Percutaneous approach to left ventricular assist device decommissioning

Francesco Moroni MD1 | Keyur B. Shah MD1 | Mohammed A. Quader MD2 | Katherine Klein MD2 | Melissa C. Smallfield MD1 | Kendall E. Parris RT1 | Zachary M. Gertz MD1

1Division of Cardiovascular Medicine, Department of Medicine, Virginia Commonwealth University, Richmond, Virginia, USA
2Division of Cardiothoracic Surgery, Virginia Commonwealth University, Richmond, Virginia, USA

Correspondence
Zachary M. Gertz, MD, VCU Pauley Heart Center, 1200 East Broad St, Box 980036, Richmond, VA 23298, USA.
Email: zachary.gertz@vcuhealth.org

Abstract
Objective: To assess the outcomes of a single-center experience with percutaneous left ventricular assist device (LVAD) decommissioning.

Background: Patients with LVADs may eventually require their removal, either due to recovery of left ventricular function or recurrent complications. Traditionally, withdrawal of LVAD support has been managed with surgical device explantation, which carries significant procedural risks. Transcatheter LVAD decommissioning, with outflow graft occlusion and driveline transection, has recently been described as an alternative to surgical removal.

Methods: Here, we report on a retrospective cohort of five consecutive cases treated with transcatheter LVAD decommissioning.

Results: The procedure was effective in all cases, and no patient experienced procedure-related complications. At midterm follow-up, the three patients who had myocardial function recovery were alive and had not experienced heart failure-related symptoms or complications.

Conclusion: Percutaneous LVAD decommissioning appears to be a safe and effective approach to LVAD treatment discontinuation.

KEYWORDS
cardiac function, cardiac remodeling, heart assist device, heart failure, ventricular assist device

1 INTRODUCTION

For patients with end-stage heart failure, left ventricular assist devices (LVADs) improve quality of life and extend survival as either a bridge to transplantation or destination therapy. In a minority of patients, approximately 1% at 1 year of support, cardiac function recovers, and the LVAD is no longer necessary. An LVAD cannot simply be turned off, as the conduit from ascending aorta to left ventricle would result in severe regurgitation. The traditional approach to withdrawal of LVAD support has been open surgical pump removal, but the inherent risks of mediastinal dissection and apical ventriculoplasty result in an operative mortality of around 10%. A less invasive approach, LVAD decommissioning, involves LVAD deactivation and outflow graft ligation, which is generally achieved via a right thoracotomy or subcostal surgical access. This is followed by severing the driveline at the exit site and leaving the device in place. To date, the experience with
decommissioning is limited, but follow-up data suggest comparable midterm outcomes with respect to LVAD explantation.\(^5\) More recently, a transcatheter approach to LVAD decommissioning, with occlusion of the outflow graft followed by surgical driveline removal, has been described.\(^6\) This approach avoids chest re-entry altogether, potentially increasing procedural safety. The published experience of transcatheter LVAD decommissioning is limited, with few cases reported worldwide.\(^7\)–\(^9\) Here, we present our experience with transcatheter LVAD decommissioning and report acute and midterm follow-up of a cohort of five patients treated at a single institution. Since beginning this approach, it was used in all patients referred for LVAD decommissioning.

2 | METHODS

All procedures were carried out under conscious sedation. Right common femoral artery access was obtained under ultrasound guidance and an 8F × 90 cm sheath was advanced to the aortic insertion of the outflow graft. A Judkins Right 4 or Multipurpose diagnostic catheter was used to direct a Glide wire into the outflow graft, and the sheath was advanced over the diagnostic catheter and the wire up to the proximal end of the outflow graft, where it exits the pump housing. The catheter and wire were then removed. After turning off the LVAD, the outflow graft was occluded with Amplatzer vascular plugs (AVPs; Abbott Cardiovascular—), typically an AVP-1 for the proximal and mid portion, but an AVP-2 for the aortic anastomosis, to be as flush as possible with the aorta and minimize any residual pouch (Figure 1). Initially, we tried two plugs per patient (at the proximal and distal graft insertions) with an occasional third plug (mid graft), but because some outflow grafts took longer to thrombose, we began routinely using three plugs per case. A fourth plug was used in one case where the third plug was not flush with the aortic anastomosis. In a single patient, the sheath could not reach the proximal outflow graft hence the devices were deployed through an 8F Multipurpose guide catheter. No systemic anticoagulation was administered during the procedure, to ensure rapid vascular plug occlusion; no wire remained exposed in the aorta to minimize the risk of thrombosis, and the sheath was regularly flushed with heparinized saline. Three subjects underwent concomitant right

![Image](image-url)
heart catheterization, from an internal jugular vein approach, to monitor the hemodynamic effects of each stage of the procedure (Figure 2). At the end of the procedure, sheaths were removed with deployment of Angioseal closure devices.

All subjects underwent driveline truncation and surgical debridement and closure of the driveline tract after a median of 4 days from graft occlusion (range: 1–8 days). All procedures were successful, with no acute complications. All patients were discharged from the hospital alive after a median of 5 days (range: 3–14 days). Antithrombotic treatment of choice was warfarin (target international normalized ratio between 2 and 3) in three cases, and apixaban 5 mg twice daily in two cases.

3 | RESULTS

3.1 | Patient population

Between October 2018 and August 2021, five patients underwent percutaneous LVAD decommissioning at our institution.

Median age at time of decommissioning was 53 years (range: 45–64) and three patients were male. LVAD support duration ranged between 203 and 2096 days at time of decommissioning. Reason for LVAD decommissioning was heart function recovery in three cases, resistant, intractable LVAD thrombosis treated with failed pump exchange in one case, and end-stage heart failure with patient preference for palliative care in another case. Table 1 reports the clinical characteristics of the patients, and Table 2 reports the devices employed in each subject.

3.2 | Clinical follow-up

Clinical follow-up was available for all subjects. Patient 1, who had been discharged to hospice care after LVAD decommissioning in the setting of severe, end-stage biventricular heart failure, died 27 days after the intervention. Patient 3, who underwent LVAD decommissioning for refractory LVAD thrombosis, was discharged on outpatient milrinone but experienced progressive worsening of...

**FIGURE 2** Aortic and pulmonary artery pressure tracings during different stages of percutaneous left ventricular assist device (LVAD) decommissioning. (A) Baseline tracings while the LVAD is active. Aortic pressure (red tracing) displays physiologic pulsatility, which is consistent with recovered native heart function. Periodic pressure dips superimposed on physiological pulsatile flow (arrowheads) are consistent with HeartMate 3 intrinsic pulsatility. (B) Upon deactivating the LVAD (white arrow), there is an abrupt decrease in mean aortic pressure (MAP) and increase in mean pulmonary artery pressure (PAP, azure tracing). (C) After LVAD deactivation but before outflow graft occlusion is complete, the MAP remains below baseline, while aortic pulse pressure increases and mean PAP rises. These findings are consistent with withdrawal of LVAD unloading and aortic regurgitation-like effect of retrograde flow through the device. (D) Upon effective occlusion of the outflow graft, aortic pulse pressure decreases and MAP increases, while PAP decreases, yet neither returns to baseline levels. This likely reflects the increased workload of the left ventricle in the absence of LVAD support. [Color figure can be viewed at wileyonlinelibrary.com]
biventricular heart failure leading to total artificial heart implant 162 days after the procedure. The remaining three subjects, who underwent decommissioning due to cardiac function recovery, were free from heart failure symptoms on the last follow-up (follow-up duration: Patient 2: 563 days; Patient 4: 173 days; and Patient 3: 138 days).

### DISCUSSION

Current experience with LVAD decommissioning is limited, with few reports of using a transcatheter approach. Previously published case reports and series of percutaneous LVAD decommissioning are summarized in Table 3. Our experience, applied to consecutive patients, builds on this previous experience and expands the evidence regarding this technique in several ways. First, we described a standardized, streamlined approach, using conscious sedation and a single arterial access. Previous reports described the use of bi-femoral access and transesophageal echocardiographic monitoring, which required general anesthesia. Second, we were able to show the safety of a heparin-free approach, which has the potential to reduce time to outflow graft occlusion, thereby minimizing hemodynamic instability. Third, we describe in detail the hemodynamic changes in both aortic and pulmonary artery pressures during the procedure, which can be used to monitor for adequate outflow graft occlusion. Finally, we showed that apixaban may be safe for thromboprophylaxis after LVAD decommissioning. The use of direct oral anticoagulants is not standard of care among patients with LVADs, yet they were shown to be safe and effective in a small series of patients at high bleeding risk. Further study is clearly necessary before recommending this approach for routine use.

LVAD decommissioning appears to be a lower risk procedure as compared to LVAD explantation, with no procedural mortality currently reported in the literature. In addition, leaving the device in place could maintain the apical orifice if reimplantation were eventually needed, while avoiding the development of adhesions that would occur after surgical explantation, making subsequent procedures even more difficult. Both considerations make transcatheter LVAD decommissioning an attractive option, especially for those subjects who are considered at high risk of heart failure recurrence. On the downside, leaving the device in place may expose patients to systemic thromboembolism due to the potential thrombogenicity of the inflow cannula, requires continued anticoagulation, and may pose an infection risk. Long-term follow-up involving more patients is required to evaluate the optimal treatment course in these patients.

### TABLE 1  Clinical characteristics of the patients

| Patient, n | Sex | Age | LVAD type | Baseline heart disease | Duration of support (days) | Reason for decommissioning |
|------------|-----|-----|-----------|------------------------|---------------------------|----------------------------|
| 1          | F   | 53  | Heartmate 2 | Cardiac sarcoidosis    | 2096                      | Severe, refractory, biventricular heart failure. Transition to palliative care as per patient preference. |
| 2          | M   | 57  | HVAD       | Idiopathic dilated cardiomyopathy | 203                    | Heart function recovery |
| 3          | F   | 51  | Heartmate 2 | Post-partum cardiomyopathy | 1659                   | Intractable pump thrombosis, failed thrombolysis, and pump exchange |
| 4          | M   | 64  | Heartmate 3 | Idiopathic dilated cardiomyopathy | 631                    | Heart function recovery |
| 5          | M   | 45  | Heartmate 3 | Alcohol-induced cardiomyopathy | 657                     | Heart function recovery |

Abbreviations: F, female; LVAD, left ventricular assist device; M, male.

### TABLE 2  Device type, size, and location used to occlude the outflow graft during the percutaneous stage of the left ventricular assist device (LVAD) decommissioning.

| Patient | LVAD type | Total number of AVPs | Proximal outflow graft | Mid outflow graft | Distal outflow graft | Antithrombotic treatment |
|---------|-----------|----------------------|------------------------|------------------|---------------------|-------------------------|
| 1       | Heartmate 2 | 3 | • AVP-1 16 mm | • AVP-1 16 mm | • AVP-2 16 mm | Warfarin |
| 2       | HVAD      | 2 | • AVP-1 16 mm | • AVP-2 16 mm | Warfarin |
| 3       | Heartmate 2 | 3 | • AVP-1 14 mm | • AVP-1 16 mm | • AVP-2 14 mm | Warfarin |
| 4       | Heartmate 3 | 3 | • AVP-1 14 mm | • AVP-1 16 mm | • AVP-2 14 mm | Apixaban |
| 5       | Heartmate 3 | 4 | • AVP-1 14 mm | • AVP-1 16 mm | • AVP-2 14 mm | Apixaban |

Abbreviation: AVP, Amplatzer vascular plug.
| Author, year       | LVAD Type | Indication for decommissioning | Closure devices | Device size | Antithrombotic medication | Outcomes                      |
|-------------------|-----------|--------------------------------|----------------|-------------|---------------------------|------------------------------|
| Zeigler et al., 2014 | HM II     | Recovery                       | 1× AVP II       | 22 mm       | Warfarin                  | Alive at 3.5 years           |
| Sainte et al., 2014 | CircuLite | Recovery                       | 1× AVP II       | Not reported | Not reported              | Alive at 2 years             |
| Pettit et al., 2015 | HVAD      | Recovery                       | 2× AVP II       | 14 mm       | Not reported              | Alive at 2 years             |
| El Sayed Ahmed, 2016 | HM II    | Recovery                       | 2× AVP II       | 16 and 14 mm| Warfarin                  | Alive at 6 months            |
| Grinstein et al., 2016 | HVAD    | Infection and thrombosis      | 1× ASO          | 14 mm       | None                      | Alive at 8 months (palliative care) |
| Soon et al., 2017  | HVAD      | Recovery                       | 2× AVP II       | 14 and 16 mm| Aspirin 100 mg daily      | Alive at 5 months            |
| Pendyal et al., 2017 | HM II    | Thrombosis                     | 1× AVP II       | 20 mm       | Warfarin                  | Transplant at 3 months       |
| Kidambi et al., 2018 | HM II    | Infection                      | 1× AVP II       | 22 mm       | Not reported              | Not reported                 |
| Chowdhury et al., 2020 | HM II    | Thrombosis (palliation)       | 3× ASO          | 20 and 2× 18 mm| Warfarin + ASA 325 mg    | In-hospital death            |
| HM II              | Recovery  | 2× AVP II                      | 14 mm           | Warfarin    | Alive at 1.5 years        |
| HVAD               | Thrombosis (palliation)  | 2× ASO                        | 12 mm           | Warfarin + ASA 325 mg    | In-hospital death            |
| HM II              | Recovery  | 2× AVP II                      | 20 and 18 mm    | Warfarin + ASA 325 mg    | Alive at 2 years             |
| HVAD               | Thrombosis | 3× AVP II                     | Not reported    | Not reported | Transplant at 2 months    |
| Albulushi et al., 2020 | HM II    | Thrombosis and recovery       | 3× AVP II       | Not reported | Not reported              | Alive at time of reporting   |
| HM II              | Recovery  | 3× AVP II                      | Not reported    | Not reported | Alive at time of reporting|
| HM III             | GI bleed and recovery  | 3× AVP II                     | Not reported    | Not reported | Transplant at 3 months    |
| HM II              | Recovery  | 3× AVP II                      | Not reported    | Not reported | Alive at time of reporting|
| HM III             | GI bleed and recovery  | 3× AVP II                     | Not reported    | Not reported | Alive at time of reporting|
| HM III             | Recovery  | 3× AVP II                      | Not reported    | Not reported | Alive at time of reporting|
| Alkattan et al., 2021 | HM III   | Recovery                       | 2× AVP II       | AVP II: 20 and 16 mm | Warfarin                  | Alive at 2 months            |
|                   |           |                               | 2× AVSD         | AVSD: 2× 12 mm |                            |                              |

Note: Relevant articles were identified through systematic search of MEDLINE using a combination of the following keywords: "LVAD", "left ventricular assist device," "heart function recovery," "discontinuation," "withdrawal," "decommissioning," and "percutaneous decommissioning."

Abbreviations: ASA, aspirin; ASO, Amplatzer septal occluder; AVP, Amplatzer vascular plug; AVSD, Amplatzer ventricular septal defect muscular occluder; HM II, HeartMate II; HM III, HeartMate III; HVAD, Medtronic HVAD Support system.
CONCLUSION

Transcatheter LVAD decommissioning is a promising approach to mechanical circulatory support withdrawal. Further studies are needed to assess the long-term result of transcatheter decommissioning.

CONFLICTS OF INTEREST
Dr. Shah reports receiving consulting honoraria from Akcea Therapeutics. Other authors report no conflicts of interest.

DATA AVAILABILITY STATEMENT
The data that support the findings of this study are available on request from the corresponding author.

ORCID
Francesco Moroni https://orcid.org/0000-0002-6101-1403
Zachary M. Gertz https://orcid.org/0000-0002-9613-5246

REFERENCES
1. Guglin M, Zucker MJ, Borlaug BA, et al. Evaluation for heart transplantation and LVAD implantation: JACC council perspectives. J Am Coll Cardiol. 2020;75(12):1471-1487.
2. Wever-Pinzon O, Drakos SG, McKellar SH, et al. Cardiac recovery during long-term left ventricular assist device support. J Am Coll Cardiol. 2016;68(14):1540-1553.
3. Phan K, Huo YR, Zhao DF, Yan TD, Tchantchaleishvili V. Ventricular recovery and pump explantation in patients supported by left ventricular assist devices: a systematic review. ASAIO J. 2016;62(3):219-231.
4. Khvilivitzky K, Mountis MM, Gonzalez-Stawinski GV. Heartmate II outflow graft ligation and driveline excision without pump removal for left ventricular recovery. Proc (Bayl Univ Med Cent). 2012;25(4):344-345.
5. Choi JH, Weber MP, Horan DP, et al. Left ventricular assist device decommissioning compared with explantation for ventricular recovery: a systematic review. ASAIO J. 2020;66(1):17-22.
6. Soon JL, Tan JL, Lim CP, et al. Percutaneous decommissioning of left ventricular assist device. Heart Lung Circ. 2018;27(7):853-855.
7. Chowdhury MA, Lindenfeld J, Shah AS, et al. Percutaneous ventricular assist device exclusion: institutional case series and review of literature. ASAIO J. 2020;66(4):e60-e61.
8. Gerhard EF, Wang L, Singh R, et al. LVAD decommissioning for myocardial recovery: Long-term ventricular remodeling and adverse events. J Heart Lung Transplant. 2021;40(12):1560-1570.
9. Albulushi A, Goldswieg AM, Stoller D, et al. Percutaneous deactivation of left ventricular assist devices. Semin Thorac Cardiovasc Surg. 2020;32(3):467-472.
10. Parikh VY, Parikh UM, Moctezuma-Ramirez A, et al. Factor Xa inhibitors in patients with continuous-flow left ventricular assist devices. Gen Thorac Cardiovasc Surg. 2020;68(11):1278-1284.
11. Zeigler SM, Sheikh AY, Lee PH, et al. A novel, catheter-based approach to left ventricular device deactivation after myocardial recovery. Ann Thorac Surg. 2014;98(2):710-713.
12. Sainte S, Gewillig M, Droogne W, et al. Exploration of a CircuLite left ventricular assist device without removal of the inflow cannula: how to do it? Interact Cardiovasc Thorac Surg. 2014;18(3):393-395.
13. Pettit SJ, Shapiro LM, Lewis C, Parameshwar JK, Tsui SS. Percutaneous withdrawal of HeartWare HVAD left ventricular assist device support. J Heart Lung Transplant. 2015;34(7):990-992.
14. El-Sayed Ahmed MM, Jones MB, Kanter JP, et al. Hybrid exclusion of HeartMate II left ventricular assist device after bridge to recovery. Ann Thorac Surg. 2016;101(6):e193-e194.
15. Grinstein J, Estrada J, Sayer G, et al. Left ventricular assist device deactivation via percutaneous closure of the outflow graft. J Card Fail. 2016;22(8):653-655.
16. Pendyal A, Chien CV, Mudd JO, Gelew JM. Minimally invasive LVAD deactivation in a 65-year-old man with recurrent pump thrombosis and left ventricular recovery. Tex Heart Inst J. 2017;44(1):70-72.
17. Kidambi S, Shudo Y, Dake MD, Woo YJ, Ha RV. Percutaneous, minimally invasive approach to implantable left ventricular assist device deactivation. J Thorac Cardiovasc Surg. 2018;155(2):653-654.
18. Alkattan HN, Kjellman U, Selimovic N, Alomrani A, Alghamdi AA. Deactivation of left ventricular assist device (LVAD) after recovery of cardiac function: a case report. J Card Surg. 2021;36(8):2974-2978.

How to cite this article: Moroni F, Shah KB, Quader MA, et al. Percutaneous approach to left ventricular assist device decommissioning. Catheter Cardiovasc Interv. 2022;100:169-174. doi:10.1002/ccd.30230