

Pratiques satellites de la CEC

DU Circulation Extra-Corporelle en chirurgie cardiaque et en suppléances d'organes
10/01/2025

Gaspard Cadier, Service d'Anesthésie-Réanimation GH Sud

Conflit d'intérêt

- Aucun

Plan

- Système de récupération sanguine autologue/Récupération de Sang Péri-Opératoire (RSPO)
- Hémofiltration per CEC
- Adsorption per CEC
- Adjonction de CO₂ per CEC

Récupération de sang périopératoire (RSPO)

Permet d'administrer par voie intraveineuse au patient son propre sang récupéré dans le site chirurgical ou la plaie postopératoire lors d'une intervention chirurgicale hémorragique.
Elle fait partie des techniques d'économie du sang qui permettent d'éviter de recourir au sang homologué.

HAS - Service évaluation médico-économique et santé publique - Juillet 2006

Rationnel

- 1) Chirurgie sous CEC: hémorragique
- 2) Transfusion sanguine nocive en elle même, couteuse, stocks limités.
- Limitation des pertes sanguines (et donc de la transfusion) si récupération de celles ci en per opératoire

-Henke PK, McKeown E, et al. A case-cohort study of postoperative myocardial infarction: impact of anemia and cardioprotective medications. Surgery 2014;156
-SHANDER A, HARE GMT. What is really dangerous: anaemia or transfusion? Br J Anaesth 2011

Principe RSPO per CEC

- **1) Aspiration « directe »**, non lavée ni centrifugée:
 - Cannule du chirurgien directement reliée au bol de CEC, pas de traitement du sang autre qu'une filtration passive (20 à 40µm).
 - Reperfusion du sang total (avec facteurs de coag, plaquettes mais aussi débris cellulaires+++ et facteurs de l'inflammation)

Principe RSPO per CEC

- 1) Aspiration « directe », non lavée ni centrifugée
- 2) Aspiration « traitée », lavée et centrifugée: déplasmatisation du sang aspiré avec concentration des Erythrocytes, pertes des plaquettes, des facteurs de coagulations mais aussi d'une partie des débris cellulaires.

- Dualité

1) Aspiration directe

- Sang total
- Principalement pour l'extravasation « immédiate »
- Eviter d'aspirer de la « soupe inflammatoire » (ex: sang stagnant dans le médiastin)
- Aspiration sur tubulure de petit calibre responsable en elle même d'une inflammation importante

2) Aspirations lavées/centrifugées Principe

<https://www.youtube.com/watch?v=OIQ63oev6kc>

2) Aspirations lavées/centrifugées Recommandations

2017 EACTS/EACTA Guidelines on patient blood management for adult cardiac surgery ^{FREE}

Domenico Pagano ✉, Milan Milojevic, Michael I Meesters, Umberto Benedetto, Daniel Bolliger, Christian von Heymann, Anders Jeppsson, Andreas Koster, Ruben L Osnabrugge, Marco Ranucci ... Show more
Author Notes

European Journal of Cardio-Thoracic Surgery, Volume 53, Issue 1, January 2018, Pages

The routine use of cell salvage should be considered to prevent transfusions.

IIa

B



RECOMMANDATIONS FORMALISEES D'EXPERTS

R **Question 2.** L'utilisation d'un système de récupération sanguine préopératoire pendant la chirurgie a-t-elle un impact sur la survenue de complications postopératoires ou sur la durée d'hospitalisation ?
Experts : Guillaume LEBRETON (SFCTCV, Paris), Bertrand ROZEC (SFAR, Nantes)

R5.2 – Il est probablement recommandé d'utiliser un système de récupération sanguine peropératoire pour limiter la transfusion érythrocytaire en chirurgie cardiaque.

GRADE 2+ (accord FORT)

RFE commune SFAR - SFCTCV

Société Française d'Anesthésie et de Réanimation (SFAR)
Société Française de Chirurgie Thoracique et Cardiovasculaire (SFCTCV)

2) Aspirations lavées/centrifugées Transfusion

- Nombreux essais cliniques

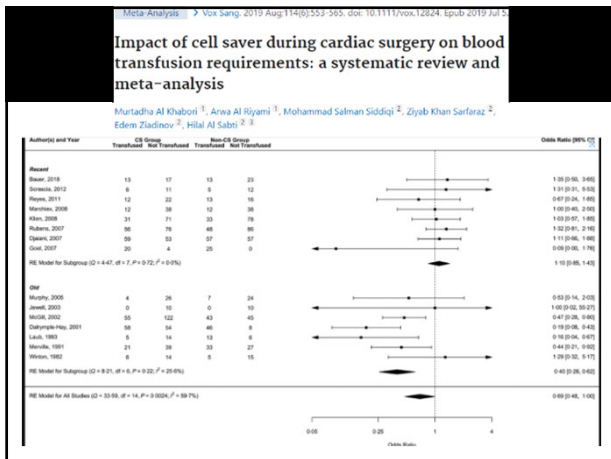
Cochrane Database of Systematic Reviews

Cell salvage for minimising perioperative allogeneic blood transfusion

Cochrane Systematic Review - Intervention | Version published: 14 April 2010 [see what's new](#)
<https://doi.org/10.1002/14651858.CD001888.pub4>

- 33 essais de chirurgie cardiaque
- Cell salvage réduirait la transfusion de CGR
- Sans effets secondaires marqués (pas + de reprise chir)
- Etudes méthodologiquement pauvres et hétérogène

- Quelle utilisation per op? de toutes les aspirations? Mediastinales? Circuit de CEC?



2) Aspirations lavées/centrifugées Transfusion

Interact Cardiovasc Thorac Surg. 2011 Feb;12(2):189-93. doi: 10.1510/ivota.2010.261538. Epub 2010 Nov 30.

Cell saving systems do not reduce the need of transfusion in low-risk patients undergoing cardiac surgery.

Reyes G¹, Prieto M, Alvarez P, Orts M, Bustamante J, Santos G, Sarra A, Planas A.

63 patients
Retransfusion CS: 461 +/- 174mL
40 vs 46% de transfusion
Protocoles de transfusion? Taux d'Hb finaux?

Interact Cardiovasc Thorac Surg. 2011 May;12(5):824-6. doi: 10.1510/ivota.2010.249136. Epub 2011 Feb 5.

The use of cell salvage in routine cardiac surgery is ineffective and not cost-effective and should be reserved for selected cases.

Maran S¹, Molloy D, Fisher BM, Pullan MD.

n <7% of cases the volume of blood loss was sufficient enough to be washed and returned. We conclude that the routine use of cell savers in all cardiac operations affords no benefit and consumes additional revenue.

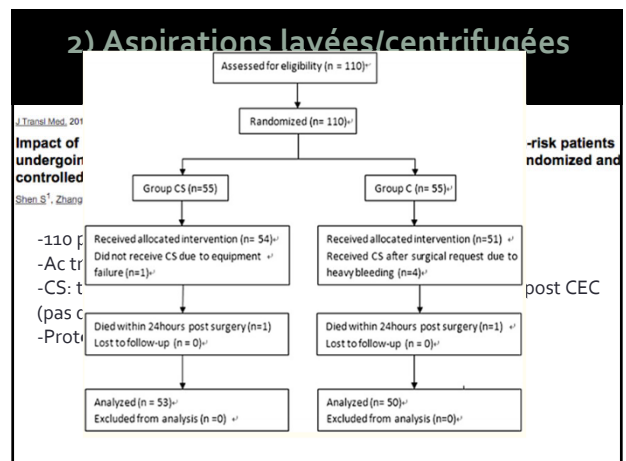
2) Aspirations lavées/centrifugées Transfusion

J Transl Med. 2016 Jul 29;14(1):228. doi: 10.1186/s12967-016-0986-6.

Impact of intra-operative cell salvage on blood coagulation in high-bleeding-risk patients undergoing cardiac surgery with cardiopulmonary bypass: a prospective randomized and controlled trial.

Shen S¹, Zhang J², Wang W³, Zheng J⁴, Xie Y⁵.

- 110 patients Cell Saver vs Control
- Ac tranex 30mg/kg
- CS: traitement uniquement du sang médiastinal et pre et post CEC (pas de traitement de l'aspi de cardiectomie) VS Contrôle : aspirations jetées
- Protocole de transfu de CGR et PFC et plq (TEG)



2) Aspirations lavées/centrifugées Transfusion

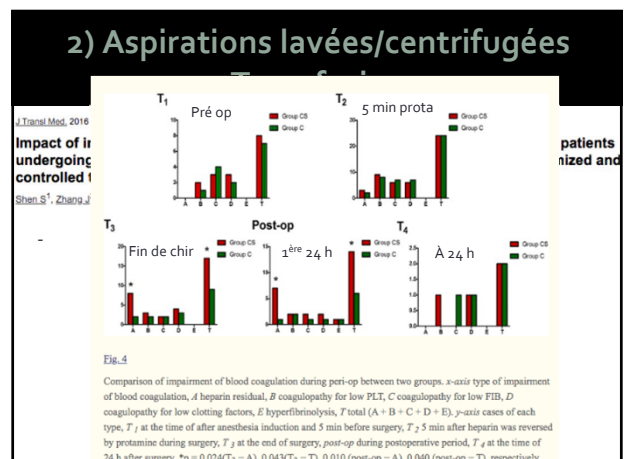
J Transl Med. 2016 Jul 29;14(1):228. doi: 10.1186/s12967-016-0986-6.

Impact of intra-operative cell salvage on blood coagulation in high-bleeding-risk patients undergoing cardiac surgery with cardiopulmonary bypass: a prospective randomized and controlled trial.

Shen S¹, Zhang J², Wang W³.

Variable	Group CS (n = 53)	Group C (n = 50)	P value
RBC			
Proportion	22 (41.51)	39 (78.00)	0.0002
Volume (U)	2.11 (2.66)	5.40 (3.48)	<0.0001
FFP			
Proportion	10 (18.87)	9 (18.00)	0.910
Volume (mL)	118.76 (253.82)	129.00 (284.92)	0.953
PLT			
Proportion	12 (22.64)	10 (20.00)	0.744
Volume (U)	1.81 (3.56)	1.92 (3.94)	0.916

Data are presented as mean (SD) or number (percentage)



2) Aspirations lavées/centrifugées

Summary of adverse events during post-op

Variable	Group CS (n = 53)	Group C (n = 50)	P value
Excessive bleeding	17 (32.08)	8 (16.00)	0.038
Resternotomy	3 (5.66)	2 (4.00)	1.000
Cardiovascular failure	6 (11.32)	7 (14.00)	0.682
Severe arrhythmias requiring treatment	4 (7.55)	3 (6.00)	1.000
Myocardial infarction	1 (1.89)	4 (8.00)	0.196
Infection	7 (13.21)	6 (12.00)	0.948
Wound	3 (5.66)	2 (4.00)	1.000
Others	4 (7.55)	4 (8.00)	1.000
Renal failure	5 (9.43)	4 (8.00)	1.000
Respiratory failure	3 (5.66)	2 (4.00)	1.000
Epileptic syndrome	1 (1.89)	1 (2.00)	1.000
Cognitive decline	0 (0.00)	3 (6.00)	0.111
Death	0 (0.00)	0 (0.00)	

Donnée manquante: volume traité??

Data are presented as number (percentage)

2) Aspirations lavées/centrifugées Transfusion

Editorial

More Is Not Always Better: Effects of Cell Salvage in Cardiac Surgery on Postoperative Fibrinogen Concentrations

Journal of Cardiothoracic and Vascular Anesthesia 34 (2020) 2383–2385

“potential harms on the coagulation system when using intraoperative cell salvage. In complex cases, excess cell salvage can only maintain hemoglobin levels but significantly alters hemostatic function via the elimination of platelets and soluble coagulation factors”

2) Aspirations lavées/centrifugées Transfusion

■ Futur?

Editorial > Anesthesiology. 2021 Aug 1;135(2):2
doi: 10.1097/ALN.0000000000003820.

Combined Platelet and Erythrocyte Salvage: Evaluation of a New Filtration-based Autotransfusion Device

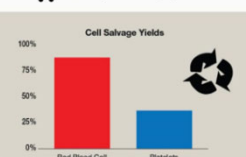
Alexandre Mansour, Benoit Decouture, Mikail Rouss
Véronique Picard, Alexandre Ouattara, Christilla Bac
Nicolas Nesseler, Isabelle Gouin-Thibault

Combined Platelet and Erythrocyte Salvage: Evaluation of a New Filtration-based Autotransfusion Device

Laboratory evaluation of a new medical device

Device salvages and washes red blood cells and platelets for autotransfusion

• Processed 30 units of healthy human whole blood using the device
• Analyzed cell integrity and function



Device salvaged red blood cells and platelets with intact cell integrity and function.
Mansour A, et al. ANESTHESIOLOGY, 2021.

2) Aspirations lavées/centrifugées Transfusion

ANESTHESIOLOGY

Combined Platelet and Red Blood Cell Recovery during On-pump Cardiac Surgery Using same™ by i-SEP Autotransfusion Device: A First-in-human Noncomparative Study (i-TRANSEP Study)

Alexandre Mansour, M.D., Ph.D., Antoine Beurlon, M.D., Anne Godier, M.D., Ph.D., Bertrand Rozec, M.D., Ph.D., Diane Zlotnik, M.D., Fabienne Nédélec, Pharm.D., Pascale Gausson, Pharm.D., Ph.D., Mathieu Fiore, M.D., Ph.D., Etienne Bessier, Pharm.D., Ph.D., Nicolas Nesseler, M.D., Ph.D., Alexandre Ouattara, M.D., Ph.D.
ANESTHESIOLOGY 2023; 139:287–97

Combined Platelet and Red Blood Cell Recovery during On-pump Cardiac Surgery using same™ by i-SEP Autotransfusion Device (i-TRANSEP Study)

First-in-human noncomparative study of a blood and platelet recovery device in 50 adult patients undergoing cardiac surgery with cardiopulmonary bypass

Hypothesis:
• Red blood cell recovery > 80%
• Posttreatment hematocrit > 40%
• Removal of > 90% heparin
• Removal of > 75% free hemoglobin

Observed 50 patients:
• 18 CABG
• 20 valve surgery
• 6 aortic root surgery

	Median (25–75th percentile)
RBC recovery/cycle (%)	86.1 (80.8–91.6)
Posttreatment Hct (%)	41.8 (39.7–44.2)
Platelet recovery (%)	52.4 (44.2–60.1)
Posttreatment platelet concentration (10 ⁹ /L)	116 (93–146)

In this first-in-human study, the blood and platelet recovery device was able to simultaneously recover and wash both platelets and red blood cells for transfusion during cardiac surgical procedures.
Mansour A, et al. ANESTHESIOLOGY, 2023.

2) Aspirations lavées/centrifugées inflammation/débris

Ann Thorac Surg. 2010 May;89(5):1511–7. doi: 10.1016/j.athoracsur.2010.02.003.

Cell saver for on-pump coronary operations reduces systemic inflammatory markers: a randomized trial.

Damgaard S¹, Nielsen CH, Andersen LW, Bendtsen K, Tvede M, Steinbrüchel DA.

Eur J Cardiothorac Surg. 2013 Sep;44(3):506–11. doi: 10.1093/ejcts/etz019. Epub 2013 Feb 12.

Cell salvage of cardiomyocyte suction blood improves the balance between pro- and anti-inflammatory cytokines after cardiac surgery.

Gäbel J¹, Westerberg M, Bengtsson A, Jernsson A.

Interact Cardiovasc Thorac Surg. 2016 Mar;22(3):298–304. doi: 10.1093/icvts/ivw355. Epub 2015 Dec 23.

Intraoperative cell salvage during cardiac surgery is associated with reduced postoperative lung injury.

Espels GE¹, van Klarenbeek J², Gu Y³, van Oeveren W⁴, de Vries AJ⁵.

Eur J Cardiothorac Surg. 2003 Apr;23(4):633–6.

A prospective randomised comparison of cardiomyocyte suction and cell saver for recycling shed blood during cardiac surgery.

Jewell AE¹, Akoum EF, Susanna SK, Braidley D, Hookinson D, Cooper G.

2) Aspirations lavées/centrifugées Contre indication

- Produits que l'on ne voudrait pas retrouver dans la circulation sanguine...
- Antiseptiques iodés, eau oxygénée, alcool, eau stérile, adjuvants de la coagulation, Liquide amniotique ou gastrique....
- Endocardite?
- Néoplasie?

2) Aspirations lavées/centrifugées Endocardite

- Concept de réalisation CEC....
- Que change l'utilisation ou non du cell saver?

2) Aspirations lavées/centrifugées Néoplasie

2011 Update to The Society of Thoracic Surgeons and the Society of Cardiovascular Anesthesiologists Blood Conservation Clinical Practice Guidelines**

Class IIb

Jump to Section

Go

- 1 In high-risk patients with known malignancy who require CPB, blood salvage using centrifugation of blood salvaged from the operative field may be considered since substantial data support benefit in patients without malignancy, and new evidence suggests worsened outcome when allogeneic transfusion is required in patients with malignancy. (Level of evidence B)

2) Aspirations lavées/centrifugées Néoplasie

Medicine (Baltimore), 2019 Jul 56(27):e16040. doi: 10.1097/MD.00000000000016040.

Survival analysis of intraoperative blood salvage for patients with malignancy disease: A PRISMA-compliant systematic review and meta-analysis.

Nu WW¹, Zhang WY¹, Zhang WH², Yang L¹, Deng XQ¹, Ou MC¹, Yang YX¹, Liu HB¹, Zhu T¹.

- 17 études incluses (1 seule prospective)
- Pas de différence entre utilisation CS vs No CS

2) Aspirations lavées/centrifugées Néoplasie

Medicine (Baltimore), 2019 Jul 56(27):e16040. doi: 10.1097/MD.00000000000016040.

Survival analysis of intraoperative blood salvage for patients with malignancy disease: A PRISMA-compliant systematic review and meta-analysis.

Nu WW¹, Zhang WY¹, Zhang WH², Yang L¹, Deng XQ¹, Ou MC¹, Yang YX¹, Liu HB¹, Zhu T¹.

Mortalité:

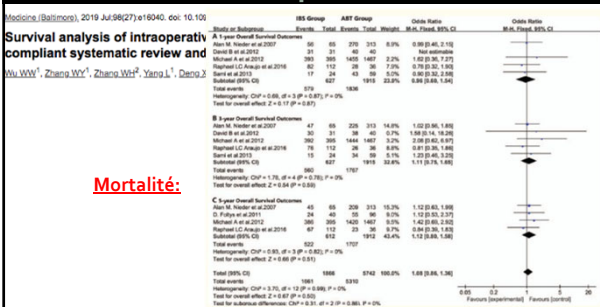


Figure 1

Meta-analysis forest plot of the overall survival outcomes. (A, 1-year overall survival outcome, B, 3-year overall survival outcome, C, 5-year overall survival outcomes).

2) Aspirations lavées/centrifugées Néoplasie

Medicine (Baltimore), 2019 Jul 56(27):e16040. doi: 10.1097/MD.00000000000016040.

Survival analysis of intraoperative blood salvage for patients with malignancy disease: A PRISMA-compliant systematic review and meta-analysis.

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RECIDIVE:

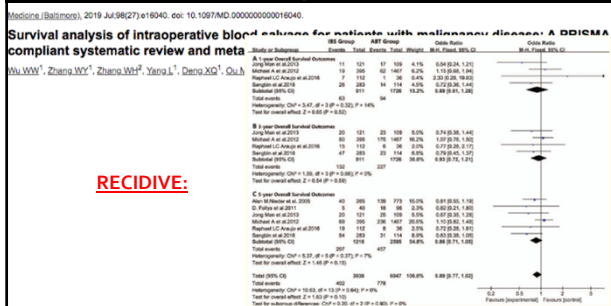


Figure 3

Meta-analysis forest plot of the recurrence rate. (A, 1-year recurrence rate, B, 3-year recurrence rate, C, 5-year recurrence rate).

En pratique Cell saver

- S'intègre dans une pratique globale d'épargne sanguine
- Possible coagulopathie pour hauts volumes traités (pauvres en plaquettes et en facteurs de coag et possible quantité non négligeable d'héparine)
- Cancer/endocardites: non contre indiqué
- Utilisation pour tous les patients? **Quelle gestion de aspiration sur le champ?**

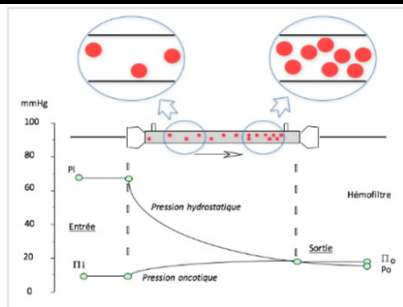
Hémofiltration



Principe

- Principe de convection, le plasma sanguin est filtré à travers une membrane semi-perméable grâce à une différence de pression transmembranaire.
- Les solutés de taille inférieure aux pores de la membrane (électrolytes, cytokines...) sont ainsi éliminés avec l'eau plasmatisque, tandis que les cellules sanguines et les protéines de grande taille, sont retenues par la membrane.

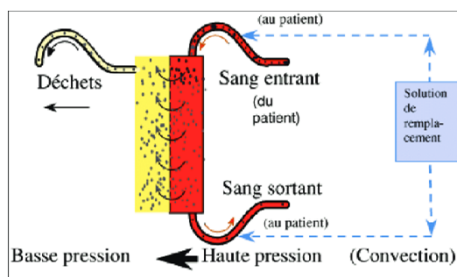
Principe



Principe

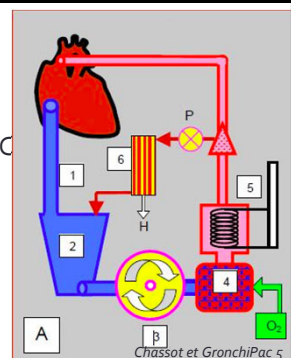
- Élimination de l'eau, des électrolytes et des petites molécules fonction:
- 1) Pression hydrostatique
- 2) taille des pores=point de coupure (poids moléculaire < 30-50 k Daltons)
- 3) débit sanguin
- Quantité de volume ainsi éliminé est nommé: « UF » pour UltraFiltrat
- Branchement du filtre sur pompe dédiée, intégré au circuit de CEC

Principe



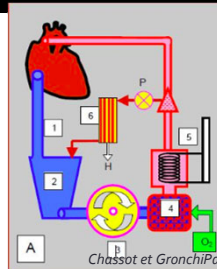
A) Hémofiltration conventionnelle

- Per CEC
- Perte d'eau et électrolytes
- hémococoncentration per CEC
- Perte de poids



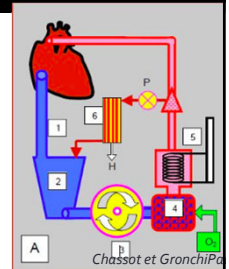
A') Hémofiltration « balance Zéro »

- Per CEC
- Perte d'eau et électrolytes
- Compensée par ajout d'un soluté cristalloïde (ou autre)



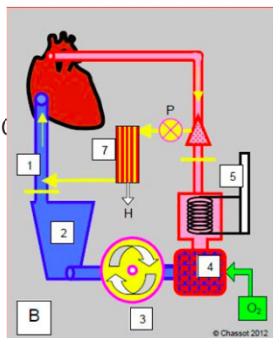
A') Hémofiltration « balance Zéro »

- Per CEC
- Perte d'eau et électrolytes
- Compensée par ajout d'un Soluté cristalloïde
- épurait de hauts volumes
- Pas d'hémoconcentration
- Pas de perte de poids
- « Renouvellement » plasma « débarrassé » de petites molécules potentiellement délétères.



B) Hémofiltration modifiée (MUF)

- En fin de CEC
- Perte d'eau et électrolytes
- hémoconcentration post CEC
- débit sang 150mL/min
- stratégie diurétique



17. Groom RC, Akl BF, Albus RA, et al. Alternative method of ultrafiltration after cardiopulmonary bypass. *Ann Thorac Surg.* 1994;58:573-574.

Indications hémofiltration

- Evidentes: Surcharge volémique avec hémodilution et œdème interstitiel (souffrance multi organe), désordre électrolytique ou élimination d'un toxique hémofiltrable (taille et liaison protéiques)
- Supposée: Bénéfice à l'élimination de molécules pro inflammatoires (majeures parties des cytokines pro inf ont un poids moléculaire < point de coupure)

Littérature hémofiltration

A) Hémofiltration conventionnelle

2011 Update to The Society of Thoracic Surgeons and the Society of Cardiovascular Anesthesiologists Blood Conservation Clinical Practice Guidelines^{*,*}

Conventional ultrafiltration is a method of ultrafiltration used during CPB. One meta-analysis [347], four randomized trials [348, 349, 350, 351], and two cohort studies [352, 353] report the use of conventional ultrafiltration in patients undergoing cardiac procedures using CPB. Subgroup analysis of five studies included in the meta-analysis by Boodhwani [347] demonstrated no advantage in terms of red cell usage or blood loss with conventional ultrafiltration alone [349, 350, 351, 352, 353].

- Pas d'avantage en terme de transfusion

A) Hémodiffiltration conventionnelle

Ann. Card. Anesth. 2016 Jan-Mar;16(1):45-51. doi: 10.4103/0971-8764.173018.

Conventional hemofiltration during cardiopulmonary bypass increases the serum lactate level in adult cardiac surgery.

Soliman B¹, Fouad E, Belghith M, Abdelmassoud T.

-Etude observationnelle, CEC prog, hypothermie modérée -138 patient avec HF VS 145 sans (pts similaires)
-HF après départ en CEC pour obj Ht entre 25 et 30% -instabilité hémodynamique et hyperlactatémie en per et au sevrage de la CEC....

Table 2: Intraoperative data of patients

Dopamine			
Number of patients	96	123	0.001
Dose (µg/kg/min)	5.23±1.63	7.95±1.97	0.001
Epinephrine			
Number of patients	42	40	0.001
Dose (µg/kg/min)	0.08±0.02	0.12±0.05	0.001
Norepinephrine			
Number of patients	16	33	0.001
Dose (µg/kg/min)	0.03±0.02	0.05±0.03	0.008
IABP (number of patients)	17	38	0.015
Packed red blood cells (units)	1.55±0.28	2.78±0.78	0.005
Urine (L)	1.49±0.28	3.62±0.46	0.001
Fluid removal (L)	2.56±0.22	-	0.001

Table 4. Postoperative data of patients.

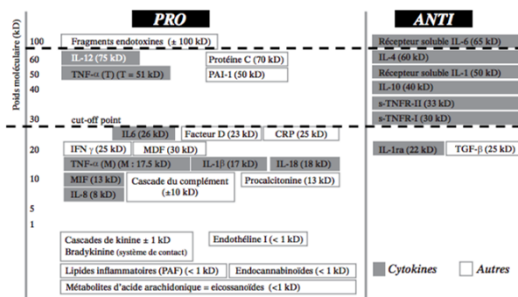
A) Hémodiffiltration « balance zero »

2011 Update to The Society of Thoracic Surgeons and the Society of Cardiovascular Anesthesiologists Blood Conservation Clinical Practice Guidelines*

With ZBUF, the ultrafiltrate fluid is replaced with an equal volume of balanced electrolyte solution during CPB. Patients may benefit from ZBUF through the removal of mediators and products of systemic inflammatory response syndrome, rather than as a result of fluid removal [356]. Randomized controlled trials investigating ZBUF in adult cardiac procedures did not demonstrate a reduction in blood loss or transfusion with the application of ZBUF [357, 358].

■ Pas d'avantage en terme de transfusion

A) Hémodiffiltration « balance zero »



Silverstein W. Mediator removal with CRRT (continuous renal replacement therapy): complement and cytokines. Am J Kidney Dis 1997;30(Suppl 4):S38-43.

A) Hémodiffiltration « balance zero »

Perfusion, 2002 Mar;17(2):111-5.

Inflammatory mediator removal by zero-balance ultrafiltration during cardiopulmonary bypass.

Talman RD¹, Dumond M, Brown D.

Author information

1. Cardiovascular Anesthesiology, University of Michigan Medical Center, Ann Arbor, Michigan, USA.

Table 2 Mean (std. dev.) values taken immediately before surgery (T1), immediately after surgery (T2), and 12 h post-surgery (T3)

	T1		T2		T3	
	Control	ZBUF	Control	ZBUF	Control	ZBUF
Hematocrit (%)	36.36 (3.92)	36.48 (3.86)	29.10 (4.86)	29.03 (2.81)	29.94 (3.49)	30.95 (4.01)
Hemoglobin (g/dl)	12.64 (1.43)	12.51 (1.26)	10.16 (1.90)	10.03 (1.05)	10.69 (1.90)	10.83 (1.37)
White Cell Count (x10 ⁹ /dL)	6.85 (1.80)	6.47 (1.29)	11.30 (2.46)	10.96 (2.46)	11.97 (2.90)	11.23 (1.83)
IL-1 (pg/ml)	0.62 (1.33)	0.24 (0.22)	0.61 (0.95)	0.63 (0.59)	0.79 (1.08)	0.51 (0.62)
TNF-α (pg/ml)	2.81 (1.93)	3.23 (2.66)	61.62 (32.51)	64.43 (68.88)	160.83 (103.62)	145.07 (74.00)
C5a (pg/ml)	2.01 (2.42)	3.46 (3.50)	2.91 (2.40)	9.43*(8.68)	1.86 (2.13)	5.27 (4.92)
Prothrombin time (s)	111.04 (1026.90)	1417.57 (862.31)	1686.91 (1448.48)	3282.09*(2407.75)	528.27 (600.78)	837.59 (585.10)
Partial thromboplastin time (s)	278.30 (317.65)	158.37 (98.48)	1007.35 (338.83)	1299.17 (277.13)	110.24 (53.05)	175.75 (126.01)
SVR	13.53 (0.77)	13.23 (0.61)	19.14 (3.17)	19.38 (4.87)	15.02 (1.37)	15.06 (1.05)
PVR	33.87 (12.45)	41.37 (22.96)	37.60 (10.03)	30.18 (7.96)	32.60 (3.83)	33.07 (4.92)
Cardiac output (l/min)	1095.80 (272.64)	1307.53 (421.56)	731.40 (301.70)	732.33 (238.65)	844.93 (193.37)	894.67 (377.75)
Cardiac index (l/min/m ²)	105.33 (42.10)	112.47 (69.76)	75.53 (29.19)	77.13 (46.09)	87.93 (34.19)	73.67 (41.34)
Cardiac output (l/min)	5.03 (1.29)	4.79 (1.37)	7.91 (2.22)	7.30 (1.53)	6.61 (1.62)	6.93 (1.84)
Cardiac index (l/min/m ²)	2.63 (0.59)	2.29 (0.51)	4.08 (1.06)	3.59 (0.65)	3.39 (0.66)	3.29 (0.64)

SVR=systemic vascular resistance, PVR=peripheral vascular resistance. *Indicates a significant difference from corresponding control values as determined by Kruskal-Wallis test of the ANCOVA residuals (p<0.05).

A) Hémo

Time since ICU admission, h	12.9 (11.7)	12.7 (11.8)
Patients on ECMO	47 (42%)	52 (46%)
Patients on IABP	19 (17%)	15 (13%)
Patients on IABP or ECMO	58 (52%)	60 (54%)
SAPS II	54.0 (12.3)	55.1 (12.3)
SOFA score	11.5 (2.8)	12.0 (2.9)
Glasgow coma score	13.5 (3.0)	13.2 (3.4)
Systolic blood pressure, mm Hg	113 (24)	109 (25)
Diastolic blood pressure, mm Hg	63 (14)	62 (14)
Mean blood pressure, mm Hg	79 (14)	77 (14)
Epinephrine dose, µg/kg/min	0.23 (0.39)	0.27 (0.46)
Norepinephrine dose, µg/kg/min	0.23 (0.45)	0.32 (0.70)
Inotropic score ¹	45.6 (53.9)	55.1 (80.7)
Creatinine, µmol/L	148 (81)	162 (77)
Blood urea nitrogen, mmol/L	11.1 (5.7)	12.0 (5.6)
Urine output, ml		
<500	36 (31)	46 (41)
500-999	35 (32)	26 (23)
>1,000	41 (37)	40 (36)
Lactate, mmol/L	5.0 (3.9)	4.8 (3.8)
Bicarbonate, mmol/L	19.5 (4.9)	19.1 (3.8)

Am J Respir Crit Care Med. 2015 Nov 15;192(11):1511-1517.

Early High-Volume Hemofiltration Study.

Cornblath A¹, Bratchov B¹, Amour J², Gao J³, A¹², Trouillet JL¹, Mallet A³, Chastain J¹, Les

1. University of Michigan Medical Center, Ann Arbor, Michigan, USA.

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A) Hémo

Variable	Early HVHF (n = 712)	Standard Care (n = 712)	Odds Ratio (95% CI)	P Value
Mortality				
Day 30	40 (36%)	40 (36%)	1.00 (0.58-1.73)	1.00
Day 60	48 (43%)	42 (38%)	1.25 (0.73-2.05)	0.82
Day 90	51 (46%)	43 (38%)	1.34 (0.79-2.28)	0.28
ICU	49 (44%)	44 (39%)	1.20 (0.71-2.05)	0.50
In-hospital	50 (45%)	44 (39%)	1.25 (0.73-2.12)	0.42
ICU length of stay, d				
For ICU survivors	13 (7-25)	13 (8-29)	—	0.78
For patients who died in the ICU	11 (2-21)	6 (2-14)	—	0.15
Hospital length of stay, d				
For survivors	37 (22-54)	29 (20-49)	—	0.31
For patients who died in-hospital	11 (2-22)	6 (2-14)	—	0.12
Day 1-60 ICU-free days	21 (0-48)	22 (0-48)	—	0.66
Day 1-60 hospital-free days	0 (0-24)	0 (0-31)	—	0.75
Days with catecholamines	5 (3-9)	5 (3-9)	—	0.50
Day 1-30 catecholamine-free	3 (0-10)	3 (0-8)	—	0.64

Am J Respir Crit Care Med. 2015 Nov 15;192(11):1511-1517.

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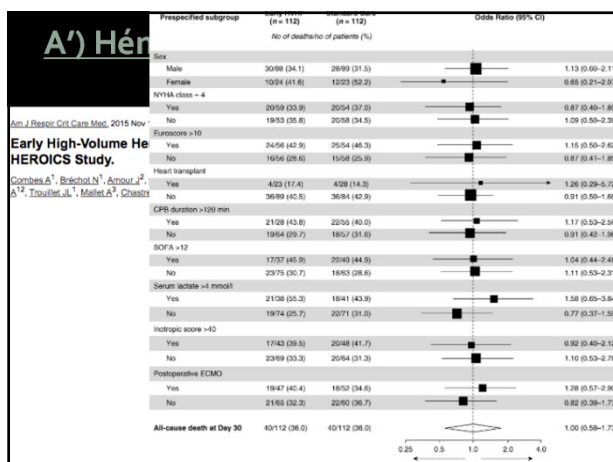
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B) Hémodiafiltration modifiée (MUF)

2017 EACTS/EACTA Guidelines on patient blood management for adult cardiac surgery

2011 Update to The Society of Thoracic Surgeons and the Society of Cardiovascular Anesthesiologists Blood Conservation Clinical Practice Guidelines*

2017 EACTS/EACTA Guidelines on patient blood management for adult cardiac surgery

Domenico Pagano, Milan Milosevic, Michael I. Meesters, Umberto Benedetto, Daniel Bolliger, Christian von Heymann, Anders Jeppesen, Andreas Koster, Ruben L. Ovenshige, Marco Ranucci, Show more

European Journal of Cardio-Thoracic Surgery, Volume 53, Issue 1, January 2018, Page

2017 EACTS/EACTA Guidelines on patient blood management for adult cardiac surgery

2011 Update to The Society of Thoracic Surgeons and the Society of Cardiovascular Anesthesiologists Blood Conservation Clinical Practice Guidelines*

Boothwan M, Williams K, Babaei A, Gill G, Saleem N, Rubens FD

Ultrafiltration reduces blood transfusions following cardiac surgery: A meta-analysis.

- Méta analyse sur HF:
- Diminution transfusion en terme de CGR, PFC et plaquettes dans les études MUF
- Etudes « Anciennes » fin go début 2000, hypothermie, cardioplégie importante en Volume, méthode imparfaite

Indications hémodiafiltration per CEC

- Probablement pas d'intérêt en systématique
- en fonction du patient:
 - Surcharge volémique avec hémodilution et œdème interstitiel (souffrance multi organe)
 - Elimination d'un médicament épurable (ex: Dabigatran) ou d'un « déchet métabolique »
 - Gestion du volume dans le réservoir de cardiologie: un diurétique peut aussi faire l'affaire...
- Non prouvée: Bénéfice à l'élimination de molécules pro inflammatoires Et anti inflammatoires

Hémodiafiltration

Section through an adsorber, Adsorber bead, Internal structure

Hémodiafiltration Cytosorb /Jafron

Section through an adsorber, Adsorber bead, Internal structure

- Cartouche remplie de billes de polymère poreuses
- Adsorption de surface des molécules entre 5 et 60 kDa
- Sur un principe uniquement « physique »
- Adsorption concentration dépendante
- surface d'échange de 4 terrains de foot
- Saturation possible

Intégration circuit

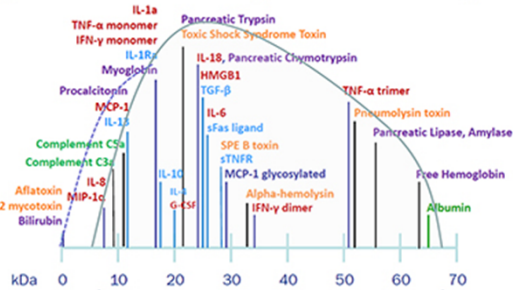
- Faible résistance d'écoulement
- Débit sanguin recommandé 150-500 ml/min
- Préchargé/primé avec une solution saline isotonique

C. Michael Gibson et al

Standard integration of DrugSorb-ATB within a CEC circuit: DrugSorb is integrated as a parallel shunt circuit to the main CEC circuit. Blood flow enters to the parallel circuit is after the pump in the main CEC circuit, and blood flow returns from the parallel circuit is to the blood reservoir of the main CEC circuit.

Principe Hémoadsorption

CytoSorb® Active in Cytokine Sweet Spot



Littérature



Cytokines

Crit Care. 2019 Apr 3;23(1):108. doi: 10.1186/s13054-019-2399-4.

Cytokine clearance with CytoSorb® during cardiac surgery: a pilot randomized controlled trial.

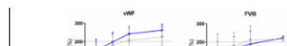
Poel EC¹, Alberici L^{2,3}, Bauer-Doering A⁴, Mancuso C^{4,5}, Roumy A⁶, Kirsch M^{6,3}, De Stefano E⁶, Lissauer L^{1,3}, Schneider AG^{6,7}.

15 patients cytosorb 15 controles, chirurgie réglée
Pas d'effet secondaire délétère
Pas de variation des taux de cytokines ni facteurs coag

Cardio-pulmonary bypass characteristics

Median bypass duration—IQR) min	138 (87–207)	145 (130–183)
Median cross-clamp duration (IQR) min	115 (68–159)	122 (97–146)
Centrifugal pump—no. (%)	5 (33.3)	4 (26.7)

CONCLUSIONS: CytoSorb® HA during CPB was not associated with a decrease in pro- or anti-inflammatory cytokines nor with an improvement in relevant clinical outcomes. The procedure was feasible and safe. Further studies should evaluate the efficacy of CytoSorb® HA in other clinical contexts.



Cytokines

The effect of perioperative hemadsorption in patients operated for acute infective endocarditis—A randomized controlled study

Silke Asch¹, Tobias Peter Kaufmann, Michaela Walter, Marcus Leistner, Bernd C. Danner, Thorsten Perl, Ingo Kutschka, Heidi Niehaus

First published: 21 June 2021 | <https://doi.org/10.1111/aor.14019>

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Patients with confirmed IE of any type (native or prosthetic valve) and localization (affected valve) undergoing cardiac surgery were included in this study and randomly assigned to either the HA group or the control group. Exclusion criteria were a lack of informed consent,

The primary endpoint of the study was the postoperative course of cytokine levels (IL-6, TNF-α, IL-1β) and infection parameters (CRP, PCT, leucocytes). Secondary endpoints were the development of the severity-of-the disease (estimated by the SOFA, SAPS II and APACHE II score), postoperative catecholamine and fluid requirement, the incidence of adverse events as well as in-hospital mortality.

Cytokines

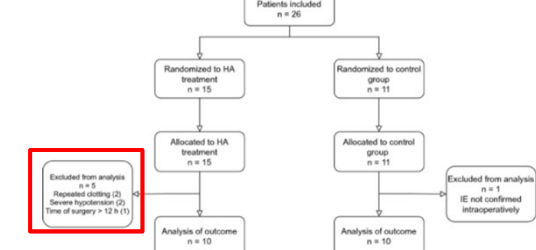
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Silke A

Ingo K

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	HA group n = 10	Control group n = 10	P value
Baseline parameters			
Age [years]	65 (53-70)	69 (56-81)	.315
Female gender [n]	3	1	.582
Body mass index [kg/m ²]	26.0 (21.5-30.6)	25.0 (20.7-31.8)	.853
LVFE [%]	50 (50-56)	60 (54-62)	.105
EuroSCORE II [%]	8.5 (2.7-16.4)	3.6 (2.6-11.8)	.393
Re-operation [n]	3	2	1.000
Emergency surgery [n]	2	1	1.000
Endocarditis-related parameters			
Single valve IE [n]	7	9	1.000
Multiple valves IE [n]	3	1	1.000
Septic embolism [n]	4	4	1.000
Cardiogenic shock [n]	1	0	1.000
Co-morbidities			
Arterial hypertension [n]	4	7	.370
Insulin-dependent diabetes [n]	1	3	.582
Peripheral artery disease [n]	2	0	.474
Renal replacement therapy [n]	1	1	1.000
COPD [n]	2	0	.474
Cardiac surgery			
AV surgery [n]	5	2	.350
MV surgery [n]	2	4	.628
AV and MV surgery [n]	2	3	1.000
MV and TV surgery [n]	1	1	1.000
Cross clamp time (minutes)	90 (65-150)	118 (60-148)	.888
CPB time (minutes)	126 (94-276)	180 (93-219)	.963

The effect of
for acute

Silke Asch
Ingo Kutsch

First published

Cytokines

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(n = 7). All patients survived to discharge. No significant differences concerning median cytokine levels (IL-6 and TNF-α) were observed between both groups. CRP and PCT baseline levels were significantly higher in the HA group (59.5 vs. 26.3 mg/dL, $P = .029$ and 0.17 vs. 0.05 µg/L, $P = .015$) equalizing after surgery. Patients in the HA group required significantly higher doses of vasopressors (0.093 vs. 0.025 µg/kg/min norepinephrine, $P = .029$) at 12 hours postoperatively as well as significantly more overall volume replacement (7217 vs. 4185 mL at 12 hours, $P = .015$; 12 021 vs. 4850 mL at 48 hours, $P = .015$). HA therapy did neither result in a reduction of inflammatory parameters nor result in an improvement of hemodynamic parameters in patients operated for IE. For a more targeted use of HA therapy, appropriate selection criteria are required.

Cytokines

Lancet Respir Med. 2021 Jul; 9(7): 755–762.

Published online 2021 May 14. doi: [10.1016/S2213-2600\(21\)00177-8](https://doi.org/10.1016/S2213-2600(21)00177-8)

PMCID: PMC8121541

PMID: 34002036

Cytokine adsorption in patients with severe COVID-19 pneumonia requiring extracorporeal membrane oxygenation (CYCOV): a single centre, open-label, randomised, controlled trial

Alexander Soudry, MD, PhD, Eva Weber, PhD, Marina Rieder, MD, PhD, Achim Lohrer, MD, PhD, Tim Niehaus, BA, PhD

The CYCOV trial was a single-centre, randomised, controlled, parallel group, open-label, superiority trial. All adult patients (≥18 years of age) admitted to the participating intensive care units (ICUs) of the Freiburg University Medical Center with reverse transcriptase (rt) PCR-confirmed SARS-CoV-2 infection who were selected to receive venovenous ECMO were eligible. Exclusion criteria were a known patient

In the intervention group, a CytoSorb adsorber was incorporated into the ECMO system and replaced every 24 h for a total treatment duration of 72 h. Routinely, the adsorber was installed in the ECMO as part of the system setup before connecting it to the patient circuit, but at the latest within 4 h after initiation of the

The primary endpoint was serum IL-6 after 72 h of ECMO with or without cytokine adsorption. Secondary endpoints were ICU survival and 30-day survival, days on ECMO, days on mechanical ventilation, serum lactate, Willebrand factor, D-dimers, vasopressor dosage, amount of fluid substitution, fluid balance after 72 h, and Sequential Organ Failure Assessment score after 24, 48, and 72 h. [21](#)

Cytokines

Lancet Respir Med.

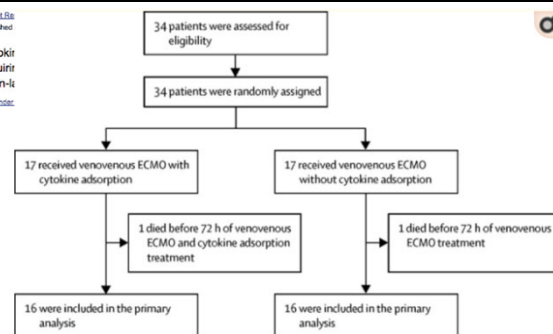
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	Cytokine adsorption group (n=17)	Control group (n=17)
Age, years	62.0 (54.0-70.0)	59.0 (43.0-66.0)
Sex		
Female	5 (29%)	4 (24%)
Male	12 (71%)	13 (76%)
Body mass index, kg/m ²	29.4 (24.6-39.2)	29.6 (24.4-35.4)
Laboratory values		
Interleukin-6, ng/mL	357.0 (127.4-1036.0)	289.0 (84.7-787.0)
C-reactive protein, mg/L	25.8 (9.0-68.0)	35.9 (12.6-142.2)
Procalcitonin, ng/mL	0.73 (0.10-5.84)	1.34 (0.17-5.98)
Troponin, ng/mL	20.0 (0.00-5.00)	14.0 (0.00-5.00)
D-dimer, µg/mL	10.0 (0.0-30.0)	14.0 (0.0-30.0)
Neutrophils, %	9.4 (0.0-40.4)	14.8 (0.0-40.4)
Lymphocytes, %	14.0 (0.0-40.4)	0.0 (0.0-40.4)
Monocytes, %	0.0 (0.0-40.4)	0.0 (0.0-40.4)
Willebrand factor antigen, %	60.0 (40.0-80.0)	39.0 (40.0-50.0)
D-dimer, ng/mL	0.0 (0.0-10.0)	4.0 (0.0-10.0)
Spies		
SOFA	0.0 (0.0-10.0)	0.0 (0.0-10.0)
APACHE II	0.0 (0.0-10.0)	0.0 (0.0-10.0)
PEEP	0.0 (0.0-10.0)	0.0 (0.0-10.0)
Comorbidities		
Hypertension	9 (53%)	7 (41%)
Diabetes	3 (18%)	3 (18%)
Coronary heart disease	3 (18%)	1 (6%)
Chronic lung disease	1 (6%)	1 (6%)
Liver disease	0	0
Haematological malignancy	1 (6%)	1 (6%)
Solid malignant tumour	0	0
Immunosuppressive therapy	1 (6%)	0
Active smoker	2 (12%)	1 (6%)
Any comorbidity	12 (71%)	10 (59%)
Pre-ECMO treatment		
Time from hospital admission to ECMO, days	6.0 (0.0-13.0)	8.0 (0.0-14.0)
Time from intensive care unit admission to ECMO, days	5.0 (0.0-13.0)	6.0 (0.0-14.0)
Duration of mechanical ventilation	6.0 (0.0-13.0)	5.0 (0.0-14.0)

Cytokines

Lancet Respir Med. 2021 Jul; 9(7): 755–762.

Published

Cytokine

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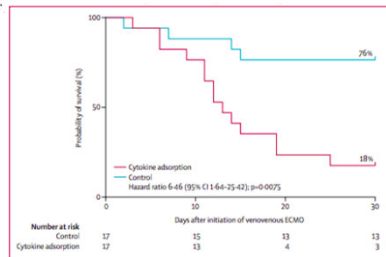
	Cytokine adsorption group (n=17)	Control group (n=17)	P value
Primary endpoint			
Serum interleukin-6 after 72 h	98.6 (71.0 to 192.8)*	112.0 (48.7 to 198.5)*	0.541
Other endpoints			
30-day survival	1 (6%)	1 (6%)	0.00164
Discharged from intensive care unit until day 30	0	3 (18%)	0.231
Serum lactate after 72 h, mmol/L	1.35 (1.05-1.58)*	1.25 (0.93-1.85)*	0.805
Willebrand factor antigen after 72 h, %	40.6 (19.6-101.0)*	31.5 (18.7 to 40.8)*	0.015
D-dimers after 72 h, mg/L FEU	8.77 (3.90 to 35.19)*	15.2 (5.79 to 34.23)*	0.485
SOFA score after 24 h	7.0 (6.0 to 9.5)	8.0 (6.0 to 10.0)	0.595
SOFA score after 48 h	8.0 (6.0 to 9.5)	8.0 (6.0 to 10.0)	0.955
SOFA score after 72 h	7.5 (6.0 to 10.8)*	8.5 (6.0 to 10.0)*	0.815
Norepinephrine support at 72 h, µg/kg per min	0.07 (0.03 to 0.13)*	0.00 (0.00 to 0.10)*	0.045
Cumulative fluid balance for 72 h after initiation of ECMO, mL	2665.0 (563.5 to 5152.0)	2145.0 (192.5 to 3002.0)	0.295
Fluid substitution during the first 72 h after implementation of venovenous ECMO, mL	11 773 (8955 to 13 458)	8344 (7304 to 10 866)	0.00685

Cytokines

Lancet Respir Med. 2021 Jul; 9(7): 755-762.
Published online 2021 May 14. doi: 10.1016/S2213-2600(21)00177-6
PMCID: PMC8121541
PMID: 34000296

Cytokine adsorption in patients with severe COVID-19 pneumonia requiring extracorporeal membrane oxygenation (CYCOV): a single centre, open-label, randomised, controlled trial

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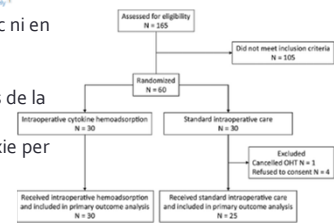


PLoS Med. 2022 Dec; 18(12): e1004262. Online ahead of print.

Use of intraoperative haemoadsorption in patients undergoing heart transplantation: a proof-of-concept randomized trial

Andra Nemeth ^{1,2}, Adam Szilvesz ^{1,2}, Endre Kovacs ^{1,2}, Zsolt Székely-Tóth ¹, Eszter Tamasik ^{1,2}, Hajna Katona ^{1,2}, Kriszta Rácz ^{1,2}, Gergely Csikós ^{1,2}, Viktor Beneszy ^{1,2}, Szabolcs Fehér ^{1,2}, Zoltanna Ulfölsz ^{1,2}, Csilla Tamas ¹, Beata Nagy ¹, Martina Varga ¹, Beáta Merkely ¹

-Greffes « simple » (ni en choc ni en aigue ni redox)
-Cyrosorb VS standard, uniquement durant le temps de la CEC au bloc
-majoration anibioprofylaxie per op



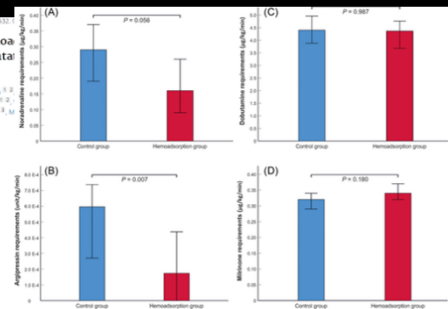
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- Critère de jugement principal: vasoactive-inotropic score (VIS)
- Patients in the haemoadsorption group had significantly lower VIS than patients in the control group during the first post-operative 24 h (median VIS: 27.2 [14.6–47.7] vs. 41.9 [22.4–63.2], $P = 0.046$, respectively)

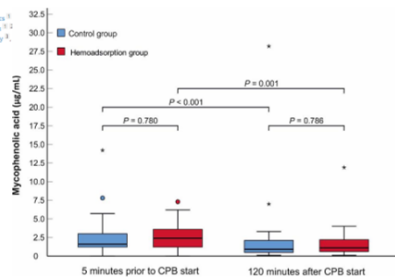
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Table 3 Comparative analysis of secondary outcome parameters

Parameters	Control group (N = 25)	Haemoadsorption group (N = 30)	P-value
ECMO, n	3 (12.0%)	0 (0%)	0.08
ECMO bleeding, mL	570 [385–1305]	565 [350–1130]	0.54
ECMO for bleeding, n	2 (8.0%)	0 (0%)	0.20
ECMO CPB 24 h, unit	4.0 [0–5.5]	2.0 [0–4.0]	0.24
ECMO CPB 24 h, unit	2.0 [0–3.0]	2.0 [0–3.0]	0.57
ECMO CPB 24 h, unit	12.0 [0–16.0]	12.0 [8.0–16.0]	0.59
ECMO CPB 24 h, unit	65 [23–287]	25 [19–58.8]	0.02
ECMO CPB 24 h, unit	15 (60.0%)	9 (30.0%)	0.02
ECMO CPB 24 h, unit	0 (0%)	1 (3.3%)	1.00
ECMO CPB 24 h, unit	4 (16.0%)	1 (3.3%)	0.10
ECMO CPB 24 h, unit	19 (76.0%)	11 (36.7%)	0.00
ECMO CPB 24 h, unit	4 (16.0%)	0 (0%)	0.03
ECMO CPB 24 h, unit	72.1 [11.2–191.4]	2.5 [–24.6–71.1]	0.00
ECMO CPB 24 h, unit	1 (4.0%)	0 (0%)	0.45
ECMO CPB 24 h, unit	12 [8.5–18.0]	8.5 [8.0–10.3]	0.02
ECMO CPB 24 h, unit	28 [24–38.5]	25 [22–34.3]	0.23
ECMO CPB 24 h, unit	2 (8.0%)	0 (0%)	0.20
ECMO CPB 24 h, unit	0 (0%)	0 (0%)	1.00
ECMO CPB 24 h, unit	5 (20.0%)	5 (16.7%)	1.00
ECMO CPB 24 h, unit	6 (24.0%)	10 (33.3%)	0.44
ECMO CPB 24 h, unit	1 (4.0%)	0 (0%)	0.45
ECMO CPB 24 h, unit	1 (4.0%)	2 (6.7%)	1.00
ECMO CPB 24 h, unit	1 (4.0%)	3 (10.0%)	0.61
ECMO CPB 24 h, unit	2 (8.0%)	1 (3.3%)	0.58

ECMO are presented as number of patients (frequency) and median [interquartile range]. N = 55. Of 36 registered cellular rejections, 5 (13.9%) were confirmed as grade 1R and 1 (2.8%) was confirmed as grade 2R (ISHLT 2005). The registered antibody-mediated rejection were confirmed as pAMR 11 (ISHLT 2013).

ECMO, extracorporeal membrane oxygenation; ECMO, endomyocardial bypass; FFP, fresh frozen plasma; ICU, intensive care unit; MV, mechanical ventilation; pAMR, pathologic antibody-mediated rejection; PLT, platelet transfusion; PRC, packed red cell; RRT, renal replacement therapy.

ECMO CPB 24 h, unit was classified according to Kidney Disease Improving Global Outcomes creatinine-based definition criteria over the

Adsorption médicament

Ann Thorac Surg. 2019 Jul;108(1):45-51. doi: 10.1016/j.athoracsu.2018.12.032. Epub 2019 Jan 23.

Cytosorb Adsorption During Emergency Cardiac Operations in Patients at High Risk of Bleeding.
Hassan S¹, Kannanachari R², Wohlmuth P³, Buddie U⁴, Schnepf M⁵, Geisel S⁶

55 patients avec CEC urgente traités par Ticagrelor ou Rivaroxaban
39 patients prospectifs traités par cytosorb VS 16 contrôles rétrospectifs

Variables	CA Group		WA Group		P Value
	Ticagrelor (n = 32)	Rivaroxaban (n = 7)	Ticagrelor (n = 11)	Rivaroxaban (n = 5)	
ECMO-treated	21 (65.6)	4 (57.1)	11 (100)	5 (100)	0.4780
ECMO rate	0 (0)	0 (0)	4 (36.4)	2 (40)	0.0003
ECMO volume in 24 hours	350 (300–450)	390 (310–430)	890 (630–1,025)	600 (590–1,000)	0.0037
ECMO care, days	2 (1–3)	2 (2–3)	3 (2–4)	6 (5–6)	0.0141
Total length of stay, days	11 (9–12)	11 (10–13)	14 (10–16)	18 (18–20)	0.0244

Saignement, transfusion et reprises plus importantes dans groupe controle

Adsorption médicament

> Am Heart J. 2021 Nov 1;245:19–28. doi: 10.1016/j.ahj.2021.10.188. Online ahead of print.

Rationale and design of the safe and timely antithrombotic removal – ticagrelor (STAR-T) trial: A prospective, multi-center, double-blind, randomized controlled trial evaluating reductions in postoperative bleeding with intraoperative removal of ticagrelor by the drugsorb™-ATR device in patients undergoing cardiothoracic surgery within 48 hours from last ticagrelor dose

C Michael Gibson¹, Michael J Mack², Victoria T Lee³, David J Schneider⁴, Frank W Selke⁵, E Magnus Ohman⁶, Vinod H Thourani⁷, Gheorghe Doros⁸, Hans Kroger³, Donald E Cutlip⁹, Efthymios N Deliargyris³

Adsorption antibiotiques

Ann Intensive Care. 2022; 12: 44. PMID: PMC9124739
Published online 2022 May 23. doi: 10.1186/s13613-022-01017-5
PMID: 35599248

Does the cytokine adsorber CytoSorb® reduce vancomycin exposure in critically ill patients with sepsis or septic shock? a prospective observational study

Christina Scharf¹, Ferdinand Weinelt^{2,3}, Ines Schroeder¹, Michael Paal⁴, Michael Weigand⁴, Michael Zoller¹, Michael Irbeek¹, Charlotte Klotz², Josef Briegel¹, and Uwe Liebchen^{2,1,2}

Conclusion

The use of CytoSorb® leads to a clinically significant adsorption of vancomycin (max. 572 mg) in critically ill patients with sepsis or septic shock. We recommend the administration of an additional dose of 500 mg vancomycin over 2 h to avoid subtherapeutic vancomycin exposure.

Conclusion hémoadsorption

- Technique récente
- Littérature encore faible pour bénéfice lié à l'immunomodulation, en chirurgie cardiaque et chez patients « médicaux », études randomisées discordantes
- Potentiellement intéressant sur adsorption ticagrelor et rivaroxaban (apixaban?)
- Adsorption de médicaments utiles?

Ventilation per CEC

Cours à venir

Insufflation de CO₂

Rationnel

-Remplacer l'air ambiant dans la cavité thoracique par du CO₂, +lourd + soluble dans le sang et + rapidement absorbé par les tissus.

-Vise à minimiser la présence de bulles d'air dans le sang susceptibles de provoquer des complications de type embolie gazeuse

Rationnel

J Thorac Cardiovasc Surg. 2006 Nov;132(5):1119-25.

Short-term changes in cerebral activity in on-pump and off-pump cardiac surgery defined by functional magnetic resonance imaging and their relationship to microembolization.

Abu-Omar Y¹, Cader S, Guentert Wolf L, Pigott D, Matthews PM, Teagart DP.

- 25 patient avec PAC CEC et 25 cœurs battant
- Association entre quantité d'embolies gazeux (doppler TC) et activation préfrontale à l'IRM fonctionnel à 1 mois

Rationnel

Circulation. 2004 Mar 9;109(9):1127-32. Epub 2004 Feb 23.

Effect of CO₂ insufflation on the number and randomized clinical trial.

Svenaeus P¹, Persson M, van der Linden J.

10 patients CO₂ VS 10 contrôles, détection micrebols par ETO ou déclampage (lues en aveugle)

TABLE 2. No. of Microemboli According to Transesophageal Echocardiographic Evaluation of the Left Atrium and Ventricle and the Proximal Part of the Ascending Aorta

	No. of Microemboli			
Study Period/Phase of Interest	Group Control (n=10)	Group CO ₂ (n=10)	P	
From release of cross clamp until 30 minutes after end of CPB				
LA	340 (200/300)	40 (20/10)	<0.01	
LV	204 (170/30)	40 (20/10)	<0.01	
Ao	194 (150/30)	30 (10/20)	<0.01	
LA+LV+Ao	738 (520/218)	110 (70/40)	<0.01	
From 10 minutes after release of cross clamp				
LA	224 (130/90)	30 (10/20)	<0.01	
LV	131 (110/20)	40 (20/10)	<0.01	
Ao	81 (70/10)	20 (10/10)	<0.01	
LA+LV+Ao	436 (310/126)	100 (70/30)	<0.01	
Last 10 minutes of CPB				
LA	70 (30/40)	17 (10/7)	<0.01	
LV	30 (20/10)	21 (10/10)	<0.01	
Ao	47 (30/17)	10 (10/0)	<0.01	
LA+LV+Ao	147 (100/47)	48 (30/18)	<0.01	
From 20 minutes after end of CPB				
LA	94 (60/30)	9 (0/9)	<0.01	
LV	70 (40/30)	10 (0/10)	0.01	
Ao	56 (30/20)	10 (0/10)	<0.01	
LA+LV+Ao	220 (130/90)	30 (0/30)	<0.01	

Values are given as median (25th/75th percentile). LA indicates left atrium; LV, left ventricle; and Ao, aorta.

Values are given as median (25th/75th percentile). LA indicates left atrium; LV, left ventricle; and Ao, aorta.

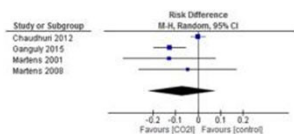
Littérature

J Thorac Cardiovasc Surg. 2017 Autumn;29(3):301-310. doi: 10.1053/j.semvs.2017.05.002. Epub 2017 May 23.

Carbon Dioxide Insufflation During Cardiac Surgery: A Meta-analysis of Randomized Controlled Trials.

Benedetto U¹, Caruso M², Guida G², Buccarelli-Ducci G², Thali R², Bryan A², Angeles G².

- 8 études à la méthodologie différentes+++
- Pas de différence sur le déclin cognitif



Direct visualization of carbon dioxide field flooding: Optical and concentration level comparison of diffuser effectiveness

Stijn Vandenberghe, PhD, David Iseli, BSc, and Stefanos Demertzis, MD

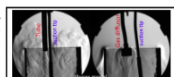
ABSTRACT

Objective: Carbon dioxide field flooding during open-heart surgery is intended to avoid blood-air contact, bubble formation, and embolism, and therefore potential neurologic and other ischemic complications. The inert gas is invisible, and thus its use and effectiveness are heavily debated. We intended to provide better insight in the behavior of the gas via direct concentration measurements and visualization of the gas cloud.

Methods: A transparent rectangular model of the open thorax was created, foreseen with carbon dioxide concentration sensors in 2 locations (atrial and aortic incisions), and placed in an optical test bench that amplifies the diffraction gradients. Six different commonly used carbon dioxide diffusers (3 commercial, 3 improvised) were tested with different flow rates of gas delivery (1, 4, 7, 10 standard liter per minute (SLPM)) and combined with the application of suction.

Results: The imaging reveals that commercially available diffusers generally create less turbulent flow than improvised diffusers, which is supported by the concentration measurements where improvised diffusers cannot generate a 100% carbon dioxide atmosphere at the aortic incision location. The atrial incision is easier to protect: 0% air with all commercial devices for all flow rates greater than 1 SLPM. A flow rate of 1 SLPM does not create an inert atmosphere with any device.

Conclusions: The optically observed carbon dioxide atmosphere is unstable and influenced by many factors. The device used for diffusion and the flow rate are important determinants of the maximum gas concentration that can be achieved, as is the location where this is measured. (J Thorac Cardiovasc Surg 2020;159:958-68)



CO₂ flow at 10 SLPM from an improvised delivery tube (L) versus a commercial diffuser (R).

Central Message

CO₂ field flooding is not well understood. We provided insight in diffuser effectiveness via visualization and concentration measurements and demonstrated an unstable CO₂ atmosphere.

Perspective

Many misunderstandings exist about CO₂ field flooding, mainly because the gas cannot be observed. This engineering study literally reveals the gas behavior, and the observations are corroborated by instantaneous concentration measurements. The reported findings should be a study of interest to cardiac surgeons, who can easily improve central protection of their next patient.

See Commentaries on pages 969 and 970.

Littérature

[Randomized Controlled Trial](#) > [J Cardiovasc Surg \(Torino\)](#). 2022 Jun;63(3):369-375.
doi: 10.23736/S0021-9509.22.12004-5. Epub 2022 Mar 28.

Warm humidified CO₂ insufflation improves pericardial integrity for cardiac surgery: a randomized control study

Reny Segal^{1,2}, Paul M Mezzavia¹, Roni B Krieser¹, Shienny Sampurno³, Michael Taylor³, Robert Ramsay^{2,3}, Michael Kluger¹, Keat Lee^{1,2}, Francis L Loh¹, James Tatoulis^{1,2}, Michael O'Keefe^{1,2}, Yinwei Chen¹, Teresa Sindoni¹, Irene Ng^{4,2}

Conclusions: Humidified warm CO₂ insufflation significantly reduced pericardial epithelial damage when compared to dry cold CO₂ insufflation in open-chamber cardiac surgery. Further studies are warranted to investigate its potential clinical benefits.

Littérature

[BMJ Open](#). 2023; 13(5): e074221.

Published online 2023 May 17. doi: [10.1136/bmjopen-2023-074221](#)
Protocol

PMCID: PMC10193051
PMD: 37197819

Efficacy and safety of carbon dioxide insufflation for brain protection for patients undergoing planned left-sided open heart valve surgery: protocol for a multicentre, placebo-controlled, blinded, randomised controlled trial (the CO₂ Study)

Bachel Todd^{2,1}, Chris A Rogers², Maria Pufulete², Lucy Culliford², Pieter Protorius³, Natalie Voets³, Enoch Akowuah⁴, Rana Sayeed⁵, Michelle Lazaroo², Surinder Kaur¹, Gianni D Angelini^{6,7} and Ben Gibbison^{6,7}

Conclusion insufflation CO₂

- couramment utilisée pour réduire le risque d'embolie gazeuse
- preuves de son efficacité en termes de protection neurologique restent limitées.
- nécessite une adaptation du balayage (pendant et après)
- perturbe l'utilisation du CO₂ gap per CEC

Merci !