Tissue-Engineered Myocardial Patch Derived From Extracellular Matrix Provides Regional Mechanical Function

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Background—Extracellular matrix (ECM), a tissue-engineered scaffold, recently demonstrated cardiomyocyte population after myocardial implantation. Surgical restoration of myocardium frequently uses Dacron as a myocardial patch. We hypothesized that an ECM-derived myocardial patch would provide a mechanical benefit not seen with Dacron.

Methods and Results—Using a canine model, a full thickness defect in the right ventricle was repaired with either Dacron or ECM. A third group had no surgery and determined baseline RV function.Eight weeks later, global systolic function was assessed by the preload recruitable stroke work relationship. Regional systolic function was measured by systolic area contraction (SAC), calculated by high density mechanical mapping. Tau was used to assess global diastolic function. Recoil rate and diastolic shear were used as measures of regional diastolic function. After functional data acquisition, tissue was fixed for histological evaluation. Global systolic and diastolic functions were similar at baseline and after ECM and Dacron implantation. Regional systolic function was greater in the ECM group compared with the Dacron group (SAC: 4.1±0.9% versus −1.8±1.1%, P<0.05). Regional diastolic function was also greater in the ECM group (recoil rate (° sec⁻¹): −44±7 versus −17±2, ECM versus Dacron; P<0.05). Immunohistochemical analysis revealed cardiomyocytes in the ECM implant region, a finding not seen with Dacron.

Conclusion—At 8 weeks, an ECM-derived tissue-engineered myocardial patch provides regional mechanical function, likely related to cardiomyocyte population. These results are in sharp contrast to Dacron, a commonly used myocardial patch. (Circulation. 2005;112[suppl I]:I-144–I-149.)

Key Words: heart failure | surgery | mechanics | remodeling | myocytes

Heart failure is a notoriously progressive disease, despite medical management.1 The increasing gap between the incidence of end-stage heart failure and surgical treatment is due, in great part, to the shortage of donor organs.2 In 2002, only 2154 heart transplants were performed, despite an annual average of 50000 deaths related to New York Heart Association stage IV congestive heart failure.3 In addition to transplantation, other surgical modalities may play an increasingly important role in treatment of end-stage heart failure. These include ventricular assist devices, surgical restoration, passive constraint devices, coronary artery bypass grafting, and mitral valve repair.4

Another surgical approach to heart failure underscores the importance of the geometry of the ventricle. In most cases of heart failure, the ventricle assumes a dilated, spherical shape, decreasing the efficiency of ventricular contraction. In this instance, revascularization alone would not be sufficient to improve systolic function. This alternate approach is based on the premise of restoring the normal ellipsoidal shape of the left ventricle from its decompensated, globular form.5-9 This principle was illustrated in the Surgical Anterior Ventricular Endocardial Restoration trial, which combined surgical restoration of an akinetic or dyskinetic anterior wall with Dacron and concomitant coronary artery bypass grafting or mitral valve repair.10 This strategy demonstrated an immediate improvement in ejection fraction, with sustained benefit and minimal morbidity.10

Tissue engineering offers possibilities beyond Dacron.11 Specifically, it seeks to create organic scaffolds that both restore the geometry of the ventricle and attract or house cellular elements better suited for myocardial function.12-14 Thus, the development of a tissue-engineered myocardial patch (TEMP) represents the next frontier in the evolving surgical treatment of heart failure. Recently, Badyak et al15 investigated the use of porcine urinary bladder extracellular matrix (ECM), a biologically latent membrane...
devoid of cells, as a TEMP. Scattered islands of cardiomyocytes were identified on the implant site after 8 weeks.

In this study, we hypothesized that ECM would confer mechanical benefit as a result of cardiomyocyte population 8 weeks after myocardial implantation. We tested this hypothesis by implanting ECM in the canine heart, and compared its mechanical function to Dacron, a biologically inactive patch frequently used in surgical restoration of the heart.

**Methods**

**Experimental Animals**

All animals received humane care in compliance with the Principles of Laboratory Animal Care formulated by the National Society for Medical Research and the Guide for the Care and Use of Laboratory Animals prepared by the National Academy of Sciences and published by the National Institutes of Health (NIH Publication No. 85-23, revised 1985). Dogs were divided into 3 groups: Baseline, ECM, and Dacron. The baseline group was subject only to the surgery necessary to establish normal right ventricular (RV) function (n=4). The ECM and Dacron groups each consisted of 5 dogs. Four dogs in each group were subject to both mechanical function evaluation and histological examination. One dog in each group was used solely for histological analysis to confirm that mechanical measurement techniques did not affect histological findings.

**Initial Surgery**

Adult mongrel dogs weighing 20 to 30 kg were sedated using ketamine, intubated, and placed under general anesthesia using isofluorane. The right chest was entered using a right anteromedial incision over the fifth intercostal space. A pericardial cradle was constructed. Next, a portion of the posterolateral right ventricular free wall was isolated using a Satinsky clamp. A full thickness portion of myocardium approximately 15 by 10 mm was excised. To test the adequacy of the defect, a suture was placed above and below its long axis, and the Satinsky clamp was slowly released. A commensurate portion of ECM (ACell) or Dacron (USCI) was used to repair the defect with a running 5.0 prolene suture (Figure 1A). The clamp was released. Once adequate hemostasis was obtained, the chest was closed in the standard fashion.

**Terminal Surgery**

Eight weeks later, the animals were returned to the operating room, sedated, intubated, and placed under general anesthesia. A catheter was placed in the femoral artery for hemodynamic monitoring. The anterior chest wall was removed via bilateral thoracotomy. Adhesions were carefully dissected off the heart to expose the patch and its suture line. A pressure transducer (Millar Instruments) was placed into the RV. Finally, the inferior vena cava (IVC) was isolated, and a length of umbilical tape was placed around it above the level of the diaphragm.

**Global Systolic Function**

Global function of the RV was studied by determining the preload recruitable stroke work relationship as previously described. Briefly, ultrasonic crystals were secured along 3 axes of the heart: Base to apex, anterior to posterior, and septal to RV free wall. Data were recorded during gradual occlusion of the IVC. At the end of the experiment, the ellipsoidal shell subtraction method was used to calculate RV volume. Volume data were used to create pressure-volume loops for each heartbeat subject to IVC occlusion. Stroke work, defined as the integral of pressure with respect to volume change, was calculated for each heartbeat and plotted against end-diastolic volume, establishing a linear relationship. The slope of this line represents right ventricular contractility, a measure of systolic function independent of heart rate and afterload. This relationship was established using at least 7 consecutive points that decreased stroke work by at least 50%.

**Global Diastolic Function**

Tau, the time constant for isovolumic relaxation, was calculated based on a method described by Weiss et al and used as a measure of global diastolic function. Right ventricular tau was calculated at 8 weeks for baseline, ECM, and Dacron groups. Data were obtained at the onset of each experiment by recording RV pressure waveforms at a rate of 250 samples per second. At the end of the experiment, RV pressure waveforms during isovolumic relaxation were fit to an exponential decay, allowing for determination of tau.

**Regional Systolic Function**

Regional function was determined by high density mapping (HDM), a technique developed in our laboratory and described in detail elsewhere. Briefly, a region of interest (ROI) was selected on the epicardial surface of the heart and covered with speckles. The speckles, composed of silicon carbide particles, created a random high contrast light intensity distribution used to measure epicardial surface deformation. Next, a complimentary metal-oxide semiconductor camera (Photron) was focused on the ROI and images were taken at 250 frames per second. Using a subpixel extended-phase correlation algorithm, high-resolution displacement vector maps of epicardial surface deformation during the cardiac cycle were obtained. The vector arrays were used to calculate area change via subject to IVC occlusion. Stroke work, defined as the integral of pressure with respect to volume change, was calculated for each heartbeat and plotted against end-diastolic volume, establishing a linear relationship. The slope of this line represents right ventricular contractility, a measure of systolic function independent of heart rate and afterload. This relationship was established using at least 7 consecutive points that decreased stroke work by at least 50%.

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Green’s theorem between consecutive images and over an entire heartbeat. Systolic area contraction (SAC) was calculated as the difference between the end-diastolic area and the end-systolic area, normalized to the end-diastolic area. To compare hearts with different ROI size, the end-diastolic area was set to 1.

Regional Diastolic Function
The rate and total amount of epicardial shear during diastole were used as measures of diastolic function. Vector maps obtained from HDM were used to calculate epicardial shear through infinitesimal Eulerian strain equations. These values were then converted to Lagrangian strain, and the change in shear angle during diastole was computed. Recoil rate, recently shown to be a correlate of tau, was computed by differentiating maximum change in diastolic shear angle with respect to time.

Histology
Specimens of the patch area in all 3 groups were placed in 4% paraformaldehyde and later transferred to 30% sucrose solution. The specimens were then placed in an embedding matrix and sectioned at a thickness of 10 μm. Measurements of thickness were taken from the middle of each implant. These sections were stained with hematoxylin and eosin and alpha sarcomeric actinin.

Statistical Analysis
All data are expressed as the mean±SEM. Statistical analyses between groups were performed by 1-way analysis of variance with a post-hoc Duncan test. Differences were considered significant for probability values less than 0.05.

Results
Mortality and Macroscopic Results
Overall, 17 dogs were included in this study. There were 3 operative deaths, 1 relating to the fragility of the ECM and 2 attributable to arrhythmia. However, mortality dropped to less than 4% in subsequent series with adjustments in protocol, ie, adding a weight-based dosage of lidocaine before placing and releasing the Satinsky clamp.

Mediastinal adhesions were present for both ECM and Dacron groups. On gross appearance, however, there were substantially more adhesions in the Dacron group. Furthermore, a dense layer of connective tissue was evident over the epicardial surface of all Dacron patches. After the removal of overlying connective tissue, the Dacron patch was clearly identifiable (Figure 1D). Aside from a wrinkled appearance, the patch was unchanged (Figure 1E). Conversely, the ECM implants appeared well integrated with adjacent myocardium (Figure 1B and 1C). A comparable layer of connective tissue was not evident. There was no evidence of aneurysmal dilation in either ECM or Dacron groups.

The average size of the defects for both groups was 16±3 by 10±2 mm. The thickness of ECM at implantation was approximately 150 μm. Eight weeks later, the average thickness was 2.2±1.1 mm. The average thickness of the Dacron patch at implantation was 250 μm. Eight weeks later, the average thickness was 2.4±1.0 mm. The average thickness of native myocardium was 5.4±0.8 mm.

Microscopic Results
Histological examination of sections was conducted in both groups (Figure 2). The Dacron group displayed fibroblast proliferation with abundant collagen deposition organized in a dense manner (Figure 2E). The Dacron patch was intact; there was no infiltration by cellular elements. Cardiomyocytes were not evident on the Dacron patch (Figure 2F).

ECM displayed a variety of cell types. Clusters of cardiomyocytes were seen along the endocardial surface (Figure 2B). These cells were organized in a diminishing gradient from the outer border to the center of the implant site, with few cardiomyocytes in the center of the patch. These cells stained positive for alpha sarcomeric actinin (Figure 2C) and were striated and approximately the same size and shape as adjacent native myocytes (Figure 3). The cardiomyocytes on
The average size of the ROI for all groups was 8 mm by 8 mm. Average SAC in baseline was 21.6 ± 1.8%, whereas the Dacron implants demonstrated essentially no SAC (SAC = -1.8 ± 1.1%). Conversely, the SAC in the 8-week ECM patch region was 4.1 ± 0.9% (Figure 4). This was significantly lower than baseline (P < 0.05), yet significantly higher than Dacron (P < 0.05). A summary of regional systolic function is seen in the Table.

**Regional Diastolic Function**

Regional diastolic function, assessed by spatiotemporal shear deformation, differed between the Dacron and ECM groups. Dacron caused a reversed shear pattern and a lower recoil rate during isovolumic relaxation (Figure 5). The Dacron group had an average diastolic shear of 0.0 ± 0.3°, compared with 5.6 ± 1.3° at baseline (P < 0.01). The average diastolic shear of the ECM group was 3.0 ± 0.1°, lower than baseline (P < 0.05) and higher than Dacron (P < 0.05). The regional recoil rate was -52 ± 8° sec⁻¹, -44 ± 7° sec⁻¹, and -17 ± 2° sec⁻¹ in the baseline, ECM, and Dacron groups, respectively (P < 0.05 for both Dacron versus baseline and Dacron versus ECM).

**Discussion**

The novel finding of our report is that a TEMP derived from ECM contributes to regional function 8 weeks after implantation in the canine heart. In addition, we confirmed cardiomyocyte population of ECM. The etiology of these cells has been under investigation, with possible explanations including the deposition of circulating bone marrow-derived progenitor cells and the fusion of cardiac progenitor cells with host cells.²⁴⁻²⁵ Beltrami et al²⁶ suggested that Lin-C-kit positive stem cells act as key players in myocardial reconstitution. The origin of these cardiomyocytes remains an unanswered question, worthy of future research, but is not the focus of this study.

The regional mechanical benefit in our report is based on a technique that correlates epicardial surface deformation with pressure change. The technique used allows for determination of regional deformation within the boundaries of the patch implantation. Examination of the area changes seen in ECM during the cardiac cycle identifies 2 important findings. First, there is active contraction of the ECM. The pressure-area loop clearly demonstrates regional systolic contraction in the ECM implant. This pattern of deformation can only result

![Figure 4](image-url)
from active contraction in the implant region and not passive elastic recoil. Second, this contraction is in synchrony with native myocardium. Although definitive electrophysiological studies were not performed, simultaneously recorded data from ECM and adjacent native myocardium in selected animals clearly demonstrate that the contraction in the implant region is in phase with that of the normal, un-operated on myocardium (Figure 4C). Microscopic evaluation of Dacron patches did not demonstrate the presence of cardiomyocytes nor do the mechanical data indicate that Dacron implantation contributes to regional function. These findings suggest that cardiomyocyte population of ECM is likely responsible for its regional systolic function.

Right ventricular contractility, a measure of global systolic function, was decreased 15% in the ECM group and 25% in the Dacron group when compared with baseline. Although the differences were not statistically significant, power analysis suggests an additional 8 dogs per group would be needed to demonstrate a statistically significant difference. The lack of outright statistical significance may also be related to the small size of the defect created in the RV in relation to its total surface area, a clear limitation of our study. While investigating the relationship of infarct size to cardiac function, Pfeffer et al\(^\text{27}\) concluded that a significant change in global function would not be evident until at least 20% of the ventricle was compromised. In the present study, approximately 5% of the RV was removed and patched. Additionally, it is unlikely that the single layer of ECM used in our study would withstand left ventricular pressure, thus limiting its applicability in its current state. Increasing the number of ECM layers could be an alternative, but it is unclear if the physiological benefit, ie, cardiomyocyte population, would still be evident.

Grossly, Dacron elicited far greater fibrosis than ECM, correlating with more mediastinal adhesions and epicardial connective tissue deposition. On placement, the Dacron patch was clearly under tension. Surprisingly, 8 weeks after implantation, all Dacron patches had a wrinkled appearance, suggesting the patch was no longer under tension (Figure 1D and 1E). The “wrinkle effect” is likely the result of wound contraction around Dacron. In sharp contrast, ECM triggered far less fibrosis. The patch was neither wrinkled nor aneurysmal and appeared to share the same surface tension as adjacent native myocardium. Finally, after removal of adhesions, it was difficult to grossly distinguish ECM from native myocardium (Figure 1C). The quantitative and qualitative differences between ECM and Dacron could be explained by an inherent ability of ECM to house cellular elements that facilitate remodeling.

Our results examining diastolic function were similar to those we obtained for systolic function. Tau, the time constant for isovolumic relaxation, was similar in all 3 groups. The differences in regional diastolic function between ECM and Dacron are likely the result of the cells found on the implant region and the compliance mismatch between native myocardium and the patch. The modulus of elasticity of Dacron is at least 4 orders of magnitude greater than healthy myocardium, ie, Dacron is stiffer than myocardium. Thus, the use of Dacron as a myocardial patch may have a “tethering effect” that would reduce the mechanical function of surrounding myocardium. Furthermore, the cellular response to Dacron was primarily diffuse fibroblast proliferation, an observation also seen with remodeling after myocardial infarction. In contrast, ECM stimulated less fibrosis and was populated by different cell types, including cardiomyocytes. Atkins et al\(^\text{28}\) have shown that the reduction of infarct stiffness via cell transplantation leads to increased diastolic function. Similarly, Quarterman et al\(^\text{29}\) created a detailed finite element model to show that cell transplantation alone will result in changes in compliance that result in mechanical benefit.

The potential clinical applications of ECM as a scaffold are many and would have a powerful impact on the management of cardiac disease. These would include instances in which Dacron is presently used as a myocardial patch: repair of ventricular aneurysms, repair of congenital heart defects, and most recently, surgical restoration of a dyskinetic or akinetic ventricle. By its contribution to regional systolic function, ECM provides true restoration of the ventricle rather than nonfunctional substitution of defective tissue, as is the case.
with Dacron. Furthermore, it already has Food and Drug Administration approval for several indications, potentially facilitating its use in humans as a TEMP.

Acknowledgments
This study was supported by grants from the National Institute of Health HL20558 (Dr Cohen), HL28958 (Dr Cohen), HL67101 (Dr Cohen), and EB000261 (Dr Badylak), and the New York State Office of Science, Technology, and Academic Research (Dr Gaudette). We greatly appreciated the effort of Peter V. Kochupura, MD, for his vigorous review of this manuscript.

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Circulation. 2005;112:I-144-I-149
doi: 10.1161/CIRCULATIONAHA.104.524355

Circulation is published by the American Heart Association, 7272 Greenville Avenue, Dallas, TX 75231
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Print ISSN: 0009-7322. Online ISSN: 1524-4539

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