Does closed-loop automated oxygen control reduce the duration of mechanical ventilation? A randomised controlled trial in ventilated preterm infants

Ourania Kaltsogianni1, Theodore Dassios1,2* and Anne Greenough2,3,4

Abstract

Background: Many preterm infants require supplemental oxygen in the newborn period but experience frequent fluctuations of their oxygen saturation levels. Intermittent episodes of hypoxia or hyperoxia increase the risk of complications. Compliance with achievement of oxygen saturation targets is variable, and the need for frequent adjustments of the inspired oxygen concentration increases workload. Closed-loop automated oxygen control systems (CLAC) improve achievement of oxygen saturation targets and reduce both episodes of hypoxia and hyperoxia and the number of manual adjustments. This study investigates whether CLAC compared with manual oxygen control reduces the duration of mechanical ventilation in preterm infants born at less than 31 weeks of gestation.

Methods: This randomised controlled trial performed at a single tertiary neonatal unit is recruiting 70 infants born at less than 31 weeks of gestational age and within 48 h of initiation of mechanical ventilation. Infants are randomised to CLAC or manual oxygen control from recruitment until successful extubation. The primary outcome is the duration of mechanical ventilation, and secondary outcomes are the percentage of time spent within target oxygen saturation ranges, the time spent in hypoxia or hyperoxia, the number of manual adjustments required, the number of days on oxygen, the incidence of bronchopulmonary dysplasia and the length and cost of neonatal unit stay. The study is performed following informed parental consent and was approved by the Yorkshire and the Humber-Sheffield Research Ethics Committee (protocol version 1.1, 13 July 2021).

Discussion: This trial will investigate the effect of CLAC on the duration of mechanical ventilation, which is an important clinical outcome as prolonged mechanical ventilation is associated with important adverse outcomes, such as bronchopulmonary dysplasia.

Trial registration: ClinicalTrials.Gov NCT05030337. Registered on 17 August 2021

Keywords: Closed loop automated oxygen control, Preterm infants, Mechanical ventilation, Intermittent hypoxemia and hyperoxemia

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**Funder(s)** The equipment was provided by SLE.

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**Committees** n/a

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**Trial registration dataset (Continued)**

| **Primary registry and trial identifying number** | ClinicalTrials.Gov, NCT05030337 |
| **Date of registration** | 17/08/2021 |
| **IRAS no** | 297749 |
| **Source of monetary or material support** | King’s College Hospital (KCH)/King’s College London (KCL) |
| **Follow-up duration (if applicable)** | Participants will be followed up until their discharge from the neonatal unit |
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| **Public title** | Optimising ventilation in preterms with closed-loop oxygen control |
| **Scientific title** | Does closed-loop automated oxygen control reduce the duration of mechanical ventilation? A randomised controlled trial in ventilated preterm infants |
| **Country of recruitment** | United Kingdom |
| **Health condition(s) or problem(s) studied** | Prematurity, mechanical ventilation, hypoxia, hyperoxia |
| **Interventions** | Closed-loop automated oxygen control (intervention group) |

**Role of study sponsor and funder**

King’s College London will take primary responsibility for ensuring the study design meets appropriate standards and uses proper conduct and reporting. King’s College London also provides insurance cover to provide for payment of damages or compensation in respect of any claim made by a research subject for bodily injury arising out of participation in a clinical trial or healthy volunteer study with certain restrictions.

King’s College Hospital takes responsibility for arranging the initiation and management of the research and will ensure that appropriate standards, conduct and reporting are adhered to with regards to its facilities and staff involved with the study. King’s College Hospital will also undertake the governance review for the project and provide cover for clinical negligence by any of its staff undertaking the research.

**Roles and responsibilities of study management committees, groups, and individuals**

There are no committees or steering groups involved in the management of this study, but all the neonatal consultants have oversight of the study. The chief investigator and sponsor will monitor and audit the conduct of this research and will ensure that appropriate standards and reporting are adhered to (p6, 59–60). All the neonatal consultants have oversight of the study (p6, 70–71). In addition to this, an independent researcher not involved in the project will review trial data upon recruiting half of planned patients. The researcher will evaluate differences in the primary outcome to decide whether enough data have been gained at that point. The research team will remain blinded to the results of
the interim analysis unless there are any significant findings (p12, 204–208). This is a non-blinded, randomised, superiority trial.

Protocol contributors
This study has been designed by Professor Anne Greenough and Dr Theodore Dassios. Dr Ourania Kaltsogianni, neonatal grid trainee and research fellow, will conduct the study and primarily perform data analysis and interpretation, manuscript writing and dissemination of results with support from both supervisors.

Background
Seven to eight per cent of all infants are born prematurely [1] and many require respiratory support in the newborn period [2]. Although such support can be life-saving, preterm infants who require mechanical ventilation and oxygen therapy frequently develop complications [3]. The most common adverse outcome is bronchopulmonary dysplasia (BPD), and importantly prematurely born infants can suffer chronic respiratory morbidity including troublesome respiratory symptoms, lung function abnormalities and exercise intolerance even in adolescence and adulthood [4]. Other complications include intracerebral haemorrhage which can result in cerebral palsy [5] and retinopathy of prematurity (ROP) which can cause blindness [6].

Ventilated neonates frequently require supplemental oxygen, but its use must be carefully monitored due to associated complications. Hyperoxia leads to the development of reactive oxide species (ROS) and increases the risk of bronchopulmonary dysplasia and retinopathy of prematurity [7, 8]. On the other hand, hypoxia increases morbidity and mortality [9, 10]. Therefore, oxygen saturation levels (SpO₂) are continuously monitored in clinical practice and used to guide adjustments to the inspired oxygen concentration (FiO₂), which are traditionally made manually by neonatal practitioners. Maintaining oxygen saturations in the target range can be difficult as preterm neonates experience frequent fluctuations with as many as 600 intermittent hypoxic episodes in a week documented in one study [11]. Moreover, prolonged hypoxic episodes have been associated with increased risk of death after 36 weeks’ postmenstrual age or disability at 18 months corrected age including motor impairment, cognitive or language delay, severe hearing loss or bilateral blindness [12]. Compliance with achievement of SpO₂ targets is variable within the same patient over time, as well as between patients and centres [13]. Furthermore, target achievement decreases as the number of patients per nurse increases [14].

Closed-loop automated oxygen control (CLAC) systems monitor SpO₂ values in real-time to calculate and make an adjustment to the FiO₂ patient delivery without any human intervention [15]. These systems may provide a solution for the low compliance with achievement of target oxygen levels and reduce the need for manual adjustments and hence the workload, as well as reducing complications. Several algorithms exist to calculate the FiO₂ adjustment in relationship to the SpO₂ changes in infants with different severities of respiratory disease [16]. An example is the Oxygenie Auto-O₂ software (SLE) that uses a proportional-integral-derivative control algorithm, and it has been shown to be effective in improving compliance with target achievement in preterm infants, reducing episodes of hypoxia or hyperoxia and the need for manual adjustments to supplementary oxygen [17]. In a randomised crossover study, using that device we demonstrated that preterm ventilated infants experienced fewer prolonged desaturations during the closed-loop automated oxygen control period and spent an increased percentage of time within their target SpO₂ range with fewer manual adjustments to the inspired oxygen concentration [18]. A review of 16 studies highlighted that the studies of CLAC were very heterogenous for design, population size and device used [15]. Moreover, the majority of those studies had a crossover design and limited data available for analysis. Two reviews emphasised that none of the previous studies had demonstrated whether the clinical outcomes of preterm infants were improved [15, 19]. A subsequent literature review including 18 studies [19] highlighted that CLAC was consistently associated with an increased percentage of time spent within the target oxygen saturation range with fewer manual adjustments to the FiO₂ and was effective in infants on non-invasive respiratory support or mechanically ventilated at a range of postnatal ages. The results appear to be consistent for all the control algorithms [19] and across different SpO₂ target ranges [20, 21]. In addition, previous studies demonstrated that CLAC could facilitate earlier weaning of the inspired oxygen concentration when compared to manual control [22, 23], which is a major determinant of an infant’s readiness for extubation. It is, therefore, possible and our hypothesis that use of CLAC would reduce the duration of mechanical ventilation

The aim of this study is to explore if in prematurely born ventilated infants, the use of closed loop automated oxygen control compared to standard care reduced the duration of mechanical ventilation. That is an important clinical outcome as prolonged mechanical ventilation increases the risk of BPD and long-term comorbidities [24]. Our primary outcome is the duration of mechanical ventilation. Secondary outcomes are the percentage of
time spent within target oxygen saturation ranges, the
time spent in hypoxia or hyperoxia, the number of man-
ual adjustments required by clinical staff, a diagnosis of
BPD at 36 weeks post menstrual age and the length of
intensive care stay.

Methods
This is a non-blinded, randomised controlled trial. In-
fants are allocated to parallel groups in a 1:1 ratio.

Setting
The setting is a single tertiary neonatal unit at the King’s
College Hospital NHS Foundation Trust, London, UK.
The unit has previous experience on the use of closed-
loop automated oxygen control in the context of trials.

Inclusion criteria
Inclusion criteria are as follows:

1. Infants delivered at less than 31 weeks of gestational
   age requiring mechanical ventilation
2. Recruitment within 48 h of initiation of mechanical
   ventilation

Exclusion criteria
1. Infants delivered above 31 weeks of gestational age
2. Infants with major congenital abnormalities

Recruitment
Parents or legal guardians of all eligible infants are
initially approached by the clinical team taking care of
the infant and if they agree, by a researcher, within 24 h
of initiation of mechanical ventilation. The parents are
provided with an information sheet about the study. The
researchers answer questions and respond to any
concerns in a face-to-face meeting and obtain written in-
formed consent. The study was approved by the Health
Research Authority and by the Yorkshire and the
Humber-Sheffield NHS Research Ethics Committee.

Randomisation
The randomisation sequence is generated by using an
online randomisation generator and concealed in sealed
opaque envelopes. This is done by a person independent
of the research team who is not involved in the study.

Enrolment
Infants will be enrolled in the study by the research
team within 48 h of initiation of mechanical ventilation
following parental consent (supplied as attachment).
Infants are randomised to one of the study arms by
opening the next envelope.

Blinding
The study is not blinded. Decisions regarding ongoing
care, for example ventilator settings and interventions
such as blood gases and chest radiographs, are made by
the clinical team as per the neonatal unit’s guidelines.

Intervention
At enrolment, participants are randomised to receiving
either closed-loop automated oxygen control (interven-
tion group) or manual control of the inspired oxygen
concentration. All infants are ventilated using the
SLE6000 ventilators and ventilation settings are manu-
ally adjusted by the clinical team as per the unit’s proto-
col. In addition to standard care, infants in the
intervention group are connected to the Oxygenie Auto-
O2 software (SLE). This software uses oxygen saturation
levels from the SpO2 probe attached to the neonate,
which are fed into an algorithm, to automatically adjust
the percentage of inspired oxygen to maintain oxygen
saturations within the target range. Manual adjustments
including the percentage of inspired oxygen are allowed
at any point during the study, if deemed appropriate by
the clinical team.

The nurse-to-patient ratio is according to the unit
protocol that is determined on the patient’s acuity. Re-
spiratory care will be provided for all infants by the clinical
team as per the unit’s guidelines. All infants will be
ventilated using the SLE6000 ventilators and ventilation settings
will be adjusted manually following the unit’s protocol. The intervention will be performed in addition to
standard care and manual adjustments will be allowed
at any point during the study, if deemed appropriate by
the clinical team (p10–11, 173–180).

Infants are studied from enrolment until successful
extubation [25]. For infants who fail extubation and
require reintubation within 48 h, they continue in their
initial study arm if they are less than 28 days old. At
KCH NICU, preterm infants with non-chronic respira-
tory pathology are extubated from a targeted tidal vol-
ume of 5 ml/kg, a peak inspiratory pressure at or below
18 cm H2O and a backup rate of 40 breaths per minute.
FiO2 requirement should be less than 0.3. The infant
should have good respiratory drive as well as blood
gases. Therefore, for infants randomised to the interven-
tion group, closed-loop oxygen delivery resumes. Pre-
term infants that remain ventilated beyond day 28 of life
continue at their study arm (closed-loop automated oxy-
gen control or manual oxygen control) till their first
extubation attempt (see Fig. 1).

Primary outcome measure
The primary outcome measure is the duration of
mechanical ventilation. The primary outcome measure is
the duration of mechanical ventilation from enrolment
to the study until successful extubation. Extubation failure is defined as the need for reintubation within 48 h. For infants who fail extubation, they continue in their initial study arm if they are less than 28 days old, and the time they remain ventilated is counted towards the total duration of mechanical ventilation. For infants that remain ventilated beyond day 28 of life, the study continues till their first extubation attempt, and that period represents the primary outcome measure.

Currently at KCH, the CLAC algorithm is used only for mechanically ventilated infants and in the context of clinical research studies. Hence, automatic oxygen control will be discontinued following extubation. Non-invasive ventilation modes will be delivered via the SLE6000 ventilators, and the control of the inspired oxygen concentration will be performed manually.

Secondary outcomes
The secondary outcomes are the percentage of time spent within target oxygen saturation range, the time spent in hypoxia or hyperoxia, the number of manual adjustments required, the number of days on oxygen, the incidence of bronchopulmonary dysplasia at 36 weeks post menstrual age and the length and cost of neonatal unit stay. In line with the latest published evidence [10], our pulse oximetry oxygen saturation targets in preterm infants born at less than 31 weeks gestational age are 91 to 95% with alarm limits in the range of 90 to 98%. A researcher who is blinded to which arm the infant was randomised will undertake the analysis of the data. Retinopathy of prematurity will not be recorded as a secondary outcome, as our study is not powered to detect any significant differences in ROP outcomes between the two participating groups, but this outcome will be reported.

The order of study events is detailed in the Standard Protocol Items: Recommendations for Interventional Trials (SPIRIT) figure in Fig. 2. A SPIRIT checklist is also provided as an additional file and the protocol is based on the SPIRIT reporting guidelines [26].
Sample size
In a cohort of 368 preterm infants born between 24 to 30 weeks of gestation, the standard deviation for the duration of mechanical ventilation was 3.83 days [27]. The median duration of ventilation in our NICU was 12 days (mean 19 days). Randomisation of 70 infants allows us to detect a decrease in the duration of mechanical ventilation of three days with a 90% power at the 5% significance level.

At the request of the ethics committee, an interim analysis will be carried out after half the sample has been recruited to review if enough data have been gained at that point. The analysis will be performed by an independent researcher not involved in the project to avoid introducing bias to our data. The research team will be blinded to the results of the interim analysis unless there are any significant findings.

Data collection
Demographic and outcome data will be collected from the clinical records and recorded under a unique study identifier number. Basic epidemiologic parameters such

| TIMEPOINT | ENROLMENT: | INTERVENTIONS: | ASSESSMENTS: |
|-----------|------------|----------------|--------------|
|           | Eligibility screen | [Closed-loop automated oxygen control] | [Duration of MV] |
|           | Informed consent | [Manual oxygen control] | [Percentage of time spent within target SpO2 range, episodes of hypoxia or hyperoxia] |
|           | Allocation |               | [Number of manual adjustments to the FiO2] |
|           |           |               | [Days on O2, BPD, length of ICU stay] |

| STUDY PERIOD | Enrolment | Allocation | Intervention | Follow up |
|--------------|-----------|------------|--------------|-----------|
|              | Within 24 hours of initiation of MV | Within 48 hours of initiation of MV | 48 hours Extubation | 36 weeks PMA or discharge from NICU |

Fig. 2 SPIRIT figure of trial interventions and timings
as gestational age at birth, birth weight, corrected gestational age, postnatal age and weight at study enrolment, use of antenatal steroids, mode of delivery, doses of surfactant administered, birth plurality and associated comorbidities that may impact on the duration of mechanical ventilation, for example patent ductus arteriosus and infection, will be recorded for each infant. Oxygen saturation and FiO₂ data will be downloaded from the ventilators. Paper data will be stored in a locked filling cabinet. Patient de-identified electronic data will be stored in password protected computers and on encrypted devices.

Statistical analysis
Data will be compared between infants in the intervention group (closed-loop automated oxygen control) and infants in the control group (manual oxygen control). The data will be assessed for normality; Student’s t test will be used for normally distributed data and Mann-Whitney U test for skewed data. Categorical data will be assessed using the Fisher two-tailed exact test. Analysis will be conducted on an intention-to-treat basis, and missing data will not be imputed. Analysis will be undertaken to a researcher blind to which arm the infants was randomised.

Safety
All participants will receive standard care and monitoring by clinical staff. Ventilator settings will be adjusted and additional investigations such as blood gases and chest radiographs will be performed at the discretion of the clinicians. Therefore, there should not be any risk imposed to the intervention group with the addition of the closed-loop automated oxygen delivery system. The system does not mask large increased oxygen requirements between manual observation recordings, as it activates an alarm when there is an increase in the oxygen requirement ≥ 30% from the basal level alerting the clinical team. All serious adverse events will be recorded on a serious adverse event (SAE) form and will be emailed to the sponsor within one working day of the chief investigator (CI) becoming aware of the event. When the event is unexpected and thought to be related to the use of the device, this will be reported by the CI/ sponsor to the ethics and Health Research Authority within 15 days.

Protocol compliance
Any accidental protocol deviations will be appropriately documented and reported to the chief investigator and sponsor immediately. The chief investigator and sponsor will monitor and audit the conduct of this research. For any amendment to the study, the chief investigator or designee, in agreement with the sponsor, will submit information to the appropriate body in order for them to issue approval for the amendment (p13, 246–249).

Research Ethics Committee (REC) and other regulatory review and reports
Before the start of the study, a favourable opinion was sought from a REC for the study protocol and other relevant documents (informed consent forms and patient information leaflets).

Regulatory review and compliance
For any amendment to the study, the chief investigator or designee, in agreement with the sponsor, will submit information to the appropriate body in order for them to issue approval for the amendment.

Data protection and patient confidentiality
The General Data Protection Regulation and Data Protection Act 2018 will be adhered to. Data will be de-identified before being entered in a secure database. A unique study identifier number will be issued to each participant on enrolment into the study. That number will be used in all subsequent data collection forms. Patient paper data will be stored in a locked filing cabinet in a room that is only accessible to the research team and that is based at the Neonatal Intensive Care Unit facilities at King’s College Hospital. Patient de-identified electronic data will be stored on encrypted university computers or memory stick devices, both of which require user identification and password verification.

Archiving
At the end of the trial, all essential documentation will be archived securely by the CI for a minimum of 25 years from the declaration of the end of the trial. Essential documents include those which enable both the conduct of the trial and the quality of the data produced to be evaluated and show whether the site complied with all applicable regulatory requirements. All archived documents will continue to be available for inspection by appropriate authorities upon request.

Indemnity
King’s College London indemnity applies for insurance/indemnity to meet the potential legal liability of the sponsor for harm to participants arising from the design and management of the research. NHS indemnity scheme applies for insurance/indemnity to meet the potential legal liability of the investigators arising from harm to participants in the conduct of the research.
Access to the final study dataset
The individuals in the study will be notified of the outcome of the study as below. The investigators will have access to the final study dataset. Participants will be informed that anonymised data may be shared with other researchers for research purposes only.

Dissemination policy
On completion of the study, the data will be analysed and a final study report will be prepared that could be accessed via the sponsor. The study will be presented at research meetings at the Neonatal Intensive Care Unit at King’s College Hospital as well as university meetings at King’s College London. Anonymised study data will be presented at conferences and published by the investigators in peer-reviewed journals. Participants will be notified of the outcome of the study via provision of the publication and an accompanying newsletter.

Authorship eligibility guidelines and any intended use of professional writers
The final study authors will include Dr Ourania Kaltsogianni, Dr Theodore Dassios and Professor Anne Greenough. Professional writers will not be used.

Intellectual property
All intellectual property rights and know-how in the protocol and in the results arising directly from the study shall belong to KCL.

Discussion
This study will compare the effectiveness of closed-loop automated oxygen control to manual oxygen control in preterm ventilated infants born at less than 31 weeks completed gestation. Previous studies in preterm infants demonstrated that CLAC improved the percentage of time spent within an assigned oxygen saturation target range [18, 20–23], but it remains unknown whether that improvement has any positive long-lasting effects on clinical outcomes. A retrospective cohort study comparing outcomes pre and post implementation of CLAC as standard care for preterm infants did not demonstrate any significant effect on morbidity or mortality at hospital discharge [28] but was associated with a significant reduction in the duration of mechanical ventilation. The study, however, was limited by its retrospective design and with several changes to neonatal care during the study span. Furthermore, it was not sufficiently powered to detect outcomes including BPD, ROP, necrotising enterocolitis (NEC), intraventricular haemorrhage (IVH) and periventricular leukomalacia (PVL), and the time infants received CLAC was not recorded, and the authors concluded that their results may have been biased by several factors not accounted for. This randomised controlled trial will provide more robust evidence on the effect of CLAC on the duration of mechanical ventilation that is related to long-term complications [24] and therefore may demonstrate if CLAC could improve long-term respiratory outcomes for preterm infants. That will be essential knowledge before the intervention is implemented into standard care.

Participants are enrolled in our study within 48 h of initiation of mechanical ventilation. This enables a reasonable time frame to approach parents and obtain informed consent on participation. Infants are randomised to CLAC or manual oxygen control from recruitment to the study until they are successfully extubated. In comparison with the previous studies that mostly had a randomised crossover design and a short duration [17, 20, 29, 30], our study will provide more data points available for analysis and will explore the effect of CLAC during the different stages of evolving lung disease in preterm infants.

Over the past decade, increased knowledge of the risks related to mechanical ventilation and advances in non-invasive respiratory support have resulted in the reduction in the duration of mechanical ventilation [31]. Nevertheless, historical data from our unit for the period 2015–2020 demonstrated that the median duration of mechanical ventilation for preterm infants born less than 31 weeks gestation was 12 days. Hence, our study is powered to detect a decrease in the duration of mechanical ventilation of 3 days between the two treatment groups.

In conclusion, if this study demonstrates that CLAC is associated with earlier extubation and a shorter duration of mechanical ventilation, that could have significant impact on long-term respiratory outcomes of preterm infants.

Trial status
At the time of submission, this trial has been approved by the NHS Research Ethics Committee and the Health Research Authority and is recruiting participants.

Abbreviations
BPD: Bronchopulmonary dysplasia; CI: Chief investigator; CLAC: Closed-loop automated oxygen control; FiO2: Fraction of inspired oxygen concentration; IVH: Intraventricular haemorrhage; KCH: King’s College Hospital; KCL: King’s College London; NEC: Necrotising enterocolitis; PVL: Periventricular leukomalacia; ROP: Retinopathy of prematurity; ROS: Reactive oxide species; SpO2: Oxygen saturation levels

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s13063-022-06222-y.

Additional file 1.
Additional file 2.
Additional file 3: SPIRIT checklist for trials
Acknowledgements
We are grateful to Mrs. Deirdre Gibbons for secretarial assistance.

Authors’ contributions
AG devised the study. AG, OK and TD wrote the protocol. OK wrote the first draft of this manuscript; all authors have contributed to and approved the final manuscript and will be involved in the interpretation and analysis of the data. The author(s) read and approved the final manuscript.

Funding
The equipment was provided by SLE.

Availability of data and materials
OK, TD and AG will have access to the final trial dataset. There are no contractual agreements that limit such access for all three investigators.

Declarations

Ethics approval and consent to participate
This study was reviewed and approved by the Health Research Authority (HRA) and the Yorkshire and the Humber-Sheffield Research Ethics Committee (Reference 21/YH/0139). The co-sponsors are King’s College London and King’s College Hospital NHS Foundation Trust.
Written informed parental consent is obtained for recruitment to the study.

Consent for publication
N/A

Competing interests
The authors declare that they have no competing interests.

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Received: 24 September 2021 Accepted: 26 March 2022

Published online: 08 April 2022

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