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Journal Title: JACC: Basic to Translational Science
Volume: Volume 2, Number 5
Publisher: Elsevier: Creative Commons Licenses | 2017-10-01, Pages 601-609
Type of Work: Article | Final Publisher PDF
Publisher DOI: 10.1016/j.jacbts.2017.06.003
Permanent URL: https://pid.emory.edu/ark:/25593/t6gn9

Final published version: http://dx.doi.org/10.1016/j.jacbts.2017.06.003

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Accessed February 5, 2019 9:44 PM EST
A Minimally Invasive, Translational Method to Deliver Hydrogels to the Heart Through the Pericardial Space

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HIGHLIGHTS

- The pericardial space is an unexploited anatomic location for hydrogel delivery.
- Hydrogels can be delivered to the pericardial space in a localized, minimally invasive manner, without detectable hemodynamic effects.
- Pericardial hydrogel delivery is a new strategy to direct therapeutics to the heart with reduced systemic delivery and off-target effects.
Despite pharmacological and technologic advances, cardiovascular diseases (CVD) remain the leading cause of morbidity and mortality in the United States, costing $215.6 billion per year (1). More patients are surviving, but with heart failure, arrhythmias, and poor quality of life. Micro-ribonucleic acid (miRNA), gene therapy, stem cells, cytokines, and other biologics are new treatments that have shown promise in preclinical and early phase clinical trials (2–4). Many of these therapies require focused delivery of the therapeutic to the heart, or even localization to particular anatomic areas, such as the peri-infarction region. Dilution of these therapeutics by systemic administration increases cost and risks off-target effects. For example, poor retention of stem cells in the heart is thought to limit efficacy in clinical trials (5–7). The proangiogenic cytokine vascular endothelial growth factor encourages neoangiogenesis and cardiac regeneration (8) but can also accelerate tumor metastasis (9). Efficient, targeted, and temporally appropriate delivery of therapeutics to the heart are keys to their successful translation into clinical use.

Early phase clinical trials are underway using hydrogels as therapeutic agents for cardiac repair (10–12). Both solid patches and injectable gel materials are under investigation and may have benefits for different applications. Cardiac patches and solid materials have been tested as structural support for the heart in a clinical trial (13) and therapeutic delivery platforms in numerous preclinical studies (14). Their widespread use is limited by the need for surgical placement. Injectable materials with liquid or gel phases, such as decellularized matrix, alginate, and engineered hydrogels, can provide scaffolds, tactile signals, and structural support for cardiac regeneration and repair (10,11,15,16). Biomaterial gels are particularly suited to deliver stem cells to the heart and retain viable cells at the site of delivery (12,17). Other materials are in preclinical trial for delivery and sustained release of miRNAs, cytokines, and other therapeutics (4,18,19). Whereas biocompatible materials may be beneficial for the treatment of CVD, there are no dedicated delivery methods that are safe and minimally invasive.

There are challenges inherent to delivering biomaterials to the heart. Open heart surgery, although feasible, is less desirable from a cost and patient perspective. Catheter delivery using commercially available single-lumen coronary catheters or Noga XP Cardiac Navigation System (Biosense Webster, Diegem, Belgium) cannot keep material components separate as they travel to the heart and thus cannot control the timing of material interaction and gelation. Premature gelation causes clogging within catheter lumen. Delayed gelation can lead to embolization, stroke, and failure to deliver material to targeted area. Another challenge with biomaterial delivery to the heart is the potential for inducing arrhythmias if the electrical conductivity of the material creates a substrate for a re-entrant circuits as it interdigitates between cardiomyocytes. Therefore the development of material-specific strategies is necessary for the safe, precise, and practical delivery of biomaterial to the heart.

The pericardium is a novel site for therapeutic delivery that has been shown in animal studies to act as a reservoir for drug delivery to the heart (20–22). The advent of epicardial ablations and external left atrial appendage ligation has demonstrated the feasibility of accessing a “dry” pericardial space for therapeutic purposes (23–25). Herein we describe a minimally invasive device to deliver biomaterials to the heart by using the pericardial space as a novel anatomic site for biomaterial delivery. Our device uses the existing anatomic structures to form a temporary compartment for gel delivery. Features of the device eliminate the risk of premature gelation and embolization and allow
precise placement of biomaterial over the area of interest. Pericardial hydrogel delivery with our device circumvents many of the obstacles to vascular or intracardiac delivery, facilitating rapid translation of biomaterial gels into clinical use.

**METHODS**

**DEVICE DESIGN.** We built a device to deliver biomaterial hydrogels to the heart through the pericardial space in a large animal model (pig). The hydrogel delivery device is constructed from varying durometers of polyether block amide (PEBAX) (Temecula Custom Extrusions, Temecula, California) biocompatible polymeric resin, using custom multilumen tooling, extrusion, and fusing processes. Two internal lumens for biomaterials keep components separated throughout the length of the device. The core is comprised of a super-elastic shape-memory nickel titanium alloy (nitinol) that facilitates fence deployment and retraction. Suction and gel ports were cut in the desired location using precision skiving. The sheath is comprised of a laminated composite shaft with an imbedded coil. Key locations of the device and sheath (distal tip, fence apparatus) are fit with radiopaque markers to enable visualization with fluoroscopy. The device is constructed in a cleanroom and sterilized using ethylene oxide before survival procedures.

**HYDROGEL GEL DESIGN AND DELIVERY.** Polyethylene glycol (PEG) hydrogels were based on 4-arm PEG macromere (20 kDa, Laysan Bio, Arab, Alabama) with maleimides at each terminus cross-linked with dithiothreitol. This platform provides structurally defined hydrogels with stoichiometric incorporation of ligands and improved cross-linking efficiency (18). Hydrogel components (macromer, cross-linker) were delivered into the pericardial space in cadaver and live pigs (n = 9) using the delivery device. The hydrogel components and cross-linker were delivered through separate lumens into the fenced area in the pericardial space for in situ mixing and cross-linking. Hydrogel components were adjusted to yield a 5-ml, 4.0% wt/vol PEG hydrogel. For nonsurvival studies, gel was labeled with radiopaque contrast agent iohexol (Omnipaque, GE Healthcare, Princeton, New Jersey) and in others trypan blue (Sigma, St. Louis, Missouri).

**LARGE ANIMAL MODEL.** Farm pigs (n = 9; 45 to 55 kg) were obtained from a commercial supplier and raised on swine feed. On the procedure day, animals were sedated with intramuscular telazol (4.4 mg/kg) and xylazine (0.5 mg/kg) and maintained on inhaled isoflurane (2% to 4%). Animals were intubated, ventilated, and continuously monitored. We accessed the pericardial space through a fluoroscopically guided percutaneous subxiphoid approach using 12-cm 21-gauge micropuncture needle. Intrapericardial location was confirmed by contrast injection and wire confinement to the outer cardiac silhouette. Arterial and venous pressures were monitored using a Swan-Ganz and pigtail catheter, respectively, connected to a MacLab Hemodynamic Recording System (GE Healthcare).

A flexible 10-F catheter sheath was placed into the pericardial space. The hydrogel delivery device was positioned over the desired anatomic area of the heart and temporarily secured in place by negative suction. PEG macromere and dithiothreitol (cross-linker) were delivered through two distinct lumens and combined within the fenced region. After allowing 5 min for gelation, the fence was retracted from around the hydrogel. Invasive hemodynamics were measured before pericardial instrumentation, immediately after hydrogel delivery device removal, and at time of sacrifice 4 to 6 weeks later. Animals received intrapericardial methylprednisolone, 1 mg/kg, and oral colchicine, 0.6 mg, by mouth twice daily.

**STATISTICAL ANALYSIS.** Statistical analysis of hemodynamic data from different time points was compared using repeated measures one-way analysis of variance using PRISM version 6 (GraphPad, San Diego, California). A priori power analysis was performed using G*Power version 3.0.10 (Dusseldorf, Germany) with the assumption of α 0.05, power 0.6, and an effect size calculated to 1.0.

**RESULTS**

**HYDROGEL DELIVERY DEVICE DESIGN AND CONSTRUCTION.** The feasibility and safety of pericardial access in the absence of pericardial effusion (i.e., dry tap) have been shown by other devices, such as the Lariat device (Sentre Heart, Redwood City, California) (23,24). In this study, we engineered a system consisting of a hydrogel biomaterial delivery device and sheath to access the pericardial space using standard micropuncture technique (Figure 1). Device features include a shape memory core, a deployable and retractable fence that creates the lateral wall of the temporary hydrogel compartment, two separated lumens for gel components, a suction mechanism to temporarily secure the device in place, electrical sensors, and an atraumatic tip (Figure 1).

As the delivery device is advanced out of the sheath, it forms a circular fence that physically isolates a 5-cm diameter area of epicardium (Supplemental Video 1). The device forms a lateral border, or fence, while the epicardium acts as a floor, and pericardium acts as a roof for the temporary gel compartment. Gentle suction further seals the
compartment and secures the device in place on the moving heart (Supplemental Video 1, Figure 1). Radiopaque markers allow precise placement over desired anatomic area using biplane fluoroscopy. Inside the device hydrogel components are kept within separate internal lumens until delivery through ports arrayed around the circular fenced area (Figure 1C). After gelation, the shape memory core allows retraction of the delivery device without disruption of hydrogel architecture (Supplemental Video 1).

**VISUALIZATION OF HYDROGEL DELIVERY.** Four-arm PEG maleimide macromer was cross-linked with dithiothreitol to form hydrogels (18). This hydrogel system is a flexible platform for therapeutic delivery of stem cells, growth factors, and pancreatic islets in various preclinical models (16,17,26). Bench studies showed that gelation occurred within 1 min after combining both components at physiological pH. We tested the delivery device in live pigs \((n = 9)\). One animal was excluded from analysis because of infection unrelated to gel delivery. The delivery procedure was minimally invasive and conducted using standard cardiac catheterization laboratory equipment (Supplemental Figure 1). There were no acute complications with pericardial access, device placement in the pericardial space, or hydrogel deployment. All animals had successful hydrogel placement and solidification. No sustained arrhythmias occurred and there were only occasional premature ventricular contractions during the pericardial access procedure. Retraction of the device respected the hydrogel boundary and did not disrupt the hydrogel by visual inspection (Supplemental Video 1). Pericardial access and delivery procedure took approximately 35 min.

We visualized gelation within the temporary compartment created by the delivery device directly and using fluoroscopy with radiopaque gel (Figures 2A and 2B). Sixty minutes after gel delivery, heart was excised and a well-circumscribed trypan blue gel was present under the pericardium and localized to the inferoposterior wall (Figure 2C). There were no instances of premature gelation within the device in 12 live and cadaver studies. After 4 weeks hearts were excised and examined for gel. On 2 hearts areas of possible gel were observed but could not be confirmed with PEG-directed antibody.

**HEMODYNAMIC ASSESSMENT OF EPICARDIAL HYDROGEL DELIVERY.** Complications of pericardial access procedures can include tamponade or constriction causing hypotension, tachycardia, and elevated diastolic filling pressures (27). We measured acute and chronic changes in cardiac hemodynamics (4 to 6 weeks) after gel placement. There were no detectable changes in heart rate, blood pressure, right atrial pressure, wedge pressure, or left ventricular end diastolic pressure (Figure 3, Supplemental Table 1). Other hemodynamic features of pericardial constriction, such as ventricular discordance, were not detected (Figure 3C). There was no periprocedural mortality. One animal was excluded from analysis because of infection unrelated to pericardial procedure but showed no hemodynamic abnormality.
INFLAMMATORY EFFECTS OF PERICARDIAL PROCEDURE AND GEL DELIVERY. Pericarditis is a well-known consequence of pericardial procedures and we evaluated animals for signs of systemic or local inflammation (23–25). We measured white blood cell count, differential, creatinine, and liver function tests, which remained normal in all animals (Supplemental Table 2). Total white blood cell count decreased in 6 of 8 animals (mean decrease by 2.1 \( \times 10^9 \) \( \pm \) 2.1 cells/l) (Supplemental Table 2). Neutrophil and lymphocyte counts were also stable (Supplemental Table 2).

FIGURE 2 Successful Hydrogel Delivery to the Pericardial Space Using the Hydrogel Delivery Device Visualized Directly and by Fluoroscopy

(A) The delivery device was successfully placed over the left anterior descending artery in a porcine cadaver. Sternum was removed to allow direct visualization. (B) Contrast-labeled hydrogel delivery by the device (arrow) was visualized in situ by fluoroscopy. (C) In a live animal, trypan blue-labeled gel was delivered with the device to the posterior wall and gelled within the temporary compartment created by the device, epicardium, and pericardium. Sixty minutes after device removal, the gel remained localized over delivery location.

FIGURE 3 Invasive Hemodynamics Were Unchanged After Pericardial Gel Placement

(A and B) Right atrial pressure and left ventricle end-diastolic pressure did not change in pigs in the immediate post-procedure period or after 4 to 6 weeks (n = 8; error bars \( \pm \) SD). (C) Simultaneous left and right ventricle pressure showed no accentuated respiratory variation and no ventricular interdependence (paper speed 25 mm/s, representative image).
On gross examination, pericardium was thin and translucent 4 to 6 weeks after hydrogel delivery (Figure 4A). Cardiac samples from each chamber were preserved in formalin and stained with hematoxylin and eosin (Figures 4B to 4D). Histologic evaluation showed mostly normal pericardial thickness and cellularity (Figures 4B and 4C). In some animals, in a localized area close to the site of pericardial access near the left ventricular apex, pericardial thickening was observed (Figure 4C). Mononuclear cellular infiltration of the parietal pericardium in this area suggests an inflammatory response or fibroblast proliferation at the site of pericardial puncture. No abnormalities were seen in the myocardial architecture.

**DISCUSSION**

Biocompatible materials are an emerging category of therapeutics to treat CVD. They come in a variety of physical forms (solid, gels) and can be designed to act alone or as carriers for other therapeutics, such as stem cells, miRNAs, and cytokines. The hydrogel used in this study has been shown to improve cardiac function by increasing retention of transplanted mesenchymal stem cells in the heart in small animal studies (17). Although this material does not directly incorporate into the myocardium, other materials nearing or in phase 1 clinical trials work by other mechanisms of action (10–12,15,19). To date, these materials are delivered by techniques or procedures designed for other purposes, such as intracoronary catheters, Noga XP Cardiac Navigation System (Biosense Webster), and open heart surgery. Biomaterial delivery using these techniques may increase the risk of complications, such as arrhythmia, coronary thrombosis, and embolic events, and device failure, such as catheter occlusion. Applying materials to the heart at the time of open heart surgery is feasible but expensive and invasive. There is an unmet need for safe and precise minimally invasive delivery methods for patients not undergoing surgical procedures.

Biomaterial delivery to the pericardial space has several potential advantages over traditional intracoronary, intramyocardial, and intravenous delivery. First, the pericardium is not a vascular space, eliminating the risk of embolization, stroke, and infarction. Second, therapeutics delivered to this space remain focused and concentrated onto the heart by the pericardium, limiting off-target delivery (21,22). Third, pericardial delivery allows a more liberal gelation time because gel components are not subjected to immediate vascular washout. Fourth, biomaterials may degrade slower in the pericardial...
localized to a 1-cm² area that was the site of pericardial puncture and was likely the result of the puncture injury and healing. Other areas of the pericardium and the myocardium appeared normal. **STUDY LIMITATIONS.** For other pericardial procedures, such as the Lariat device, prophylactic anti-inflammatory medications, such as colchicine, are commonly used. Although reported incidence of pericarditis is low, most patients are prophylactically treated with anti-inflammatory agents, specifically colchicine (28). In one study of the Lariat device, the reported incidence of pericarditis was 5% but 54% of patients were treated with colchicine (23,28). In the current study, we chose to treat our animals with colchicine based on a preliminary study that showed significant pericardial inflammation when anti-inflammatory agents were not used. Colchicine was selected over systemic steroids because of its efficacy at treating pleuritis and lack of significant side effects compared with steroids or other nonsteroidal anti-inflammatory agents. Future studies are needed to confirm the long-term safety of pericardial biomaterials.

There are other concerns specific to accessing and working within the pericardial space. Our technique is unable to form the delivery compartment in patients without a pericardium (post-surgical or congenitally absent). Large pericardial effusions could potentially be drained before biomaterial delivery, but the safety and device functionality would need to be investigated. The risks of this technique in the acute myocardial infarction period may be elevated because of ischemia-reperfusion-induced inflammation of the pericardium and myocardium. Despite these potential concerns, the safety of pericardial access has been demonstrated by the development of epicardial ablations, the Lariat device, and other procedures. Interventionalists and electrophysiologists have gained experience in this technique (23–25). Alternatives to percutaneous routes of pericardial access, such as right atrial exit, are under clinical investigation and may be safer in some patients (29). We had no difficulties with percutaneous pericardial access and no instances of myocardial puncture or coronary artery damage.

Many biomaterials close to translation for clinical use have a liquid or gel phase and could be delivered using our device. However there are some biomaterials in development, such as cell sheets and patches, which do not and would not be amenable to our delivery by our method. Our device is also not capable of delivering ultra violet light as required for gelation for some biomaterials, but could be modified to do so. Lastly, most materials have only been tested by intramyocardial or intracoronary injection, through which they are instilled into the myocardium. They may not be as effective if delivered to the epicardium. Although it is unlikely a single technique could accommodate all biomaterials for cardiac...
application, we believe our technique and device are applicable to many biomaterials. We also wish to advocate for early consideration of the technique for material delivery to facilitate rapid and safe translation of biomaterials into clinical use.

CONCLUSIONS

We have developed a novel method for delivering biomaterial gels to the heart through the pericardial space. Our platform can be used with many types of gels and is a new treatment strategy for CVD that may be particularly useful for emerging therapeutics, such as cytokines, miRNAs, and stem cells. Our minimally invasive epicardial hydrogel delivery technique takes advantage of the proximity of the pericardial space to the heart, and its relatively protected status. Minimally invasive, precise, and safe delivery techniques help facilitate translation of biomaterials into clinical use for CVD.

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REFERENCES


**KEY WORDS** biomaterials, device, hydrogel, pericardial delivery

**APPENDIX** For a supplemental figure, tables, and video, please see the online version of this article.