Mechanical Ventilation
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Mechanical ventilation

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Second Edition

**Acute Respiratory Failure**

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Learning Objectives

After studying this module on Mechanical ventilation, you should:

- List the indications of mechanical ventilation
- Describe the physiological basis of mechanical ventilation
- Describe the basic modes of mechanical ventilation
- Discuss ventilator settings in different modes and basic waveform interpretation
- Outline the basic principles of weaning from mechanical ventilation
- List the complications of mechanical ventilation
- Describe how to apply this knowledge in clinical practice.

eModule Information

Expiry date:

COBATrIKe competencies covered in this module:

Competencies

- Monitors and responds to trends in physiological variables
- Describes the use of devices for circulatory or respiratory assist

Faculty Disclosures:
The authors of this module have not reported any disclosures.

Duration: 9 hours

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1. Introduction

The mechanical ventilator is an essential part of life support in the intensive care unit (ICU). It was the need to use the mechanical ventilators outside the operating theatre during the poliomyelitis epidemic in 1950s that gave birth to the ICUs of today. Clinicians working in ICU have to become familiar quickly, with how to handle use a ventilator, and fulfil the tasks of initiating, maintaining and weaning patients from mechanical ventilation. This eCourse deals with the physiology underlying ventilatory support, and the basic principles of applying mechanical ventilation in clinical practice.

Although mechanical ventilator is an essential, life-saving device, its use is not free of harm. A solid knowledge of the principles of mechanical ventilation and of the potential harms associated with its use are of paramount importance for the successful and safe use of ventilators in the ICU.
2. Delivery of mechanical ventilation

2.1. Indications for mechanical ventilation

Mechanical ventilation is indicated to manage respiratory failure of any cause and to permit pharmaceutic depression of the respiratory centre during anaesthesia. As shown in Figure 1 the causes of respiratory failure can be classified into:

- Failure to maintain a patent airway
- Failure to maintain adequate ventilation
- Failure to maintain adequate oxygenation

![Figure 1: Causes of respiratory failure. Mechanical Ventilation module, ESICM Academy, 2018](image)

**Figure 1:** The causes of respiratory failure can be classified as: failure to maintain a patent airway, due to obstruction, as in case of facial trauma or laryngeal edema, or due to loss of reflexes, as in coma or deep sedation; failure to ventilate, due to central nervous system (CNS) problems affecting the respiratory centre, such as CNS trauma or pharmaceutical depression, due to failure in nerve conduction, such as in cervical spine injury and neuromuscular blockade, due to respiratory muscle failure, such as myopathies, or due to problems of chest wall, such as flail chest or pneumothorax; failure to oxygenate, due to V/Q mismatch, as in ARDS, due to shunt, as in congenital heart disease, due to dead space ventilation, as in exacerbation of obstructive lung diseases, or due to diffusion abnormalities, as in fibrotic lung diseases.

The choice of timing and interface to deliver mechanical ventilation depends on the underlying cause of respiratory failure.

The failure to maintain a patent airway is a common and relatively straight-forward
indication for endotracheal intubation and mechanical ventilation. Common causes include:

- Decreased level of consciousness (a GCS<9 is used for trauma patients)
- Need for deep sedation
- Upper airway obstruction (trauma, hematoma, oedema)
- Inability to manage secretions

In patients with hypercapnic (failure to maintain adequate ventilation) or hypoxemic (failure to maintain adequate oxygenation) respiratory failure, the choice of timing and patient-ventilator interface to deliver mechanical ventilation is more complicated, and depends on the underlying cause of respiratory failure (see eCourse on [Acute Respiratory Failure](#) and [COPD and Asthma](#)).

The presence of hypercapnic acidosis, hypoxaemia despite administration of oxygen (PO₂< 60mmHg, with FiO₂= 50%), or dyspnea with increased work of breathing are broadly accepted indications for mechanical ventilation. Increased work of breathing, a common feature of acute exacerbation of obstructive lung disease, pneumonia, or ARDS, may induce muscle fatigue, and is associated with high energy expenditure of the respiratory muscles and a significant increase in regional blood flow.

Clinical signs of increased work of breathing and respiratory distress include:

- Nasal flaring
- Mouth breathing
- Recruitment of accessory and expiratory muscles
- Tracheal tug
- Intercostal recession
- Paradoxical abdominal movement
- Tachypnea
- Tachycardia
- Hypertension or hypotension
- Diaphoresis

\[\text{Note}\]

initiating mechanical ventilation during shock in experimental setting caused a 7-fold decrease in the percentage of cardiac output received by the respiratory muscles

\[\text{In text References}\]

(Tobin and Alex. 1994; Viires et al. 1983)

\[\text{References}\]
2. 2. Aims of mechanical ventilation

The overall goal of mechanical ventilation is to support gas exchange and sustain life until the cause of respiratory failure is resolved. As Hippocrates stated, an important goal of medical practice is to “do no harm”, and this is an important aspect to consider when setting the goals during mechanical ventilation.

The aims of mechanical ventilation according to the initiating indications include:

- Achieve adequate ventilation - CO₂ elimination
- Improve oxygenation
- Relieve respiratory distress - offload respiratory muscles

Ventilatory support is now recognised as a necessary, often life saving, intervention that can be harmful. Indeed, mechanical ventilation can induce more harm than benefit to the patient by overstretching the lungs while trying to maintain normal blood gases, or by inducing respiratory muscle injury or atrophy from prolonged unloading. These complications of mechanical ventilation, termed ventilator-induced lung injury (VILI) and ventilator-induced diaphragmatic dysfunction, are discussed in part 6 of this eCourse. The choice of targets during mechanical ventilation requires a careful balance between the costs and benefits of achieving the specific target in the specific clinical situation. While targeting normal blood gases in patients with healthy lungs, as often the case is during anesthesia, is relatively safe and easy. In patients with lung injury or airflow limitation the cost of achieving normal blood gases may be higher than the benefit.

2. 2. 1. PCO₂, pH targets:

A normal PCO₂ and pH is commonly targeted in patients with healthy lungs in the peripertative period, as well as in patients with any brain injury, aiming to prevent an increase in intracranial pressure. However, a normal PCO₂ is not mandatory in all patients at any cost. Permissive hypercapnia is a ventilatory management approach targeting higher than normal PaCO₂. This allows the application of a low tidal volume (Vt) (and lower than required minute ventilation MV) to prevent or minimise lung over-distention. Lung over-distention may cause either VILI, in patients with reduced FRC,
or dynamic hyperinflation, in patients with flow limitation (further details are discussed in part 6 of this eCourse, and in eCourse on COPD and Asthma). Permissive hypercapnia is therefore a sophisticated name for a compromise in ventilation targets, when lung pathology does not easily allow achieving normal PaCO₂ values (Figure 2, Figure 3).

![Figure 2: PaCO2 targets: costs and benefits. Mechanical Ventilation module, ESICM Academy, 2018](image)

**Figure 2:** Achieving a normal PaCO₂ may be challenging in the presence of lung pathology. Although it prevents the increase in respiratory drive caused by hypercapnia and acidosis, if it results in high distending pressures during breathing, it may promote VILI or dynamic hyperinflation. On the other hand, limiting tidal volume and minute ventilation in such patients will protect the lungs. This is at the cost of hypercapnia and acidosis, which increase respiratory drive. Here patients often require deep sedation and/or paralysis to tolerate mechanical ventilation.

![Figure 3: Correlation of respiratory drive with PaCO2. Mechanical Ventilation module, ESICM Academy, 2018](image)

**Figure 3:** Changes in minute ventilation in response to changes in alveolar PCO₂, under normal conditions, and in the presence of hypoxia (left) and acidosis (right).

2.2.2. Oxygenation targets

In most patients targeting a SpO₂ above 90-92% is a reasonable target. Remembering the Hb dissociation curve and oxygen delivery equation (Figure 4), it becomes clear that there is no significant benefit for the patient from a supraphysiological PaO₂.
Recent studies demonstrated that targeting hyperoxia had no benefit and might be associated with adverse outcomes.

![Image](image.png)

**Figure 4:** The sigmoidal shape of Hb dissociation curve indicates that increases in Hb saturations above 90% have a small effect on PaO₂. Moreover, increasing PaO₂ above 100 mmHg has minimal effects in oxygen delivery, as the amount of oxygen dissolved in blood is very small.

2. 2. 3. Offloading the respiratory muscles

The benefits of offloading the respiratory muscles are very important. These include the relief of respiratory distress, prevention of muscle fatigue, and avoiding the increase in their blood supply (important in shock states). Moreover, avoiding rigorous breathing may prevent self-induced lung injury. Unfortunately, prolonged respiratory muscle rest may also induce structural alterations and muscle weakness (ventilator-induced diaphragmatic dysfunction, part 6).

In text References

(Tobin and Alex. 1994; Laffey et al. 2004; Helmerhorst et al. 2015; Roussos. 1990; Dres et al. 2017; Yoshida et al. 2013)

References

2.3. Means to deliver positive pressure ventilation

The currently used positive pressure ventilators permit the delivery of oxygen mixture to the lungs by application of positive pressure at the airway opening. To effectively deliver the positive pressure, an interface that guarantees a reasonably effective pneumatic seal is required. This can be accomplished by bypassing the oropharyngeal airway (invasive), or by applying a tight fitting facial mask (non-invasive). The two kinds of interfaces (Figure 5) used are:

- the endotracheal or tracheostomy tubes for invasive mechanical ventilation
- the non-invasive ventilation masks for non-invasive mechanical ventilation (NIMV)

![Figure 5: Interfaces to deliver positive pressure ventilation. Mechanical Ventilation module, ESICM Academy, 2018](image)

**Figure 5:** From left to right: Endotracheal Tube and Non-invasive Ventilation Masks, Total face, Helmet (above), and nose-mouth mask (below).

Each interface and approach (invasive or non-invasive) has specific advantages, disadvantages and limitations which have to be carefully balanced in every patient and aligned with the goals of treatment.

<table>
<thead>
<tr>
<th>Table 1: Advantages, disadvantages and limitations for Endotracheal Tube and Non-invasive Ventilation Masks</th>
</tr>
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<tbody>
<tr>
<td><strong>Endotracheal Tube</strong></td>
</tr>
<tr>
<td>Protection from major aspiration</td>
</tr>
<tr>
<td>Relief of fixed upper airway obstruction*</td>
</tr>
<tr>
<td>Permit suction of secretions</td>
</tr>
<tr>
<td>Stable, leak-proof connection with the ventilator</td>
</tr>
<tr>
<td>Protection from infections of lower respiratory tract by reflexes of the upper airway</td>
</tr>
<tr>
<td>Comfort</td>
</tr>
<tr>
<td>Need for sedation</td>
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<tr>
<td>Need for patient cooperation</td>
</tr>
<tr>
<td>Permit effective cough</td>
</tr>
<tr>
<td>Ability to speak</td>
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<tr>
<td>Ability to speak</td>
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- Variable airway obstruction, as occurs in obstructive sleep apnea is successfully managed with NIMV.

The main benefit of NIMV is avoiding endotracheal intubation, and thus preventing all related complications, including all device-related ICU-acquired infections (ventilator-associated pneumonia, catheter-related bloodstream infections, urinary catheter-related infections). Non-invasive ventilation can achieve all aims of ventilation described above. It can be used in a large number of patients with respiratory failure in absence of contraindications. The contraindications for NIMV are

- Cardiac or respiratory arrest
- Inability to protect airway
- Inability to manage secretions
- Facial or upper airway trauma, surgery, or (potential of) obstruction
- Intractable emesis or GI bleeding
- Shock (relative contraindication, common cause of failure of NIMV)

**Note**

As NIMV does not provide a stable, leak-proof connection to the ventilator it is essential that the patient tolerates periods of disconnection. Respiratory
insufficiency severe enough to require high levels of PEEP and/or FiO² is a relative contraindication for NIMV.

The use of NIMV for acute respiratory failure has been increasing steadily over the past years, as has the success rate the prevention of intubation. Increased experience and improvements in patient-ventilator interfaces account for this success. Another important factor for the success of NIMV is appropriate patient selection.

Conditions in which NIMV has been studied and proven helpful are:

- Acute Exacerbation of COPD
- Cardiogenic Pulmonary edema
- Discontinuation of invasive mechanical ventilation
- Preventing postoperative respiratory failure

In patients with acute exacerbation of COPD and hypercapnic acidosis, the use of NIMV has been shown to decrease the need for intubation and mortality by almost 50%. In patients with hypoxemic respiratory failure, from pneumonia or ARDS, studies have not consistently shown benefit. NIMV failure is high in patients with moderate and severe ARDS with some studies demonstrating an increase in mortality.

Other conditions in which NIMV has been studied, but there is currently limited evidence include:

- Chest trauma
- Asthma
- Neuromuscular diseases

Factors associated with NIMV failure include:

- Severity of disease
- Comorbidities
- Shock
- Impaired level of consciousness
- ARDS
- Severe acidosis
- Failure to improve after 1-2h of NIMV

For further details about the use of NIMV in patients with acute exacerbation of COPD, please see COPD and Asthma.

In text References

(Rochwerg et al. 2017)
3. Basic Principles of Mechanical Ventilation

3. 1. Basic physiology of respiratory system related to mechanical

The main reasons for instituting mechanical ventilation are to support gas exchange and decrease the work of breathing, at the same time, allowing other interventions to reverse the cause of respiratory failure. In this section we will discuss the physiologic mechanisms by which mechanical ventilation supports gas exchange and reduces work of breathing.

3. 1. 1. Equation of motion

The respiratory control system consists of a motor arm, which executes the act of breathing, a control centre located in the brain medulla, and several pathways that convey information to the control centre. Based on the input, the control centre activates spinal motor neurons, and subsequently peripheral nerves, to activate respiratory muscles (inspiratory and expiratory muscles) which generate pressure (Pmus). This pressure drives the inflation of lungs, which is used to overcome the elastic recoil pressure, generated as the lungs are inflated above their passive functional residual capacity (FRC), and the resistance to flow, (inertia is assumed to be negligible) as described in the equation of motion. According to the equation of motion equation 1 Pmus at time t is dissipated to overcome the resistive (Pres) and elastic (Pel) pressure of the respiratory system, where Rrs and Ers are resistance and elastance of respiratory system, respectively, ΔV(t) is instantaneous volume relative to passive functional residual capacity (FRC) and V'(t) is instantaneous flow.
Figure 6: Pressure volume (PV) curves of lung, chest wall, and respiratory system. Note that FRC is defined as the point where the elastic recoil pressure is equal and opposite to the elastic recoil pressure of the chest wall, and the respiratory system PV curve crosses the X axis at zero. Above passive FRC, the elastic recoil pressure of the respiratory system becomes positive, and thus lung inflation requires the application of external pressure. Notice also the relative flattening of the pressure-volume curve of the respiratory system near TLC, indicating that larger increases in pressure are required for smaller increase in volume.

\[
P_{\text{mus}} = P_{\text{res}} + P_{\text{el}} = R_{\text{rs}} \cdot V'_{\text{rs}} + E_{\text{rs}} \cdot \Delta V_{\text{rs}} \quad \text{(Eq. 1)} \\
P_{\text{TOT}} = P_{\text{aw}} + P_{\text{mus}} = R_{\text{rs}} \cdot V' + E_{\text{rs}} \cdot \Delta V \quad \text{(Eq. 2)}
\]

Figure 7: Equation of motion. Mechanical Ventilation module, ESICM Academy, 2018

Figure 7: Pmus at time t is dissipated to overcome the resistive (Pres) and elastic (Pel) pressure of the respiratory system, where Rs and Er are resistance and elastance of respiratory system, respectively, \( \Delta V(t) \) is instantaneous volume relative to passive functional residual capacity (FRC) and \( V'(t) \) is instantaneous flow (Equation 1). During mechanical ventilation, the pressure provided by the ventilator (Paw or Pvent) is incorporated into the system, and the total pressure applied to respiratory system (PTOT) is the sum of Pmus and Paw (Equation 2).

During mechanical ventilation, the pressure provided by the ventilator (Paw or Pvent) is incorporated into the system. The ventilator and the respiratory muscles can be viewed as pressure generators arranged in series. Therefore, in mechanically ventilated patients the total pressure applied to the respiratory system (PTOT) is the sum of Pmus and Paw (Equation 2, Figure 7)

- **Note**

  The resistance and elastance of the respiratory system, particularly in the presence of lung or chest wall disease, are not constant during the breath but exhibit a considerably low flow and volume dependency.

The combination of PTOT, Rs, Er, and respiratory rate, determines minute ventilation. The partial pressures of arterial blood gases will be based on minute ventilation and the gas exchange properties of the lungs.

3. 1. 2. PCO elimination

The levels of arterial CO2 depend on the balance between CO2 production (V′CO2) and elimination through alveolar ventilation (V′A), as described in Figure 8, Equation 3. Total, or minute ventilation (VE) is the amount of air moved in or out of the lungs per minute. It is the product of tidal volume (VT) and respiratory rate (RR), Figure 8
Equation 4. Alveolar ventilation (V'A) represents the fraction of minute ventilation involved in gas exchange, thus excluding ventilation of airways and non-perfused alveoli, the ‘dead space’ ventilation (VD), as described in Figure 8 Equation 5-6. CO₂ removal, therefore, depends on respiratory rate and the ratio of tidal volume to dead space. These are the factors that can be manipulated during mechanical ventilation.

\[
\text{PaCO}_2 = 0.86 \times V'\text{CO}_2 / V'A \quad \text{Eq. 3}
\]

\[
V'\text{E} = VT \times RR \quad \text{Eq. 4}
\]

\[
V'A = V'E - V'D \quad \text{Eq. 5}
\]

\[
V'A = VT \times RR \times (1 - \text{VD/VT}) \quad \text{Eq. 6}
\]

Figure 8: Equations determining PaCO₂.
Mechanical Ventilation module, ESICM Academy, 2018

- Equation 3 PaCO₂ is determined by the ratio of CO₂ production (V'CO₂) and elimination through alveolar ventilation (V'A), where 0.86 is a constant used for unit harmonisation.
- Equation 4 Total, or minute ventilation (VE) is the amount of air moved in or out of the lungs per minute, thus the product of tidal volume (VT) times respiratory rate (RR),
- Equation 5 Alveolar ventilation (V'A) represents the fraction of minute ventilation involved in gas exchange, thus excluding ventilation of airways and non-perfused alveoli, the ‘dead space’ ventilation (V'D)
- Equation 6 Modifying eq.5, to express VD as a ratio over VT, we see that alveolar ventilation depends on VT, respiratory rate, and the ratio of VT to VD.

**Note**

Based on Equation 6, it is clear that changes in VT will have a greater impact on alveolar ventilation than changes in respiratory rate. Practice solving the equation using actual numbers: RR\text{Initial} = 10, VT\text{Initial} = 500ml, Vd = 150ml. Re-calculate V'A after a 50% increase in either RR or VT.

**Note**

The Heat and Moisture Exchangers (HME) used in clinical practice have a volume of 30-50 ml, which is added to the dead space, and thus changing from a HME to a humidifier (no additional dead space volume) will increase V'A without changing minute ventilation.

3. 1. 3. Oxygenation
The mechanisms of hypoxemia are:

- Ventilation/perfusion (V/Q) mismatch
- Shunt
- Diffusion limitation
- Alveolar hypoventilation
- Low FiO2

In critically ill mechanically ventilated patients, only the first two mechanisms are clinically relevant. Significant limitations in diffusion are only observed in patients with advanced fibrosis, alveolar hypoventilation is prevented, and obviously low FiO2 is never present. Moreover, a sixth mechanism is decreased mixed venous oxygen content, present in patients with shock.

In mechanically ventilated patients hypoxemia can be corrected by increasing FiO₂ and application of positive end expiratory pressure (PEEP). The increase in FiO₂ increases alveolar PAO₂ according to the equation:

\[
PAO₂ = (P_{atm} - PH_{2O}) - \frac{PCO₂}{R}
\]

Because the need for mechanical ventilation arises when oxygen administration alone is not enough to correct hypoxemia, and because hypoxemia in critically ill patients is caused by V/Q mismatch and shunt, enriching the inspired gas in oxygen is usually not enough to correct hypoxemia.

3. 1. 4. Positive End Expiratory Pressure

PEEP improves oxygenation by permitting the reopening of alveoli closed due to atelectasis or oedema. The application of PEEP can cause redistribution of extravascular lung water from the alveoli to the peribronchial and perivascular spaces and thus open the collapsed alveoli.

Effects of PEEP:

- Increase in FRC
  - Improved oxygenation
  - Shift of tidal breathing upward in the pressure-volume curve (more compliant)
- Decrease in intrapulmonary shunt
  - Improved oxygenation
- Reduction of alveolar opening and closing
  - Prevention of lung injury
- Increase in intrathoracic pressure
- Decrease in venous return
- Decrease in left ventricular transmural pressure
- Distention of normally aerated alveoli
  - Increase of dead space
  - Causing lung injury
- In the presence of airflow obstruction, there is a substitution of the pressure that must be generated by the inspiratory muscles to overcome intrinsic PEEP and generate inspiratory flow.

The hemodynamic effects of PEEP result from intrathoracic pressure increase (Figure 9). If the cardiac output is dependent on preload, then PEEP may decrease cardiac output by decreasing venous return. In cases of heart failure and volume overload, when cardiac output is less dependent on preload, PEEP can improve left ventricular function by decreasing the transmural pressure of the left ventricle. This effect is direct, due to the increase in intrathoracic pressure and indirect, by improving FRC, thus reducing work of breathing, and diminishing large inspiratory efforts which cause an increase in left ventricular afterload.

![Figure 9: The hemodynamic effects of PEEP.](image)

Mechanical Ventilation module, ESICM Academy, 2018

Figure 9: PEEP increases intrathoracic pressure, thus decreases venous return, and right and left ventricular preload. Preload dependant cardiac output will be decreased by the increased intrathoracic pressure. The increase in intrathoracic pressure will also decrease left ventricular afterload and the transmural pressure of the left ventricle, which may improve left ventricular (dys)function. Therefore, in cases of heart failure and volume overload, when cardiac output is less dependent on preload, PEEP can increase cardiac output. Indirectly, by improving FRC, which may improve gas exchange and reduce work of breathing, this diminishes large inspiratory efforts and prevents the related increases in left ventricular afterload.

Overall the beneficial effects of PEEP include:

- Improved oxygenation by increasing FRC, improving V/Q match, and decreasing shunt
• Reduced work of breathing by shifting tidal breathing towards the more compliant part of the pressure-volume curve, decreasing respiratory drive by improving oxygenation, and, in the presence of intrinsic PEEP, by substituting the pressure that must be generated by the inspiratory muscles to generate inspiratory flow
• Prevent ventilator-induced lung injury (VILI) by preventing cyclic opening and closing of alveoli
• Improve left ventricular function by decreasing its transmural pressure

The detrimental effects of PEEP include:

• Overdistention of alveoli, increasing dead space thus decreasing the efficiency of CO₂ removal, and promoting VILI and barotrauma
• Decrease in cardiac output

It is not possible to pre-defined a specific range of PEEP for a specific patient at a specific time to ensure the beneficial effects and minimise the development of the detrimental effects of PEEP.

In text References

(Kondili, Prinianakis and Georgopoulos. 2003; Feihl and Broccard. 2009; Hess 2015; Tobin and Alex. 1994; West, Luks and 2011)

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• Feihl F, Broccard AF., Interactions between respiration and systemic hemodynamics. Part II: practical implications in critical care., 2009, PMID:18825366
• Hess DR, Recruitment Maneuvers and PEEP Titration., 2015, PMID:26493593

3. 2. Basic functions of positive pressure ventilators

The mechanical ventilators currently used in Intensive Care are equipped with:
- pressurised gas source and an air/oxygen blender
- inspiratory and expiratory valves and external ventilator circuit connecting to the patient
- controllers regulating the opening and closing of inspiratory and expiratory valves, and pressurised air delivery, as well as monitoring and alarm functions

The proper function of a positive pressure ventilator requires the operator to provide the ventilator control system with information on how the breath will be initiated and terminated, and what amount of air will be delivered during inspiration. These three variables are:

- the triggering variable, which is the signal initiating the delivery of positive pressure
- the control variable, which is the algorithm that determines the delivered pressure during the mechanical inspiration
- the cycling off variable, which is the signal of terminating the pressure delivery

**Triggering variable:** the signal initiating the delivery of positive pressure leading to the initiation of mechanical inspiratory phase.

The most commonly used triggering variables are time, pressure and flow. Other triggering variables include flow waveform, and diaphragmatic electrical activity, used in non-invasive ventilation and neurally-adjusted ventilation, respectively.

According to the set triggering mode, the delivery of positive pressure will initiate when the specified condition, as described below, is met.

**Time:** when a respiratory rate (RR) is set, a timed inspiration will begin at the specified interval, for example if RR is set at 20 br/min, positive pressure will be delivered every 3 sec.

**Pressure:** pressure triggering requires the patient to decrease the pressure in the ventilator circuit to a pre-set level (Figure 10).

![Figure 10: Pressure and Flow triggering method](image)
Figure 10: Triggering with pressure (left) and flow (V’, right); notice the drop of pressure (pink arrows) in pressure triggering, and the absence of flow at the same time, as opposed to the change in flow without the pressure drop in flow-triggering (pink line, and purple arrows, right) (Georgopoulos, Prinianakis and Kondili. 2006).

Flow: similar to pressure triggering, the patient is required to decrease flow in the circuit to a pre-set level (Figure 10).

Note

when pressure triggering is used, the respiratory muscles are being subjected to isometric contractions during triggering, as no airflow is yet allowed in the ventilator circuit.

Variable that controls the pressure delivery
The most commonly used variables determining the delivery of positive pressure are:

- Volume: where the ventilator, once triggered, delivers a pre-set (tidal) volume with a pre-set flow-time profile (volume-targeted ventilation)
- Pressure: where the ventilator delivers a pre-set pressure (pressure targeted ventilation)

Other variables used to determine the delivery of positive pressure are:

- Instantaneous flow and volume (reflecting Pmus): in Proportional Assist Ventilation (PAV) where the ventilator delivers pressure which is proportional (the proportionality is pre-set) to instantaneous flow and volume (see Proportional Assist Ventilation)
- Electrical activity of the diaphragm: in Neurally-Adjusted Ventilation (NAVA) the ventilator delivers pressure which is proportional (the proportionality is pre-set) to diaphragmatic electrical activity, monitored by an appropriate esophageal catheter (see Neurally-Adjusted Ventilation Assist).

Note

modern ventilators have the ability to combine different modes of triggering and pressure delivery control, presented in Modes of ventilation with combinations of triggering and pressurisation variables.

Cycling off variable: the signal of terminating pressure delivery.
The most commonly used cycling off criteria, determining the end of pressure delivery and thus of the mechanical inspiratory phase, are time, pressure, and flow.

- Time: the set inspiratory time, determined directly or indirectly by setting respiratory rate and inspiratory to expiratory ratio, initiates the opening of the exhalation valve.
• Pressure: in this case the ventilator terminates the pressure delivery when, due to expiratory muscle activity, Paw increases above a certain threshold (usually 1.5-3 cmH₂O).
• Flow: in this case the exhalation valve opens when inspiratory flow decreases to a pre-set flow criterion which may be either a percentage of peak inspiratory flow or a fixed value.

As a safety feature the ventilator may terminate the pressure delivery when the duration of inspiratory flow is considerably long (i.e. above 3 sec).
In PAV mode, pressure delivery terminates when inspiratory flow decreases to zero, and in NAVA when diaphragmatic activity drops below a preset level.

**Note**
the ‘mechanical’ inspiratory time is the one chosen by the user by setting the triggering and cycling off variables. This differs from the patient’s neural inspiratory time. The aim of ventilator settings is to minimise this difference.

The mode of ventilation is characterised as either ‘control’ (operator-controlled) or ‘assisted’ (patient-controlled) based on the way the breath is initiated and terminated. In other words, if it is controlled by the operator (triggering and cycling off being time) or the patient (all other signals). Based on the variable that regulates pressure delivery a mode is characterised as ‘volume’ or ‘pressure’ targeted.

**Note**
The proper function of a ventilator requires the use of the appropriate external circuit with an inspiratory and expiratory arm, and filters. Additionally air humidification should be performed with either a ‘heat and moisture exchanger’ (HME) mounted at the Y-piece or a heated humidifier mounted within the inspiratory tubing. Routine change of ventilator circuit as a mean to prevent ventilator-associated pneumonia is not considered necessary.

**Note**
ventilators, as any other ICU equipment, require maintenance for proper function. In addition to professional maintenance, every user should be able to perform the basic calibration procedures and tests each ventilator in use.

In text References

(Branson, Hess and Chatburn 2000)

**References**

- Georgopoulos D, Prinianakis G, Kondili E., Bedside waveforms interpretation as a tool to identify patient-ventilator asynchronies, 2006, PMID:16283171
4. Modes of Positive Pressure Ventilation

As discussed above, the parameters which define the mode of positive pressure ventilation are the triggering (and cycling off) variable and the control variable (Figure 11). ‘Controlled’ refers to when the modes in which the triggering variable is time. When a patient’s effort triggers the delivery of positive pressure, the mode is referred to as ‘assisted’. The delivery of positive pressure can be subsequently regulated by a pre-set volume or a pre-set pressure. Modern ventilators use several combinations of trigger and control variables in the ‘dual modes’, as presented in Figure 12, but the basic principles of ventilation remain the same.

![Table](image)

**Figure 11**: Basic modes of ventilation, set and derived parameters. Mechanical Ventilation module, ESICM Academy, 2018

**Figure 11**: The variables set by the operator and derived from the patient’s respiratory system mechanics and muscle function are presented for the most commonly used modes of ventilation.

- **Blue**: variables set by the operator
- **Orange**: variables derived based on the mechanical properties of the respiratory system and respiratory muscle function
- **Pink**: parameters controlled by the patient (but influenced through control-of-breathing feedback mechanisms from the set parameters).

![Table](image)

**Figure 12**: Dual modes of ventilation, set and derived parameters. Mechanical Ventilation module, ESICM Academy, 2018

**Figure 12**: List of common dual modes and relevant ventilator variables. In dual modes the operator defines the variables for both controlled and spontaneous (assisted) breaths. The parameters set are set are dependant on the specific mode/ventilator brand.

**Abbreviations:**
VC: volume control
PC: pressure control
APRV: airway pressure release ventilation
SIMV: spontaneous intermittent mandatory ventilation
MMV: mandatory minute ventilation

- In Bi-Level mode, by defining the time (duration) of P-high and P-low a mechanical respiratory rate is set.

Note

there are more than 50 different modes in the commercially available ventilators
and a complete presentation is beyond the scope of this module.

4. 1. Basic controlled modes volume and pressure control

In both controlled modes, the respiratory rate, inspiratory to expiratory ratio, presence
or absence of inspiratory plateau time, as well as tidal and minute volume are selected
by the operator. This is done directly in volume control mode and indirectly in pressure
control mode.

In volume control mode (Figure 13), the set, or independent variable is volume (tidal
volume) and the dependent variable is pressure.

Based on the equation of motion:

\[ \text{Paw} = V' \times R + VT \times E \]

Figure 13: Waveforms in Volume Control
Ventilation. Mechanical Ventilation module, ESICM
Academy, 2018

Figure 13: Pressure, Flow, and Volume waveforms in volume control ventilation with
constant flow. Notice the gradual increase in pressure during inflation (pink arrow), as
a result of the gradual increase in elastic recoil pressure caused by the increase in
lun volume above FRC. Notice the decrease to the set level of PEEP at end-inspiration (yellow arrow). Notice the shape of the passive expiratory flow-time curve - this is modified in obstructive lung diseases in the same way we see in pulmonary function tests, with an abrupt decrease of expiratory flow.

As the tidal volume is set, the pressure that will develop in the respiratory system at end-inspiration is dependent on the mechanical properties of the respiratory system. These are resistance and elastance (assuming that the patient does not show any spontaneous breathing activity). Moreover, by setting a tidal volume and an inspiratory time, the inspiratory flow becomes also defined (set volume over set time). Most ventilators offer a choice between constant and decelerating flow shape. A constant flow allows measurement of the mechanical properties of the respiratory system. A simple visual analysis of the pressure waveform provides useful information on the mechanical properties of the respiratory system (Figure 14, Figure 15).

![Figure 14: Waveform interpretation in Volume Control. Mechanical Ventilation module, ESICM Academy, 2018](image)

Figure 14: In the absence of respiratory muscle activity, the instant pressure, Paw(i), at any time depends on the volume that has entered the system VTi, the set flow (V’), and the mechanical properties of the respiratory system:

\[ Pawi = V' \times Ri + VTi \times Ei \]

The initial increase in Paw (purple arrow) is caused by the resistance of the circuit and the large airways, as only little volume has entered the system, thus the component VTi\*E is small. Subsequently, both the resistance to flow and the gradually increasing elastic recoil pressure contribute to the increase in Paw (blue arrow). The maximum pressure reached, termed Peak inspiratory pressure (PIP), is when all volume is delivered, just before the inspiratory flow terminates. When inspiratory flow becomes zero, an abrupt decrease in Paw is observed, (green arrow). If no inspiratory plateau time is chosen and expiratory valve opens, pressure drops to the set level of PEEP. If there is a set inspiratory plateau time (inspiratory flow stops, but exhalation valve does not open), we see the plateau pressure (Pplat) determined by the elastic recoil pressure:
\[ Paw = (0 \times Ri) + VT \times E \]

A closer look at the green arrow point may reveal a gradual decrease from the point of zero flow to Ppl caused by redistribution of air in inhomogeneous lungs. As the drop of Paw from Ppeak to Ppl is caused by the elimination of flow in the system it becomes apparent that this is the amount of pressure required to overcome resistance to flow, (red arrow) thus

\[ Ppeak - Ppl = V' \times R \]

After measuring Ppeak and Ppl, and knowing the set, constant flow, one may calculate the resistance of the respiratory system. At the end of inspiratory time, the opening of expiratory valve results in a new, abrupt decrease in Paw, to the set level of PEEP. Following the same reasoning as above, the drop of Ppl to PEEP is caused by the elimination of volume in the system. Therefore, this is the amount of pressure required to overcome the elastic recoil caused by the tidal volume inflation (orange arrow). That is:

\[ Ppl - PEEP = VT \times E \]

and therefore respiratory system elastance, or compliance (C=1/E) can be measured. If there are increases in the initial pressure rise (purple arrow) we have to check for conditions causing increased circuit and large airways resistance. If we see increases in peak pressure and the difference to plateau pressure (red arrow), we have to check for conditions causing increased resistance, and if we see increased Pplat (orange arrow) we have to check for conditions resulting in decreased compliance.

**Figure 15:** Measuring respiratory system resistance and compliance in volume control mode with constant flow. Mechanical Ventilation module, ESICM Academy, 2018

Figure 15: Note that valid measurements require absence of patient's effort. With airway occlusion at the end of inspiration, flow drops to zero, and the proximal airway pressure immediately decreases from a peak value (Ppeak) to a lower initial value (the pressure at zero flow (Pinit)), followed by a gradual decrease until a plateau (Pplat) is achieved after approx. 3 sec. With airway occlusion at the end of expiration, airway pressure may remains unchanged at the level of set PEEP, or increases, indicating the
presence of intrinsic PEEP (PEEPi). The presence of PEEPi should be accounted for when measuring respiratory system compliance.

Compliance of the respiratory system at end-inspiration is defined as the ratio of tidal volume and the difference between plateau pressure (Pplat) and total PEEP (set PEEP or PEEPi)

\[ C = VT / P_{\text{plat}} - P_{\text{EEP tot}} \]

Resistance of large airways determines the difference between peak inspiratory pressure (Peak) and Pinit. The difference between Pinit and Pplat is determined by time constant heterogeneity within the lungs and the viscoelastic behavior of the stress relaxation of the pulmonary tissue.

\[ R_{\text{tot}} = P_{\text{peak}} - P_{\text{Plat}} / V' \]

- \( R_{\text{tot}} \): total respiratory resistance
- \( V' \): preset inspiratory

\[ C = R_{\text{min}} = P_{\text{peak}} - P_{\text{init}} / V' \]

Rmin reflects the resistance from conducting airways and endotracheal tube

In pressure control mode, the independent variable is pressure and the dependent variables are flow and volume, as defined by the equation of motion:

\[ P_{\text{aw}} = V'(i) \times R_i + VT_i \times E_i \]

Thus in pressure control ventilation the instant flow depends on the mechanical properties of the respiratory system. Similarly when the pressure is set, the volume that will be delivered is dependent on the mechanical properties of the respiratory system and the set inspiratory time. (Figure 16)
Figure 16: Pressure, Flow, and Volume waveforms in pressure control ventilation. Notice the differences with VC, the (almost) rectangular shape of the pressure waveform, which in this case is the set value. Also notice the different shape of the flow waveform (pink arrow), which in this case is the dependent variable, gradually decreasing, as the elastic recoil pressure progressively increases with lung inflation (remember the equation of motion $Paw = V'(i) \times R_l + V_Ti \times Ei$). Finally notice that the shape of the passive expiratory flow-time curve is the same irrespective of the mode of inspiration.

![Figure 17: Waveforms in Pressure Control Ventilation – B. Dean R Hess, Respiratory Mechanics in Mechanically Ventilated Patients, RESPIRATORY CARE. NOVEMBER 2014 VOL 59 NO 11](image)

Figure 17: B. In pressure control mode, the independent variable is pressure, and the dependent variables are flow and volume. Therefore, because the instant flow depends on the mechanical properties of the respiratory system, the flow waveform may provide information on these properties. Increasing resistance will depress and flatten the inspiratory flow, while decreasing compliance will decrease peak flow and shorten the time of inspiratory flow (the lower the compliance, or higher elastance, the faster the component $V*T$ will reach the set value of $Paw$, thus make $V'i$ become zero).

![Figure 18: Waveforms in Pressure Control Ventilation - C. Mechanical Ventilation module, ESICM Academy, 2018](image)

Figure 18: C. Measuring respiratory system compliance in pressure control mode can be done similarly to volume control mode, provided that flow at end-inspiration is zero:
\[ C = V T / P_{\text{plat}} - PEEP_{\text{tot}} \]

4. 2. Setting the ventilator in control modes

When choosing ventilator settings, the physician is required to manage two problems, oxygenation and ventilation for CO2 elimination. Management of oxygenation relies mostly on PEEP and FiO₂, while CO₂ elimination depends on tidal volume and respiratory rate, although significant interactions should be considered. Apart from achieving arterial blood gas targets, ventilator settings should focus on preventing or minimising ventilator-induced lung injury. The final choice of ventilator settings is usually a compromise between achieving blood gases targets and protecting the lungs from VILI.

Ventilator parameters to set in Volume/Pressure Control Mode

- PEEP
- FiO₂
- Tidal Volume (Volume Control)
- Inspiratory pressure (Pressure Control)
- Respiratory Rate
- I:E ratio
- Inspiratory plateau (Volume control)

4. 2. 1. PEEP

The application of PEEP aims to open and maintain patent alveoli which tend to close during tidal breathing, thus improving V/Q matching and oxygenation while preventing alveolar overdistention. Before applying PEEP we must first assess whether the lungs are recruitable and then select the appropriate PEEP. The methods to assess lung recruitability are:

- Oxygenation improvement with PEEP. This signifies lung recruitment and shunt reduction.
- Imaging techniques (CT, helium dilution or nitrogen washout techniques, electrical impedance tomography).
- Construction of airway pressure-volume curves during tidal ventilation at different PEEP levels.
4.2.2. How to adjust PEEP at the bedside

In the absence of severe obstructive lung disease with airflow limitation and presence of intrinsic PEEP, a minimum initial PEEP of 5 cmH₂O is used in most patients. Although it is generally accepted that lung injury leading to hypoxemic respiratory failure is best managed by PEEP higher than 5 and that a personalised approach is preferred, there is no consensus as to what physiologic variable should be used as an end-point for PEEP titration. The level of PEEP to be expected in patients with mild ARDS is in the range of 5-10 cmH₂O, 10-15 cmH₂O in moderate ARDS and above 15 cmH₂O in severe ARDS.

The main methods used to titrate PEEP are:

- PEEP-FiO₂ table of the ARDS network
- PEEP above the lower inflection point of the pressure-volume curve
- Respiratory system compliance optimisation
- Driving pressure below 14 cmH₂O
- Stress index <1
- End-expiratory transpulmonary pressure PL > 0 cmH₂O
- Imaging techniques: Computerised tomographic scanning at different levels of PEEP, electrical impedance tomography, end-expiratory lung volume (EELV) increase from a lower to a higher PEEP level.

Titrating PEEP according to oxygenation is widely used and PEEP is selected based on the PEEP-FiO₂ tables proposed by the ARDS Network. This method is easily implemented in clinical practice but it lacks any personalisation and cannot be recommended as a way of PEEP titration, but rather as a starting point. Keep in mind that PaO₂ is also influenced by factors unrelated to lung recruitment, such as cardiac output.

The lower inflection point is considered to indicate the pressure at which a large number of alveoli are recruited. Selecting a PEEP level above the lower inflection point of the static respiratory system pressure-volume curve is rarely employed in clinical practice due to a number of limitations: the patient must be heavily sedated or paralysed, there are occasions where the lower inflection point cannot be identified and lung recruitment continues above the lower inflection point.

The value of PEEP that is associated with the best respiratory system compliance (Crs) can be easily applied, it is non-invasive and allows the assessment of lung recruitment. A limitation is that cyclic opening and collapse of lung units with tidal insufflations may cause an increase in measured tidal compliance not related to the PEEP-induced end-expiratory recruitment.

The concept of driving pressure measurement to adjust PEEP shares the same concept with Crs optimisation. If PEEP is raised and driving pressure decreases, at the same VT, the Crs has increased, suggesting that the higher PEEP caused lung recruitment. In contrast, if PEEP is raised and driving pressure increases, Crs has
decreased, suggesting that higher PEEP caused overdistention of aerated lung. Thus, adjusting PEEP to minimise driving pressure may permit a personalised approach to minimise VILI but shares the same limitation with Crs optimisation.

The slope of the airway pressure-time curve (stress index) gives information about the changes in Crs during inspiration: if the slope increases during inspiration (stress index > 1), Crs is decreasing, while a decreasing slope (stress index<1) indicates increasing Crs. PEEP can be adjusted to a level at which the stress index equals 1. However, this approach requires specialised monitoring equipment to record and analyse the pressure-time relationship, limiting its adoption in clinical practice at present.

The transpulmonary pressure to guide PEEP targets a positive end-expiratory transpulmonary pressure (end expiratory PL) of 0-10cmH₂O. This approach led to higher PaO₂/FiO₂, better Crs and a trend towards improved 28-day survival than the PEEP-FiO₂ table approach. It requires the insertion of an esophageal catheter to estimate pleural pressure.

Imaging techniques to guide PEEP include CT imaging of the lung, lung ultrasound and electrical impedance tomography. Further studies are required to incorporate these techniques as a method to adjust PEEP in clinical practice.

**Summary:** No single method of PEEP titration has been shown to improve clinical outcomes compared to others. Future trials should focus on identifying patients who respond to higher PEEP with recruitment and on clinically important outcomes such as mortality. PEEP may be initially set using the PEEP-FiO₂ table and then individualised to optimise compliance and minimise driving pressure. In selected patients, such as those with suspected high chest wall elastance or severe ARDS, transpulmonary pressure can be used to ensure that applied PEEP is adequate (stop end-expiratory collapse, as suggested by an end-expiratory PL above 0 cmH₂O) and not harmful (no overdistension, end-inspiratory PL < 20cmH₂O).

*Note*

It is important to remember that lung recruitability is a critical factor determining whether the effect of PEEP is predominantly recruitment or overdistention.

*Note*

In patients with obstructive lung disease and airflow limitation, intrinsic PEEP is often present (Figure 19), and the aim of the ventilatory strategy is to eliminate it. As the effects of PEEP are the same whether intrinsic or external, it is important in such patients to monitor for the presence of PEEPi and add external PEEP cautiously, monitoring the resulting total PEEP, titrating to the desired level *(read more in Module on COPD)*.
Figure 19: In the presence of airflow limitation lung emptying may be incomplete at the end of the mechanical expiratory time. The volume remaining trapped increases pressure and can be revealed with an end-expiratory hold manoeuvre.

4.2.3. FiO

Once the appropriate level of PEEP is chosen the minimum FiO₂ is set to achieve the set target of SpO₂. The detrimental effects of hypoxemia are well established, but recent studies indicate that hyperoxia may also induce patient harm through absorption atelectasis and oxygen toxicity. While more studies are needed, it is reasonable to avoid supra-normal PaO₂ when choosing the FiO₂.

4.2.4. Tidal Volume

The tidal volume is one of the main components determining PCO₂ removal, as shown in:

\[ \text{PaCO}_2 = 0.86 \times \frac{\text{V'}CO_2}{VA} \]

and

\[ VA = VT \times RR \times (1 - \frac{VD}{VT}) \]

The choice of tidal volume will be based on the desired levels of PCO₂. In healthy subjects normal tidal volume has been measured in the range of 6-10 ml/kg ideal body weight (IBW). When choosing tidal volume in patients ventilated in controlled modes, it should be remembered that tidal volume is the main determinant of tidal inflation. This, should not be excessive in order to prevent lung injury, especially if the underlying lung pathology has reduced the number of lung units participating in gas exchange. In obstructive lung diseases, tidal volume is limited by the volume that can be exhaled in every breath, so that dynamic hyperinflation is not exacerbated. Thus, in patients with flow limitation, it becomes often necessary to limit tidal volume, even below 6ml/kg, and accept a higher than normal PCO₂, to minimise hyperinflation and PEEPi (see Figure 19, and module on COPD). In patients with healthy lungs, and even more so
in patients with pulmonary edema or ARDS, the main aim when choosing the tidal volume is to minimise end-inspiratory distending pressures which cause VILI. Several ways have been reported to evaluate end-inspiratory inflation and corresponding safety limits. Indices of high distending pressures at end-inspiration and thresholds associated with increased risk of VILI and mortality are:

1. Inspiratory plateau pressure of 30cmH₂O or above: inspiratory plateau pressure, measured after an end-inspiratory hold, is an indicator of end-inspiratory inflation. A threshold of 30cmH₂O has been shown in several studies to be associated with VILI and increased mortality.

2. Transpulmonary end-inspiratory pressure of  >24cmH₂O: transpulmonary pressure can be measured using an esophageal catheter and equals Ppl - PEEP.

3. Driving pressure >14cmH₂O: driving pressure (DP) is the difference between Plat and PEEP (DP=Plat-PEEP). It equals the ratio of delivered VT to respiratory system compliance (Crs), DP = VT/Crs, and thus could better represent tidal inflation in relation to the aerated lung volume. A recent meta-analysis found that DP was the strongest predictor of outcome among the various ventilator variables and proposed a safe threshold of ≤14cmH₂O.

Other indices of high distending pressures at end-inspiration include the stress index calculated from the slope of inspiratory pressure waveform (Figure 20), and the inspiratory mechanical power (Figure 21). This remains to be validated in large clinical trials. Similar to the choice of PEEP, the choice of tidal volume is a compromise between achieving blood gas targets and protecting the lung from VILI.

![Image](Figure 20: The stress index, Respiratory Mechanics in Mechanically Ventilated Patients, RESPIRATORY CARE • NOVEMBER 2014 VOL 59 NO 11)

Figure 20: The stress index mathematically describes the slope of the pressure-time curve during volume control (VC) ventilation with constant flow. If, during tidal inflation, compliance remains constant (linear part of P-V curve, B, ) an upward concavity will be observed, which corresponds to a stress index >1. If compliance progressively increases, suggesting recruitment during tidal inflation, a downward concavity will
appear, corresponding to a stress index <1. When an upward concavity is observed in the
pressure waveform of a patient ventilated in VC, hyperinflation should be
suspected, and tidal volume and/or PEEP adjusted to limit it.

Figure 21: Ventilator-related causes of lung injury.
Intensive Care Medicine, October 2016, Volume
42, Issue 10, pp 1567-1575

Figure 21: This graph is composed of a large triangle (green plus blue), to which a
parallelogram (Resistive, yellow) is added on the right. The left side of the big triangle
represents the total volume (i.e. TV + PEEP volume), while the upper side represents
the plateau pressure. The slope of the hypotenuse represents the compliance of the
system. The area of this large triangle is the total elastic energy present at plateau
pressure. This total elastic energy has two components: the smaller triangle (Elastic
Static, green), which represents the energy delivered just once when PEEP is applied,
and the larger rectangle trapezoid (Elastic Dynamic, blue), whose area represent the
elastic energy delivered at each tidal breath. Note that the rectangle trapezoid results
from the sum of two components (both blue): a rectangle, whose area is TV × PEEP
(third component of the power equation) and a triangle, whose area is TV × ΔPaw ×
1/2, equal to ELrs × TV × 1/2 (first component of the power equation). The third
component of the power equation is the area described by the Resistive parallelogram
(yellow), whose area is equal to (Ppeak − Pplat) × TV. From Gattinoni et al Intensive
care medicine 2016;10:1567-1575

4.2.5. Respiratory rate, Inspiratory/expiratory ratio, inspiratory plateau

The choice of respiratory rate (RR) complements the choice of VT with the aim of
reaching a minute ventilation which will achieve a target PCO₂. The choice of RR
includes also the choice of inspiratory to expiratory ratio (I:E) as well as the addition or
not of inspiratory plateau. In patients ventilated in controlled modes, there is no need
to try to match the mechanical to the neural respiratory rate or inspiratory time, as
these modes are used in patients lacking spontaneous breathing activity.
As several constraints tend to limit VT, there is usually a need for RR higher than the
normal 12-18 br/min (for adults). Clearly, RR cannot be increased indefinitely, as
sufficient time for passive expiration has to be allowed, otherwise dynamic
hyperinflation will develop. The time required for passive expiration is proportional to
the respiratory system resistance and compliance. Therefore, patients with ARDS, who
have low resistance and compliance require short expiratory time, while patients with
obstructive lung disease, who have high resistance and normal compliance require prolonged expiratory time. Observation of the expiratory flow waveform provides information on the adequacy of expiratory time (Figure 19). Manipulation of expiratory time can be achieved, not only through changes in RR, but also in I:E ratio (Figure 22). The normal I:E ratio is between 1:2 and 1:1.5, but in patients with obstructive lung disease, much lower ratios are preferred to allow lung emptying (see module on COPD).

![Image](image.png)

Figure 22: Inspiratory to Expiratory ratio, Plateau time, RR, Inspiratory Flow. Mechanical Ventilation module, ESICM Academy, 2018

Figure 22: In volume control mode I:E ratio, Plateau time, RR, Flow and VT are interconnected by their mathematical relationship and not dependent on the patient. A setting of RR of 25 br/min means that each breath will last Ttot= 60/25 =2.4sec.

The user may choose an inspiratory time (for example 0.75sec) which will result in an expiratory time of Te = Ttot-Ti = 2.4 - 0.75 = 1.65sec and an I:E ratio of 0.75:1.65 = 1:2.2. If the user would like to increase expiratory time, and depending on the type of ventilator, either inspiratory time or I:E ratio has to be modified accordingly. An inspiratory plateau time increases mean intrathoracic pressure and acts as a second, higher level, of PEEP that helps improve oxygenation. The addition of inspiratory plateau time could benefit patients with ARDS, but should be avoided in patients with obstructive lung diseases where the aim is to provide maximum expiratory time. Again, depending on the ventilator brand, the inspiratory plateau is set as an absolute time or as a percent of inspiratory time. In this case, flow is computed based on the set total inspiratory time and volume. In some ventilators the operator chooses the inspiratory time, tidal volume and inspiratory flow. To achieve an inspiratory plateau, the chosen flow has to deliver the set tidal at a shorter time than the set inspiratory time. Luckily in all ventilator brands these calculations are presented on the ventilator setup screen, so the user does not have to perform them manually.

The addition of inspiratory plateau will provide additional time of lung inflation which provides a second higher level of PEEP. Thus, an inspiratory plateau would be desired to improve oxygenation. On the contrary, if flow limitation is the main problem, then an inspiratory plateau would necessitate a decrease in expiratory time for the same RR, which would compromise lung emptying. Additionally, the inspiratory plateau increases mean airway/intrathoracic pressure affecting hemodynamics just as high PEEP does.
In patients ventilated in pressure control mode, the inspiratory time will determine the tidal volume for a given inspiratory pressure setting as shown in Figure 21. Therefore in patients ventilated in pressure control mode, a change in tidal volume can be made by changing either inspiratory pressure or time.

**Note**

at end-inspiration the plateau and driving pressure (Pplat-PEEP) will be the same for the same tidal volume, whether the patient is ventilated in volume or pressure control mode. This is determined by respiratory system compliance:

\[
CRS = \frac{VT}{P_{Pl} - PEEP}
\]

**In text References**

(Tobin and Alex. 1994; Cavalcanti et al. 2017; Sahetya, Goligher and Brower 2017; Franchineau et al. 2017; Hess 2015; Talmor et al. 2008; Amato et al. 2015; Chiumello et al. 2008; Terragni et al. 2013; Cressoni et al. 2016)

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4. 3. Assisted modes of ventilation

As discussed above, when a patient's effort triggers the delivery of positive pressure, the mode is termed 'assisted'. The choice of ventilating a patient in an assisted mode implies that the patient is able to trigger the ventilator at an acceptable rate. However, the patient is unable to maintain adequate ventilation to achieve the target PCO₂. The ventilator is therefore needed to contribute to the work of breathing. There a variety of modes of delivery of positive pressure after triggering. The modes of assisted ventilation include:

• Pressure support: a constant pressure is provided during inspiration. The level of assist is constant and not related to patient’s effort. The delivered volume depends on the level of assist, patient’s effort and total respiratory system compliance
• Volume support: a pre-set volume is provided after triggering. The level of assist decreases with increasing patient’s effort. The delivered volume is constant and does not depend on patient's effort
• PAV and NAVA: a pressure is delivered, proportional to patient's effort or diaphragmatic activity, respectively. Both the level of assist and the delivered volume increase with increasing patient effort (Figure 23, Figure 24, Figure 25).
Figure 23: A. Assisted modes of ventilation – Level of assist in relation to patient’s effort – In Pressure Support.

Figure 24: B. Assisted modes of ventilation – Level of assist in relation to patient’s effort – In Assist-Volume.

Figure 25: C. Assisted modes of ventilation – Level of assist in relation to patient’s effort – In Proportional-assist Ventilation.

Figure 23, Figure 24, Figure 25: Response of ventilator-provided support in experimentally increased patient effort by increase in PETCO₂. In Pressure Support (A) assist level remains unchanged at the set level of pressure. In Assist-Volume (B) the level of assist decreases so that the delivered volume remains unchanged. In Proportional-assist ventilation (PAV, C), the level of assist increases proportionally to

In all assisted modes, the necessary ventilator settings required for proper function of the ventilator include:

- Inspiratory trigger mode and corresponding setting
- Mode and level of assist
- Pressurisation rate or rising time (in pressure support mode)
- Expiratory trigger mode and corresponding setting
- PEEP and FiO₂

**Note**

Setting the ventilator in assisted modes is more challenging than in controlled modes because of the complex interactions between the patient’s system of breathing control and the ventilator-delivered breaths.

**Note**

Although VILI has only been studied during controlled mechanical ventilation, there is no reason to believe that high distending pressures are not just as harmful during assisted ventilation. Thus, it appears reasonable at the moment to avoid high tidal volumes and distending pressures during assisted ventilation.

**Note**

In proportional ventilation the assist provided by the ventilator is proportional to the inspiratory effort of the patient. The ventilator follows the patient’s effort and amplifies it to a degree determined by the physician. Two proportional ventilator modes are currently available, the proportional assist ventilation (PAV) and the Neurally Adjusted Ventilatory Assist (NAVA).

4.3. Pressure Support

Pressure support is the most commonly used mode of assisted ventilation in clinical practice. With this mode the ventilator, once triggered, provides a pre-set level of constant pressure. The level of assist is independent of patient’s effort. This pressure (Pvent) is added to patient’s muscle pressure (Pmus) and as a result the patient’s effort is augmented. Thus, the patient’s muscles and the ventilator can be considered pressure generators arranged in series where

\[ P_{tot} = P_{vent} + P_{mus} \]

Similarly to Pressure Control mode, during inspiration the equation of motion indicates that

\[ P_{toti} = P_{vent} + P_{mus(i)=V'\ast Ri+VTi\ast Ei} \]
Therefore the resulting tidal volume and inspiratory flow–time profile depends on 1) Pmus, 2) Pvent and 3) mechanical properties (i.e.Rrs and Ers) of respiratory system. The basic waveforms during Pressure Support are shown in Figure 26.

![Waveforms in Pressure Support](image)

Figure 26: Waveforms in Pressure Support
Ventilation. Mechanical Ventilation module, ESICM
Academy, 2018

Figure 26: Pressure, Flow and volume waveforms during Pressure Support: note that the shape of waveforms is similar to pressure control ventilation, but because Pmus differs from breath to breath, the waveforms of each breath also have small differences.

**Inspiratory trigger**

Flow or Pressure triggering requires the patient to decrease the flow or pressure respectively in the ventilator circuit to the pre-set level (Figure 10). When pressure triggering is selected, a part of the patient’s effort is ‘consumed’ or ‘wasted’ in decreasing circuit pressure to the set threshold before the ventilator inspiratory valve opens and inspiratory flow begins. Flow triggering is thus prefered and the usual setting is 2-3 l/min.

**Problems during triggering phase**

1. Triggering delay and Ineffective effort

At the beginning of a breath in mechanically ventilated patients with obstructive lung disease and dynamic hyperinflation, the inspiratory muscles start contracting at volumes above passive FRC where alveolar pressure is positive (PEEPi). In this case, in order to trigger the ventilator the patient must first counterbalance PEEPi in order to decrease alveolar pressure below external PEEPe. Therefore, a portion of Pmus is dissipated to counteract PEEPi (elastic threshold load) and as a consequence there is a delay between the beginning of inspiratory effort and the triggering. In some cases the inspiratory effort of the patients is not able to counterbalance PEEPi resulting in the inability to trigger the ventilator (ineffective effort). Ineffective efforts may occur in the absence of dynamic hyperinflation in patients with severe respiratory muscle weakness. Triggering delay and ineffective efforts can be best detected by recording esophageal pressure, but careful inspection of flow tracings may provide the same information (Figure 27).
Figure 27: Flow, airway pressure and oesophageal pressure recordings from a patient ventilated in Pressure Support mode. Small arrows on the oesophageal pressure tracing indicate patient’s inspiratory efforts. Note that several efforts (dotted circle) are not followed by inspiratory flow and increase in airway pressure (ineffective efforts). Note also the delay between the initiation of the patient’s effort and the initiation of mechanical breath (ovals, and magnified tracing on the right). Georgopoulou D, Prinianakis G, Kondili E. Bedside waveforms interpretation as a tool to identify patient-ventilator asynchronies Intensive Care Med. 2006 Jan;32(1):34-47.

Provided proper ventilator function, the strategies aiming to decrease the triggering delay and the number of ineffective efforts are 1) measures that decrease the magnitude of dynamic hyperinflation (decrease tidal volume by decreasing pressure support or decrease expiratory resistance by bronchodilators), 2) interventions which increase Pmus during the triggering phase (decrease sedation), 3) application of external PEEP and 4) decrease the threshold for triggering and/or use flow triggering.

**Note**

Flow distortion due to cardiac oscillation might be confused with ineffective efforts, particularly if the stroke volume of the patient is relatively high. The short duration (<0.3 sec) and the rapid frequency (close to heart rate) of flow distortion suggest cardiac oscillation rather than ineffective efforts.

**Auto-triggering**

Auto triggering refers to the phenomenon whereby ventilator is triggered in the absence of patient effort (Figure 28). This phenomenon may be caused by a random noise in the circuit, circuit leaks, presence of water in the circuit, cardiogenic oscillators or a low threshold for triggering. Hiccoughs may also trigger the ventilator. Auto-triggering may occur with any triggering mode when zero end-expiratory flow remains for some time before the next inspiration, usually when respiratory drive and rate are low. Interventions to manage auto-triggering include correction of circuit issues if present and increase in triggering sensitivity (mainly for cardiogenic oscillation-trigger). Hiccoughs usually creates significant intrathoracic pressure swings which cannot be managed by increasing triggering sensitivity, so only pharmacological interventions are helpful.
Figure 28: Flow, airway pressure and EMG recordings from a patient ventilated in Pressure Support mode. Arrows indicate auto-triggered breaths.

**Setting the Level of Assist in Pressure Support**

The ideal level of assist is such that patient's needs for ventilation are satisfied while his/her inspiratory muscles are active during inspiration, but not excessively to prevent distress and fatigue. The choice of assist level (in cmH2O of pressure) will be based on the derived ventilatory variables such as VT, RR and derived VE, and the clinical evaluation of patient's effort and comfort. This level is adjusted by trial and error. As the level of assist is stable during pressure support ventilation, the ventilator cannot adapt to changing the patient's needs for ventilation. This is why over- or under-assist may occur. In case of under-assist, the compensatory mechanisms of the patient's control of breathing system will lead to an increase in muscle pressure and/or respiratory rate. When this is insufficient, signs of respiratory distress and/or hypercapnia will develop. In case of over-assist, the patient will either relax the respiratory muscles during inspiration and/or develop hypocapnia and respiratory alkalosis. In patients with obstructive lung disease, high levels of assist resulting in high tidal volume may induce dynamic hyperinflation and unassisted efforts. Over-assist may also promote periodic breathing. Recent studies indicate that high assist, which can lead to complete respiratory muscle relaxation after triggering, may also induce diaphragmatic dysfunction.

**Rising time**

With new generation ventilators, the user may modify the time in which pressure delivered by the ventilator reaches the pre-set value (Figure 29). The equation of motion predicts that the rising time has a profound influence on flow-time waveform. An instantaneous increase of pressure to the desired level is associated with a sharp increase in inspiratory flow. A slower increase in rising time eliminates the sharp increase in inspiratory flow. This is because when pressure reaches its final value the opposing effect of elastic recoil (due to volume increase during the rising time) decreases the driving pressure (i.e. Paw+Pmus-Pel) for flow. Very low rising time may cause a round shape of the inspiratory flow graph. Note that a similar pattern (rounded shape) is observed if the patient continues to increase his/her inspiratory muscle pressure during mechanical inspiration.
Figure 29: Flow, airway pressure and oesophageal pressure recordings from a patient ventilated in Pressure Support mode, with long (A) and short (B) rising time. A slow increase in rising time (smaller angle at pressure waveform) is associated with a rounded shape of inspiratory flow because when pressure reaches its final value the opposing effect of elastic recoil has increased. On the contrary an instant increase of pressure to set level (angle at pressure waveform close to 90°) is associated with a sharp increase in inspiratory flow. Georgopoulos D, Prinianakis G, Kondili E. Bedside waveforms interpretation as a tool to identify patient-ventilator asynchronies Intensive Care Med. 2006 Jan;32(1):34-47.

**Expiratory trigger or Cycling off**

The aim of the expiratory trigger setting is to match the mechanical inspiratory time to the neural inspiratory time. The most commonly used expiratory trigger, or cycling off criterion, is when inspiratory flow has decreased to a pre-set percentage of the peak inspiratory flow. This threshold can be modified by the user, thus changing the duration of mechanical inspiration (Figure 30). It is important for the physician to recognise this expiratory asynchrony during PS ventilation.

Figure 30: Pressure and flow waveforms during pressure support ventilation with cycling off set at different % of peak inspiratory flow. Note the progressive increase in mechanical inspiratory time as the set cycling off % is decreased.

**Problems during cycling off**

1. Premature termination of mechanical inspiration
Irrespective of modes of ventilation, the opening of the exhalation valve is followed by reversal of flow from inspiratory to expiratory. Expiratory flow instantly achieves its highest value as dictated by the elastic recoil pressure at end-inspiration, airway pressure (i.e. PEEP) and expiratory resistance (respiratory system + ventilator circuit). Thereafter, flow decreases exponentially to zero following the corresponding decrease in elastic recoil pressure. If the patient continues to generate negative pressure using inspiratory muscle, the expiratory flow pattern will be affected, as shown in Figure 31. Zero or small inspiratory flow for some time after opening the exhalation valve indicates that inspiratory muscles continue to contract after the end of mechanical inspiration. Because volume (and thus elastic recoil pressure) decreases while inspiratory muscles continue to contract, a progressively increasing opposing pressure to expiratory flow ensues, causing a corresponding decrease in expiratory flow and deceleration of volume emptying. Relaxation of inspiratory muscles eliminates this opposing pressure and expiratory flow increases as determined by the elastic recoil pressure and resistive properties of the patient and expiratory circuit. Inspiratory effort may sometimes be adequate to decrease expiratory flow to zero and initiate the triggering process, so one inspiratory effort triggers the ventilator more than once, a phenomenon called ‘double triggering’ (Figure 32). In pressure support the duration of the second breath is usually relatively short because the inflation of the lung stars at lung volumes well above the passive FRC and as a result the driving pressure for inspiratory flow is relatively low. On the contrary, in assist-volume the ventilator once triggered delivers the pre-set volume, which adds to the remaining volume from the previous breath resulting in high distending pressures. The premature termination of pressure delivery is usually caused by low levels of pressure support, short time constant of respiratory system, set of cycling off at relatively high flow threshold and dynamic hyperinflation. These can be managed by increasing the level of pressure support and/or setting the cycling off at a lower percentage of peak flow.

![Flow, airway pressure, gastric, oesophageal and transdiaphragmatic pressure recordings from a patient ventilated in Pressure Support mode. Inspiratory muscle pressure continues after the cycling off criterion is reached and the mechanical](image)

Figure 31: Cycling off - premature opening of exhalation valve

Figure 31: Flow, airway pressure, gastric, oesophageal and transdiaphragmatic pressure recordings from a patient ventilated in Pressure Support mode. Inspiratory muscle pressure continues after the cycling off criterion is reached and the mechanical
inspiration is terminated, resulting in distortion of expiratory flow.


Figure 32: Double triggering. Mechanical Ventilation module, ESICM Academy, 2018

Figure 32: Flow, airway pressure and oesophageal pressure recordings from a patient ventilated in Pressure Support mode. A prolonged inspiratory effort decreases flow and triggers the ventilator twice in the same breath.

2. Delayed opening of exhalation valve

Identification of the delayed opening of the exhalation valve in relation to neural inspiration using the basic waveform is difficult, particularly if the patient does not use expiratory muscles. Relaxation of inspiratory muscles well before the end of mechanical inspiration results in pressure, flow and volume waveforms similar to those obtained with passive inflation, with a rather sharp decrease in inspiratory flow followed by an exponential decline (Figure 33). Delayed opening of the exhalation valve is usually caused by excessive support, long time constant of the respiratory system of the patient and cycling off set at a relatively low percentage of peak flow.
Figure 33: Airway pressure, flow, and muscle pressure recordings from a patient ventilated in Pressure Support mode. Note the termination of inspiratory effort soon after the initiation of the mechanical breath (line), and the duration of mechanical inspiration well after the decline in inspiratory muscle pressure. Note also the smooth exponential decline of flow. Georgopoulos D, Prinianakis G, Kondili E. Bedside waveforms interpretation as a tool to identify patient-ventilator asynchronies Intensive Care Med. 2006 Jan;32(1):34-47.

In text References


4. 3. 2. Proportional Assist Ventilation (PAV)

Proportional Assist Ventilation (PAV) delivers assistance in proportion to the patient’s respiratory drive and mechanical obstacles to the inspiratory breath (i.e. resistance and elastance).

Triggering in PAV+ is identical to conventional assisted ventilator modes. After triggering, the ventilator monitors inspiratory flow and volume and generates pressure which is, at any time during inspiration, the sum of the instantaneous flow and volume multiplied by a predetermined gain factor (% assist).

\[ \text{Paw} = (V' \times Rrs + VT \times Ers) \]

where Paw is instantaneous airway pressure, \(V'\) is instantaneous inspiratory flow, \(VT\)
is instantaneous lung volume above end-expiratory level, Rrs the respiratory system resistance and Ers the respiratory system elastance. The % percentage of offloading is set by the caregiver. The maximum % assist is 95% of the measured values of elastance and resistance. The pressure muscles need to develop (Pmus) is reduced by the % assist. When assist is set to zero, breathing is unassisted and Pmus undertakes the whole work of inspiration. When assist is set to 50%, half of the inspiratory work is performed by the patient and the other half by the ventilator. As Pmus declines towards the end of neural inspiration, inspiratory flow gradually decreases and when it reaches a preselected threshold, the ventilator terminates pressure generation.

PAV+ has utilises software that allows automatic and noninvasive measurement of respiratory system mechanics. At random intervals of 4 to 7 breaths, the ventilator applies a 300 msec pause maneuver at the end-inspiration (Figure 34).

Figure 34: Flow, Airway Pressure (Paw) and transdiaphragmatic pressure (Pdi) vs. time curves in a patient ventilated with PAV+. Notice an end-inspiratory pause maneuver during the second breath (shaded area). The maneuver is performed near the end of the neural breath when Pdi declines to zero and, therefore, there is no patient effort. Flow is zero, so that respiratory system resistance is zero. Paw measured at the end of the maneuver is the Paw occlusion pressure used to calculate respiratory system compliance.

Airway pressure at the end of the occlusion (Pawocclusion) is measured and Ers and compliance (Crs=1/Ers) are calculated as follows:

\[ E_{rs} = \frac{P_{awocclusion} - P_{PEEP}}{VT} \]

and

\[ C_{rs} = \frac{VT}{P_{awocclusion} - P_{PEEP}} \]

where PEEP is positive end–expiratory airway pressure. In the presence of PEEPi (dynamic hyperinflation), the calculated value of Ers overestimates respiratory system elastance (and the calculated Crs underestimates respiratory system compliance).

The measurement of Rrs is performed during the expiration that follows the pause
maneuver. Assuming that the expiratory flow early in exhalation is driven by the elastic recoil pressure (i.e. alveolar pressure, Palv), the software identifies three points on the expiratory flow-time curve corresponding to peak flow and 5 msec and 10 msec later. At these points Palv and total expiratory resistance (RTOT) are calculated as follows:

$$\text{Palv} = \text{Paw occlusion} - \Delta V \times Ers$$

$$RTOT = \frac{(\text{Palv} - \text{Paw})}{V'}$$

where $\Delta V$ is the exhaled volume up to the point of interest and $V'$ and Paw are the corresponding expiratory flow and airway pressure, respectively. The values of RTOT at these points are averaged and an estimate of RTOT is obtained. RTOT is the sum of the flow-dependent resistance of the endotracheal tube (Rtube) and that of the respiratory system (RrsPAV). Rtube is calculated using the following equation:

$$Rtube = a + bV'$$

where a and b are constants, depending on tube length and diameter, estimated using in vitro testing. RrsPAV is derived by subtraction of Rtube from RTOT.

**Advantages and limitations of PAV+**

PAV+ may be particularly helpful in several conditions but there are also contraindications to its use (Table 2).

The main advantages of PAV+ are:

- Improved patient-ventilator synchronisation as ventilator assist follows the patient’s effort both in terms of timing and ventilator support necessary.
- Continuous measurement and display of respiratory system mechanics.
- Less risk of lung overdistension, since this mode does not interfere with the operation of lung protective innate reflexes (i.e. Herring-Breuer).

Conditions that may result in inappropriate assist level with PAV+ are:

- The presence of a large difference between inspiratory and expiratory resistance (i.e. patients with obstructive lung disease). PAV+ measures expiratory resistance assuming that the difference between inspiratory and expiratory resistance is small. If the difference is large, the calculated Rrs is inaccurate and so is the assist level.
- Conditions that depress the respiratory centre (i.e. central nervous system diseases, sedatives, metabolic or respiratory alkalosis etc), affect respiratory muscle output (polyneuropathy, muscle fatigue, Guillain Barre, myasthenia gravis,
muscle relaxing drugs etc) or reduce the magnitude of the force generated by the diaphragm (e.g. in hyperinflation). As PAV+ follows patient effort, there is a risk of hypoventilation in the above conditions.

- Excessive assist (runaway). Runaway occurs when % assist is greater than the sum of Ers and Rrs at a particular point during inflation. As a result, the ventilator continues to deliver volume despite the fact that the patient has terminated his/her inspiratory effort. The volume will continue to increase until an alarm limit (pressure or volume) is activated, the compliance of the respiratory system is decreased because the respiratory system approaches total lung capacity or when expiratory muscles are recruited by the patient. With PAV+, runaway occurs rarely and only when the % of assist approaches 90%.

- Dynamic hyperinflation. Following triggering, pressure delivery in PAV+ is driven by patient effort. In the presence of dynamic hyperinflation, triggering delay will reduce the fraction of the patient’s effort that is being assisted. In addition, the calculated values of Crs and RTOT underestimate the actual values if PEEPi is present.

- The presence of leaks. In this case, the ventilator misinterprets the flow and volume escaping the circuit as a continuous patient effort and extends its assist delivery into exhalation. The effect of a large bronchopleural fistula is similar.

<table>
<thead>
<tr>
<th>Conditions where PAV+ is particularly beneficial</th>
<th>Conditions where PAV+ might be problematic</th>
<th>Conditions where PAV+ is contraindicated</th>
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<tbody>
<tr>
<td>Considerable patient-ventilator asynchrony</td>
<td>Dynamic hyperinflation</td>
<td>Deep ventilation, central apnea, CNS deficits</td>
</tr>
<tr>
<td>Monitoring of lung mechanics helpful</td>
<td>Severe neuromuscular disease</td>
<td>Muscle paralysis</td>
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<tr>
<td>High or variable ventilator demand</td>
<td>Marked hemodynamic compromise</td>
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<tr>
<td>Concern for lung overdistention</td>
<td>Bronchopleural fistula</td>
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<tr>
<td>Difficult to wean patients</td>
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<tr>
<td>Periodic breathing</td>
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Table 2: Applicability of PAV+ in different clinical conditions. Mechanical Ventilation module, ESICM Academy, 2018

Table 2: In the left column the physician should specifically prefer PAV+ or switch other assisted modes to PAV+. In the middle column PAV+ is not contraindicated and should be tried but specific supervision of patients is warranted.

**Setting the ventilator in PAV+**

The following algorithm to set PAV+ has been proposed by Xiouchaki et al in a randomised controlled study of 208 critically ill patients:

- Correct ideal body weight, endotracheal tube size and maximum airway pressure (40 cmH2O) must be set. Setting airway pressure limit is important, since it will protect the lungs from overdistention.
• Triggering, initial PEEP and fraction of inspired oxygen (FiO2) are set using common criteria. PEEP can be titrated according to changes in compliance: PEEP should be increased as long as compliance improves due to lung recruitment. In the absence of further Crs change, PEEP escalation should be stopped.

• The initial % level of assist must be sufficiently high to support the patient but not too high to cause over-assist and runaway. Starting with a 70% assist is a reasonable approach. If there is distress at 70% assist, possible reasons are the presence of delayed triggering due to dynamic hyperinflation or very low compliance. PEEP increase may improve distress in both conditions. If distress at 70% assist continues, % assist can be increased in steps of 5% up to 90% provided that there is no runaway phenomena. A high respiratory rate or a low VT (i.e. 3-4 ml/kg) do not necessitate action if there are no other signs of respiratory distress. Weaning with gradual decrease of % assist as long as the patient shows no signs of distress and blood gas targets are met follows the same rationale as in other modes of assisted ventilation.

• When changing from a conventional mode to PAV+, we may observe: no change in breathing pattern, rapid shallow breathing (an indication of the presence of significant number of ineffective efforts on previous modes) or apnoea (an indication of over-assist on previous modes). Waiting a few minutes is important to see how the breathing pattern evolves before deciding on the next step.

**Note**

An increase in PEEP from 0.9 cm H2 O to 3.5 cmH2 O in PAV+ has been shown to decrease the portion of supported inspiratory effort from 86% to 66%.

**In text References**

(Akoumianaki, Kondili and Georgopoulos 2012; Kondili et al. 2006; Xirouchaki et al. 2008)

4. 3. 3. Neuromuscular Ventilatory Assist (NAVA)

Neurally Adjusted Ventilatory Assist (NAVA) is based on the continuous recording of the electrical activity of the diaphragm (EAdi). The EAdi signal is proportional to the intensity of the diaphragmatic contraction. The more the diaphragm contracts, the greater the level of support delivered by the ventilator. The EAdi signal is obtained through a dedicated feeding tube with a mounted distal array of multiple electrodes. This signal is processed to provide the highest possible quality of signal (Figure 35). An EAdi increase above a predetermined value triggers the opening of the inspiratory valve. The ventilator then provides pressure (Paw) which is an amplification of the spontaneous EAdi:

\[ Paw = NAVAlevel \times EAdi \]
where Paw is the pressure provided by the ventilator in cmH₂O, NAVA level (in cmH₂O/µV) is the amplification factor of EAdi and EAdi the electrical activity of the diaphragm in µVolt. When EAdi decreases by a percentage in relation to its maximum value, the inspiratory phase is terminated and the exhalation valve opens. The triggering and cycling off criteria as well as the NAVA level are determined by the physician (Figure 35).

For safety reasons, the triggering of the ventilator in NAVA follows a ‘first come first serve’ approach. This means that the caregiver, besides EAdi triggering, determines a pneumatic (flow or pressure) triggering criterion, as in conventional assisted modes. The criterion that will be satisfied first (EAdi or pneumatic) will trigger the ventilator. Moreover, in the presence of a problem in the EAdi signal (noise, malfunction, removal of the catheter etc.), the ventilator will be automatically switched to pressure support ventilation with settings selected by the physician.

![Figure 35: Recording of diaphragm electrical activity during NAVA. Mechanical Ventilation module, ESICM Academy, 2018](image)

Setting the ventilator in NAVA

FiO₂ and PEEP settings in NAVA follow the same principles as in other ventilator modes. The EAdi inspiratory trigger is usually set to a default value of 0.5 µVolts, meaning that an EAdi increase of 0.5 µVolts above its minimum value will trigger the inspiratory phase. A 70% decrease in EAdi from its maximum value cycles-off the ventilator. The most challenging issue is the titration of the NAVA level. The NAVA level has units of cmH₂O/µV and the available range is between 0-15 cmH₂O/µV. A NAVA level of 0 cmH₂O/µV is considered similar to CPAP. Several approaches have been proposed to adjust the NAVA level at the bedside:
• NAVA level can be set through the ‘NAVA preview’ function of the ventilator Servo-i screen. This function estimates the NAVA level required to obtain the same peak pressure as during pressure support ventilation.
• Brander et al described a two-phase response during NAVA level escalation: in the first phase, NAVA level escalation leads to Paw and VT increase. Above that, any further increase will be associated with a relatively unchanged Paw and VT due to the activation of feedback control mechanisms. The ideal NAVA level is considered to be the inflection point from the first to the second phase.
• An alternative approach is to determine the EAdi during a spontaneously breathing trial (no ventilator assist). Then NAVA level can be titrated to decrease the EAdi recorded during the spontaneous breathing trial by 60%.

Irrespective of the initial NAVA level selected, NAVA level should be subsequently titrated according to patient comfort, signs of respiratory distress and observation of the flow, pressure and EAdi waveforms. If the support provided is too high, the nerve centres receive negative feedback leading to less support. This is detection by a difference in flow or pressure in the system from one cycle to the next. If the diaphragmatic contraction is insufficient, positive feedback will cause a more powerful EAdi signal and thus more support. As the patient respiratory function improves, EAdi decreases at the same NAVA level and NAVA level can be then reduced in a stepwise manner.

In text References

(Brander et al. 2009; Rozé et al. 2011)

References

• Georgopoulos D, Prinianakis G, Kondili E., Bedside waveforms interpretation as a tool to identify patient-ventilator asynchronies, 2006, PMID:16283171
• Kondili E, Akoumianaki E, Alexopoulou C, Georgopoulos D., Identifying and relieving asynchrony during mechanical ventilation., 2009, PMID:20477318
4. 4. Modes of ventilation with combinations of triggering and pressurisation variables

In modern commercially available ICU ventilators there are over fifty different ‘modes’ of ventilation. These do not represent completely different ways of delivering positive pressure ventilation, but rather using several combinations of triggering/cycling off and pressurisation variables in the basic modes described above. Knowledge of the basic principles of ventilation modes allows the user to understand any such ‘new’ mode. Several modes permit the combination of controlled and assisted breaths. In modern ventilators, all ‘control’ modes are ‘assist-control’. This means a triggering threshold is selected, and if the patient triggers the ventilator, the breath will be assisted in the same way the control breath is delivered. Spontaneous intermittent mandatory ventilation (SIMV) is similar to volume assist-control, but if and how (by pressure or volume) the spontaneous breaths will by supported is set separately by the user. Bi-level ventilation modes allow patients to breath spontaneously during any phase of the ventilator cycle, as the ventilator switches at pre-set intervals from a low level of pressure (P-low) to a high level of pressure (P-high). Again, spontaneous breaths can be supported (pressure support) or not. Airway pressure release ventilation (APRV) is similar to Bi-level ventilation using an inverse I:E ratio.

In several modes, the user chooses a target minute ventilation and the modes to provide the assisted and the mandatory breaths. These can be any combination of pressure and volume controlled. The ventilator continuously calculates the delivered volume and estimates if mandatory breaths are required to achieve the set minute
ventilation. A commonly used such mode is mandatory minute ventilation (MMV). The choice of pressure as controller leads to a delivered volume that is dependent on the mechanical properties of the respiratory system which may vary. In several pressure control or support modes, a minimum ‘safe’ tidal volume is set. If this is not reached, the ventilator adjusts the pressure delivered to meet the set ‘safe’ tidal volume. Finally, as artificial intelligence evolves, new ventilator software have become available. These have algorithms that continuously change ventilator settings (for example decrease/increase pressure support) after measuring the dependent variables of ventilation. The aim is to substitute the physician in selecting setting in well-described situations. For example, the gradual decrease of pressure support during weaning. It is questionable at the moment whether a given patient will have a course that is predictable enough to allow ventilator management by these algorithms.

In text References

(Tobin and Alex. 1994; Chatburn and Mireles-Cabodevilla. 2011)

References

- Chatburn RL, Mireles-Cabodevilla E., Closed-loop control of mechanical ventilation: description and classification of targeting schemes., 2011, PMID:21235841
5. Weaning the patient from mechanical ventilation

Although often life-saving, mechanical ventilation is associated with several life-threatening complications. Accordingly, it is important to discontinue mechanical ventilation and extubate the patient at the earliest possible opportunity.

5. 1. Definitions

**Weaning** is defined as the entire process of liberating the patient from mechanical support and the endotracheal tube.

**Weaning failure** is defined as the failure to pass spontaneous-breathing trial or the need for reintubation or NIV support within 48 hours (for some authors within 72 hours) following extubation.

**Extubation failure** is defined as the need for reintubation.

The majority of patients can be successfully extubated after one spontaneous breathing trial and are categorised as having undergone ‘simple weaning’. Patients who can be successfully extubated after the second or third spontaneous breathing trial within seven days from the first attempt are categorised as having undergone ‘difficult weaning’. Patients who require more than three spontaneous breathing trials or longer than seven days for successful extubation are categorised as having undergone ‘prolonged weaning.’ The proportion of patients in each weaning group varied between different studies and ranged from 30-59% in the simple weaning, 26-40% in the difficult weaning and 6-30% in prolonged weaning groups.

Weaning failure significantly affects patient outcomes. Numerous studies have shown that patients who undergo prolonged weaning have significantly higher ICU mortality rates compared to those with simple and difficult weaning. Patients failing extubation and needing reintubation have a much higher mortality rates than patients successfully extubated ranging from around 30 to 50% compared to 5-10% for successful extubation.

5. 2. The course of weaning
First step:
The weaning process starts at the time that the illness that led to the need for mechanical ventilation has (at least partially) resolved.

Second step:
Readiness-to-wean should be suspected early in the course of mechanical ventilation and assessed by objective criteria.

<table>
<thead>
<tr>
<th>Table 3: Readiness to wean criteria</th>
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<tbody>
<tr>
<td>Satisfactory oxygenation: e.g., PaO₂ / FiO₂ &gt;200 with PEEP ≤ 5 cm H₂O</td>
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<tr>
<td>Hemodynamic stability: e.g. no continuous vasopressor infusion</td>
</tr>
<tr>
<td>Adequate level of consciousness: Patient awake or easily aroused</td>
</tr>
<tr>
<td>Adequate Cough &amp; secretion management: Patient able to cough effectively, as roughly assessed by the presence of coughing in response to endotracheal aspiration</td>
</tr>
<tr>
<td>Respiratory physiology criterion: Rapid shallow breathing index RSBI &lt; 100 after 2 minutes of a spontaneous breathing trial</td>
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</table>

The RSBI index is the ratio of respiratory rate to tidal volume after 2 minutes of spontaneous breathing trial. RSBI was first introduced by Yang and Tobin and represents a sensitive screening test for early detection of readiness-to-wean. It can identify those who have a chance of passing a confirmatory SBT. It does not identify those who actually pass the SBT.

In text References
Third step:
Spontaneous Breathing Trial (SBT)
Once the readiness to wean has been confirmed in the presence of the criteria mentioned above, an SBT should be conducted. The SBT is needed to confirm the patient’s ability to breathe without assistance.

How to perform an SBT
Weaning guidelines suggest performing the SBT with no (T-piece strategy) or little ventilator assistance (low levels of inspiratory pressure support or continuous positive airway pressure). Ideally, SBT should be performed using the T-piece method, as that is the method that most accurately simulates the post-extubation physiological conditions.
Performing an SBT by applying low inspiratory pressure support (up to 7 cmH₂O) with or without continuous positive airway pressure has been shown to significantly decrease the work of breathing and may underestimate the patient’s ability to handle post-extubation workload.
Duration of SBT
In the majority of the patients, a 30 min trial has been shown to be adequate in identifying a successful or failed SBT. However, be aware that SBT might have to last for longer (up to 120 min) in patients at high-risk for reintubation such as elderly patients with COPD, heart failure, or neuromuscular disorders.

Note
During the initial few minutes of the SBT, the patient should be monitored attentively, before judgment is made to continue the SBT.

Criteria defining the success of SBT (Table 4)

<table>
<thead>
<tr>
<th>Table 4: Criteria of successful SBT</th>
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</thead>
<tbody>
<tr>
<td>Respiratory rate &lt; 35 breaths/minute</td>
</tr>
<tr>
<td>Good tolerance to spontaneous breathing trials</td>
</tr>
<tr>
<td>Heart rate &lt; 140 /minute or heart rate variability of &gt;20%</td>
</tr>
<tr>
<td>$\text{SaO}_2$&gt;90% or $\text{PaO}_2$ &gt; 60 mmHg on FiO₂&lt;0.4</td>
</tr>
<tr>
<td>Systolic blood pressure &gt;80 and $&lt;$180 mmHg or $&lt;$20% change from baseline</td>
</tr>
<tr>
<td>No signs of increased work of breathing or distress *</td>
</tr>
</tbody>
</table>

Accessory muscle use, paradoxical or asynchronous rib abdominal cage movements, intercostal retractions, nasal flaring, profuse diaphoresis, agitation
Failed spontaneous breathing Trial

Figure 38: Clinical signs of SBT failure. Deepak Talwar, Vikas DograJ Assoc Chest Physicians 2016;4:43-9

Table 5: Criteria of failure of SBT (see Figure 38)

<table>
<thead>
<tr>
<th>Clinical criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diaphoresis</td>
</tr>
<tr>
<td>Nasal flaring</td>
</tr>
<tr>
<td>Increasing respiratory effort</td>
</tr>
<tr>
<td>Tachycardia (increase in Heart rate &gt;40 bpm)</td>
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<tr>
<td>Cardiac arrhythmias</td>
</tr>
<tr>
<td>Hypotension</td>
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<tr>
<td>Apnea</td>
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<table>
<thead>
<tr>
<th>Gas exchange criteria</th>
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<tbody>
<tr>
<td>Increase of PetCO$_2$&gt;10 mm Hg</td>
</tr>
<tr>
<td>Decrease of arterial pH &lt;7.32</td>
</tr>
<tr>
<td>Decline in arterial pH&gt;0.07</td>
</tr>
<tr>
<td>PaO$_2$&lt;60 mmHg with an FiO$_2$&gt;0.40 (PaO$_2$/FiO$_2$ ratio &lt;150)</td>
</tr>
<tr>
<td>Fall in SpO2&gt;5%</td>
</tr>
</tbody>
</table>

Fourth step: Extubation.
Following a successful SBT, the patient should undergo assessment for and removal of the endotracheal tube.

In text References

5. 3. Diagnostic approach to weaning failure

The majority of patients can be extubated after the first SBT. A patient failing a weaning test or extubation is automatically allocated to the difficult-to-wean group. The most common causes of failing a SBT are:

- Incomplete resolution of critical illness
- Errors in assessing readiness to wean
- Presence of a new problem

Numerous studies have investigated the risk factors for weaning failure. Patients at a high risk for extubation failure are those >65 years of age and those with underlying
chronic cardiovascular or respiratory disease.

Following a failed SBT and before performing a new SBT, the physician should determine the reason of failure and subsequently develop an appropriate treatment strategy.

5. 3. 1. Pathophysiological determinants of weaning failure

The pathophysiology of patients suffering weaning failure is complex and multifactorial. Determination of the pathophysiological factors that cause weaning failure requires a dedicated clinician with in-depth knowledge of the pathophysiology of weaning failure (Figure 39).

![Figure 39: Adapted with permission from Perren A Brochard L Managing the apparent and hidden difficulties of weaning from mechanical ventilation. Intensive Care Med (2013) 39:1885–1895 with permission](image)

The primary pathophysiologic mechanisms related to weaning failure include:

- Respiratory pump insufficiency
- Cardiovascular dysfunction
- Neuromuscular disorders
- Psychological factors
- Metabolic/ endocrine diseases, alone or combined.

5. 3. 1. 1. Respiratory pump insufficiency

Respiratory pump insufficiency is probably the most common cause of weaning failure and may result in an imbalance between respiratory muscle workload and respiratory neuromuscular capability (Table 6).
Increased mechanical workload

An increase in the load on the respiratory muscle pump may result from increased ventilatory requirements and/or increased mechanical load (Table 7). Physiological studies indicate that compared to patients who are successfully extubated, patients with COPD who failed an SBT exhibit substantially higher respiratory workload, expressed either as higher inspiratory airway resistance and/or higher elastance and PEEPi.

Laryngeal injuries (both structure injury and abnormal vocal cord mobility) may significantly increase mechanical workload and are considered as a cause of post-extubation stridor and extubation failure. A recent prospective study including patients extubated after more than 24 h, showed a high incidence of laryngeal injuries. Moreover, in patients who needed reintubation, laryngeal granulation and vocal cord abnormalities were frequently observed.

**Note**

In patients who meet extubation criteria and judged at high risk for post-extubation stridor, a cuff leak test should be performed before extubation.

**Impaired respiratory neuromuscular capability**

Adequate respiratory capability requires a structurally intact respiratory pump and adequate signal transmission from the respiratory centre to the inspiratory muscles. The leading causes that may result in impaired respiratory capability are summarised...
Respiratory centre depression may rarely be the cause of weaning failure and is usually due to sedatives and opioids overdose. On the other hand, respiratory muscle dysfunction is believed to be a predominant mechanism of weaning failure, in particular in patients with COPD. Geometrical disarrangement of the respiratory muscles due to dynamic hyperinflation is considered the primary causative factor. Critical illness neuromyopathy (Critical illness polynmyopathy and/or myopathy) may also be consider as a common cause of weaning failure. This is seen particularly in patients with sepsis, COPD or in those who have received treatment with corticosteroids, and/or neuromuscular blockers. Of particular importance is the presence of Ventilator-Induced Diaphragmatic Dysfunction (VIDD) as the cause of weaning failure. A recent study reported that compared to those with simple weaning, patients who underwent prolonged and difficult weaning had a significantly higher prevalence of VIDD. The impaired respiratory neuromuscular capability may also present as a result of the development of respiratory muscle fatigue. Tension Time Index (TTI) is a physiological variable which quantifies the magnitude and duration of inspiratory muscle contraction (mainly the diaphragm). Studies in patients during SBT, have shown that TTI increased over the course of the trial in patients who failed whereas it remained unchanged in those who were successfully extubated. Moreover, some of the patients who failure weaning developed a TTI greater than 0.15. This value has been associated with respiratory muscle fatigue.

In text References


5. 3. 1. 2. Cardiovascular Dysfunction

Cardiovascular impairment is considered an important cause of weaning failure in patients with known or previously unrecognised left heart disease. The transition from mechanical ventilation to spontaneous breathing imposes an additional load on the cardiovascular system. This is because a decrease in intrathoracic pressure significantly affects preload and afterload of both right and left ventricle and is associated with increased oxygen consumption by the respiratory muscles. Several studies have shown that in patients with COPD, with or without preexisting cardiac disease, weaning is associated with a significant reduction in left ventricle ejection fraction, increased left ventricular afterload, increased adrenergic tone and decreased compliance of left ventricle (mainly in patients with preexisting heart disease). As a result, some patients may present with myocardial ischemia, acute heart decompression, and pulmonary oedema. Fluid overload in patients with or without pre-existing left ventricular dysfunction has been recognised as a cause of weaning and extubation failure. Positive fluid balance the day before extubation is identified as a
strong risk factor for extubation failure. Multiple studies have demonstrated that baseline values and changes in Brain Natriuretic Peptide (BNP), an indirect index of ventricular expansion and volume overload, are significantly higher in patients with weaning failure than in patients with successful extubation. See Figure 40.

Figure 40: Cardiocirculatory mechanisms seemingly leading to failure of the T-tube test

Figure 40: Cardiocirculatory mechanisms leading to failure of the T-tube test. IPS inspiratory pressure support, PEEP positive end-expiratory pressure, WOB work of breathing, ITP intrathoracic pressure, PaO₂ arterial oxygen tension, PaCO₂ arterial carbon dioxide tension. Adapted with permission from Perren A, Brochard L. Managing the apparent and hidden difficulties of weaning from mechanical ventilation. Intensive Care Med (2013) 39:1885–1895 with permission

In text References


5. 3. 1. 3. Other causes of weaning failure

Brain dysfunction and psychological disturbances
Brain dysfunction in patients who have weaning failure is mainly caused by delirium. Delirium, as assessed by the CAM-ICU has been significantly associated with difficult weaning and a higher risk of failed extubation. Psychological disturbances other than delirium, such as anxiety and depression have also associated with failed extubation.

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5. 4. Diagnostic workup and management of weaning failure
Diagnostic approach and management of the most common causes of weaning failure are summarised in Figure 41

5. 4. 1. The role of NIV in the management of weaning failure

NIV has been used at three different time points during the weaning process (Figure 42)

- To facilitate extubation
- To prevent reintubation in high-risk patients
- To treat postextubation acute respiratory failure

**NIV To Facilitate extubation**

NIV is used as a weaning method in patients who usually do not meet standard extubation criteria and cannot withstand a weaning test. A recent meta-analysis
revealed that in patients with COPD with hypercapnia this form of NIV application is associated with a reduced duration of invasive mechanical ventilation, decreased length of ICU stay and lower incidence of nosocomial pneumonia.

**NIV for Prevention of post-extubation respiratory failure**

Early NIV application after extubation prevents post-extubation respiratory failure, decreases re-intubation rates and ICU mortality in patients at high risk for extubation failure. High risk for extubation failure was defined as patients >65 years old with underlying chronic cardiovascular or respiratory disease.

**Treatment of post-extubation respiratory failure**

Two large randomised studies not only failed to prove any benefit of NIV as rescue therapy in post-extubation respiratory failure, but showed an increase in ICU mortality. This was mainly attributed to delayed reintubation. Nevertheless, in a very selected group of patients, mainly with COPD, a trial of NIV could be considered as long as it does not delay reintubation in case of failure.

**In text References**


5. 4. 2. The role of tracheostomy

Performing a tracheostomy is currently common in patients with COPD requiring prolonged mechanical ventilation. However, the right timing and the impact on outcome remains debatable. Some studies in patients with prolonged weaning have shown that a tracheostomy did not favorably influence ICU survival. Other studies reported that a tracheostomy performed in ICU for long-term mechanically ventilated patients was associated with lower ICU and in-hospital mortality rates. From a physiological point of view, compared to the endotracheal tube, the tracheostomy in these patients may significantly reduce airway resistance and dead space. Hence the work of breathing and ventilation requirements are reduced. Efforts should be made to identify patients who might benefit from a tracheostomy to avoid unnecessary prolonged mechanical ventilation.

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(Durbin CG 2010; Diehl et al. 1999)

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6. Complications of Mechanical Ventilation

6. 1. Ventilator Induced Lung Injury (VILI)

Mechanical ventilation per se can aggravate or even induce lung injury leading to an entity histopathologically indistinguishable from ARDS. This is called ‘Ventilator Induced Lung Injury’ (VILI).

6. 1. 1. Mechanisms of ventilator-induced lung injury

The main mechanisms of VILI include:

- Increased filtration: Ventilation with high distending pressures increases filtration through an increase in transmural vascular pressure and/or a decrease in lung interstitial pressure. Moreover, mechanical ventilation can lead to surfactant depletion or inactivation. It can lead to an increase in alveolar surface tension along with decreases in the peri-microvascular pressure (pressure in the perivascular space surrounding extra-alveolar vessels).

- High alveolo-capillary permeability: Lung overinflation may damage the alveolo-capillary barrier increasing the epithelial and endothelial permeability. The combination of increased filtration and high alveolo-capillary permeability is responsible for the interstitial and alveolar oedema.

- Leukocyte activation and release of proinflammatory mediators which contribute to pulmonary and systemic inflammatory response known as ‘biotrauma’. Proinflammatory mediators and cytokines implicated in the pathogenesis of biotrauma include: tumor necrosis factor (TNF-a), interleukin IL-1β, IL-6, IL-8, macrophage inflammatory protein (MIP-2), nuclear factor-κB (NF-κB).

6. 1. 2. Physiologic determinants of VILI

Mechanisms related both to ventilator settings and to lungs’ pathology are associated with alveolar injury and activation of inflammation.

**Mechanisms related to ventilator settings**
Mechanical ventilation resulting in high distending pressures and volumes can lead to barotrauma or volutrauma. ‘Stress’, instead of barotrauma and ‘strain’, instead of volutrauma, describe more accurately the mechanisms of lung injury. Stress is the transpulmonary pressure (PL), which is the pressure difference across the lung (PL = Pairway – Ppleural). Strain can be static, which equals VT, or dynamic, which is computed by the ratio of VT to functional residual capacity (FRC). Because static strain fails to incorporate the size of the aerated lung (the lower the size of the aerated lung the higher the strain applied at the same VT), dynamic strain better assesses the risk of VILI caused by VT. Lung stress and strain are two faces of the same coin, as shown by the following equation:

\[
PL = EL_{spec} \times \frac{VT}{FRC_{or\ Stress}} = EL_{spec} \times Strain
\]

where ELspec is the tissue elastance of the lung and averages 12cmH2O.

Respiratory rate and the inspiratory flow rate have also been implicated the pathogenesis of VILI. They express how many times (respiratory rate) and how fast (flow rate) a potential harmful stress and strain is applied to the lung.

All of these mechanical factors (volume, pressure, rate and flow) can be considered parts of a single physical entity: the mechanical power. Mechanical power is derived by multiplying the classical equation of motion by the tidal volume and respiratory rate.

**Mechanisms related to lungs pathophysiology**

The same applied stress, strain and/or mechanical power may have considerably different effects depending on the heterogeneity of the lung parenchyma. Neighbouring alveoli are mechanically interdependent. Alveolar units located at the margins of atelectatic lung regions can collapse and reopen with each tidal breath. This phenomenon, known as atelectrauma, subjects these alveoli to high shear strain with each breath. Heterogeneously inflated regions of the lung may act as stress raisers, amplifying local mechanical stress.

6. 1. 3. Strategies to protect from VILI

The main strategies to protect from VILI are summarised in Table 8. They can be divided into ventilator and non-ventilator strategies.
6. 1. 3. 1. Ventilator strategies

**Strategies to protect from excessive stress and strain**

To prevent excessive lung stress and strain during MV, tidal volume and pressure can be selected as follows:

- VT to target a predicted body weight (PBW) of approximately 6 ml/kg (ARDS Network protocol). Further reductions of VT (4-5 ml/kg/PBW) are proposed to maintain an inspiratory plateau pressure (Pplat) ≤ 30cmH₂O. However, the same VT is associated with different levels of strain depending of the size of the aerated lung.
- VT adjusted to reduce dynamic strain (VT/FRC). Studies have shown that VILI occurs when dynamic strain is higher than 1.5-2.0. Nevertheless, dynamic strain calculation requires the measurement of FRC.
- VT adjusted by calculation of driving pressure (DP), which is the difference between the airway Pplat and the PEEP (DP=Pplat-PEEP). Although mortality increases steeply at DPs above 14 cmH₂O, a safe DP threshold has not been identified yet.
- Estimation of PL, although reliable targets for safe limit of end-inspiratory PL are lacking: a study identified a PL of 27cmH₂O as a safe upper limit while other investigators suggest considerably lower end-inspiratory PL values, < 13-15 cmH₂O.

[Note]

A recent meta-analysis of >3000 patients detected higher survival in patients with DP ≤14cmH₂O and found that a lower DP was the strongest predictor of improved outcome among the various ventilator variables. Changes in VT, Pplat and PEEP did not influence survival at a constant DP (Amato et al).

**Ventilator strategies to reduce lung heterogeneity**

The main ventilator strategy to reduce lung heterogeneity is the application of positive end-expiratory pressure (PEEP). PEEP has two main effects:

(a) Prevents end-expiratory collapse of unstable alveolar units (units that open and close with every breath). This minimises the risk of atelectrauma.
(b) Increases the number of aerated alveoli. This improves lung compliance and homogeneity. Hence, shear strain and injury at the margins between aerated and collapsed lung tissue are further reduced.

Patients with ARDS do not respond uniformly to an increase in PEEP: some exhibit lung reductions in collapsed lung while others exhibit minimal lung recruitment. Increases in PEEP will raise end-inspiratory stress in both groups. However, in the first group the recruitment of lung units will reduce mechanical strain and atelectrauma, decreasing dead space ventilation and shunt. In the second group the increased stress without recruitment will worsen mechanical strain, causing additional distension of already aerated alveoli and dead space ventilation.
Therefore, before applying PEEP we must first assess whether the lung is recruitable and then select the adequate PEEP. The ways to assess lung recruitability and adjust PEEP at the bedside are discussed in section 4.2.

**High-frequency Oscillatory Ventilation (HFOV)**

High-frequency Oscillatory Ventilation (HFOV) consists of a simple circuit where oxygenated, humidified gas (bias flow) is passed across the path of an oscillating membrane at a set frequency (usually 3-15 Hz) generating VT well below anatomic dead space (1–3 mL/kg PBW). The fraction of inspired oxygen (FiO2) and the mean airway pressure determines the PaO2 while the frequency of oscillations, their pressure amplitude and the inspiratory time determine the PaCO2. Theoretically, the very low VT limits overdistension while the high mean Paw may both recruit collapsed alveoli and prevent cyclic recruitment-derecruitment of unstable lung units. HFOV should be considered in severe patients with ARDS who have failed conventional ventilator strategies and should be performed in centres with expertise in using this mode.

**Note**

in two recent large randomised trials, ventilation with HFOV was associated with difference in mortality and possible harm compared to conventional ventilation. However, there are limitations in each of these trials e.g. higher doses of sedation, fluids, vasopressors and neuromuscular blocking agents in the HFOV groups.

6. 1. 3. 2. Non ventilator strategies to protect from VILI

Non ventilator strategies to protect from VILI include:

- Prone positioning
- Neuromuscular blockade
- Extracorporeal dioxide removal (ECCO₂R) and the extracorporeal membrane oxygenation (ECMO)

Prone positioning alters the distribution of lung ventilation and lung perfusion. In the normal lung, ventilation favors ventral lung regions because alveolar size decreases from ventral to dorsal regions. In contrast to ventilation, pulmonary perfusion preferentially distributes to the dorsal lung regions. In patients with ARDS, lung oedema further diminishes the aeration of dorsal regions and aggravates shunt in these regions (high perfusion, less aeration). Prone position increases the aeration of dorsal lung units. The redistribution of aeration does not impair gas exchange in the ventral region while pulmonary perfusion remains preferentially distributed to the dorsal lung regions. Therefore, alveolar/perfusion matching improves, shunt fraction decreases and gas exchange is improved. Moreover, the more homogenous lung ventilation and the increased size of the aerated lung protect the lung from abnormal stress and strain and mitigate the risk of VILI. A recent multicentre randomised
controlled trial (PROSEVA) found that placing patients with severe ARDS (\(\text{PaO}_2/\text{FiO}_2<150\) mmHg) in the prone position for at least 16 hours/day improved survival compared to semirecumbent supine position.

**Note**

Prone positioning sessions should be initiated early in the course of severe ARDS, applied for long periods (>10-12 hours/session) and must be combined with other lung protective strategies (small VT, low P\(_{\text{plat}}\) and driving pressures, individualized PEEP titration etc).

In a recent multicentre randomised controlled trial in 340 patients with moderate-to-severe ARDS, early neuromuscular blockade decreased 90-day mortality. The exact mechanism is unknown and it is hypothesised that a lower transpulmonary pressure along with improved patient-ventilator interaction (less patient-ventilator asynchrony) facilitated lung-protective ventilation. Another hypothesis is that cisatracurium has anti-inflammatory properties by blockade of nicotine-acetylcholine receptor a1 signaling.

Extracorporeal strategies include extracorporeal dioxide removal (ECCO\(_2\)R) and extracorporeal membrane oxygenation (ECMO). ECCO\(_2\)R can be used when the application of lung protective ventilation is impeded by hypercapnia and acidosis. By lowering \(\text{PaCO}_2\), the veno-venous ECCO\(_2\)R may permit an ultra-protective ventilation strategy with very low VTs (3-4 ml/kg/PBW). ECMO may be considered in persistent hypoxemia. Ongoing multicentre controlled studies are investigating the role of ECCO\(_2\)R and ECMO in patients with severe ARDS.

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6. 2. Ventilator Induced Diaphragmatic Dysfunction (VIDD)

**Definition**

Ventilator-induced diaphragmatic dysfunction (VIDD) is defined as a loss of diaphragmatic force-generating capacity specifically related to the use of mechanical ventilation (MV). VIDD is specifically related to MV and cannot be solely explained by other factors such as sepsis, drugs or metabolic derangements, although these factors may further exacerbate diaphragmatic weakness. Unlike ICU-acquired weakness, in VIDD phrenic nerve signal transmission and signal transduction at the neuromuscular junction is normal or even increased.

The exact prevalence of VIDD is unknown. Studies have shown that diaphragmatic dysfunction is two times more frequent than ICU-acquired weakness, while 80% of
patients with ICU-acquired weakness have diaphragmatic dysfunction. VIDD is associated with difficult weaning and weaning failure and may increase ICU and hospital mortality.

**Pathophysiology**

Diaphragmatic weakness in VIDD is due to both atrophy and contractile dysfunction. The main histopathological and biochemical changes are:

- muscle fibre atrophy,
- muscle fibre remodeling and
- muscle fibre injury (disrupted myofibrils, increased numbers of vascular structures and abnormal mitochondria).

Diaphragm fibre atrophy results from MV-induced increase in proteolysis coupled with depressed protein synthesis. Proteolysis is activated within 6 hrs of controlled MV and associated with increased oxidative stress. Diaphragmatic contractile dysfunction is caused by: MV-induced oxidative modifications to contractile proteins, resulting in depressed fibre sensitivity to calcium, protease activation resulting in sarcomere disruption, and loss of myosin heavy chain protein.

**Note**

Studies have shown that 12 hrs of controlled MV results in a 15% reduction of diaphragmatic fibres and that this reduction approaches 30% after 18-24 hrs of MV. Both controlled and assisted MV (with high assist) can induce diaphragmatic atrophy albeit at a slower rate.

**Diagnostic approach**

The presence of VIDD should be considered in any mechanically ventilated patient experiencing difficulties in weaning. The diagnostic approach of VIDD includes:

- Exclusion of other causes of weaning failure
- Assessment of the respiratory muscle strength
- Assessment of diaphragmatic strength
- Diaphragmatic ultrasonography

Causes of weaning failure other than VIDD such as underlying lung diseases, drugs, electrolyte disturbances, malnutrition, congestive heart failure, central nervous system disorders, neuromuscular disorders should be considered and evaluated accordingly. Respiratory muscle strength is most commonly assessed through the measurement of maximum inspiratory pressure (MIP). MIP is easily measured at the bedside but it assesses all inspiratory muscles and not specifically the diaphragm. It is affected by lung diseases and it is effort dependent. The gold standard for diaphragmatic strength evaluation is the measurement of
maximal transdiaphragmatic pressure (Pdimax, the difference between esophageal and gastric pressure) in combination with transdermal phrenic nerve (magnetic) stimulation. This technique requires oesophageal and gastric balloon placement and equipment for magnetic stimulation.

Ultrasonography of the diaphragm is easy to perform and non-invasive. It allows assessment of diaphragmatic function and structure and may exclude other causes of weaning failure (e.g. heart failure, pneumonia etc). Two ultrasonography parameters are evaluated: diaphragmatic excursion and thickening fraction of the diaphragmatic muscle during inspiration. Thickening fraction is defined as thickness at end-inspiration minus thickness at end-expiration divided by thickness at end-expiration. A diaphragmatic excursion less than 1cm and a thickening fraction < 30% have been associated with weaning failure.

**Note**

Patients may suffer from VIDD even if they don't have weaning difficulties.

**Management**

VIDD develops rapidly after MV implementation and the longer the duration of MV the more serious the damage to the diaphragm. Diaphragmatic inactivity is the primary cause of VIDD. Therefore, limiting the duration of MV and particularly controlled MV is the key to reducing the chance of VIDD. Pharmacological interventions, such as antioxidants and protease inhibitors are currently under investigation.

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6. 3. Ventilator Associated Pneumonia (VAP)

**Definition-Epidemiology**
Ventilator-associated pneumonia (VAP) is defined as pneumonia that occurs more than 48 hours following endotracheal intubation and invasive mechanical ventilation. It represents half of all cases of hospital-acquired pneumonia and is estimated to occur in 10-27% of all mechanically ventilated patients. It is the second most common infection in the Intensive Care Unit (ICU). The risk of VAP is higher in the first 5 days following mechanical ventilation (3% risk/day) and declines thereafter (2%/day between days 5-10 and 1%/day after the 10th day of ventilation).

**Clinical impact**
VAP is the leading cause of death related to nosocomial infection in critically ill patients. All-cause mortality rates associated with VAP range from 20% to 50% and mortality directly related to VAP is estimated at 9-13%. Mortality depends on the causative organism and the severity of the underlying medical illness. VAP prolongs the length of mechanical ventilation and hospitalisation.

**Prevention**
Measures proved to prevent VAP include:

- Head-of-bed elevation
- Thromboprophylaxis
- Alcohol-based hand washing policy
- Early discontinuation of MV

The use of NIV as a means to prevent intubation and avoid re-intubation and to decrease the duration of mechanical ventilation (early weaning and extubation) plays an important role in decreasing the incidence of VAP.

The role of other suggested measures (stress ulcer prophylaxis, early tracheostomy,
subglottic secretion drainage, conical and polyurethane endotracheal tube cuffs, silver/antibiotic coated endotracheal tubes, prophylactic probiotics, selective digestive decontamination and oral care with chlorhexidine) remain controversial.

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