Multidisciplinary Approach to Hemodynamic Management During High-Risk Ventricular Tachycardia Ablation

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ABSTRACT

Percutaneous ventricular assist devices have been used for high-risk ventricular tachycardia ablation when hemodynamic decompensation is expected. Utilizing a case example, we present our experience with development of a coordinated, team-based approach focused on periprocedural management of patients with high-risk ventricular tachycardia.

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Ventricular tachycardia ablation (VTA) is increasingly performed in patients with advanced heart failure.1 Ablation often requires induction and mapping of ventricular arrhythmias and may be associated with acute hemodynamic decompensation (AHD). Mechanical circulatory support (MCS) during VTA has increased, and observational studies using the PAINESD score suggest a mortality benefit.2 The PAINESD score estimates the risk of AHD during VTA. Among patients with non-ischemic cardiomyopathy and a PAINESD score ≥15, MCS during VTA decreased 30-day rehospitalization, repeat ablation, recurrent implantable cardioverter-defibrillator (ICD) therapy, and 3-month mortality.3 Moreover, prophylactic implementation of MCS in high-risk patients is associated with a 3.5-fold reduction in mortality or need for heart transplantation.4,5

Although evidence supports MCS during high-risk VTA (HR-VTA), pathways for case selection, pre-procedural assessment, and multidisciplinary coordination have not been elucidated. Currently, there are no established weaning protocols to guide postprocedure care. We present a multidisciplinary approach for HR-VTA requiring MCS, including recommendations for weaning.

LEARNING OBJECTIVES

- Acute hemodynamic decompensation may occur during ablation of ventricular tachycardia irrespective of the mapping/ablation strategy.
- Objective measures of cardiac function and tissue perfusion can be used to assess the efficacy of mechanical circulatory support and guide weaning.
- A multidisciplinary approach to periprocedural hemodynamic management with objective measures of perfusion and a framework for weaning mechanical support ensures the best clinical outcome for the patient.
CASE PRESENTATION

A 66-year-old man with heart failure with reduced ejection fraction (20%-25%) due to ischemic cardiomyopathy, left ventricular (LV) aneurysm, and history of coronary artery bypass grafting, mitral valve repair, and closure of an atrial septal defect presented for evaluation of ventricular tachycardia (VT), resulting in appropriate ICD therapy. He had a history of VT storm treated with catheter ablation using a substrate-based approach and extracorporeal membrane oxygenation support 5 months prior, and had previously failed treatment with amiodarone, sotalol, and dofetilide. The patient presented after multiple ICD shocks for sustained monomorphic VT. He was evaluated by electrophysiology and considered a candidate for repeat VT ablation; however, the patient preferred medical therapy. He was discharged on sotalol and re-admitted 11 days later after receiving 4 shocks for monomorphic VT with a single morphology. At this point, the patient elected for repeat catheter ablation.

A multidisciplinary conference with electrophysiology and a heart failure cardiologist, interventional cardiologist, and cardiac anesthesiologist was convened. Electrophysiology recommended an activation and entrainment mapping strategy rather than a substrate-based approach given the recurrence of VT within 6 months of ablation. Pre-emptive MCS with a percutaneous ventricular assist device (pVAD) was planned because the patient was deemed high risk (PAINESD score of 17).

ANESTHESIA. Before induction, American Society of Anesthesiologists monitors were placed, cerebral oximetry was initiated, and the radial artery was cannulated. General anesthesia was induced with etomidate and succinylcholine, followed by endotracheal intubation. A pulmonary artery catheter was placed via the right internal jugular vein. Norepinephrine and epinephrine infusions were titrated to maintain blood pressure within 20% of baseline, cerebral saturation >60% and within 20% of baseline, as well as cardiac index >2 L/min/m², mixed venous oxygen saturation (SVO₂) >60%, cardiac power output (CPO) >0.6 W, and serum lactate <2 mmol/L.

PERCUTANEOUS VENTRICULAR SUPPORT. The right common femoral artery was accessed under ultrasound and fluoroscopic guidance, and angiography confirmed position before placement of a 6-F sheath. A pigtail catheter was positioned at the abdominal aortic bifurcation, and an angiogram was performed to evaluate for significant peripheral artery disease. Heparin was administered to maintain activated clotting time >300 seconds. The arteriotomy site was progressively dilated before placing a 14-F peel-away sheath. The pigtail catheter was advanced into the

FIGURE 1 Left Ventricular Mapping: Clinical Ventricular Tachycardia

(A) Left ventricular activation map of the clinical ventricular tachycardia. (B) Left ventricular voltage map showing large anterior scar extending to the septum along with ablation lesions. (C) Left bundle morphology, superior axis, precordial transition in V₅, and positive in lead I.
left ventricle, and a pVAD was placed with resultant 3.8 L/min flow.

**ABLATION.** Right femoral vein access was obtained under ultrasound guidance. A long, fixed curve sheath was used to perform a transseptal puncture and was exchanged for a large curl deflectable sheath. An irrigated force-sensing ablation catheter was advanced through the sheath into the left ventricle. A detailed bipolar voltage map of the left ventricle showed a large anterior wall scar corresponding to the location of the aneurysm (Figure 1). VT was induced and mapped using a multielectrode splined mapping catheter; the activation map identified a critical isthmus-based activation pattern and presence of mid-diastolic potentials. Ablation was performed, and the clinical VT was no longer inducible after targeted ablation. Additional ablation was performed around and within the scar with a core isolation approach. Pacing at high output confirmed the scar was electrically unexcitable postablation.

Induction was attempted after scar homogenization, and a second, morphologically distinct VT was induced. An activation map of the left ventricle was created, but the full cycle length of the tachycardia was not captured in the LV endocardium, suggesting an epicardial component of the circuit (Figure 2). Additional ablation was performed at the exit site based on findings of the activation map. After extensive endocardial ablation, a sustained nonclinical VT with an epicardial component remained inducible, but the initial clinical VT remained noninducible.

**WEANING MCS AND POSTPROCEDURE CARE.** Postablation, the CPO was 0.9 W, consistent with baseline, and was maintained as the pVAD was weaned from “Auto” to P2 over 40 minutes. On P2, the arterial pulsatility was preserved and mean arterial pressure was 70 mm Hg on norepinephrine 3 μg/kg/min, SVO2 was >60%, and serum lactate was 1.4 mmol/L. The pVAD was removed, and hemostasis was obtained with suture-based closure devices. The patient was extubated, and his care was transferred to the cardiac intensive care unit where norepinephrine was weaned, and his home heart failure medication regimen was restarted. He was discharged home 2 days’ postablation. At 1-year follow-up, he remains free of sustained VT and has not required ICD therapy.

**QUESTION 1: WHAT IS THE BENEFIT OF MCS DURING HR-VTA?**

MCS improves hemodynamics and continuously unloads the left ventricle; we believe this action translates into improved end-organ perfusion during HR-VTA. Indeed, cerebral desaturation has been observed during fast VT (tachycardia cycle length <300 milliseconds), but with MCS, the incidence of cerebral desaturation decreases significantly. Prophylactic MCS is superior to a rescue strategy, and 30-day mortality is higher with rescue compared with pre-emptive pVAD implantation in high-risk patients experiencing AHD during VTA. Even with successful rescue and improved hemodynamics, 40.2% of patients with AHD during VTA died within 30 days. Furthermore, pre-emptive MCS may
reduce inotropic and vasopressor usage and avoid the associated myocardial oxygen demand and impairment in tissue perfusion. This underscores the importance of preprocedure risk stratification and pre-emptive MCS. Pre-emptive MCS should be considered irrespective of VTA strategy, as activation and entrainment techniques as well as substrate mapping may precipitate AHD. We prefer pVAD because of the ease of placement and its ability to unload the left ventricle. The PAINESD score has been validated as an effective way to risk stratify patients; our use of objective perfusion measures provides additional guidance when MCS may be warranted and can guide weaning.

**QUESTION 2: WHAT IS THE APPROACH TO MULTIDISCIPLINARY CARE?**

Multidisciplinary care in a dedicated VT unit favorably affects VT recurrence and hospitalization. However, multidisciplinary management of HR-VTA is not well established. Ideally, HR-VTA includes interdisciplinary expertise from interventional cardiology, a heart failure specialist, cardiac anesthesiologist, and occasionally, a cardiac/vascular surgeon, in addition to the treating electrophysiologist (Table 1). MCS implantation and explantation should be performed by an experienced interventional cardiologist who is proficient with large-bore vascular access and adheres to best practices (Table 2). The interventionalist and electrophysiologist coordinate the timing of MCS insertion as well as need for epicardial access. The cardiac anesthesiologist is tasked with maintaining end-organ perfusion even in the presence of recurrent VT and AHD (Figure 3). The heart failure cardiologist assists with optimizing volume status before the procedure, and, at our center, heart failure medications are typically held to reduce periprocedural hypotension. They also assist with weaning MCS, provide postprocedure care, and are instrumental in

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**TABLE 1 Multidisciplinary Periprocedural Care**

| Role                        | Responsibilities                                                                 |
|-----------------------------|----------------------------------------------------------------------------------|
| Electrophysiologist         | - Case identified as “high-risk” VTA requiring hemodynamic support              |
|                             | - Determination of PAINESD score and assessment of current hemodynamics          |
|                             | - Ablation strategy (mapping vs substrate ablation only)                         |
| Heart failure cardiologist  | - Discuss anticipated need for delayed weaning                                   |
|                             | - Establish plan for durable LVAD/heart transplantation if persistent hemodynamic support is required |
| Interventional cardiologist | - Preprocedure risk stratification                                              |
|                             | - Hemodynamic monitoring (ASA monitors, arterial line, PA catheter, cerebral oximetry, and TEE) and evaluation of end-organ perfusion |
|                             | - Vasopressor/inotrope management                                               |
| Cardiac anesthesiologist    | - Postprocedure care                                                             |
|                             | - Assist with delayed weaning of MCS                                            |
|                             | - Wean vasopressors/inotropes                                                   |
| Cardiac intensivist         | - Postprocedure care                                                             |
|                             | - Assist with delayed weaning of MCS                                            |
|                             | - Wean vasopressors/inotropes                                                   |

ASA = American Society of Anesthesiologists; LVAD = left ventricular assist device; MCS = mechanical circulatory support; PA = pulmonary arterial; TEE = transesophageal echocardiography; VTA = ventricular tachycardia ablation.

**TABLE 2 Best Practices for Implant and Explant of MCS**

| Category                          | Practice                                                                 |
|-----------------------------------|--------------------------------------------------------------------------|
| Vascular access                   | Ultrasound and fluoroscopy-guided access                                   |
|                                  | Access with micropuncture kit and confirmation with femoral angiography   |
|                                  | Abdominal aortogram to assess for peripheral artery disease               |
| Intraprocedural monitoring        | Periodic assessment for hematoma or oozeing around the sheath given prolonged nature of VTA and high ACTs (>300 s) |
|                                  | Assessment of distal limb perfusion, recommend ipsilateral or contralateral femoral bypass in case of occlusive large-bore sheath |

**ACT** = activated clotting time; other abbreviations as in Table 1.
assessing candidacy for advanced heart failure therapies. When epicardial access is planned, cardiothoracic surgery is also involved with multidisciplinary care.

During HR-VTA, we monitor CPO and objective measures of end-organ perfusion (Figure 3) as perfusion delineates the efficacy of MCS and defines our weaning criteria (Figure 4). Weaning begins with assessment of CPO, as this is an important marker of end-organ perfusion in patients with acute myocardial infarction and cardiogenic shock. A CPO $>0.6$ W
suggests adequate intrinsic cardiac function. Next, the arterial waveform is evaluated; loss of pulsatility indicates dependence on MCS (Figure 5). Pulsatility should be preserved without significant vasopressor or inotropic support. A SVO₂ >60% and lactate levels <2 mmol/L also suggest that tissue perfusion is adequate. We suggest decreasing device flows and reassessing weaning criteria before withdrawing MCS. Delayed weaning should be considered when baseline LV ejection fraction is <20% and/or CPO is <0.6 W. In our experience, objective assessment of perfusion helps avoid AHD during HR-VTA even without MCS, as these parameters guide vasoactive support and suggest when ventricular arrhythmias should be terminated and/or mapping discontinued.

CONCLUSIONS

We present an approach to HR-VTA with MCS and highlight the importance of multidisciplinary coordination and objective hemodynamic assessment. This potentially paradigm-shifting approach to HR-VTA with MCS should be interrogated systematically to evaluate clinical outcomes.

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