



Everything you need to know about deresuscitation

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Why deresuscitation?

While fluid administration is one of the most common therapeutic interventions in critical care, it carries high potential for harm as overzealous fluid resuscitation may lead to glycocalix degradation and endothelial injury. Fluids should be regarded as drugs with specific indications, contraindications, and potential adverse effects. The renewed concept of “fluid stewardship” [1], analogous to antibiotic stewardship, focusses on the 4 D’s (drug, dose, duration, and de-escalation), the 4 questions (when to start and when to stop fluid therapy, and when to start and when to stop fluid removal), the 4 indications (resuscitation, maintenance, replacement and nutrition), and the conceptual ROSE model describing 4 fluid phases (resuscitation, optimization, stabilization and evacuation, Fig. 1, Panel A) [2].

What is deresuscitation?

The term deresuscitation was coined in 2014 [3] and defined as active fluid removal in patients with fluid overload using drugs and/or ultrafiltration (UF). The term ‘de-escalation’ relates more to reducing the amount and rate of previously started fluid administration [4]. There is no clear or unique definition of fluid overload in the literature and some suggest that it is an all or nothing phenomenon associated with worse outcomes [3, 5], defined as 10% body weight fluid accumulation (dividing the cumulative fluid balance in liters by the patient’s baseline body weight and multiplying by 100). We suggest to avoid the term fluid overload as this may misleadingly refer

to intravascular hypervolemia [6] and propose the term fluid accumulation (FA) instead. FA denotes a pathologic state of overhydration associated with clinical impact and worse outcomes which may vary depending on age, comorbidity and phase of illness. It can be estimated by different methods: change in daily body weight, cumulative fluid balance, or bio-electrical impedance analysis with body composition. It may occur with concomitant intravascular hypovolemia, normovolemia and hypervolemia and may or may not be associated with interstitial edema. Fluid accumulation describes a continuum and no specific threshold of fluid balance alone can define fluid accumulation across all individuals. Fluid accumulation syndrome (FAS) describes the presence of any % of fluid accumulation or fluid overload with negative impact on end-organ function (Fig. 1, Panel B) which may or may not be associated with global increased permeability syndrome (GIPS) [7].

When to start deresuscitation?

As soon as salvage resuscitation has been completed, de-escalation should begin, aiming for a zero fluid balance (passively, by decreasing fluid intake). Deresuscitation should be considered when FA negatively impacts organ function. Close monitoring is key to identifying patients who benefit from fluid removal. Triggers include clinical signs (increased weight, positive fluid balance), laboratory parameters (hemodilution), radiological signs (B-lines, pleural effusion, low inferior vena cava collapsibility index) [8] altered cardiopulmonary function (absence of fluid responsiveness, increased filling pressures and volumetric preload, low pulse pressure variation) and measures of end-organ function.

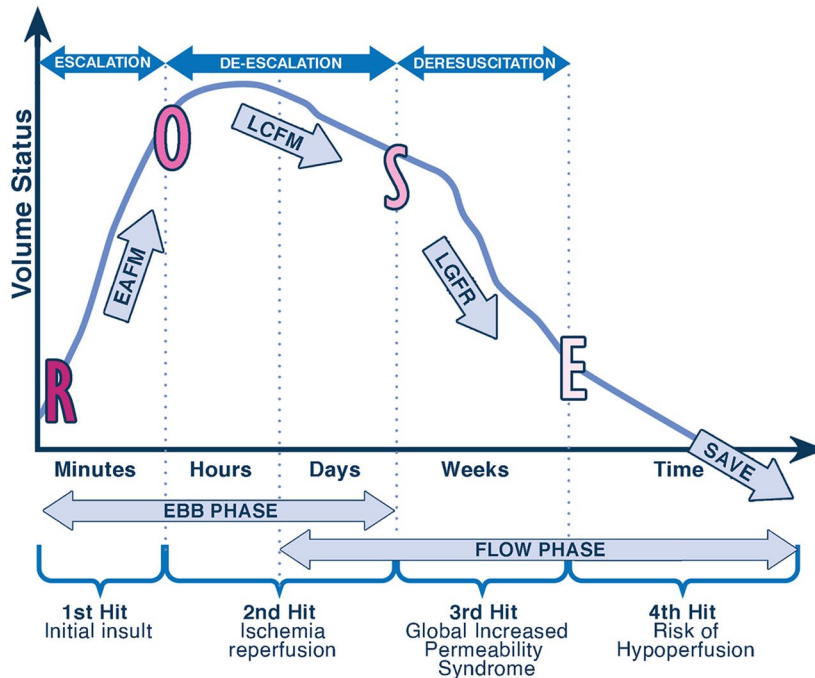
Potentially promising techniques exist that may serve as triggers and safety limits for deresuscitation. In critically ill patients receiving renal replacement therapy (RRT), the absence of preload dependence pre-RRT, as

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A



R

Life saving **Resuscitation** phase with focus on **patient rescue** and early adequate fluid management (EAFM), eg 30ml/kg/1hr according to SSCG or a fluid challenge/bolus of 4ml/kg given in 5-10 minutes.

Triggers to start IV fluids: shock

- MAP < 65mmHg
- GEDVI < 640ml/m²
- (RVEDVI < 80ml/m²)#
- (CVP < 8mmHg)*
- (PAOP < 10mmHg)*
- CI < 2,5L/min/m²
- PPV or SVV > 12-15%
- PLR test positive
- Lactate > 3mmol/L (shock)
- IVCCI > 50%

O

Optimization phase with focus on **organ rescue** (maintenance) and avoiding fluid overload (fluid creep). Aiming for neutral fluid balance.

Triggers to stop IV fluids: unresponsiveness

- MAP/APP > 65/55mmHg
- GEDVI 640-800ml/m²
- CI > 2,5 L/min/m²
- PPV or SVV < 12%
- PLR test negative
- PPV or SVV > 12-15%
- Normal lactate < 2mmol/L
- LVEDAI 8-12cm²/m²
- IAP < 15mmHg

S

Stabilization phase with focus on **organ support** (homeostasis). Late conservative fluid management (LCFM) is defined as two consecutive negative FB within 1st week after initial insult.

Triggers to start fluid removal: FAS/GIPS

- MAP/APP > 65/55mmHg
- GEDVI > 850ml/m²
- EVLWI > 10-12ml/kg PBW
- PVPI > 3 and PF ratio < 150
- PPV or SVV < 12%
- PLR test negative
- LVEDAI > 14cm²/m², high VExUS score
- IAP > 12-15mmHg
- COP < 16-18mmHg; CLI > 60
- BIA: ECW/ICW > 1; V_e > 5%

E

Evacuation phase with focus on **organ recovery** and resolving fluid overload (in case of no flow state) with active late goal directed fluid removal (LGFR) and negative FB.

Triggers to stop fluid removal: hypoperfusion

- MAP/APP < 55/45mmHg**
- PPV or SVV > 15%
- PLR test positive
- Lactate > 2,5mmol/L
- S_vO₂ < 70-75%
- S_{cv}O₂ < 65-70%
- ICG PDR < 14-16%

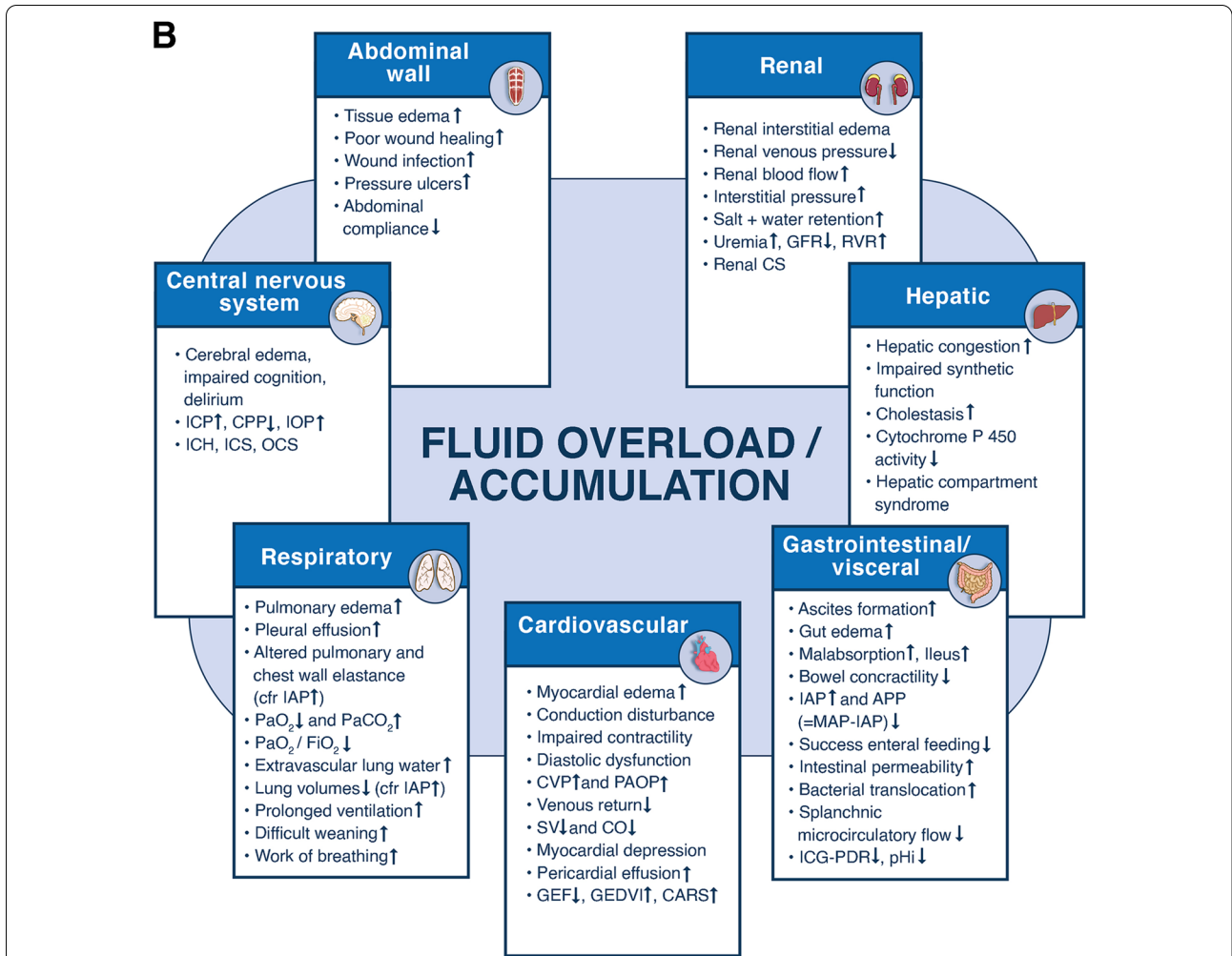


Fig. 1 The 4 phases conceptual model and deleterious effects of fluid accumulation syndrome. Panel **A** The four-hit model of shock with evolution of patients' cumulative fluid volume status over time during the five phases of resuscitation: Resuscitation (R), Optimization (O), Stabilization (S), and Evacuation (E) (ROSE), followed by a possible risk of Hypoperfusion in case of too aggressive deresuscitation. On admission patients are often hypovolemic, followed by normovolemia after fluid resuscitation (escalation or EAFM, early adequate fluid management), and possible fluid overload, again followed by a phase returning to normovolemia with de-escalation via achieving zero fluid balance or late conservative fluid management (LCFM) and followed by late goal directed fluid removal (LGFR) or deresuscitation. In case of hypovolemia, O₂ cannot get into the tissue because of convective problems, in case of hypervolemia O₂ cannot get into the tissue because of diffusion problems related to interstitial and pulmonary edema, gut edema (ileus and abdominal hypertension). *Volumetric preload indicators like GEDVI, LVEDAI, or RVEDVI are preferred over barometric ones like CVP or PAOP. **Vasopressor can be started or increased to maintain MAP/APP above 55/45 during deresuscitation phase. #Can only be measured via Swan-Ganz pulmonary artery catheter (PAC) and became obsolete. Panel **B** Potential consequences of fluid accumulation syndrome (formerly known as fluid overload) and GIPS (global increased permeability syndrome) on end-organ function. Figures adapted from Malbrain et al. with permission, according to the Open Access CC BY Licence 4.0 [2, 7]. ACS abdominal compartment syndrome, APP abdominal perfusion pressure = MAP-IAP, BIA bio-electrical impedance analysis, CARs cardio-abdominal-renal syndrome, CI cardiac index, CLI capillary leak index (serum CRP divided by serum albumin), CO cardiac output, COP colloid oncotic pressure, CPP cerebral perfusion pressure, CS compartment syndrome, CVP central venous pressure, ECW/ICW extracellular/intracellular water, EVLWI extra-vascular lung water index, FAS fluid accumulation syndrome, GEDVI global end-diastolic volume index, GEF global ejection fraction, GFR glomerular filtration rate, IAH intra-abdominal hypertension, IAP intra-abdominal pressure, ICG-PDR indocyanine green plasma disappearance rate, ICH intracranial hypertension, ICP intracranial pressure, ICS intracranial compartment syndrome, IOP intra-ocular pressure, IVCCI inferior vena cava collapsibility index, LVEDAI left ventricular end-diastolic area index, MAP mean arterial pressure, OCS ocular compartment syndrome, PAOP pulmonary artery occlusion pressure, PF P_aO₂ over FiO₂ ratio, pH_i gastric tonometry, PLR passive leg raising, PPV pulse pressure variation, PVPI pulmonary vascular permeability index, RVEDVI right ventricular end-diastolic volume index, RVR renal vascular resistance, S_{cv}O₂ central venous oxygen saturation, SSCG surviving sepsis campaign guidelines, S_vO₂ mixed venous oxygen saturation, SV stroke volume, SVV stroke volume variation, V_E volume excess (from baseline body weight), VExUS venous congestion by ultrasound

determined by a negative passive leg raising (PLR) test, predicted tolerance of fluid removal [9]. In 123 critically ill patients, increased extra-vascular lung water index (EVLWI) and failure to lower EVLWI were associated with increased mortality and longer duration of ventilation [10]. Use of a protocol including blood volume analysis with calculation of total circulating blood volume by albumin-marked isotope dilution technique resulted in a 66% reduction in mortality, 20% reduction in length of stay (LOS), 36 h earlier treatment decisions and change of treatment strategy in 44% (reduced fluid and diuretic administration in 50% of cases) [11]. A 5% increase in volume excess from baseline, measured by bio-electrical impedance analysis, was associated with increased mortality in a retrospective study of 101 critically ill patients [12]. Another study in 125 patients showed that the hydration status measured by bio-electrical impedance vector analysis (BIVA) seems to predict mortality risk in intensive care unit (ICU) patients better than the conventional method of fluid balance recording [13]. Recently, the venous congestion by point-of-care-ultrasound (VExUS) score gained attention as a technique to assess fluid status at the bedside [14]. Venous congestion as a contributor to organ failure can also be demonstrated by the impact of right ventricular dysfunction on inferior vena cava, portal, hepatic, and renal veins using echocardiography [15].

How to deresuscitate?

In principle, parameters achieved during the stabilization phase should be maintained during deresuscitation. Prevention (fluid restriction) and de-escalation come first and clinicians should be aware that most fluids given are not given for resuscitation purposes (only 6%) but rather for maintenance (25%), nutrition (33%) and drug administration (fluid creep) (33%) [16]. Measures to remove excess fluid can be pharmacologic (drugs) and non-pharmacologic (net UF), combined with fluid restriction. Provided some kidney function is preserved, diuretics are usually tried first, either as monotherapy or in combination. The options are summarized in supplementary Table 1 and include loop diuretics (furosemide, bumetanide), carbonic anhydrase inhibitor (acetazolamide), mineralocorticoid receptor antagonist (spironolactone), thiazides or thiazide-like drugs (indapamide). In case of low serum albumin levels (<30 g/L) or low serum total protein levels (<60 g/L), hyper-oncotic albumin 20–25% can be added for synergistic effect [17, 18]. The combination of positive end-expiratory pressure (PEEP, in cmH₂O, set at the level of intra-abdominal pressure (IAP) in mmHg), followed by hyper-oncotic albumin (up to albumin levels of 30 g/L) and furosemide resulted in a negative cumulative fluid balance, a reduction of EVLWI

and IAP, and improved clinical outcomes in a matched cohort of 114 critically ill patients with hyperpermeability pulmonary edema [19]. However, caution is necessary when using hyper-oncotic human albumin solution to facilitate restrictive fluid therapy. Current evidence stems mostly from observational studies, and more randomized trials are needed to better establish a personalized approach to fluid management [20].

Diuretic failure is defined as inability to generate adequate diuresis to achieve the target negative fluid balance (diuretic resistance) or as the occurrence of important side effects. Diuretic resistance is primarily based on variable gastrointestinal absorption, poor kidney function or interactions with other drugs. Mechanical fluid removal should be considered when fluid overload has not responded to diuretics, cannot be corrected safely or in situations where diuretics are unlikely to be effective. Different types of mechanical fluid removal can be utilized for deresuscitation, including slow continuous ultrafiltration (SCUF) and intermittent or continuous RRT. The primary modality for fluid removal in all techniques is UF. In 11 critically ill patients receiving RRT, 1.9 L net fluid removal lowered IAP, EVLWI and global end-diastolic volume index (GEDVI) significantly [21].

Although excess fluid should be removed without hemodynamic compromise, occasionally it may be necessary to accept an increase in catecholamine support to facilitate fluid removal and improve critical physiological parameters such as oxygenation.

When to stop deresuscitation?

Deresuscitation should be stopped when the goal is met (i.e. benefit has been achieved) or when safety concerns arise. This goal can be fluid-related (neutral balance), physiologic (e.g. central venous pressure, CVP) or clinical (improved oxygenation, extubation). However, there is no one single valid marker that indicates when euvolaemia has been achieved.

During UF, hypotension may occur when the rate of removal of plasma exceeds the refilling capacity of compensatory fluid movement from the extra-vascular compartments. In stable patients undergoing intermittent haemodialysis, the fluid refill rate is 2 to 6 ml/kg per hour but may exceed 10 ml/kg per hour at high rates of ultrafiltration [22]. Since transcapillary refill rate depends on oncotic pressure, vascular integrity and blood pressure, it is reduced during critical illness [23]. A recent study showed that in critically ill adults receiving continuous veno-venous hemodiafiltration for acute kidney injury, the rate of net UF at which mortality is increased seemed to be from 1.75 ml/kg/h and certainly from 2.8 ml/kg/h upwards [24]. The decision to change from mechanical fluid removal to diuretics or vice versa should be guided

by clinical volume status, organ function, spontaneous diuresis and perceived trajectories.

So what is next?

Research is required to investigate strategies to maintain and improve tissue perfusion and organ function while mitigating FAS. This includes the need to identify sensitive and accurate markers of intravascular volume status. Further research should also identify the optimal rates of fluid removal in specific patient groups, tools to predict the consequences of fluid removal, and investigate the role of adjunctive therapies (e.g. albumin) and vasopressors to support fluid removal versus the harm from prolonged fluid accumulation. Aggregation and integration of large-scale (realworld) physiological and clinical data will lend itself to data science techniques such as artificial intelligence and machine learning that can in real-time accurately measure FAS and account for patient characteristics such as illness severity and chronic co-morbidities to identify the optimal method, timing and rate of fluid removal for individual patients.

Wrap it up—take home messages

One of the most common interventions in critically ill patients is assessment of fluid status and prescription of fluid administration or removal. Both interventions have potential for harm and benefit, thus they should be regarded as drugs and managed according to the principles of fluid stewardship [25]. While stewardship and deresuscitation are important, we must avoid the pendulum swinging back to lead hypovolemic patients to inappropriately receive diuretics (as was the case during the first wave of the coronavirus disease 2019 (COVID-19) pandemic). Deresuscitation should never be too fast, too long or too aggressive. Deciding when to start and stop deresuscitation is key to improving patient outcomes, and research is ongoing to identify the best parameters to guide fluid removal in critically ill patients [26, 27]. While waiting for the results of the RADAR-2 and CLAS-SIC trials, prevention of FA and de-escalation of fluid therapy remain the most effective strategies to avoid deresuscitation.

Supplementary Information

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Declarations

Conflicts of interest

MLNGM is co-founder, past-President and current Treasurer of WSACS (The Abdominal Compartment Society, <http://www.wsacs.org>). He is member of the medical advisory Board of Pulsion Medical Systems (part of Getinge group), Serenno Medical, Potrero Medical, and Baxter, and consults for BBraun, Becton Dickinson, ConvaTec, Spiegelberg, and Holtech Medical. He is co-founder and President of the International Fluid Academy (IFA). The IFA (<http://www.fluidacademy.org>) is integrated within the not-for-profit charitable organization iMERIT, International Medical Education and Research Initiative, under Belgian law. MO has received speaker honoraria and research funding from Fresenius Medical and Baxter and is a member of an advisory board for NxStage and Pfizer. GM has served on research or medical advisory boards or conducted funded research from Genentech, Grifols, Regeneron and Siemens.

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