Reducing Atelectasis during General Anaesthesia – the Importance of Oxygen Concentration, End-Expiratory Pressure and Patient Factors

A Clinical Study Exploring the Prevention of Atelectasis in Adults

LENNART EDMARK
Dissertation presented at Uppsala University to be publicly examined in Vårdskolans aula, Ingång 21, Västmanlands sjukhus Västerås, Västerås, Wednesday, 18 December 2013 at 13:00 for the degree of Doctor of Philosophy (Faculty of Medicine). The examination will be conducted in Swedish. Faculty examiner: Professor emeritus Sten Lindahl (Karolinska institutet, Institutionen för fysiologi och farmakologi).

Abstract

Background: The use of pure oxygen during preoxygenation and induction of general anaesthesia is a major cause of atelectasis. The interaction between reduced lung volume, resulting in airway closure, and varying inspiratory fractions of oxygen (F\(_{IO_2}\)) in determining the risk of developing atelectasis is still obscure.

Methods: In this thesis, computed tomography (in studies I and II during anaesthesia, in studies III and IV postoperatively) was used to investigate the area of atelectasis in relation to F\(_{IO_2}\) and varying levels of continuous positive airway pressure (CPAP) or positive end-expiratory pressure (PEEP).

Study I investigated the short-term influence of reducing F\(_{IO_2}\) during preoxygenation and induction of general anaesthesia, and the time to hypoxia during apnoea.

Study II focused on the long-term effect of an F\(_{IO_2}\) of 0.8 for preoxygenation.

Study III applied CPAP/PEEP with an F\(_{IO_2}\) of 1.0 or 0.8 for pre- and postoxygenation until extubation. After extubation, CPAP with an F\(_{IO_2}\) of 0.3 was applied before the end of mask ventilation.

Study IV compared two groups given CPAP/PEEP during anaesthesia and an F\(_{IO_2}\) of 1.0 or 0.3 during postoxygenation, but without CPAP after extubation.

Results: Study I showed a reduction in atelectasis with an F\(_{IO_2}\) of 0.8 or 0.6, compared with 1.0, but the time to hypoxia decreased. In study II, atelectasis evolved gradually after preoxygenation. In study III, atelectasis was reduced with an F\(_{IO_2}\) of 1.0 and CPAP/PEEP compared with an F\(_{IO_2}\) of 1.0 without CPAP/PEEP. The intervention failed in the group given an F\(_{IO_2}\) of 0.8, this group had more smokers. Atelectasis and age were correlated. In study IV, no difference was found between the groups. Post hoc analysis showed that smoking and ASA class increased the risk for atelectasis.

Conclusion, the effect of reducing F\(_{IO_2}\) during preoxygenation to prevent atelectasis might be short-lived. A lower F\(_{IO_2}\) shortened the time to the appearance of hypoxia. Increasing lung volume by using CPAP/PEEP also decreased the risk of atelectasis, but the method might fail; for example in patients who are heavy smokers. In older patients care must be taken to reduce a high F\(_{IO_2}\) before ending CPAP.

Keywords: Anaesthesia, general. Lung: Atelectasis; CPAP; Oxygen; PEEP; Ventilation, mechanical; Tomography, X-ray computed.

Lennart Edmark, Centre for Clinical Research, County of Västmanland, Centrallasarettet, Uppsala University, SE-72189 Västerås, Sweden. Department of Medical Sciences, Clinical Physiology, Akademiska sjukhuset, Uppsala University, SE-75185 Uppsala, Sweden.

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urn:nbn:se:uu:diva-209714 (http://urn.kb.se/resolve?urn=nbn:se:uu:diva-209714)
To my parents Roland and Bergljot
And to my daughter Emma
List of Papers

This thesis is based on the following papers, which are referred to in the text by their Roman numerals.


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### Abbreviations

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<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>ASA</td>
<td>American Society of Anesthesiologists physical status</td>
</tr>
<tr>
<td>BMI</td>
<td>Body mass index (calculated as weight/(height in meters)²)</td>
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<tr>
<td>CC, CV</td>
<td>Closing capacity, closing volume</td>
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<tr>
<td>COPD</td>
<td>Chronic obstructive pulmonary disease</td>
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<tr>
<td>CPAP</td>
<td>Continuous positive airway pressure</td>
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<tr>
<td>CT</td>
<td>Computed tomography</td>
</tr>
<tr>
<td>DAWD</td>
<td>Duration of apnoea without desaturation</td>
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<tr>
<td>EELV</td>
<td>End expiratory lung volume</td>
</tr>
<tr>
<td>ERV</td>
<td>Expiratory reserve volume</td>
</tr>
<tr>
<td>F&lt;sub&gt;ET&lt;/sub&gt;O₂</td>
<td>Fraction of end tidal oxygen</td>
</tr>
<tr>
<td>F&lt;sub&gt;I&lt;/sub&gt;O₂</td>
<td>Fraction of inspired oxygen</td>
</tr>
<tr>
<td>FRC</td>
<td>Functional residual capacity</td>
</tr>
<tr>
<td>HPV</td>
<td>Hypoxic pulmonary vasoconstriction</td>
</tr>
<tr>
<td>HU</td>
<td>Hounsfield unit</td>
</tr>
<tr>
<td>IBW</td>
<td>Ideal body weight</td>
</tr>
<tr>
<td>PCV</td>
<td>Pressure controlled ventilation</td>
</tr>
<tr>
<td>PEEP</td>
<td>Positive end expiratory pressure</td>
</tr>
<tr>
<td>PSV</td>
<td>Pressure support ventilation</td>
</tr>
<tr>
<td>RM</td>
<td>Recruitment manoeuvre</td>
</tr>
<tr>
<td>RV</td>
<td>Residual volume</td>
</tr>
<tr>
<td>SpO₂</td>
<td>Peripheral arterial oxygen saturation</td>
</tr>
<tr>
<td>TCI</td>
<td>Target control infusion</td>
</tr>
<tr>
<td>TIVA</td>
<td>Total intravenous anaesthesia</td>
</tr>
<tr>
<td>V&lt;sub&gt;A&lt;/sub&gt;</td>
<td>Alveolar minute ventilation</td>
</tr>
<tr>
<td>V&lt;sub&gt;A/Q&lt;/sub&gt;</td>
<td>Alveolar ventilation/perfusion ratio</td>
</tr>
<tr>
<td>VCV</td>
<td>Volume controlled ventilation</td>
</tr>
<tr>
<td>V&lt;sub&gt;T&lt;/sub&gt;</td>
<td>Tidal volume</td>
</tr>
<tr>
<td>Q</td>
<td>Perfusion</td>
</tr>
<tr>
<td>ZEEP</td>
<td>Zero end expiratory pressure</td>
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Preface

The first two studies in this thesis were conceived more than ten years ago, although the second study was published only recently. These two studies were relevant at the time they were designed. However, when the opportunity to expand the research focus arose, the ever changing nature of science and knowledge, as well as a little bit on a personal level, made the last two studies turn in what might appear as quite a new direction. Therefore, it may seem that the two parts come from different perspectives. However, from different angles these studies tell a common story, and put together a main theme emerges.
Introduction

A crucial question for the anaesthetist responsible for the high-risk patient during surgery involving general anaesthesia and mechanical ventilation is: what type of lung do we deliver to the post-operative ward?\textsuperscript{1} The recent identification of ventilator-induced lung injury as a possible mechanism, starting perioperatively\textsuperscript{2}, has made the concept of “lung protective ventilation” a serious concern both in the intensive care unit and the operating theatre. Applying strategies of lung protective ventilation during general anaesthesia has produced mixed results. However, according to a recent meta-analysis\textsuperscript{3}, lung protective intraoperative ventilatory settings have “the potential to protect against postoperative pulmonary complications”. The importance of reducing the risk of such complications relates to their incidence, and severity\textsuperscript{4}, and the vast number of major surgical procedures undertaken worldwide\textsuperscript{5} every year. In 2013, a small randomised trial of protective mechanical ventilation during general anaesthesia showed improved postoperative pulmonary function\textsuperscript{6}. The first multicentre study of the benefits of using lung-protective ventilation during anaesthesia for abdominal surgery emerged in the summer of 2013\textsuperscript{7}. The lung-protective strategy used in these two trials rests on three pillars: low tidal volumes ($V_T$), positive end-expiratory pressure (PEEP), and the repeated use of a recruitment manoeuvre (RM). It is interesting to note that even though the results in the studies are impressive, we do not really know why the lung-protective strategy is effective.

In the context of this thesis on reducing atelectasis formation during general anaesthesia in adults, we have to ask if it is possible that some of the positive effects seen in the mentioned studies originated from less “atelectrauma”, i.e. repetitive opening and closing of dependent lung tissue during anaesthesia with mechanical ventilation, or from reduced area of atelectasis postoperatively?

Etymologically, the term atelectasis refers to incomplete expansion, and in practice, the word has come to indicate collapse of alveoli in the lungs. Atelectasis is a pathological condition that is not observed in healthy people but occurs in about 90% of patients given general anaesthesia. It occurs predominately in the most dependent parts of the lungs\textsuperscript{8} and impairs oxygenation\textsuperscript{9}. The clinical significance of atelectasis might be substantial\textsuperscript{10}. An

One of the pioneers studying atelectasis was the 19th century German physician Ludwig Lichtheim. He observed that if a lung unit was closed off completely, this unit, if perfused, would collapse in a time frame that was dependent on the gas mixture in the unit. This observation is of critical importance, and the mechanism described by Lichtheim is referred to as “static atelectasis” in this thesis. Static atelectasis evolves if, for example, mucous, a tumour, swelling, oedema, pressure, or a foreign body blocks the communication to a lung unit.

In 1933, Beecher was the first to show the importance of decreased lung volume after abdominal surgery in the origin of postoperative respiratory failure, and the functional residual capacity (FRC) was identified as “the most important of all volumes”. Static atelectasis might also evolve if closing capacity (CC) in the lungs exceeds the sum of the functional residual capacity and the tidal volume (VT) for some critical time. As long as CC > (FRC + VT) there is widespread closure of small airways, and parts of the lungs are unable to participate in ventilation even during inspiration. The phenomenon of airway closure explains probably most of the age-dependent deterioration in arterial oxygenation, and occurs as soon as CC > FRC. The negative effect will increase as more and more of the fraction of the VT is needed to open up the airway during inspiration. In the supine position FRC is reduced, which is why airway closure is first revealed in the recumbent position.

Another way of expressing the impact of airway closure is to address it as a disturbance in the alveolar ventilation (VA) to lung perfusion (Q) relationship (VA/Q). Greater and more prominent airway closure reduces ventilation, and the VA/Q ratio declines, although the negative influence can be reduced by hypoxic pulmonary vasoconstriction (HPV). A simple way of treating hypoxemia caused by the occurrence of regions in the lungs with low VA/Q is to give extra oxygen. Applying the right amount of extra oxygen, depending on the degree of VA/Q disturbance, will normalise oxygenation completely. In everyday clinical work, we do not normally calculate the degree of VA/Q mismatch, and there is a risk of overdosing oxygen when treating hypoxemia. For every fraction of inspired oxygen (FIO2) there exists a critical inspiratory VA/Q where the expiratory volume decreases to zero. It has been calculated, that this critical VA/Q is increases from about 0.001 to 0.1, as the inspired gas is changed from air to 100 % oxygen. The lungs comprise of units with different VA/Q. Units that reach their critical level will not eliminate gas, but they may continue gas uptake until atelectasis develops. The partial pressure difference between gases in the alveoli and the lung
Capillaries govern the movement of these gases. With 100 % oxygen, also units with very low \( V_{A}/Q \) will contribute to oxygenation because of the low partial pressure in mixed venous blood and the function of the haemoglobin dissociation curve. These intricate events, leading to atelectasis, were first suggested by Briscoe et al.\textsuperscript{19} and elucidated further by Dantzker et al.\textsuperscript{18}, and these mechanisms leading to alveolar collapse are referred to as “dynamic atelectasis” in this thesis.

Thus, the theoretical backgrounds for understanding both static and dynamic atelectasis have been known for some time.

Atelectasis also reflects disturbance in the morphology of the lungs, but it was not until the technical advantages offered by computed tomography (CT), that it became possible to diagnose in detail the occurrence of atelectasis and to combine this information with the results from other investigations or procedures. Since 1980, the use of CT\textsuperscript{20} for measuring atelectasis has revealed much new information. In 1985, transmitted compression on the lower part of the lungs by the abdominal viscera was thought to play a crucial role in the genesis of atelectasis in adults\textsuperscript{21}. Later, the importance of the oxygen concentration was “rediscovered” and established firmly\textsuperscript{22,23}.

In 2003, the importance of opposing airway closure during anaesthesia was demonstrated. Counteracting airway closure before and during induction of general anaesthesia with the use of continuous positive airway pressure (CPAP), followed by a positive end-expiratory pressure (PEEP), completely stopped early atelectasis from developing in spite of preoxygenation with 100 % oxygen. This was first demonstrated in patients with normal weights\textsuperscript{24} and then in obese patients\textsuperscript{25}. Using CPAP/PEEP in this way also prolonged the period of non-hypoxic apnoea\textsuperscript{26,27}. It looked like the problem might be solved; i.e., pure oxygen and open lungs working together.

Applying CPAP was not a common feature in anaesthesia ventilators 10 years ago, but is now common in modern ones. With this background in mind, the first two studies should be seen in the perspective of what could be done and what was known before 2003. The last two studies aimed to investigate the importance of oxygen concentration combined with the practise of keeping the lungs open by increasing FRC with CPAP/PEEP before, during, and after anaesthesia.
Aims of the studies

Computed tomography was used, when applicable, in adults:

1. To examine the effect of 60 and 80 % oxygen compared with 100 % oxygen during preoxygenation and induction of anaesthesia on the early formation of atelectasis (Study I).

2. To measure the duration of apnoea without desaturation after induction of anaesthesia (Study I).

3. To follow the effects of 60 and 80 % oxygen compared with 100 % oxygen during preoxygenation and induction of anaesthesia on formation of atelectasis 4, 7 and 14 min after the start of preoxygenation (Study II).

4. To follow the time course of atelectasis formation using 80 % oxygen during preoxygenation and induction of anaesthesia 14, 21, 28, and 45 minutes after the start of preoxygenation (Study II).

5. To compare the area of postoperative atelectasis after pre- and postoxygenation with 100 % oxygen and a ventilatory strategy during anaesthesia, comprising the coherent use of CPAP/PEEP and a reduced fraction of end-tidal oxygen (FETO2) before ending CPAP mask ventilation after extubation with that of a control group after pre- and postoxygenation with 100 % oxygen but without CPAP/PEEP and a high FETO2 before ending mask ventilation (Study III).

6. To compare the influence of 80 or 100 % oxygen for pre- and postoxygenation in combination with this ventilatory strategy on the area of postoperative atelectasis (Study III).

7. To compare the influence of 30 or 100 % oxygen for postoxygenation in combination with coherent use of CPAP/PEEP until extubation on the area of postoperative atelectasis (Study IV).

8. To investigate whether using peripheral arterial oxygen saturation (SpO2) as an oxygenation index perioperatively can be used to identify patients with presumed differences in the area of atelectasis (Studies III and IV).
Materials and Methods

Patients

The Regional Ethics Committee (Uppsala, Sweden) approved the studies. A total of 135 adult patients gave their written informed consent and participated. The last study was registered internationally at ClinicalTrials.gov.

In studies I and II, only non-smoking women with American Society of Anesthesiologists physical status (ASA) class I or II and a body mass index (BMI) of ≤ 31 were enrolled. A total of 36 females were randomised to one of three preoxygenation groups with 60, 80, or 100 % oxygen (in nitrogen if applicable). Preoperative evaluation included haemoglobin level, electrocardiogram (ECG), arterial blood gases, spirometry, lung volumes and metabolic rate. All preoperative values were normal.

In study II, 27 of the patients from study I were included (early group) and 10 women with the same preoperative demographic characteristics as the patients in the early group were studied in a consecutive order (late group).

In study III, 30 non-smoking or smoking patients, excluding those with chronic obstructive pulmonary disease (COPD), with ASA class I-III and a BMI of ≤ 31 were included and randomised to one of three groups. Only routine preoperative evaluation as dictated by departmental policy was performed.

In study IV, 60 non-smoking or smoking patients (excluding COPD patients) with ASA class I-III and a BMI of < 35 were included and randomised to one of two groups. Only routine preoperative evaluation as dictated by departmental policy was performed.

Apparatus and monitoring

In all patients, ECG and SpO2 were continuously monitored during anaesthesia, as were end-tidal CO2 concentration and end-expiratory oxygen fraction (FETO2). Blood pressure was measured non-invasive and periodically with a CS 3 (Datex-Ohmeda, Helsinki, Finland), Infinity Delta XL (Draeger Medical Ag & Co. KG, Lubeck, Germany), or Philips IntelliVue MP Anesthesia (Philips Medizin Systeme, Boeblingen, Germany) monitor.
In studies I and II, the Mentell system was used for ventilation and spontaneous breathing (Anmedic, Vallentuna, Sweden). In studies III and IV, the Primus Draeger (Draeger Medical Ag & Co. KG) and the Datex-Ohmeda S/5 Avance (GE Healthcare, Datex-Ohmeda, Madison, WI, USA) were used for mechanical ventilation and spontaneous breathing, respectively, with both ventilators functioning as circle systems.

Neuromuscular function was monitored using a neuromuscular train-of-four (TOF) monitor from Draeger (Medical Ag & Co. KG) or the TOF-Watch S (Organon Ltd., Dublin, Ireland).

In studies I and II, a target control infusion (TCI) of propofol was delivered by a Diprifusor target-controlled infusor pump (Astra Zeneca, Macclesfield, Cheshire, United Kingdom). In studies III and IV, a TCI of propofol and remifentanil were delivered by a Care Fusion (Alaris Medical UK Ltd., Hampshire, UK) or the Agilia Injectomat TIVA (Fresenius Vial, Brezius, France).

SpO₂ was measured continuously postoperatively with an Infinity Masimo SET (Masimo Corporation, Irvine, CA, USA) or Ohmeda TuffSat (GE Healthcare Finland Oy, Kuortaneenkatu, Helsinki, Finland).

Anaesthesia

As a general principle, patients did not receive sedative premedication. However, on request, some patients were given a benzodiazepine, but the number of such patients did not differ between the randomisation groups.

In all four studies patients were investigated during (studies I and II) or after (studies III and IV) total intravenous anaesthesia (TIVA) with a TCI of propofol. In studies I and II, repeated doses of fentanyl and alfentanil complemented propofol, whereas in studies III and IV, a TCI of remifentanil was added. TIVA was administered in accordance to clinical signs of the depth of anaesthesia. To facilitate endotracheal intubation, in studies I, II, and III, rocuronium was given at a dose of ~0.5 mg/kg ideal body weight (IBW). All patients were checked for residual muscle relaxation block.

In study IV, a laryngeal mask airway was used without a muscle relaxant. To eliminate the postoperative need for opiates in all patients in studies III and IV, local anaesthesia complemented TIVA.

Ventilation

Preoxygenation lasted for 1 min before any anaesthetic drugs were given in all studies. Thereafter, induction of anaesthesia started, and the patients lost consciousness progressively. When spontaneous breathing ceased gradually,
ventilation was assisted through the mask. During preoxygenation and induction of anaesthesia a fresh gas flow of 6-8 L/min was used to avoid rebreathing and improve denitrogenation. In studies I and II, patients were randomised to one of three preoxygenation groups of 60, 80, or 100 % oxygen. The late group in study II comprised 10 consecutive patients who received 80 % oxygen for preoxygenation.

In studies I and II, patients breathed and were mechanically ventilated in volume-controlled ventilation (VCV) mode through the Mentell system, a hybrid low-flow anaesthetic circuit. This circuit had a built-in PEEP of 3 cmH₂O; no other level of PEEP was used. During maintenance of anaesthesia, a standard ventilatory setting was used, comprising 40 % inspiratory oxygen in nitrogen, a V̇ of 6-7 mL/kg and a respiratory rate of 12-11 breaths/min. Normocapnia, defined as an end-tidal CO₂ concentration of 4-6 %, was achieved by adjusting the fresh gas flow.

In studies III and IV, ventilators based on a circle system with advanced features were used in different modes and different levels of CPAP/PEEP according to the randomisation. During maintenance of anaesthesia a standard ventilatory setting was used, which comprised 30-40 % inspiratory oxygen in nitrogen, a V̇ of 6-7 mL/kg IBW and a respiratory rate of 8-11 breaths/min. Normocapnia (CO₂ concentration 4-6 %) was achieved by adjusting the minute ventilation.

In study III, two intervention groups were placed on 6 cmH₂O CPAP/PEEP from the start of preoxygenation to the end of mask ventilation after extubation; one group received 80 and the other 100 % oxygen during pre- and postoxygenation. In both groups, breathing by mask (CPAP 6 cmH₂O) with 30 % oxygen was started after extubation and aimed at an F̄ETO₂ ≤ 0.30. When patients in these intervention groups where spontaneously breathing, the ventilator was in the pressure-support ventilation (PSV) mode and during controlled ventilation in the pressure-controlled ventilation (PCV) mode. In the control group in study III, no CPAP/PEEP was applied, and 100 % oxygen was used for pre- and postoxygenation, including the period by mask breathing after extubation. When spontaneous breathing ceased in the control group, ventilation was assisted manually, and during controlled ventilation the ventilator was set to VCV mode.

In study IV, 100 % oxygen was used for preoxygenation in both groups. The ventilator was in PSV mode with CPAP at 6-8 cmH₂O during spontaneous breathing. During maintenance of anaesthesia, a standard ventilatory setting was used as described with the ventilator in the PCV-volume guaranteed mode and a PEEP of 6-8 cmH₂O. CPAP/PEEP was set to 6 cmH₂O when BMI was <25, to 7 cmH₂O when BMI was 25-30, and to 8 cmH₂O when BMI was ≥30. In study IV patients were randomised to postoxygenation with 30 or 100 % oxygen before extubation. After extubation all patients immediately breathed air.
Duration of apnoea without desaturation (DAWD)

DAWD was investigated in study I. Before recording DAWD, anaesthesia was induced after 1 min of strict preoxygenation. The recording of DAWD started at the time mask ventilation ended before intubation, and stopped when SpO₂ reached 90%. Anaesthesia was induced with the patient in a strict horizontal position on the CT table. Endotracheal intubation was performed at the beginning of the apnoea, and in all patients, the tube could be seen passing the vocal cords. All patients were intubated on the first attempt and correct placements were confirmed on the CT scan. During the recording of DAWD, the endotracheal tube was left open to air.

Computed Tomography of the lungs

Throughout the studies, the lungs were investigated with CT using a standardised procedure. All CT scans were transverse single-slice with 5 mm thickness. All studies included a basal scan 10-20 mm above the dome of the right diaphragm. The patient was placed in the supine position with the arms parallel to the body. All scans were performed in apnoea at the end of expiration.

In studies I and II, a control scan was taken in alert patients to exclude any atelectasis before anaesthesia. Remaining scans in studies I and II were performed during anaesthesia at different time points as measured from the start of preoxygenation. In studies III and IV, CT scans were performed only postoperatively.

The radiologist was unaware of the group affiliation of individual patients when the examining the CT scans. The atelectasis area in the scans was calculated using the region of interest (ROI) program included in the CT computer software. To trace the atelectasis, the window level and width were set at -500 and ±1500 Hounsfield units (HU), respectively. Atelectasis was defined as area with attenuation values between -100 and +100 HU and was expressed in cm² and as a percentage of the total area of the lung in a particular scan. The dorsal border was drawn manually between the atelectasis and the pleura. The ventral border was drawn manually between atelectasis and adjacent aerated lung area. The atelectasis area was calculated using the software program. Each ROI was redrawn twice, and the mean value was used.

When presenting results for the area of atelectasis, the aim is to display data in both cm² and per cent, however this is not always appropriate. When comparing data with those of earlier studies, the choice of data presentation in earlier studies governs the presentation in the present studies. A simple rule of thumb is to remember that atelectasis expressed in cm² corresponds approximately to half the value when expressed in percentage of the lung.
area, and consequently atelectasis expressed in percentage corresponds to twice the value in cm².

Another source of inconsistency relates to presenting descriptive statistical data as mean ± SD or as median and range (minimum to maximum), the former is used when indicated by comparisons with earlier studies.

The smallest possible value of atelectasis area that can be detected by CT has been assumed to be 0.1 cm². The theoretical resolution of CT is most likely greater than the inter-observer variability. However, even with this in mind, radiologists might report atelectasis as differing in size, but seldom disagree on the fact that atelectasis is present or not.

Arterial Oxygenation

In studies III and IV, oxygenation was estimated by measuring SpO₂ with the patient breathing air before the start of preoxygenation. In study III, SpO₂ was measured twice during stable anaesthesia with the patients ventilated with air. In studies III and IV, oxygenation was estimated continuously post-operatively after the end of mask ventilation.

Statistical analysis

The average and spread of data are described by median and range. In studies I, II, and III, the sample size was based on the assumption that a difference in the area of atelectasis of 50 % or more would be of clinical importance. In study IV the secondary dependent variable, SpO₂, determined the sample size. In all instances, α = 0.05 and 1 - β = 0.8 were assumed. This implied that 10-12 patients would be needed in each group in studies I-III and 30 patients in each group in study IV. In study IV the number of patients also included a margin for a high variability.

Few patients were included in studies I, II, and III, and the main variables were not assumed to have a normal distribution; therefore non-parametric tests were used to compare groups. The Kruskal-Wallis one-way analysis of variance was used to compare data between the three groups in study I. The Wilcoxon matched-pairs signed-rank test or the Mann-Whitney U test was used to compare two groups. In study III the Spearman rank correlation coefficient was used to analyse relationships between variables, and the Willett’s residual method for post hoc adjustment of the results was used to control for the influence of effect modifiers. For post hoc comparisons, Fisher’s exact test was used to analyse the occurrence of uneven distribution of potential effect modifiers.
In the post hoc analysis, because of the use of repeated comparisons, the results in the late group in study II were modified using Bonferroni correction. The correction was based on calculation of the cumulative Type I error and adjusting for the number of comparisons. Assuming an unadjusted $\alpha = 0.05$ and performing four comparisons, the adjusted $\alpha$ was 0.0125. Data in the late group was also analysed using a mixed model regression analysis with random effects for patients, and common effects for time and time squared, with $\alpha = 0.05$.

**Keeping track of time**

A professional sport watch with built-in memory was used during all studies, and the timer was started exactly as preoxygenation began. Specific peri-operative events were stored in sequential order as laps, and the time corresponding to the events could be retrieved from the watch memory.
Results

General

No adverse effects were observed in connection to the studies.

General demographic data for the 135 patients are presented in Table 1.

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<thead>
<tr>
<th>(\Sigma N=135)</th>
<th>Controls</th>
<th>Intervention group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients (n)</td>
<td>62</td>
<td>73</td>
</tr>
<tr>
<td>Age (years)</td>
<td>50±12</td>
<td>52±10</td>
</tr>
<tr>
<td>Sex (M/F)</td>
<td>13/49</td>
<td>23/50</td>
</tr>
<tr>
<td>Height (cm)</td>
<td>169±8</td>
<td>170±9</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>72±12</td>
<td>77±15</td>
</tr>
<tr>
<td>BMI (kg/m(^2))</td>
<td>25.4±3.2</td>
<td>26.4±3.7</td>
</tr>
<tr>
<td>ASA (n) I, II, III</td>
<td>41, 16, 5</td>
<td>48, 23, 2</td>
</tr>
<tr>
<td>Smokers</td>
<td>15/47</td>
<td>14/59</td>
</tr>
</tbody>
</table>

In this table, the patients are grouped into the control or intervention group. Sixty-two patients were classified into the control group, which comprised 12 patients from study I (100 % oxygen during preoxygenation), 10 patients from study II (80 % oxygen during preoxygenation, late group), 10 patients from study III (100 % oxygen during preoxygenation, no CPAP/PEEP), and 30 patients from study IV (100 % during both pre- and postoxygenation). Seventy-three patients were classified into the intervention group, which comprised 24 patients from study I (60 or 80 % oxygen during preoxygenation), 20 patients from study III (80 or 100 % oxygen during preoxygenation, 6 cmH\(_2\)O CPAP/PEEP), and 29 patients from study IV (100 % oxygen during preoxygenation, 30 % oxygen during postoxygenation).

The rationale for this division was based on the findings of the primary studies, in which patients had been randomly assigned to either a control or an intervention group. However, as explained later, the patients in the late group in study II were not classified into the combined intervention group but were assigned to the control group. The data from the 24 patients from study I, who were classified into the combined intervention group, were not
included in the analysis of the effect of intervention on the area of atelectasis; this is also explained later.

Duration of apnoea without desaturation (DAWD)

Figure 1. Shows the pattern of SpO₂ decrease during DAWD corresponding to the different oxygen concentrations.

The median DAWD was 389 s (range 239-528 s), 318 s (171-380 s), and 210 s (128-360 s) in the groups given 100, 80, or 60 % oxygen for preoxygenation, respectively, \((P < 0.01)\). The two patients in the 80 and 60 % groups with the shortest DAWD both had a BMI of 31. The patient in the 100 % group with the shortest DAWD had copious mucus production as found later during surgery.

Computed Tomography of the lungs and the area of atelectasis

In all patients investigated during anaesthesia (studies I and II), the control scan taken in conscious patients before anaesthesia showed normal lungs without atelectasis. The number of patients with atelectasis and the different oxygen concentrations, different levels of CPAP/PEEP, and different modes of ventilation at different time points perioperatively are presented in Table 2.
Table 2. Summary of oxygen fraction, the area of atelectasis (cm², median), time of CT, and interventions in studies I-IV

<table>
<thead>
<tr>
<th>Study</th>
<th>FIO₂ preoxy</th>
<th>Atelectasis cm²</th>
<th>CT time min</th>
<th>Apnoea ZEEP</th>
<th>FIO₂ before extubation</th>
<th>FIO₂ after extubation</th>
<th>CPAP induction</th>
<th>PEEP maintenance</th>
<th>CPAP after extubation</th>
<th>Ventilation mode</th>
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<tbody>
<tr>
<td>n=2</td>
<td>2</td>
<td>60</td>
<td>0</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
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<tr>
<td>n=4</td>
<td>2</td>
<td>80</td>
<td>0.1</td>
<td>4</td>
<td>NA</td>
<td>NA</td>
<td>0</td>
<td>3</td>
<td>NA</td>
<td>VCV</td>
</tr>
<tr>
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<td>4</td>
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<td>NA</td>
<td>0</td>
<td>3</td>
<td>NA</td>
<td>VCV</td>
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<tr>
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<td>2</td>
<td>60</td>
<td>0.1</td>
<td>7</td>
<td>NA</td>
<td>NA</td>
<td>0</td>
<td>3</td>
<td>NA</td>
<td>VCV</td>
</tr>
<tr>
<td>n=10</td>
<td>2</td>
<td>80</td>
<td>0.2</td>
<td>7</td>
<td>NA</td>
<td>NA</td>
<td>0</td>
<td>3</td>
<td>NA</td>
<td>VCV</td>
</tr>
<tr>
<td>n=9</td>
<td>2</td>
<td>100</td>
<td>0.5</td>
<td>7</td>
<td>NA</td>
<td>NA</td>
<td>0</td>
<td>3</td>
<td>NA</td>
<td>VCV</td>
</tr>
<tr>
<td>n=12</td>
<td>1</td>
<td>60</td>
<td>0.2</td>
<td>10.5</td>
<td>3.5</td>
<td>NA</td>
<td>0</td>
<td>3</td>
<td>NA</td>
<td>VCV</td>
</tr>
<tr>
<td>n=12</td>
<td>1</td>
<td>80</td>
<td>0.7</td>
<td>12</td>
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<td>3</td>
<td>NA</td>
<td>VCV</td>
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<tr>
<td>n=6</td>
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<td>NA</td>
<td>0</td>
<td>3</td>
<td>NA</td>
<td>VCV</td>
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<td>1</td>
<td>80</td>
<td>0.4</td>
<td>14</td>
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<td>NA</td>
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<td>3</td>
<td>NA</td>
<td>VCV</td>
</tr>
<tr>
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<td>100</td>
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<td>8.5</td>
<td>+25</td>
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</tr>
<tr>
<td>n=29</td>
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<td>30</td>
<td>21</td>
<td>6-8</td>
<td>6-8</td>
<td>0</td>
</tr>
</tbody>
</table>

FIO₂ preoxy = fraction of inspired oxygen during preoxygenation. CT = computed tomography. ZEEP = zero end expiratory pressure. CPAP = continuous positive pressure. PEEP = positive end expiratory pressure. VCV = volume controlled ventilation. PCV = pressure controlled ventilation. PCV-VG = pressure controlled ventilation – volume guaranteed.
Performing the scanning procedure, including positioning the patients in studies I and II, took 1.5 min. Intubation was complete after 5.5 min after the start of preoxygenation (i.e., 4.5 min after start of induction in studies I and II.) The results from 4 to 45 min apply to the patients in studies I and II investigated during anaesthesia.

4 min: A total of nine patients with a PEEP of 3 cmH₂O were studied 4 min after the start of preoxygenation (i.e., only 3 min after start of induction. Regardless of oxygen concentration, atelectasis was negligible.

7 min: A total of 26 patients with a PEEP of 3 cmH₂O were studied 7 min after the start of preoxygenation (i.e., 6 min after start of induction and 1.5 min after intubation). Tiny areas of atelectasis were found in all oxygenation groups, but the areas were slightly larger with 100 % oxygen.

10.5-14 min: A total of 36 patients with a PEEP of 3 cmH₂O were studied 10.5, 12, and 14 min after the start of preoxygenation. This corresponded to 9.5, 11, and 13 min after the start of induction and 5, 6.5 and 8.5 min after intubation in the groups given 60, 80, and 100 % oxygen for preoxygenation, respectively (n = 12 patients in each group). The scans taken at these occasions were performed after a period of apnoea (3.5, 5, and 7 min) corresponding to the different oxygenation levels used during preoxygenation. Small areas of atelectasis were found in the groups given 60 or 80 % oxygen during preoxygenation, and large areas of atelectasis were present in the 100 % group (P < 0.001).

14 min: A total of 12 patients, six given 60 % and six given 80 % oxygen during preoxygenation, with a PEEP of 3 cmH₂O, were studied 14 min after the start of preoxygenation (i.e., 13 min. after start of induction and 8.5 min. after intubation.) These 12 patients were also investigated immediately after a period of apnoea, as reported in the preceding paragraph, which allowed us to compare the area of atelectasis after apnoea with the area after some more minutes with controlled ventilation in the same patients. This also made it possible to compare atelectasis at the same time point, 14 min after the start of preoxygenation, in all three oxygenation groups. Only small areas of atelectasis of similar size were found regardless of whether the oxygen concentration was 60 or 80 %, or the timing of the CT scans.

14-45 min: A total of 10 non-randomised, consecutive patients, all given 80 % oxygen during preoxygenation and induction, and 40 % oxygen during maintenance of anaesthesia with a PEEP of 3 cmH₂O, were studied 14, 21, 28, and 45 min after the start of preoxygenation. In contrast to the patients
reported in the preceding paragraphs, apnoea tolerance time was not investigated in this trial.

The median areas of atelectases at the referred time points were 4.1 cm$^2$ (range 1.3-12.3 cm$^2$), 5.4 cm$^2$ (1.7-16 cm$^2$), 6.3 cm$^2$ (1.8-18 cm$^2$), and 7.3 cm$^2$ (2.0-21 cm$^2$), respectively. The median areas of atelectasis expressed as a percentage of the total lung area were 2.3 % (range 0.8-7.7 %), 2.9 % (1.0-9.8 %), 3.5 % (1.1-11.0 %), and 4.2 % (1.1-12.8 %), respectively.

Wilcoxon signed-ranks tests to compare the areas of atelectasis between awake and 14 min, between 14 and 21 min, between 21 and 28 min, and between 28 and 45 min, showed that, after a Bonferroni correction, the two first comparisons (both $P = 0.005$) were significant, but the last two were not significant (both $P = 0.028$), $P = 0.0125$. In a post hoc analysis of a common time effect for all patients, we found a quadratic function of time, with a linear term showing an increase of atelectasis of 0.3 cm$^2$ per min, ($P = 0.002$), while the quadratic term showed that this effect gradually decreased with 0.004 cm$^2$ per min squared ($P = 0.002$).

**Study III**, 25 min postoperatively: A total of 30 patients were randomised according to protocol as described under “Ventilation”. All patients breathed air during the CT investigation. The median areas of atelectasis were 5.2 cm$^2$ (range 1.6-12.2 cm$^2$) and 8.5 cm$^2$ (3-23.1 cm$^2$) in the groups given an $F_1O_2$ of 1.0 with or without CPAP/PEEP (median areas expressed in %, and range, were 2.4; 0.7-5.1, and 3.9; 1.7-10.5, respectively). In the group given an $F_1O_2$ of 0.8, in which seven patients were ex- or current smokers, the median area of atelectasis was 8.2 cm$^2$ (range 1.8-14.7 cm$^2$) (area in % was 4.4; 1.1-5.8). Fisher’s exact test showed that smoking was significantly more common in the the group given an $F_1O_2$ of 0.8 compared to the control group, $P < 0.05$.

In the primary analysis, no statistical significances were found between groups in the area of atelectasis. In the post hoc analysis, after correction for BMI and age, the difference between the two groups given an $F_1O_2$ of 1.0 was significant ($P = 0.016$). The Spearman correlations coefficients were 0.46 ($P < 0.05$) between atelectasis and age, and 0.16 ($P = 0.39$) between atelectasis and BMI.

**Study IV**, 14 min postoperatively: A total of 60 patients were randomised according to the protocol described under “Ventilation”. The total of patients investigated and analysed were 59, 1 patient was excluded because of a technical problem with the CT scanner. All patients breathed air during the CT investigation. The median areas of atelectasis were 5.5 cm$^2$ (range 0-16.9 cm$^2$) and 6.8 cm$^2$ (0-27.5 cm$^2$) in the groups given an $F_1O_2$ of 0.3 or 1.0 before extubation, respectively; this difference was not significant (median areas expressed in %, and range, were 2.7; 0-9.9, and 3.9; 0-14, respectively)
The post hoc analysis using the exact Wilcoxon test showed a significant difference ($P = 0.038$) in the square root of the area of atelectasis between non-smokers, defined as patients ($n = 41$) smoking less than 6 pack-years compared with smoking patients ($n = 18$), defined as smoking more than 6 pack-years (median 22; range 7-43 pack-years). The median areas of atelectasis were 5.4 cm$^2$ (range 0-19.1 cm$^2$) in non-smokers, and 9.0 cm$^2$ (1.3-27.5 cm$^2$) in smokers.

The median atelectasis areas were 5.3 cm$^2$ (range 0-11.5 cm$^2$) in ASA class I patients ($n=34$) and 8.7 cm$^2$ (1.3-27.5 cm$^2$) in ASA class II-III patients ($n=25$). The difference in the square root area of atelectasis between these two groups was also significant ($P = 0.015$). Finally, we performed post hoc analysis of the interaction between being a non-smoker or smoker/ex-smoker and ASA class. The areas of atelectasis in the group of smoker/ex-smoker did not differ from the group of non-smoker, neither if classified as ASA class I, nor for those classified as ASA II-III.

Oxygenation

Table 3 shows the SpO$_2$ of patients breathing air before the start of preoxygenation in the different groups in studies III and IV, during test 1 and 2 with the patients ventilated with air during anaesthesia in study III, and postoperatively in the different groups in studies III and IV.

Table 3. Peripheral oxygen saturation, (SpO$_2$) in % (median and range) in the study groups measured with $FIO_2 = 0.21$.

<table>
<thead>
<tr>
<th>$FIO_2$ x, y, z = fraction of inspired oxygen for preoxygenation, postoxygenation, and after extubation, respectively ZEEP = zero end expiratory pressure. CPAP = continuous positive airway pressure. PEEP = positive end expiratory pressure. preop = preoperatively in the supine position before anaesthesia. postop = postoperatively in the supine position after anaesthesia. CT = computed tomography.</th>
<th>$FIO_2$</th>
<th>$FIO_2$</th>
<th>$FIO_2$</th>
<th>$FIO_2$</th>
<th>$FIO_2$</th>
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<tbody>
<tr>
<td>Study</td>
<td>III, n=10</td>
<td>III, n=10</td>
<td>III, n=10</td>
<td>IV, n=30</td>
<td>IV=29</td>
</tr>
<tr>
<td>$SpO_2$ preop</td>
<td>99; 95-100</td>
<td>98; 94-99</td>
<td>99, 96-100</td>
<td>98; 95-100</td>
<td>99; 95-100</td>
</tr>
<tr>
<td>Test 1</td>
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<td>96; 92-98*</td>
<td>95; 93-97*</td>
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<td>NA</td>
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<td>95; 93-96*</td>
<td>95; 92-98*</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>$SpO_2$ postop</td>
<td>9/10 ≥ 94</td>
<td>9/10 ≥ 94</td>
<td>9/10 ≥ 94</td>
<td>94; 87-98</td>
<td>93; 87-98</td>
</tr>
<tr>
<td>$SpO_2$ at CT</td>
<td>All ≥ 94</td>
<td>All ≥ 94</td>
<td>All ≥ 94</td>
<td>All ≥ 94</td>
<td>All ≥ 94</td>
</tr>
</tbody>
</table>
Test 1 and 2 were performed during anaesthesia. * P < 0.025, Wilcoxon signed-rank test (limited because of Bonferroni correction) comparing SpO₂ values in patients in the same group preoperatively with values during anaesthesia. No differences between groups at any time.

The SpO₂ when breathing air did not differ significantly between groups before, during or after anaesthesia. The median differences in SpO₂, observed by comparing values immediately before anaesthesia with values measured twice during ventilation with air during anaesthesia in study III, were significant in both groups.

Post hoc analysis of pooled data from studies I-IV

As mentioned earlier, the results from the 24 patients from study I (given 60 or 80 % oxygen during preoxygenation), classified in the combined intervention group, is not included in this overview. The reason is that the results from these patients in study I cannot be regarded as providing any information about the final development of atelectasis, because the CT-scan was performed early.

The irrelevance of early CT-investigations using low oxygen concentrations during preoxygenation was demonstrated in study II, which showed that using 80 % oxygen during preoxygenation is probably not better than conventional preoxygenation (i.e., using 100 % oxygen, assuming no CPAP/PEEP). Therefore, the results from the late group in study II was transferred to the control group, which now included a “failed intervention”. Another indication that 80 % oxygen is a failed intervention comes from the result in study III, in which the group given 80 % oxygen for pre- and post-oxygenation had as much atelectasis as the control group. However, in this trial, 80 % oxygen was used with CPAP/PEEP, and therefore, it was still defined as an intervention. As noted earlier, 7/10 patients in this group were ex- or current smokers.

An overview, showing the number of patients and the area of atelectasis as a percentage of total lung area 1 cm above the right diaphragm, is presented in Figure 2.

The reason for showing n = 111 (135-24) in the histogram was explained in the first sentence of the preceding paragraph. The median areas of atelectasis were 4.2 % (range 0-13.2 %) in the control group (n=62) and 2.8 % (0-9.9 %) in the intervention group (n=49) (P = 0.021, Mann-Whitney U test) (Figure 3).
Figure 2. Number of patients and the area of atelectasis as a percentage of total lung area, 1 cm above the right diaphragm. For definitions, see the text.

Figure 3. Area of postoperative atelectasis as a percentage of total lung area, 1 cm above the right diaphragm in the combined control group and the combined intervention group. The box represents the first to the third quartile. The dark line is the median. Dots indicate outliers. *P = 0.021.
A histogram showing the distribution of patients in relation to age is presented in Figure 4. A significant linear correlation coefficient for atelectasis and age of 0.38 ($P = 0.008$) was found in the intervention group but not in the control group.

**Figure 4.** Number of patients according to age in patients with atelectasis that was presumed to be fully developed. The area of atelectasis is presented as a percentage of total lung area 1 cm above the right diaphragm.
Discussion

Duration of apnoea without desaturation (DAWD)

An expected result was that the DAWD was significantly shortened using less than 100 % oxygen for preoxygenation. In general, the DAWD in this study was shorter compared with the value in more optimal procedures because we used only normal breathing for 60 s, in contrast to normal breathing for 3 min\textsuperscript{32} or with the eight deep-breaths technique for 60 s\textsuperscript{33}. The horizontal position of the patient on the CT-table probably also contributed to the reduced DAWD.

Methodological consideration: Why a short duration of preoxygenation?

The vital importance of the oxygen concentration during preoxygenation and its duration on the development of atelectasis has been demonstrated in theoretical and clinical studies\textsuperscript{34-36}. Consistently, strict preoxygenation (i.e., before any drug was administered), lasted only 1 min in all four studies. After 1 min of preoxygenation with 100 % oxygen at normal tidal ventilation, denitrogenation is about 80 % complete\textsuperscript{37}. We chose to study this duration for preoxygenation, because it was the routine in our department. The consequence was a systematically reduced $F_{ET}O_2$ at the time of starting induction compared with the recommended duration of preoxygenation of 3 min.

The more important reason for adopting this relatively short time for preoxygenation was to avoid exaggerating the extent of atelectasis in our studies compared with what can occur during routine anaesthesia, by assuming that a shorter duration of preoxygenation is more common in practise than a longer duration. That said, we measured $F_{ET}O_2$ continuously but recorded the values only at the end of mask ventilation.
Computed Tomography of the lungs and the area of atelectasis

The area of atelectasis in the basal CT scans, general observations from study I and II

The early and late atelectasis formation, as found in studies I and II, clearly showed that atelectasis formation is an oxygen- and time-dependent process. The development of static atelectasis was dependent on the $F_{1}O_2$, especially during preoxygenation, but also during induction (i.e., the timing of total airway closure, as proposed in a mathematical model and in a clinical trial). In the mathematical model, the airway to the dependent parts of the lungs closes with induction of anaesthesia, and these areas then behave as a closed collapsible cavity. The model assumed the full effect of preoxygenation on denitrogenation (3 min) and that a closed collapsible cavity formed immediately at induction.

In our clinical studies, even though preoxygenation was for only 1 min, the appearance of total airway closure in the dependent parts of the lungs was not instantaneous, rather more gradual, probably giving some “preoxygenation effect” also during the induction of anaesthesia. With 100% oxygen for preoxygenation and induction, the model predicts alveolar collapse in 7.9 (no HPV) or 8.7 (intact HPV) min. Only small areas of atelectasis were seen in patients immediately after intubation, 6 min after the start of induction, which fits with the model. We know that atelectasis formation increased markedly between 6 and 13 min after the start of induction in the group given 100% oxygen, but the exact time of development could not be identified. Thus, we cannot rule out that the atelectasis in this group was complete some minutes before the time when it was observed. On the other hand, we cannot rule out another possibility: that the atelectasis area increased beyond 13 min. However, the area of atelectasis in this work is similar to that reported by earlier studies.

It should be observed that when presenting the area in percentage, some earlier studies calculated this in proportion of the total inner thoracic area. In the current studies the calculations were in proportion of the total lung area.

The model also predicts that alveolar collapse will ensue after 8.7 (no HPV) or 9.3 (intact HPV) min after preoxygenation with 100% oxygen, followed by 80% oxygen during induction of anaesthesia. The model does not incorporate the effect of preoxygenation with 80% or 60% oxygen, but the importance of the duration of preoxygenation was demonstrated, and as stated most of the effect occurred during the first three minutes.

Atelectasis in our late group given 80% oxygen increased progressively during anaesthesia but more slowly as time passed and appeared as static atelectasis. This mechanism was demonstrated in 1989.
In a post hoc analysis, a common time effect on the development of atelectasis during the study period in the late group could be established, using a mixed model regression analysis. The exact values to describe the linear progression of atelectasis and the exponential opposing effect, expressed in cm$^2$, is of no practical use, but it illustrates the nature of the development of static atelectasis as an oxygen- and time dependent process.

The areas of atelectasis in the 60 and 80 % groups in study I found after 14 min were probably influenced by the preceding prolonged period of apnoea, which is discussed more in detail in the following paragraph.

**How did the investigation of atelectasis and DAWD in the same occasion in study I affect the comparisons between the early (study I) and late 80 % group (study II)?**

In study I, the patients were apnoeic until desaturation before atelectasis was measured, whereas in study II, the late group was studied without any prolonged period of apnoea. The median areas of atelectasis 14 min after the start of preoxygenation were 0.6 cm$^2$ (range 0.1–3.2) in the patients given 80 % oxygen in study I and 4.1 cm$^2$ (1.3–12.3 cm$^2$) in the patients given 80 % oxygen in the late group in study II. The patients’ characteristics, level of PEEP, and other ventilatory variables did not differ between these groups of patients.

This raises the question of what could explain the difference in atelectasis formation between these two groups. Putting aside the possibility that the result of study I could represent a type 1 error or that the result of study II might represent a type 2 error, the exposure to prolonged apnoea probably explained the difference between groups. In study I patients were apnoeic with an endotracheal tube in place, which was not connected to the ventilator during the DAWD, leaving the lungs open to apnoeic oxygenation at ZEEP with air$^{40,41}$. A fresh gas flow of 6 L/min with 40 % oxygen in nitrogen was used when ventilation was started after completion of the apnoea period. Analysing this situation in terms of the risk of inducing static or dynamic atelectasis formation, ZEEP might increase the area of static atelectasis on the one hand, but air and 40 % oxygen might reduce the area of dynamic atelectasis on the other hand.

Without prolonged apnoea, ventilation in the late group started immediately after intubation with a PEEP of 3 cmH$_2$O. During the time when the lungs were auscultated for bilateral breaths sounds and when securing the endotracheal tube, a fresh gas flow of 6 L/min with 80 % oxygen in nitrogen was used. As soon as a normal capnography curve was identified and bilateral breaths sounds were confirmed, the fresh gas flow was set to 1 L/min with
40 % oxygen in nitrogen. These findings suggest that a PEEP of 3 cmH₂O instead of ZEEP might reduce the area of static atelectasis, but the period with 80 % oxygen after intubation and the time for the inspiratory oxygen concentration to decrease from about 80 % to 40 % with a fresh gas flow of 1 L/min might increase the area of dynamic atelectasis. The duration of the period with 80 % oxygen after intubation was not recorded, neither the time it took to decrease the oxygen concentration from 80 % to 40 %, but it might be estimated as several minutes. Thus, it appears that there may have been opposing effects in the two groups regarding the tendency towards static or dynamic atelectasis, which might have cancelled each other. However, one reasonable hypothesis is that the exposure to a prolonged period of 80 % oxygen after intubation in the late group was more important than ZEEP in the early group.

The results during anaesthesia 45 min after starting preoxygenation in the late group showed that the median atelectasis area had increased to 7.3 cm² (range 2.0-21 cm²). During the interval from 14 to 45 min, ventilation was unchanged with 40 % oxygen and a PEEP of 3 cmH₂O. In an earlier study, that presumably used 100 % oxygen during preoxygenation, the area of atelectasis during anaesthesia was stable during an interval of about 50 min with 40 % oxygen at ZEEP. Therefore, it is unlikely that the increased area of atelectasis can be explained by the ventilation variables used between 14 to 45 minutes in our late group.

Rather, the increased area of atelectasis might indicate that closed sections in the dependent parts of the lungs had become more atelectatic as oxygen left more quickly than nitrogen entered. In other words, most of the final atelectasis was potentially static and the full extent of atelectasis occurred as time passed without opening the airway with a high enough PEEP or a high enough inspiratory pressure. However, it is uncertain whether this line of reasoning can fully explain the finding of smaller atelectasis area in the 80 % group submitted to a prolonged apnoea. Certainly, a prolonged period of apnoea does not happen often, and the increased area of atelectasis with 80 % oxygen for preoxygenation that develops over time offers no real benefit, but a reduced DAWD is always negative.
Is there a window of opportunity?

As seen in figure 5 the atelectasis area immediately after intubation was small in most patients, regardless of the oxygen concentration during pre-oxygenation.

![Figure 5. Increase in the area of atelectasis 1 cm above the right diaphragm during anaesthesia in individual patients in the early group treated with 60, 80, or 100 % oxygen for preoxygenation from awake to 4, 7, and 14 min after the start of preoxygenation.](image)

This raises the question of whether there is a time window during which different methods for preventing further atelectasis formation can be implemented, and whether a lower F\textsubscript{1}O\textsubscript{2} during preoxygenation extends this time window. Atelectasis can be reversed by a RM\textsuperscript{43-45}, but performing a full scale RM (up to 40 cmH\textsubscript{2}O in non-obese) immediately after intubation might be unwise because it can accentuate hypotension. Any intervention at this time point of anaesthesia should be simple and secure.

Interestingly, a recently published study on the use of “protective ventilation”\textsuperscript{6} during anaesthesia implemented a RM with 40 % oxygen up to a plat-
eau pressure of 30 cmH$_2$O immediately after induction, which must have been very early after intubation and perhaps within the time frame outlined in study II.

**The area of atelectasis in the basal CT scans, general observations from study III**

The time point of investigation is of obvious concern when interpreting different studies. The “normal” amount of atelectasis during anaesthesia was proposed as 3-4 % of the total lung area in a basal scan$^1$. It is unknown what percentage should be considered a normal amount of postoperative atelectasis. A study in 13 patients who were investigated 2 hours after lower abdominal surgery$^{46}$ showed that 11 patients developed atelectasis and that ~1 % of the intrathoracic area (~2 % of the lung area) was atelectatic.

In a 2002 study$^{47}$ of high relevance, a vital capacity manoeuvre was performed, i.e a RM up to 40 cmH$_2$O, 10 min before the presumed end of surgery, the patients were randomly assigned to an F$_1$O$_2$ of 1.0 or 0.4 before extubation, and no PEEP was used. Twenty min after extubation, the atelectatic areas comprised 2.6 % ± 1.1 % (mean ± SD) in the group given an F$_1$O$_2$ of 0.4, and 6.8 % ± 3.4 % in the group given an F$_1$O$_2$ of 1.0. This study also included a control group without a vital capacity manoeuvre but with an F$_1$O$_2$ of 1.0 before tracheal extubation. Atelectasis in this group was 8.3 % ± 6.2 %. The patients in this study were all preoxygenated with 100 % oxygen for 3 min, and BMI was <30 in all patients.

At least two important conclusions can be drawn from the information discussed above. First, the area of atelectasis can be almost eliminated by using a vital capacity manoeuvre and low oxygen concentration before extubation. Second, the area of atelectasis postoperatively is probably larger after both pre- and postoxygenation with 100 % oxygen than after preoxygenation with 100 % oxygen but administration of 40 % oxygen during anaesthesia, i.e. before postoxygenation and extubation. Study III investigated the area of atelectasis about the same time point postoperatively as in the study discussed above, and the results were similar. However, probably because of differences in methodologies and study populations between these studies, the results in study III were significant only after a post hoc adjustment for age, or BMI, or both.

The control group in study III had an area of atelectasis of only 5.0 % ± 2.9 % (mean ± SD), compared with 8.3 % ± 6.2 % (mean ± SD) in the study discussed above, suggesting that the short period of preoxygenation may have reduced atelectasis formation. The results for the intervention groups are similar in the two studies: 2.6 % ± 1.1 % (mean ± SD) in the mentioned trial and 2.6 % ± 1.5 % (mean ± SD) in study III. The result in
study III suggest that even without a vital capacity manoeuvre, the coherent use of CPAP/PEEP and low \( F_{102} \) after extubation is as effective in reducing atelectasis as a vital capacity manoeuvre and a low \( F_{102} \) before extubation.

In terms of patient safety, the technique in study III omitted the vital capacity manoeuvre and included CPAP initially with on-mask 100 % oxygen after extubation; i.e.; \( F_{102} \) decreased only after securing a free upper airway and a satisfactory spontaneous breathing. This means that immediately after extubation, the safe apnoea time in the event of a total failure of ventilation should be compared between patients with an \( F_{102} \) of 1.0 and increased FRC because of CPAP, and patients with an \( F_{102} \) of 0.4 and no CPAP.

The area of atelectasis in the basal CT scans, general observations from study IV

The primary study objective was to investigate whether reducing the oxygen concentration from 100 to 30 % during emergence from anaesthesia would reduce the area of atelectasis. This objective could not be achieved. However, it is of interest to look at the results in the context of the preceding paragraph. In study IV, the area of atelectasis was 3.4 % ± 2.4 % (mean ± SD) in the intervention group, and 4.4 % ± 3.4 % (mean ± SD) in the control group. Although the results in the intervention groups were similar between studies III (2.6 % ± 1.5 %), IV (3.4 % ± 2.4 %), and the study discussed above from 2002 (2.6 % ± 1.1 %), very different results were found in the control groups, in which the area of atelectasis ranged from 4.4 % ± 3.4 % (IV), 5.0 % ± 2.9 % (III) to 8.3 % ± 6.2 % in the study from 2002. This result in the control group in study IV probably reflects the fact that CPAP/PEEP was used during preoxygenation and anaesthesia; the only difference from the intervention group was the use of 100 % instead of 30 % oxygen during postoxygenation.

The ventilation strategies used in studies III and IV did not achieve the goal of delivering a “clean” lung to the postoperative ward. The level of CPAP/PEEP might have been insufficient for the task of keeping the small airways open in dependent parts of the lungs during anaesthesia; 10 cmH\(_2\)O has been proposed as the optimal PEEP in normal-weight patients after an RM\(^{48}\).

More on the topic of lung volumes and small airway closure

The end-expiratory lung volume and airway closure

One important aspect in the nomenclature of lung volumes differs from ordinary use, going from a conscious to an anaesthetic state. The term denoting
resting volume in the lungs after expiration is called the functional residual
capacity (FRC) during normal spontaneous breathing and end expiratory
lung volume (EELV) during anaesthesia. This is in order to highlight the fact
that the balancing forces involved are changed. In 1963\textsuperscript{49}, the first report
demonstrating decreased FRC during anaesthesia was published. FRC in-
creases slightly by age, it is linearly related to height, and the supine position
will reduce FRC compared with the upright position in the range of 500 to
1000 ml\textsuperscript{50}. Obesity can induce significant reduction in FRC\textsuperscript{51}. With the ex-
ception of ketamine, all anaesthetic drugs have been confirmed to reduce
FRC in the awake supine position by 400 to 500 ml, or by 15-20 % of the
awake value. Thus, anaesthesia induction has an immediate effect on the
respiratory system\textsuperscript{52}. The combined effect of the supine position and anaes-
thesia will give an EELV close to residual volume (RV). EELV is the sum of
RV and end expiratory reserve volume (ERV), which means that ERV is
decreased first by the supine position and later by anaesthesia.

A reduction in ERV and therefore in EELV must be considered along
with the closing volume (CV) and the CC (RV + CV); i.e. the vol-
ume/capacity at which small airways in the dependent parts of the lungs will
close. CC increases by age and equals FRC in the supine position at the age
of 45-50 years and in the standing position at the age of 65-70 years.

Whereas a reduced resting volume in the lungs from FRC to EELV is
firmly established, a consistent reduction in CV or CC during anaesthesia is
disputed\textsuperscript{30, 53-55}. The conflicting results might be explained by differences in
the maximal inspiratory pressure used to inflate the lungs before the meas-
urements: 30 or 40 cmH\textsubscript{2}O. Only the latter pressure will reliably open up the
lungs to vital capacity\textsuperscript{45}. Thus, overall, the data favours the interpretation
that CV will be larger than ERV in many patients during anaesthesia, which
implies partial or total airway closure during the respiratory cycle. The de-
gree of airway closure will depend on the actual difference between CV and
ERV in relation to the size of the V\textsubscript{T}. Use of PEEP immediately increases
ERV and thus reduces the risk of airway closure.

In this context, we note that FRC are higher in the lungs of patients with
asthma or COPD than in normal lungs. However, the increased FRC in these
diseases occurs because of increased RV, not increased ERV. Because CV
also increases in asthma and COPD, these patients exhibit more extensive
airway closure than what is normally found.

The missing links

These data suggest that a correlation between atelectasis and airway closure,
and between atelectasis and age should have been found in earlier studies,
especially concerning the fact that airway closure is the main explanation for
the observed deterioration in arterial oxygenation with age and that airway
closure increases even more during anaesthesia. However, no such correla-
tions have been found. Based on the findings in the present studies, we present tentative theoretical explanations for this inconsistency. To begin, these explanations are explored by looking at possible implications related to patient factors.

**Patient factors**

*Age and atelectasis*

In 1987, Strandberg et al.\(^{56}\) published a study of 38 patients on the constitutional factors correlated with the increased extent of atelectasis. They found no indications of atelectasis being more abundant because of gender, age or smoking habits. Calculation of BMI was not reported. Instead, they found Broca’s index (ideal weight in kg=height in cm - 100) and weight to be correlated to atelectasis, the linear correlation coefficients reported as 0.34 and 0.28 \((P < 0.05)\). These patients were ventilated with a Servo 900C, the level of PEEP was not reported.

The results had not been refuted until tentative findings in studies III and IV, in which CPAP/PEEP was used to counteract for the reduced ERV during anaesthesia. Moreover, in study IV some adjustments of CPAP/PEEP were made according to BMI. Linear correlation coefficients between atelectasis and BMI were only \(~0.2\) in studies III and IV. This is a reasonable finding, indicating that the applied CPAP/PEEP was effective against ERV reduction related to overweight. In study III, the linear correlation coefficient for atelectasis and age was 0.40 \((P = 0.03)\) and the Spearman correlation coefficient for atelectasis and age was 0.46 \((P < 0.05)\). In study IV, in which CPAP/PEEP was higher than in study III, no linear correlation was found between age and atelectasis for the whole study group. A significant linear correlation was found \((r = 0.37, P = 0.048)\) in the group given 30 % oxygen but not for the group given 100 % oxygen \((r = 0.14)\). However, there were significantly more patients classified as smokers in 100 % group.

Thus, with the use of CPAP/PEEP, what is the possible explanation behind the preliminary finding of a correlation between age and atelectasis in study III?

*Age and the duration of preoxygenation*

To explain this preliminary finding, we start from the very beginning - during preoxygenation and induction. If we acknowledge that airway closure increases with age and is more likely to occur in the supine position than in the sitting position, the time to achieve a final \(F_{ET}O_2\) should be longer in older patients than in the young ones. Studies on \(F_{ET}O_2\) confirm this.\(^{57-59}\) If alveolar minute ventilation is depressed, more time is needed for an older patient to reach the same \(F_{ET}O_2\) as in the young.
Another aspect is that the areas of low $V_A/Q$ that result from airway closure, respond completely to oxygen treatment. Thus, after some time, there should be no difference in the $F_{ETO_2}$ between older and younger patients. Studies of preoxygenation suggest that the critical time lies 1-2 min from start of preoxygenation. Premedication that reduces minute ventilation and leaks around the face mask will both prolong the critical time.

Knowing this, and the pivotal role of the oxygen concentration behind closed or semi-closed airways for events to come, any study on the early formation of atelectasis conducted without control of these parameters will probably have a systematic error. Inducing anaesthesia without having exact control of the duration of preoxygenation and the timing of $F_{ETO_2}$, in relation to the fundamental effects of anaesthetic drugs on the respiratory muscle tone and thus ERV, probably creates a situation where younger patients will have higher $F_{ETO_2}$ than older patients. In younger patients, higher $F_{ETO_2}$ will increase the early formation of atelectasis, and in older patients, lower $F_{ETO_2}$ will reduce the early formation of atelectasis.

**Age and CPAP/PEEP**

Application of CPAP/PEEP will fundamentally change the interaction between $F_{ETO_2}$ and airway closure, but most studies of atelectasis have been performed without CPAP during preoxygenation and induction, and in many studies even without (or not reporting) the use of PEEP during anaesthesia. Looking at atelectasis formation over time will yield different results depending on the conditions. However, it may be proposed that at the beginning of anaesthesia, a low oxygen concentration behind permanently closed airways will not have sufficient time to promote atelectasis. Consistent use of CPAP/PEEP during preoxygenation, induction and maintenance of anaesthesia will have a profound effect on the events described in the preceding paragraph. The difference in $F_{ETO_2}$ related to age will be much reduced, but the age-dependent airway closure will also be reduced as long as CPAP/PEEP is maintained.

Returning to the correlation between atelectasis and age found in study III, we propose that the patients of different ages in study III were more homogeneous regarding $F_{ETO_2}$ during induction of anaesthesia. After induction and intubation atelectasis develops according to the level of $F_{ETO_2}$ and the amount of airway closure. If the level of PEEP is insufficient in relation to $F_{ETO_2}$, static and/or dynamic atelectasis will follow. In study III and IV, the levels of CPAP/PEEP were fixed at 6-8 cmH\textsubscript{2}O and these levels might be too low to fully avoid airway closure. An insufficient level of PEEP will affect older patients more as closing volume increases with age. Assuming no difference in $F_{ETO_2}$ at the time of intubation, older patients will have more airway closure after intubation and therefore an increased risk of atelectasis.
The time it takes exchanging high $F_{ET}O_2$ to low will probably influence the risk.
Thus, applying a CPAP/PEEP technique in older patients could become a double-edged sword. On the one hand, better oxygenation and less airway closure should occur, provided that CPAP/PEEP is working; on the other hand, if left without this protection, the risk of formation of atelectasis would increase if $F_{ET}O_2$ remained high.

As studies III and IV investigated the occurrence of postoperative atelectasis, more than one hour had passed since induction of anaesthesia, letting more of the final development of atelectasis to reveal itself. The effect of reducing the $F_{ET}O_2$ after extubation with an $F_iO_2$ of 0.3, as in study III, will not be uncovered until after ~3 hours. We did not include a group in which 100% oxygen was given with CPAP on mask after extubation. Inclusion of such a group might have given insights into the importance of high $F_{ET}O_2$ after ending CPAP, but a similar group was studied by Lumb et al. Lumb et al used the alveolar-arterial $PaO_2$-difference as a surrogate marker for atelectasis.

Their data were collected postoperatively after a RM and the use of CPAP at 10 cmH$_2$O until extubation but not thereafter. The surrogate marker measured 1 hour after extubation did not differ between the group treated with a RM and CPAP, and a control group not given RM. Therefore, a high $F_iO_2$ when CPAP ends, was proved ineffective. In study III, a CPAP of 6 cmH$_2$O was used during the period in which $F_{ET}O_2$ was deliberately reduced. Even if there were leaks that diminished the effectiveness, it is unlikely that an $F_{ET}O_2$ of less than 0.5 was not achieved, and static atelectasis caused by this level of $F_{ET}O_2$ would not have appeared in the CT scan after 14-25 minutes.

To summarise, and to answer the question about the possible link found between age and atelectasis, using CPAP during preoxygenation and induction in older patients might expose them to higher $F_{ET}O_2$ during induction of anaesthesia. If the level of PEEP is insufficient during anaesthesia, and airway closure follows, this might later lead to increased areas of atelectasis compared with younger patients.

**Smoking and atelectasis**

Another new finding of the post hoc analysis from study IV was the correlation coefficient of 0.3 ($P = 0.022$) between smoking more than 6 pack-years and atelectasis formation. Most of the smokers (14/18) were in the control group, and as noted earlier, atelectasis and age correlated significantly in the intervention group but not in the control group. Added to this is the fact that, in study III, the intervention failed in the group given 80% oxygen with CPAP/PEEP. Overall, smoking was more common in this group compared with the other two groups in study III. Circumstantial evidence about smoking as a contributor to atelectasis is thus gaining relevance. Clinically, an
association between smoking and atelectasis makes sense. However, this association can be overlooked because of the great variability in the effects of smoking in patients relating to different exposure in pack-years and individual susceptibility, especially when studying the early formation of atelectasis in a small number of patients. Both studies III and IV observed the late formation of atelectasis, and study IV included the largest number of patients in any study using CT to detect atelectasis.

It would seem that when smoking has an effect, this occurs early during anaesthesia, and it is questionable whether a fixed level of CPAP/PEEP will suffice to prevent atelectasis formation. It might have been more relevant to look at CPAP/PEEP as a possible double-edged sword in terms of smoking and its effects on airway closure, $F_{ET}O_2$, and the degree of respiratory disturbance. In 1991, Gunnarson et al$^{61}$ studied 10 patients with COPD and found only minor areas of atelectasis during anaesthesia in three of the patients. The $V_A/Q$ relationship was also studied with the multiple inert gas elimination technique, and the amount of true shunting found (mean 1 %) was consistent with only minor atelectasis. Gas trapping because of airway closure was proposed as a possible mechanism. The pivotal role of the oxygen concentration as a determinant of atelectasis had not established in 1991, and the duration of preoxygenation was not specified in the article.

In 2000, Samain et al$^{62}$ studied 10 patients with COPD (FRC 5.6 ± 1.1 L; mean ± SD) and 10 patients without COPD (FRC 3.1 ± 0.4 L) during preoxygenation. After 3 min of preoxygenation $F_{ET}O_2$ was 0.83 and 0.91 in the group with and without COPD, respectively. Thus, if one does not take into account the effect of increased FRC and increased CC in COPD for achieving a predefined level of $F_{ET}O_2$ during preoxygenation, the gas trapped in the closed-off units in the lungs during anaesthesia will contain more nitrogen. More nitrogen protects against atelectasis formation, and this protection is increased by concomitant HPV. Using CPAP during preoxygenation and induction probably reduces this protection. The smokers included in studies III and IV had not been diagnosed clinically with COPD, but they might be suspected to have been in different stages in the progression to COPD. What these stages might possibly look like is impossible to say, other than that it somehow increased the tendency to develop atelectasis. We suspect that, as in older patients, the level of PEEP during anaesthesia was insufficient in regard to the tendency of airway closure.
Comments on the post hoc analysis of pooled data from studies I-IV

There was a significant effect on the area of atelectasis in the combined intervention group as compared with the combined control group. Even though the definitions of the groups might be discussed, the post hoc analysis showed that the interventions were a step in the right direction. The pooling of data gained the statistical power that was missing in study III. The average area of atelectasis in the control group was similar to that reported in other compilations.

Can lessons be learned from outliers?

As mentioned, atelectasis appears in ~90 % of anaesthetised patients. The frequency and the area of atelectasis will depend on many different factors; the time point after the start of preoxygenation during or after anaesthesia is one key factor. In the four studies of this thesis, the radiologist reported only 3/135, or 2 % of patients as completely free of atelectasis. In one of these patients, only 60 % oxygen had been given during preoxygenation, and this patient was studied early during anaesthesia. Of more interest are the two other patients who exhibited no atelectasis even at 14 min after awakening. One patient who had been given 100 % oxygen during pre- and postoxyge- 
nation was 46 years old and had a BMI of 19.6. The other patient who had been given 100 % oxygen during preoxygenation and 30 % oxygen for post-oxygenation was 41 years old and had a BMI of 33.3. A BMI of 19.6 might be a light burden for the lungs but this cannot be said of a BMI of 33.3. Thus, the results for two positive outliers are contradictory.

However, there were also patients on the negative side of the scale, who developed much more atelectasis than others in their group. If we arbitrarily identify patients with increased area of atelectasis as outliers when the area of atelectasis is more than 19 cm$^2$ (as used in study I; i.e., almost twice the normal amount, irrespective of the time of investigation), six patients were exhibited. All six patients belonged to a control group. The characteristics of these patients are given in Table 4.
Table 4. Characteristics of patients (controls) with an area of atelectasis > 19 cm²

<table>
<thead>
<tr>
<th></th>
<th>Study 1</th>
<th>Study 2</th>
<th>Study 3</th>
<th>Study 4</th>
<th>Study 4</th>
<th>Study 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atelectasis</td>
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<td>21.0</td>
<td>23.1</td>
<td>19.1</td>
<td>21.8</td>
<td>27.7</td>
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<tr>
<td>Time (min)</td>
<td>14 periop</td>
<td>45 periop</td>
<td>18 postop</td>
<td>21 postop</td>
<td>13 postop</td>
<td>11 postop</td>
</tr>
<tr>
<td>Age (years)</td>
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<td>52</td>
<td>63</td>
<td>40</td>
<td>60</td>
<td>58</td>
</tr>
<tr>
<td>BMI kg/m²</td>
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<td>26.8</td>
<td>26.0</td>
<td>33.5</td>
<td>26.1</td>
<td>28.1</td>
</tr>
<tr>
<td>Smoking</td>
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<td>No</td>
<td>No</td>
<td>No</td>
<td>23 P-years</td>
<td>43 P-years</td>
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<td>1</td>
<td>2</td>
<td>2</td>
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<td>2</td>
</tr>
<tr>
<td>Event</td>
<td>Mucous</td>
<td>-</td>
<td>Prolonged intubation</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

BMI = body mass index, calculated as weight/(height)². ASA = physical status according to the American Society of Anesthesiologists. periop = perioperatively during anaesthesia. postop = postoperatively after anaesthesia. P-years = pack years.

In two of the six patients, special events occurred, which probably influenced the area of atelectasis. One patient had severe mucous production and, during surgery, had to be disconnected from the ventilator several times to perform suction of the trachea. In one patient there was a prolonged intubation procedure, necessitating three intubation attempts before intubation was successful. In the remaining four patients, potential modifying factors could be identified in three patients. Two of the patients were heavy smokers and one patient had a BMI of 33.5. The effect of BMI may have been balanced overall by the use of CPAP/PEEP but it is possible that in some patients CPAP/PEEP might fail to prevent atelectasis.

We propose that using an expression of the effect of overweight that concentrates more on the distribution of this extra burden (e.g., to the abdomen) may correlate more strongly with atelectasis than does BMI.

Oxygenation

*SpO₂ as an oxygenation index and correlation to atelectasis*

Using SpO₂ as an oxygenation index did not reveal any differences between groups at any time perioperatively. In all groups, SpO₂ was significantly lower during anaesthesia (with air) than before anaesthesia. Postoperatively, only 3/89 patients needed extra oxygen temporarily to maintain an SpO₂ ≥ 94. The other patients responded promptly to physical stimulation if SpO₂ < 94 %.

SpO₂ has become the standard for monitoring arterial oxygenation perioperatively.⁶³ A recent meta-analysis confirmed that pulse oximetry could detect hypoxemia and related events.⁶³ Desaturations or hypoxemia can be
defined as mild (SpO$_2$ 86-90 %), moderate (SpO$_2$ 81-85 %), or severe (SpO$_2$ < 81 %)\textsuperscript{64}. Assessment of the efficiency of pulmonary oxygenation is often influenced by F$_{1}$O$_2$.

When data for mixed venous blood gases are lacking, an estimate of the venous admixture based on arterial blood gas analysis and an assumption of the arteriovenous O$_2$ content difference can be used\textsuperscript{65}. During anaesthesia, F$_{1}$O$_2$ is usually 0.3-0.4, which is a good way of hiding or treating desaturations that would have been seen with ventilation with air.

In fact, using an F$_{1}$O$_2$ of 0.4 and assuming “normal” values for the variables in the shunt equation and a haemoglobin value of 120 g/L gives an arterial oxygen saturation (SaO$_2$) of 94 % with a true shunt of ~23 %\textsuperscript{66}. With air, a true shunt of 23 % would give an SaO$_2$ of ~90 %, and a true shunt of 13 % would give an SaO$_2$ of ~94 %. During total intravenous anaesthesia, the true shunt has been correlated to the area of atelectasis\textsuperscript{30} and is expressed as shunt (\% CO) = 2.9 + 0.55 \times \text{atelectasis area in cm}^2. Using this formula, a true shunt of 23 % would correspond to an atelectasis area of 36.5 cm$^2$, and a true shunt of 13 % would correspond to an atelectasis area of 18.4 cm$^2$. Based on studies\textsuperscript{67, 68} showing correlations between SaO$_2$ and SpO$_2$, in the following paragraph, SpO$_2$ is used as a surrogate variable for SaO$_2$.

**SpO$_2$ as an oxygenation index perioperatively**

In study III, SpO$_2$ with air was measured before anaesthesia, twice during anaesthesia using air and continuously postoperatively with air. The absolute value of SpO$_2$, 99-100 %, before anaesthesia indicates a systematic error, because, according to the alveolar equation of ideal alveolar PO$_2$, breathing air at normal atmospheric pressure with normal alveolar ventilation cannot produce such high values. SpO$_2$ measured using the same pulse oximeter during anaesthesia revealed values of 95-96 % with no difference between groups. Assuming that the error was unchanged, one may conclude that SpO$_2$ decreased significantly during anaesthesia.

This might look like trivial information, but there is a point to be made. If we assume the systematic error to be +2 %, SpO$_2$ before anaesthesia would be 96-97 % if corrected for this error, and 93-94 % during anaesthesia (with air). Thus, systematic error in the SpO$_2$ reading that produces values that are too high, will underestimate an oxygenation problem. In addition, giving extra oxygen without caution, for example in the recovery ward after surgery, will make it even harder to diagnose the existence of an oxygenation problem.

These shortcomings were addressed some years ago in the guidelines of the British Thoracic Society for emergency oxygen use in adult patients\textsuperscript{69}. For normal patients, the guidelines recommend that the lower limit for giving extra oxygen should be set at an SpO$_2$ of 94 %, and that pulse oximeters
should have an accuracy of ± 1-2 %. The guidelines also acknowledge that a deterioration of 3 % in SpO\textsubscript{2} is a warning sign.

To summarise the point discussed, when patients are monitored with SpO\textsubscript{2} and the readings are judged as correct, an SpO\textsubscript{2} < 94 % might indicate a serious shunt, equivalent to more than 23 % of cardiac output and the existence of large areas of atelectasis of more than 36 cm\textsuperscript{2}.

The discussion so far has assumed that the full decrease in SpO\textsubscript{2} represents only true shunt, which is seldom the case. There are many possible reasons for impaired oxygenation during and after anaesthesia. Perfusion of areas with low V\textsubscript{A}/Q is almost obligatory, adding venous admixture to the equations. As much as 75 % of the impairment of arterial oxygenation during anaesthesia might be explained by the combined effects of atelectasis and airway closure\textsuperscript{30}. Other causes for impaired oxygenation include hemodynamic disturbances or increased metabolic demands. In the recovery ward, hypoventilation is a frequent feature that is effectively disguised by oxygen treatment.

Postoperatively, with a few brief exceptions, all patients in studies III and IV had an SpO\textsubscript{2} ≥ 94 %. The median area of atelectasis was 8.5 cm\textsuperscript{2} in the control group in study III. Using the formula given above, this would correspond to a true shunt of ~8 % and an SpO\textsubscript{2} of 95 % during air breathing. Thus, it is not surprising that in the clinical situation with “normal” amounts of atelectasis, pulse oximetry alone will not alert us to the situation.

One control patient in study III and three control patients in study IV had an area of atelectasis of more than 19 cm\textsuperscript{2}. Theoretically, using the formula as cited, atelectasis in these patients might have been detected by pulse oximetry if SpO\textsubscript{2} had declined below 94 %, but they were not. There could be several explanations for this. A systematic false high SpO\textsubscript{2} value is one possibility. Using the formula linking atelectasis to true shunt as derived from conditions during anaesthesia postoperatively might be inaccurate. Finally, we know very little about how alterations in HPV affect these calculations. It is reasonable to assume that HPV is more effective in the wake patient than during anaesthesia\textsuperscript{70, 71}; therefore, stronger HPV might reduce the effect of atelectasis, especially when breathing air.

To summarise, pulse oximetry is not a suitable tool for revealing normal-size atelectasis that often accompanies anaesthesia. Even greater amounts of de-aerated lung tissue, up to ~20 cm\textsuperscript{2}, may escape detection because of:

- inaccurately high SpO\textsubscript{2} readings up to +2 percentage points;
- inappropriate use of oxygen supplementation, which can mask impaired oxygenation;
- an effective HPV response in the postoperative period;
• confounding factors such as disturbance of \( V_A/Q \) or postoperative hypoventilation.

**Clinical implications**

The standard routine of using 100 % oxygen during preoxygenation is still the optimal procedure. Using 80 % oxygen during preoxygenation shortens the time available in a “cannot ventilate, cannot intubate” situation and the size of the area of atelectasis will eventually be similar to that with 100 % oxygen. Avoiding airway closure with CPAP/PEEP increases the effect of preoxygenation and simultaneously reduces the tendency for atelectasis formation, even with 100 % oxygen.

However, precautionary measures might be considered if CPAP/PEEP is used combined with 100 % oxygen for pre- and postoxygenation, especially in a high-risk older patient. First, immediately before extubation, a RM can be performed, eliminating any atelectasis evolved hitherto. Second, after extubation, ventilation with CPAP, or bi-level positive airway pressure, on-mask should start, and before ending maskventilation, the original high \( F_{I\text{O}_2} \) should be exchanged to a low value using 30-40 % oxygen in nitrogen.

In the postoperative ward, deep-breathing exercises might be implemented. In patients unable to maintain \( \text{SpO}_2 \) by breathing air at \( \geq 94 \% \), a more detailed diagnosis might be needed, including arterial blood gas analysis. In selected cases, postoperative use of non-invasive ventilation might be an appropriate treatment.

The use of CPAP/PEEP probably needs titrating to individual demands when the pulmonary physiology deviates from normal limits. A history of heavy smoking, especially with a high ASA class, should always raise a warning flag.
Conclusions

1. The early formation of atelectasis was halted by the use of 60 or 80 % oxygen compared with 100 % oxygen during preoxygenation and induction of general anaesthesia.

2. The duration of apnoea without desaturation shortened markedly when the oxygen concentration was reduced from 100 % during preoxygenation.

3. During induction of anaesthesia, only small areas of atelectasis appeared around the time of intubation, irrespective of oxygen concentration. In the time window starting after intubation, from 7 until 14 min after start of preoxygenation, the area of atelectasis expanded substantially with 100 but not with 80 or 60 % oxygen used during preoxygenation. These processes were probably influenced by the fact that the duration of apnoea without desaturation was investigated at the same occasion.

4. The benefit of using 80 % oxygen during preoxygenation seemed to be short lived because the area of atelectasis evolved gradually from 14 to 45 min after start of preoxygenation; the area approached the size normally found after 14 min with 100 % oxygen during preoxygenation.

5. A ventilation strategy using 100 % oxygen during pre- and postoxygenation with a combination of CPAP or PEEP and a reduced F<sub>ET</sub>O<sub>2</sub> before ending mask ventilation with CPAP after extubation reduced the area of atelectasis compared with a group with no CPAP/PEEP and a high F<sub>ET</sub>O<sub>2</sub> after ending mask ventilation after extubation. In a post hoc analysis, after correction for modifying effects of age and BMI, this reduction became statistically significant.

6. The areas of atelectasis did not differ between ventilation strategies using 80 or 100 % oxygen during pre- and postoxygenation with a combination of CPAP or PEEP and a reduced F<sub>ET</sub>O<sub>2</sub> before ending mask ventilation with CPAP after extubation.

7. The area of atelectasis after a ventilation strategy using 100 % oxygen during preoxygenation with a combination of CPAP or PEEP and 30 % oxygen during postoxygenation before extubation, did not differ from that observed in a group treated with the same strategy but with 100 % oxygen during postoxygenation.

8. The use of SpO<sub>2</sub> in patients when ventilated with, or breathing, air to detect possible differences in the area of atelectasis between different treatment groups was discouraged in relation to the area of atelectasis that had developed.
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