Mitral chordae tendineae force profile characterization using a posterior ventricular anchoring neochordal repair model for mitral regurgitation in a three-dimensional-printed ex vivo left heart simulator

Michael J. Paulsen a, Annabel M. Imbrie-Moore a,b, Hanjay Wang a, Jung Hwa Bae b, Camille E. Hironaka a, Justin M. Farry a, Haley J. Luciani a, Akshara D. Thakore a, John W. MacArthur Jr a, Mark R. Cutkosky b and Y. Joseph Woo a,c,*

a Department of Cardiothoracic Surgery, Stanford University, Stanford, CA, USA
b Department of Mechanical Engineering, Stanford University, Stanford, CA, USA
c Department of Bioengineering, Stanford University, Stanford, CA, USA

* Corresponding author. Departments of Cardiothoracic Surgery and Bioengineering, Stanford University, Falk Cardiovascular Research Building CV-235, 300 Pasteur Drive, Stanford, CA 94305-5407, USA. Tel: +1-650-7253828, fax: +1-650-7360901; e-mail: yjwoo@stanford.edu (Y. J. Woo).

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Abstract

OBJECTIVES: Posterior ventricular anchoring neochordal (PVAN) repair is a non-resectional technique for correcting mitral regurgitation (MR) due to posterior leaflet prolapse, utilizing a single suture anchored in the myocardium behind the leaflet. This technique has demonstrated clinical efficacy, although a theoretical limitation is stability of the anchoring suture. We hypothesize that the PVAN suture positions the leaflet for coaptation, after which forces are distributed evenly with low repair suture forces.

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METHODS: Porcine mitral valves were mounted in a 3-dimensional-printed heart simulator and chordal forces, haemodynamics and echocardiography were collected at baseline, after inducing MR by severing chordae, and after PVAN repair. Repair suture forces were measured with a force-sensing post positioned to mimic in vivo suture placement. Forces required to pull the myocardial suture free were also determined.

RESULTS: Relative primary and secondary chordae forces on both leaflets were elevated during prolapse (P < 0.05). PVAN repair eliminated MR in all valves and normalized chordae forces to baseline levels on anterior primary (0.37 ± 0.23 to 0.22 ± 0.09 N, P < 0.05), posterior primary (0.62 ± 0.37 to 0.14 ± 0.05 N, P = 0.001), anterior secondary (1.48 ± 0.52 to 0.85 ± 0.43 N, P < 0.001) and posterior secondary chordae (1.42 ± 0.69 to 0.59 ± 0.17 N, P = 0.005). Repair suture forces were minimal, even compared to normal primary chordae forces (0.08 ± 0.04 vs 0.19 ± 0.08 N, P = 0.002), and were 90 times smaller than maximum forces tolerated by the myocardium (0.08 ± 0.04 vs 6.9 ± 1.3 N, P < 0.001).

DISCUSSION: PVAN repair eliminates MR by positioning the posterior leaflet for coaptation, distributing forces throughout the valve. Given extremely low measured forces, the strength of the repair suture and the myocardium is not a limitation.

Keywords: Mitral valve repair • Chordae tendineae forces • Biomechanics • Mitral valve surgery

ABBREVIATIONS

FBG Fibre Bragg Grating
LV Left ventricular
MR Mitral regurgitation
PTFE Polytetrafluoroethylene
PVAN Posterior ventricular anchoring neochordal
SAM Systolic anterior motion

INTRODUCTION

Since being reported in the 1950s, mitral valve repair has become a highly reproducible and effective treatment for degenerative mitral regurgitation (MR) in even the most complex valvular lesions [1–6]. Many techniques exist ranging from extensive resections to elaborate geometric reconstructions to completely non-resectional techniques [7–9]. Resection is highly effective, but minor drawbacks include being irreversible and time-consuming for complex repairs. In addition, overly aggressive resection may result in malcoaptation function, whereas insufficient resection may result in systolic anterior motion (SAM) [10–12]. Neochordal replacement with polytetrafluoroethylene (PTFE) suture is oftentimes faster, but requires precise measurements to ensure a proper coaptation plane and prevention of SAM [13, 14]. More extensive and complex techniques can also be more challenging in minimally invasive approaches. To remedy some of these drawbacks, we have previously described a simplified, non-resectional leaflet-remodelling technique whereby redundant, prolapsing leaflet tissue is imbricated to create a smooth coaptation surface to repair degenerative MR [15–17]. Because this repair keeps potentially diseased chordal tissue in place, it carries the theoretical risk of continued chordal elongation resulting in SAM or late prolapse. A further iteration on this technique that was first described in the ‘European Journal of Cardio-Thoracic Surgery’ in 2013—the posterior ventricular anchoring neochordal (PVAN) repair—remedied this risk by using a single suture to anchor the remodelled posterior leaflet to the posterior left ventricular (LV) wall (Fig. 1A) [18]. This technique has proven to be highly effective and particularly useful in minimally invasive approaches, though some concern has been raised about the stability of the suture anchored in the posterior myocardium as opposed to the typical neochord anchoring location in the fibrous portion of a papillary muscle. We hypothesize, however, that the PVAN repair suture serves primarily to position the leaflet posteriorly for proper coaptation and to prevent SAM (Fig. 1B), after which point systolic forces will be evenly distributed throughout the mitral valve apparatus due to the structural optimization provided by imbrication of excess leaflet tissue, resulting in low peak forces on the PVAN repair suture and the underlying myocardial anchoring point (Fig. 1C and D). In this study, we used a novel 3-dimensional (3D)-printed ex vivo left heart simulator with high-fidelity fibre-optic force sensors with embedded Bragg gratings to validate our theory.

MATERIALS AND METHODS

Left heart simulator

We designed a customized left heart chamber and prototyped the device using 3D printing (Carbon M2, Carbon3D Inc., Redwood City, CA, USA) and milling, which we have previously described (Fig. 2A) [19, 20]. Briefly, the chamber was mounted to a pulsatile linear actuator (VIVitro Superpump, VIVitro Labs, Victoria, BC, Canada) and outfitted with left atrial, ventricular and aortic pressure transducers (Utah Medical Products Inc., Midvale, UT, USA), as well as electromagnetic flow probes (Carolina Medical Electronics Inc., East Bend, NC, USA) in the mitral and aortic positions. Normal saline was used as our test fluid to ensure proper flow probe transduction. Compliance chambers in the aortic root and aortic positions, as well as a viscoelastic impedance adapter, allowed precise tuning of the waveforms. In the aortic position, a 29-mm mechanical aortic valve (St. Jude Regent, Abbott Vascular, Lake Bluff, IL, USA) was placed. Using a 28-mm leafless disc valve (VIVitro) in the mitral position as our reference valve, the system was tuned to generate physiological pressure and flow waveforms with systolic pressure of 120 mmHg, diastolic pressure of 80 mmHg and cardiac output of 5 l/min.

Sample preparation

Fresh porcine hearts were obtained from a local abattoir and the mitral valve apparatus was dissected free, including the papillary muscles and chordae tendineae. Only valves with intercommissural distances of 34–36 mm were included (n = 8). A 5-mm cuff of left atrium was left in place and used to sew the valves to 3D-printed elastomeric sewing rings that modelled the shape and material properties of the native mitral annulus.
Figure 1: Rendered illustration of the posterior ventricular anchoring neochordal (PVAN) repair and the hypothesized forces experienced on the repair suture throughout the cardiac cycle. During diastole (A), the PVAN suture is relaxed and experiences no tension. As systole begins (B), the closing posterior leaflet is held in position for optimal coaptation by the PVAN repair suture. Forces on the PVAN suture peak immediately before leaflet coaptation. Once the anterior and posterior leaflets coapt (C), the forces on the PVAN suture begin to fall as the leaflet forces are distributed throughout the 2 opposing leaflets (D). Images modified with permission and licensed from 3D4Medical Ltd.

Figure 2: (A) Schematic of the Stanford Left Heart Simulator with each major component labelled. Reprinted with permission from [21]. Copyright 2019 Elsevier. (B) Computer-aided design rendering of the force-sensing post with embedded FBG sensor for measuring strain on the PVAN repair suture. (C) Ex vivo experimental set-up showing the PVAN repair suture anchored to the force-sensing post with embedded FBG sensor, which has been positioned to mimic the correct in vivo placement of the PVAN repair suture. FBG: Fibre Bragg Grating; PTFE: polytetrafluoroethylene; PVAN: posterior ventricular anchoring neochordal; TOE: transoesophageal echocardiography.
Importantly, the native mitral annulus was not directly attached to the sewing ring, so as not to overly limit the motion of the native annulus, hence the reason for using the cuff of the left atrium to attach the valves to the sewing rings. The valves were first tacked to the sewing ring in the proper position with 6–8 interrupted horizontal mattress sutures using 2-0 braided polyester suture, followed by a continuous running 2-0 polypropylene suture line sealing the left atrial cuff and sewing ring. The papillary muscles were sewn to silicone sewing pads with pledgeted 2-0 braided polyester suture; the papillary muscle pads were inserted onto the ends of carbon fibre positioning rods, which passed into the chamber through spherical compression gaskets allowing for adjustment.

**Posterior ventricular anchoring neochordal and chordae tendineae force measurements**

We developed customized chordae tendineae force sensors with high accuracy and a small footprint using Fibre Bragg Grating (FBG) sensors, which we have described previously [20]. Briefly, FBGs are optical fibres with a series of spatial period gratings that reflect a particular wavelength of light transmitted by an optical interrogator. When tensile or compressive strain is applied, the space between the gratings changes, altering the reflected wavelength. The wavelength shift is directly proportional to strain, which can be calibrated to force applied. Using FBGs as a base structure, we developed an outer shell that allows for reusable attachment of the sensors to chordae. For each valve tested, multiple chordae were instrumented with FBGs (5–6 per valve), including a range of anterior, posterior, primary and secondary chordae. PTFE suture (CV-5 or CV-7, depending on chordae size) was used to sew the sensors to the chordae proximally and distally to the sensor. Once attached, the chordae segment between the sutures was cut, allowing the sensor to transmit the entire chordae load. As the mitral apparatus was dissected from the native hearts and mounted in the simulator, no LV wall was present to use for PVAN anchoring. Instead, we created a 3D-printed force-sensing post to simultaneously measure forces on and position the PVAN suture (Fig. 2B). The anchoring post was positioned between the papillary muscles ~10 mm posterior and 10 mm inferior to the P2 segment of the posterior leaflet.

Supplementary experiments explored how measured forces on the suture correspond to the ultimate strength of the suture anchoring point in the LV wall. Porcine myocardium (n = 6) was explanted from the posterior LV wall, between the papillary muscles inferior to the annulus. A CV-5 PTFE suture was passed into the myocardium to an approximate depth of 3–4 mm and then loosely tied in accordance with the PVAN technique. The force-sensing post was positioned to mimic the location of the LV anchoring point (Fig. 2C), allowing for accurate placement of the PVAN suture whilst measuring forces. The PVAN suture was then passed through the posterior leaflet and used to imbricate the prolapsed segment before being tied. Given that the experimental valves had normal annular dimensions, ring annuloplasty was not performed to limit confounding factors influencing the repair. Data collection was repeated following repair.

**Data acquisition and analysis**

Haemodynamic data were acquired using a data acquisition system and included software (ViVitro). Strain measurements were measured with an optical interrogator (Micron Optics si255; Micron Optics, Atlanta, GA, USA) at 1000 Hz, and converted into forces using calibration plots obtained during sensor manufacture. Waveforms were imported into MATLAB (R2019a, MathWorks Inc., Natick, MA, USA) for signal processing, composite plotting and summary data generation. Summary data were imported into R for statistical analysis (R 3.6.0. with Jamovi 0.9.6.9). We used a linear mixed-model analysis fit by the restricted maximum-likelihood model with unadjusted post-hoc testing for multiple comparisons to statistically compare the groups using a repeated-measures design [22]. This technique allows for a repeated-measure design (in this case, taking measurements from the baseline condition, prolapse condition and repair condition) much like a repeated-measures ANOVA, but also allows the analysis to incorporate the clustering found from within each valve tested. Because each valve tested is different, which may influence the measurements taken from this valve, it is important to account for these differences. Lastly, because of anatomical considerations, it was not always possible to test an anterior primary, posterior primary, anterior secondary and posterior secondary chordae from each individual valve, resulting in missing values in the data. Repeated-measures ANOVA uses list-wise deletion to address missing data, which can result in significant bias, whereas linear mixed models do not. As such, bias can be reduced and power increased through the use of a linear mixed-model design. Experimental condition (grouped into baseline, prolapse and repair conditions) and chordae location (grouped into anterior primary, posterior primary, anterior secondary and posterior secondary) were the fixed effects and valve number was the random effect. Data are normal and homoscedastic unless...
noted and are reported as mean ± standard deviation with a $P$-value <0.05 being considered significant.

**RESULTS**

The PVAN technique eliminated MR in all valves ($n = 8$). Visually, PVAN repair corrected P2 prolapse (Fig. 3A and Video 1), which was confirmed with echocardiography (Fig. 3B and Video 2). For more sensitive measurement of MR, regurgitant fraction was calculated from flow waveforms generated by the electromagnetic flow meters (Fig. 3C). Regurgitant fraction decreased from $30.3 ± 6.5\%$ during prolapse to $9.1 ± 4.0\%$ following PVAN repair ($P < 0.001$), which was equivalent to baseline levels in normal valves ($8.4 ± 1.5\%$, $P = 0.79$). Gross haemodynamic parameters, including pressure and flow waveforms, are plotted in Fig. 4A and B. The shaded regions represent the standard deviation, which suggests that PVAN repair maintains baseline haemodynamic properties, as the tracings are superimposable. Other haemodynamic parameters are summarized in Table 1, demonstrating no significant difference in any parameters between baseline and following PVAN repair. PVAN did not have a restrictive effect on the mitral valve, as effective orifice area was maintained following PVAN repair versus baseline ($6.8 ± 3.5$ vs $4.5 ± 3.4$, $P = 0.12$).

Composite force tracings are shown in Fig. 5. In the prolapse condition, as expected, forces normalized to ventricular pressure were significantly elevated versus baseline force measurements in

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**Figure 3:** High-speed videography still frames of the mitral valve from the surgeons’ view during prolapse (A, top) showing isolated P2 prolapse with ruptured (cut) chordae. Valve competency is restored following PVAN repair (A, bottom). Two-dimensional colour Doppler echocardiography images demonstrate moderate-severe mitral regurgitation (MR) secondary to posterior leaflet prolapse with an eccentric jet (B, top). PVAN repair corrected MR in all experiments. The repair suture attached to the force-sensing post can be appreciated in the echocardiography images after PVAN repair, with correction of posterior leaflet prolapse (B, bottom). Regurgitant fraction (C) measured by an electromagnetic flow probe demonstrates that PVAN repair effectively corrects MR to baseline levels ($8.4 ± 1.5\%$ vs $30.3 ± 6.5\%$ vs $9.1 ± 4.0\%$). *$P < 0.001$.

AL: anterior leaflet; LA: left atrium; LV: left ventricle; PL: posterior leaflet; PVAN: posterior ventricular anchoring neochordal.

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**Video 1:** High speed video demonstrating isolated P2 prolapse model of mitral regurgitation before and after posterior ventricular anchoring neochordal repair.

**Video 2:** Echocardiographic views of the mitral valve with color doppler prior to and after posterior ventricular anchoring neochordal repair.
Table 1: Haemodynamic parameters and chordae tendineae force measurements

| Haemodynamics | Baseline (n = 8) | Prolapse (n = 8) | PVAN (n = 8) | P-value |
|---------------|-----------------|-----------------|--------------|---------|
| Heart rate (bpm) | 70.0 ± 0.0 | 70.0 ± 0.0 | 70.0 ± 0.0 | 1.000 |
| Mean arterial pressure (mmHg) | 100.2 ± 1.2 | 77.3 ± 9.7 | 97.6 ± 3.5 | <0.001 | 0.395 | <0.001 |
| Systolic pressure (mmHg) | 120.9 ± 2.1 | 95.4 ± 10.0 | 118.4 ± 4.9 | <0.001 | 0.465 | <0.001 |
| Diastolic pressure (mmHg) | 80.7 ± 2.1 | 60.9 ± 9.8 | 77.9 ± 2.3 | <0.001 | 0.368 | <0.001 |
| Mean atrial pressure (mmHg) | 5.9 ± 3.3 | 4.9 ± 2.6 | 6.5 ± 3.9 | 0.310 | 0.512 | 0.11 |
| Mean ventricular pressure (mmHg) | 50.7 ± 3.5 | 41.0 ± 4.6 | 50.1 ± 4.1 | <0.001 | 0.696 | <0.001 |
| Maximum ventricular pressure (mmHg) | 136.9 ± 14.3 | 112.2 ± 19.8 | 136.1 ± 13.9 | <0.001 | 0.831 | <0.001 |
| Cardiac output (l/min) | 4.5 ± 0.3 | 3.7 ± 0.7 | 4.5 ± 0.3 | <0.001 | 0.923 | <0.001 |
| Effective stroke volume (ml) | 64.9 ± 4.7 | 53.4 ± 9.8 | 64.7 ± 4.3 | <0.001 | 0.923 | <0.001 |
| Pump stroke volume (ml) | 103.5 ± 13.9 | 103.6 ± 13.9 | 103.5 ± 13.9 | 0.788 | 0.969 | 0.76 |
| Mean transmural pressure (mmHg) | 0.3 ± 1.6 | 0.0 ± 1.7 | 1.3 ± 1.8 | 0.567 | 0.112 | 0.039 |
| Mean transmural back pressure (mmHg) | 114.1 ± 10.5 | 88.7 ± 15.9 | 111.8 ± 9.9 | <0.001 | 0.425 | <0.001 |
| Mitral forward flow time (s) | 0.53 ± 0.02 | 0.51 ± 0.01 | 0.55 ± 0.03 | 0.041 | 0.194 | 0.003 |
| Effective orifice area (cm²) | 6.8 ± 3.5 | 6.5 ± 3.3 | 4.5 ± 3.4 | 0.826 | 0.118 | 0.17 |
| Mitral valve regurgitant fraction (%) | 8.4 ± 1.5 | 30.3 ± 6.5 | 9.1 ± 4.0 | <0.001 | 0.788 | <0.001 |
| Mitral forward volume (ml) | 70.9 ± 5.2 | 76.3 ± 10.3 | 71.2 ± 4.6 | 0.021 | 0.877 | 0.029 |
| Mitral closing volume (ml) | -5.5 ± 0.6 | -7.1 ± 1.5 | -6.0 ± 1.6 | 0.029 | 0.494 | 0.11 |
| Mitral leakage volume (ml) | -0.5 ± 1.4 | -15.7 ± 5.7 | -0.5 ± 1.8 | <0.001 | 0.984 | <0.001 |
| Mitral leakage rate (ml) | -1.8 ± 5.1 | -55.5 ± 18.3 | -2.0 ± 7.1 | <0.001 | 0.974 | <0.001 |
| Transmitral leakage energy loss (mJ) | 10.6 ± 22.5 | 182.1 ± 45.6 | 11.7 ± 30.0 | <0.001 | 0.936 | <0.001 |
| Chordae tendineae forces | | | | |
| Anterior primary (N) | 0.23 ± 0.08 | 0.37 ± 0.23<sup>a</sup> | 0.22 ± 0.09 | 0.046 | 0.791 | 0.028 |
| Posterior primary (N) | 0.14 ± 0.06 | 0.62 ± 0.37<sup>a</sup> | 0.14 ± 0.05 | 0.001 | 0.978 | 0.001 |
| Anterior secondary (N) | 0.93 ± 0.38 | 1.48 ± 0.52<sup>a</sup> | 0.85 ± 0.43 | <0.001 | 0.520 | <0.001 |
| Posterior secondary (N) | 0.76 ± 0.28 | 1.42 ± 0.69<sup>a</sup> | 0.59 ± 0.17 | 0.005 | 0.376 | 0.001 |
| PVAN repair suture (N) | 0.08 ± 0.04 | | | |

<sup>a</sup>Prolapse chordae force measurements normalized to baseline ventricular pressures for comparative purposes in the setting of acute mitral regurgitation. Bold denotes P-value <0.05.

bpm: beats per minute; PVAN: posterior ventricular anchoring neochordal.

Figure 4: Composite pressure tracings (A) were indistinguishable between baseline and following PVAN repair, confirming restoration of aortic, left ventricular and left atrial pressures to baseline levels after PVAN repair. No significant difference was found in mean flow (B) between the baseline and PVAN repair groups. The shaded regions represent standard deviation. PVAN: posterior ventricular anchoring neochordal.
anterior primary (0.23 ± 0.08 vs 0.37 ± 0.23 N, P = 0.046), posterior primary (0.14 ± 0.06 vs 0.62 ± 0.37 N, P = 0.001), anterior secondary (0.93 ± 0.38 vs 1.48 ± 0.52 N, P < 0.001) and posterior secondary (0.76 ± 0.28 vs 1.42 ± 0.69 N, P = 0.005). Following PVAN repair, forces returned to baseline levels in all conditions. Forces on the PVAN repair suture itself were minimal with mean peak forces of 0.08 ± 0.04 N, which is significantly less than even forces experienced by primary chordae in healthy valves (0.19 ± 0.08 N, P = 0.002). The force required to pull the PVAN repair suture free from the posterior ventricular tissue was significantly higher at 6.9 ± 1.3 N (P < 0.001, Fig. 6).

**DISCUSSION**

The PVAN repair is an effective technique for correcting MR secondary to posterior leaflet prolapse and requiring only a single suture makes the technique especially attractive for minimally invasive approaches. By positioning the imbricated and structurally optimized leaflet posteriorly, the PVAN repair suture effectively prevents SAM and maintains ideal leaflet height for maximal coaptation. In addition to restoring haemodynamic parameters, the PVAN repair normalizes chordae forces to baseline levels; whether this translates to enhanced durability of this technique over others is unknown. Unlike traditional neochord repairs anchored in the fibrous head of a papillary muscle, the PVAN suture is anchored in the posterior ventricular myocardium. A theoretical concern of PVAN repair is the strength of the ventricular anchor given that the repair relies on this suture. While the imbrication of excess leaflet tissue structurally optimizes the leaflet for coaptation to prevent leaflet prolapse, we hypothesized that the PVAN repair suture itself does not play a significant structural role in the repair, and like native primary chordae, serves to position the leaflet for ideal coaptation, after which point systolic forces are distributed throughout the mitral apparatus [12, 23, 24]. The results of our
study support this hypothesis, with forces being nearly zero in diastole, peaking in early systole as the leaflet is positioned for coaptation, and then decreasing for the remainder of systole as forces are distributed throughout the valve apparatus (Fig. 7). We found that the force on the PVAN suture is extremely small, even compared to forces experienced by primary chordae of healthy valves, and the force required to tear the PVAN suture free from the posterior LV wall with resultant repair failure is over 90 times higher than the mean peak force on the repair suture. Although tissues experiencing cyclic stresses below their yield strength may still accumulate damage, the large discrepancy in suture force compared to pull-out force suggests that the strength of the posterior ventricular anchoring tissue and the forces exerted on the repair suture should not limit durability of this repair [25].

Other benefits of the PVAN technique include it being non-resectional; if during the operation the repair is felt to be inadequate, the repair suture can be modified, repositioned, or removed without damaging the leaflet. Resection, on the contrary, is irreversible and a suboptimal repair can result in unintended valve replacement [26]. Unlike traditional papillary muscle-based neochord repairs where precise suture length is required to prevent excessive leaflet tissue from causing SAM, the PVAN technique results in a posteriorly positioned leaflet, substantially reducing the risk of SAM [27]. No significant difference in effective orifice area when comparing baseline values to those following PVAN repair, though EOA did appear to trend downwards slightly, though not into ranges concerning for iatrogenic mitral stenosis as demonstrated by extremely low transmitral gradients in either group. Furthermore, we did not identify any significant differences in any other haemodynamic parameter when comparing the baseline and PVAN groups, which suggests that the magnitude of this downwards trend in EOA is unlikely to be large enough to have clinical significance. Lastly, the addition of an annuloplasty ring would also likely reduce EOA for any repair, regardless of surgical technique used. Future studies investigating the influence of annuloplasty ring on the PVAN repair, among many others, are currently underway.

Limitations

As an ex vivo investigation, our study does have some limitations in perfectly recreating the complex interactions between the annulus and ventricular components of the mitral valve apparatus, though our system does allow for native annular motion. The valves used in this study were also normal, healthy valves and P2 prolapse was induced by cutting chordae as opposed to natural pathological processes resulting in chordal rupture typical of degenerative valve disease. Nonetheless, this is an accepted model of MR commonly used experimentally [28, 29]. Future in vivo investigations currently underway will address some of these limitations. Lastly, we used porcine and not human valves. However, porcine and human mitral valves are remarkably similar, with the anterior leaflet attaching to approximately one-third of the annulus with a fibrous aortomitral curtain, the posterior leaflet attaching to the posterior two-thirds of the annulus, and the papillary muscles having comparable locations [30]. Chordae number and histological composition are also similar between these species [31]. As such, porcine mitral valves are realistic surrogates for human mitral valves in experiments not feasible in humans. While measuring forces on individual human valves in vivo is not currently possible, further validation using clinical outcomes data is indicated and currently underway.

CONCLUSION

This exploratory ex vivo study helps to provide biomechanical validation of the PVAN repair technique. In addition to restoring haemodynamic variables to baseline levels, PVAN repair results in normalization of chordae forces to baseline levels. By positioning the leaflet posteriorly, this technique is also extremely effective in preventing SAM. The theoretical concern regarding the strength of the repair suture and anchoring tissue in this technique should not be a limiting factor, as the forces experienced by the repair suture are extremely small relative to the maximum forces that can be supported by the posterior ventricular myocardium. Overall, the PVAN repair is an...
effective, non-resectional, single-suture mitral valve repair technique that lends itself well to a minimally invasive approach and is biomechanically sound.

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Author contributions

Michael J. Paulsen: Conceptualization; Data curation; Formal analysis; Funding acquisition; Investigation; Methodology; Project administration; Resources; Software; Supervision; Validation; Visualization; Writing – original draft; Writing – review & editing.

Annabel M. Imbrie-Moore: Conceptualization; Data curation; Formal analysis; Funding acquisition; Investigation; Methodology; Project administration; Resources; Software; Supervision; Validation; Visualization; Writing – original draft; Writing – review & editing.

Hanjay Wang: Conceptualization; Funding acquisition; Investigation; Methodology; Project administration; Resources; Software; Supervision; Validation; Visualization; Writing – original draft; Writing – review & editing.

Jung Hwa Bae: Conceptualization; Data curation; Investigation; Methodology; Resources; Software; Supervision; Validation; Visualization; Writing – original draft; Writing – review & editing.

Camille E. Hironaka: Data curation; Formal analysis; Investigation; Validation; Visualization; Writing – review & editing.

Justin M. Farry: Data curation; Formal analysis; Investigation; Methodology; Software; Writing – review & editing.

Haley J. Lucian: Data curation; Formal analysis; Investigation; Methodology; Software; Writing – review & editing.

Akhsha D. Thakore: Data curation; Formal analysis; Investigation; Methodology; Software; Writing – review & editing.

John W. MacArthur: Conceptualization; Formal analysis; Methodology; Supervision; Writing – review & editing.

Mark R. Cutkosky: Conceptualization; Data curation; Formal analysis; Investigation; Methodology; Project administration; Resources; Software; Supervision; Validation; Visualization; Writing – review & editing.

Y. Joseph Woo: Conceptualization; Data curation; Formal analysis; Funding acquisition; Investigation; Methodology; Project administration; Resources; Software; Supervision; Validation; Visualization; Writing – original draft; Writing – review & editing.

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