. When the 3 groups (IBR-MSC, IBR-Placebo, and DI-MSC) were compared, there were significant differences across the 3 groups for the change in ESV (P=0.05). Post hoc analysis revealed that the change in ESV at 4 weeks was significantly lower in the IBR-MSC group than the DI-MSC group (+2.14±4.13 mL versus +22.89±9.39 mL; P=0.01). For the change in EDV at 4 weeks, the difference across the 3 groups did not reach significance (P=0.06). There was also no significant difference across the 3 groups for change in LVEF (P=0.51).

The MRI LGE studies at 4 weeks demonstrated significant decreases in total scar (−5.44±1.43 g; P=0.04) and core scar (−3.72±1.55 g; P=0.04) extents in the DI-MSC group compared with baseline, but no significant change in gray zone extent (−1.72±1.46 g; P=0.35). Across the 3 groups, there was a significant difference for the change in gray zone extent (P=0.016), with post hoc analysis revealing a greater increase in gray zone in the IBR-MSC group than in the DI-MSC group (+2.39±0.46 g versus −1.72±1.46 g; P=0.003). There was no difference for the change in gray zone extent between the IBR-Placebo and DI-MSC groups (P=0.11), and

Table: MRI Measurements

| Measurement | IBR-MSC Baseline | IBR-MSC 4 Weeks | IBR-MSC 8 Weeks | DI-MSC Baseline | DI-MSC 4 Weeks | DI-MSC 8 Weeks |
|-------------|-----------------|----------------|----------------|----------------|----------------|----------------|
| EDV, mL     | 90.7±5.96       | 89.7±6.04      | 90.3±6.84      | 85.8±5.13      | 85.0±5.35      | 84.4±5.24      |
| ESV, mL     | 59.6±3.94       | 57.7±3.02      | 59.8±3.28      | 57.7±3.02      | 57.7±3.02      | 57.7±3.02      |
| LVEF, %     | 38.7±5.82       | 38.7±5.82      | 38.7±5.82      | 38.7±5.82      | 38.7±5.82      | 38.7±5.82      |
| Core scar mass, g | 1.9±0.21        | 1.9±0.21       | 1.9±0.21       | 1.9±0.21       | 1.9±0.21       | 1.9±0.21       |
| Gray zone mass, g | 1.4±0.21        | 1.4±0.21       | 1.4±0.21       | 1.4±0.21       | 1.4±0.21       | 1.4±0.21       |
| LV mass, g  | 77.8±4.39       | 78.2±4.39      | 78.0±4.39      | 78.2±4.39      | 78.0±4.39      | 78.0±4.39      |

Mean values ± standard error of the mean (SEM). DI indicates direct injection; EDV, end-diastolic volume; ESV, end-systolic volume; IBR, implantable bioreactor; LV, left ventricular; LVEF, left ventricular ejection fraction; MSC, mesenchymal stem cell.
also no significant difference across the 3 groups for the change in total scar mass ($P=0.39$) and core zone mass ($P=0.33$) at 4 weeks.

Necropsy at 8 weeks revealed no evidence of pericarditis in the DI-MSC animals that survived the duration of the study (0/5).

**Discussion**

There is growing evidence that the beneficial effects of stem cell administration following MI are mediated to a large extent by PFs, including growth factors and cytokines, as opposed to cell engraftment.\(^9\)–\(^12\) Here, we present proof-of-concept that a device designed to harness the beneficial effects of stem cell PFs by providing a protected environment in the vascular space, remote from the heart, from which exogenous allogeneic stem cells can release PFs over a prolonged period while being protected from washout, dilution, and immune-mediated rejection. Our findings suggest that allogeneic MSCs deployed in an IBR device can reduce adverse cardiac remodeling 4 weeks after MI compared with both placebo and the same number of directly injected MSCs. By nature of the IBR design, which allows free release of PFs but not cells, the beneficial effects observed in the IBR-MSC group are necessarily not the result of stem cell engraftment or direct cell-cell interaction, and thus provide further evidence to support the “paracrine hypothesis” of stem cell–mediated cardiac repair.\(^9\),\(^10\)

Specifically, we show that in a large animal model, the deployment and maintenance of a catheter-based IBR in the superior vena cava with contained exogenous stem cells is feasible and well tolerated for up to 4 weeks with no evidence of adverse effects from the IBR itself or from the contained cells. This is despite the fact that the cells used were xenogeneic in the preliminary biocompatibility testing, when IBRs containing human MSCs were implanted in pigs, and allogeneic in the MI efficacy study, when IBRs loaded with pig MSCs were deployed in unrelated pigs. Second, we show that a relatively small number of MSCs (2.5 x 10\(^7\)) contained in the IBR in a location remote from the heart exerts protective effects as evidenced by decreased adverse remodeling and preserved LVEF over 4 weeks following an MI. The magnitude of this benefit was significant, including a >5-fold difference in the change in ESV compared with placebo (+2.14 mL versus +11.35 mL; $P=0.036$). This result compares favorably with those reported in clinical trials of angiotensin-converting enzyme inhibitors,\(^24\),\(^25\) which are currently first-line therapy to limit adverse remodeling and left ventricular dysfunction after MI.\(^26\) These results also compare favorably with clinical trials of bone marrow–derived stem cells in acute MI, despite the use of much higher numbers of cells in those studies.\(^27\),\(^28\) In this study, there was a significantly greater increase in ESV 4 weeks after MI in animals receiving intramyocardial injection of 25 million MSCs compared with the same number of cells deployed in the IBR. Relative to prior animal studies, 25 million MSCs is a low dose,\(^29\)–\(^31\) and our results suggest

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**Figure 6.** Implantable bioreactor (IBR) based cell therapy prevents adverse remodeling compared with placebo at 4 weeks after myocardial infarction as assessed by MRI. Over the 4-week treatment phase, end-diastolic volume (EDV) (A) and end-systolic volume (ESV) (B) significantly increased in the IBR-Placebo group, but not in the IBR-mesenchymal stem cell (MSC) group. These changes were accompanied by a significant decrease in left ventricular ejection fraction (LVEF) in the IBR-Placebo group but not in the IBR-MSC group (C). Unpaired analysis demonstrated a significant difference for ESV ($P=0.036$), but not EDV ($P=0.059$) or LVEF ($P=0.40$) between the 2 groups at 4 weeks. After the final 4-week “washout” phase following IBR removal (dashed lines), animals in the IBR-MSC group were no longer protected from adverse remodeling, with no significant difference between groups for the changes in EDV, ESV, or LVEF. Colored $P$ values indicate significant differences for paired analyses where present. Black $P$ values show results of unpaired analyses at 4 weeks; there were no significant differences between the groups from week 4 to 8 ($P$ values not shown). $^*$ indicate $P$ values less than or equal to 0.05. $P$-values that are greater than 0.05 do not have an asterisk.

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**A** End Diastolic Volume  
**B** End Systolic Volume  
**C** Left Ventricular Ejection Fraction

![Graphs showing data](image-url)
that while this dose is sufficient to protect against adverse remodeling when deployed in the IBR, intramyocardial injection requires higher numbers of MSCs to protect against adverse remodeling.

The observed attenuation of adverse remodeling in animals in the IBR-MSC group was accompanied by differential tissue characteristics within the infarct as assessed by LGE. At week 4, there was a significant increase in the heterogeneous gray zone extent in the IBR-MSC group that was not observed in the IBR-Placebo or DI-MSC groups. Histology at 8 weeks after MI revealed the MRI gray zone corresponded with heterogeneous tissue composed of viable myocytes and scar tissue (see Figure 7E). The peri-infarct border zone is thought to be important in remodeling because of the presence of myocytes prone to apoptosis and dedifferentiation.\textsuperscript{32,33} The significant increase in MRI gray zone in the IBR-MSC group suggests improved salvage of viable myocytes in the peri-infarct border zone and/or prevention of their dedifferentiation and apoptosis.

In addition, there was a marked difference in the prevalence of pericarditis observed between the IBR-MSC and IBR-Placebo groups at the time of necropsy: 0\% versus 50\% ($P=0.02$). Pericarditis after MI is typically present following large infarcts and may result from a robust inflammatory response.\textsuperscript{34,35} Inflammation following MI that is prolonged,
excessive in magnitude, and/or insufficiently suppressed can lead to sustained tissue damage, promoting adverse remodeling and resultant heart failure.36,37 The absence of pericarditis in the IBR-MSC group suggests modulation of inflammation, which is a known effect of MSCs attributable to the release of soluble factors.38–41 There was also no evidence of pericarditis in the DI-MSC group, consistent with the anti-inflammatory effects of MSCs. That both the IBR-MSC and DI-MSC groups were protected from pericarditis, but only the IBR-MSC from adverse remodeling suggests that the early, time-limited anti-inflammatory effect of DI-MSCs may be sufficient to prevent pericarditis, but that sustained PF release is necessary to prevent remodeling.

While further study is needed, the protection from adverse remodeling and increase in the peri-infarct heterogeneous gray zone observed in the IBR-MSC group appear to be directly related to the presence of the IBR-MSC device, as over the 4-week “washout” phase after IBR removal there was no significant difference between the IBR-MSC and IBR-Placebo groups in terms of protection from further adverse remodeling or change in infarct size and composition. As a proof-of-concept study, the parameters tested here were limited in terms of the initiation and duration of IBR placement and of the number and type of cells used. In future studies, it will be important to examine these factors. For example, there are clinical data to suggest that delaying cell delivery by several days after MI enhances the benefit, likely due in part to poor cell survival in the inhospitable, proinflammatory environment of recently infarcted myocardium.42 In contrast, angiotensin-converting enzyme inhibitor administration is most effective when initiated early, that is, in the first 24 to 48 hours following an MI.25,26 Intravascular bioreactor-based therapy is predicated on the provision of a protected environment for contained cells in a location remote from the heart, and thus could afford a mechanism to provide the benefit of cell-based PF administration throughout the time course of infarct healing, including shortly after MI when the anti-inflammatory and antiapoptotic effects of stem cell PFs would be expected to be advantageous. Importantly, the IBR can be readily placed in acutely ill patients with the same techniques used to place a central venous line and without the potential risks associated with more invasive cell delivery techniques in the early post-MI phase. In addition, while most cell death and inflammation occur in the initial days after MI, adverse remodeling is known to continue for weeks to months.1,43–45 Analysis of the media collected from outside of the MSC-IBRs explanted after 4 weeks in vivo showed continued VEGF production, indicating survival and continued PF release by the contained MSCs. As such, if an IBR were left in place for longer than 4 weeks, further protection from adverse remodeling may be afforded, perhaps resulting in a more durable effect.

The dose and type of cells deployed in the IBR also requires further study. The dose of $2.5 \times 10^7$ MSCs used in this study was chosen on the basis of a limited assessment of in vitro PF production for a relatively simple IBR design and geometry. The IBR could be designed to accommodate a much larger number of cells and allow for dose efficacy studies to determine the optimal in vivo cell dose. Also, while MSCs were used here, other cell types may have a favorable PF profile for cardiac repair,11,12 and the protected, immune-privileged environment the IBR provides could allow for use of multiple different cell types, whether allogeneic or xenogeneic, alone, or in combinations. Deployment of the IBR via the internal jugular vein, as tested here, would also provide ready access to replace or “recharge” the contained cells, as there is evidence that repeat dosing of stem cells may be advantageous.46

Finally, an alternate strategy would be to deliver PFs alone. However, it remains unclear which of the hundreds or perhaps thousands of PFs produced by stem cells are critical for cardiac repair, whether optimal action requires a combination of particular factors, and how the ratios and types of factors vary over the time course of the healing process. This may explain why the results of clinical trials employing individual PFs administered to the heart or peripheral vasculature did not, to date, achieve all of the desired benefits.47–50 In contrast, delivery of an IBR containing cells that have the ability to produce their full complement of PFs and vary their release based on changing environmental conditions would, in theory, allow the contained cells to tailor the combination of factors released to optimize tissue repair and healing.

Study Limitations

The conclusions from this study are limited by the relatively small number of animals studied (n=8 in each IBR group and n=6 in the DI group). In addition, the animals were young and healthy, and thus may not be indicative of clinical conditions in which patients suffering an MI are typically older with concomitant conditions such as diabetes mellitus and hypertension that may adversely affect recovery and remodeling. The animals in this study were also not treated with the evidence-based medical therapies (eg, angiotensin-converting enzyme inhibitors and beta-blockers) that postinfarction patients typically receive. In addition, although we provide data indicating that MSCs release PFs from the IBR in vitro and continue to do so after being explanted from the body, we were unable to obtain in vivo measures of PF release because of an inability to differentiate IBR-released PFs from endogenous PFs attributable in part to a limited number of porcine-specific PF antibodies. We also cannot identify which PFs resulted in the beneficial effect noted in the IBR-MSC group. While we show that porcine MSCs contained in explanted
IBRs continued to release VEGF in vitro, this factor may not be representative of the many PFs released by MSCs. Likewise, while a significant reduction in pericarditis was seen in the IBR-MSC group compared with placebo, we did not collect serum samples to correlate this result to inflammatory markers such as interleukin-1, interleukin-6, or tumor necrosis factor-alpha. Finally, the hemodynamic and autopsy data were obtained from animals at 8 weeks after MI, which was 4 weeks after removal of the IBR. As a result, we do not have hemodynamic or pathologic data at 4 weeks after MI, when the favorable MRI remodeling effects were most pronounced, and thus are limited in our ability to correlate the hemodynamic and pathologic findings reported here with the observed beneficial effects on adverse remodeling.

Conclusions

In summary, we demonstrate here proof of concept that allogeneic MSCs, housed in the protected environment of an IBR in the superior vena cava, remote from the heart, can prevent adverse remodeling and alter scar composition after MI compared with placebo and to direct myocardial injection of the same number of MSCs. The IBR allows free release of soluble PFs but not of cells, and thus the beneficial effects observed in the IBR-MSC group are necessarily mediated by paracrine mechanisms as opposed to stem cell engraftment, lending further support to the paracrine hypothesis of myocardial repair that underlies this therapeutic paradigm.

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Disclosures

This study involves technology that was developed at the Johns Hopkins University and is under license by Domicell, Inc. Drs Johnston, Hwang, Schulman, Tomaselli, Weiss, and Gerstenblith are inventors of the technology. Additionally, they are founders of Domicell and hold equity in the company. Johns Hopkins University also holds equity in Domicell. This arrangement has been reviewed and approved by Johns Hopkins University in accordance with its conflict of interest policies. Ms Bogdan is an employee of Domicell, Inc. Dr Hare reports having a patent for cardiac cell–based therapy and holds equity in Vestion Inc. He maintains a professional relationship with Vestion Inc. as a consultant and member of the Board of Directors and Scientific Advisory Board. Vestion Inc. did not play a role in the design and conduct of the study. Dr Hare holds a relationship with Longeveron, LLC that includes consulting and member of the Board of Directors and Scientific Advisory Board. The remaining authors have no disclosures to report.

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