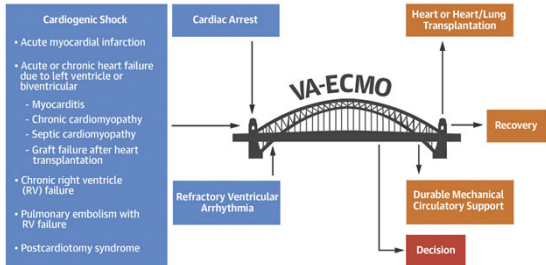
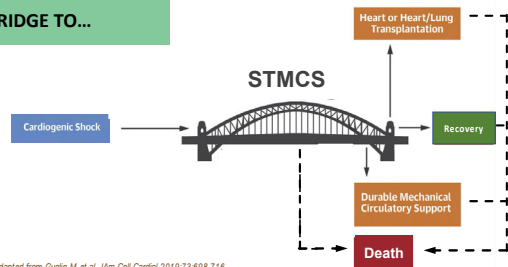


BRIDGE TO...



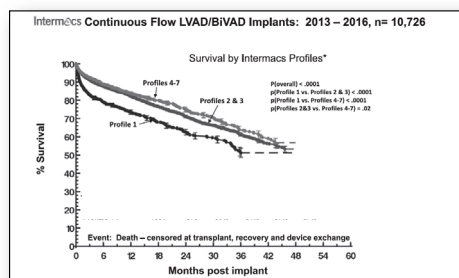
Outcomes of Short Term Mechanical Circulatory Support (STMCS)

BRIDGE TO...



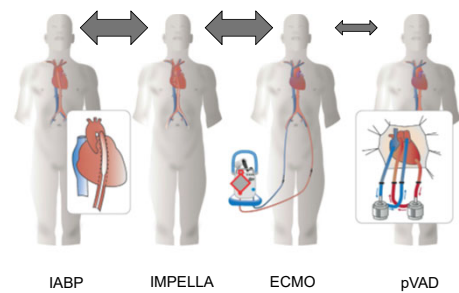
Adapted from Gugin M et al. J Am Coll Cardiol 2019;73:698-716

Short-term Mechanical Circulatory Support (STMCS) prior to Left Ventricular Assist Device (LVAD)

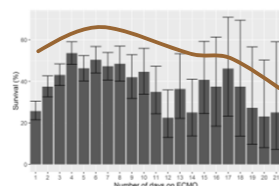


D-6

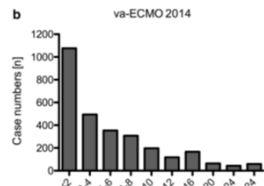
SHORT TERM MECHANICAL CIRCULATORY SUPPORT (STMCS)



ECLS DURATION AND OUTCOMES



Smith M et al. Crit Care 2017;21:45



Karagiannis C et al. Intensive Care Med 2016;42:889-96

SOLUTION

PROBLEM

	IABP (7.5-9 Fr)	IMPELLA 2.5 and CP	IMPELLA 5.0	ECLS
Size of MCS	7.5 to 9 Fr	Motor 12/14 Fr/Catheter 9 Fr	Motor 21 Fr/Catheter 9 Fr	A: 15-19 Fr/N: 23-29 Fr
Very frequent (> 10%)	Thrombocytopenia	Severe access vascular bleeding**	Severe access vascular bleeding**	Severe access vascular bleeding**
Frequent (5-10%)		- Intravascular hemolysis - Site infection	- Limb ischemia* - Site infection	Limb ischemia*
Non exceptional (1-5%)	- Device malfunction - Severe limb ischemia* - Severe access vascular bleeding**	- Limb ischemia* - Device malfunction - Pump displacement	- Intravascular hemolysis - Device malfunction - Pump displacement	- Intravascular hemolysis - Pulmonary hemorrhage
Exceptional (<1%)	- Retroperitoneal bleeding - Intravascular hemolysis - Aortic complication - Cerebral embolism - Pancreatitis - Site infection - Mesenteric ischemia - Balloon leak	- Retroperitoneal bleeding - Functional mitral stenosis - Mitral regurgitation (chordal rupture) - Left ventricular wall perforation - Intra ventricular thrombosis	- Functional mitral stenosis - Mitral regurgitation (chordal rupture) - Aortic regurgitation - Left ventricular wall perforation - Intra ventricular thrombosis	- Aortic complication - Device malfunction

Bonello L et al. Arch Cardiovasc Dis 2020 ;113:448-60

ECLS for cardiac indications

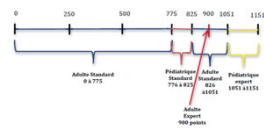
Diagnosis	No. Runs, N	Average ECLS Duration (hour)	Survival, N (%)
Adult (>16 years)			
Shock*	2,083	144	882 (42)
Cardiomyopathy	704	162	358 (51)
Myocarditis	227	188	143 (65)
Congenital defect	420	129	156 (37)

~ 6 to 8 days ~ 50%

Thiagarajan RR et al. ASAIO Journal 2017;63:60-7

New graft allocation rules for heart transplantation in France (from January 2108)

Candidate risk score



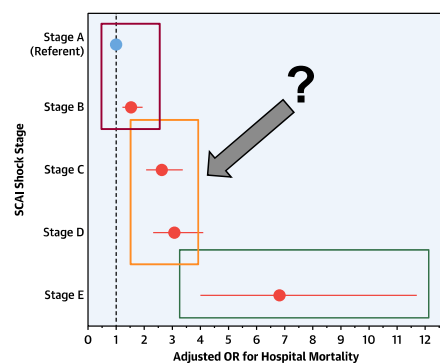
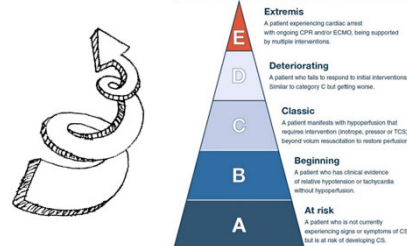
Since April 2019, the score decreases from 100% to 0% in candidates on VA-ECMO from 12 to 16 days of support duration.

Baran DA et al. Catheter Cardiovasc Interv

WHEN ? "NEITHER TOO EARLY NOR TOO LATE"

SCAI: Society for Cardiovascular Angiography and Interventions (SCAI)

The SCAI pyramid of cardiogenic shock classification¹

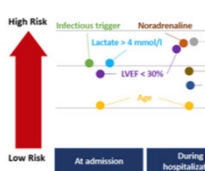


Jentzer JC et al. Am Coll Cardiol 2019; 74:2117-28

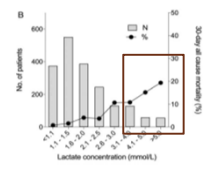
The FRENDSOCK registry

A Real Life Picture of Cardiogenic Shock in France

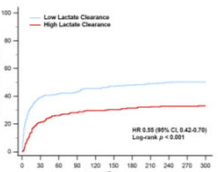
48 centers 772 patients



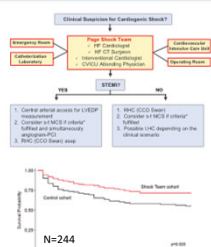
Delmas C et al. ESC Heart Failure 2022; 9:408-19



Frydland M et al. Shock 2019;51:321-7



Park IH et al. J Intensive Care 2021;9:63

[illegible]

Multidisciplinary cardiogenic shock team approach improves the long-term outcomes of patients suffering from refractory cardiogenic shock treated with short-term mechanical circulatory support

```

graph TD
    A[618 consecutive patients with valvular cardiogenic shock treated by ECMO  
between 2007 and 2010 were screened] --> B[269 patients were included in the final analysis]
    A --> C[149 patients excluded  
107 post-operative cardiogenic shock  
42 non-ECMO treated  
22 acute  
23 with no data missing]
    B --> D[Control group  
January 2007 - January 2013 (n=143)]
    B --> E[Shock state group  
April 2013 - December 2013 (n=126)]
    D --> D1[8 ECMO-treated cases]
    D --> D2[8 non-ECMO-treated cases]
    D1 --> D1a[8 ECMO-treated cases  
1 month  
4 ECMO-treated cases  
4 non-ECMO-treated cases]
    D1 --> D1b[8 ECMO-treated cases  
2 months  
4 ECMO-treated cases  
4 non-ECMO-treated cases]
    D1 --> D1c[8 ECMO-treated cases  
3 months  
4 ECMO-treated cases  
4 non-ECMO-treated cases]
    D1 --> D1d[8 ECMO-treated cases  
4 months  
4 ECMO-treated cases  
4 non-ECMO-treated cases]
    D1 --> D1e[8 ECMO-treated cases  
5 months  
4 ECMO-treated cases  
4 non-ECMO-treated cases]
    D1 --> D1f[8 ECMO-treated cases  
6 months  
4 ECMO-treated cases  
4 non-ECMO-treated cases]
    D2 --> D2a[8 non-ECMO-treated cases  
1 month  
4 non-ECMO-treated cases  
4 ECMO-treated cases]
    D2 --> D2b[8 non-ECMO-treated cases  
2 months  
4 non-ECMO-treated cases  
4 ECMO-treated cases]
    D2 --> D2c[8 non-ECMO-treated cases  
3 months  
4 non-ECMO-treated cases  
4 ECMO-treated cases]
    D2 --> D2d[8 non-ECMO-treated cases  
4 months  
4 non-ECMO-treated cases  
4 ECMO-treated cases]
    D2 --> D2e[8 non-ECMO-treated cases  
5 months  
4 non-ECMO-treated cases  
4 ECMO-treated cases]
    D2 --> D2f[8 non-ECMO-treated cases  
6 months  
4 non-ECMO-treated cases  
4 ECMO-treated cases]
    E --> E1[8 ECMO-treated cases]
    E --> E2[8 non-ECMO-treated cases]
    E1 --> E1a[8 ECMO-treated cases  
1 month  
4 ECMO-treated cases  
4 non-ECMO-treated cases]
    E1 --> E1b[8 ECMO-treated cases  
2 months  
4 ECMO-treated cases  
4 non-ECMO-treated cases]
    E1 --> E1c[8 ECMO-treated cases  
3 months  
4 ECMO-treated cases  
4 non-ECMO-treated cases]
    E1 --> E1d[8 ECMO-treated cases  
4 months  
4 ECMO-treated cases  
4 non-ECMO-treated cases]
    E1 --> E1e[8 ECMO-treated cases  
5 months  
4 ECMO-treated cases  
4 non-ECMO-treated cases]
    E1 --> E1f[8 ECMO-treated cases  
6 months  
4 ECMO-treated cases  
4 non-ECMO-treated cases]
    E2 --> E2a[8 non-ECMO-treated cases  
1 month  
4 non-ECMO-treated cases  
4 ECMO-treated cases]
    E2 --> E2b[8 non-ECMO-treated cases  
2 months  
4 non-ECMO-treated cases  
4 ECMO-treated cases]
    E2 --> E2c[8 non-ECMO-treated cases  
3 months  
4 non-ECMO-treated cases  
4 ECMO-treated cases]
    E2 --> E2d[8 non-ECMO-treated cases  
4 months  
4 non-ECMO-treated cases  
4 ECMO-treated cases]
    E2 --> E2e[8 non-ECMO-treated cases  
5 months  
4 non-ECMO-treated cases  
4 ECMO-treated cases]
    E2 --> E2f[8 non-ECMO-treated cases  
6 months  
4 non-ECMO-treated cases  
4 ECMO-treated cases]
  
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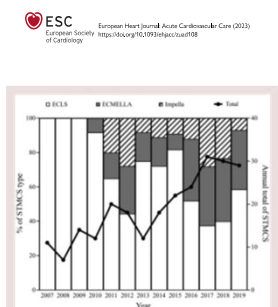
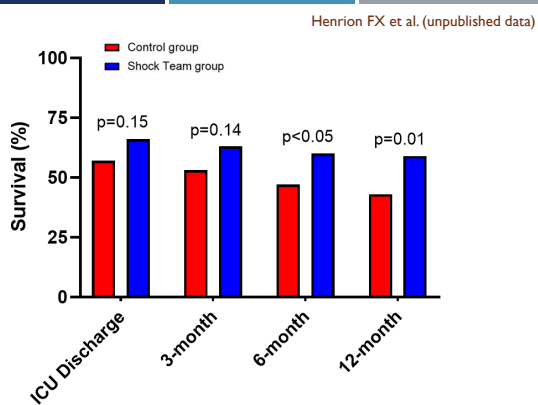
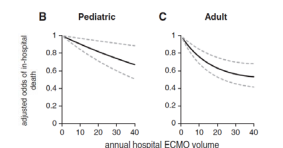


Figure 3 Trends in STMCs over the years. ECLS, extracorporeal life support; ECMELLA, Association of ECLS and Impella® 2.5, CP or 5.0; IABP, intra-aortic balloon pump; STMCs, short-term mechanical circulatory support. The "Impella®" category includes only CP and 5.0 used alone.

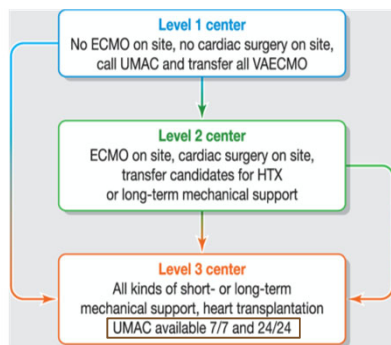


Analysis of the Extracorporeal Life Support Organization Registry

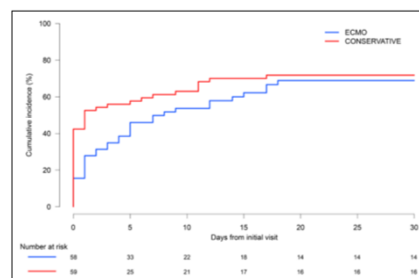
Year	Neonatal	Pediatric	Adult
1989	1500	200	100
1991	1600	300	100
1993	1700	400	100
1995	1600	500	100
1997	1500	600	100
1999	1400	700	100
2001	1300	800	100
2003	1200	900	100
2005	1300	1000	100
2007	1400	1100	100
2009	1300	1200	100
2011	1200	1200	100
2013	1100	1100	2500



Period	Annual Hospital ECMO Volume	Adjusted Mortality Odds Ratio (95% CI)		
		Neonate	Pediatric	Adult
1989-2013	1-5	Referent	Referent	Referent
	6-14	0.80 (0.75-0.86)	0.99 (0.96-1.13)	0.61 (0.46-0.995)
	15-30	0.74 (0.63-0.88)	0.86 (0.75-1.01)	0.75 (0.59-0.94)
	>30	0.69 (0.56-0.84)	0.99 (0.89-1.14)	0.61 (0.48-0.79)
2008-2013	1-5	Referent	Referent	Referent
	6-14	1.01 (0.79-1.28)	1.03 (0.84-1.26)	0.82 (0.64-1.05)
	15-30	0.94 (0.70-1.25)	0.92 (0.73-1.16)	0.72 (0.55-0.96)
	>30	0.68 (0.47-1.00)	0.85 (0.67-1.08)	0.61 (0.46-0.80)



ORIGINAL RESEARCH ARTICLE

Ostadal P et al. *Circulation* 2023; 147:454–64

ORIGINAL ARTICLE			
Extracorporeal Life Support in Infarct-Related Cardiogenic Shock			
H. Thiele, U. Zeymer, J. Akin, M. Behnes, T. Rassaf, A.A. Mahabadi, R. Lehmann, T. Endl, T. Grot, T. Sessler, A. Schuster, C. Skurk, D. Dierschke, P. Clemmensen, M. Hennerdorf, S. Fichtlscherer, I. Voigt, M. Seyfarth, S. John, S. Ewen, A. Linke, E. Tigges, P. Nordbeck, L. Bruch, C. Jung, J. Franz, P. Lauten, T. Grottel, H.-J. Teisberg, J. Pils, E. Kirschhof, T. Quatrecas, S. Schneider, S. Desch, and A. Freund, for the ECLS-SHOCK Investigators*			
Design	Characteristic	ECLS (N=209)	Control (N=208)
• Prospective, randomized multicentre study (Slovenia and Germany)	SCAI shock stage — no. (%)		
	C	104 (49.8)	111 (53.4)
	D	18 (8.6)	18 (8.7)
	E	87 (41.6)	79 (37.9)
• AMI complicated by CS (SAP<90 mmHg and lactate>3) early revascularized (PCI or CABG)	Catheterization success — no./total no. (%)		
	Percutaneous	156/208 (75.0)	148/207 (71.5)
	Coronary artery bypass grafting	52/208 (25.0)	59/207 (28.5)
• SCAI C to E requiring early use of ECLS after coronary angiography	PCI	199/208 (95.7)	189/204 (92.5)
• Crossover to ECLS was possible	CABG	1/208 (0.5)	0/204
• Primary outcome: all cause death at D30	PCI with transfer to CABG	2/208 (1.0)	0/204
	No revascularization	6/208 (2.9)	5/204 (2.5)
Results	ECLS therapy — no. (%)		
• Inclusion period 2019 to 2022 (44 centers) n=417	Initiation in catheterization laboratory	42/202 (21.3)	4/26 (15.4)
	Before revascularization	50/192 (26.1)	8/26 (30.8)
	After revascularization	100/192 (52.1)	7/26 (26.9)
• Cross over from control to ECLS group for n=26	Median duration of ECLS therapy (IQR) — days	2.7 (1.5–4.3)	2.7 (1.3–4.8)
	Revascularization before randomization — no. (%)	162 (77.5)	162 (77.9)

Thiele H et al. New Engl J Med 2023; 389:1286-97

Characteristic	ECLS (N=209)	Control (N=208)
Target temperature management — no./total no. (%)	82/209 (39.2)	109/208 (52.4)
Invasive mechanical ventilation		
Patients — no./total no. (%)	183/203 (90.1)	177/202 (87.6)
Median duration (IQR) — days	7.0 (4.0–12.0)	5.0 (3.0–9.0)
Catecholamine requirement — no./total no. (%)	203/209 (97.1)	195/208 (93.8)
Norepinephrine	181/203 (89.2)	181/195 (92.8)
Epinephrine	63/203 (31.0)	69/195 (35.4)
Dobutamine	88/203 (43.3)	59/195 (30.3)
Dopamine	1/203 (0.5)	0/195
Sepsis within 30 days after randomization — no. (%)	21 (10.0)	21 (10.1)
Intra-aortic balloon pump	—	1/28 (3.6)
Impella 2.5	—	1/28 (3.6)
Impella CP	—	24/28 (85.7)
Impella 5.0	—	1/28 (3.6)
Impella 5.5	—	1/28 (3.6)
Permanent left ventricular assist device — no./total no. (%)	1 (0.5)	1 (0.5)

Thiele H et al. New Engl J Med 2023; 389:1286-97

	ECLS (n=209)	Control (n=208)
All-cause mortality at 30 days; n/total (%)	100/209 (47.8)	102/208 (49.0)
Causes of death at 30 days		
Refractory cardiogenic shock; n/total (%)	51/100 (51.0)	56/102 (54.9)
Sudden cardiac death; n/total (%)	7/100 (7.0)	5/102 (4.9)
Recurrent myocardial infarction; n/total (%)	2/100 (2.0)	2/102 (2.0)
Mechanical complication of infarction; n/total (%)	1/100 (1.0)	1/102 (1.0)
Bleeding; n/total (%)	4/100 (4.0)	0/102
Brain injury; n/total (%)	26/100 (26.0)	27/102 (26.5)
Sepsis; n/total (%)	4/100 (4.0)	10/102 (9.8)
Unknown cause; n/total (%)	0/100	1/102 (1.0)
Other cause; n/total (%)	5/100 (5.0)	0/102

Thiele H et al. New Engl J Med 2023; 389:1286-97

	ECLS (n=209)	Control (n=208)
Active left ventricular unloading during ECLS therapy;	11/190 (5.8)	6/19 (31.6)
Type of unloading; n/total (%)		
Additional insertion of IABP	2/11 (18.2)	2/6 (33.3)
Additional insertion of percutaneous left ventricular assist device (Impella®)	9/11 (81.8)	4/6 (66.7)
Atrial septostomy with drainage of the left atrium by a pigtail catheter connected to the venous cannula of the ECLS	0/11	0/6
Pulmonary artery drainage with connection to the venous cannula of the ECLS	0/11	0/6
Transcatheter venting by pigtail catheter insertion into the left ventricle and connection to the venous cannula of the ECLS	0/11	0/6

Thiele H et al. New Engl J Med 2023; 389:1286-97

2021 ESC Guidelines for the diagnosis and treatment of acute and chronic heart failure		
Recommendations	Class ^a	Level ^b
Short-term MCS should be considered in patients with cardiogenic shock as a BTR, BTD, BTB. Further indications include treatment of the cause of cardiogenic shock or long-term MCS or transplantation.	IIa	C
IABP may be considered in patients with cardiogenic shock as a BTR, BTD, BTB, including treatment of the cause of cardiogenic shock (i.e. mechanical complication of acute MI) or long-term MCS or transplantation. ⁴⁵⁰	IIb	C
IABP is not routinely recommended in post-MI cardiogenic shock. ^{500–502}	III	B

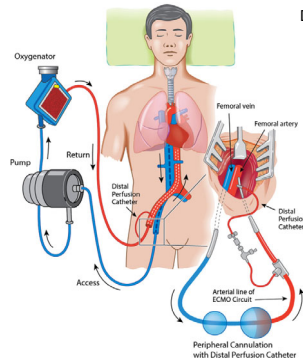
McDonagh TA et al. Eur Heart J 2021; 42:3599-3726

Extracorporeal Membrane Oxygenation in Cardiopulmonary Disease in Adults

Abrams D et al. J Am Coll Cardiol 2014; 63:2769-78

- Rescue therapy (no first line)
- Refractory cardiogenic shock
- Respiratory and/or cardiac supply
- Ensure "optimal" peripheral organs perfusion
- Cardiac function recovery (+++)
- Left ventricular loading conditions
- Coronary perfusion and myocardial working
- No influence on myocardial recovery (+++)

Dennis M JAHA 2020



Several advantages...

- Respiratory and/or cardiac supply
- Biventricular supply (including right ventricular dysfunction)
- Restore peripheral end-organ perfusion by providing high flow (up to 7 L.min⁻¹)
- Rapid insertion (cardiac arrest...)
- Peripheral cannulation (femoral access)
- Cheap and easily reliable
- Biventricular supply
- Mobile

INDICATIONS

■ Post-cardiotomy (conventional or heart transplantation)

- Immediately when the CPB weaning appears to be impossible
- Secondary in presence of refractory Low cardiac output syndromue

Bokhtari F et al. J Thorac Cardiovasc Surg 2008;135:382-8
Rastan AJ et al. J Thorac Cardiovasc Sug 2010; 139:302-11

■ Medical indications

- Myocardial infarction Infarctus du myocarde
- Dilated cardiomyopathy
- Acute Myocarditis fulminante
- CIntoxication médicamenteuse cardio-toxiques
- Pulmonary Cœur pulmonaire aigue (embolie pulmonaire, embolie amniotique)
- Accidental hypothermia (noyade)

Marasco SF et al Heart, Lung and Circulation 2008; 17:541-7
Baud F et al Crit Care 2007; 11:207

Outcomes and long-term quality-of-life of patients supported by extracorporeal membrane oxygenation for refractory cardiogenic shock*

Crit Care Med 2008; 36:1404-1411

Alain Combes, MD, PhD; Pascal Leprince, MD, PhD; Charles-Edouard Luyt, MD, PhD; Nicolas Bonnet, MD; Jean-Louis Trouillet, MD; Philippe Léger, MD; Alain Pavie, MD; Jean Chastre, MD

Table 4. Multivariable logistic-regression analysis: early independent predictors of intensive care unit death

Factor	OR (95% CI)	p
Female sex	3.89 (1.06-14.22)	.04
Myocarditis	0.13 (0.02-0.78)	.03
ECMO under CPR	20.68 (1.09-392.03)	.04
Prothrombin activity <50%	3.93 (1.11-13.85)	.03
24-hr urine output <500 mL	6.52 (1.87-22.74)	.003

OR, odds ratio; CI, confidence interval; CPR, cardiopulmonary resuscitation; ECMO, extracorporeal membrane oxygenation.

Extracorporeal Life Support Organization Registry Report 2012

MATTHEW L. PADEN,* STEVEN A. CONRAD,† PETER T. RYCLUS,‡ and RAVI R. THAGARAJAN§, ON BEHALF OF THE ELSO REGISTRY

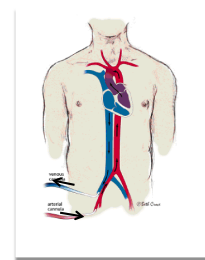
ASAIO Journal 2013;59:202-210

Table 7. Mechanical and Patient-related Complications for Cardiac ECLS

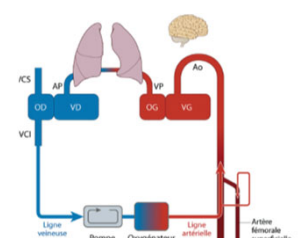
	0-30 Days	31 Days and <1 Year	1 Year and <16 Years	>16 Years
Mechanical				
Extracorporeal circuit	7.4 (24)	8.1 (29)	9 (43)	15.1 (36)
Tubing rupture	0.3 (25)	0.6 (50)	0.7 (47)	0.2 (9)
Pump malfunction	1.6 (29)	2.1 (32)	2.1 (50)	0.7 (28)
Prone to embolism	6.1 (33)	5.6 (38)	6.3 (42)	4.4 (27)
Patient-related				
ICU	11.3 (23)	5.7 (29)	3.8 (21)	1.7 (7)
Cannula site bleeding	10.4 (30)	11.9 (40)	17.6 (52)	20.9 (39)
Surgical site bleeding	31.7 (30)	33 (39)	28.9 (48)	25.5 (34)
Cardiac tamponade	6.1 (27)	5.1 (38)	5.1 (50)	5.7 (27)
Clinical seizures	7.3 (29)	9 (28)	4.5 (21)	2.1 (15)

Table entries are reported in percentage (% survival).
ECLS, extracorporeal life support; ICU, intracranial hemorrhage.

Peripheral veno-arterial ECLS

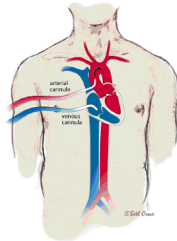


Retrograde aortic flow in total competition with native stream



Calderon J et al. Traité Anesthésie-Réanimation O. Fourcade (4^{ème} Edition)

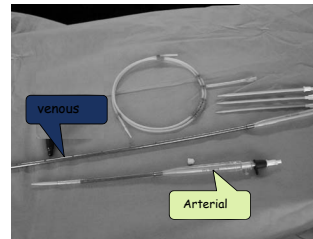
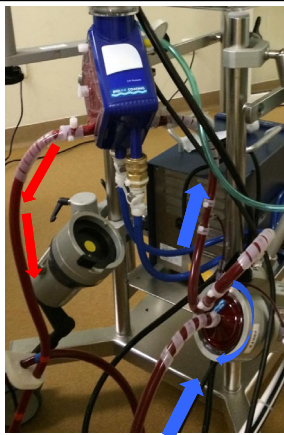
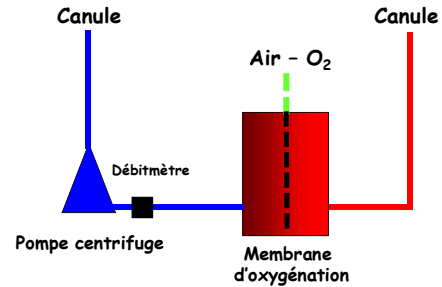
Central veno-arterial ECLS



Central ECLS
(anterograde flow)

Marasco SF et al. Heart lung and Circulation 2008

PRINCIPES DE L'ECLS



Percutaneous at the bedside (echography guided +++) or surgical way
Preferentially removed in operating room +++++

Informations de commande des canules HLS artérielles

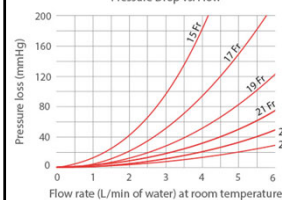
Type	Diamètre extérieur	Longueur d'insertion	Orifices latéraux	Longueur de perforation	Connection	Revêtement Bioline
PAS 1315	13 Fr (4,3 mm)	15 cm	2	1 cm	3/8" LL	BE-PAS 1315
PAS 1515	15 Fr (5,0 mm)	15 cm	2	1 cm	3/8" LL	BE-PAS 1515
PAS 1715	17 Fr (5,7 mm)	15 cm	2	1 cm	3/8" LL	BE-PAS 1715
PAS 1915	19 Fr (6,3 mm)	15 cm	2	1 cm	3/8" LL	BE-PAS 1915
PAS 2115	21 Fr (7,0 mm)	15 cm	2	1 cm	3/8" LL	BE-PAS 2115
PAS 2315	23 Fr (7,7 mm)	15 cm	2	1 cm	3/8" LL	BE-PAS 2315
PAL 1523	15 Fr (5,0 mm)	23 cm	2	1 cm	3/8" LL	BE-PAL 1523
PAL 1723	17 Fr (5,7 mm)	23 cm	2	1 cm	3/8" LL	BE-PAL 1723
PAL 1923	19 Fr (6,3 mm)	23 cm	2	1 cm	3/8" LL	BE-PAL 1923
PAL 2123	21 Fr (7,0 mm)	23 cm	2	1 cm	3/8" LL	BE-PAL 2123
PAL 2323	23 Fr (7,7 mm)	23 cm	2	1 cm	3/8" LL	BE-PAL 2323



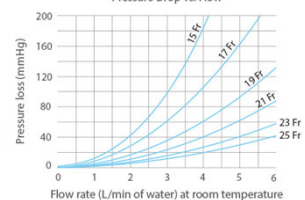
Informations de commande des canules HLS veineuses

Type	Diamètre extérieur	Longueur d'insertion	Orifices latéraux	Longueur de perforation	Connection	Revêtement Bioline
PVS 1938	19 Fr (6,3 mm)	38 cm	12	10 cm	3/8"	BE-PVS 1938
PVS 2138	21 Fr (7,0 mm)	38 cm	12	10 cm	3/8"	BE-PVS 2138
PVS 2338	23 Fr (7,7 mm)	38 cm	16	10 cm	3/8"	BE-PVS 2338
PVS 2538	25 Fr (8,3 mm)	38 cm	20	10 cm	3/8"	BE-PVS 2538
PVL 2155	21 Fr (7,0 mm)	55 cm	20	20 cm	3/8"	BE-PVL 2155
PVL 2355	23 Fr (7,7 mm)	55 cm	20	20 cm	3/8"	BE-PVL 2355
PVL 2555	25 Fr (8,3 mm)	55 cm	24	20 cm	3/8"	BE-PVL 2555
PVL 2955	29 Fr (9,7 mm)	55 cm	32	20 cm	3/8"	BE-PVL 2955

Arterial Pressure Drop vs. Flow



Venous Pressure Drop vs. Flow



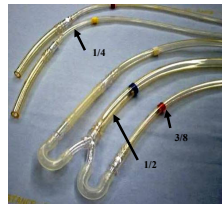
Lignes de connexion

Connexion des divers éléments de l'ECMO

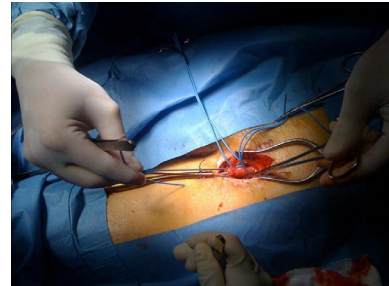
Unité de mesure : inch ou pouce (25,4 mm)

- 1/2 (≈ 12 mm) (CEC)
- **3/8 (≈ 10 mm) (ECMO +++)**
- 1/4 ($\approx 6,4$ mm) Bonne hémocompatibilité

Bonne souplesse



Surgical access



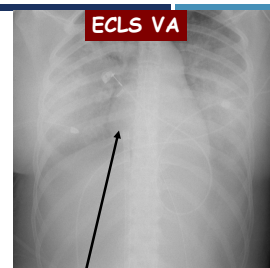
Percutaneous versus surgical femoro-femoral veno-arterial ECMO: a propensity score matched study

Table 2 VA-ECMO-related outcomes in the propensity matched population

	Surgical group n=266 (%)	Percutaneous group n=266 (%)	p-value
30-day mortality	150 (56.3)	170 (63.8)	0.034
Cannulation site infection	74 (27.8)	44 (16.5)	<0.001
Infection requiring surgical revision ^a	40 (15.0)	14 (5.3)	<0.001
Vascular complications at cannulation ^b	7 (2.6)	10 (3.8)	0.663
Limb ischemia	33 (12.4)	23 (8.6)	0.347
Cannula relocation or removal	25 (9.4)	15 (5.6)	0.250
Limb fasciotomy	10 (3.8)	6 (2.3)	0.310
Amputation	2 (0.8)	2 (0.8)	1.000
Vascular complications after cannula removal	9 (3.4)	39 (14.7)	<0.001
Surgical revision for persistent bleeding early after decannulation	4 (1.5)	25 (9.4)	<0.001
Surgical revision in the days after decannulation ^c	5 (1.9)	14 (5.3)	0.035
Lower limb sensory-motor deficit	6 (2.3)	7 (2.6)	0.779

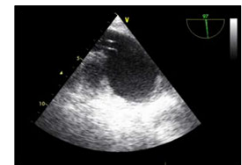
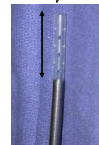
Daniel P et al. Intensive Care Med 2018; 44:2153-61

ECLS VA



Veine cave inférieure (drainage)

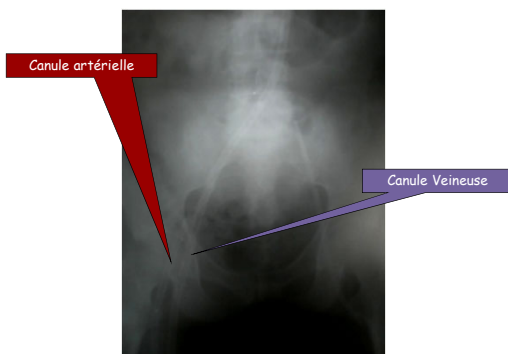
Extrémité non radio-opaque (2-3 cm)



ME bicaval

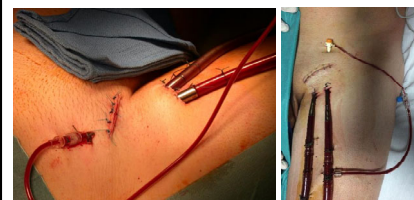
Mise en place sous ETO ou contrôle rapide!

Peripheral veno-arterial ECLS



Canule artérielle

Canule Veineuse



Solidement fixée
Contrôlée à l'angio
Fémorale superficielle

Reperfusion line
(systematically inserted)

Peripheral VA ECLS

This reperfusion line is alimented by arterial canula (risk of ischemia when ECLS flow is decreased)
Usefulness of NIRS (STO2) for detecting early lower limb ischemia

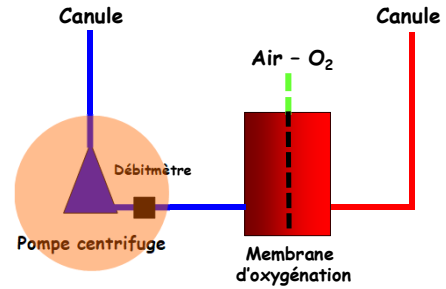
Schachner T et al. Eur J Cardiothorac Surg 2008; 34:1253-4

DÉTECTION ISCHÉMIE MI = CLINIQUE ET NIRS +++

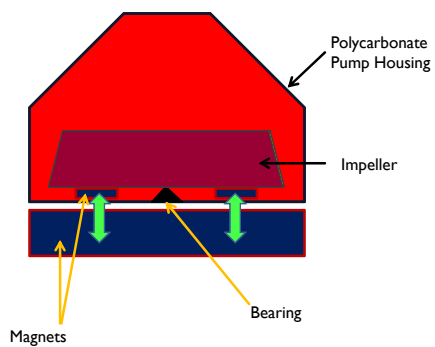


Saturométrie Transcutanée

ROTATIVE CENTRIFUGAL PUMP



Magnetically Coupled Pumps



CENTRIFUGAL ROTATIVE PUMP

- Non-occlusive
- Flow depends:
 - Gradient pressure generated by impeller (speed of rotation)
 - Size of canula, length and diameter of tubing
 - Volemia of patient (pre-load of the pump)
 - Systemic vascular resistances (after-load of the pump)
- Preload and after-load dependence +++
- Measurement of CO is mandatory (electromagnetic or ultrasonic method)
- Backflow phenomenon possible when speed of pump < 1500 TRM **Please clamp the tubing +++**



Rotaflow, MAQUET 32 ml



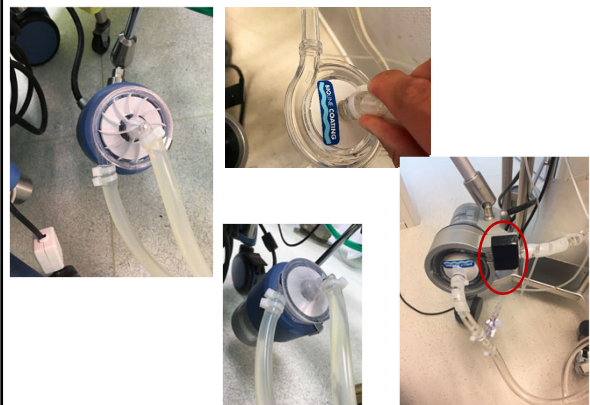
Biomedicus, Medtronic®
(cones superposés)

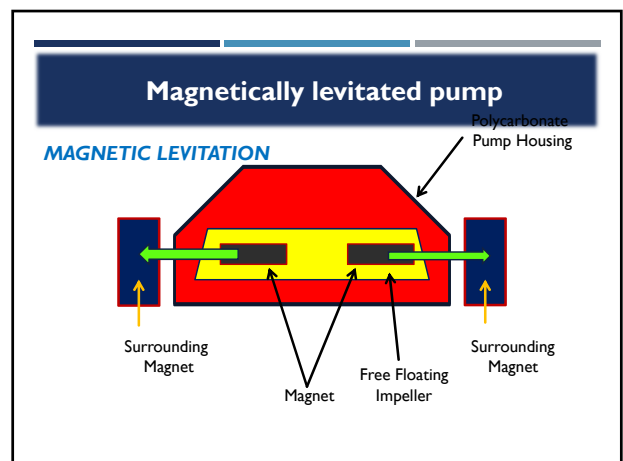
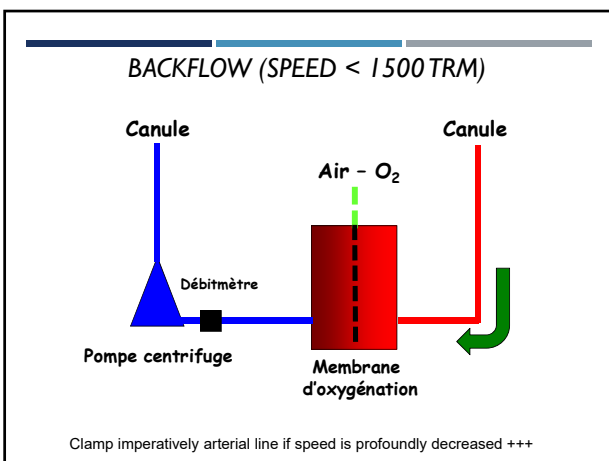
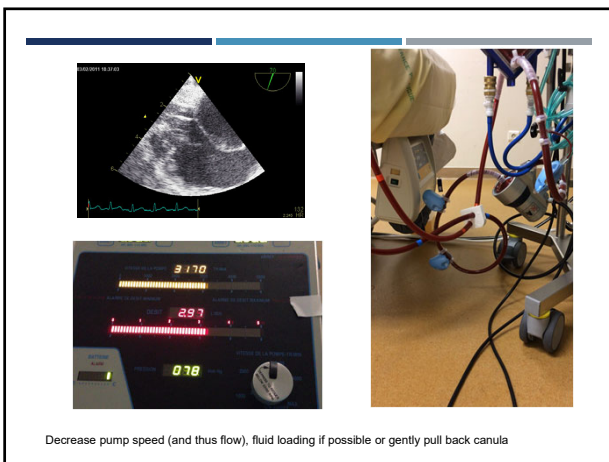
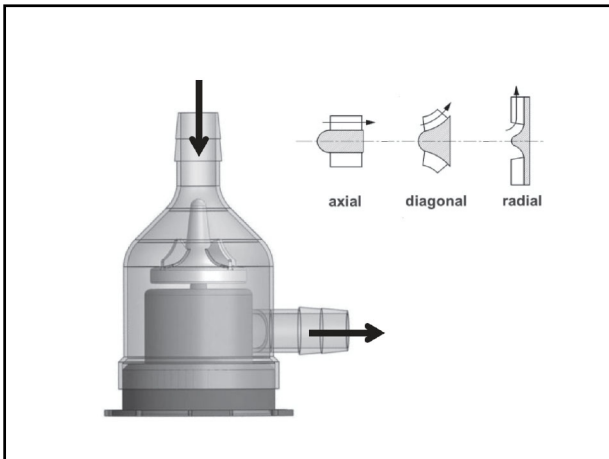


Deltastream pump, MEDOS (16 ml)

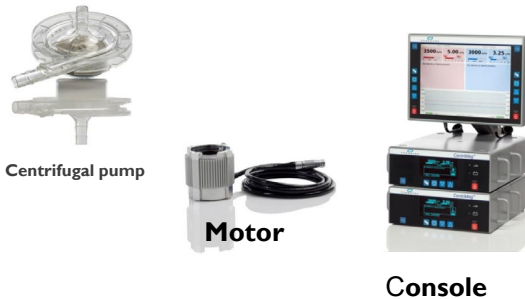


Revolution, Sorin®
(avec aubes vitesse moindre)





SYSTEM COMPONENTS – 2ND GEN

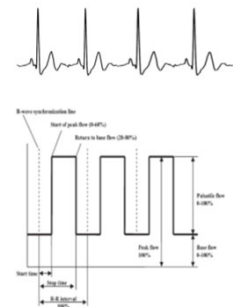


BENEFITS OF PULSATILE ECLS

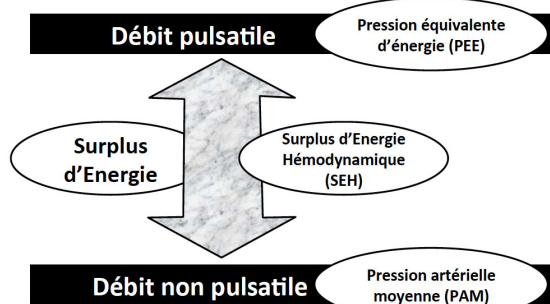
- Add diastolic pulsatility
- Improve coronary blood flow (diastolic delivery)
- ECG-synchronized pulse flow (pulse delivery in diastole)
 - Avoid conflict between pulse-wave from intrinsic activity of failing heart and ECLS-generated pulse-waves
 - Modulation of ECLS by the patient's own autonomic nervous system

R-wave ECG synchronisation

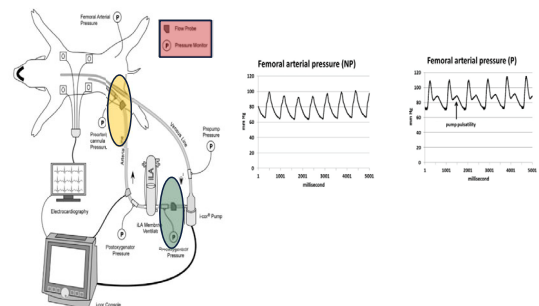
- Width of pulsatility 200 msec
- Trigger pulsatile mode (assist frequency 1:1 to 1:2)
- Differential speed value 2000-4000 trm
- Pulsatile must represent at least 20% of cardiac cycle (R-R interval)

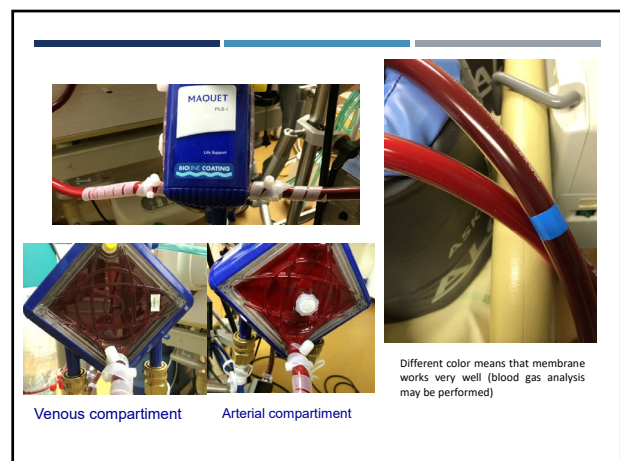
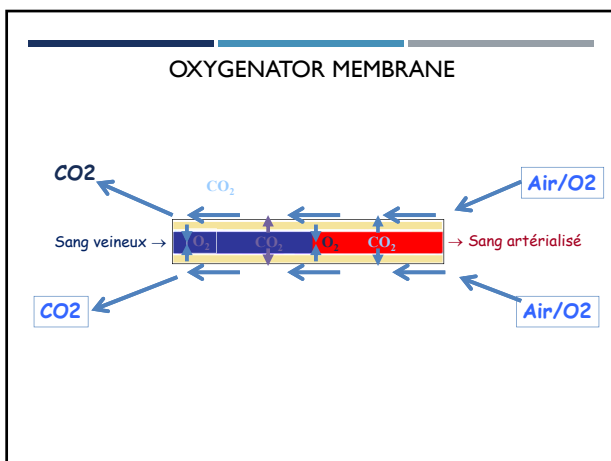
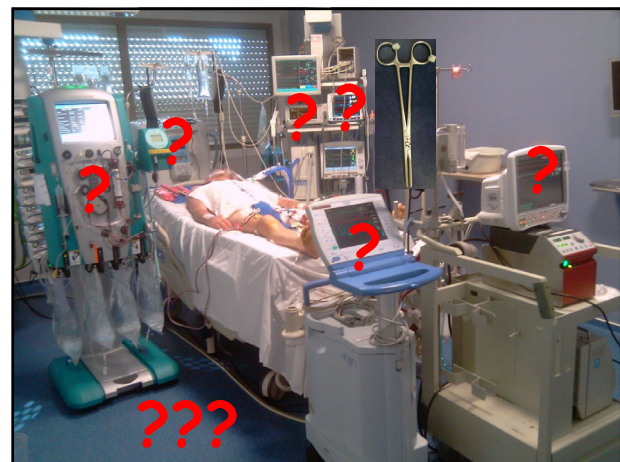
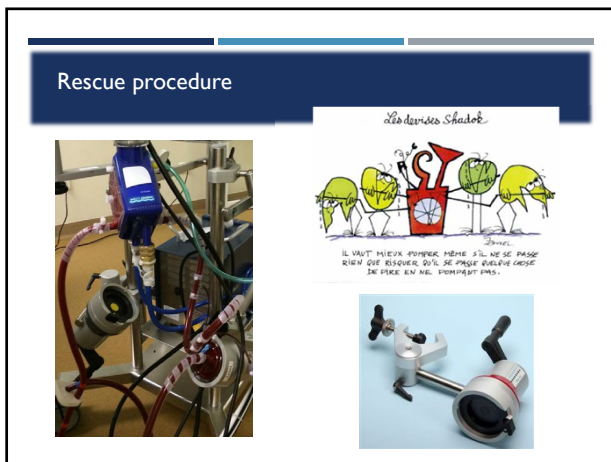
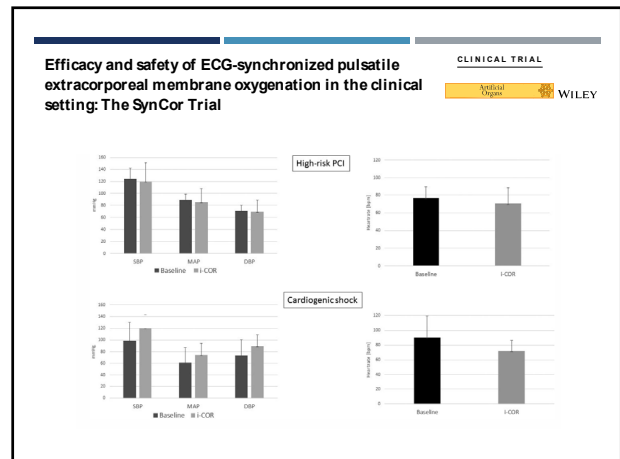
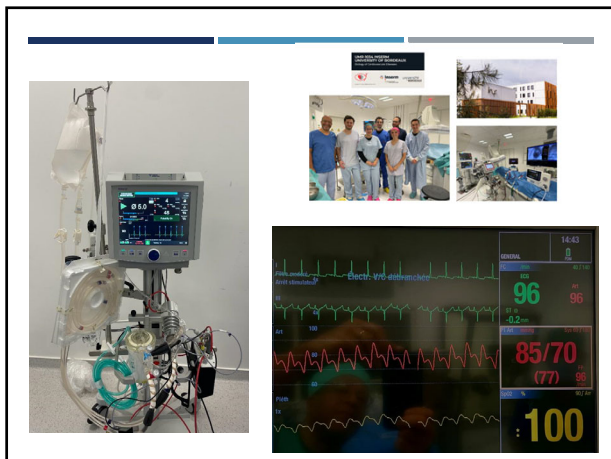


SEH = PEE - PAM

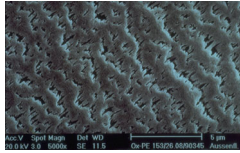


Wang S et al. *Arti Organs* 2015;39:E90-E101



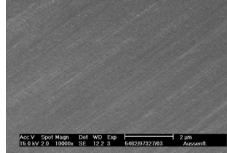


Micropores Membrane (CEC) (Polypropylène)



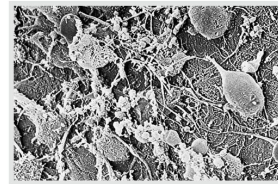
Micropores for gas exchange

Diffusion Membrane (ECMO) (Polyméthylpentène)

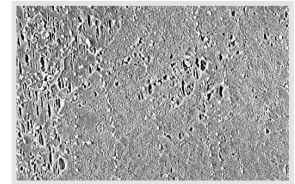


Diffusion of gas through the permeable membrane (human lung)
Better transfert
Impermeable for plasma
Possible coating
No plasma leakage
Duration of use (label CE 14 days)

Surface coating

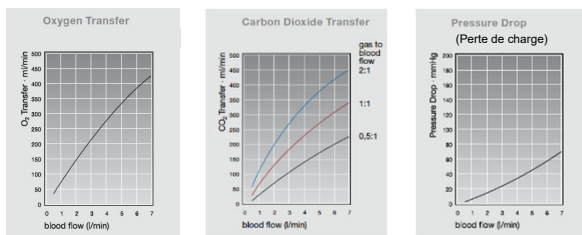


Membrane surface without SAFELINE treatment: Uncoated membrane with depositions potentially leading to high pressure drop phenomena.



QUADROX membrane surface with SAFELINE treatment: The advantage of the SAFELINE treatment under the electron microscope: significant reduction in surface deposition on the QUADROX membrane. The typical transmembrane pressure increase is minimized consistently.

Membrane's characteristics

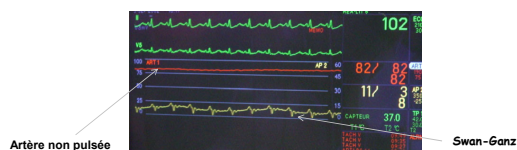


Bloc thermique



MONITORAGE HÉMODYNAMIQUE I

- Pression artérielle moyenne
 - KT radial droit +++ (sang cœur natif)
 - Oxygénation cérébral et myocardique
 - PAM 60-70 mmHg (post-charge dépendance)
 - Aspect linéaire = absence de récupération
- Pression veineuse centrale (décharge cavités droites)



Monitoring Hémodynamique II

- Cathéter de Swan-Ganz
 - Saturation veineuse mélangée O₂ (SvO₂)
 - Index cardiaque (appréciation de la qualité de décharge cavités droites)
 - PAP = maintien d'une pression diastolique minimale (flux intra vasculaire)
 - Evaluation de PAPO (PTDVG) risque d'OAP (PAPO < 18 mmHg)
 - Phase de sevrage
- Echographie (quotidienne ETO > ETT)
 - Prudence traumatisme induit +++ (ETT)
 - Positionnement canules veineuse et artérielle (si centrale)
 - Décharge cavités Dte et Gche
 - Evaluation pressions de remplissage
 - Evaluation de la fonction systolique (FE, Tei index, Strain rate,...)
 - Recherche d'un épanchement péricardique
 - Thrombus intra-cavitaire
 - Valvulopathie (IM, IA)
 - Aspect aorte descendante

MINIMAL BLOOD FLOW TO MEET THE METABOLIC DEMAND DO_2/VO_2

Mixed venous oxygen saturation (SvO₂)

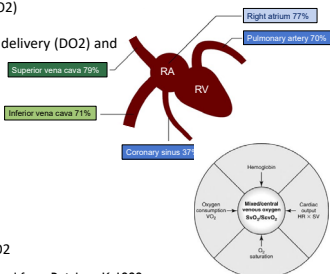
Parameter of balance between oxygen delivery (DO₂) and demand (VO₂)

Increased oxygen extraction (O₂ER)

Goal therapy $70 \pm 5 \%$

$$O_2ER = \frac{SaO_2 - SvO_2}{SaO_2} = 1 - SvO_2$$

Adapted from Reinhart K 1988



CIRCUIT D' ECLS

- Perfusionniste (/ 24 à 48h) et personnel de réanimation (MD et IDE)

- Aspect des canules et lignes

- Caillots, plicature, efficacité du sertissage
- Ligne de reperfusion +++
- Aspect des orifices
- Battement canule veineuse (mauvais drainage)

- Membrane et pompe

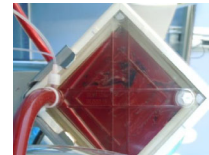
- Recherche de caillots (discuter le changement)
- Purge circuit gaz frais (condensation)

- Console

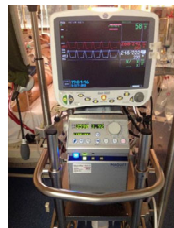
- Meilleur débit avec le moindre de TRM

- Changement de circuit si:

- Caillots,
- hémolyse importante,
- Défaut oxygénation P/F < 150-200 mmHg,
- Systématique (> 14 j)



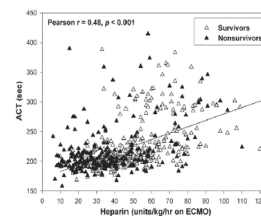
Monitrage pressions pré- et post-filtre



Intérêts:

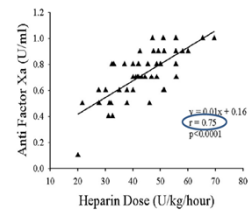
- Suspicion thrombose de filtre si gradient augmente
- Signes évocateurs récupération VG (oscillations pré-filtre)
- Valeurs max 300 mmHg

Faible corrélation ACT avec la dose HNF perfusée



Ann Thorac Surg 2007 83

L'activité antiXa est mieux corrélée à la dose



Nankervis CA et al. ASIAO Journal 2007

Objectif anti-Xa entre 0,2-0,4 UI/ml

MAIN OBJECTIVES OF SHORT-TERM MECHANICAL CIRCULATORY SUPPORT

- Restore end-organ perfusion for limiting multi-organ failure (**supply**)

- ✓ Blood flow, systemic pressure, oxygenation
- ✓ Cerebral, coronary, renal, hepatic and mesenteric perfusion...
- ✓ Poor prognostic of multi-organ failure

- Limit ventricular congestion (**assist**)

- ✓ Volume and pressure unloading +++
- ✓ Reduce risk of pulmonary edema (prolonged MV, VAP...) [**short term**]
- ✓ Reduce the risk of ventricular remodeling [**Mild or long-term**]

VENTRICULAR REMODELING

- 30% of patients after MI (infarct size is the principal risk factor)

- Main origin of chronic heart failure after MI

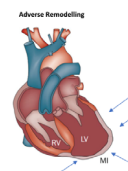
- Increased parietal stress (+++) and myocardial work

- Increased myocardial oxygen consumption

- Decreased coronary blood flow (+++)

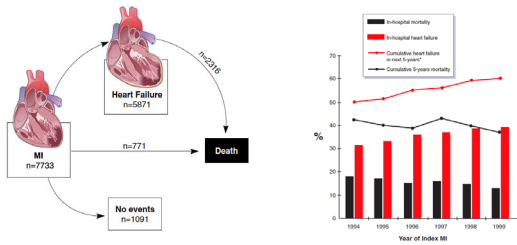
- Increased compliance of infarcted zone

- Thinning of myocardial wall
- Left ventricular dilation



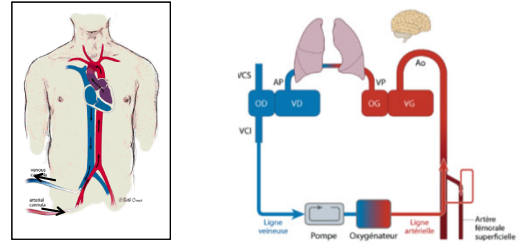
Deja MA et al. Circulation 2012; 125:2639-48
Michler RE et al. J Thorac Cardiovasc Surg 2013; 146:1139-45
Jackson BM JACC 2002; 40:1160-7

CHRONIC HEART FAILURE AFTER MYOCARDIAL INFARCTION



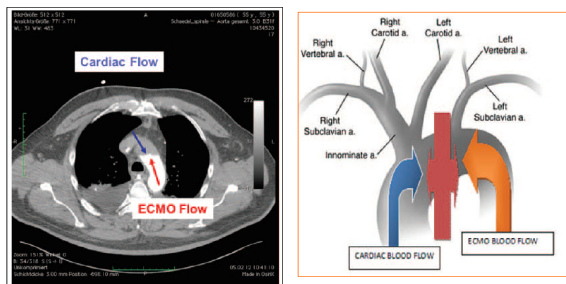
Ezekowitz JA et al. J Am Coll Cardiol 2009;53:13-20

INTERACTION BETWEEN PERIPHERAL VENO-ARTERIAL ECLS AND NATIVE FAILING HEART



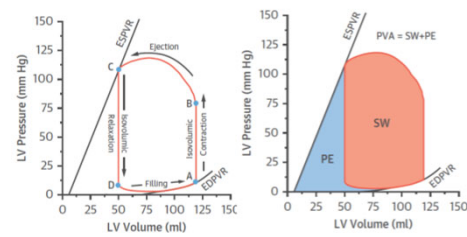
Retrograde aortic flow in total competition with native stream

Calderon J et al. Traitée Anesthésie-Réanimation O. Fourcade (4^{ème} Edition)



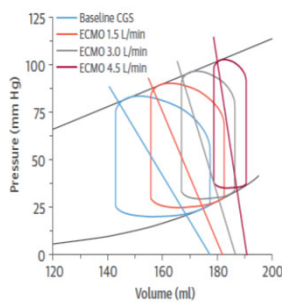
Rupprecht L et al. ASAIO Journal 2013;59:547-53

Pressure-volume area, mechanical work of LV and MvO_2

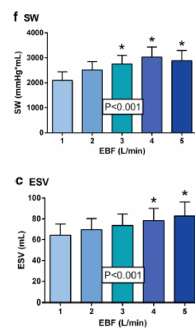


The PVA represents the total mechanical work of LV and may be assessed as the sum of Potential Energy (PE) and stroke work (SW). A relationship exists between PVA and MvO_2

Uriel N et al. J Am Coll Cardiol 2018;72:569-80



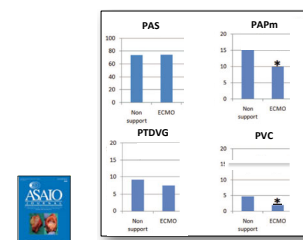
Burkhardt et al. JACC 2015;66:2663-74



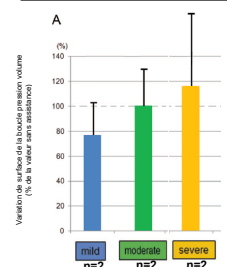
Ostadal P et al. J Transl Med 2016

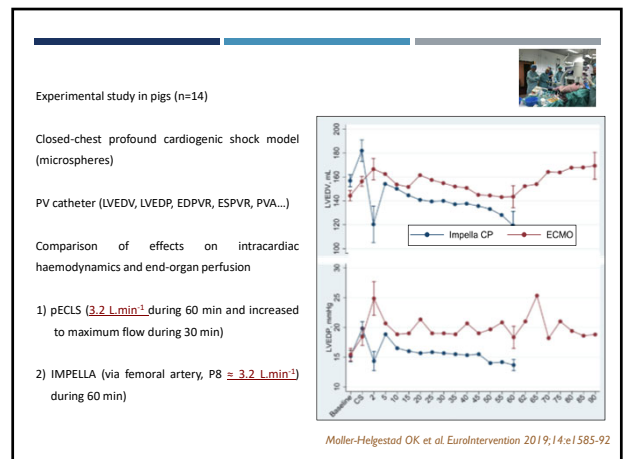
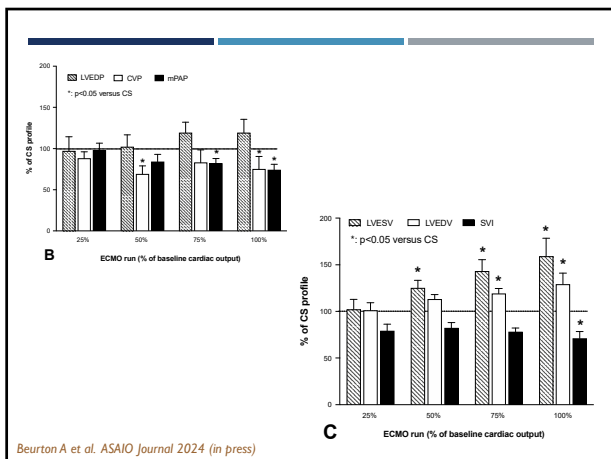
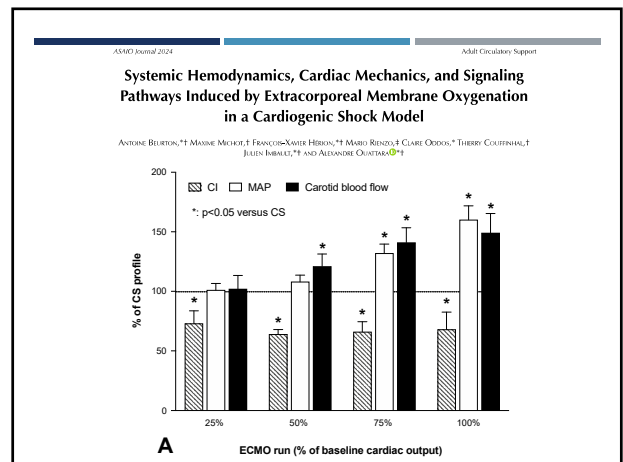
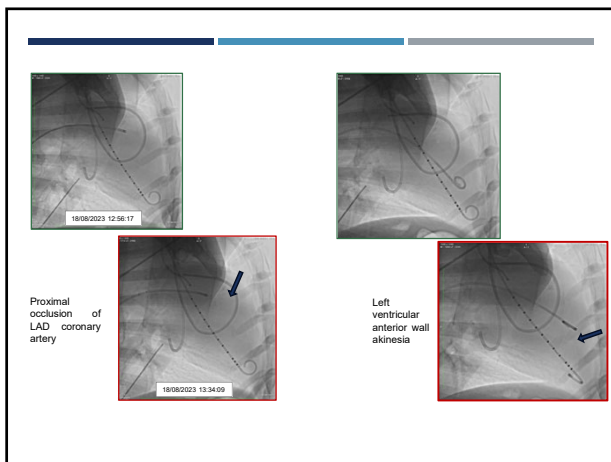
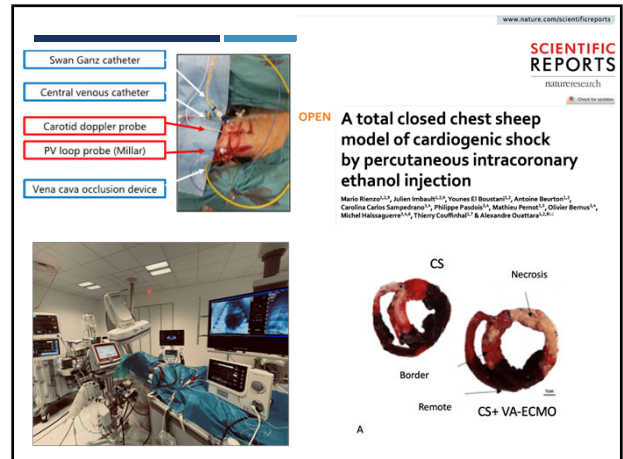
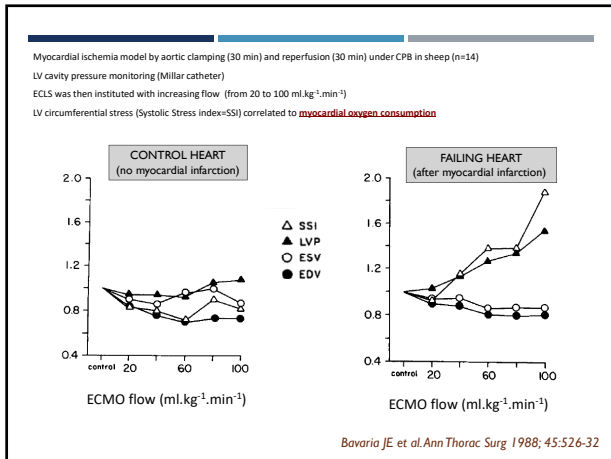
Left Ventricular Mechanical Support with Impella Provides More Ventricular Unloading in Heart Failure Than Extracorporeal Membrane Oxygenation

Basic research in dogs
Haemodynamic monitoring (AP, Swan-Ganz catheter, Millar)
Cardiogenic shock after MI by coronary banding
Peripheral ECLS (10 Fr AF/28 Fr OD)



Total myocardial work (PV curve under ECLS)

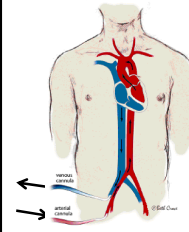




Left ventricular overloading

- Continuous filling of left cardiac cavities
 - Residual pulmonary arterial flow
 - Bronchial circulation (broncho-pulmonary shunts)
 - Thebesian vein (myocardial wall veins draining into the left ventricle)
 - Increase in LV afterload (increase in end-systolic volume)
- Overloading of left ventricle (severe failing heart+++)
- Ventricular dilation and increase in parietal stress
- Subendocardial coronary hypoperfusion

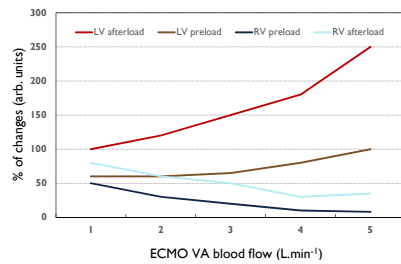
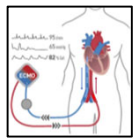
Hydrostatic pulmonary edema



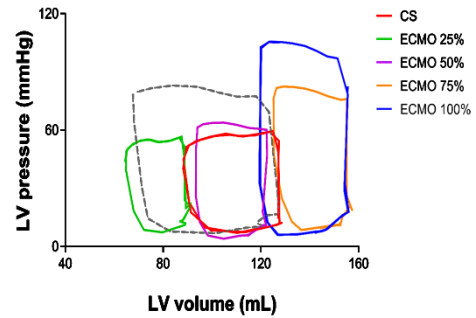
Peripheral ECLS

- Changes in ventricular loading conditions (increase afterload)
- Technical and anatomy dispositions
- Increase in parietal stress
- Risk of pulmonary edema (pulmonary morbidity)
- Alteration of sub-endocardial myocardial perfusion
- Alteration of myocardial recovery
- Increased risk when profound LV dysfunction (no « wash-out ») and in presence of MR ou AR

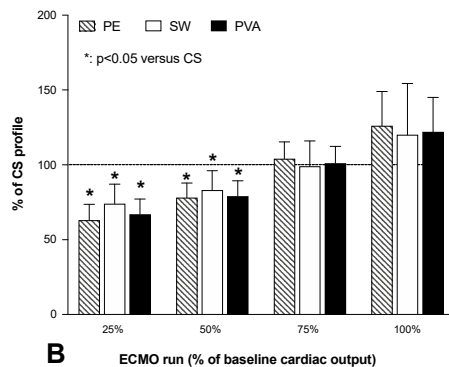
IMPACT OF VA-ECMO ON CARDIAC (LEFT AND RIGHT) LOADING CONDITIONS



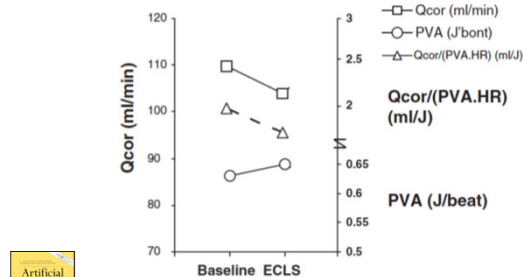
Beurton A et al. ASAIO Journal 2024 (in press)



Beurton A et al. Am J Resp Crit Care Med (in submission)



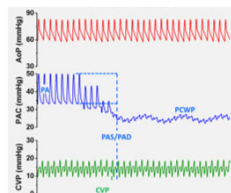
ECLS and coronary perfusion



Sauren LDC et al. Artificial Organs 2007;31:31-8

ASSESSMENT OF LV PRELOAD

- Pulmonary capillary wedge pressure (PCWP) or pulmonary artery occlusion pressure (PAOP)
- Estimation of LA pressure
- Upper limit 18 mmHg



WHITE PAPER

Value of Hemodynamic Monitoring in Patients With Cardiogenic Shock Undergoing Mechanical Circulatory Support

Abhinav Saxena¹, MD
A. Reshad Garan, MD
Navin K. Kapur, MD
William W. O'Neill, MD
JoAnn Lindenfeld, MD
Sean R. Pinney, MD
Nir Uriel, MD
Daniel Burkhardt, MD, PhD
Morton Kern, MD

SUMMARY

"PACs should be used in all patients undergoing MCS to monitor effectiveness, optimize device settings, assess the need for escalation and guide timing and rate of weaning..."

Circulation 2020; 141:1184-97

ASAIO Journal 2021



Guidelines

ELSO Interim Guidelines for Venoarterial Extracorporeal Membrane Oxygenation in Adult Cardiac Patients

« A pulmonary artery catheter should be considered during VA ECMO, as it allows detection of elevated left-sided filling pressures, which may prompt the use of adjunct LV unloading techniques... »

« ...PAC-based continuous cardiac output monitoring during VA ECMO is inaccurate but it may provide an indication of pulmonary artery flow and aid in weaning... »

Table 6. Clinical Monitoring During Venoarterial Extracorporeal Membrane Oxygenation

- Invasive arterial blood pressure monitoring/right radial artery**
 - Pulse pressure—measure of native contractility vs. ECMO blood flow
 - Oxygen saturation—measure of oxygenation in proximal aortic arch/distal aorta of differential oxygenation
- Pulse oximetry/right hand**
 - Oxygen saturation—measure of oxygenation in proximal aortic arch/distal aorta of differential oxygenation
- Pulmonary artery catheter**
 - Directly measures left-sided filling pressure
 - Support indication for adjunct LV unloading
 - Pulmonary artery flow (alternatively, residual pulmonary artery flow can be monitored by measuring end-tidal CO₂)
- Echocardiography**
 - Visualization of proper vascular access and guidance cannulation
 - Optimal tailoring of ECMO support
 - Serial assessment of hemodynamic and cardiac conditions
 - Cardiac assessment during weaning trial
- Electrocardiography**
 - Consider continuous, multi-lead electrocardiographic monitoring
 - Monitoring of limb (single and bilateral comparison) and brain perfusion

ECMO, extracorporeal life support; LV, left ventricle; NIRS, near-infrared spectroscopy; VA ECMO, venoarterial extracorporeal membrane oxygenation.

Lorusso R et al. ASAIO Journal 2021

TO ASSESS RIGHT CAVITIES UNLOADING AND RESIDUAL PULMONARY FLOW

- Central venous pressure < 10 mmHg
- Diastolic pulmonary partial pressure < 20 mmHg
- Cardiac index between 1.0 and 2.0 L.min⁻¹.m⁻²

Merkle J et al. Thorac Dis 2019; 11(Suppl 6): S946–56

CLINICAL IMPLICATIONS

- Avoid over-assistance (blood flow to meet the metabolic demand (SvO₂))
- Screening risk of pulmonary edema (TTE, Swan-Ganz catheter PCWP < 16-18 mmHg...)

STATE OF THE ART



Puajara D et al. Semin Thoracic Surg 2015;27:17-23

The State of the Art in Extracorporeal Membrane Oxygenation

Pharmacologic unloading

MANAGEMENT CONSIDERATIONS

Inotropic support and ventricular assist devices (eg, Impella, Abiomed; intra-aortic balloon pump; TandemHeart trans-septal cannula) should be maintained to facilitate the left-side chamber unloading if cardiac recovery is a possibility.

WHAT'S NEW IN INTENSIVE CARE

Cardiogenic shock in 2024: insight the complex reality of ECLS and left ventricular unloading strategies

Intensive Care Med. 2024;50:971-3

Aure Ughetto^{1,2}, Benedikt Schrage² and Clément Delmas^{1,2}

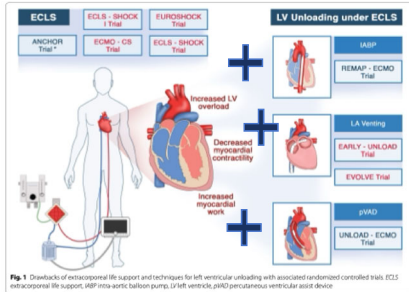
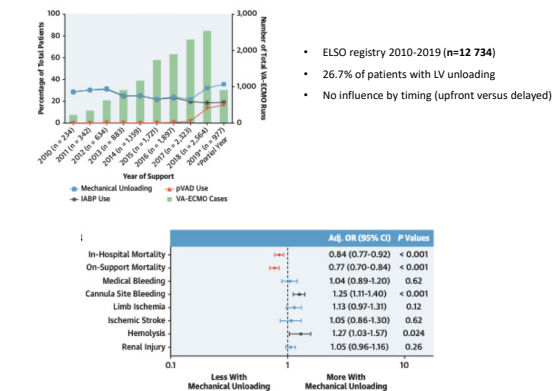


Fig. 1 Downside of extracorporeal life support and techniques for left ventricular unloading with associated randomized controlled trials. ECLS: extracorporeal life support; IABP: intra-aortic balloon pump; LV: left ventricle; PIVD: percutaneous ventricular assist device.

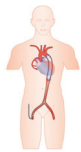
Grandin EW et al. J Am Coll Cardiol 2022;79:1239-50



Intra-aortic balloon pump (IABP)

Improvement energetic balance of myocardium

- Decrease MVO_2 (loading conditions)
- Increase myocardial perfusion



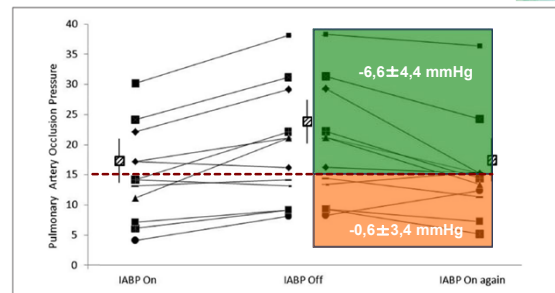
Improvement of end-organ perfusion (Pulsatility)



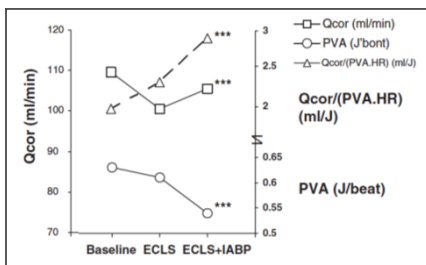
- * Increase diastolic aortic pressure 30 à 70% (rapid inflation r)
- * Decrease systolic aortic pressure of 5-15% (rapid deflation)
- * Decrease LV afterload
- * Decrease LV preload
- * Decrease HR (10%)
- * Increase SV and CO (5-10%).

deWeho S et al. Vascular Pharmacology 2014;61:30-34
Ro SK et al. Eur J Cardiothorac Surg 2014;46:186-92
Aso S et al. Crit Care Med 2016;44:1974-9

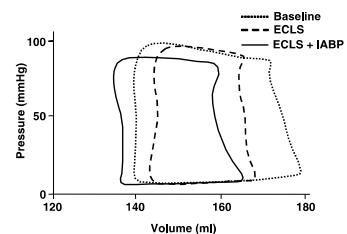
Changes in PCWP



Petroni T et al. Crit Care Med 2014;42:2075-82

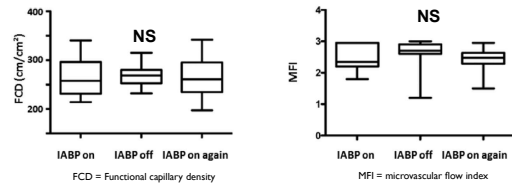


Sauren LDC et al. Artificial Organs 2007;31:31-8



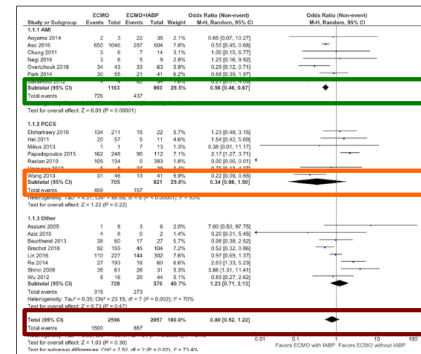
Decrease in LVEDP and LVEDV
Decrease in SV
Reduction of myocardial oxygen consumption

Sauren LDC et al. Artif Organs 2007;31:31-8

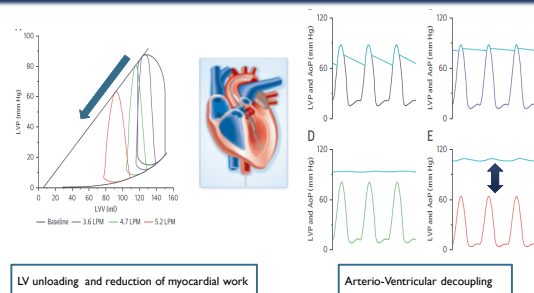


Petroni T et al. Crit Care Med 2014; 42:2075-82

• Meta-analysis of 22 studies (2000-18, n= 4653). The IABP was used in nearly 44% of VA ECMO patients



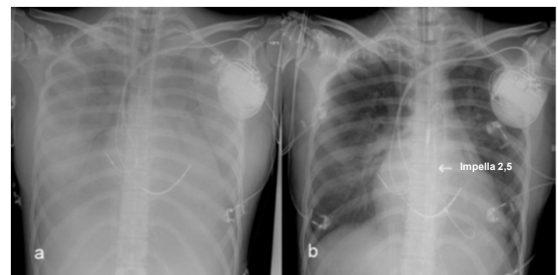
INTRA-CARDIAC HAEMODYNAMIC EFFECTS OF IMPELLA DEVICE



Uriel N et al. J Am Coll Cardiol 2018;72:569-80

Successful left ventricular decompression following peripheral extracorporeal membrane oxygenation by percutaneous placement of a micro-axial flow pump

Jouan J et al. J Heart Lung Transplant 2010;29:135-6



Concomitant implantation of Impella® on top of veno-arterial extracorporeal membrane oxygenation may improve survival of patients with cardiogenic shock

Pappalardo F et al. Eur J Heart Failure 2016

Retrospective multicenter trial (n=2) including **Propensity Score adjustment**
ECLS alone versus ECLS + IMPELLA (2.5 or CP)

Table 1 Comparison of baseline characteristics between patients treated with veno-arterial extracorporeal membrane oxygenation (ECMO) and Impella and patients treated with veno-arterial ECMO only in the original unmatched population (n=157)

Parameter	Total (n=157)	ECMO + Impella (n=34)	ECMO (n=123)	P-value	Absolute standardized difference*
Age, years	55 (46–64)	54 (47–66)	55 (45–64)	0.9	0.0217
Male, n (%)	130 (83)	28 (82)	102 (83)	0.9	0.0263
CRP, n (%)	100 (64)	14 (41)	86 (70)	0.002	0.6101
STEMI, n (%)	85 (54)	15 (44)	70 (57)	0.2	0.2622
PCL, n (%)	56 (36)	16 (47)	40 (33)	0.10	0.2887
pH	7.33 (6.98–7.39)	7.36 (7.08–7.41)	7.16 (6.95–7.37)	0.02	0.5087
Lactates, mmol/L	9.55 (4.40–15.35)	8.96 (3.10–16.25)	10.26 (4.97–15.24)	0.4	0.1173
Concomitant IABP, n (%)	54 (34)	8 (24)	46 (37)	0.10	0.2852

CPB, cardiopulmonary bypass; IABP, intra-aortic balloon pump; STEMI, ST-segment elevation myocardial infarction.
*0–0.2 small effect size; 0.2–0.5 medium effect size; 0.5–0.8 large effect size; 0.8–1 very large effect size.

Concomitant implantation of Impella® on top of veno-arterial extracorporeal membrane oxygenation may improve survival of patients with cardiogenic shock

Pappalardo F et al. Eur J Heart Failure 2016

Retrospective multicenter trial (n=2) including **Propensity Score adjustment**
ECLS alone versus ECLS + IMPELLA (2.5 or CP)

Table 3 Comparison of major outcomes between patients treated with veno-arterial extracorporeal membrane oxygenation (ECMO) and Impella and patients treated with veno-arterial ECMO only in the propensity score matching sample (n=63)

Parameter	Total (n=63)	ECMO + Impella (n=21)	ECMO (n=42)	P-value
Hospital mortality, n (%)	41 (65)	10 (48)	31 (74)	0.04
Bridge to next therapy or recovery, n (%)	28 (44)	13 (62)	15 (36)	0.048
Bridge to recovery, n (%)	19 (30)	8 (38)	11 (26)	0.3
Bridge to VAD, n (%)	8 (13)	4 (19)	4 (9.5)	0.5
Bridge to cardiac transplantation, n (%)	0	0	0	
Duration of ECMO, h	120 (34–234)	148 (72–239)	73.5 (29–217)	0.2
Duration of PVI, h	93 (29–228)	163 (90–228)	48 (17–265)	0.04
CVH, n (%)	18 (29)	10 (48)	8 (19)	0.02
Haemolysis, n (%)	30 (48)	16 (76)	14 (33)	0.004
Major bleeding, n (%)	20 (32)	8 (38)	12 (29)	0.6
Hourly bleeding, n (%)	14 (22)	4 (19)	10 (24)	0.8
LVF at weaning, %	45.3 (30–55)	52.5 (47–55.5)	37.5 (25–50)	0.13

CVH, continuous veno-venous haemofiltration; PCL, mechanical circulatory support; PVI, mechanical ventilation; VAD, ventricular assist device.

ORIGINAL RESEARCH ARTICLE

Left Ventricular Unloading Is Associated With Lower Mortality in Patients With Cardiogenic Shock Treated With Venoarterial Extracorporeal Membrane Oxygenation: Results From an International, Multicenter Cohort Study

Variable	VA-ECMO, matched (n=255)	ECMELLA (n=255)	P value
Bleeding complications			
pericardial bleeding	16/152 (8.5)	18/216 (8.3)	0.99
hemorrhagic stroke	1/191 (0.5)	7/216 (3.2)	0.38
severe bleeding	45/252 (17.8)	38/255 (14.9)	<0.01
moderate bleeding	74/192 (38.5)	112/241 (46.5)	0.01
exsanguination from bleeding	33/201 (16.4)	47/251 (18.7)	0.61
hemolysis	43/150 (28.6)	75/235 (31.9)	0.61
Ischemic complications			
ischemic stroke	22/252 (8.7)	16/230 (7.0)	0.59
interlimb ischemia because of access	31/252 (12.3)	55/255 (21.6)	<0.01
limb ischemia because of abdominal compartment	9/243 (3.7)	23/245 (9.4)	0.02
limb ischemia because of lower extremities	7/243 (2.9)	11/245 (4.5)	0.48
Other complications			
myocardial brain damage	19/179 (10.6)	29/216 (13.4)	0.49
renal replacement therapy	39/255 (15.3)	48/255 (18.8)	0.02
sepsis	44/228 (19.3)	75/251 (29.9)	<0.01

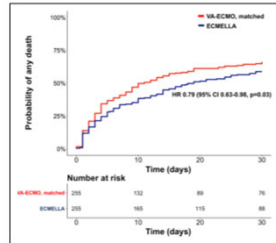


Figure 2. Kaplan-Meier curve of the matched study cohort. ECMELLA indicates Impella-extracorporeal membrane oxygenation; HR, hazard ratio; and VA-ECMO, venoarterial extracorporeal membrane oxygenation.

Schrago B et al. Circulation 2020;142:2095-106

Decompression of the left atrium during extracorporeal membrane oxygenation using a transeptal cannula incorporated into the circuit*

Raouf M. Ayyagari, MD; Albert P. Rocchini, MD; Robert T. Remington, RN; Joseph N. Graziano, MD

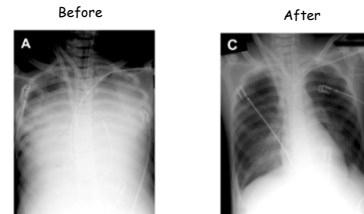
Crit Care Med 2005 Vol. 34, No. 10

Expérience d'une septotomie atriale percutanée (n=7)

Délai médian 11 heures (6-130 heures)

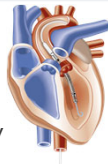
POG = 31 mmHg (22-45)

Durée de procédure 51 min (42-145)



TEST OF RV FUNCTION IN RESPONSE TO CARDIAC OUTPUT RESTORATION

- Mono-ventricular axial pump
- Without oxygenation
- Restore systemic (and anterograde) blood flow
- Increase venous return (to the right heart)
- In vivo test of RV tolerance if LVAD is planned
- Opportunity for rehabilitation program (axillary insertion)



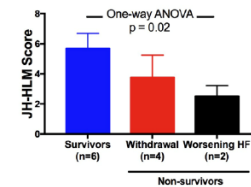
Short communication

IJAO The International Journal of Artificial Organs

Maximum level of mobility with axillary deployment of the Impella 5.0 is associated with improved survival

Michele L Esposito, Janelle Jablonski, Allison Kras, Sara Krasney and Navin K Kapur

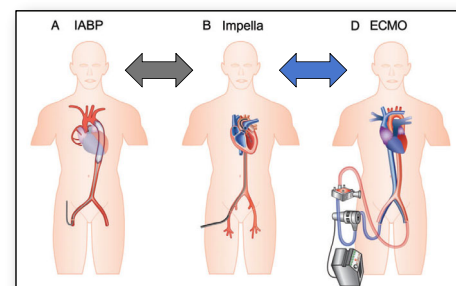
The International Journal of Artificial Organs
1-4
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DOI: 10.1177/03913186187752575
jia.sagepub.com/home/jia

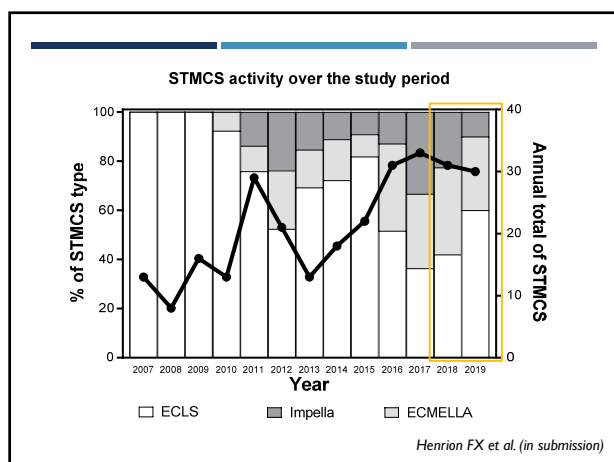


WHICH BEST STRATEGY?

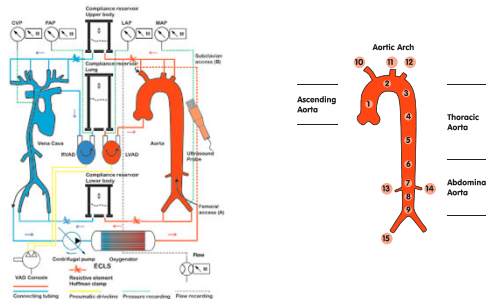


Short-term mechanical circulatory support



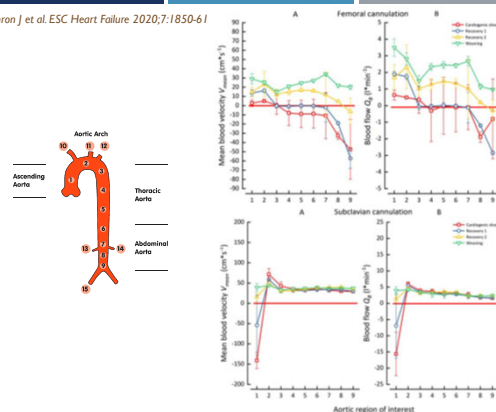


Watershed phenomena during extracorporeal life support and their clinical impact: a systematic in vitro investigation

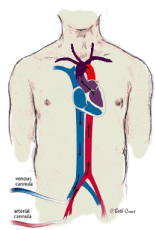


Gehron J et al. ESC Heart Failure 2020;7:1850-61

Gehron J et al. ESC Heart Failure 2020;7:1850-61



Refractory cardiogenic shock and ARDS



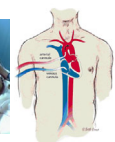
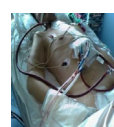
- Persistence of arterial pulmonary blood flow
- Loading of left atrium with hypoxic blood
- Perfusion of supra-aortic and coronary vessels
- Interface between hypoxemic blood and normoxic from ECLS depends:
 - Residual ventricular contractility
 - Level of supply by ECLS
 - Importance of pulmonary disease
- Arlequin Syndrome+++
- **Perform blood gas analysis from right artery radial ++++**

Marasco SF et al. Heart lung and Circulation 2008

SOLUTIONS

Centralisation

- Heavy technology
- Bleeding risk



Axillary cannulation

Stulak JM et al. Seminars Cardiothorac Vasc Anesth 2009;13:176-82



- Advantages**
- Anterograde flow
 - Cephalic perfusion
- Disadvantages**
- Upper limb ischemia



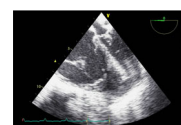
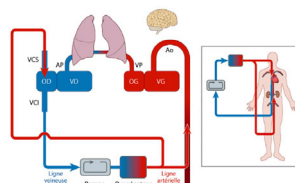
SFAR

Société Française d'Anesthésie et de Réanimation

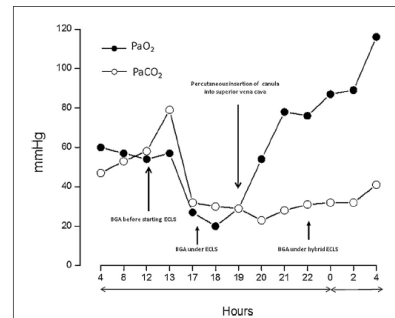
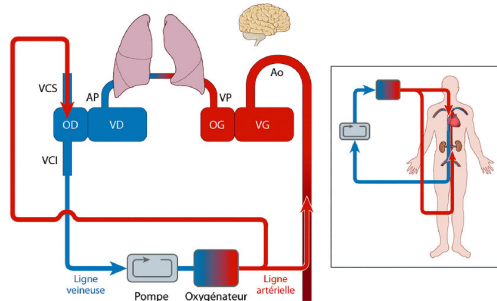
Case report

Moisan M et al. Ann Fr Anesth Réanim 2013;32:e71-75

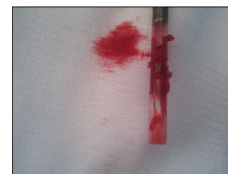
Pulmonary alveolar proteinosis requiring "hybrid" extracorporeal life support, and complicated by acute necrotizing pneumonia



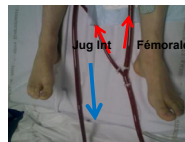
Precardiac oxygenation



Measurement of re-infusion flow by doppler

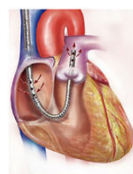
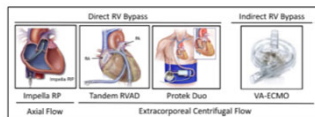


Re-infusion canula 17 Fr



Connection 3/8-1/4 (arterial return)

Mechanical Circulatory Support Devices for Acute Right Ventricular Failure



Kapur NK et al. Circulation 2017; 136:314-26

DEFINITION OF WEANING FROM ECLS

Weaning successful from ECLS is defined as: 1) device removal **and** 2) no further requirement for mechanical support because of recurring CS over the following **7 to 30 days (alive patients)**

Aissaoui N et al. Intensive Care Med 2015;41:902-5

Weaning from ECLS does not mean alive....

In-hospital mortality and successful weaning from venoarterial extracorporeal membrane oxygenation: analysis of 5,263 patients using a national inpatient database in Japan

Aso S et al. Crit Care 2016; 20:80

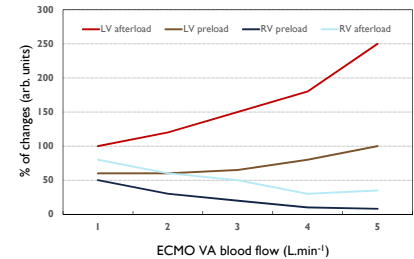
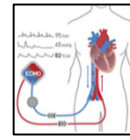
Table 2 In-hospital death and weaning from VA-ECMO among patients classified by six etiological categories

	All	Total in-hospital death	In-hospital death under VA-ECMO	Transferred to other hospitals with VA-ECMO	Weaning from VA-ECMO	Discharged after weaning from VA-ECMO	In-hospital death after weaning from VA-ECMO
All, n (%)	5263	3817	(72.5)	1823	(34.6)	51	(1.0)
Cardiogenic shock, n (%)	4658	3429	(73.6)	1554	(33.4)	44	(0.9)
Pulmonary embolism, n (%)*	353	226	(64.0)	151	(42.8)	7	(2.0)
Hypothermia, n (%)	99	65	(65.7)	49	(49.5)	0	(0.0)
Poisoning, n (%)**	50	31	(62.0)	22	(44.0)	0	(0.0)
Trauma, n (%)	103	66	(64.1)	47	(45.6)	0	(0.0)
						37	(35.9)
						19	(18.4)

*p < 0.001 for in-hospital death after weaning from venoarterial extracorporeal membrane oxygenation (VA-ECMO) vs. cardiogenic shock

**p < 0.05 for in-hospital death after weaning from VA-ECMO vs. cardiogenic shock

IMPACT OF VA-ECMO ON CARDIAC (LEFT AND RIGHT) LOADING CONDITIONS



Beurton A et al. ASAIO Journal 2024 (in press)

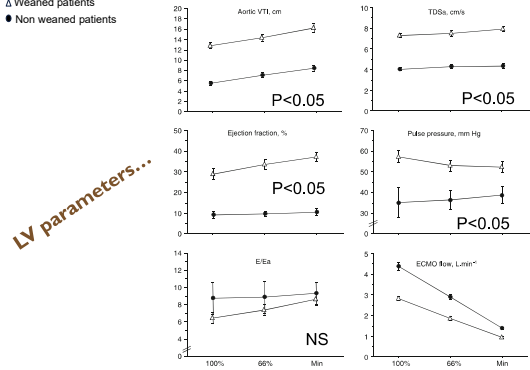
Weaning procedure from VA-ECMO

Changes in ventricular function parameters (Left and Right) in response to ECLS blood flow changes

Assessment of ventricular function (Left and right) at the minimal ECLS flow

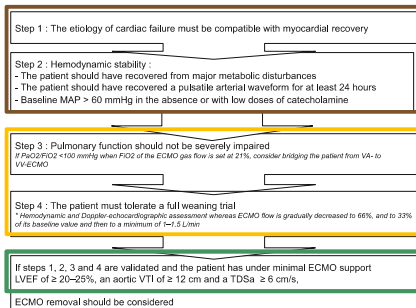
- Increase in **RV** preload
- Increase in **RV** afterload
- Decrease in **LV** afterload

Δ Weaned patients
● Non weaned patients



Aissouli N et al. Intensive Care Med 2011; 37:1738-45

Weaning test from ECLS



60-75% of patients tolerated successfully the weaning from ECLS

Prognostic Implication of RV Coupling to Pulmonary Circulation for Successful Weaning From Extracorporeal Membrane Oxygenation

Kim D et al. J Am Coll Cardiol Imag 2021;14:1523-31

Unicenter study (n=79) between 2016 and 2019

Explore if right ventricular (RV) contractile function and its coupling pulmonary circulation were able to predict successful weaning from ECLS

At full ECLS blood flow +++ (avoid harmful haemodynamic effects)

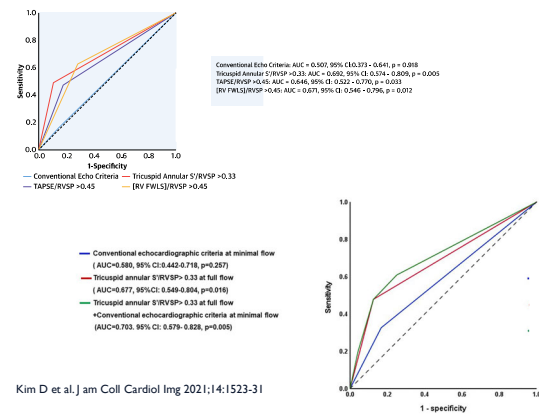
	Successful Weaning (n = 39)	Failed Weaning (n = 20)	p Value
ECMO flow at time of echocardiography, L/min	3.1 (3.0-3.3)	3.1 (3.0-3.3)	0.304
LV end-diastolic dimension, mm	52 (48-50)	56 (49-60)	0.132
LV end-systolic dimension, mm	42 (37-49)	48 (33-59)	0.232
LV ejection fraction, %	24 (19-35)	23 (15-37)	0.862
Transmitral E velocity, m/s	0.56 (0.44-0.68)	0.51 (0.35-0.63)	0.032
Transmitral A velocity, m/s	0.54 (0.36-0.69)	0.58 (0.39-0.77)	0.365
Lateral A velocity, cm/s	4.6 (3.3-6.0)	4.6 (3.3-6.0)	0.309
Lateral E velocity, cm/s	4.4 (3.2-5.7)	4.4 (3.2-5.7)	0.877
E/Ea, m/s	8.6 (6.6-10.3)	14.1 (10.3-18.3)	0.005
LVOT velocity-time integral, cm	8.5 (5.9-12.3)	7.4 (4.3-12.3)	0.338
RV end-diastolic area, cm²	15.1 (10.4-18.5)	16.2 (11.2-19.8)	0.079
RV end-systolic area, cm²	8.8 (7.3-10.7)	9.5 (7.3-11.8)	0.304
Indexed RV end-diastolic area, cm²	7.65 (6.09-9.42)	9.03 (7.07-12.3)	0.054
Indexed RV end-systolic area, cm²	5.17 (3.24-6.17)	6.46 (4.04-8.63)	0.027
TAPSE, mm	12.8 (9.3-15.0)	11.3 (7.3-15.0)	0.243
Indexed TAPSE, mm	8.7 (6.0-11.7)	6.4 (3.5-11.3)	0.054
RV FAC, %	35.7 (26.2-45.5)	28.5 (18.2-38.5)	0.038
RV FACSL, %	16.1 (8.7-24.8)	10.4 (7.3-14.6)	0.054
RVSP, mm Hg	27 (19-36)	33 (24-42)	0.040
Tricuspid annular S' (RVSP)	0.32 (0.19-0.44)	0.22 (0.10-0.42)	0.009
RV FAC/RVSP	1.04 (0.73-1.59)	0.87 (0.53-1.31)	0.134
TAPSE/RVSP	0.450 (0.29-0.68)	0.31 (0.21-0.40)	0.019
RV FACSL/RVSP	0.49 (0.33-0.87)	0.31 (0.23-0.48)	0.001

At full ECLS blood flow....!

TABLE 3 Predictors for Successful ECMO Weaning

	Univariable Analysis	
	OR (95% CI)	p Value
Age	1.02 (0.98-1.06)	0.286
Idiopathic dilated cardiomyopathy	0.15 (0.40-0.64)	0.004
Continuous renal replacement therapy	0.28 (0.11-0.75)	0.011
Extracorporeal CPR	2.81 (1.05-7.55)	0.040
LV venting	0.07 (0.02-0.19)	<0.001
Early venting (<12 h)	0.17 (0.04-0.70)	0.014
LV ejection fraction >20%	1.25 (0.46-3.42)	0.661
LVOT velocity-time integral ≥10 cm	1.25 (0.46-3.42)	0.664
Mitral annular S' ≥6 cm/s	2.37 (0.82-6.85)	0.113
FAC/RVSP ≥0.94	2.52 (0.98-6.52)	0.056
Tricuspid annular S'/RVSP >0.33	8.01 (2.15-29.92)	0.002
TAPSE/RVSP >0.45	4.09 (1.34-12.43)	0.013
[RV FWLS/RVSP] >0.45	4.21 (1.53-11.41)	0.005

Kim D et al. J Am Coll Cardiol Img 2021;14:1523-31



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Step 1: The **etiology** of cardiac failure must be compatible with myocardial recovery

Step 2: **Hemodynamic stability:**

- The patient should have recovered a pulsatile arterial waveform for at least 24 hours
- Baseline MAP >60 mmHg in the absence or with low doses catecholamine and/or pulsed pressure >
- The patient should have recovered from major metabolic disturbances

Step 3: **Pulmonary function** should not be severely impaired

If PaO2/FiO2 <100 mmHg when FiO2 of the ECMO gas flow is set at 21%, consider bridging the patient from VA- to VV-ECMO

Step 4: The patient **must tolerate a full weaning trial**

- * Hemodynamic and echocardiographic assessment whereas ECMO flow is gradually decreased to 66%, and to 33% of its baseline value and then to a minimum of 1-1.5 L/min

If steps 1, 2, 3 and 4 are validated and the patient has under minimal ECMO support:

- AVE of >20-25%, at aortic flow >3.0 cm/s and a 1DSe >6 cm/s
- 3D-RV ejection fraction (if feasible) >24.6%

ECMO removal should be considered

Tricuspid annular S'/RVSP >0.33

TAPSE/RVSP >0.45

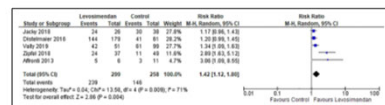
[RV FWLS/RVSP] >0.45

at full ECLS blood flow

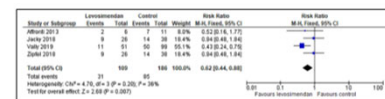
Adapted from Ortuno S et al. Ann Cardiothorac Surg 2019;8(1):E1-8

Effects of levosimendan on weaning and survival in adult cardiogenic shock patients with veno-arterial extracorporeal membrane oxygenation: systematic review and meta-analysis

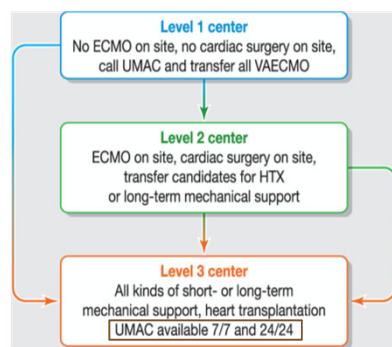
Analysis of risk of successful weaning from VA ECMO in patients treated with levosimendan



Analysis of risk of all-cause mortality in VA ECMO patients treated with levosimendan



Burgos LM et al. Perfusion 2020



Flecher E et al. Arch Cardiovasc Dis 2019; 12:441-9

Intensive Care Med 2001; 37:824-830
 DOI: 10.1007/s00134-001-2158-5

ORIGINAL

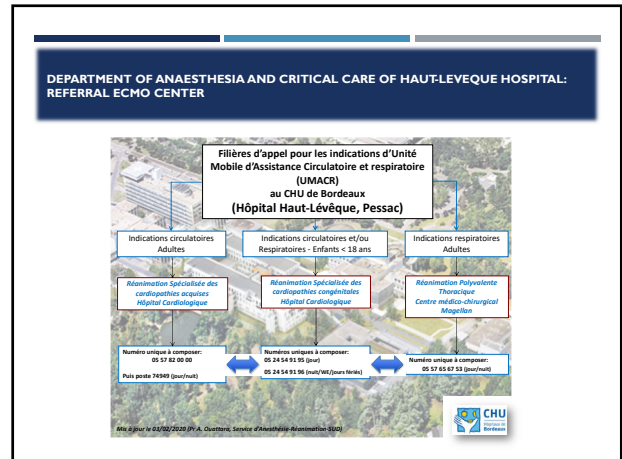
P. Forrest
 J. Ratchford
 B. Burns
 R. Herkes
 A. Jackson
 B. Plimkett
 P. Torzillo
 P. Nair
 E. Granger
 M. Wilson
 R. Pye

Retrieval of critically ill adults using extracorporeal membrane oxygenation: an Australian experience



800 000 km² (X 1,5 F)
 6 100 000 h





Mobile circulatory support unit



ROTAFLOW and
specific (smaller)
TROLLEY
(2020)



Miniaturisation...



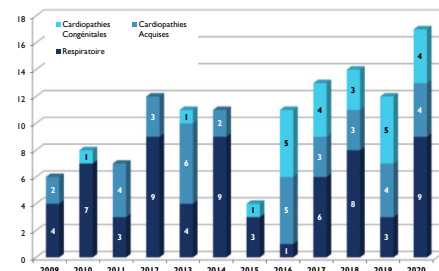
CardioHelp (Maquet)



HLS Module Advanced 7.0:

- Oxygénateur à membrane diffusion
- Pompe centrifuge innovante intégrée
- Marquage CE pour **30 J et le transport**
- Traitement de surface BIOLINE

Global activity (n=126)



Data from 126 patients managed by our Mobile assistance unit between 2009 and 2020

Age, year	33 [16-53]
Male, %	84 (67%)
Duration of ECMO, days	10 [2-23]
IGS II	52 [38-68]
In-hospital Mortality, %	46 (36.2%)
Distance, km	10 [10-146]

SHORT-TERM MECHANICAL CIRCULATORY SUPPORT...

- ✓ Rescue and temporary strategy to restore end-organ perfusion
- ✓ Intrinsic morbidity and even mortality (ischemia, bleeding,...)
- ✓ Impact on intra-cardiac haemodynamic (failing myocardium)
- ✓ Potentially harmful for myocardial recovery (ventricular remodeling)
- ✓ Multidisciplinary cardiogenic shock team approach
- ✓ Multimodal and evolutive strategy
- ✓ Further research still required to confirm and quantify outcome improvement
- ✓ And identify the best strategy regarding the severity of CS

INSERM, UMR 1034 Biology of cardiovascular diseases Systemic and cardiac haemodynamics in acute heart failure

Program leaders

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Professor Thierry COUFFINHAL MD PhD (Cardiology)

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Associate Professor Julien IMBAULT MD
Associate Professor Mario RIENZO MD PhD

PhD Students

Antoine BEURTON MD MSc

MSc Students

FX HERION MD
Claire ODDOS MD
Elara BORDIER MD
Simon VEYRET MD

