

Mechanical preconditioning enables electrophysiologic coupling of skeletal myoblast cells to myocardium

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Objective: The effect of mechanical preconditioning on skeletal myoblasts in engineered tissue constructs was investigated to resolve issues associated with conduction block between skeletal myoblast cells and cardiomyocytes.

Methods: Murine skeletal myoblasts were used to generate engineered tissue constructs with or without application of mechanical strain. After in vitro myotube formation, engineered tissue constructs were co-cultured for 6 days with viable embryonic heart slices. With the use of sharp electrodes, electrical coupling between engineered tissue constructs and embryonic heart slices was assessed in the presence or absence of pharmacologic agents.

Results: The isolation and expansion procedure for skeletal myoblasts resulted in high yields of homogeneously desmin-positive ($97.1\% \pm 0.1\%$) cells. Mechanical strain was exerted on myotubes within engineered tissue constructs during gelation of the matrix, generating preconditioned engineered tissue constructs. Electrical coupling between preconditioned engineered tissue constructs and embryonic heart slices was observed; however, no coupling was apparent when engineered tissue constructs were not subjected to mechanical strain. Coupling of cells from engineered tissue constructs to cells in embryonic heart slices showed slower conduction velocities than myocardial cells with the embryonic heart slices (preconditioned engineered tissue constructs vs embryonic heart slices: 0.04 ± 0.02 ms vs 0.10 ± 0.05 ms, $P = .011$), lower maximum stimulation frequencies (preconditioned engineered tissue constructs vs embryonic heart slices: 4.82 ± 1.42 Hz vs 10.58 ± 1.56 Hz; $P = .0009$), and higher sensitivities to the gap junction inhibitor (preconditioned engineered tissue constructs vs embryonic heart slices: 0.22 ± 0.07 mmol/L vs 0.93 ± 0.15 mmol/L; $P = .0004$).

Conclusions: We have generated skeletal myoblast-based transplantable grafts that electrically couple to myocardium. (*J Thorac Cardiovasc Surg* 2012;144:1176-84)

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Skeletal myoblast cells (SMBs) have been investigated for cardiac cell therapy of congestive heart failure since the

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1990s. Despite limited information about safety and efficacy, Hagege and colleagues¹ applied SMBs in a clinical trial. As shown in the Myoblast Autologous Grafting in Ischemic Cardiomyopathy trial, the inability of SMBs to integrate functionally and electrically to the host myocardium resulted in adverse ventricular arrhythmias.² At the same time, it was demonstrated that unmodified transplanted SMBs neither integrate into the host myocardium nor transdifferentiate into cardiomyocytes.³

On the other hand, it has been shown that SMBs, which have been preconditioned before transplantation, were able to integrate into the host myocardium.⁴⁻¹¹ The preconditioning regimens included virally mediated expression of connexin43 (Cx43), in vitro co-culturing of SMBs with cardiomyocytes, and chemical induction of Cx43 expression. Because autologous cardiomyocytes are difficult to obtain and there are safety issues associated with transgenic approaches, these methods may not be clinically relevant.

Other cell types that have been applied for cardiac cell therapy approaches include mesenchymal stem cells, endothelial progenitor cells, and cardiac cells derived from pluripotent stem cells. Although all of these cells have been

Abbreviations and Acronyms

AP	= action potential
Cx43	= connexin43
DMEM	= Dulbecco's Modified Eagle Medium
EGFP	= enhanced green fluorescent protein
EHS	= embryonic heart slice
ETC	= engineered tissue construct
FBS	= fetal bovine serum
K-ETC	= killed engineered tissue construct
NP-ETC	= nonpreconditioned engineered tissue construct
P-ETC	= preconditioned engineered tissue construct
SMB	= skeletal myoblast cell

shown to exert some myocardial effects,¹² the underlying mechanisms remain unknown. Furthermore, we have shown that the quality of autologous progenitor cells (ie, mesenchymal stem cells) is dependent on patient-specific factors.¹³ Because of ethical and safety reasons, so-called embryonic stem cells and induced pluripotent stem cells, respectively, are not considered for clinical application at this time.

Our group¹⁴ previously demonstrated a method to precondition rat SMBs in a manner that could be clinically applicable. By seeding the cells into a 3-dimensional hydrogel construct and subjecting them to tensile strain, the expression levels of proteins essential for functional and electrical coupling were significantly increased. After transplantation, the skeletal cells were able to establish an accessory conduction pathway as a potential treatment for atrioventricular conduction block.

We sought to address known electrophysiologic issues of SMB transplantation for cardiac therapy² by investigating the electrical coupling properties of SMBs, embedded in mechanically preconditioned engineered tissue constructs (ETCs), to myocardium in an ex vivo transplantation model.

MATERIALS AND METHODS**Isolation and Cell Culture of Skeletal Myoblast Cells**

Animals received humane care in compliance with the *Principles of Laboratory Animal Care*, formulated by the National Society of 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 (Publication No. 86-23, revised 1996). The Institutional Animal Care and Use Committee approved all experiments. All experiments were approved by the local institutional and governmental authorities.

SMBs were isolated from the limb muscle tissue of neonatal C57BL/6 mice as described previously.¹⁵ The tissue sample processing involved incubation in 0.2% type IV collagenase (Invitrogen, Karlsruhe, Germany), 2.4 IU/mL dispase (Invitrogen) and 3 mmol/L CaCl₂ (Sigma-Aldrich, Munich, Germany) in phosphate-buffered saline (Invitrogen). Isolation medium containing Ham's F10 medium (Invitrogen), 20% fetal bovine

serum (FBS, Invitrogen), and 2.5 ng/mL basic fibroblast growth factor (PeproTech, Hamburg, Germany) was applied for culturing the cells at a density of 1×10^5 cells/cm² on collagen-coated plates (5 μg/cm² type I collagen, Invitrogen). Preplating procedures (ie, further culture of suspension cells, while adherent cells were discarded) were applied after 1, 2, 18, and 48 hours of incubation. After a further incubation for 48 hours, the adherent cells were expanded in growth medium: 40% Ham's F10 medium, 40% Dulbecco's Modified Eagle Medium (DMEM, Invitrogen), and 20% FBS, 2.5 ng/mL basic fibroblast growth factor. Ten days after isolation at 80% confluence, differentiation of SMBs to myotubes was triggered by a change in culture conditions to differentiation medium: DMEM, 2% horse serum (PAA, Coelbe, Germany), which was changed daily for 14 days. All media were supplemented with penicillin/streptomycin (1:100) and fungizone (1:1000, both Invitrogen).

Fabrication of Engineered Tissue Constructs

ETC fabrication was based on a slightly modified method as previously published.¹⁵ The matrix scaffold for the ETC was prepared by mixing ice-cold isolation medium, type I collagen (5 mg/mL, Invitrogen), and 0.1 mol/L NaOH and Geltrex (Invitrogen) at a ratio of 3:3:1:1. SMBs were labeled with Vybrant DiI (Invitrogen) according to the manufacturer's instructions before using for ETCs. A defined number (5×10^6) of SMBs were resuspended in 200 μL isolation medium and mixed with 500 μL of the premixed matrix.

To fabricate preconditioned ETC (P-ETCs), the mixture was cast into molds composed of silicone tubing cut in half lengthwise with monofilament polyester mesh (0.331 opening; McMaster-Carr, Elmhurst, Ill) attached to each end with silicone adhesive (Rhodia, Cranbury, NJ). Constructs were warmed at 37°C to induce gelling and covered with myoblast culture media.

For the fabrication of nonpreconditioned ETCs (NP-ETCs), the suspension was poured directly on a cell culture dish (BD Biosciences, Heidelberg, Germany). The tissue constructs were first incubated at 37°C in isolation medium for 3 days followed by incubation in differentiation medium for 11 days. The complete timeline of the ETC fabrication protocol is as follows: skeletal muscle tissue preparation (day 0), preplating (days 0-3), cell culture expansion (days 3-10), ETC generation (day 10), ETC cell culture (days 10-13), ETC differentiation (days 13-24), and co-culture with embryonic heart slices (EHSs) (days 24-30).

Preparation of Embryonic Ventricular Heart Slices

Vital EHSs from day 18 C57BL/6 mice, expressing enhanced green fluorescent protein (EGFP) under control of the cardiac specific α -actin promoter, were prepared as described previously.^{16,17} Briefly, the explanted hearts were prepared and embedded in 4% low-melt agarose (Roth, Karlsruhe, Germany). These agarose blocks were subsequently sectioned into 300-μm-thick transversal sections using a VT1000s microtome (Leica, Wetzlar, Germany) and stored in preoxygenized ice-cold Tyrode's solution (composition in mmol/L: NaCl 136, KCl 5.4, NaH₂PO₄ 0.33, MgCl₂ 1, D-glucose 10, 2,3-butanedione monoxime 30, and N-2-hydroxyethylpiperazine-N'-2-ethanesulfonic acid 5; pH 7.4). After sectioning, CaCl₂ (Sigma-Aldrich) was added to a final concentration of 0.9 mmol/L, and the tissue slices were incubated for 30 minutes at 4°C. The EHSs were washed twice in phosphate-buffered saline and transferred to a 10-cm dish (BD Biosciences) containing ice-cold Iscove's modified Dulbecco's medium (Invitrogen) supplemented with 20% FBS (Invitrogen). EHSs were incubated for 1 hour in a standard cell-culture incubator and then used for co-culture with ETCs.

Engineered Tissue Construct Preparation and Co-Culture With Embryonic Heart Slices

On day 11 of differentiation cell culture, the center regions (2-mm long) of P-ETCs were prepared using a sterile scalpel. The center parts of

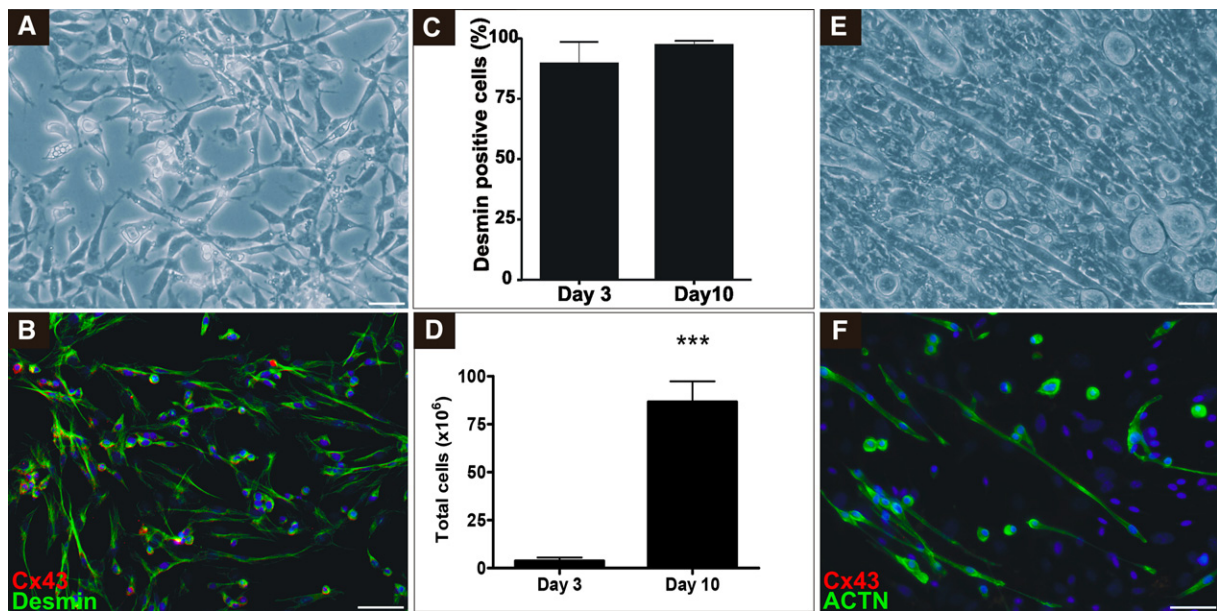


FIGURE 1. Isolation and characterization of SMBs. Bright field micrograph (A) and (B) immunocytochemical staining of SMBs after 7 days of cell culture: desmin (green), connexin43 (Cx43, red), and 4',6-diamidino-2-phenylindole-dihydrochloride (blue). C, Flow cytometric quantification of desmin expression in SMBs after preplating (day 3) and cell culture expansion (day 10). D, Quantification of total cell yield per 100 mg muscle tissue and fraction of desmin-positive cells after preplating (day 3) and cell culture expansion (day 10, *** $P < .0001$; $n = 5$). Bright field micrograph (E) and (F) immunocytochemical staining of SMBs differentiated to myotubes: α -actinin (ACTN, green), Cx43 (red), and 4',6-diamidino-2-phenylindole-dihydrochloride (blue) after 14 days of differentiation. Scale bars = 50 μ m.

NP-ETCs were cut into similarly sized and shaped pieces. Control ETCs containing dead cells (killed [K]-ETCs) were prepared by immersing P-ETCs in distilled water for 24 hours.

EHSs and ETCs were transferred to a custom-made well chamber with a funnel-shaped cavity (funnel dish) to facilitate physical contact between EHSs and ETCs. After 6 days of incubation in differentiation medium, cocultures were used for immunohistochemistry and electrophysiologic assessment.

Histology

Tissue samples were snap-frozen in TissueTek (Sakura Finetek, Staufen, Germany) and cut into 8- μ m sections with a CM1950 cryotome (Leica).

For histochemistry, the sections were fixed with acetone and stained with Masson's trichrome (both Sigma-Aldrich) following the manufacturer's instructions.

For immunohistochemistry, the sections were fixed (4% paraformaldehyde) and permeabilized (0.25% Triton X-100). Unconjugated primary antibodies used for immunocytochemistry included anti-desmin (GeneTex, Irvine, Calif), anti-connexin45 (Millipore, Schwalbach, Germany), anti-Cx43, anti-N-cadherin (both Abcam, Cambridge, UK), anti-sarcomeric- α -actinin (Sigma-Aldrich), anti-dystrophin, (Santa Cruz, Heidelberg, Germany), and anti-EGFP (Invitrogen). Primary antibodies were detected with species-specific AlexaFluor conjugated secondary antibodies. Nuclei were stained with 4',6-diamidino-2-phenylindole-dihydrochloride (both Invitrogen). Fluorescence microscopy was performed with an Eclipse Ti-U microscope using the NIS Elements BR 3.10 software package (Nikon, Düsseldorf, Germany).

Electrophysiologic Analysis

The funnel dish with the co-culture sample in 3 mL serum-free DMEM, oxygenated with O₂/CO₂ (95%/5%), was placed on a custom-made

heating plate (37°C). Electrophysiologic measurements were performed with sharp electrodes (20-40 M Ω when filled with 3 mol/L KCl) made of filament borosilicate glass capillaries (WPI, Sarasota, Fla). Signals were amplified with a SEC-10LX amplifier (NPI Electronic, Tamm, Germany) and acquired with Pulse software (HEKA, Lambrecht, Germany). A defined stimulation frequency (2-15 Hz) was applied using a square pulse stimulator (SD9; Grass Technologies, West Warwick, RI). Data were analyzed with the Mini Analysis program (Synaptosoft, Fort Lee, NJ). Pharmacologic inactivation of gap junctions was performed by titrating defined volumes (2-100 μ L) of 1-heptanol (Sigma-Aldrich) to the medium.

Statistical Analysis

Numeric data are expressed as mean \pm 1 standard deviation. Statistical analyses were performed using the Statistical Package for the Social Sciences (release 20; IBM, Somers, NY). Data were tested by 1-way analysis of variance followed by a Bonferroni post hoc test for multiple comparisons.

RESULTS

Isolation and Culture of Skeletal Myoblast Cells

The SMB isolation process yielded $3.80 \pm 1.77 \times 10^6$ cells per 100 mg ($n = 5$) of skeletal muscle tissue from neonatal mice. After 3 preplating steps and a subsequent incubation for 48 hours, the total cell number was reduced to approximately 50%. The majority of the remaining adherent cells showed the typical spindle-shaped morphology of SMBs (Figure 1, A), and immunocytochemical analysis revealed a majority of these cells to be positive for desmin (Figure 1, B). This was confirmed quantitatively by flow

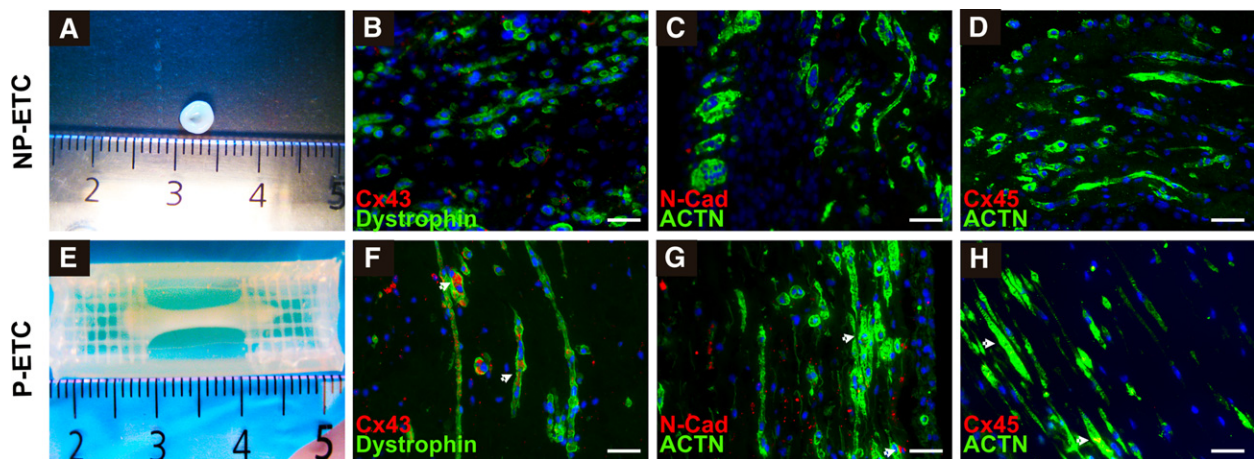


FIGURE 2. Characterization of ETCs. SMBs and myotubes from nonpreconditioned engineered tissue construct (*NP-ETC*; A-D) and preconditioned engineered tissue construct (*P-ETC*; E-H) conditions. Macroscopic images of NP-ETC (A) and P-ETC (E). Immunohistochemical staining confirms increased gap junction (*arrows*, connexin43 [*Cx43*] and connexin45 [*Cx45*]) and adhesion protein (N-cadherin [*N-Cad*]) expression for P-ETC (F-H) compared with NP-ETC (B-D). The regression pattern of myotube-specific (dystrophin) and muscle-specific structural markers (α -actinin [*ACTN*]) was similar for both conditions. Scale bars = 50 μ m.

cytometric analysis showing $89.5\% \pm 4.0\%$ ($n = 5$) desmin-positive cells. A further 10 days of cell culture expansion yielded $86.71 \pm 10.66 \times 10^6$ cells per 100 mg of tissue with even further increased purity of desmin-positive cells ($97.1\% \pm 0.1\%$, $n = 5$; Figure 1, C and D). When changing culture conditions to differentiation medium, SMBs changed morphology and fused to multinucleated, spontaneously contracting myotubes, showing expression of sarcomeric α -actinin but not retaining expression of gap junction protein Cx43 (Figure 1, E and F).

Construct Morphology and Co-Culture

SMBs from cell culture expansion were used for the fabrication of ETCs. The application of 5×10^6 cells per ETC resulted in a homogeneous and longitudinal alignment of myotubes in P-ETC, whereas NP-ETCs showed no defined orientation of cells (Figure 2, A, B, E, and F). Immunohistochemical analysis revealed ubiquitous expression of dystrophin and sarcomeric α -actinin for cells within the ETCs. Expression of gap junction proteins Cx43 and connexin45, as well as adherens junction protein N-cadherin, was detectable only in P-ETCs and not in NP-ETCs (Figure 2, B, D, F, and H).

Co-Culture and Electrophysiologic Analyses

For co-cultures, the center sections of ETCs were prepared to obtain the regions of the ETCs with the highest and most reproducible longitudinal alignment of embedded SMBs¹⁵ and carefully positioned in funnel dishes to guarantee direct contact with the EHSs. For 8 of 10 co-culture experiments, P-ETCs and EHSs remained physically attached for 6 days during co-culture (NP-ETCs: 6/10; K-ETCs: 4/10; Figure E1, A-G).

Electrophysiologic measurements were performed to investigate electrical coupling between cells in the ETCs and EHSs. Sharp glass electrodes were used to target individual cells in EGFP-positive EHSs for recording of intracellular action potentials (APs) after stimulation in ETCs or EHSs (Figure 3, A and B).

When the stimulation electrode was placed in the ETCs (S1-R), only the P-ETC group demonstrated APs in unison with EHS cells (Figure E1, H), as opposed to the NP-ETC and K-ETC groups, in which only APs from spontaneous contractions of the EHSs were observed. In all 3 conditions, stimulation within the EHSs (S2-R) resulted in APs spreading between cells in the EHS, which reflects the viability and excitability of the EHS (Figure 3, C-E).

APs from stimulation at 4 Hz from P-ETCs showed longer delays to stimulation trigger than APs from stimulation from EHSs (Figure 4, A) and resulted in a significantly lower conduction velocity when normalized to distance between location of stimulation and recording (0.04 ± 0.02 ms vs 0.10 ± 0.05 ms, $P = .011$; $n = 5$; Figure 4, B).

To further assess the quality of electrical coupling between ETCs and EHSs, increased stimulation frequencies were applied. There were significantly higher maximal stimulation frequencies for stable electrical coupling when stimulation within the EHSs was compared with stimulation from the ETCs (Figure 4, C-F; 4.82 ± 1.42 Hz vs 10.58 ± 1.56 Hz; $P = .0009$; $n = 5$). Higher stimulation frequencies led to irregular “skips” in the resulting APs (Figure 4, D).

To verify that the stimulation was propagated through electrical coupling by gap-junctions between cells, and not a conduction artifact caused by a field stimulation effect, the gap-junction inhibitor 1-heptanol was added

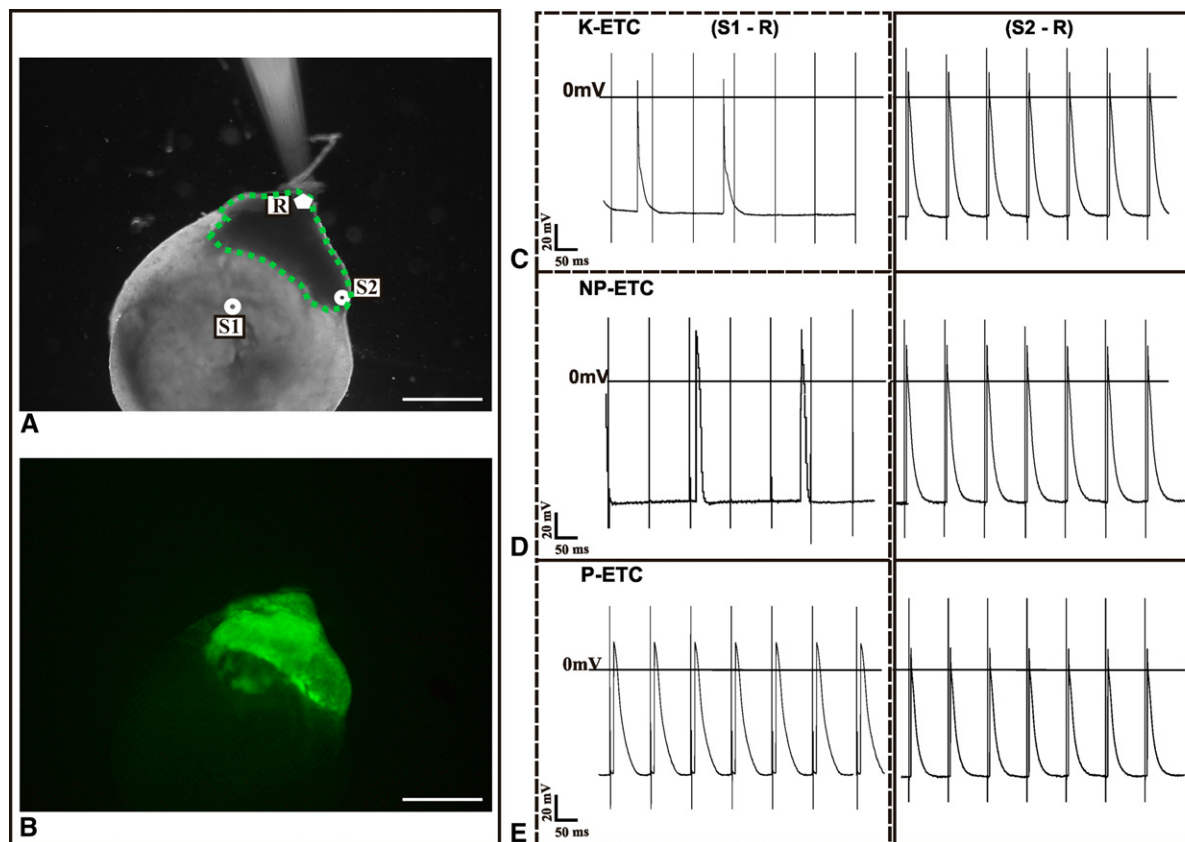


FIGURE 3. Electrophysiological measurements of EHSs co-cultured with ETCs. A, Positions of electrodes for electrophysiological measurements: stimulation in the ETC (S1), stimulation in the EHS (S2), and recording in the EHS (R). B, eGFP-positive embryonic ventricle slice (EHS) in co-culture with ETC. C-E, Representative APs measured at R from stimulation at S1 (ETC) and S2 (EHS) for killed engineered tissue construct (*K-ETC*; C), nonpreconditioned tissue construct (*NP-ETC*; D), and preconditioned engineered tissue construct (*P-ETC*; E). Traces on the *left* show AP at R from stimulation at S1 (ETC to EHS), and traces on the *right* show AP at R from stimulation at S2 (EHS to EHS). Scale bars = 1 mm.

incrementally to the media until the stimulation failed to evoke APs.

Although 1-heptanol eventually blocked AP generation in the EHSs independently of whether the stimulation originated in the ETCs or within EHSs, the concentration of inhibitor was significantly lower for ETC-to-EHS conduction block compared with EHC-to-EHS conduction block (Figure 4, *G-J*; 0.22 ± 0.07 mmol/L vs 0.93 ± 0.15 mmol/L; $P = .0004$; $n = 3$).

The observed blocking effects were completely reversible under decreasing 1-heptanol concentrations (Figure E1, *I*).

Histologic Analyses of Co-Cultures

Histochemistry (Masson's trichrome; Figure 5, *C*) of the border zone revealed close physical contact of ETCs and EHCs, but a markedly lower cell density in the ETCs compared with the EHSs. Immunohistochemistry demonstrated desmin expression across the border zone, with less expression of this protein in the ETCs compared with the EHSs (Figure 5, *D*). Cx43 expression was high in the EHSs and in the immediate border zone to the

ETCs with only minimal expression within the ETCs (Figure 5, *E*).

DISCUSSION

In the present study, we investigated the physical and electrophysiological integration of SMB-based ETCs in an *in vitro* transplantation model. Our major findings can be summarized as follows:

First, we have shown highly efficient isolation of expandable SMB populations from neonatal murine muscle tissue. Cells from these preparations nearly ubiquitously expressed the myogenic marker desmin, emphasizing the absence of contaminating fibroblasts and homogeneity of enriched cells with a functional muscle cell phenotype. Subsequent change of cell culture conditions led to differentiation of single cell SMBs to syncytial, spontaneously contracting myotubes, underlining the efficient generation of cellular structures with the ability to generate contractile force.

Second, ETCs produced with SMBs and matrix components were subjected to unidirectional strain during differentiation of the cells into contractile myotubes.

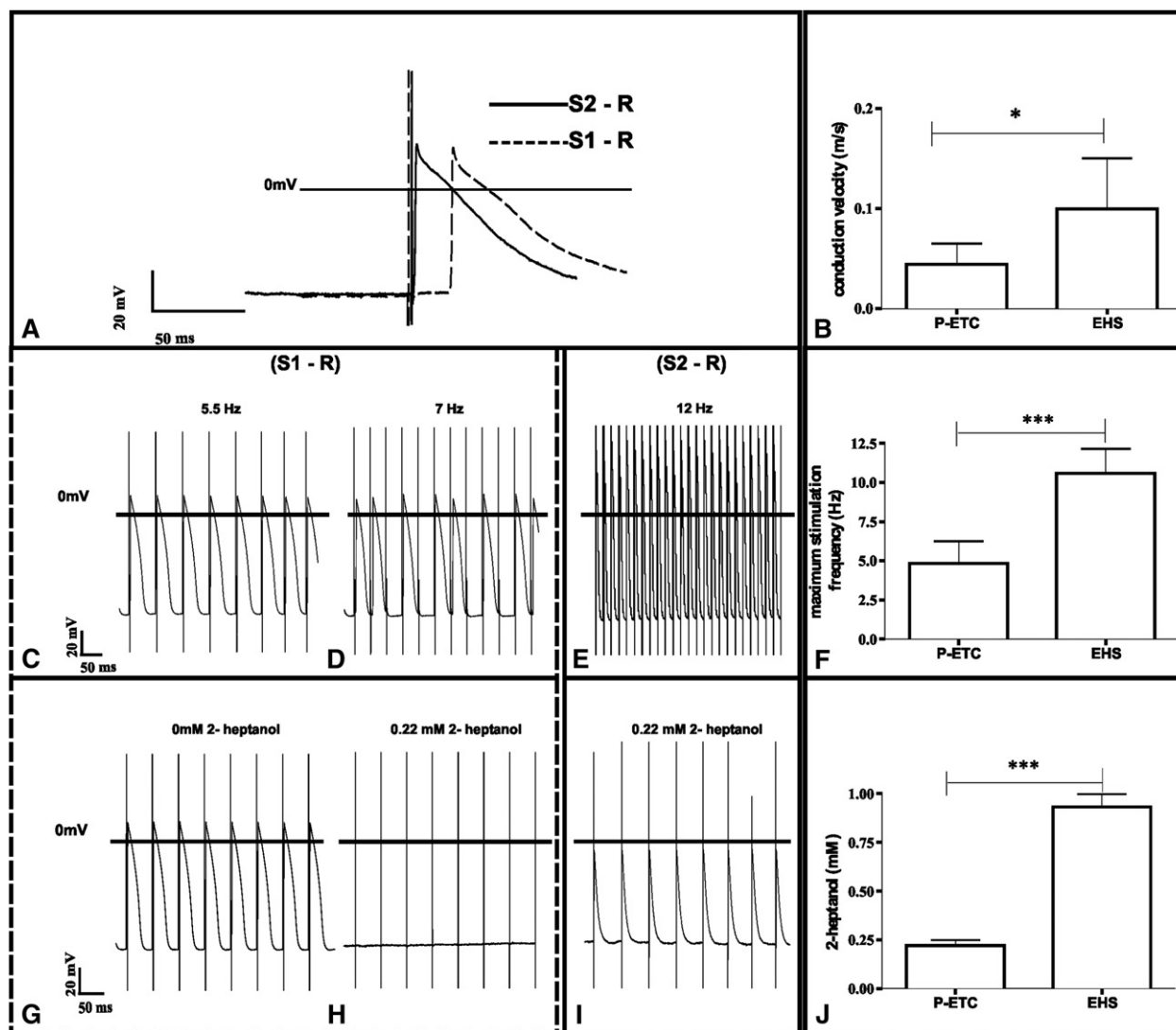


FIGURE 4. Characterization of electrophysiologic coupling between preconditioned engineered tissue constructs (*P-ETC*) and embryonic heart slice (*EHS*). A and B, Conduction velocity: overlay of exemplary *EHS* APs (A) evoked by unipolar stimulation in the *ETC* (S1) and *EHS* (S2). Significant differences in conduction velocity measurements (B) between *P-ETC* to *EHS* and *EHS* to *EHS*. C-F, Maximum stimulation frequency: *P-ETC* stimulation and *EHS* measurement show APs at 5.5 Hz stimulation (C), but resulting in conduction blocks at 7 Hz (D), whereas *EHS* stimulation follows *EHS* recording until 12 Hz without conduction blocks (E), resulting in different maximum stimulation frequency between *P-ETC* to *EHS* and *EHS* to *EHS* measurements (F). G-J, 1-Heptanol sensitivity: AP at *EHS* from 5.5 Hz stimulation at *P-ETC* before 1-heptanol treatment (G) and after addition of 0.2 mmol/L 1-heptanol (H). Stimulation at *EHS* still results in AP at *EHS* at 0.2 mmol/L 1-heptanol (I) and 1-heptanol blocking concentration (J) between *P-ETC* to *EHS* and *EHS* to *EHS* measurements. * $P < .05$. *** $P < .001$, $n = 5$.

Immunohistochemical analyses revealed continued expression of gap junction and adherens junction proteins, which are necessary for the functional integration of (transplanted) cells in the heart. In *ETCs* that were not mechanically stimulated (NP-*ETCs*), these proteins were expressed at low levels and the cells in the constructs remained unorganized.

Third, co-culture of *P-ETCs* and vital *EHSs*, which served as an *in vitro* transplantation model, showed physical engraftment of the engineered tissue with the native tissue and electrophysiologic integration of skeletal muscle-derived cells in *ETCs* to cardiomyocytes in vital *EHSs*. By

using sharp electrode measurements, we quantitatively determined the efficiency of the coupling between *ETCs* and *EHSs* and compared values for conduction velocity, maximum stimulation frequency, and sensitivity to specific pharmacologic gap junction inhibitors to native electrophysiologic coupling within the *EHS*, serving as an internal positive control. Our results clearly showed electrophysiologic coupling of cells within the *ETC* to cells within the *EHS*, but for all 3 parameters examined, this occurred to a lesser extent than the coupling of cells within the native heart tissue.

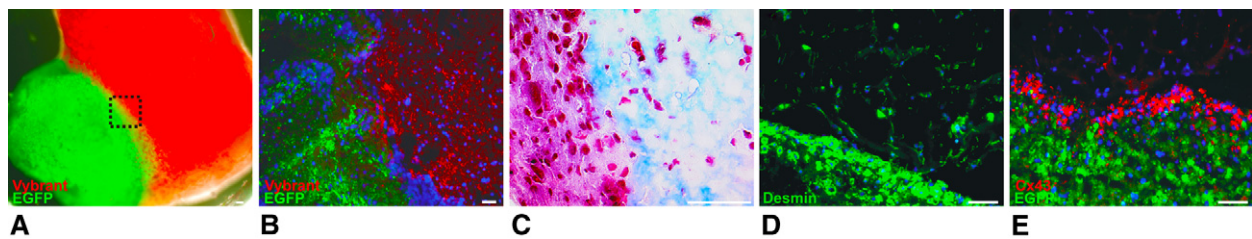


FIGURE 5. Histologic analysis of co-cultured engineered heart tissue slices and engineered heart tissue at day 6. A, Macroscopic fluorescence picture of EHS (green: enhanced green fluorescent protein [EGFP]) and P-ETC (red: Vybrant DiI [Invitrogen, Karlsruhe, Germany]). B, Fluorescence micrograph of the P-ETC/EHS border zone. C, Masson's trichrome staining of the P-ETC/EHS border zone (blue: collagen, red: cytoplasm). Immunohistochemical staining for desmin (green) (D) and EGFP (green) and Cx43 (red) (E) in P-ETC/EHS border zone. Cx43 was highly expressed in myotubes in the P-ETC/EHS border zone. Scale bars = 50 μ m.

Isolation of skeletal muscle–derived cell populations has been achieved by various methods,^{18,19} usually including preplating steps during the initial phase of culturing crude primary cell suspensions. This technique serves to separate cell populations on the basis of their attachment to plastic surfaces and is often used for the removal of unwanted fibroblasts, but also for the enrichment of specific multipotent progenitor cell populations.^{20,21} In our case, SMBs were acquired from neonatal tissue for reasons of cell culture compatibility and expansion potential,²² and culture conditions were optimized to generate high numbers of functionally competent SMBs for the subsequent fabrication of ETCs.

By applying directional mechanical strain on the tissue constructs, the embedded SMBs were oriented in a longitudinal fashion and condensation of the hydrogel led to a compact and uniform distribution of cells. Furthermore, the expression of gap junction proteins Cx43, Cx40, and N-cadherin was maintained during maturation of SMBs to myotubes. This effect of mechanical signal transduction has been described for the expression of genes involved in various mechanistic pathways,²³ including the expression of connexins,^{24,25} and is considered to be linked to integrins and cadherins,^{26,27} whereas the exact mechanism is still not fully understood. We showed that electrophysiologic integration to cardiac tissue is only possible when SMB-based ETCs are subjected to tensile strain, demonstrating the necessity for mechanical preconditioning. This characteristic of skeletal muscle–generated progenitor cells has been described²⁸ but never as a prerequisite for electrical integration.

Nevertheless, SMBs have been one of the first cell populations that have been applied in clinical trials for cardiac cell therapy,²⁹ but they showed poor reproducible therapeutic efficacy in the most recent clinical trials.^{2,30} The induction of ventricular arrhythmias after SMB transplantation has been a recurring and intensely debated issue. A direct association of immature phenotypes of cell transplants and decreased conduction velocity,³¹ as well

as improper alignment of gap junction and adherens proteins, was demonstrated in vitro and in animal models.³²

It is now generally accepted that resolving the challenges regarding electrophysiologic integration remains a major challenge in advancing the clinical utility of cell-based cardiac therapies.³³

The idea to modify gene expression from SMBs by preconditioning is well established in these approaches.⁴ For example, the transplantation of genetically modified SMBs overexpressing Cx43 has been shown to prevent postinfarction arrhythmia.¹¹ This study demonstrated the potential of tissue engineering and preconditioning of SMBs to build gap junctions and to establish electrical coupling with cardiac cells; however, genetic manipulation limits the potential use of these cells in a clinical setting.

Vital EHSs^{34,35} co-cultured with ETCs allowed us to directly assess the quality of electrical integration using quantitative electrophysiologic parameters. We have shown that SMBs within the ETCs couple electrophysiologically to the EHSs after 6 days of co-culture. However, this coupling was less robust when compared with electrophysiologic connections between cells in the native heart tissue.

Corresponding to our previous findings,¹⁴ which showed that ETC grafts could augment the cardiac conduction system in a rat model of atrioventricular conduction block, the current study confirms the expression of Cx43 within the transplanted ETC.

As a structural basis for electrical integration, expression of Cx43 at the EHS–ETC interface was shown for the mechanically P-ETCs as opposed to the control NP-ETCs, confirming the notion of Cx43 being essential for cardiac electrophysiologic coupling.

Study Limitations

Calcium transient imaging experiments were not performed because they were beyond the primary scope of the present study. However, we recognized that the detailed characterization of the calcium transients of the different tissues may help identify the origin of potential ectopic

arrhythmias.^{3,6} Future experiments will focus on using voltage- or calcium-sensitive dyes to further assess the connectivity of the tissues using optical mapping as described previously.¹⁴

Furthermore, the investigation of the contractile properties of SMBs embedded in the ETCs and their ability to couple mechanically to the host myocardium are of significant importance. In vitro determination of developed force as fatigue testing would provide further insight. In addition to the in vivo electrical integration of the tissue constructs that we confirmed previously,¹⁴ the functional in vivo integration of the tissue constructs needs to be investigated and will be the subject of future projects.

CONCLUSIONS

We demonstrated electrophysiologic coupling of skeletal myoblast-derived cells with cardiomyocytes in the context of engineered tissues. This is a decisive step forward toward the future application of autologous and functionally competent cells as therapeutic agents for cardiac cell therapy.

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Discussion

Dr Y. Joseph Woo (Philadelphia, Pa). Dr Choi, I congratulate you and your colleagues on an outstanding body of work. Your work is very innovative. The mechanical preconditioning, your in vitro transplant model of heart slices, sophisticated

electrophysiologic studies, proper biologic controls, and loss of function and restoration of function controls all should be commended. I have 2 quick questions.

Have you looked at any complementary studies of gap junction function and electrophysiologic coupling, such as direct imaging of calcium transients?

Dr Choi. We did not do that. We performed a dye transfer experiment by immersing one end into calcium AM, the dye was transferred, and the dye transfer could be stopped by additional administration of 1-heptanol.

Dr Woo. Would you comment on any ongoing in vivo studies or speculate on your work transitioning into an in vivo model? Would

multiple factors, such as the varied vectors of natural cardiac contractility, potentially decondition your preconditioned myotubules and might your engineered construct serve as a focus for a reentrant tachycardia?

Dr Choi. We also performed transplantation in vivo in mice, and by small animal magnetic resonance imaging in these mice, we observed that the size of the scar decreased and the contractility of the left ventricle increased.

In regard to the other question, we are starting a clinical trial. We are trying to translate everything to meet the Good Manufacturing Practice guidelines, and we are just waiting for approval.

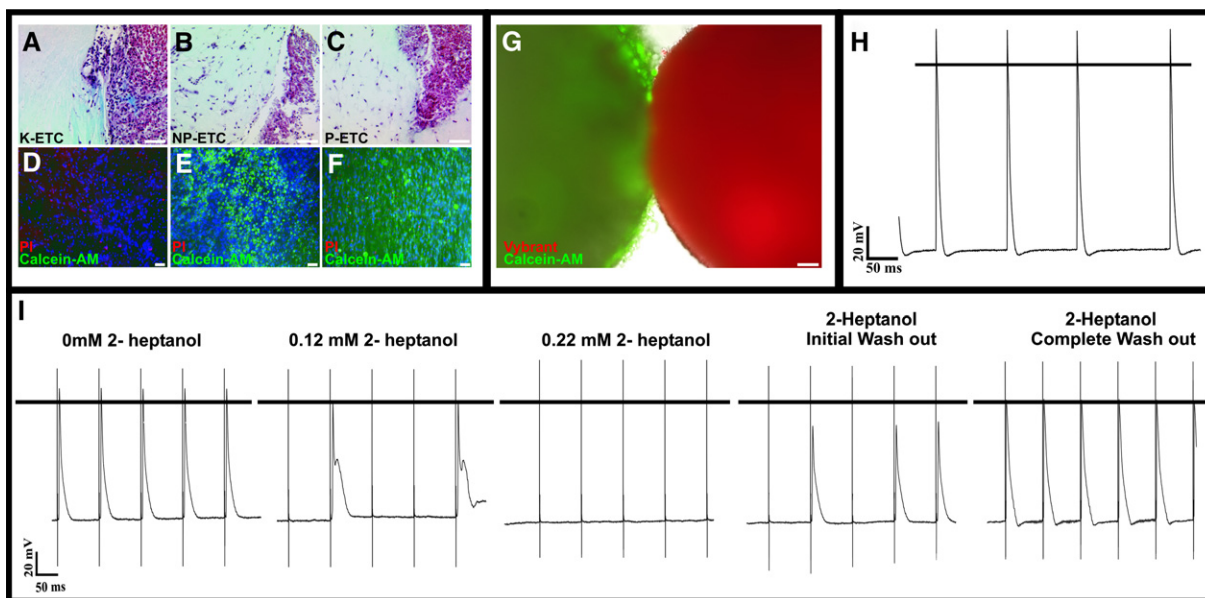


FIGURE E1. Masson’s trichrome staining of the co-culture samples at the junction of (A) killed engineered tissue construct (*K-ETC*)/EHS and (B) nonpreconditioned engineered tissue construct (*NP-ETC*)/EHS shows poor grafting, whereas (C) preconditioned engineered tissue construct (*P-ETC*)/EHS shows close contact of adjacent tissues. D-F, Staining of ETCs shows a majority of dead (propidium iodide positive) cells for *K-ETC* (D), whereas *NP-ETC* (E) and *P-ETC* (F) show mainly vital (calcein AM positive) cells. G, Macroscopic fluorescence picture of Vybrant Dil (red) labeled EHS co-culture with calcein AM (green) labeled EHS to show poor attachment of adjacent tissues. H, Spontaneous AP of EHS. I, APs of *P-ETC*/EHS co-culture during 2-heptanol treatment and after washout. Scale bars = 50 μ m.

ET/BS