Unilateral antegrade cerebral perfusion using axillary venoarterial extracorporeal membrane oxygenation during central decannulation

Suguru Ohira, MD, PhD, David Spielvogel, MD, Steven L. Lansman, MD, PhD, and Masashi Kai, MD, Valhalla, NY

In postcardiotomy cardiogenic shock (PCS), central cannulation via the ascending aorta can expeditiously establish venoarterial extracorporeal membrane oxygenation (VA-ECMO) but has a high incidence of stroke (17.4%-22.4%), bleeding, and mortality.1-3 We prefer to convert to peripheral VA-ECMO to facilitate chest closure but believe that aortic decannulation poses a high risk for embolic stroke. We report our technique of unilateral antegrade cerebral perfusion (ACP) using VA-ECMO via the right axillary artery (RAX) to minimize the risk of embolic stroke during central decannulation and to confer a smooth transition to peripheral VA-ECMO.

TECHNIQUE

This study was approved by the institutional review board at the New York Medical College, with a waiver of informed consent (institutional review board approval number: #14055 [4/14/2020]). The regional cerebral oxygen saturation is monitored. The RAX is exposed through a small infraclavicular incision,4 and, via the previous sternotomy, the brachiocephalic vessels are dissected out. An epiaortic echo is performed to evaluate the aorta, the aortic cannula, and the brachiocephalic vessels to confirm that there is no large thrombus on the aortic cannula or severe disease at clamping sites of brachiocephalic vessels.

After heparin is given (activated coagulation time >250 seconds), a 10-mm graft is anastomosed to the RAX for arterial outflow. In the case of central VA-ECMO briefly paused, the arterial line of the ECMO circuit is attached to the 10-mm RAX graft. VA-ECMO is resumed with low flow (1.0-1.5 L/min), monitoring a right radial arterial line to maintain mean arterial pressure <60 mm Hg, and the innominate artery is clamped, thus establishing unilateral ACP (Figure 1). The aortic cannula is removed, momentarily allowing the aorta to back-flush before securing a purse string suture. The innominate artery is unclamped and ECMO flow is increased to wash the cannula site in case of potential residual thrombus (Figure 2). Finally, the LCCA and...
LSCA clamps are released and full VA-ECMO flow is established. The clamping time of the LCCA and LSCA is usually less than 90 seconds.

**DISCUSSION**

Central VA-ECMO is an option in patients with profound PCS, as the great vessels can be expeditiously accessed. However, switching to peripheral VA-ECMO is desirable, given the greater incidence of stroke and bleeding complications related to central cannulation.1-3 One possible source of embolic stroke during central VA-ECMO may be thrombus formation around the indwelling aortic cannula, as use of intraoperative protamine sulfate and blood products and interruption of anticoagulation may promote clot formation.1-3 If so, aortic decannulation following central VA-ECMO may pose a high risk for embolic stroke by dislodgement of thrombus that formed around the aortic cannula.

**FIGURE 1.** Unilateral ACP using right axillary VA-ECMO for central decannulation. The left subclavian and left common carotid arteries are clamped, and with central VA-ECMO briefly paused, the ECMO arterial line is attached to a 10-mm right axillary graft. VA-ECMO is resumed with low flow (1.0–1.5 L/min), maintaining mean right radial arterial pressure <60 mm Hg. The innominate artery is clamped, establishing unilateral ACP and isolating the cerebral circulation from the cardiac ejection. ECMO, Extracorporeal membrane oxygenation; CCA, common carotid artery; VA-ECMO, venoarterial extracorporeal membrane oxygenation; ACP, antegrade cerebral perfusion.

**FIGURE 2.** The innominate artery is unclamped while increasing VA-ECMO flow to flush possible residual thrombus from the central cannulation site. The left common carotid and left subclavian arteries are unclamped and full VA-ECMO flow is established. CCA, Common carotid artery; VA-ECMO, venoarterial extracorporeal membrane oxygenation.
The decannulation strategy described herein could minimize the risk of embolic stroke in 2 ways: first, clamping the brachiocephalic vessels at the time of decannulation prevents embolic material from entering the cerebral circulation, and second, flushing the decannulation site with antegrade ECMO flow, with the left carotid and left subclavian arteries clamped.

**Hypothesis**
High risk of embolism during central decannulation in PCS on VA-ECMO

**Technique**
1. Switching Central VA-ECMO to right axillary artery cannulation
2. Left carotid, and left subclavian artery are clamped
3. VA-ECMO flow 1.0-1.5 L/min from right axillary artery, and innominate artery is clamped (Unilateral ACP)
4. Central aortic cannula is removed
5. Clamp on the innominate artery is released to flush the decannulation site (VA-ECMO flow: 3-4 L/min)
6. All clamps are released

**FIGURE 3.** ACP for stroke prevention during central ECMO decannulation. The left subclavian and left common carotid arteries are clamped, and with central VA-ECMO briefly paused, the ECMO arterial line is attached to a 10-mm right axillary graft. VA-ECMO is resumed with low flow (1.0-1.5 L/min), maintaining mean right radial arterial pressure <60 mm Hg. With the innominate artery clamped, establishing unilateral ACP and isolating the cerebral circulation from the cardiac ejection, the aortic cannula is removed, briefly allowing the aorta to back-flush before securing a purse string suture. The innominate artery is unclamped while increasing VA-ECMO flow to flush possible residual thrombus from the central cannulation site. The left common carotid and left subclavian arteries are unclamped and full VA-ECMO flow is established. VA-ECMO, Venoarterial extracorporeal membrane oxygenation; PCS, postcardiotomy cardiogenic shock; ECMO, extracorporeal membrane oxygenation; CCA, common carotid artery; ACP, Antegrade cerebral perfusion.
arteries clamped, may remove any remaining intra-aortic thrombus. While these maneuvers are performed, the cerebral circulation is maintained by unilateral ACP via the RAX (Figure 3), which is an ideal cannulation site for this technique compared to the femoral artery. During unilateral ACP, we have not seen a significant drop of the regional cerebral oxygen saturation. Although exposing the LSCA is challenging in some cases, especially with a large habitus patient, we found it less difficult on VA-ECMO support, possibly because the heart is decompressed. We prefer to isolate all the brachiocephalic vessels, but this technique could be simplified by just clamping the LCCA, if exposure of the LSCA is difficult or if there is a concern for clamping the brachiocephalic vessels due to atherosclerosis. In a patient with significant atherosclerotic disease burden, manipulations of neck vessels are not recommended (Video 1).

Our representative patient was 225 lbs and transferred to our hospital after placement of central VA-ECMO after coronary artery bypass grafting. We successfully switched to RAX VA-ECMO. He was directly transplanted off VA-ECMO several days later without any complications. We should note the limitation that the proposed technique has only a theoretical benefit of reducing stroke, as a large number of data are not available (3 patients).

In conclusion, we describe a technique of transitioning from central to peripheral VA-ECMO that uses the ECMO circuit to provide unilateral ACP via the RAX to isolate the cerebral circulation and might prevent thromboembolic stroke in patients with PCS.

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