

The unrestricted global effort to complete the COOL trial

Kirkpatrick, Andrew W.; Coccolini, Federico; Tolonen, Matti; Minor, Samuel; Catena, Fausto; Gois, Emanuel; Doig, Christopher J.; Hill, Michael D.; Ansaloni, Luca; Chiarugi, Massimo; ...

Source / Izvornik: **World Journal of Emergency Surgery, 2023, 18**

Journal article, Published version

Rad u časopisu, Objavljena verzija rada (izdavačev PDF)

<https://doi.org/10.1186/s13017-023-00500-z>

Permanent link / Trajna poveznica: <https://urn.nsk.hr/urn:nbn:hr:105:105136>

Rights / Prava: [Attribution 4.0 International](#)/[Imenovanje 4.0 međunarodna](#)

Download date / Datum preuzimanja: **2024-11-29**



Repository / Repozitorij:

[Dr Med - University of Zagreb School of Medicine
Digital Repository](#)



RESEARCH

Open Access



The unrestricted global effort to complete the COOL trial

Andrew W. Kirkpatrick^{1*}, Federico Coccolini², Matti Tolonen³, Samuel Minor⁴, Fausto Catena⁵, Emanuel Gois Jr.⁶, Christopher J. Doig⁷, Michael D. Hill⁸, Luca Ansaloni⁹, Massimo Chiarugi¹⁰, Dario Tartaglia¹⁰, Orestis Ioannidis¹¹, Michael Sugrue¹², Elif Colak¹³, S. Morad Hameed¹⁴, Hanna Lampela¹⁵, Vanni Agnoletti¹⁶, Jessica L. McKee¹⁷, Naisan Garraway¹⁸, Massimo Sartelli¹⁹, Chad G. Ball²⁰, Neil G. Parry²¹, Kelly Voght²¹, Lisa Julien²², Jenna Kroeker¹⁸, Derek J. Roberts²³, Peter Faris²⁴, Corina Tiruta²⁵, Ernest E. Moore²⁶, Lee Anne Ammons²⁷, Elissavet Anestiadou¹¹, Cino Bendinelli²⁸, Konstantinos Bouliaris²⁹, Rosemarry Carroll²⁸, Marco Ceresoli³⁰, Francesco Favi³¹, Angela Gurrado³², Joao Rezende-Neto³³, Arda Isik³⁴, Camilla Cremonini¹⁰, Silvia Strambi¹⁰, Georgios Koukoulis²⁹, Mario Testini³², Sandy Trpcic³³, Alessandro Pasculli³², Erika Picariello³⁵, Fikri Abu-Zidan³⁶, Ademola Adeyeye³⁷, Goran Augustin³⁸, Felipe Alconchel³⁹, Yuksel Altinel⁴⁰, Luz Adriana Hernandez Amin⁴¹ , José Manuel Aranda-Narváez⁴², Oussama Baraket⁴³, Walter L. Biffi⁴⁴, Gian Luca Baiocchi⁴⁵, Luigi Bonavina⁴⁶, Giuseppe Brisinda⁴⁷, Luca Cardinali⁴⁸, Andrea Celotti⁴⁹, Mohamed Chaouch⁵⁰, Maria Chiarello⁵¹, Gianluca Costa⁵², Nicola de'Angelis⁵³, Nicolo De Manzini⁵⁴, Samir Delibegovic⁵⁵, Salomone Di Saverio⁵⁶, Belinda De Simone^{57,58}, Vincent Dubuisson⁵⁹, Pietro Fransvea⁶⁰, Gianluca Garulli⁶¹, Alessio Giordano⁶², Carlos Gomes⁶³, Firdaus Hayati⁶⁴, Jinjian Huang⁶⁵, Aini Fahriza Ibrahim⁶⁶, Tan Jih Huei⁶⁷, Ruhi Fadzlyana Jailani⁶⁸, Mansoor Khan⁶⁹, Alfonso Palmieri Luna⁷⁰, Manu L. N. G. Malbrain^{71,72}, Sanjay Marwah⁷³, Paul McBeth⁷⁴, Andrei Mihailescu⁷⁵, Alessia Morello⁷⁶, Francesk Mulita⁷⁷, Valentina Murzi⁷⁸, Ahmad Tarmizi Mohammad⁷⁹, Simran Parmar²⁴, Ajay Pak⁸⁰, Michael Pak-Kai Wong⁸¹, Desire Pantalone⁸², Mauro Podda⁸³, Caterina Puccioni⁸⁴, Kemal Rasa⁸⁵, Jianan Ren⁶⁵, Francesco Roscio⁸⁶, Antonio Gonzalez-Sanchez⁴², Gabriele Sganga⁸⁴, Maximilian Scheiterle⁸⁷, Mihail Slavchev⁸⁸, Dmitry Smirnov⁸⁹, Lorenzo Tosi⁹⁰, Anand Trivedi⁹¹, Jaime Andres Gonzalez Vega⁹², Maciej Waledziak⁹³, Sofia Xenaki⁹⁴, Desmond Winter⁹⁵, Xiuwen Wu⁶⁵, Andee Dzulkarnean Zakaria⁹⁶ and Zaidi Zakaria⁹⁶

Abstract

Background Severe complicated intra-abdominal sepsis (SCIAS) has an increasing incidence with mortality rates over 80% in some settings. Mortality typically results from disruption of the gastrointestinal tract, progressive and self-perpetuating bio-mediator generation, systemic inflammation, and multiple organ failure. A further therapeutic option may be open abdomen (OA) management with negative peritoneal pressure therapy (NPPT) to remove inflammatory ascites and attenuate the systemic damage from SCIAS, although there are definite risks of leaving the abdomen open whenever it might possibly be closed. This potential therapeutic paradigm is the rationale being

*Correspondence:

Andrew W. Kirkpatrick

Andrew.kirkpatrick@albertahealthservices.ca

Full list of author information is available at the end of the article



© The Author(s) 2023. **Open Access** This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit <http://creativecommons.org/licenses/by/4.0/>. The Creative Commons Public Domain Dedication waiver (<http://creativecommons.org/publicdomain/zero/1.0/>) applies to the data made available in this article, unless otherwise stated in a credit line to the data.

assessed in the Closed Or Open after Laparotomy (COOL trial) (<https://clinicaltrials.gov/ct2/show/NCT03163095>). Initially, the COOL trial received Industry sponsorship; however, this funding mandated the use of a specific trade-marked and expensive NPPT device in half of the patients allocated to the intervention (open) arm. In August 2022, the 3 M/Acelity Corporation without consultation but within the terms of the contract canceled the financial support of the trial. Although creating financial difficulty, there is now no restriction on specific NPPT devices and removing a cost-prohibitive intervention creates an opportunity to expand the COOL trial to a truly global basis. This document describes the evolution of the COOL trial, with a focus on future opportunities for global growth of the study.

Methods The COOL trial is the largest prospective randomized controlled trial examining the random allocation of SCIAS patients intra-operatively to either formal closure of the fascia or the use of the OA with an application of an NPPT dressing. Patients are eligible if they have free uncontained intraperitoneal contamination and physiologic derangements exemplified by septic shock OR severely adverse predicted clinical outcomes. The primary outcome is intended to definitively inform global practice by conclusively evaluating 90-day survival. Initial recruitment has been lower than hoped but satisfactory, and the COOL steering committee and trial investigators intend with increased global support to continue enrollment until recruitment ensures a definitive answer.

Discussion OA is mandated in many cases of SCIAS such as the risk of abdominal compartment syndrome associated with closure, or a planned second look as for example part of “damage control”; however, improved source control (locally and systemically) is the most uncertain indication for an OA. The COOL trial seeks to expand potential sites and proceed with the evaluation of NPPT agnostic to device, to properly examine the hypothesis that this treatment attenuates systemic damage and improves survival. This approach will not affect internal validity and should improve the external validity of any observed results of the intervention.

Trial registration: National Institutes of Health (<https://clinicaltrials.gov/ct2/show/NCT03163095>).

Keywords Intraperitoneal sepsis, Septic shock, Peritonitis, Open abdomen, Multiple organ dysfunction, Laparotomy, Randomized controlled trial, Global health

Background

Sepsis is an increasing cause of death worldwide [2, 3], with an incidence estimated between 18 and 31 million cases worldwide per year [3–7]. Sepsis mortality approaches 30–40% when a shock is present [8–10], and may be higher in the developing world [2]. The incidence and mortality of sepsis can be compared to other critical global health problems such as COVID-19 with 6.5 million deaths worldwide over more than 2 years [11], or 4.4 million deaths from trauma each year [12]. Sepsis was the single most expensive medical condition in the USA in 2016, with 22.2 billion dollars spent just on in-hospital stays [13]. Intra-abdominal sepsis (IAS) is the 2nd most common form of sepsis, and may be particularly severe because of the unique anatomic, physiologic, and microbiologic characteristics of hollow viscera within the abdominal cavity [14]. IAS occurs within a semirigid anatomic container that is exquisitely affected by raised intra-compartmental pressure that quickly induces abdominal visceral malperfusion and ischemia [15, 16]. Further, the extensive flora of the human microbiome is contained within the abdominal container exacerbating any pathology in a multitude of ways that are yet only minimally understood [17, 18]. Thus, it has been reported that hospital mortality is highest for patients who have intra-abdominal infection

secondary to ischemic bowel or disseminated infection [19].

Severe complicated intra-abdominal sepsis (SCIAS) represents a subset of IAS sepsis but is perhaps the most challenging clinical situation. Sartelli and the World Society of Emergency Surgery have defined IAS as severe when associated with organ dysfunction [9, 20–22], and as complicated when the inflammation or contamination spreads beyond a single organ, causing either localized or diffuse peritonitis [20, 23]. SCIAS may be distinguished from other causes of severe sepsis through a requirement for surgical abdominal exploration to address disruption in the gastrointestinal (GI) tract and provide source control.

Patients with SCIAS require early hemodynamic support, source control, and antimicrobial therapy [23]. Despite advances in diagnosis, surgery, and antimicrobial therapy, mortality rates associated with complicated intra-abdominal infections and IAS remain very high [22]. Failure to obtain adequate source control is often considered the driving cause of SCIAS and has been identified as an independent predictor of mortality [24]. Even with prompt appropriate therapy, SCIAS may progress to septic shock and multiple organ dysfunction, presumed as consequences of peritoneal and systemic inflammation. There is significant variability in the human immune

response to an infectious focus, whereas some individuals produce a massive bio-mediator storm propagating multisystem organ failure and death, whereas other individuals may be anergic with little or no response to the same stimuli.

In patients with SCIAS, repeat laparotomy may be necessary to eliminate persistent peritonitis or new infectious foci [25–27]. Differentiating “failed source control” [28, 29] from a self-propagating bio-mediator storm is difficult or impossible without abdominal re-exploration. In a Dutch multicenter randomized controlled trial (RCT), 42% of those randomized to expectant management after laparotomy for IAS, underwent relaparotomy for suspected or proven persistent peritonitis [25]. Interestingly, 31% of the repeat laparotomies were negative. The results of the Dutch study concluded a previously long-standing debate concerning two closed surgical approaches to ensuring source control in the peritoneal cavity; that of “laparotomy on demand – (LOD)” versus “planned relaparotomy” (PRL) [25, 30, 31]. The relative merits of either approach were widely debated until the conduct of the above RCT [25]. Although this trial noted no difference in mortality between the two methods, the LOD strategy reduced direct medical costs by 23% [25]. This equivalence in outcomes, coupled with apparent cost-savings, resulted in the generation of consensus guidelines recommending that LOD after laparotomy for SCIAS be adopted as the standard of care [32]. However, neither LOD nor PRL arm included an open abdomen or negative peritoneal pressure therapy (NPPT). The mortality in this RCT of severe secondary peritonitis illustrates the devastating nature of this disease having a mortality of approximately 1/3 of all enrolled patients regardless of treatment allocation. This observed mortality rate calls out for ongoing examination of alternative approaches to manage SCIAS.

Pharmacologic approaches do not currently offer hope in SCIAS as studies of promising agents directed to combat post-infective inflammation have not shown evidence of significantly improved patient outcomes, and when suggested as having a role, have been incredibly expensive [33, 34]. Alternatively, OA is increasingly recommended as an option to control intraperitoneal contamination and to ameliorate the propagation of inflammatory bio-mediators in SCIAS [35–37].

The use of the OA for non-trauma general surgery is increasingly being reported in uncontrolled series as an option for patients with SCIAS [20, 28, 29, 38–40]. The use of the OA approach in SCIAS may increase drainage of residual infection, allow early identification and control of persistent infection, increase the removal of bio-mediator-rich peritoneal fluid, prophylaxis against the development of the abdominal compartment syndrome,

and allow for the deferral of gastrointestinal anastomoses, with a potentially safer exit at the index operation [20]. However, compared to trauma patients, OA management for IAS patients has been reported to have a greater risk of complications, including enteroatmospheric fistula (EAF), intra-abdominal abscess, and a lower rate of primary fascial closure (i.e., fascia-to-fascia closure within the index hospitalization) [20, 21, 41] [42, 43]. Thus, there remains clinical equipoise in the regular use of the OA in SCIAS, with benefits and risks to adopting or avoiding its use.

Metanalyses and randomized controlled studies of the open abdomen in trauma and sepsis

Although the use of Damage Control and an OA concept was once liberally embraced and assumed to be the ideal therapy for major trauma [44], sober critique has questioned the need for this approach and suggested that the treatment paradigm and actual intervention may be over-used [45–47]. These concerns are germane when discussing non-trauma emergency surgical patients subjected to OA therapy as in IAS patients’ comorbidities are more common and more severe, closure rates are lower, and patients tend to be older and less able to withstand OA complications should they occur. Thus, it is important to have data unique to IAS patients to inform clinical decision-making.

Unfortunately, although case series on OA after non-trauma laparotomies have been reported, there are no contemporary RCTs. A recent meta-analysis on the use of Damage Control in perforated acute colonic diverticulitis [48], found no RCTs and ultimately the conclusions reverted back to opinions, the weakest level of Evidence in the World Society of Emergency Surgery Consensus Guidelines [49, 50]. In 2022, Cheng published a Cochrane Review on the use of negative pressure wound therapy for the non-trauma open abdomen and concluded that no recommendations could be made as there was no meaningful data [51]. Only one other RCT, conducted prior to 2006, has randomized 40 patients to a closed or open strategy, but the technique of OA management utilized then is inadequate according to current guidelines, as the NPPT apart from other aspects of OA management has evolved in technique and technology. This earlier RCT randomized patients with severe secondary peritonitis to an open or closed strategy after laparotomy, using a non-absorbable polypropylene (Marlex™) mesh in an interposed position between the open fascia, exposing the underlying bowel to the risk of enterocutaneous or enteroatmospheric fistula formation [52]. The study was stopped at the first interim analysis for futility. The risk of death was higher with the OA, but did not reach statistical significance, again leaving uncertainty as to how

to treat patients [52]. Otherwise, there is no prospective randomized data and results other than that which will be collected in the COOL trial.

Negative pressure peritoneal therapy (NPPT)

Newer non-commercial and commercial negative pressure peritoneal therapy (NPPT) systems are now available for OA and may reduce the risks of enterocutaneous fistula and facilitate enhanced delivery of negative peritoneal pressure to the peritoneal cavity [14, 32, 53]. In one of the largest contemporary OA databases, no difference in enterocutaneous fistula rates was noted related to the type of temporary abdominal closure dressing used [54]. However, there is a suggestion that more efficient peritoneal drainage may fundamentally impact the systemic complications of SCIAS. Animal studies [55] and in silica modeling of these animal studies [56] demonstrate that NPPT provides negative pressure and clearance of fluid throughout the peritoneum in contrast to simply leaving the fascia open with a temporary closure device. NPPT may reduce plasma bio-mediator levels compared to passive peritoneal drainage. Systemic inflammation (TNF- α , IL-1 β , IL-6) in a single animal study was significantly reduced in the NPPT group and was associated with significant improvement in intestine, lung, kidney, and liver histopathology [55].

Ugh—You want our advice? We don't really know!

Many of the current investigators in the COOL trial also conducted the largest prospective randomized controlled trial addressing the question of differing NPPT in open abdomen management, the Intra-Peritoneal Vacuum Trial [35]. Patients were enrolled in the operating room after an attending surgeon made the decision that an OA approach was required in critically ill/injured patients. Serum bio-mediator levels were measured every 24 h in the initial post-laparotomy phase of critical care [35, 57]. Although standard systemic bio-mediator levels were not statistically different nor was peritoneal fluid drainage, the 90-day survival rate was higher in the NPPT group ($P=0.04$) [35]. A valid critique of this trial was the heterogeneous mix of trauma and non-trauma patients [35]. A reasonable interpretation of this study's results is that the study's suggestion of a survivable benefit at minimum supports further investigation into therapeutic benefit in patients affected by severe SCIAS. In summary, great clinical equipoise remains as to whether the abdomen should be left open or closed after laparotomy in patients with SCIAS and warrents continuing to conduct the COOL Trial [38, 58].

The globalization of COOL

The original intent of the COOL trial investigators was to examine an OA-NPPT technique that could be used

anywhere [59]. The vision is to provide clinical operative guidance to surgeons with severe complicated abdominal sepsis as to whether they should close or not when the abdominal cavity is physically closeable. At the Inaugural Investigators COOL trial Meeting in Parma, Italy, the COOL trial Steering Committee endorsed the requirement to utilize an AbThera dressing (3 M, 3 M Center St. Paul, MN 55,144–1000). This decision was quite controversial and was fundamentally tied to financial trial support/sponsorship from the device manufacturer. It is important to clarify that apart from the use of the AbThera dressing, the sponsor was independent of the design or conduct of the study. The investigators assumed that the manufacturers of the AbThera would welcome the opportunity for an unbiased Global network of scientists to validate the efficacy of their proprietary device. This reflected the fact that the AbThera was only approved for use by the United States Food and Drug Administration, based on a so-called 510 K “loophole” that recognizes a substantial equivalence of the AbThera to 1976 predicate technologies, and not that the AbThera has ever been validated as better in any patient focused in rigorous human trials. Thus, the initial COOL Protocol required the use of a 3 M/Acelity AbThera dressing for any patient enrolled in the OA (intervention) arm of the Trial. This protocol stipulation was not without consequence as it precluded a “global” approach as many centers could not participate as the device was either not available and/or affordable.

The potential to utilize other non-commercial negative peritoneal pressure abdominal dressings in the COOL Trial

On August 19, 2022, the 3 M Company, who had acquired the Acelity Corporation canceled support for the COOL trial [59]. The sponsorship contract for the trial did permit the Corporate Sponsor to cancel support any time without cause. While a major logistical problem for the COOL trial Investigators, an unanticipated benefit is the removal of the requirement for use of the specific AbThera dressing in the OA arm. The COOL trial was always designed to be pragmatic, and the original protocol upon which ethics approval was obtained was generic regarding OA and NPPT management. The intervention arm of the trial has simply required NPPT administered to an OA defined by the fascia not being formally closed following all four intraperitoneal quadrants washed until macroscopically clear [32]. Thus, any manner of mechanical devices [60, 61], or potential instillation therapies [62], are permitted adjuncts as long as the primary requirement for an open fascia with NPPT is met.

Methods/Design

The current document is based upon the previously published COOL trial concise protocol [1], and outlines the evolution and lessons learned during the initial conduct of the COOL trial. Prompt resuscitation and the earliest possible appropriate antibiotic administration are critical for optimal outcomes in SCIAS. The COOL trial is pragmatic and will not stipulate specific protocols for such care, but emphasizes the importance of this for all patients whether not enrolled or enrolled into either arm of COOL.

Objective/Aims

The aim of the COOL trial is to test the null hypothesis that there will be no difference in survival when an OA management strategy administering NPPT is utilized compared to a primary fascial closure strategy in patients with SCIAS. The study is designed as a prospective, single-blinded, multicenter, international RCT. A SPIRIT Diagram overview of the trial is presented in Table 1. The complete protocol as well as a rich library of study-related documentation is available at www.coolstudy.ca.

Setting

The COOL trial is being conducted in operating rooms around the world where critically ill patients with SCIAS undergo source control laparotomy. The lead study center is the Foothills Medical Centre, a Quaternary Care Academic Medical Centre located in Calgary, Alberta, Canada. To date, thirteen hospitals on four continents have enrolled patients in the COOL trial, although more centers are open for recruitment.

Inclusion/exclusion criteria

Potential patients will first be identified in the emergency departments, in-patient ward, and critical care units of the participating centers. Eligibility can only be completely determined after the abdomen is explored in the operating room during the conduct of a laparotomy for source control. Patients will be eligible for inclusion if they have SCIAS, as operationally defined by the COOL trial (Fig. 1).

The inclusion criteria are conceptually a two-part assessment to ascertain if patients clearly fulfill the definition of both severe and complicated IAS (SCIAS) while undergoing source control laparotomy. Thus, during the laparotomy it will become apparent to the operating surgical team that peritonitis is complicated, which will be reproducibly demonstrated by uncontained or unconfined purulent, feculent, or enteric spillage. In addition to being complicated, the inclusion criteria require that patients have severe IAS. For the purpose of the COOL trial, severe will be defined by any of: septic shock as

defined by Sepsis 3 Consensus Guidelines [8], a World Society of Emergency Surgery Sepsis Severity Score >8 [9], or a Calgary Predisposition-Infection-Response-Organ Dysfunction Score >3 [63]. An elaborated explanation of the thought processes and identification attributes of these criteria modeled on a trial population of SCIAS patients was previously published by the COOL trial Investigators [64].

The exclusion criteria for the COOL trial include: a) pregnancy, b) perceived physical inability to physically close the fascia primarily without undue tension or concerns for inducing severe IAH/ACS, c) intra-operatively determined absolute or relative requirement for “damage control” laparotomy including intraperitoneal packing or non-anatomic postsurgical anatomy (i.e., surgically placed permanent packing or bowel that the operating surgeon believes must be left in discontinuity after resection), d) the patient is expected to die shortly after operation because of their condition in the operating room and there is no intention of providing ongoing care (i.e., the treating team wishes to close the abdomen to leave the operating room with the sole intention of withdrawing aggressive measures and providing only “comfort care” in the ICU; an example of where this could occur would be complete transmural midgut ischemia/necrosis), e) laparoscopic surgery (no laparotomy), f) pancreatitis as the source of peritonitis, g) acute superior mesenteric artery occlusion as the primary pathology, h) co-enrollment in another investigational study, i) peritoneal carcinomatosis, j) traumatic injury within 24 h of the development of SCIAS, k) age <18, or l) uncontrolled bleeding. It will be important for surgeons considering recruiting a patient to recognize before enrolling and randomizing a patient that fascial closure is not possible, as recognizing this after allocation to closure will constitute a protocol violation.

In current practice, it is likely that the most common reason for non-eligibility will be a surgeon-based decision to resect a hollow viscus and due to the perceived critical nature of the patient decide not to re-anastomose the bowel but to instead perform damage control and return the bowel ends into the peritoneal cavity without a diverting stoma. As this is an absolute indication for a future reoperation these patients will be ineligible for randomization.

Randomization

Treatment arm allocation is randomly allocated from a central, password-protected, randomization Web site (www.coolstudy.ca) (Fig. 2). This can be done from any computer or smartphone and accessing the enrollment site for randomization need not be conducted by the attending surgeon. The ability to enroll a patient

Table 1 SPRIT Diagram describing schedule of enrollment, interventions, and assessments

	Study Period							
	Assessment	Enrollment	Allocation	Open Abdomen Management	Fascial Closure	In hospital care	Post-Discharge Follow-up	
							90 days	1 year
Timepoint	Any critical In hospital patient	Intra-operative	Intra-operative					
Identification Tool	Suspect based on qSOFA $\geq 2(8)$	Septic shock OR WSESSSS > 8(9) OR CPIRO > 3(64) AND uncontained sepsis, purulence, and feces	Septic shock OR WSESSSS > 8(9) OR CPIRO > 3(64) AND uncontained sepsis, purulence, and feces					
Enrollment								
Eligibility Screen	X							
Informed Consent or Delayed Consent	X depending on local ethics	X depending on local ethics						
Randomization		X						
Allocation		X						
Intervention								
Fascial Closure			X	Reopen "on-demand"	Reopen "on-demand"			
Active Negative Pressure Peritoneal Therapy			██					
Assessments								
Mortality							90 days	
ICU Resource Consumption (potential)			██					
Logistical Outcomes			██					
Physiological Outcomes			██					
Health Care Spending (potential)			██					

1. Singer et al. [8]
 2. Sartelli et al. [9]
 3. Posadas-Calleja et al. [64]

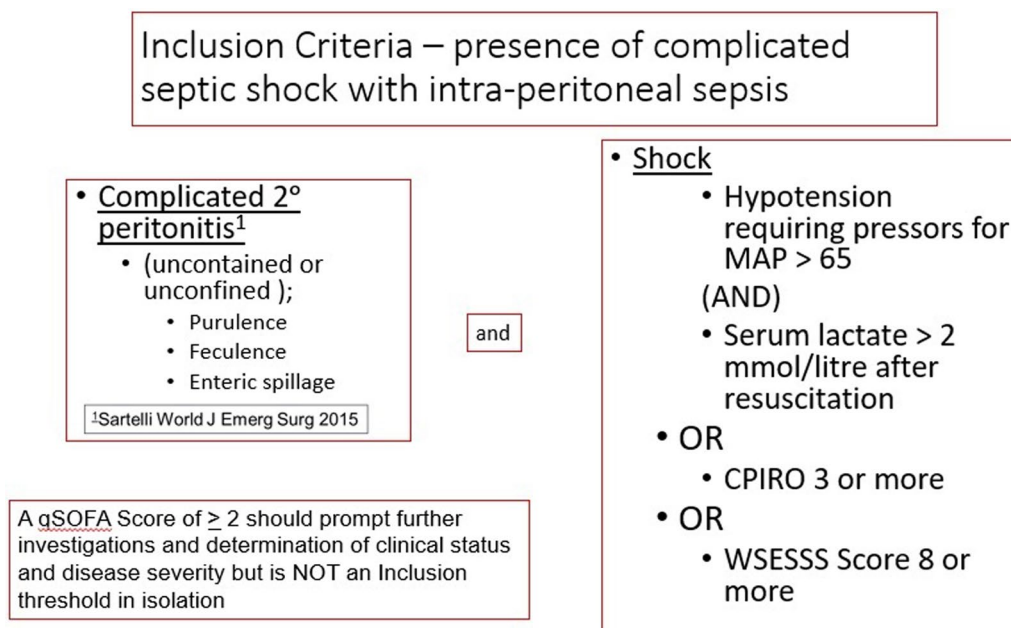


Fig. 1 Inclusion criteria for COOL

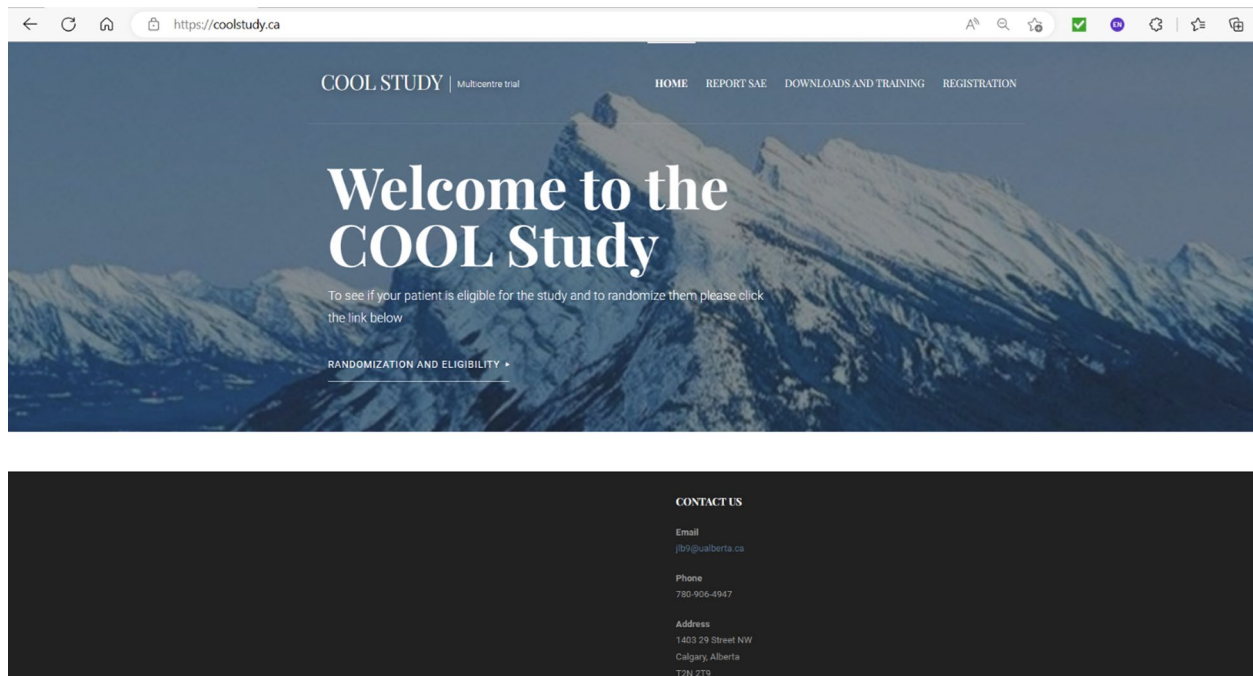


Fig. 2 COOL trial enrollment site

can be accessed with a password by any member of the surgical/anesthesia/critical care medicine/nursing team. When an appropriate patient is recognized, the research Web site will be accessed, simple identifiers of the patient will be entered, and treatment allocation (CLOSED with fascial closure or OPEN with an NPPT TAC dressing being applied) associated with this entry will be generated. To ensure the close balance of the numbers in each of the two treatment groups, permuted block randomization by site will be used. If the operating team is uncertain regarding the potential stratified severity according to either the WSESSSS or CPIRO methods, online decision support software simplifies these calculations regarding any potential enrollment.

Primary closure—CLOSED allocation (Control arm)

This strategy consists of primary closure of the fascia using any technique or suture material as chosen by the attending surgical team. Closure of the skin and the method for preventing surgical site infections is left to the discretion of the attending surgeon. There is no expectation for relaparotomy. Postoperative diagnostic imaging and all other aspects of postoperative care including any decision to perform a relaparotomy shall be at the discretion of the treating critical care/surgical teams. A decision to perform a relaparotomy will constitute a study outcome. If at any subsequent laparotomy the attending and responsible surgeon selects an open abdominal strategy (crossover to the intervention arm), the outcomes will be analyzed based on the intention-to-treat allocation at the time of original enrollment. Any application of any wound suction or negative pressure device to the soft tissue above a closed fascia is permitted within the control arm (closed abdomen).

Open Abdomen with Negative Pressure Peritoneal Therapy—OPEN allocation (Intervention arm)

Once the patient has been allocated to an OA, the trial protocol does not mandate the interval until fascial closure although the intention is that closure will occur expeditiously once clinically determined safe by the treating surgeon. The COOL trial protocol does not mandate any length of OA therapy, although the principle of the earliest safe formal closure is expected. The time that the temporary abdominal closure dressing will be left in place, will be left to the discretion of the attending surgeon, but typical practice guidelines mandate either formal abdominal closure or dressing changes at 24–72 h if formal abdominal closure cannot be completed [49]. For both arms of the trial, it will be expected that attending surgeons are involved in either the direct supervision

and/or intra-operative participation with either facial closure or temporary abdominal closure. The trial is considered pragmatic in allowing a variety of techniques as long as NPPT is being administered to an OA defined by the fascia not being formally closed and that all four intraperitoneal quadrants have been washed until macroscopically clear [32]. A suitable NPPT dressing must provide a complete visceroprotective layer, a means of the controlled egress of intraperitoneal fluid, and negative pressure within the peritoneal cavity. Thus, any manner of mechanical traction devices [60, 61, 65], or potential instillation therapies [62], will be permitted adjuncts as long as the primary requirement for an open fascia with NPPT is met. Terminology related to closure strategies and explicit methods used will be consistent with the terminology detailed in both the World Society of the Abdominal Compartment Syndromes most recent consensus definitions and practice management guidelines and those of the Open Abdomen Advisory Panel [32, 66, 67]. When the COOL trial was initiated, the commercial AbThera dressing was mandated, but this requirement was amended on August 2022 following 3 M's termination of the contract and sponsorship. Thus, other centers from countries that choose to use any other negative pressure dressing will be permitted; the type of NPPT will be considered in a subgroup analysis.

Outcomes

The primary outcome will be survival at 90 days from enrollment. Recent systematic reviews reveal higher mortality for both sepsis and septic shock populations examining 90- versus 30-day mortality rates and thus the 90 days of time frame was felt to provide a more comprehensive understanding of SCIAS mortality [68]. Secondary outcomes will be logistical and physiologic (Table 2). Logistical outcomes will include Days Free Of (DFO); ICU, ventilation, renal replacement therapy, and hospital at 90 days from the Index Laparotomy. The physiological secondary outcomes will include a change in APACHE II, SOFA, and ARDS scores after laparotomy. The COOL trial inclusion criteria concerning intraperitoneal contamination will be recorded, and the index source control laparotomy and every subsequent laparotomy will be graded according to the OA classification system from the 2013 World Society of Abdominal Compartment Syndrome (WSACS) grading scale for OA [32, 69, 70]. Surgical complications occurring after the index laparotomy will be graded according to Clavien-Dindo (Grade I=any deviation from normal postoperative course, including wound infections opened at the bedside but not treated with antibiotics; Grade II=requiring pharmacological treatment, e.g., antibiotic treatment, blood transfusion or parenteral nutrition; Grade IIIa=requiring

Table 2 Overview of study outcomes

	Indicator	Timeline
Primary outcome	Mortality	90 days
Secondary outcomes		
Logistical	Days free of ICU	30 days
	Days free of ventilation	30 days
	Days free of RRT ^a	30 days
	Days free of hospital	30 days
Physiological	APACHE II ^b scoresup to	30 days ^c
	SOFA ^d scoresup to	30 days ^c
	PaO ₂ /FI ₀₂ ^e ratiosup to	30 days ^c
	ARDS ^f scoresup to	30 days ^c
Safety	enterocutaneous fistula	30 days
	ACS ^g and/or severe IAH ^h	30 days
	Intra-abdominal abscess	30 days
	Need for reoperation	30 days

^a RRT = Renal Replacement Therapy

^b Acute Physiology and Chronic Health Evaluation Score

^c Measured daily using the worst value of that day

^d SOFA = Sequential organ Failure Assessment

^e PaO₂/FI₀₂ = Partial pressure of oxygen over inspired fraction of oxygen

^f ARDS = Acute Respiratory Distress Syndrome

^g ACS = Abdominal Compartment Syndrome

^h IAH = Intraabdominal Hypertension

surgical, endoscopic or radiologic intervention without general anesthesia and IIIb under general anesthesia; Grade IVa = life-threatening complication requiring IC/ICU management with single organ dysfunction and IVb with multiorgan dysfunction; Grade V = death of patient) [71, 72].

All data are entered into a secure web application for building and managing online surveys and databases (REDCap) maintained by the University of Calgary. While the COOL Trial Case Report form is available in paper format (Fig. 3), investigators are encouraged to submit data directly into the online format securely hosted in REDCap (Research Electronic Data Capture). The Case Report Form (CRF) was also recently simplified to become more pragmatic in anticipation of an increasingly global participation with less dedicated research administration. Although an immensely detailed and exhaustive COOL trial database would facilitate future “spin-off” studies, this should not be constructed at the expense of exhausting global collaborators dedicated to participate, but with limited research resources.

The Evolution of COOL over COVID and other World Crises

The initial protocol for the COOL trial envisioned multiple nested studies examining all aspects of OA management, of which an adequately powered trial of

mortality was the centerpiece [1]. Thus, any hospital providing emergency surgical services with intensive care support can participate if they are committed to recruit and randomize patients with SCIAS fulfilling the eligibility criteria during source control laparotomies. Contributing toward this main outcome will require only collection of the clinical outcome data. Prospective sub-studies that were envisioned to augment this main goal included COOL-Max (Bio-mediators), COOL-Mic (Microbiology), COOL-Cells (cellular defense mechanism), and COOL-Costs (economics). After the initiation of the clinical COOL trial, it became apparent that realistic operational demands and economic limitations precluded the conduct of these sub-studies, although a retrospective COOL trial economic analysis of open versus closed treatment is still a practical future analysis [73]. Thus, the dedicated focus of the current COOL trial efforts is completing the clinical outcome analysis powered on mortality.

Sample size calculations

The COOL trial is overall powered to detect a significant difference in the primary outcome, 90-day survival. Although imperfect, the preceding Intra-Peritoneal Vacuum Trial study revealed an Intention-to-treat 90-day mortality of 21.7% in the ABThera group versus 50.0% in the Barker’s vacuum pack group [HR, 0.32; 95% confidence interval (CI), 0.11– 0.93; P=0.04] [74]. This 30% reduction in mortality was considered too dramatic to be practically replicated and a much more conservative effective 10% reduction in mortality was chosen. Thus, given a mortality rate of 33% in the general population of those with severe intra-abdominal sepsis, and considering a power of 80% and an alpha of 0.05, the number needed to recruit was calculated as 275 patients in each arm.

Statistical analyses

The effectiveness of randomization will be displayed through a detailed presentation of patient demographic characteristics as outlined in Table 3. The analysis of the primary outcome, mortality, will be on an intention-to-treat basis related to the allocation of initial intra-operative therapy. There will be a planned subgroup analysis of the mortality stratifying patients into those with and without the presence of septic shock (defined as Sepsis-3 Consensus Guidelines) during the first 48 h after onset of peritonitis (if known and 24 h before and 24 h after 1st laparotomy if not known). There will also be a planned subgroup analysis looking for any difference in outcomes within the intervention arm of the study between patients managed with the AbThera commercial dressing and any other NPPT dressing.

COOL Study: CRF Version 3.2 Page | 1 Pt ID Number: [] [] [] [] [] - [] [] [] [] [] - [] [] [] [] []

Demographics		Surgical History Data Has the patient had any intra-abdominal surgery in the 30 days prior to the index laparotomy? Yes <input type="checkbox"/> No <input type="checkbox"/> If yes, please describe below* Date of Surgery (dd/mm/yyyy): [] [] [] [] [] []
Age (years)		
Gender	<input type="checkbox"/> M <input type="checkbox"/> F	
Height (cm)	.	
Weight (kg)	.	

*If there has been more than one intra-abdominal surgery in the 30 days prior to the index laparotomy, please use a separate surgical history data collection form for each. See Appendix A.

Index ICU Admission Data (24 hrs of admission to ICU. Even if Prior to Index Laparotomy)		Admission Date (dd/mm/yyyy): [] [] [] [] [] []	Admission Time: [] [] [] [] [] [] AM/PM
Vasopressors received: Yes <input type="checkbox"/> No <input type="checkbox"/> If Yes, please enter highest value below*		Vitals and Laboratory Results. If not available enter 'NTA' APACHE II: [] [] [] [] [] []	
Dopamine (ug/kg/min):	Dobutamine (ug/kg/min):	PaO2/FiO2** (mmHg) (Lowest):	GCS*** (Lowest):
Norepinephrine (ug/kg/min):	Milrinone (ug/kg/min):	Platelets x (10 ⁹ /L) (Lowest):	Bilirubin (umol/L) (Highest):
Vasopressin (ug/kg/min):	Epinephrine (ug/kg/min):	MAP (mmHg) (Lowest):	Creatinine (umol/L) (Highest):
Noesynephrine (Total ugs):	Ephedrine (Total ugs):	Respiratory Rate (BPM) (Highest):	Systolic BP (mmHg) (Lowest):

*If a vasopressor was not given enter '0' in the field ** if no ABG, use SaO2/FiO2 Ratio ***If on sedatives or intubated estimate assumed GCS off sedation or intubated

Study Enrollment Data / Shock		Yes	No
Did the patient have complicated secondary peritonitis at laparotomy including purulence, feculence and/or enteric spillage?			
Did the patient have hypotension requiring pressors of MAP > 65 mmHg?			
Did the patient have a serum lactate > 2 mmol/L after resuscitation?			
CPIRO			
Age > 65 years?			
Any of the following comorbidities*: Liver Insufficiency, Cardiovascular, Respiratory, Renal, Immunosuppression?			
Did they have a WBC <4000			
Did they have a temperature <36 degrees Celsius			
Mean arterial pressure <90 OR administration of any vasopressor medications			
PaO2/FiO2 (mmHg) <300			
Creatinine > 171 umol/L (2 mg/dl) OR < 500 ml urine day			
Glasgow Coma Scale ≤ 12			
WSESS			
Did the patient have sepsis with organ dysfunction at admission*			
At admission was the patient in acute septic shock requiring vasopressor agents*			
Does/did the patient have health-care associated infection			
Was Colonic non-diverticular perforation peritonitis the origin of intra-abdominal infection?			
Was Small Bowel perforation peritonitis the origin of intra-abdominal infection?			
Was Diverticular diffuse peritonitis the origin of intra-abdominal infection?			
Was Post-operative diffuse peritonitis the origin of intra-abdominal infection?			
Was there a delayed initial intervention of >24 hours for source control for your patient?*			
Age > 70 years:			
Does your patient have any of the following: Chronic glucocorticoids, immunosuppressive agents, chemotherapy, lymphatic disease, virus, immunosuppression from chemotherapy, radiation therapy, long-term or recent high-dose steroids, immunodeficiency (eg, leukemia, lymphoma, AIDS)?			

*For further clarification on this question see Appendix B

PRE- Operative Data*	Date (dd/mm/yyyy): [] [] [] [] [] []	Patient Location	Ward	ICU	ED	CVICU	Transferred from another Hospital	Step-down unit
Vasopressors received: Yes <input type="checkbox"/> No <input type="checkbox"/> If Yes, please enter highest value below*		Vitals and Laboratory Results. If not available enter 'NTA'						
Dopamine (ug/kg/min):	Dobutamine (ug/kg/min):	PaO2/FiO2** (mmHg) (Lowest):	GCS*** (Lowest):					
Norepinephrine (ug/kg/min):	Milrinone (ug/kg/min):	Platelets x (10 ⁹ /L) (Lowest):	Bilirubin (umol/L) (Highest):					
Vasopressin (ug/kg/min):	Epinephrine (ug/kg/min):	MAP (mmHg) (Lowest):	Creatinine (umol/L) (Highest):					
Noesynephrine (Total ugs):	Ephedrine (Total ugs):	Respiratory Rate (BPM) (Highest):	Systolic BP (mmHg) (Lowest):					
Charlson Co-morbidity**		Yes	No	Charlson Comorbidity**		Yes	No	Randomization Data
Myocardial infarct				Diabetics with end organ damage				Randomization Date (dd/mm/yyyy)
Congestive heart failure				Cancer (without metastasis)				Randomization Time (HH : MM)
Cerebrovascular disease				Cancer with metastasis				AM
Hemiplegia				Leukemia/lymphoma				PM
Peripheral vascular disease, AAA				Dementia				Randomization: Open Closed
Connective tissue disease				Severe Organ Dysfunction, Kidney				If Open, AbThera Product Number:
Immunosuppression/steroids use/chemotherapy				Severe organ dysfunction, Liver				Notes:
Smoking				Presence of ostomy/incisional hernia				
Pulmonary disease mild				HIV infection				
Pulmonary disease modern				HIV infection with complications				
Pulmonary disease severe				Liver disease mild				

Fig. 3 COOL study case report form. The Case Report Form is an extensive document that can be accessed online at Study Documents – COOL Study, but Investigators are encouraged to complete the form on-line where it will be securely entered into the University of Calgary REDCap (Research Electronic Data Capture) database.

Table 3 Baseline demographic characteristics of the study patients

Male/female
Age, median (IQR) ^a , years
Septic shock ^b
World Society of Emergency Surgery Sepsis Severity Score ^c
Calgary PIRO Score ^d
GCS ^e , median (IQR)
APACHE II ^f , median (IQR)
Arterial pH, mean (95% CI)
Base deficit, median (IQR)
Lactate, median (IQR)
INR ^g , median (IQR)
Temperature, mean (95% CI)
APACHE II score ^f , mean \pm SDd
SOFA score ^h , mean \pm SDe
Charlson Comorbidity Index score ⁱ , median (IQR)
Worst physiologic measurements prior to randomization, median (IQR)
Systolic blood pressure, mmHg
Temperature (injured patients), °C
Temperature (sepsis patients), °C
pH
Lactate, mmol/L
Base deficit, mmol/L
INR
Fluid administration prior to randomization, median (IQR)
PRBC ^j , units
FFP ^k , units
PRBC/FFP ratio
Crystalloid, L
Patient location prior to OR admission—no. (%)
Emergency Department
Hospital ward
Intensive Care Unit
Vasopressors required prior to randomization—no. (%)
Hours from sepsis diagnosis to laparotomy, median (IQR)
If allocated to OPEN
Type of Negative pressure peritoneal therapy (NPPT) temporary abdominal closure (TAC) device

^a IQR, interquartile range^b septic shock as defined by SESPS-3 Guidelines[8]^c WSESS [9]^d CPIRO [63]^e GCS—Glasgow coma score^f Acute Physiology And Chronic Health Evaluation II^g INR—international normalized ratio^h SOFA—Sequential Assessment of Organ Failure [137]ⁱ Charlson Comorbidity Index[138]^j PRBC—packed red blood cells^k FFP—fresh frozen plasma

There will be a single interim analysis planned after the recruitment of 275 patients, which will analyze the difference in 90-day mortality between allocated therapies. The COOL trial Investigators appreciate the general reluctance to stop randomized trials early due to benefit, due to the frequent over-estimating of treatment effects [75–77]. Despite this, if a profoundly significant difference is found ($p < 0.01$) the trial will be stopped, otherwise it will continue to full recruitment.

Ethical concerns

There is clinical equipoise concerning the operative management of SCIAS. Thus, the COOL trial Investigators feel a moral imperative to conduct this research to provide the best evidence to counsel bedside critical care physicians and surgeons [78]. The COOL trial is currently approved by the Conjoint Health Research Ethics Board of the University of Calgary (REB-16-1588) to proceed with a delayed consent process given the time-sensitive critical nature of decision-making. Research ethics will vary throughout the world and it is anticipated that various local policies concerning community consent, waiver of consent, or informed consent of significant patient proxies may be considered. All participating Institutions will thus be required to obtain Ethical Approval appropriate and applicable to their Institutions.

Research in critically ill incapacitated patients is important to advance care. Conducting research among SCIAS is complicated due to the severity of illness, need for emergent interventions, diagnostic criteria confirmed only at laparotomy, and obtundation from anesthesia. In other circumstances involving critically ill patients, clinical experts have worked closely with ethicists to apply principles that balance the rights of patients while simultaneously permitting inclusion in research. COOL Investigators have collaborated with both current and past Chairs of REB's to review and interpret the science and ethics for surgical investigators globally [79, 80]. The ultimate goal is to balance respect for patient participants and to permit participation with a reasonable opportunity for improved outcome and minimal risk of harm.

Discussion

Randomized surgical trials, especially those not supported by industry are notoriously few, hard to complete, and increasingly poorly supported by traditional granting agencies [81–83]. Yet these trials are desperately required. In general, the overall quality of surgical research can be criticized as being grossly inadequate despite being the purported basis of surgeons making

evidence-informed decisions with an impact which may affect a patient's outcome including death or being permanently impaired [79, 80, 84, 85]. One famous commentary compared surgical research to "comic opera" [86], lamenting the reliance on retrospective case series as a methodology, and another referred to the typical retrospective case series (that constitute the near totality of research concerning SCIAS) as "research waste" [84]. Unfortunately, retrospective case series predominate, potentially because they are vastly easier to conduct, are free of regulatory hurdles that accompany conducting an RCT, are publishable in journals and offer career advancement to investigators. However, why surgical RCTs are so few may also relate to fundamental differences in the regulatory approval process between medicines and medical devices. Whereas the level of confidence in pharmaceutical safety has risen substantially since the Thalidomide debacle [87], comparable changes in the safety bar to approve medical devices are less well developed. Thus, RCTs are often not required by device manufacturers or regulators to allow market entry [84], and thus research funding for devices demonstrating a beneficial effect on outcome is often lacking.

Nonetheless, the COOL trial has been designed to answer a critical clinical question that faces clinicians worldwide on a daily basis for which there is important clinical equipoise and potential severe consequences for patients in regards to outcomes [38, 58]. Thus, this question has been identified as one requiring urgent study [49]. The COOL trial has continued to be supported by not-for-profit Scientific Organizations with a vested interest in the best care of the critically ill patient including the Abdominal Compartment Society and the World Society of Emergency Surgery. The trial design and vision follow directly from the preceding single-center study of differing modalities of NPPT conducted at the Foothills Medical Centre [57, 74]. When the Intra-Peritoneal Vacuum Trial investigators considered following up the pilot study and enrolling more patients in a multicenter fashion, it became apparent after peer-to-peer discussion that any differing effectiveness of NPPT techniques was not the most relevant question concerning the OA [88]. With an evolution in resuscitation practices involving balanced resuscitation, more and more trauma patients who previously become so edematous they required OA therapy, are no longer being over-resuscitated with crystalloids, and can be primarily closed [89–91]. This change in at least the trauma care paradigm has justified questions regarding the whole premise of damage control surgery for trauma [92], and IAS [45].

As over-resuscitation becomes less common [93, 94], it is intuitive that there will be more abdomens in non-trauma IAS patients which may be technically closed

without inducing intra-abdominal hypertension (IAH). However, although these abdomens *may* be closed, *should* they be closed? As has been recently emphasized, there are profound differences in the basic science of sepsis and traumatic injury [95], with the previously unifying concepts of non-infectious Systemic Inflammatory Response Syndrome (SIRS) being effectively discarded as a clinically helpful construct [8, 96, 97]. The one nebulous, poorly defined "holy grail" of the optimal management of SCIAS is adequate "source control." It is suggested that even if an abdomen can be physically closed that there may be an advantage to leaving it open for a brief period to allow better drainage of intraperitoneal contamination, a concept that is supported by animal data suggesting the ability of NPPT to mitigate the elaboration of the inflammatory bio-mediator cascade [55, 56, 98], although this has not been demonstrated in humans [74].

The peritoneal cavity as a reservoir for systemic inflammation

There is a complex relationship between pressure, ischemia, and inflammation within the peritoneal cavity [14, 16]. Independently the damaged gut seems to act as a continued source of inflammation propagating SIRS and potentiating MODS [99–103]. Basic, predominantly animal laboratory research, from the last decade suggests an exciting potential. Visceral ischemia characteristically generates multiple immunological mediators with the pro-inflammatory cytokines tumor necrosis factor-alpha (TNF- α), and interleukin six (IL-6), as well as inhibitive cytokines such as interleukin ten (IL-10) [104–107]. Post-operative complications are associated with increasing levels of systemic IL-6, and peritoneal TNF- α [106, 108]. Jansson and colleagues thus postulated that peritoneal cytokines in humans respond more extensively compared to systemic cytokine, and that a normal postoperative course is characterized by decreasing levels of peritoneal cytokines based on studies of both elective and emergency surgery [109]. Overall, the peritoneal cytokine response is much higher than the systemic response in peritonitis [107, 110–112]. Hendriks and colleagues demonstrated that peritoneal cytokine levels (especially IL-6, TNF- α [113], and IL-10) were dramatically different in rats who either survived or succumbed to an IAS model in the 24 h after cytokine determination [110]. Finally, recent work suggests that blood filters designed to hemofiltrate blood endotoxins and cytokines may improve hemodynamics, organ dysfunction and even mortality in the critically ill [114–117].

The biologic rationale for COOL is that if safe, removing intraperitoneal bio-mediators may mitigate their local effects and prevent their being absorbed systemically.

Although early work suggested a benefit of simple continuous peritoneal lavage after either gross peritoneal contamination in secondary peritonitis or in the setting of necrotizing pancreatitis [118, 119], subsequent studies could not confirm a benefit [120–122]. Studies using hemofiltration to remove inflammatory mediators from the blood have been associated with reduced elevations of inflammatory cytokines (as assessed by blood IL-6 levels), early improvements in hemodynamic state and decreased lactate levels [123–125]. However, it has not yet been demonstrated that extracorporeal filtration of inflammatory mediators improves clinical outcomes [126, 127]. One possible explanation for this is that after the mediators have left the peritoneal cavity and become systemic the “horse is out of the barn.”

NPPT therapy may be a more direct, earlier, and focused solution to this complicated problem, and one that will be complementary to the other benefits of OA. Whether improved postoperative courses can be obtained through this relatively simpler approach of actively removing peritoneal cytokines with a more efficient and comprehensive VAC therapy in humans is therefore part of the biologic rationale of the COOL trial.

Another potential benefit of NPPT after severe infection may be the attendant decompression of the abdominal compartment and prevention of even modest IAH. Patients with intra-abdominal infections are at risk of elevated IAP both because of the primary intraperitoneal disease, and as a consequence of the use of large-volume crystalloid resuscitation often used to maintain organ perfusion [128–130]. Recent studies have demonstrated a high prevalence of IAH following aggressive volume resuscitation of septic patients. IAH is present in as many as 80% of septic medical and surgical ICU patients [131, 132]. Reintam also reported that septic patients with IAH had a 50% mortality rate compared to 19% without IAH, making IAH a significant marker for an increased risk of death [133]. Within the lead COOL Institution rates of IAH were over 87% of septic ICU patients and further 61% of these patients had severe IAH at levels commensurate with ACS, despite the fact that IAP was only measured in 10% of the patients in whom guidelines recommend monitoring [134]. Although direct translation to humans is uncertain, even modest degrees of IAH (often clinically ignored) have been found to have profound far-reaching effects on propagating multiple organ failure in animals with ischemia/intraperitoneal infections [135–137].

COOL trial recruitment

Like many, especially investigator-initiated randomized trials, recruitment has lagged behind original predictions for the COOL trial. Poor participant recruitment

is the most frequent cause for premature discontinuation of randomized clinical trials [138, 139]. The COOL trial has competed with the COVID pandemic as a novel challenge apart from other established causes for poor trial enrollment such as inadequate funding, a narrow (but necessary) eligibility criteria, and a de-emphasis of research priorities even in University hospitals [138]. The financial burden of Clinical Trial Insurance has been a particularly challenging burden to the COOL trial. The difficulty in financing was made worse by 3 M canceling its contract to support the COOL trial. However, recruitment is measured against an arbitrary prediction, so the true adequacy of recruitment will only be assessable when the outcome data is formally analyzed. Although this is not planned until 275 patients have been recruited, it is relevant that at this time COOL is nearly twice as large as the most relevant RCT previously reported [52]. Thus, as new centers are added (as they have been monthly) the COOL trial will continue and should be successful in meeting its enrollment goals.

Conclusions

The COOL trial is designed to examine if a mortality difference exists in this highly lethal and morbid condition, and to ensure critically ill patients are receiving the best care possible and not being harmed by inappropriate interventions or devices based on opinion only. The COOL trial Investigators now welcome truly global participation for all interested surgical scientists and their supporters.

Abbreviations

COOL	Closed Or Open after Laparotomy trial (https://clinicaltrials.gov/ct2/show/NCT03163095)
SCIAS	Severe complicated intra-abdominal sepsis
OA	Open abdomen
NPPT	Negative pressure peritoneal therapy
IAS	Intra-abdominal sepsis
RCT	Randomized Controlled Trial
LOD	Laparotomy on demand
PRL	Planned relaparotomy
EAF	Enteroatmospheric fistula
CPIRO	Calgary Predisposition, Infection, Response, and Organ Dysfunction
APACHE	Acute Physiology and Chronic Health Evaluation
SOFA	Sequential Organ Failure Assessment
ARDS	Acute respiratory distress syndrome
MODS	Multiple Organ Dysfunction Score
SIRS	Systemic inflammatory response syndrome
qSOFA	Quick SOFA score
WSES/SSS	World Society of Emergency Surgery Sepsis Severity Score
APC	Activated protein C
mtDNA	Mitochondrial DNA
C5a	Complement factor 5 activated
C3a	Complement factor 3 activated
REDCap	(Research Electronic Data Capture) database
REB	Research Ethics Board
TAC	Temporary abdominal closure
IAH	Intra-abdominal hypertension

IAP Intra-abdominal pressure
ACS Abdominal compartment syndrome

Acknowledgements

Not applicable.

Author contributions

AWK, LA, both FCs, both MSs, MT, JLM, CD, and MDH, as well as the General membership of the Abdominal Compartment Society and the World Society of Emergency Surgery conceptualized the study. AWK drafted the initial manuscript. CD, MH, AG, JLM, and DJR extensively rewrote the initial manuscript. All authors read and approved the final manuscript.

Funding

No funding was received for the conduct of this analysis or the preparation of this manuscript. The COOL Investigators as an entity were supported to conduct a Protocol Development Meeting in Parma Italy on November 26, 2017, through an unrestricted grant from the Acelity Corporation in which Inclusion Criteria for Closed Or Open after Laparotomy (COOL) for Source Control in Severe Complicated Intra-Abdominal Sepsis (<https://clinicaltrials.gov/ct2/show/NCT03163095>) trial was discussed. The 3 M/Acelity Corporation provided limited financial support for the COOL trial under contract 032519-000/2050 until unilateral cancellation on August 19, 2022. The COOL trial has also received unrestricted support from the Abdominal Compartment Society and the Andrew W Kirkpatrick Professional Corporation.

Availability of data and materials

All results and data from the COOL trial will be available from Dr. Andrew Kirkpatrick (andrew.kirkpatrick@albertahealthservices.ca) on reasonable request, as well as from the Study Web site (www.coolstudy.ca).

Declarations

Ethics approval and consent to participate

The COOL trial has been ethically approved at the lead and pilot center by the Conjoint Health Research Ethics Board (CHREB) of the University of Calgary (REB16-1588). The study has also been registered with the National Institutes of Health (ClinicalTrials.gov Identifier: NCT03163095).

Consent for Publication

Not Applicable.

Competing interests

Andrew W Kirkpatrick serves as the PI of the COOL trial, as a member of the Canadian Forces Medical Services, and has consulted for the 3 M/Acelity Corporation, Zoll Medical, Innovative Trauma Care, CSL Behring, and the Statesman's Group. Federico Coccolini, Matti Tolonen reported no declarations. Samuel Minor reported receiving research support and speaking honoraria from COOK Biotech. Emanuel Gois Jr. reported no declarations. Fausto Catena, Christopher J Doig, Michael D Hill, Luca Ansaloni, Massimo Chiurigi, Dario Tartaglia, Orestis Ioannidis reported no declarations. Michael Sugrue reported consultancy for 3 M/Acelity and Novus Scientific. Elif Colak reported no declarations. S Morad Hameed reported being the Founder of T6 Health Systems. Hanna Lampela, Vanni Agnoletti reported no declarations. Jessica L McKee reported consultancies with the Aceso, Innovative Trauma Care, Andrew W Kirkpatrick, and Zoll Corporations, as well as consulting with the Geneva Foundation and South Trail Psychology. Naisan Garraway, Massimo Sartelli, Chad G Ball reported no declarations. Neil G Parry reported being a medical advisor for Front Line Medical Technologies – Cobra REBOA. Kelly Voght, Lisa Julien, Jenna Kroeker reported no declarations. Derek J Roberts, Peter Faris, Corina Tiruta, Ernest E Moore, Lee Ann Ammons, Elissavet Anestiadou, Cino Bendinelli, Konstantinos Bouliaris, Rosemary Carroll, Marco Ceresoli, Francesco Favi, Angela Garrado, Joao Rezende-Neto, Arda Isik, Camilla Cremonini, Silivia Strambi, Georgios Konstantoudakis, Mario Testini, Sandy Trpcic, Alessandro Pasculli, Erika Picariello, Fikri Abu-Zidan, Ademola Adeyeye, Goran Augustin, Felipe Alconchel, Yuksel Altinel, Luz Adriana Hernandez Amin, José Manuel-Narváez, Oussama Baraket, Walter L Biffi, Gian Luca Baiocchi, Luigi Bonavina, Giuseppe Brisinda, Luca Cardinali, Andrea Celotti, Mohamed Chaouch, Maria Michela Chiarello, Gianluca Costa, Nicola deAngelis, Nicolo de Manzini, Samir Delibegovic, Salomone Di Saverio, Belinda De Simone reported

no declarations. Dr Vincent Dubuisson received speaking honoraria from 3 M-Acelity in 2021 for a conference about how to manage an OA, at the congress of the French Association of Surgery. Pietro Fransvea, Luca Garulli, Alessio Giordano, Carlos Gomes, Firdaus Hayati, Jinjian Huang, Aini Fahriza Ibrahim, Tan Jih Huei, Ruhi Fadzlyana Jailani, Mansoor Khan, Alfonso Palmieri Luna reported no declarations. Manu L.N.G. Malbrain reported he is co-founder, past-President and current Treasurer of WSACS (The Abdominal Compartment Society, <http://www.wsacs.org>). He is a member of the medical advisory Board of Pulsion Medical Systems (part of Getinge group), Serenno Medical, Potrero Medical, Sentinel Medical and Baxter. He consults for B Braun, Becton Dickinson, ConvaTec, Spiegelberg, and Holtech Medical and received speaker's fees from PeerVoice. He holds stock options for Serenno and Potrero. Sanjay Marwah, Paul McBeth, Andrei Mihailescu, Alessia Morello, Francesk Mulita, Valentina Murzi, Ahmad Tarmizi Mohammad, Simran Parmar, Ajay Pal, Michael Pak-Kai Wong, Desire Pantalone, Mauro Podda, Caterina Puccioni, Kemal Raza, Jianan Ren, Francesco Roscio, Antonio Gonzalez-Sanchez, Gabriele Sganga, Maximilian Scheiterle, Mihail Slavchev, Dmitry Smirnov, Lorenzo Tosi, Anand Trivedi, Jaime Andres Gonzalez Vega, Maciej Waledziak, Sofia Xenaki, Desmond Winter, Xiuwen Wu, Andee Dzulkarnean Zakaria reported no declarations.

Author details

¹Departments of Surgery and Critical Care Medicine, University of Calgary, Foothills Medical Centre, Calgary, AB EG23T2N 2T9, Canada. ²General, Emergency and Trauma Surgery Department, Pisa University Hospital, Pisa, Italy. ³Abdominal Center, Helsinki University Hospital and University of Helsinki, Helsinki, Finland. ⁴Departments of Critical Care Medicine and Surgery, Dalhousie University, Halifax, NS, Canada. ⁵Department of Surgery, Bufalini Hospital, Cesena, Italy. ⁶Department of Surgery, Londrina State University, and National COOL Coordinator for Brazil, Londrina, Brazil. ⁷Departments of Critical Care Medicine and Community Health Sciences, Cumming School of Medicine, University of Calgary, Calgary, AB, Canada. ⁸Department of Clinical Neuroscience and Hotchkiss Brain Institute, Cumming School of Medicine, University of Calgary and Foothills Medical Centre, Calgary, AB, Canada. ⁹General Surgery I, San Matteo Hospital Pavia, University of Pavia, Pavia, Italy. ¹⁰Emergency Surgery and Trauma Center, University of Pisa, Pisa, Italy. ¹¹4th Department of Surgery, Medical School, Aristotle University of Thessaloniki, General Hospital "George Papanikolaou", Thessaloniki, Greece. ¹²Letterkenny University Hospital, Donegal, Ireland. ¹³University of Samsun, Samsun Training and Research Hospital, Samsun, Turkey. ¹⁴Department of Surgery, University of British Columbia, Vancouver, BC, Canada. ¹⁵Department of Gastroenterological Surgery, Helsinki University Hospital and University of Helsinki, Espoo, Finland. ¹⁶Chief Anesthesiology, Bufalini Hospital, Cesena, Italy. ¹⁷Global Project Manager, COOL Trial and the TeleMentored Ultrasound Supported Medical Interventions Research Group, Calgary, AB, Canada. ¹⁸Departments of Surgery and Critical Care Medicine, University of British Columbia, Vancouver, BC, Canada. ¹⁹Department of Surgery, Macerata Hospital, Global Alliance for Infections in Surgery, Macerata, Italy. ²⁰Trauma and Acute Care Surgery, Foothills Medical Center, Calgary, AB, Canada. ²¹Departments of Surgery and Medicine, Schulich School of Medicine and Dentistry, Western University, London, ON, Canada. ²²Department of Surgery, NSHA-Queen Elizabeth II Health Sciences Center, Dalhousie University, Halifax, NS, Canada. ²³Division of Vascular and Endovascular Surgery, Department of Surgery and School of Epidemiology and Public Health, Faculty of Medicine, University of Ottawa, Ottawa, ON, Canada. ²⁴University of Calgary, Calgary, AB, Canada. ²⁵Alberta Health Services, Calgary, AB, Canada. ²⁶Ernest E. Moore Shock Trauma Center, University of Colorado, Denver, CO, USA. ²⁷Department of Surgery, Denver Health, Denver, CO, USA. ²⁸John Hunter Hospital, Newcastle, NSW, Australia. ²⁹General Surgery Department of Koutlimbaineio, Triantafylleio General Hospital of Larissa, Larissa, Thessaly, Greece. ³⁰General and Emergency Surgery, School of Medicine and Surgery, Milano-Bicocca University, Monza, Italy. ³¹Chirurgia Generale E d'Urgenza, Ospedale M. Bufalini - Cesena, AUSL Della Romagna, Cesena, Italy. ³²Department of Precision and Regenerative Medicine and Ionian Area, Unit of Academic General Surgery "V. Bonomo", University of Bari "A. Moro", Bari, Italy. ³³Trauma and Acute Care Surgery, General Surgery, St. Michael's Hospital, Toronto, ON, Canada. ³⁴General Surgery Department, Istanbul Medeniyet University School of Medicine Istanbul, Istanbul, Turkey. ³⁵General Surgery Unit, Ospedale M. Bufalini Di Cesena, Cesena, Italy. ³⁶College of Medicine and Health Sciences, United Arab Emirates University, Al-Ain, United Arab Emirates. ³⁷Division of Surgical Oncology, Afe Babalola

University Multisystem Hospital, Ado-Ekiti, Nigeria. ³⁸University Hospital Centre Zagreb, School of Medicine University of Zagreb, Zagreb, Croatia. ³⁹Virgen de la Arrixaca University Hospital IMIB-Arrixaca, Ctra. Madrid-Cartagena, S/N, Murcia, Spain. ⁴⁰Bagcilar Research and Training Hospital, Istanbul, Turkey. ⁴¹Nurse Master of Nursing, Professor and Coordinator of the teaching-service relationship, Faculty of Health Sciences, University of Sucre, Sincelejo, Colombia. ⁴²Trauma and Emergency Surgery Unit, General, Digestive and Transplantation Surgery Department, University Regional Hospital of Málaga, Malaga, Spain. ⁴³Department of Surgery, Bizerte Hospital, Bizerte, Tunisia. ⁴⁴Scripps Clinic Medical Group, La Jolla, CA, USA. ⁴⁵Department of Clinical and Experimental Sciences, University of Brescia, Brescia, Italy. ⁴⁶Department of Surgery, University of Milan Medical School, Milan, Italy. ⁴⁷Department of Surgery, Fondazione Policlinico Universitario A Gemelli IRCCS, Rome, Italy. ⁴⁸Department of Surgery, General Hospital Madonna del Soccorso, San Benedetto del Tronto, Italy. ⁴⁹General Surgery Unit, UO Chirurgia Generale - Ospedale Maggiore Di Cremona, Cremona, Italy. ⁵⁰Department of Visceral and Digestive Surgery, Monastir University, Monastir, Tunisia. ⁵¹Department of Surgery, Azienda Sanitaria Provinciale Di Cosenza, Cosenza, Italy. ⁵²Fondazione Policlinico Campus Bio-Medico, University Campus Bio-Medico of Rome, Rome, Italy. ⁵³Colorectal and Digestive Surgery Unit-DIGEST Department, Beaujon University Hospital AP-HP, University Paris Cité, Clichy, France. ⁵⁴Department of General Surgery, Cattinara University Hospital, Trieste, Italy. ⁵⁵Department of Proctology, Clinic for Surgery, University Clinical Center Tuzla, Tuzla, Bosnia and Herzegovina. ⁵⁶Department of General Surgery, University of Insubria, University Hospital of Varese, ASST Sette Laghi, Regione Lombardia, Italy. ⁵⁷Unit of Digestive and Metabolic Minimally Invasive Surgery, Clinique Saint Louis, Poissy, Poissy, Ile de France, France. ⁵⁸Unit of Emergency and General Surgery, Guastalla Hospital, AUSL Reggio Emilia, Guastalla, Italy. ⁵⁹Chirurgie Digestive, Service de Chirurgie Vasculaire Et, Générale University Hospital of Bordeaux FR, Bordeaux, France. ⁶⁰Fondazione Policlinico A, Gemelli IRCCS, Rome, Italy. ⁶¹Department Head of Surgery, Infermi Hospital, Rimini, Italy. ⁶²Emergency and General Consultant Surgeon, Nuovo Ospedale "S. Stefano", Azienda ASL Toscana Centro, Prato, Italy. ⁶³Surgery Unit, Hospital Universitário Terezinha de Jesus, SUPREMA, Juiz de Fora, Brazil. ⁶⁴Department of Surgery, Faculty of Medicine and Health Sciences, Universiti Malaysia Sabah, Kota Kinabalu, Sabah, Malaysia. ⁶⁵Research Institute of General Surgery, Jinling Hospital, Medical School of Nanjing University, Nanjing, Jiangsu, China. ⁶⁶Hospital Umum Sarawak/Universiti Malaysia, Sarawak, Malaysia. ⁶⁷Hospital Sultanah Aminah, Johor Bahru, Malaysia. ⁶⁸Universiti Sains Islam Malaysia, Nilai, Malaysia. ⁶⁹General Surgery, University Hospitals, Sussex, UK. ⁷⁰Universidad de Sucre, Medicine Program, Clínica Santa María, Sincelejo, Colombia. ⁷¹First Department of Anaesthesiology and Intensive Therapy, Medical University of Lublin, Lublin, Poland. ⁷²International Fluid Academy, Lovenojoel, Belgium. ⁷³Postgraduate Institute of Medical Sciences, Rohtak, Haryana, India. ⁷⁴University of Calgary, Calgary, Canada. ⁷⁵Tameside and Glossop Integrated Care NHSFT, Ashton-under-Lyne, UK. ⁷⁶Department of General Surgery, Madonna del Soccorso Hospital - San Benedetto del Tronto, Italy, Italy. ⁷⁷Department of Surgery, General University Hospital of Patras, Rio, Greece. ⁷⁸Department of Surgical Science, Cagliari State University, Cagliari, Italy. ⁷⁹Hospital Angkatan Tentera Tuanku Mizan, Kuala Lumpur, Malaysia. ⁸⁰Department of General Surgery, King George's Medical University, Lucknow, UP, India. ⁸¹School of Medical Sciences & Hospital, Universiti Sains Malaysia, Kelantan, Malaysia. ⁸²University Hospital Careggi, Florence, Italy. ⁸³Department of Emergency Surgery, Cagliari University Hospital, Cagliari, Italy. ⁸⁴Fondazione Policlinico Universitario A. Gemelli IRCCS, Catholic University of Sacred Heart, Rome, Italy. ⁸⁵Department of General Surgery, Hüseyin Kemal Raşa, Anadolu Medical Center, Kocaeli, Turkey. ⁸⁶Division of General and Minimally Invasive Surgery, ASST Valle Olona, Busto Arsizio, Italy. ⁸⁷Emergency Surgery Unit and Trauma Team, Careggi University Hospital, Florence, Italy. ⁸⁸Plovdiv Medical University, Plovdiv, Bulgaria. ⁸⁹Department of Surgery, South Ural State Medical University, Chelyabinsk City, Russia. ⁹⁰Department of General Surgery, University of Bologna, Bologna, Italy. ⁹¹Fiona Stanley Hospital, Perth, WA, Australia. ⁹²Clinica Santa Maria, Sincelejo, Colombia. ⁹³Military Institute of Medicine, Warsaw, Poland. ⁹⁴Department of General Surgery, University Hospital of Heraklion, Crete, Greece. ⁹⁵St Vincent's University Hospital, Dublin, Ireland. ⁹⁶Department of Surgery, School of Medical Sciences and Hospital USM, Universiti Sains Malaysia, Georgetown, Malaysia.

Received: 21 February 2023 Accepted: 13 April 2023
Published online: 11 May 2023

References

- Kirkpatrick AW, Coccolini F, Ansaloni L, Roberts DJ, Tolonen M, McKee JL, et al. Closed or open after source control laparotomy for severe complicated intra-abdominal sepsis (the COOL trial): study protocol for a randomized controlled trial. *World J Emerg Surg.* 2018;13:26.
- Jawad I, Luksic I, Rafnsson SB. Assessing available information on the burden of sepsis: global estimates of incidence, prevalence and mortality. *J Glob Health.* 2012;2(1): 010404.
- Fleischmann C, Scherag A, Adhikari NK, Hartog CS, Tsaganos T, Schlattmann P, et al. Assessment of global incidence and mortality of hospital-treated sepsis. Current estimates and limitations. *Am J Respir Crit Care Med.* 2016;193(3):259–72.
- Angus DC, Linde-Zwirble WT, Lidicker J, Clermont G, Carillo J, Pinsky MR. Epidemiology of severe sepsis in the United States: analysis of incidence, outcomes, and associated costs of care. *Crit Care Med.* 2001;29:1303–10.
- Slade E, Tamber PS, Vincent JL. The surviving sepsis campaign: raising awareness to reduce mortality. *Crit Care.* 2003;7:1–2.
- Dellinger RP, Levy MM, Carlet JM, Bion J, Parker MM, Jaeschke R, et al. Surviving Sepsis Campaign: international guidelines for management of severe sepsis and septic shock: 2008. *Intensive Care Med.* 2008;34(1):17–60.
- Dellinger RP, Levy MM, Carlet JM, Bion J, Parker MM, Jaeschke R, et al. Surviving Sepsis Campaign: international guidelines for management of severe sepsis and septic shock: 2008. *Crit Care Med.* 2008;36(1):296–327.
- Singer M, Deutschman CS, Seymour CW, Shankar-Hari M, Annane D, Bauer M, et al. The third international consensus definitions for sepsis and septic shock (sepsis-3). *JAMA.* 2016;315(8):801–10.
- Sartelli M, Abu-Zidan FM, Catena F, Griffiths EA, Di Saverio S, Coimbra R, et al. Global validation of the WSES Sepsis Severity Score for patients with complicated intra-abdominal infections: a prospective multicentre study (WISS Study). *World J Emerg Surgery.* 2015;10:61.
- Szakmany T, Lundin RM, Sharif B, Ellis G, Morgan P, Kopczyńska M, et al. Sepsis prevalence and outcome on the general wards and emergency departments in wales: results of a multi-centre, observational, point prevalence study. *PLoS ONE.* 2016;11(12): e0167230.
- WHO Coronavirus (COVID-19) Dashboard: World Health Organization; 2022 [cited 2022 Dec 21 2022]. Available from: <https://covid19.who.int/data>.
- World Health Organization. Injuries and violence: World Health Organization; 2022 [Available from: <https://www.who.int/news-room/factsheets/detail/injuries-and-violence#:~:text=Key%20facts%201%20injuries%20E2%80%93%20both%20unintentional%20and,of%20all%20years%20lived%20with%20disability%20More%20items>].
- Torio CM, Moore BJ. National Inpatient Hospital Costs: The Most Expensive Conditions by Payer, 2013. Rockville, MD; 2016. Contract No: 27359025.
- Roberts DJ, Ball CG, Kirkpatrick AW. Increased pressure within the abdominal compartment: intra-abdominal hypertension and the abdominal compartment syndrome. *Curr Opin Crit Care.* 2016;22(2):174–85.
- Kirkpatrick AW, Hamilton DR, McKee JL, MacDonald B, Pelosi P, Ball CG, et al. Do we have the guts to go? The abdominal compartment, intra-abdominal hypertension, the human microbiome and exploration class space missions. *Can J Surg.* 2020;63(6):E581–93.
- Clements TW, Tolonen M, Ball CG, Kirkpatrick AW. Secondary peritonitis and intra-abdominal sepsis: an increasingly global disease in search of better systemic therapies. *Scand J Surg.* 2021;110(2):139–49.
- Alverdy JC, Hoyoju SK, Weigerinck M, Gilbert JA. The gut microbiome and the mechanism of surgical infection. *Br J Surg.* 2017;104(2):e14–23.
- Guyton K, Alverdy JC. The gut microbiota and gastrointestinal surgery. *Nat Rev Gastroenterol Hepatol.* 2017;14(1):43–54.
- Leligdowicz A, Dodek PM, Norena M, Wong H, Kumar A, Kumar A, et al. Association between source of infection and hospital mortality in patients who have septic shock. *Am J Respir Crit Care Med.* 2014;189(10):1204–13.
- Sartelli M, Abu-Zidan FM, Ansaloni L, Bala M, Beltran MA, Biffi WL, et al. The role of the open abdomen procedure in managing severe abdominal sepsis: WSES position paper. *World J Emerg Surg.* 2015;10:35.

21. Sartelli M, Catena F, Ansaloni L, Coccolini F, Corbella D, Moore EE, et al. Complicated intra-abdominal infections worldwide: the definitive data of the CIAOW Study. *World J Emerg Surg.* 2014;9:37.
22. Sartelli M, Catena F, Ansaloni L, Moore E, Malangoni M, Velmahos G, et al. Complicated intra-abdominal infections in a worldwide context: an observational prospective study (CIAOW Study). *World J Emerg Surg.* 2013;8(1):1.
23. Sartelli M, Viale P, Catena F, Ansaloni L, Moore E, Malangoni M, et al. 2013 WSES guidelines for management of intra-abdominal infections. *World J Emerg Surg.* 2013;8(1):3.
24. Tellor B, Skrupky LP, Symons W, High E, Micek ST, Mazuski JE. Inadequate source control and inappropriate antibiotics are key determinants of mortality in patients with intra-abdominal sepsis and associated bacteremia. *Surg Infect (Larchmt).* 2015;16(6):785–93.
25. van Ruler O, Mahler CW, Boer KR, Reuland EA, Gooszen HG, Opmeer BC, et al. Comparison of on-demand vs planned relaparotomy strategy in patients with severe peritonitis: a randomized trial. *JAMA, J Am Med Assoc.* 2007;298(8):865–72.
26. Nathens AB, Rotstein OD. Therapeutic options in peritonitis. *Surg Clin North Am.* 1994;74(3):677–92.
27. Bosscha K, van Vroonhoven TJ, van der Werken C. Surgical management of severe secondary peritonitis. *Br J Surg.* 1999;86(11):1371–7.
28. Leppaniemi A, Kimball EJ, De Laet I, Malbrain ML, Balogh ZJ, De Waele JJ. Management of abdominal sepsis—a paradigm shift? *Anaesthesiol Intensive Therapy.* 2015;47(4):400–8.
29. De Waele JJ. Abdominal sepsis. *Curr Infect Disease Rep.* 2016;18(8):23.
30. Lamme B, Boermeester MA, Reitsma JB, Mahler CW, Obertop H, Gouma DJ. Meta-analysis of relaparotomy for secondary peritonitis. *Br J Surg.* 2002;89(12):1516–24.
31. Opmeer BC, Boer KR, van Ruler O, Reitsma JB, Gooszen HG, de Graaf PW, et al. Costs of relaparotomy on-demand versus planned relaparotomy in patients with severe peritonitis: an economic evaluation within a randomized controlled trial. *Crit Care.* 2010;14(3):R97.
32. Kirkpatrick AW, Roberts DJ, De Waele J, Jaeschke R, Malbrain ML, De Keulenaer B, et al. Intra-abdominal hypertension and the abdominal compartment syndrome: updated consensus definitions and clinical practice guidelines from the World Society of the Abdominal Compartment Syndrome. *Intensive Care Med.* 2013;39(7):1190–206.
33. Opal SM, Dellinger RP, Vincent JL, Masur H, Angus DC. The next generation of sepsis clinical trial designs: what is next after the demise of recombinant human activated protein C? *Crit Care Med.* 2014;42(7):1714–21.
34. Gentile LF, Moldawer LL. HMGB1 as a therapeutic target for sepsis: it's all in the timing! *Expert Opin Ther Targets.* 2014;18(3):243–5.
35. Kirkpatrick AW, Roberts DJ, Faris PD, Ball CG, Kubek P, Tiruta C, et al. Active negative pressure peritoneal therapy after abbreviated laparotomy: the intraperitoneal vacuum randomized controlled trial. *Ann Surg.* 2015;262(1):38–46.
36. Cheatham ML, Demetriades D, Fabian TC, Kaplan MJ, Miles WS, Schreiber MA, et al. Prospective study examining clinical outcomes associated with a negative pressure wound therapy system and Barker's vacuum packing technique. *World J Surg.* 2013;37(9):2018–30.
37. Malig MS, Jenne CN, Ball CG, Roberts DJ, Xiao Z, Kirkpatrick AW. High mobility group box-1 protein and outcomes in critically ill surgical patients requiring open abdominal management. *Mediators Inflamm.* 2017;2017:6305387.
38. Khan A, Hsee L, Mathur S, Civil I. Damage-control laparotomy in non-trauma patients: review of indications and outcomes. *J Trauma Acute Care Surg.* 2013;75(3):365–8.
39. Bruns BR, Ahmad SA, O'Meara L, Tesoriero R, Lauerman M, Klyushnikova E, et al. Nontrauma open abdomens: a prospective observational study. *J Trauma Acute Care Surg.* 2016;80(4):631–6.
40. Goussous N, Jenkins DH, Zielinski MD. Primary fascial closure after damage control laparotomy: sepsis vs haemorrhage. *Injury.* 2014;45(1):151–5.
41. Coccolini F, Biffi W, Catena F, Ceresoli M, Chiara O, Cimbanassi S, et al. The open abdomen, indications, management and definitive closure. *World J Emerg Surg.* 2015;10:32.
42. Atema JJ, Gans SL, Boermeester MA. Systematic review and meta-analysis of the open abdomen and temporary abdominal closure techniques in non-trauma patients. *World J Surg.* 2015;39(4):912–25.
43. Quyn AJ, Johnston C, Hall D, Chambers A, Arapova N, Ogston S, et al. The open abdomen and temporary abdominal closure systems—historical evolution and systematic review. *Colorectal Dis.* 2012;14(8):e429–38.
44. Johnson JW, Gracias VH, Schwab CW, Reilly PM, Kauder DR, Shapiro MB, et al. Evolution in damage control for exsanguinating penetrating abdominal injury. *J Trauma.* 2001;51:261–71.
45. Harvin JA, Podbielski J, Vincent LE, Fox EE, Moore LJ, Cotton BA, et al. Damage control laparotomy trial: design, rationale and implementation of a randomized controlled trial. *Trauma Surg Acute Care Open.* 2017;2:1–5.
46. Hatch QM, Osterhout LM, Ashraf A, Podbielski J, Kozar RA, Wade CE, et al. Current use of damage-control laparotomy, closure rates, and predictors of early fascial closure at the first take-back. *J Trauma.* 2011;70(6):1429–36.
47. Harvin JA, Wray CJ, Steward J, Lawless RA, McNutt MK, Love JD, et al. Control the damage: morbidity and mortality after emergent trauma laparotomy. *Am J Surg.* 2016;212(1):34–9.
48. Zizzo M, Castro Ruiz C, Zanelli M, Bassi MC, Sanguedolce F, Asciani S, et al. Damage control surgery for the treatment of perforated acute colonic diverticulitis: a systematic review. *Medicine (Baltimore).* 2020;99(48):e23323.
49. Coccolini F, Montori G, Ceresoli M, Catena F, Moore EE, Ivatury R, et al. The role of open abdomen in non-trauma patient: WSES Consensus Paper. *World J Emerg Surg.* 2017;12:39.
50. Sartelli M, Catena F, Ansaloni L, Coccolini F, Griffiths EA, Abu-Zidan FM, et al. WSES Guidelines for the management of acute left sided colonic diverticulitis in the emergency setting. *World J Emerg Surg.* 2016;11:37.
51. Cheng Y, Wang K, Gong J, Liu Z, Gong J, Zeng Z, et al. Negative pressure wound therapy for managing the open abdomen in non-trauma patients. *Cochrane Database Syst Rev.* 2022;5:1013710.
52. Robledo FA, Luque-de-Leon E, Suarez R, Sanchez P, de-la-Fuente M, Vargas A, et al. Open versus closed management of the abdomen in the surgical treatment of severe secondary peritonitis: a randomized clinical trial. *Surg Infect.* 2007;8(1):63–72.
53. Roberts DJ, Zygun DA, Grendar J, Ball CG, Robertson HL, Ouellet JF, et al. Negative-pressure wound therapy for critically ill adults with open abdominal wounds: a systematic review. *J Trauma Acute Care Surg.* 2012;73(3):629–39.
54. Coccolini F, Ceresoli M, Kluger Y, Kirkpatrick A, Montori G, Salvetti F, et al. Open abdomen and entero-atmospheric fistulae: an interim analysis from the International Register of Open Abdomen (IROA). *Injury.* 2019;50(1):160–6.
55. Kubiak BD, Albert SP, Gatto LA, Snyder KP, Maier KG, Vieau CJ, et al. Peritoneal negative pressure therapy prevents multiple organ injury in a chronic porcine sepsis and ischemia/reperfusion model. *Shock.* 2010;34(5):525–34.
56. Emr B, Sadowsky D, Azhar N, Gatto LA, An G, Nieman G, et al. Removal of Inflammatory Ascites is Associated with Dynamic Modification of Local and Systemic Inflammation along with Prevention of Acute Lung Injury: In Vivo and In Silico Studies. *Shock.* 2014.
57. Roberts DJ, Jenne CN, Ball CG, Tiruta C, Leger C, Xiao Z, et al. Efficacy and safety of active negative pressure peritoneal therapy for reducing the systemic inflammatory response after damage control laparotomy (the Intra-peritoneal Vacuum Trial): study protocol for a randomized controlled trial. *Trials.* 2013;14:141.
58. Person B, Dorfman T, Bahouth H, Osman A, Assalia A, Kluger Y. Abbreviated emergency laparotomy in the non-trauma setting. *World J Emerg Surg.* 2009;4:41.
59. Huang J, Ren J, McKee JL, Kirkpatrick AW. 2023 updates on the closed or open after-source control for severe complicated intraabdominal sepsis (COOL) trial. *World J Surg Infect.* 2022;1(2):47–9.
60. Acosta S, Bjarnason T, Petersson U, Palsson B, Wanhainen A, Svensson M, et al. Multicentre prospective study of fascial closure rate after open abdomen with vacuum and mesh-mediated fascial traction. *Br J Surg.* 2011;98(5):735–43.
61. Mukhi AN, Minor S. Management of the open abdomen using combination therapy with ABRA and ABThera systems. *Can J Surg.* 2014;57(5):314–9.
62. Smith JW, Matheson PJ, Franklin GA, Harbrecht BG, Richardson JD, Garrison RN. Randomized controlled trial evaluating the efficacy of

- peritoneal resuscitation in the management of trauma patients undergoing damage control surgery. *J Am Coll Surg.* 2017;224(4):396–404.
63. Posadas-Calleja JG, Stelfox HT, Ferland A, Zuege DJ, Niven DJ, Berthiaume L, et al. Derivation of a PIRO score for prediction of mortality in surgical patients with intra-abdominal sepsis/severe sepsis/septic shock. *Am J Crit Care.* (in press).
 64. Tolonen M, Coccolini F, Ansaloni L, Sartelli M, Roberts DJ, McKee JL, et al. Getting the invite list right: A Discussion of sepsis severity scoring systems in severe complicated intra-abdominal sepsis and randomized trial inclusion criteria. *World J Emerg Surg.* (in press).
 65. Mahoney EJ, Bugaev N, Appelbaum R, Goldenberg-Sandau A, Baltazar GA, Posluszny J, et al. Management of the open abdomen: a systematic review with meta-analysis and practice management guideline from the Eastern Association for the Surgery of Trauma. *J Trauma Acute Care Surg.* 2022;93(3):e110–8.
 66. Campbell A, Chang M, Fabian T, Franz M, Kaplan M, Moore F, et al. Management of the open abdomen: from initial operation to definitive closure. *Am Surg.* 2009;75(11 Suppl):S1–22.
 67. Sharrock AE, Barker T, Yuen HM, Rickard R, Tai N. Management and closure of the open abdomen after damage control laparotomy for trauma. *A Syst Rev Meta-anal Injury.* 2016;47(2):296–306.
 68. Bauer M, Gerlach H, Vogelmann T, Preissing F, Stiefel J, Adam D. Mortality in sepsis and septic shock in Europe, North America and Australia between 2009 and 2019- results from a systematic review and meta-analysis. *Crit Care.* 2020;24(1):239.
 69. Bjorck M, Kirkpatrick AW, Cheatham M, Kaplan M, Leppaniemi A, De Waele JJ. Amended classification of the open abdomen. *Scand J Surg.* 2016;105(1):5–10.
 70. Bjarnason T, Montgomery A, Acosta S, Petersson U. Evaluation of the open abdomen classification system: a validity and reliability analysis. *World J Surg.* 2014;38(12):3112–24.
 71. Clavien PA, Vetter D, Staiger RD, Slankamenac K, Mehra T, Graf R, et al. The comprehensive complication index (CCI(R)): added value and clinical perspectives 3 years "Down the Line." *Ann Surg.* 2017;265(6):1045–50.
 72. Dindo D, Demartines N, Clavien PA. Classification of surgical complications: a new proposal with evaluation in a cohort of 6336 patients and results of a survey. *Ann Surg.* 2004;240(2):205–13.
 73. Ng-Kamstra JS, Rennert-May E, McKee J, Lundgren S, Manns B, Kirkpatrick AW. Protocol for a parallel economic evaluation of a trial comparing two surgical strategies in severe complicated intra-abdominal sepsis: the COOL-cost study. *World J Emerg Surg.* 2020;15(1):15.
 74. Kirkpatrick AW, Roberts DJ, Faris PD, Ball CG, Kubes P, Tiruta C, et al. Active negative pressure peritoneal therapy after abbreviated laparotomy: the intraperitoneal vacuum randomized controlled trial. *Ann Surg.* 2014.
 75. Briel M, Lane M, Montori VM, Bassler D, Glasziou P, Malaga G, et al. Stopping randomized trials early for benefit: a protocol of the Study Of Trial Policy Of Interim Truncation-2 (STOPIT-2). *Trials.* 2009;10:49.
 76. Bassler D, Briel M, Montori VM, Lane M, Glasziou P, Zhou Q, et al. Stopping randomized trials early for benefit and estimation of treatment effects: systematic review and meta-regression analysis. *JAMA.* 2010;303(12):1180–7.
 77. Bassler D, Montori VM, Briel M, Glasziou P, Guyatt G. Early stopping of randomized clinical trials for overt efficacy is problematic. *J Clin Epidemiol.* 2008;61(3):241–6.
 78. Freedman B. Equipoise and the ethics of clinical research. *N Engl J Med.* 1987;317(3):141–5.
 79. Doig CJ, Page SA, McKee JL, Moore EE, Abu-Zidan FM, Carroll R, et al. Ethical considerations in conducting surgical research in severe complicated intra-abdominal sepsis. *World J Emerg Surg.* 2019;14:39.
 80. Doig CJ, Page SA, McKee JL, Moore EE, Abu-Zidan FM, Carroll R, et al. Correction to: Ethical considerations in conducting surgical research in severe complicated intra-abdominal sepsis. *World J Emerg Surg.* 2019;14:47.
 81. Solomon MJ, Laxamana A, Devore L, McLeod RS. Randomized controlled trials in surgery. *Surgery.* 1994;115(6):707–12.
 82. Smithson M, McLeod MC, Chu DI, Kennedy G, Morris M, Chen H, et al. NIH funding of researchers in surgery: decreased career development awards over time. *J Surg Res.* 2021;266:6–12.
 83. Hu Y, Edwards BL, Brooks KD, Newhook TE, Slingluff CL Jr. Recent trends in National Institutes of Health funding for surgery: 2003 to 2013. *Am J Surg.* 2015;209(6):1083–9.
 84. McCulloch P, Feinberg J, Philippou Y, Koliass A, Kehoe S, Lancaster G, et al. Progress in clinical research in surgery and IDEAL. *Lancet.* 2018;392(10141):88–94.
 85. McCulloch P, Taylor I, Sasako M, Lovett B, Griffin D. Randomised trials in surgery: problems and possible solutions. *BMJ.* 2002;324(7351):1448–51.
 86. Horton R. Surgical research or comic opera: questions, but few answers. *Lancet.* 1996;347:984–5.
 87. Johnson M, Stokes RG, Arndt T. The Thalidomide Catastrophe: How it happened, who was responsible and why the search for justice continues after more than six decades. Cranbrook, Exeter: Onwards and Upwards Publishers; 2018. p. 2018.
 88. Kirkpatrick AW, Sugrue M, McKee JL, Pereira BM, Roberts DJ, De Waele JJ, et al. Update from the Abdominal Compartment Society (WSACS) on intra-abdominal hypertension and abdominal compartment syndrome: past, present, and future beyond Banff 2017. *Anaesthesiol Intensive Ther.* 2017.
 89. Cantle PM, Cotton BA. Balanced resuscitation in trauma management. *Surg Clin North Am.* 2017;97(5):999–1014.
 90. Chang R, Holcomb JB. Optimal fluid therapy for traumatic hemorrhagic shock. *Crit Care Clin.* 2017;33(1):15–36.
 91. Joseph B, Zangbar B, Pandit V, Verccruyse G, Aziz H, Kulvatyouyoun N, et al. The conjoint effect of reduced crystalloid administration and decreased damage-control laparotomy use in the development of abdominal compartment syndrome. *J Trauma Acute Care Surg.* 2014;76(2):457–61.
 92. Schreiber MA. The beginning of the end for damage control surgery. *Br J Surg.* 2012;99(Suppl 1):10–1.
 93. Malbrain ML, Marik PE, Witters I, Cordermans C, Kirkpatrick AW, Roberts DJ, et al. Fluid overload, de-resuscitation, and outcomes in critically ill or injured patients: a systematic review with suggestions for clinical practice. *Anaesthesiol Intensive Therapy.* 2014;46(5):361–80.
 94. Jacobs R, Wise RD, Myatchin I, Vanhonacker D, Minini A, Mekeirele M, et al. Fluid management, intra-abdominal hypertension and the abdominal compartment syndrome: a narrative review. *Life (Basel).* 2022;12(9).
 95. Loftus TJ, Jordan JR, Croft CA, Smith RS, Efron PA, Mohr AM, et al. Temporary abdominal closure for trauma and intra-abdominal sepsis: different patients, different outcomes. *J Trauma Acute Care Surg.* 2016.
 96. Vincent JL. Dear SIRS, I'm sorry to say that I don't like you. *Crit Care Med.* 1997;25(2):372–4.
 97. Shankar-Hari M, Deutschman CS, Singer M. Do we need a new definition of sepsis? *Intensive Care Med.* 2015;41(5):909–11.
 98. Kubiak BD, Albert SP, Gatto LA, Vieau CJ, Roy SK, Snyder KP, et al. A Clinically Applicable Porcine Model of Septic and Ischemia/Reperfusion-Induced Shock and Multiple Organ Injury. *J Surg Res.*
 99. Marshall JC. Inflammation, coagulopathy, and the pathogenesis of multiple organ dysfunction syndrome. *Crit Care Med.* 2001;29:599–106.
 100. Johnson D, Mayers I. Multiple organ dysfunction syndrome: a narrative review. *Can J Surg.* 2001;48:502–9.
 101. Fink MP, Delude RL. Epithelial barrier dysfunction: a unifying theme to explain the pathogenesis of multiple organ dysfunction at the cellular level. *Crit Care Clin.* 2005;21:177–96.
 102. Kirkpatrick AW, Xiao J, Jenne CN, Roberts DJ. Inflammatory mediators in intra-abdominal sepsis. In: Bassetti M, Martin-Loeches I, editors. *Sartelli M. Abdominal Sepsis. Hot Topics in Acute Care Surgery and Trauma.* Cham: Springer; 2018. p. 15–28.
 103. Kirkpatrick AW, Coccolini F, McDonald BR, Roberts DJ. Definition, Pathophysiology, and Pathobiology of Intra-Abdominal Hypertension and the Abdominal Compartment Syndrome. In: Coccolini F, Malbrain MLNG, Kirkpatrick AW, Gamberini E, editors. *Compartment Syndrome. Hot Topics in Acute Care Surgery and Trauma.* Cham: Springer; 2021. p. 51–62.
 104. Rongione AJ, Kusske AM, Ashley SW, Reber HA, McFadden DW. Interleukin-10 prevents early cytokine release in severe intraabdominal infection and sepsis. *J Surg Res.* 1997;70(2):107–12.
 105. Yao YM, Redl H, Bahrami S, Schlag G. The inflammatory basis of trauma/shock-associated multiple organ failure. *Inflamm Res.* 1998;47(5):201–10.

106. Wortel CH, van Deventer SJ, Aarden LA, Lygidakis NJ, Buller HR, Hoek FJ, et al. Interleukin-6 mediates host defense responses induced by abdominal surgery. *Surgery*. 1993;114(3):564–70.
107. Scheingraber S, Bauerfeind F, Bohme J, Dralle H. Limits of peritoneal cytokine measurements during abdominal lavage treatment for intraabdominal sepsis. *Am J Surg*. 2001;181(4):301–8.
108. van Berge Henegouwen MI, van der Poll T, van Deventer SJ, Gouma DJ. Peritoneal cytokine release after elective gastrointestinal surgery and postoperative complications. *Am J Surg*. 1998;175(4):311–6.
109. Jansson K, Redler B, Truedsson L, Magnuson A, Matthiessen P, Andersson M, et al. Intraperitoneal cytokine response after major surgery: higher postoperative intraperitoneal versus systemic cytokine levels suggest the gastrointestinal tract as the major source of the postoperative inflammatory reaction. *Am J Surg*. 2004;187(3):372–7.
110. Hendriks T, Bleichrodt RP, Lomme RM, De Man BM, van Goor H, Buyne OR. Peritoneal cytokines predict mortality after surgical treatment of secondary peritonitis in the rat. *J Am Coll Surg*. 2010;211(2):263–70.
111. Holzheimer RG, Schein M, Wittmann DH. Inflammatory response in peritoneal exudate and plasma of patients undergoing planned reoperation for severe secondary peritonitis. *Arch Surg*. 1995;130(12):1314–9; discussion 9–20.
112. Martineau L, Shek PN. Peritoneal cytokine concentrations and survival outcome in an experimental bacterial infusion model of peritonitis. *Crit Care Med*. 2000;28(3):788–94.
113. Marshall JC, Innes M. Intensive care unit management of intra-abdominal infection. *Crit Care Med*. 2003;31(8):2228–37.
114. Antonelli M, Fumagalli R, Cruz DN, Brienza N, Giunta F. PMX endotoxin removal in the clinical practice: results from the EUPHAS trial. *Contrib Nephrol*. 2010;167:83–90.
115. Nakamura M, Oda S, Sadahiro T, Hirayama Y, Watanabe E, Tateishi Y, et al. Treatment of severe sepsis and septic shock by CHDF using a PMMA membrane hemofilter as a cytokine modulator. *Contrib Nephrol*. 2010;166:73–82.
116. Ratanarat R, Brendolan A, Piccini P, Dan M, Salvatori G, Ricci Z, et al. Pulse high-volume haemofiltration for treatment of severe sepsis: effects on hemodynamics and survival. *Crit Care*. 2005;9(4):R294–302.
117. Caronna R, Benedetti M, Morelli A, Rocco M, Diana L, Prezioso G, et al. Clinical effects of laparotomy with perioperative continuous peritoneal lavage and postoperative hemofiltration in patients with severe acute pancreatitis. *World J Emerg Surg*. 2009;4:45.
118. Jennings WC, Wood CD, Guernsey JM. Continuous postoperative lavage in the treatment of peritoneal sepsis. *Dis Colon Rectum*. 1982;25(7):641–3.
119. DeWaele JJ, Hesse UJ, Patryn P, Decruyenaere J, de Hemptinne B. Postoperative lavage and on demand surgical intervention in the treatment of acute necrotizing pancreatitis. *Acta Chir Belg*. 2000;100(1):16–20.
120. Schwarz A, Bolke E, Peiper M, Schulte am Esch J, Steinbach G, van Griensven M, et al. Inflammatory peritoneal reaction after perforated appendicitis: continuous peritoneal lavage versus non lavage. *Eur J Med Res*. 2007;12(5):200–5.
121. Buanes TA, Andersen GP, Jacobsen U, Nygaard K. Perforated appendicitis with generalized peritonitis. Prospective, randomized evaluation of closed postoperative peritoneal lavage. *Eur J Surg*. 1991;157(4):277–9.
122. Hallerback B, Andersson C, Englund N, Glise H, Nihlberg A, Solhaug J, et al. A prospective randomized study of continuous peritoneal lavage postoperatively in the treatment of purulent peritonitis. *Surg Gynecol Obstet*. 1986;163(5):433–6.
123. Nakada TA, Oda S, Matsuda K, Sadahiro T, Nakamura M, Abe R, et al. Continuous hemodiafiltration with PMMA Hemofilter in the treatment of patients with septic shock. *Mol Med*. 2008;14(5–6):257–63.
124. Hoffmann JN, Faist E, Deppisch R, Hartl WH, Inthorn D. Hemofiltration in human sepsis: evidence for elimination of immunomodulatory substances. *Contrib Nephrol*. 1995;116:76–9.
125. Horner C, Schuster S, Plachky J, Hofer S, Martin E, Weigand MA. Hemofiltration and immune response in severe sepsis. *J Surg Res*. 2007;142(1):59–65.
126. Payen DM, Guilhot J, Launey Y, Lukaszewicz AC, Kaaki M, Veber B, et al. Early use of polymyxin B hemoperfusion in patients with septic shock due to peritonitis: a multicenter randomized control trial. *Intensive Care Med*. 2015;41(6):975–84.
127. Schadler D, Pausch C, Heise D, Meier-Hellmann A, Brederlau J, Weiler N, et al. The effect of a novel extracorporeal cytokine hemoabsorption device on IL-6 elimination in septic patients: a randomized controlled trial. *PLoS ONE*. 2017;12(10): e0187015.
128. Malbrain ML, Chiumello D, Pelosi P, Wilmer A, Brienza N, Malcagni V, et al. Prevalence of intra-abdominal hypertension in critically ill patients: a multicentre epidemiological study. *Intensive Care Med*. 2004;30:822–9.
129. Plantefeve G, Hellman R, Pajot O, Thirion M, Bleichner G, Mentec H. Abdominal compartment syndrome and intraabdominal sepsis. *Acta Clin Belg*. 2007;62:5240.
130. Balogh Z, McKinley BA, Cocanour CS, Kozar RA, Holcomb JB, Ware DN, et al. Secondary abdominal compartment syndrome is an elusive early complication of traumatic shock resuscitation. *Am J Surg*. 2002;184:538–44.
131. Regueira T, Bruhn A, Hasbun P, Aguirre M, Romero C, Llanos O, et al. Intra-abdominal hypertension: incidence and association with organ dysfunction during early septic shock. *J Crit Care*. 2008;23(4):461–7.
132. Regueira T, Hasbun P, Rebollo R, Galindo J, Aguirre M, Romero C, et al. Intra-abdominal hypertension in patients with septic shock. *Am Surg*. 2007;73(9):865–70.
133. Reintam A, Parm P, Kitus R, Kern H, Starkopf J. Intra-abdominal hypertension as a risk factor of death in patients with severe sepsis or septic shock. *Crit Care*. 2007;11:S130.
134. McBeth PB, Leger C, Ball CG, Ouelett JF, Tiruta C, Laupland KB, et al. Intra-abdominal hypertension and intra-abdominal sepsis: critical concepts and possibilities. *Int J Intensive Care*. 2011;19–26.
135. Kirkpatrick AW, Roberts DJ, De Waele J, Laupland K. Is intra-abdominal hypertension a missing factor that drives multiple organ dysfunction syndrome? *Crit Care*. 2014;18(2):124.
136. Cheng J, Wei Z, Liu X, Li X, Yuan Z, Zheng J, et al. The role of intestinal mucosa injury induced by intra-abdominal hypertension in the development of abdominal compartment syndrome and multiple organ dysfunction syndrome. *Crit Care*. (in press).
137. Leng Y, Zhang K, Fan J, Yi M, Ge Q, Chen L, et al. Effect of acute, slightly increased intra-abdominal pressure on intestinal permeability and oxidative stress in a rat model. *PLoS ONE*. 2014;9(10): e0109350.
138. Briel M, Elger BS, McLennan S, Schandelmaier S, von Elm E, Satalcar P. Exploring reasons for recruitment failure in clinical trials: a qualitative study with clinical trial stakeholders in Switzerland, Germany, and Canada. *Trials*. 2021;22(1):844.
139. Briel M, Olu KK, von Elm E, Kasenda B, Alturki R, Agarwal A, et al. A systematic review of discontinued trials suggested that most reasons for recruitment failure were preventable. *J Clin Epidemiol*. 2016;80:8–15.
140. Posadas-Calleja JG, Stelfox HT, Ferland A, Zuege DJ, Niven DJ, Berthiaume L, et al. Derivation of a PIRO score for prediction of mortality in surgical patients with intra-abdominal sepsis. *Am J Crit Care*. 2018;27(4):287–94.
141. Vincent JL, Moreno R, Takala J, Willatts S, De Me A, Bruining H, et al. The SOFA (Sepsis-related Organ Failure Assessment) score to describe organ dysfunction/failure. *Intensive Care Med*. 1996;22:707–10.
142. Charlson ME, Pompei P, Ales KL, Mackenzie CR. A new method of classifying prognostic comorbidity in longitudinal studies: development and validation. *J Chronic Dis*. 1987;40(5):373–83.

Publisher's Note

Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Ready to submit your research? Choose BMC and benefit from:

- fast, convenient online submission
- thorough peer review by experienced researchers in your field
- rapid publication on acceptance
- support for research data, including large and complex data types
- gold Open Access which fosters wider collaboration and increased citations
- maximum visibility for your research: over 100M website views per year

At BMC, research is always in progress.

Learn more biomedcentral.com/submissions

