Estimated oxygen extraction versus dynamic parameters of fluid-responsiveness for perioperative hemodynamic optimization of patients undergoing non-cardiac surgery: a non-inferiority randomized controlled trial

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Abstract

Background: Goal directed therapy (GDT) is able to improve mortality and reduce complications in selected high-risk patients undergoing major surgery. The aim of this study is to compare two different strategies of perioperative hemodynamic optimization: one based on optimization of preload using dynamic parameters of fluid-responsiveness and the other one based on estimated oxygen extraction rate (O₂ER) as target of hemodynamic manipulation.

Methods: This is a multicenter randomized controlled trial. Adult patients undergoing elective major open abdominal surgery will be allocated to receive a protocol based on dynamic parameters of fluid-responsiveness or a protocol based on estimated O₂ER. The hemodynamic optimization will be continued for 6 h postoperatively. The primary outcome is difference in overall postoperative complications rate between the two protocol groups. Fluids administered, fluid balance, utilization of vasoactive drugs, hospital length of stay and mortality at 28 day will also be assessed.

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**Background**

Any surgical intervention is a trauma for the organism and a stress response is activated to cope the external insult. This stress response is responsible of an increase in oxygen consumption. Shoemaker et al. [1] showed that oxygen debt starts intraoperatively in high-risk surgical patients. If patient is not able to overcome the deficit in oxygen consumption (VO\textsubscript{2}) during the first hours postoperatively, he/she will go toward complications (in case of delay to meet metabolic demand) or death (in case of persistent VO\textsubscript{2} deficit). The same authors showed that the incidence of organ failure and mortality were reduced when oxygen debt was rapidly compensated using a protocol of hemodynamic optimization aiming to reach the same hemodynamic targets recorded in survived patients [1]. Therefore, several protocols have been developed to optimize hemodynamic parameters with the aim to reduce tissue hypoperfusion coming from maldistribution or inadequate perfusion and meet the increased metabolic need as soon as possible.

Every patient that probably will not be able to face the surgical stress himself might benefit from modulation of hemodynamic parameters. Actually, goal directed therapy (GDT) is able to improve survival only in high-risk surgical patients [2]. On the other hand, the reduction of complications rate has been shown also in intermediate-risk population [3–7].

Originally, hemodynamic optimization protocols were developed to reach supranormal value for cardiac output (CO) and oxygen delivery (DO\textsubscript{2}) [1]. DO\textsubscript{2} is locally regulated according to tissue metabolism and an adequate global DO\textsubscript{2} may coexist with local hypoperfusion. An inadequate regional DO\textsubscript{2} is responsible for an increase of oxygen extraction, a reduction of mixed venous oxygen saturation (SvO\textsubscript{2}) and central venous oxygen saturation (ScvO\textsubscript{2}) as its surrogate and finally an increase of lactate. Based on the concept that oxygen extraction rate (O\textsubscript{2}ER) reflects the balance between DO\textsubscript{2} and VO\textsubscript{2} a GDT protocol based on O\textsubscript{2}ER estimation (O\textsubscript{2}ERE) calculated as (SaO\textsubscript{2}-ScvO\textsubscript{2})/SaO\textsubscript{2} (where SaO\textsubscript{2} is arterial oxygen saturation) has been proposed showing a significantly lower number of organ failure postoperatively compared with control group [8].

The major determinants of DO\textsubscript{2} are CO, hemoglobin (Hb) level and SaO\textsubscript{2}. An inadequate CO may be optimized using fluids as first-line therapy and then inotropes. In mechanically ventilated patients, heart-lung interaction is useful to recognize in which portion of the Frank-Starling curve the heart of the patient is working and then if CO is able to rise after fluid administration aimed to increase preload. Several parameters based on mini-invasive monitor systems are available to assess fluid responsiveness such as pulse pressure variation (PPV) and stroke volume variation (SVV) [9]. Optimization of functional parameters allows titration of fluid administration and personalization of therapy for each patients [10]. Thus, several perioperative GDT protocols are based on fluids administration as long as PPV or SVV is above a pre-defined cut-off, aiming to preload maximization.

Although positive effects of perioperative GDT have been clearly demonstrated, currently studies comparing different strategies of hemodynamic optimization are lacking. To date, only one trial investigated the effect of a protocol based on O\textsubscript{2}ERE [8], whereas many studies were based on functional parameters of fluid-responsiveness (PPV or SVV). Thus, we believe useful a trial comparing different strategies of hemodynamic optimization.

**Methods/design**

The protocol adheres to the Standard Protocol Items: Recommendations for Interventional Trials (SPIRIT) 2013 guidelines [11].

**Study aim**

The aim of the study is to compare a perioperative hemodynamic optimization protocol based on O\textsubscript{2}ERE referred to a protocol based on normalization of dynamic parameters of fluid-responsiveness (PPV or SVV).
Objectives
The primary objective is to evaluate the difference in developing overall complications during hospital stay between the two group of patients managed with two GDT protocols using a non-inferiority approach.

The secondary outcomes are the difference between the total amount of fluid administered, the total fluid balance, the needs of vasopressor/inotropes, the hospital length of stay and the mortality at day 28.

Study design and setting
This is a multicenter randomized controlled trial. As both PGDT protocols showed to reduce postoperative complications compared to standard therapy and they are routinely applied for this kind of patients in common clinical practice [8, 12], a non-inferiority trial will be performed as the effect on primary outcome is expected to be maintained in both groups and it does not seem reasonable for ethical reason to deprive a study group of an established strategy for hemodynamic management during perioperative period. The study will be performed in 15 centers in Italy (Additional file 1).

Subjects and study population

Inclusion criteria

- Adult patients (age > 18 years) undergoing to general anesthesia and mechanical ventilation for elective major open abdominal surgery (gastrointestinal, urologic, gynecologic and vascular surgery)
- Expected duration of surgical procedure higher than 120 min
- ASA-PS Classification II-III-IV
- Planned postoperative ICU/HDU admission

Exclusion criteria

- Pregnancy
- Cardiac arrhythmia
- Non-correctable arterial curve alterations
- Undergoing palliative surgery
- Denial of consent to participate

Randomization

After screening for inclusion and exclusion criteria, patients suitable to be enrolled will be allocated with 1:1 ratio to randomly receive one of the two protocol of perioperative hemodynamic optimization. A computer-based system will be used to create a block randomization list (block size of 6). The allocation sequence concealment will be guaranteed by sealed opaque envelopes. Patients and researchers that perform postoperative follow up will be blinded to the allocated protocol.

General care and procedures

Patients undergoing elective non-cardiac surgery will be screened to participate to the study. They will be enrolled and allocated to one of the two groups if they meet all inclusion criteria and none of exclusion ones. All patients will be undergone to general anesthesia and mechanically ventilated with a tidal volume of 8 ml/kg and a positive end expiratory pressure (PEEP) of 5 cmH2O. FiO2 will be set at discretion of attending anesthesiologist to maintain normoxia. An arterial cannula will be used for continuous arterial monitoring. The arterial transducer will be zeroed at the level of the fourth intercostal space in the middle axillar line. The square-wave test will be used to assess the presence of over-/under-dumping on the arterial signal. A CO monitoring system will be used to record hemodynamic parameters. A central venous catheter will be placed after induction of anesthesia. A balanced crystalloid solution at the rate of 1 ml/kg/h will be administered intraoperatively. Any further fluid bolus will be administered according to the protocol in which the patient is allocated. Balanced crystalloid solutions are the only fluids permitted. Packed red blood cells will be transfused to maintain Hb ≥ 8 g/dl, at discretion of attending anesthesiologist. Mean arterial pressure will be maintained between 60 and 100 mmHg modulating anesthetic drugs on surgical stimulus or using vasopressor/vasodilator drugs, at discretion of attending anesthesiologist.

Protocol a

Patients allocated to protocol A will received a GDT protocol based on dynamic parameters of fluid responsiveness for optimization of fluid status (Fig. 1). Dobutamine will be used with discretion of the attending anesthesiologist as clinically needed. The protocol will be continued postoperatively up to 6 h when dynamic parameters remain applicable (controlled mechanical ventilation, absence of arrhythmia, etc.). On the other hand, fluid challenge technique with 250 ml of crystalloid over 5 min will be used for fluid optimization considering 10% increase in CO as positive response.

PPV is calculated as:

$$PPV(\%) = \frac{PPmax - PPmin}{(PPmax + PPmin)/2} \times 100$$

where PP is pulse pressure.

SVV is calculated as:

$$SVV(\%) = \frac{SVmax - SVmin}{(SVmax + SVmin)/2} \times 100$$

where SV is stroke volume.
Protocol B
Patients allocated to protocol B will received a GDT protocol based on O\textsubscript{2}ER\textsubscript{e} intraoperatively and for the first 6 h postoperatively (Fig. 2).

O\textsubscript{2}ER\textsubscript{e} is estimated at the start of the surgery and then hourly as:

\[ O2ER_e = \frac{SaO_2 - ScvO_2}{SaO_2} \]

where SaO\textsubscript{2} is arterial oxygen saturation and ScvO\textsubscript{2} is central venous oxygen saturation.

Data collection
At randomization, demographic and clinical data will be recorded:

- Age, sex, weight, height
- Admission diagnosis
- Comorbidities (e.g. diabetes mellitus, chronic kidney injury, chronic obstructive pulmonary disease, cardiac dysfunction, etc.)

During perioperative period, main clinical parameters will be recorded:

- Ventilation modality and parameters (spontaneous, assisted/controlled, tidal volume, respiratory rate, airway pressures, FiO\textsubscript{2})
- Anesthetic/analgesic drugs
- Hemodynamic parameters (blood pressure, heart rate, CO, SV, SVR, PPV, SVV)
- Arterial blood gas
- Central venous blood gas
- Fluids (crystalloids and colloids), transfusion and fluid balance
- Inotrope/vasoactive drugs
- Daily SOFA score

Finally, the following outcomes will be recorded:

- Complication during hospital stay (Table 1)
- Hospital length of stay
- Mortality at day 28

Severity of postoperative complications will be defined according to Clavien-Dindo classification [13]. Complication grades I and II will be considered as minor complications while grades III and IV will be considered as major ones.

The Case Report Forms (CRFs) will not bear the participant’s name or other directly identifiable data. The participant’s trial Identification Number (ID) only, will be used for identification.

The flow diagram according to CONSORT guidelines [14] is provided as Fig. 3. Figure 4 shows an overview of all outcome measures in accordance with SPIRIT guidelines [11].

Adverse events
All adverse events (AE) will be recorded in the hospital notes in the first instance. A record of all AEs, whether related or unrelated to the treatment will also be kept in the CRF. If the Investigator suspects that the disease or condition has progressed faster due to the intervention, then he will report this as an unexpected adverse event to the sponsor. Only deaths that are assessed to be
caused by the trial intervention will be reported to the Sponsor. This report will be immediate.

Considering the safety profile of the interventions applied in this study (PGDT protocols that showed a benefit to reduce postoperative complication rate without significant adverse effects), we believe that institution of a safety committee is not mandatory.

Sample size calculation
From literature analysis, we estimate a complication rate for control and experimental group of 28 and 35% respectively [2, 15]. With a non-inferiority margin of 10%, 184 patients (92 for each group) are needed to have a power of 80% and an alfa error of 0.05 [16]. Considering a 10% losing in follow-up, we enroll 200 patients (100 for each group).

Statistical analysis
Analysis will be performed on intention-to-treat basis.

Data will be checked for normal distribution using Kolmogorov-Smirnov test and presented as mean and standard deviation or as median and interquartile range as appropriate.

The primary end point is the difference of postoperative complication rate between the two group of patients. It will be evaluated using Fisher’s exact test or Pearson’s Chi square test as appropriate. Non-inferiority of experimental protocol respect to control group will be confirmed if the difference (including 95% confidence interval) will be lower than non-inferiority margin established at 10%. The 10% margin was chosen because it represents a two-thirds proportion of lower expected beneficial effect of PGDT (considering higher 95%CI for RR of 0.85) to reduce postoperative complications in comparison with conventional fluid therapy (RR 0.76 [CI95% 0.68–0.85]) [17].

Student t test or Mann–Whitney U test will be used to analyzed secondary outcomes as appropriate. Mortality between groups will be analyzed using the Kaplan-Meier method. The Log Rank test will be used to evaluate the statistical significance. Cox regression model using forward selection method for entering explanatory variables will be used to perform multivariate analysis. The age, type of surgery, ASA-PS classification and comorbidities will be considered as variables. A p value of F-test < 0.05 will be considered for variable inclusion. The amount of multicollinearity in the model will be estimated by the variance inflation factor (VIF). A VIF higher than 4 will be considered a sign of multicollinearity. Predicted $R^2$ will be used to assess overfitting.

An interim analysis will be performed by independent statistician when 50% of information will be available (100 patients recruited). To monitor for harm or futility, the one-sided $p$ value is calculated for testing the

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**Table 1** Complications assessed during postoperative period

| Complication                              | Other
|-------------------------------------------|----------------------------------|
| Acute myocardial ischemia/infarction      | Multiorgan failure               |
| Cardiac arrhythmia                        | Delirium/psychosis               |
| Cardiac/respiratory arrest                | Urinary tract infection          |
| Cardiogenic pulmonary edema               | Bacteremia                       |
| Pulmonary embolism                        | Surgical site infection          |
| Acute respiratory distress syndrome       | Nosocomial pneumonia             |
| Gastrointestinal bleeding                 | Unknown origin infection         |
| Intestinal infarction                     | Stroke                           |
| Anastomotic leakage                      | Post-operative hemorrhage        |
| Paralytic ileum                           | Surgical re-intervention          |
| Acute kidney injury                       |                                  |

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**Fig. 2** Protocol B: Hemodynamic optimization protocol based on estimated oxygen extraction rate. CVP: central venous pressure; O$_2$ERE: oxygen extraction rate estimate; PPV: pulse pressure variation; SVV: stroke volume variation
hypothesis HR = 1 versus the alternative HR > 1 (meaning the experimental treatment is doing worse than the standard treatment). If the \( p \) value is < 0.0394 at a monitoring time, then the trial would stop with the conclusion that non-inferiority cannot be claimed [18, 19]. Stopping rule for efficacy has not been defined [20]. For each statistical test, a \( p \) value < 0.05 is considered statistically significant.

**Discussion**
The role of perioperative hemodynamic optimization has been clearly shown by literature and the current spreading of mini-invasive monitoring systems able to estimate CO permits an easier application of these protocols than occurring in the past when only invasive techniques (e.g. pulmonary artery catheter) were available. As a predefined target of CO or DO\(_2\) seems to be not adequate for every patient, a personalized therapy is likely more appropriate. Following this concept, dynamic parameters of fluid-responsiveness (PPV and SVV) allow to titrate fluid administration aiming an increase of CO as a consequence of preload optimization but avoiding fluid overload. This approach has the advantage of personalized fluid therapy, but it does not consider if CO is adequate or not because preload optimization occurs independently from parameters of tissue perfusion and oxygen consumption. A protocol based on O\(_2\)ER considers this second important aspect. An O\(_2\)ER higher than a predefined cutoff is reflecting an inadequate DO\(_2\) to tissue and consequently an inadequate CO. This condition, but not fluid-responsiveness per se, is the trigger to start hemodynamic optimization. Moreover, in this
contest we believe that CVP may be useful to limit fluid administration independently from PPV or SVV value. In fact, CVP depends on several conditions including preload status but also on cardiac function and ventricular compliance. Thus, this trial aims to demonstrate that this latest strategy of perioperative hemodynamic optimization is not inferior to the former in term of postoperative complications, but it may be responsible for a lower fluid administration. We decided to conduct a non-inferiority study because PGDT is routinely applied in this kind of patients due to the demonstrated benefits on outcome. Thus, small difference in reduction of postoperative complication rate between two different PDGT protocols seems reasonable.

Methodologically, the main limitation of this trial is the inability to blindly administer the interventions. However, this point is not clinically resolvable in this research field. Moreover, if participation of several centers may increase enrollment rate and may potentially extend the applicability of results, on the other hands too few patients per center may increases heterogeneity.

**Trial status**
Protocol version 2.0 of 26.06.2019. At the date of manuscript submission, the patients’ recruitment process is ongoing. It is expected that the recruitment will be complete within December 2021. The results of this study have not already been published or submitted to any journal.

**Supplementary information**
Supplementary information accompanies this paper at https://doi.org/10.1186/s12871-020-01011-z.

**Additional file 1.** Participating centers list.

**Abbreviations**
ASA-PS: American Society of Anesthesiologists Physical Status; CO: Cardiac output; DO₂: Oxygen delivery; GDT: Goal directed therapy; Hb: Hemoglobin; HDU: High dependency unit; ICU: Intensive care unit; O₂ER: Oxygen
The authors declare that they have no competing interests. The study has Competing interests Not applicable.

Consent for publication
participate and they can ask the elimination of acquired data.

patients. The participation to the study is free and patients can ask for the consent. The consent will be asked to legal representative for interdicted the trial. Patients will be enrolled after the subscription of written informed approval before the start of enrollment.

Marche has been approved by local ethical committee Ethics approval and consent to participate

The datasets used and/or analyzed during the current study are available Ethics approval and consent to participate

Funding
This trial will be conducted with no external funding.

Availability of data and materials
The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Ethics approval and consent to participate
This trial will be conducted according to Good Clinical Practice, Helsinki declaration and national regulation for clinical trial conduction. This protocol has been approved by local ethical committee “Comitato Etico Regione Marche” (Prot. 2019 141, Prot. 2019 141/5682) for the coordinator center. The satellite centers will submit the protocol to their Ethical Committee for approval before the start of enrollment.

Potentially eligible patients will receive printed and verbal information about the trial. Patients will be enrolled after the subscription of written informed consent. The consent will be asked to legal representative for interdicted patients. The participation to the study is free and patients can ask to abandon the study at any time. They can withdraw the consent to participate and they can ask the elimination of acquired data.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests. The study has not received funding/assistance from any commercial organization.

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