Wean earlier and automatically with new technology: a multicentre, pilot randomised controlled study

Submission date Recruitment status [X] Prospectively registered 03/01/2006 No longer recruiting [X] Protocol

Registration date Overall study status 30/01/2006 Completed

Last Edited Condition category 31/07/2013 Respiratory

[X] Protocol[Statistical analysis plan[X] Results[Individual participant data

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s)

Scientific

Contact name

Dr Karen E. A. Burns

Contact details

St Michael's Hospital 30 Bond street 3-075E Queen wing Toronto Canada M5B 1W8

Additional identifiers

Protocol serial number N/A

Study information

Scientific Title

Acronym

The WEAN Study

Study objectives

The primary objective of this multicentre pilot study is to compare automatic and protocolised weaning and to evaluate compliance with the proposed weaning and sedation protocols

Ethics approval required

Old ethics approval format

Ethics approval(s)

Not provided at time of registration

Study design

Multicentre randomised controlled trial

Primary study design

Interventional

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Weaning intubated adults from invasive mechanical ventilation

Interventions

Automatic (computer-driven) weaning versus protocolized weaning

Intervention Type

Other

Phase

Not Specified

Primary outcome(s)

Compliance with the weaning and sedation protocols

Key secondary outcome(s))

- 1. Acceptance of the weaning and sedation protocols
- 2. Weaning and Intensive Care Unit (ICU) related-complications
- 3. Time to successful completion of a Single Breath Test (SBT) and extubation

Completion date

01/06/2006

Eligibility

Key inclusion criteria

Patients who are invasively ventilated for at least 24 hours

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Sex

All

Key exclusion criteria

- 1. Age < 16 years
- 2. Tracheotomy
- 3. Pregnancy
- 4. Do not intubate (DNI) or do not resuscitate (DNR) patients
- 5. Planned extubation on the day of study inclusion
- 6. Planned surgery or procedures within 48 hours of randomization

Date of first enrolment

30/05/2006

Date of final enrolment

01/06/2006

Locations

Countries of recruitment

Canada

Study participating centre St Michael's Hospital

Toronto Canada M5B 1W8

Sponsor information

Organisation

Physician Services Incorporated (Canada)

ROR

https://ror.org/0385yzn06

Funder(s)

Funder type

Charity

Funder Name

Physician Services Incorporated (Canada)

Funder Name

Draeger Canada (in kind)

Results and Publications

Individual participant data (IPD) sharing plan

IPD sharing plan summary

Not provided at time of registration

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article	results	01/06/2013		Yes	No
Protocol article	protocol	04/09/2009		Yes	No