Ventilation strategies with different inhaled Oxygen concentration during Cardiopulmonary Bypass in cardiac surgery (VONTCPB): study protocol for a randomized controlled trial

Meng-Qiu Zhang, Yu-Qi Liao, Hong Yu, Xue-Fei Li, Liang Feng, Xiao-Yun Yang* and Hai Yu*

Abstract

Background: There is no consensus on the ventilation management during cardiopulmonary bypass (CPB), and the anesthesiologists or the surgeons usually ventilate the lungs with different ventilation strategies or keep them static. Better outcomes are more likely to occur when the ventilation is administered during CPB according to the existing literatures. However, the use of high fraction of inspired oxygen (FiO2) is debatable in cardiac surgery. And the potential effects of strategies combining low tidal volume (VT) ventilation with different FiO2 during CPB on postoperative pulmonary complications (PPCs) are unclear.

Design: The VONTCPB trial is a single-center, prospective, double-blinded, randomized, controlled trial. We are going to recruit total 420 elective cardiac surgery patients with median sternotomy under CPB, who will be equally randomized into three different ventilation strategy groups: NoV, LOV and HOV. (1) The NoV group receives no mechanical ventilation during CPB; (2) the LOV group receives a low VT of 3-4 ml/kg of ideal body weight (IBW) with the respiratory rate (RR) of 10–12 acts/min, and the positive end-expiratory pressure (PEEP) of 5–8 cmH2O during CPB; the FiO2 is 30%; (3) the HOV group receives a low VT of 3-4 ml/kg of IBW with the RR of 10–12 acts/min, and the PEEP of 5–8 cmH2O during CPB; the FiO2 is 80%. The primary endpoints are the incidence of the composite of PPCs and the PPCs score. The secondary endpoints refer to the incidence of the oxygenation index (PaO2/FiO2 ratio) < 300 mmHg at three time points (the moment arriving in the ICU, 6 and 12 h after arrival in the ICU), the surgical incision healing grade, the intubation time, the stay of ICU, the length of hospital stay, and mortality at 30 days after the surgery.

Discussion: The VONTCPB trial is the first study to assess the effects of strategies combining low tidal volume (VT) ventilation with different FiO2 during CPB on patients’ outcomes.

Trial registration: ChiCTR1800015261. Registered on 20 March 2018.

Keywords: Cardiopulmonary bypass, Low tidal volume, Mechanical ventilation, Oxygen concentration, Postoperative pulmonary complications
Background
Surgical and anesthesia techniques of heart surgery have been greatly improving. However, the postoperative pulmonary complications (PPCs) are still the main complications after surgery, resulting in increasing mortality in cardiac surgery with the incidence of 25% [1]. The PPCs have the forms of respiratory failure, respiratory infection, pleural effusion resulting in thoracentesis, atelectasis, pneumothorax, bronchospasm, aspiration pneumonitis [2]. These complications extend the length of mechanical ventilation and intensive care unit (ICU) stay, indicating worse prognosis [3, 4].

The etiologies of the PPCs are quite complicated. Lungs are isolated from the artificial extracorporeal circulation, which results in blood supply to the lungs only from the bronchial arteries. While two sets of independent circulation systems composing the blood delivery system of the lungs, the bronchial arteries supply only about 3–5% of the pulmonary blood flow system, which may be reduced to one-tenth of the original upon experimental conditions of CPB in pig model. Those reduced blood flow may be difficult to meet the physiological metabolic needs of the lungs [5]. The tissue of the lungs is also very loose, vascular fluid can be easily transferred to the interstitial tissue under pathologic conditions. Those factors, in addition to the strikes of mechanical ventilation during CPB, and finally we could make a conclusion of optimal ventilation mode during CPB on the outcomes in heart surgery patients [12–27]. All these RCTs have their own limits. Though the evidence is weak, it still suggests that maintaining mechanical ventilation during CPB is beneficial to lung protection.

The recent meta-analysis made by our research team shows that the ventilation of the resting lung during CPB is indeed effective in improving the oxygenation (PaO2/FiO2 ratio) and gas exchange. This approach can also reduce the alveolar-arterial oxygen partial pressure difference and the intrapulmonary shunt. But it is unclear of its effect on the long-term prognosis and outcomes for patients. The authors pointed that different FiO2 and other respiratory parameters settings may impact the long-term outcomes [28]. An earlier meta-analysis summarized the publications of different ventilation strategies during CPB. Using continuous positive airway pressure (CPAP) and lung recruitment maneuvers can increase oxygenation immediately after CPB to a certain extent, but there is no evidence of a sustained effect [29]. A just published systematic review and meta-analysis concluded that CPAP during CPB may improve the oxygenation and gas exchange while ventilation did not. The authors mentioned that the FiO2 had been ignored for a long period [30]. Heinrichs et al. [31] reviewed the RCTs to investigate the impact that hyperoxia has on postoperative organ dysfunction, length of stay, and mortality during adult cardiac surgery. They concluded that the impact of hyperoxia on these outcomes seems to be minimal because of the high heterogeneity of include studies. Similarly, McGuinness et al. [32] failed to demonstrate any difference in AKI, markers of organ damage, or length of stay by comparing the avoidance of arterial hyperoxemia versus usual care during CPB.

There is no generally accepted ventilation management model due to weak evidence of heterogeneity researches [10, 33, 34]. Therefore, we designed a prospective RCT combining the different regimens of ventilation settings during CPB with low or high FiO2 to explore the optimal ventilation mode during CPB.

Objectives
We designed this RCT to compare the clinical effects of three mechanical ventilation strategies on the total incidence of PPCs during postoperative hospitalization. We chose the incidence of composite PPCs as the primary outcome because it is a hard parameter. While in the past many studies only focused on observing biochemical indicators or indirect parameters. You could never tell whether those indicators in the blood have sustained influences in the long run or not. It could be just a transient change after surgery. We hope we would reach a conclusion of optimal ventilation management during CPB, and finally we could make a difference in the clinical practice.
Methods/design
The VONTCPB study is a single-center, double-blinded, three-armed, randomized controlled trial comparing three mechanical ventilation strategies (NoV, LOV and HOV) during CPB in adult patients undergoing cardiac surgery. The protocol structure is written according to the Consolidated Standards of Reporting Trials (CONSORT) 2010 Statement guidelines and follows the Standard Protocol Items: Recommendations for Interventional Trials (SPIRIT) Statement. The SPIRIT checklist can be found in Additional file 1. The SPIRIT figure is illustrated in Fig. 1.

Ethical approval and trial registration
This clinical study protocol has been approved by the Biomedical Research Ethics Committee of West China Hospital of Sichuan University on 12 March 2018 (Approval number 2017 (400)) and registered in the Chinese Clinical Trials Registry (ChiCTR) (www.chictr.org.cn) with the registration number ChiCTR1800015261 on 19 March 2018.

Participants
The patients included in the trial must be more than 18 years old and undergo elective cardiac surgery with median sternotomy under CPB. Simultaneous other non-cardiac surgery procedure is excluded. The inclusion and exclusion criteria are shown in Table 1.

Endpoints
The primary endpoints are the incidence of a composite of PPCs and the PPCs score during hospital stay [35–37]. The detailed definitions of PPCs are shown in Table 2. The score is presented in Table 3. Secondary endpoints are listed as follows: the incidence of oxygen index < 300 mmHg at three
time points (the moment arriving in the ICU, 6 and 12 h after arrival in the ICU), the surgical incision healing grade, the intubation time, the length of ICU stay and hospital stay, and postoperative 30-day mortality rate.

Randomization/Blinding
The random number table was generated by Microsoft Excel in a 1:1:1 treatment ratio. The investigator who is responsible for grouping visits the patients the day before surgery and gets the consents. Once the patient is qualified, he determines the serial number and random number of the patient and writes on the Case Report Form (CRF). He will also inform the anesthesia care providers who do not participate in the study the treatment of the patient. The patient, the data collector and the intensivist are not aware of the treatment information in the whole process. Any intraoperative event (such as pleura incision) or deviation from the protocol is recorded on the CRF.

Intervention
The participants will be randomly divided into the following three groups (Fig. 2):

1) The NoV arm: no ventilation during CPB.
2) The LOV arm: participants will receive a low V\textsubscript{T} of 3–4 ml/kg of IBW with the RR of 10–12 acts/min, and the PEEP of 5–8 cmH\textsubscript{2}O during CPB; the FiO\textsubscript{2} is 30% with a flow of 2 L/min.
3) The HOV arm: participants will receive a low V\textsubscript{T} of 3–4 ml/kg of IBW with the RR of 10–12 acts/min, and the PEEP of 5–8 cmH\textsubscript{2}O during CPB; the FiO\textsubscript{2} is 80% with a flow of 2 L/min.

FiO\textsubscript{2} 80% will be applied during oxygenation for denitrogenation 5 min before endotracheal intubation. The lung protective ventilation strategy \cite{33,38} will be applied in the participants before and after CPB: V\textsubscript{T} = 6–8 ml/kg IBW, PEEP = 5–8 cmH\textsubscript{2}O, inspiration/expiration time ratio(I:E) = 1:2, RR = 10–12 to maintain end-tidal carbon dioxide partial pressure (P\textsubscript{ET}CO\textsubscript{2})

Table 1 Eligibility criteria

| Inclusion criteria                        | Exclusion criteria                              |
|-------------------------------------------|-------------------------------------------------|
| • Age ≥ 18 years                          | • Pregnant women                                |
| • Signed informed consent                 | • Heart transplantation,                        |
| • Elective cardiac surgery patients with  | • Pulmonary thromboendarterectomy               |
|   median sternotomy under CPB.            | • Aortic arch and other deep                    |
|                                            |   hypothermic circulatory arrest surgery        |
|                                            | • Long-term use of hormone therapy              |
|                                            |   before surgery (≥23 months)                   |
|                                            | • Participated in other similar researches       |
|                                            |   3 months ago                                  |

Table 2 Definitions of post-operative pulmonary complications*

| Complication                | Definition                                                                                                                                 |
|-----------------------------|-------------------------------------------------------------------------------------------------------------------------------------------|
| Respiratory failure         | At least one of the following criteria after surgery:                                                                                    |
|                             | • SpO\textsubscript{2} <90% and requiring oxygen therapy                                                                                   |
|                             | • PaO\textsubscript{2}/FiO\textsubscript{2} <300                                                                                            |
|                             | • PaO\textsubscript{2} <60 mmHg                                                                                                           |
| Respiratory infection       | Postoperative antibiotic therapy administered for suspected respiratory infection with at least one of the following criteria:              |
|                             | • New or changed sputum                                                                                                                   |
|                             | • New or changed lung opacity on x-ray                                                                                                      |
|                             | • Fever                                                                                                                                   |
|                             | • Leukocyte count >12,000/mL                                                                                                               |
| Pleural effusion            | Evidence at chest x-ray of:                                                                                                               |
|                             | • Blunting of the costophrenic sinus                                                                                                       |
|                             | • In upright position: loss of the sharp silhouette of the ipsilateral hemidiaphragm                                                      |
|                             | • In supine position: hazy opacity in 1 hemithorax with preserved vascular shadows                                                        |
|                             | • Displacement of adjacent anatomic structures                                                                                             |
| Atelectasis                 | Evidence at chest x-ray of:                                                                                                               |
|                             | • New parenchymal opacification                                                                                                            |
|                             | • Shift of the median structures (mediastinum, hilum or hemidiaphragm) toward the affected area                                             |
|                             | • Compensatory overinflation of contralateral lung                                                                                          |
| Aspiration pneumonitis      | Inhalation of gastric content in the perioperative period with subsequent acute lung injury                                               |
| Broncho spasm               | New expiratory wheezing responsive to treatment with bronchodilators                                                                      |
| Pneumothorax                | At chest x-ray, presence of air within pleural space, with no vascular bed surrounding the visceral pleura                                     |

*Adapted from Bignami et al. \cite{2}
Table 3 Postoperative pulmonary complications score

| Grade | Definition |
|-------|------------|
| Grade 1 | Cough, dry |
| Grade 1 | Bronchospasm: new wheezing or pre-existent wheezing resulting in a change in therapy |
| Grade 1 | Hypoxemia (SpO2 ≤ 90%) at room air |
| Grade 1 | Atelectasis: gross radiological confirmation (concordance of 2 independent experts) plus either temperature > 37.5 °C or abnormal lung findings |
| Grade 1 | Hypercarbia (PaCO2 > 50 mmHg), requiring treatment |
| Grade 2 | Pleural effusion, resulting in thoracentesis |
| Grade 2 | Pneumonia: radiological evidence (concordance of 2 independent experts) plus clinical symptoms (two of the following: leucocytosis or leucopenia, abnormal temperature, purulent secretions), plus either a pathological organism (by Gram stain or culture), or a required change in antibiotics |
| Grade 2 | Pneumothorax |
| Grade 2 | Noninvasive ventilation, strictly applied to those with all of the following: a) oxygen saturation (SpO2) lower than 92% under supplemental oxygen; b) need of supplemental oxygen > 5 L/min; and RR ≥ 30 bpm |
| Grade 2 | Re-intubation postoperative or intubation, period of ventilator dependence (non-invasive or invasive ventilation) ≤ 48 h |
| Grade 4 | Ventilatory failure: postoperative ventilator dependence exceeding 48 h, or reintubation with subsequent period of ventilator dependence exceeding 48 h |
| Grade 5 | Death before hospital discharge |

Perioperative management and monitoring

All these participants will be managed and monitored following the general standard of the practices of the Department of Anesthesiology, West China Hospital of Sichuan University no matter which group they belong to. Intraoperative standard monitoring includes capnography, electrocardiogram (ECG), pulse oximeter, invasive arterial blood pressure, central venous pressure, urine output, nasopharyngeal temperature, transesophageal echocardiogram (TEE), activated clotting time (ACT). Other managements are optional, including bispectral index (BIS) and cerebral oxygen saturation monitoring.

The anesthesiologists give the participants heparin (3 mg/kg) to obtain an ACT of 280 s for aortic cannulation and 480 s before starting the CPB. At the end of the CPB, protamine is given to antagonize heparin with 1:1. The target ACT is about the same as the baseline value. Mean arterial pressure will be kept between 50 and 70 mmHg during CPB.

Data collection

All the related data are collected on the CRF. Preoperative data include age, height, weight, EuroSCORE II, smoking status, diabetes mellitus, peripheral arterial disease, pulmonary comorbidity, pulmonary infection in the past 30 days and the classification of American Society of Anesthesiologists (ASA). Intraoperative data include type and duration of surgery procedure, duration of CPB and aortic cross clamp, units of allogeneic red blood cell transfusions, volume and type of fluids administered. Postoperative data include time to extubation, need for non-invasive ventilator, re-intubation, postoperative hospital stay, temperature, leukocyte count and chest X-ray. Other data at three time points (the moment arriving in the ICU, and 6 and 12 h after arrival in the ICU) include PaO2, FiO2, and partial pressure of carbon dioxide (PaCO2). PPCs events are traced throughout the hospitalization.

Statistical considerations

Sample size

Sample-size calculation is based on two-sided alpha error of 0.05 and 80% power. We performed a pilot-study, showing that the Pmax is 40% in the LOV group (15 patients) and the Pmin is 22% in the NoV group (14 patients). Therefore, we expect a reduction of 45% in the incidence of this parameter, and then calculate the sample size of 127 patients per group (381 in total). Considering a dropout fraction of 10%, we will finally need a total of 420 participants.

Data analysis

Data will be analyzed following the intention-to-treat principle using SPSS version 22.0 (SPSS, Chicago, IL, USA). Comparisons in the dichotomous variables will be
made by the chi-square tests (or Fisher exact tests where numbers are small). Comparisons in the continuous variables will be made by analysis of variance or the non-parametric Kruskal-Wallis H test. Two-sided significance tests will be used throughout. The Results will be presented as n (%), mean (SD), or median (interquartile range). Bonferroni correction will be used when doing post hoc analysis.

Discussion
About 1–1.25 million patients undergo cardiac surgery worldwide each year and the number continues to increase [40]. CPB is an essential part of major cardiac surgery which may induce PPCs after operation. Since the PPCs are leading causes of mortality and morbidity [37], the effective preventive measures are particularly important and urgently needed. Various researches have been performed to reach this goal. The MECANO trial (NCT03098524) aims to assess the effects of mechanical ventilation (V̇ₜ: 3 ml/kg, PEEP: 5 cmH₂O, RR: 5 bpm) on all-cause mortality and respiratory failure comparing with no ventilation during CPB [41]. It does not provide the detailed usage of oxygen though the ambitious number of patients (n = 1500). The CPBVENT trial (NCT02090205) is a three-armed multicenter study [42], which compares two different ventilation strategies (CPAP: 5 cmH₂O or V̇ₜ: 2-3 ml/kg of IBW, PEEP: 3-5 cmH₂O, RR: 5 bpm) with no ventilation during CPB. This trial’s primary endpoint (PaO₂/FiO₂ ratio) can be easily identified, observed, and calculated, but indirect.
That is why we use it as a secondary endpoint. Besides, the inclusion and exclusion criteria are relatively strict, making the conclusion only applicable to a specific group of populations. The purpose of the VONTCPB study is to determine the impact that the different FiO₂ has on the PPCs when combining the ventilation strategy with low VT (3-4 ml/kg of IBW, only about half of the normal tidal volume), routine respiratory rate (10–12 bpm) and PEEP (5–8 cmH₂O) during CPB. This trial is the first study to investigate the effect of oxygen under different ventilation strategy during CPB. The number of the sample size (n = 420) is adequate. The primary endpoints are clinically relevant. The purpose of setting ventilator parameters in this way is to make the ventilation closer to the general. And this ventilation management during CPB may potentially benefit patients’ outcomes, but at the same time it has an imperceptible impact on the operating field.

There are limitations of the trial due to the inherent properties such as its single-center design. It is questionable whether the conclusion drawn by the VONTCPB trial can be applied in other hospitals because of the different clinical practice in other hospitals. Besides, only the volume controlled ventilation mode is applied, so it is unknown whether it is also applicable in other ventilation mode, such as pressure controlled mode.

In general, the VONTCPB trial would answer the question that whether maintaining mechanical ventilation during CPB is more preferable than no ventilation. Better still, which level of FiO₂ is beneficial to patients will be determined from this trial.

**Trial status**

The trial was started after we obtained the approval of local ethics committee and registered in the Chinese Clinical Trials Registry. We plan to spend six months to include cases and complete the trial in November 2018.

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**Funding**

Not applicable.

**Availability of data and materials**

Not applicable.

**Authors’ contributions**

XYY and HY provided the original idea of this randomized controlled trial. HY and MQZ contributed to the study concept and design and wrote the manuscript. YQL and LF participated patients' recruitment and data collection. XFL contributed to the sample size design and data analysis. All authors read and approved the final manuscript.

**Authors’ information**

See the title page.

**Ethics approval and consent to participate**

This clinical study protocol has been approved by the West China Hospital of Sichuan University Biomedical Research Ethics Committee on 12 March 2018 (Approval number 2017 (400)). We will obtain informed consent from patients when recruiting before the experiment. The patients can withdraw from the trial at any time.

**Consent for publication**

All authors agree to its submission to *Trials* and publish it.

**Competing interests**

The authors declare that they have no competing interests.

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**Additional file**

**Additional file 1:** The SPIRIT 2013 checklist of this trial. (DOCX 53 kb)
