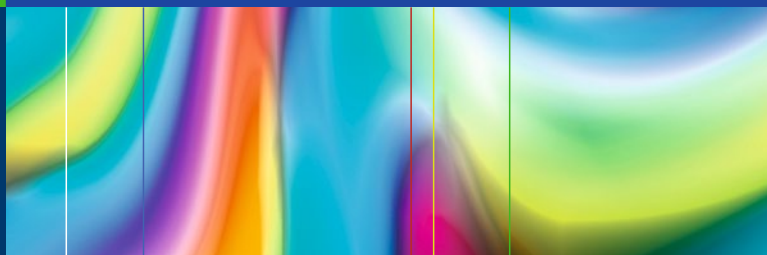


Susan R. Wilcox
Ani Aydin
Evie G. Marcolini



Mechanical Ventilation in Emergency Medicine

 Springer

Mechanical Ventilation in Emergency Medicine

Susan R. Wilcox • Ani Aydin
Evie G. Marcolini

Mechanical Ventilation in Emergency Medicine



Springer

Susan R. Wilcox
Department of Emergency
Medicine
Massachusetts General Hospital
Boston, MA
USA

Ani Aydin
Departments of Surgery and
Neurology
University of Vermont Medical
Center
Burlington, VT
USA

Evie G. Marcolini
Departments of Surgery and
Neurology
University of Vermont Medical
Center
Burlington, VT
USA

ISBN 978-3-319-98409-4 ISBN 978-3-319-98410-0 (eBook)
<https://doi.org/10.1007/978-3-319-98410-0>

Library of Congress Control Number: 2018957093

© Springer Nature Switzerland AG 2019

This work is subject to copyright. All rights are reserved by the Publisher, whether the whole or part of the material is concerned, specifically the rights of translation, reprinting, reuse of illustrations, recitation, broadcasting, reproduction on microfilms or in any other physical way, and transmission or information storage and retrieval, electronic adaptation, computer software, or by similar or dissimilar methodology now known or hereafter developed.

The use of general descriptive names, registered names, trademarks, service marks, etc. in this publication does not imply, even in the absence of a specific statement, that such names are exempt from the relevant protective laws and regulations and therefore free for general use.

The publisher, the authors, and the editors are safe to assume that the advice and information in this book are believed to be true and accurate at the date of publication. Neither the publisher nor the authors or the editors give a warranty, express or implied, with respect to the material contained herein or for any errors or omissions that may have been made. The publisher remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

This Springer imprint is published by the registered company Springer Nature Switzerland AG

The registered company address is: Gewerbestrasse 11, 6330 Cham, Switzerland

Contents

1	Introduction	1
2	Terminology and Definitions	5
	Ventilator Basics	5
	Physiology Terms	6
	Phases of Mechanical Breathing	6
	Ventilator Settings	8
	Ventilator Modes	11
	Conventional Modes of Ventilation	11
	Suggested Reading	13
3	Review of Physiology and Pathophysiology	15
	Gas Exchange	15
	Issues with Oxygenation	17
	Hypoxemia	17
	Hypoxic Vasoconstriction	25
	Atelectasis and Derecruitment	27
	Issues with Ventilation	27
	Compliance and Resistance	29
	Suggested Reading	34
4	Noninvasive Respiratory Support	35
	Oxygen Support	35
	High Flow Nasal Cannula	35
	Noninvasive Positive Pressure Ventilation	37
	References	40
5	Modes of Invasive Mechanical Ventilation	43
	Modes of Invasive Ventilation	43
	Pressures on the Ventilator	49
	Reference	52
	Suggested Reading	52

6	Understanding the Ventilator Screen	53
	Suggested Reading	59
7	Placing the Patient on the Ventilator	61
	Anticipating Physiologic Changes	61
	Setting the Ventilator.	62
	After Initial Settings	66
	Suggested Reading	66
8	Specific Circumstances: Acute Respiratory Distress Syndrome (ARDS)	69
	Recruitment Maneuvers	73
	Neuromuscular Blockade	75
	References.	77
9	Specific Circumstances: Asthma and COPD	79
	COPD	84
	Suggested Reading	88
10	Specific Circumstances: Neurologic Injury	89
	Traumatic Brain Injury	89
	Ischemic Stroke.	92
	Intracranial Hemorrhage	93
	Status Epilepticus	94
	References.	94
11	Troubleshooting the Ventilated Patient	97
	Suggested Reading	99
12	Case Studies in Mechanical Ventilation	101
	Case 1	101
	Case 2	102
	Case 3	104
	Case 4	105
	Case Study Answers	107
	Case 1	107
	Case 2	108
	Case 3	110
	Case 4	112
	Suggested Reading	114
13	Conclusions and Key Concepts	115
	Index	119

About the Authors

Susan R. Wilcox attended medical school at Washington University School of Medicine and trained in Emergency Medicine in the Harvard Affiliated Emergency Medicine Residency. After residency, she completed an Anesthesia Critical Care Fellowship at Massachusetts General Hospital (MGH). She has since divided her time between the Emergency Department and Intensive Care Units, including working in surgical, medical, and cardiac critical care. She is currently an Assistant Professor of Emergency Medicine at Harvard Medical School, and she is the Chief of the Division of Critical Care in the Department of Emergency Medicine at MGH.

Ani Aydin is an Assistant Professor of Emergency Medicine at Yale School of Medicine. She completed a Trauma-Surgical Critical Care Fellowship at the R Adams Cowley Shock Trauma Center in Baltimore, Maryland. She currently works as an attending physician in the Emergency Department and Surgical Intensive Care Unit at Yale-New Haven Hospital. Dr. Aydin is also the founder and Immediate Past Chairperson of the Society for Academic Emergency Medicine (SAEM) Critical Care Medicine Interest Group.

Evie G. Marcolini is an Assistant Professor in Emergency Medicine and Neurocritical Care at the University of Vermont College of Medicine. She completed a Surgical Critical Care Fellowship at the R Adams Cowley Shock Trauma Center in Baltimore and now divides her clinical time at UVM between Emergency Medicine and Neurocritical Care. Evie is on the Board of Directors for the American Academy for Emergency

Medicine. She is a member of the Ethics Committees for the American College of Critical Care, Neurocritical Care Society, and the University of Vermont Medical Center. She is also active in wilderness medicine and teaches for Wilderness Medical Associates International. In her spare time, she loves to skijore with her husband and two Siberian huskies.

Chapter 1

Introduction



Mechanical ventilation is a procedure often performed in patients in the emergency department (ED) who present in respiratory distress. The indications of mechanical ventilation include airway protection, treatment of hypoxemic respiratory failure, treatment of hypercapnic respiratory failure, or treatment of a combined hypoxic and hypercapnic respiratory failure. On some occasions, patients are also intubated and placed on mechanical ventilation for emergent procedures in the ED, such as the traumatically injured and combative patient who needs emergent imaging. However, intubation and initiation of mechanical ventilation requires a great degree of vigilance, as committing to this therapy can affect the patient's overall course.

Traditionally, mechanical ventilation has not been taught as a core component of Emergency Medicine practice, instead, principles of ventilation have been left to intensivists and respiratory therapists. However, with increasing boarding times in the ED and increased acuity of our patients, emergency physicians are frequently caring for mechanically ventilated patients for longer and longer periods of time. Additionally, the data supporting the importance of good ventilator management in all critically ill patients continues to increase.

Compared to many of the other procedures and assessments emergency physicians perform, management of basic mechanical ventilation is relatively simple. While there are

occasionally patients who are very difficult to oxygenate and ventilate and require specialist assistance, the vast majority of patients can be cared for by applying straightforward, evidence-based principles. Ventilator management can seem intimidating due to varied and confusing terminology (with many clinicians using synonyms for the same modes or settings), slight variation among brands of ventilators, unfamiliarity, or ceding management to others. The objectives of this chapter are to:

1. Familiarize ED clinicians with common terms in mechanical ventilation.
2. Review key principles of pulmonary physiology, relevant to mechanical ventilation.
3. Discuss the basic principles of selecting ventilator settings.
4. Develop strategies for caring for the ventilated ED patients with acute respiratory distress syndrome (ARDS), asthma, chronic obstructive pulmonary disease (COPD), and traumatic brain injury.
5. Assess and respond to emergencies during mechanical ventilation.

A few words about the style and function of these educational materials are in order. First, the authors assume that the readers are knowledgeable, experienced clinicians who happen to be new to mechanical ventilation. The explanations of ventilation are deliberately simplified in response to other manuscripts and texts, which may at times overcomplicate the subject. Second, the principles herein are deliberately repeated several times throughout the text, working on the educational principle that presenting the same information in different ways enhances understanding and recall. Third, the goal of these materials is to present key concepts. Readers should know that with sophisticated modern ventilators, some may have backup modes or other safeguards that allow for automated switching of modes or other adaptations for patient safety. The details of this complex ventilation function are beyond the scope of this text. However, it is the authors' contention that a thorough understanding of core

principles will allow any emergency clinician to provide evidence-based critical care to their ventilated patients, as well as communicate effectively with their colleagues in critical care and respiratory therapy. As with many aspects of medicine, there are multiple correct ways to present data about mechanical ventilation. In this course, we will use the same method repeatedly to facilitate recall.

For the sake of brevity, this text will not focus on details of clinical management beyond mechanical ventilation, assuming that clinicians are familiar with the medical management of the conditions discussed. Additionally, while interpreting blood gases is essential for providing good care for ventilated patients, a detailed discussion of blood gas analysis is beyond the scope of this text.

Chapter 2

Terminology and Definitions



Ventilator Basics

Control (target) variables are the targets that are set based on the mode of mechanical ventilation chosen. For example, there are *pressure-controlled* and *volume-controlled* modes of ventilation.

Conditional variables are the dependent variable in mechanical ventilation. For example, in volume controlled modes of ventilation, the tidal volume is a set parameter, while the pressure is a conditional variable and can vary from breath to breath.

Trigger The factor that initiates inspiration. A breath can be pressure triggered, flow triggered, or time triggered.

Cycle The determination of the end of inspiration, and the beginning of exhalation. For example, the mechanical ventilator can be volume, pressure, or time cycled.

Physiology Terms

Airway resistance refers to the resistive forces encountered during the mechanical respiratory cycle. The normal airway resistance is ≤ 5 cmH₂O.

Lung compliance refers to the elasticity of the lungs, or the ease with which they stretch and expand to accommodate a change in volume or pressure. Lung with a low compliance, or high elastic recoil, tend to have difficulty with the inhalation process and are colloquially referred to as “stiff” lungs. An example of poor compliance would be a patient with a restrictive lung disease, such as pulmonary fibrosis. In contrast, highly compliant lungs, or ones with a low elastic recoil, tend to have more difficulty in the exhalation process, as seen in obstructive lung diseases.

Derecruitment is the loss of gas exchange surface area due to atelectasis. Derecruitment is one of the most common causes of gradual hypoxemia in intubated patients and can be minimized by increasing PEEP.

Recruitment is the restoration of gas exchange surface area by applying pressure to reopen collapsed or atelectatic areas of lung.

Predicted body weight is the weight that should be used in determining ventilator settings, never actual body weight. Lung volumes are determined largely by sex and height, and therefore, these two factors are used in determining predicted body weight. The formula for men is: $PBW \text{ (kg)} = 50 + 2.3 \text{ (height (in)} - 60)$, and for women is: $PBW \text{ (kg)} = 45.5 + 2.3 \text{ (height (in)} - 60)$.

Phases of Mechanical Breathing

Initiation phase is the start of the mechanical breath, whether triggered by the patient or the machine. With a patient initiated breath, you will notice a slight negative deflection (negative pressure, or sucking) (Fig. 2.1).

Inspiratory phase is the portion of mechanical breathing during which there is a flow of air into the patient's lungs to achieve a maximal pressure, the peak airway pressure (PIP or P_{peak}), and a tidal volume (TV or V_T) (Fig. 2.2).

Plateau phase does not routinely occur in mechanically ventilated breaths but may be checked as an important diagnostic maneuver to assess the plateau pressure (P_{plat}). With cessation of air flow, the plateau pressure and the tidal volume (TV or V_T) are briefly held constant (Fig. 2.3).

FIGURE 2.1 Waveform illustrating initiation phase or triggering

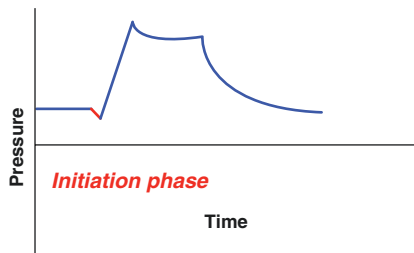


FIGURE 2.2 Waveform illustrating inspiratory phase

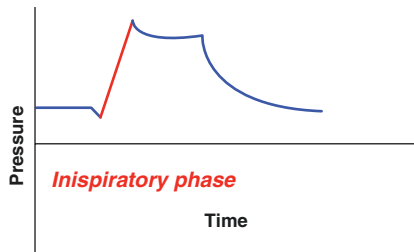


FIGURE 2.3 Waveform illustrating plateau phase

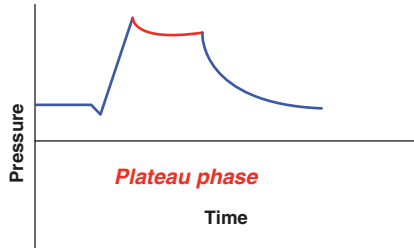
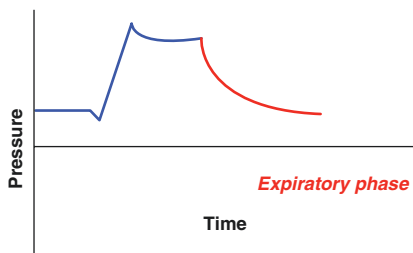


FIGURE 2.4 Waveform illustrating expiratory phase



Exhalation is a passive process in mechanical breathing. The start of the exhalation process can be either volume cycled (when a maximum tidal volume is achieved), time cycled (after a set number of seconds), or flow cycled (after achieving a certain flow rate) (Fig. 2.4).

Ventilator Settings

Peak inspiratory pressure (PIP or Ppeak) is the maximum pressure in the airways at the end of the inspiratory phase. The value is often displayed on the ventilator screen. Since this value is generated during a time of airflow, the PIP is determined by both airway resistance and compliance. By convention, all pressures in mechanical ventilation are reported in “cmH₂O.” It is best to target a PIP <35 cmH₂O.

Plateau pressure (Pplat) is the pressure that remains in the alveoli during the plateau phase, during which there is a cessation of air flow, or with a breath-hold. To calculate this value, the clinician can push the “inspiratory hold” button on the ventilator. The plateau pressure is effectively the pressure at the alveoli with each mechanical breath and reflects the compliance in the airways. To prevent lung injury, the Pplat should be maintained at <30 cmH₂O.

Positive end-expiratory pressure (PEEP) is the positive pressure that remains at the end of exhalation. This additional applied positive pressure helps prevent atelectasis by preventing the end-expiratory alveolar collapse. PEEP is usually set at 5 cmH₂O or greater, as part of the initial ventilator

settings. PEEP set by the clinician is also known as *extrinsic PEEP*, or *ePEEP*, to distinguish it from the pressure that can arise with air trapping. By convention, if not otherwise specified, “PEEP” refers to *ePEEP*.

Intrinsic PEEP (iPEEP), or *auto-PEEP*, is the pressure that remains in the lungs due to incomplete exhalation, as can occur in patients with obstructive lung diseases. This value can be measured by holding the “expiratory pause” or “expiratory hold” button on the mechanical ventilator.

Driving pressure (ΔP) is the term that describes the pressure changes that occur during inspiration, and is equal to the difference between the plateau pressure and PEEP ($P_{\text{plat}} - \text{PEEP}$). For example, a patient with a P_{plat} of 30 cmH₂O and a PEEP of 10 cmH₂O would have a driving pressure of 20 cmH₂O. In other words, 20 cmH₂O would be the pressure that exerted to expand the lungs.

Inspiratory time (iTime) is the time allotted to deliver the set tidal volume (in volume control settings) or set pressure (in pressure control settings).

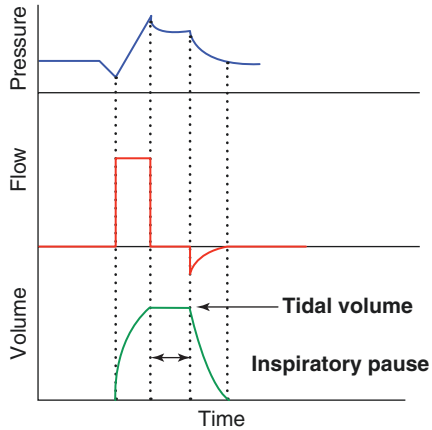
Expiratory time (eTime) is the time allotted to fully exhale the delivered mechanical breath.

I:E ratio, or the inspiratory to expiratory ratio, is usually expressed as 1:2, 1:3, etc. The I:E ratio can be set directly or indirectly on the ventilator by changing the inspiratory time, the inspiratory flow rate, or the respiratory rate. By convention, decreasing the ratio means increasing the expiratory time. For example, 1:3 is a decrease from 1:2, just like 1/3 is less than 1/2.

Peak inspiratory flow is the rate at which the breath is delivered, expressed in L/min. A common rate is 60 L/min. Increasing and decreasing the inspiratory flow is a means of indirectly affecting the I:E ratio. A patient with a respiratory rate set at 20, who is not overbreathing, has 3 s for each complete cycle of breath. If you increase the inspiratory flow, the breath is given faster, and that leaves more time for exhalation. Thus, inspiratory flow indirectly changes the I:E ratio.

Tidal volume (TV or VT) is the volume of gas delivered to the patient with each breath. The tidal volume is best

FIGURE 2.5 Typical ventilator waveforms illustrating volume, flow, and pressure



expressed in both milliliters (ex: 450 mL) and milliliters/kilogram (ex: 6 mL/kg) of predicted body weight, much as one might describe a drug dosage in pediatrics. Clinicians can choose to set the ventilator in a volume control mode, where the tidal volume will be constant for each breath. In pressure control modes, the pressure is constant, but the tidal volume is an independent variable, and will vary slightly with each breath. Regardless, every mode of ventilation delivers a tidal volume. Figure 2.5 illustrates the correlation between the tidal volume, the flow of air, and the pressure waveforms. This is similar to what may be seen on a ventilator screen. For a clinical example of similar waveforms from a patient's ventilator screen, refer to Fig. 6.1.

Respiratory rate (RR or f) is the mandatory number of breaths delivered by the ventilator per minute. However, it is important to be mindful that the patient can breathe over this set rate, and therefore one must report both your set RR and the patient's actual RR; both of these values can be found on the ventilator screen. In addition, it is important to remember that the RR is a key factor in determining time for exhalation. For example, if a patient has a RR of 10 breaths per minute (bpm), he will have 6 s per breath: $((60 \text{ s/min}) / 10 \text{ bpm} = 6 \text{ s/breath})$. A RR of 20 bpm only allows 3 s for the entire respiratory cycle.

Minute ventilation (\dot{V}_E , $V\dot{e}$, or MV) is the ventilation the patient receives in 1 min, calculated as the tidal volume multiplied by the respiratory rate ($TV \times RR$), and expressed in liters per minute (L/min). Most healthy adults have a baseline minute ventilation of 4–6 L/min, but critically ill patients, such as those attempting to compensate for a metabolic acidosis, may require a minute ventilation of 12–15 L/min, or even higher, to meet their demands.

Fraction of inspired oxygen (F_iO_2) is a measure of the oxygen delivered by the ventilator during inspiration, expressed as a percentage. Room air contains 21% oxygen. A mechanical ventilator can deliver varying amounts of oxygen, up to 100%.

Ventilator Modes

Conventional Modes of Ventilation

Assist control (AC) is a commonly used mode of ventilation and one of the safest modes of ventilation in the emergency department. Patients receive the same breath, with the same parameters as set by the clinician, with every breath. They may take additional breaths, or over-breathe, but every breath will deliver the same set parameters. Assist control can be volume-targeted (volume control, AC/VC) where the clinician sets a desired volume, or pressure-targeted (pressure control, AC/PC) where the clinician selects a desired pressure.

Synchronized intermittent mandatory ventilation (SIMV) is a type of intermittent mandatory ventilation, or IMV. The set parameters are similar to those in AC, and the settings can be volume controlled (SIMV-VC) or pressure controlled (SIMV-PC). Similar to AC, each mandatory breath in SIMV will deliver the identical set parameters. However, with additional spontaneous breaths, the patient will only receive pressure support or CPAP. For example, in SIMV-VC, we can set a TV, and as long as the patient is not breathing spontane-

ously, each delivered mechanical breath will achieve this tidal volume. However, spontaneous breaths in this mode of ventilation will have more variable tidal volumes, based on patient and airway factors.

Pressure regulated volume control (PRVC) is a type of assist control that combines the best attributes of volume control and pressure control. The clinician selects a desired tidal volume, and the ventilator gives that tidal volume with each breath, at the lowest possible pressure. If the pressure gets too high and reaches a predefined maximum level, the ventilator will stop the air flow and cycle into the exhalation phase to prevent excessive airway pressure and resulting lung injury. In this mode of ventilation, the pressure target is adjusted based on lung compliance, to help achieve the set tidal volume.

Pressure support is a partial support mode of ventilation in which the patient receives a constant pressure (the PEEP) as well as a supplemental, “supporting” pressure when the ventilator breath is triggered. In this mode, the clinicians can set the PEEP and the additional desired pressure over the PEEP. However, the peak inspiratory airflow, the respiratory rate, and the tidal volume are all dependent variables and determined by the patient’s effort. The patient triggers every breath, and when the patient stops exerting effort, the ventilator stops administering the driving pressure, or the desired pressure over PEEP. Therefore, patients placed on this mode of ventilation must be able to take spontaneous breaths.

Noninvasive positive pressure ventilation (NIPPV) refers to two noninvasive modes of ventilation, in which the patient’s airway is not secured with an endotracheal tube. Rather, these modes of ventilation are delivered through a tight-fitting facemask or nasal prongs. There are several indications, and clear contraindications to these modes of ventilation, please see Noninvasive positive pressure ventilation (NIPPV) in Chap 4. Both CPAP and BPAP are noninvasive modes of ventilation.

Continuous positive airway pressure (CPAP) is a partial support mode of ventilation, in which the patient received a constant airway pressure throughout the respiratory cycle.

The peak inspiratory airflow, respiratory rate, and tidal volume are all dependent variables and determined by the patient's effort. Therefore, the patient must be awake, minimally sedated, and able to take spontaneous breaths during this mode of ventilation.

Bilevel positive airway pressure (BPAP or BiPAP) is a partial support mode of ventilation, in which the patient receives two levels of airway pressure throughout the respiratory cycle. A high *inspiratory pressure (iPAP)* is similar to the peak airway pressure setting. The lower *expiratory pressure (ePAP)*, similar to PEEP, is clinically apparent at the end of expiration and helps to maintain alveolar distention. The patient must be awake, minimally sedated, and able to take spontaneous breaths during this mode of ventilation.

Unconventional modes of ventilation There are other modes of ventilation occasionally used in specific circumstances in ICUs, including airway pressure release ventilation (APRV), also referred to as bi-level or bi-vent, high frequency oscillatory ventilation, proportional assist ventilation (PAV), and neurally adjusted ventilator assist (NAVA), but these modes are not appropriate in the ED without expert consultation.

Suggested Reading

1. Crimi C, Hess D. Principles of mechanical ventilation. In: Bigatello LM, editor. The critical care handbook of the Massachusetts General Hospital. 5th ed. Philadelphia: Lippincott Williams & Wilkins; 2010a.
2. Crimi E, Hess D. Respiratory monitoring. In: Bigatello LM, editor. The critical care handbook of the Massachusetts General Hospital. 5th ed. Philadelphia: Lippincott Williams & Wilkins; 2010b.
3. Singer BD, Corbridge TC. Basic invasive mechanical ventilation. *South Med J*. 2009;102(12):1238–45.
4. Wood S, Winters ME. Care of the intubated emergency department patient. *J Emerg Med*. 2011;40(4):419–27.

Chapter 3

Review of Physiology and Pathophysiology



Gas Exchange

The diagram in Fig. 3.1 represents normal cluster of alveoli with a normal capillary, delivering carbon dioxide (CO_2) and picking up oxygen (O_2).

Figure 3.1 is highly simplified for conceptual emphasis. However, a slightly more detailed diagram illustrating the role of hemoglobin is important to understand the fundamental concepts of gas exchange (Fig. 3.2).

Carbon dioxide travels dissolved in blood, as carbonic anhydrase and as hydrogen and bicarbonate. The components of CO_2 transport are indicated in Fig. 3.2 as green dots in the serum. Approaching the alveolus, the CO_2 easily crosses through the blood, across the capillary wall, and into the alveolus. CO_2 dissolves quite readily, about 20 times faster than oxygen.

Because CO_2 crosses so readily into the alveolus from the serum, ventilation occurs readily.

Conversely, the path for oxygen is less simple (Fig. 3.3). Oxygen is transported largely bound to hemoglobin inside the red blood cells. The hemoglobin in this schematic demonstrates the four binding sites per hemoglobin molecule inside the red blood cells. Oxygen is represented by small blue dots. The concentration of oxygen is high in the alveoli, and it diffuses down the concentration gradient, into the capillary, into the RBC, and binds with Hgb.

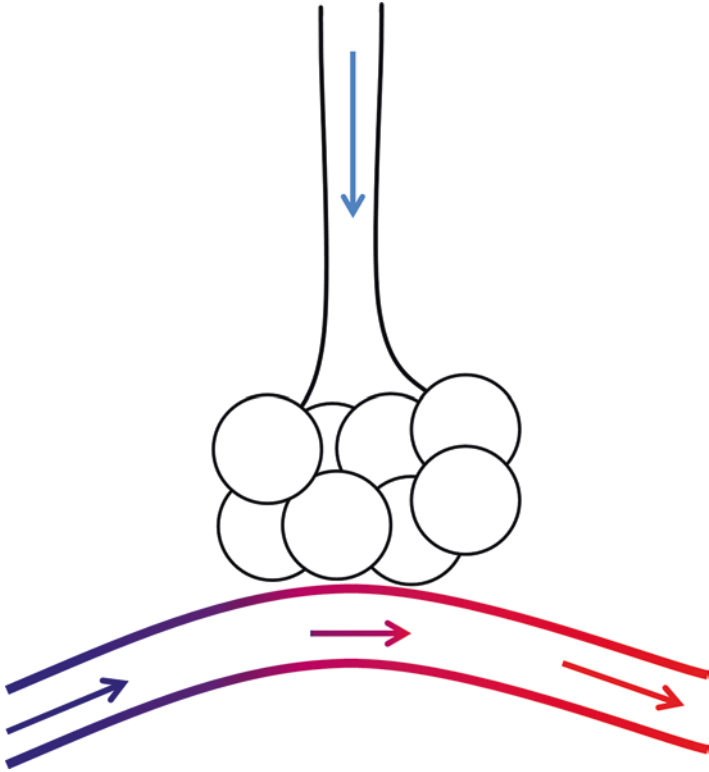


FIGURE 3.1 Schematic of normal alveoli and capillary

While this binding allows for great efficiency in carrying oxygen, oxygen's solubility is much lower, leading to a slower transit time for oxygen to cross the capillary-alveolar interface.

A small amount of oxygen is carried dissolved in the plasma, but compared to the amount bound to hemoglobin, this amount is trivial. The oxygen-carrying capacity of the blood is described by the equation:

$$\begin{aligned} \text{Delivery of Oxygen} = & \text{Cardiac Output} \\ & \times (\text{Hgb} \times 1.39 \times \text{Oxygen Saturation}) \\ & + (\text{PaO}_2 \times 0.003). \end{aligned}$$

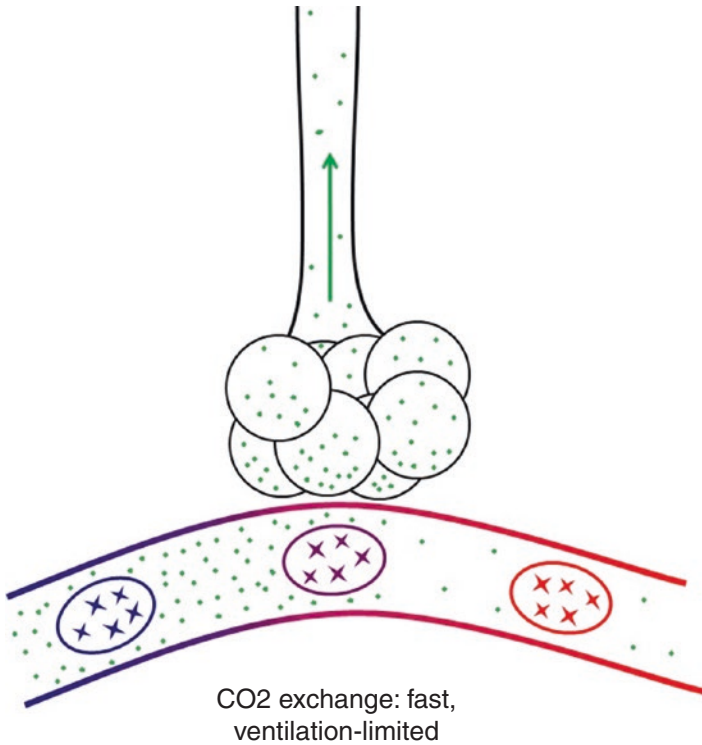


FIGURE 3.2 Carbon dioxide uptake by the alveoli. Green dots = carbon dioxide

This equation intuitively makes sense, as the more Hgb available to carry oxygen, the more oxygen that can be delivered.

Issues with Oxygenation

Hypoxemia

There are five broad physiologic causes of hypoxemia: shunting, V/Q mismatch, alveolar hypoventilation, decreased partial pressure of oxygen, and decreased diffusion. Understanding these mechanisms allows the clinician at the bedside to quickly develop a differential diagnosis for

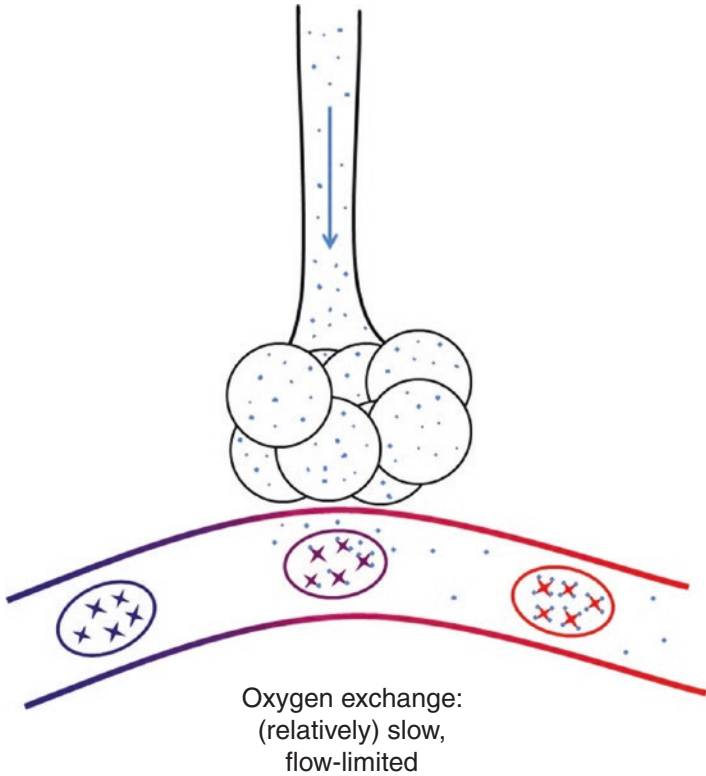


FIGURE 3.3 Oxygen uptake by capillary and hemoglobin. Small blue dots = oxygen

hypoxemia and target diagnostics to assess for the precise etiology. We will review each mechanism in detail.

V/Q mismatch is a broad term that indicates that the ventilation and perfusion of lung units are not optimally aligned. At the two extremes, lung units can have perfusion without ventilation, or shunts, and ventilation without perfusion, or dead space. With commonly encountered clinical insults, such as pneumonia or acute respiratory distress syndrome (ARDS), patients will have both and exhibit a range in-between on a micro-level. It can be helpful to consider them each in more detail, however.

Shunts can also occur on a more macro-level. When an area of the lung is perfused, but not ventilated, such that the inspired oxygen cannot reach the alveoli for gas exchange, that results in an intrapulmonary *shunt*. Examples of shunts are depicted in Figs. 3.4 and 3.5.

There are several different causes of intrapulmonary shunts, including atelectasis, pneumonia, pulmonary edema, acute respiratory distress syndrome (ARDS), hemothorax or pneumothorax, hyperinflation, or auto-PEEPing. All of these pathological processes prevent effective gas exchange at the alveoli. Intrapulmonary shunts can also occur with normal

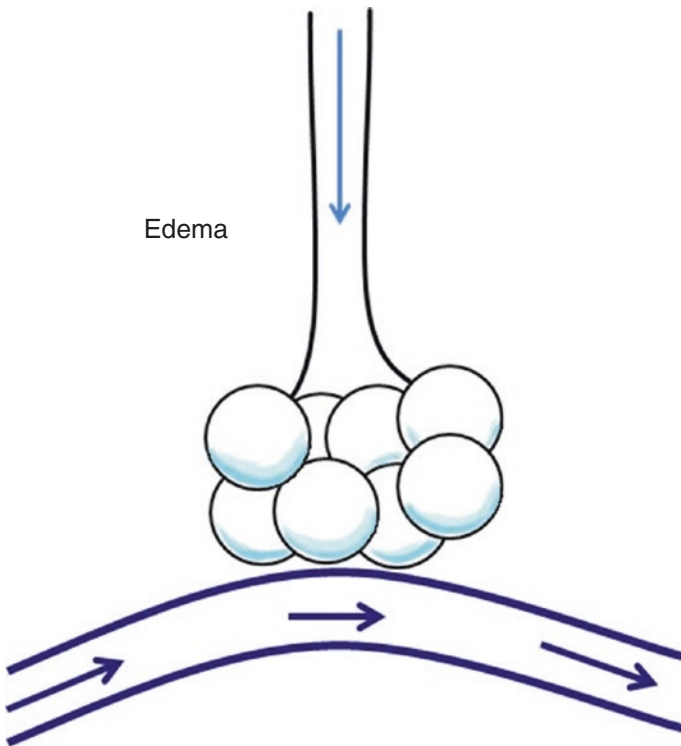


FIGURE 3.4 Fluid-filled alveoli inhibiting gas exchange

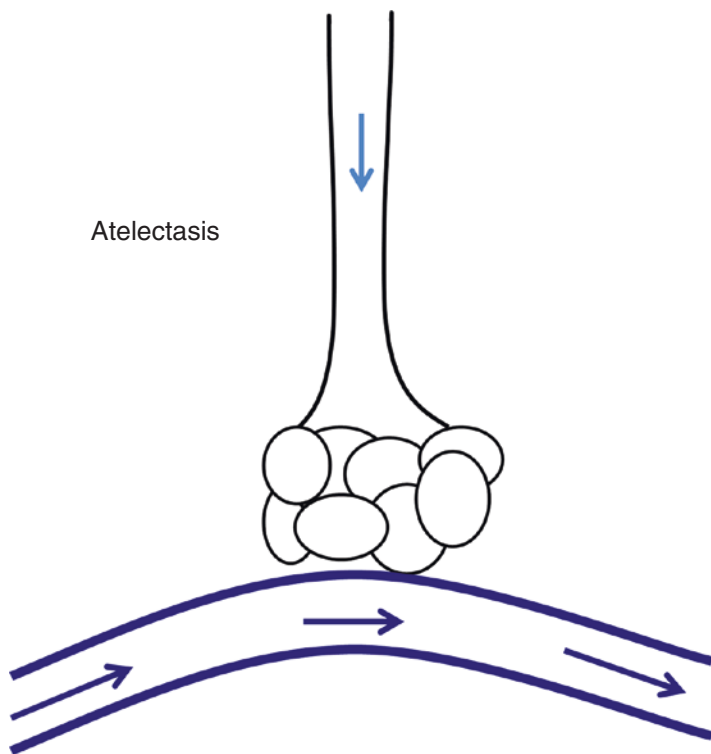


FIGURE 3.5 Collapsed alveoli inhibiting gas exchange

lungs. As an example, in patients with cirrhosis, vasodilation can lead to large volumes of blood bypassing the alveoli with resulting hypoxemia.

Shunt can also occur in the cardiac system, with patent foramen ovals (PFOs) or other congenital or acquired connections between the right and left circulations. At times, the increased stress on the right heart and/or increased intrathoracic pressure from mechanical ventilation may cause a right to left shunt to develop through a previously clinically silent connection, like a PFO (Fig. 3.6).

When an area has ventilation, but no perfusion, this is *dead space* (Fig. 3.7). In other words, the airways are functioning

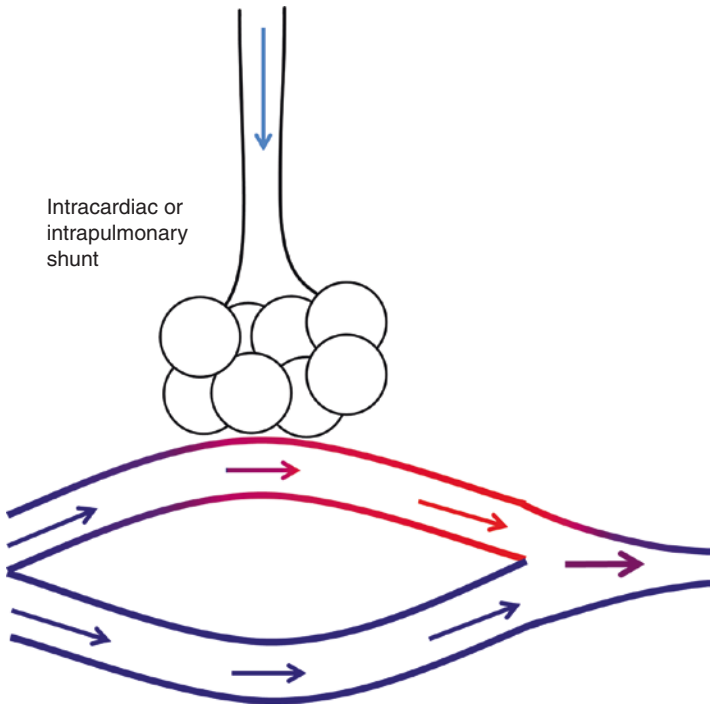


FIGURE 3.6 Shunting can occur at the organ level, with shunts in the heart or lungs

normally, but there is disease process in the vasculature. The best example would be a patient is cardiac arrest who is intubated and ventilated, but there is an interruption of chest compressions. Dead space can be anatomic and physiologic, such as oxygenation but lack of gas exchange that occurs in the upper airways, like the trachea. There can also be pathological causes of dead space, such as this diagram of a pulmonary embolism.

Other examples of dead space include low cardiac output and hyperinflation, as occurs in obstructive lung disease. In diseases such as chronic obstructive pulmonary disease (COPD), there can be a significant level of hyperinflation or auto-PEEP, which can lead to vasoconstriction of the capillaries involved in gas exchange, thereby leading

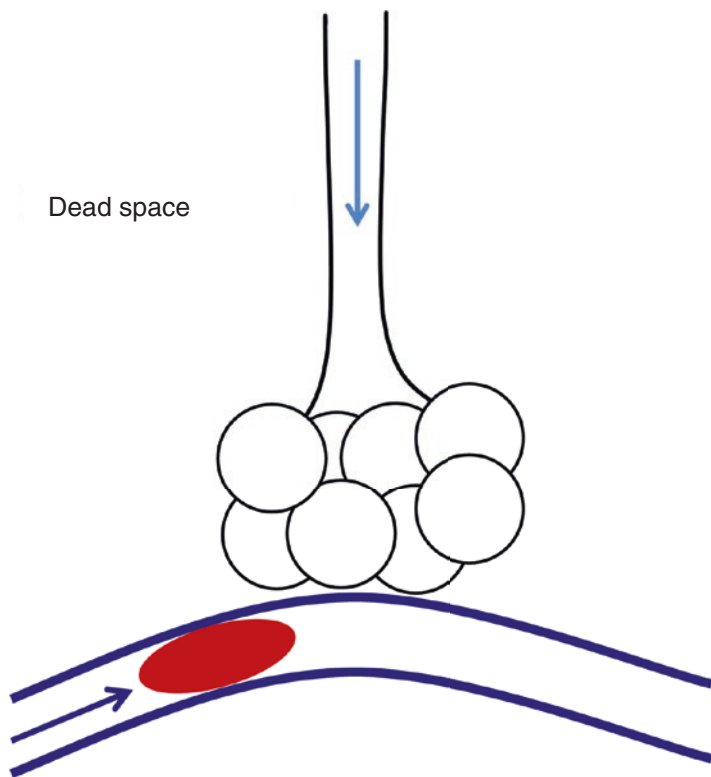


FIGURE 3.7 Decreased perfusion inhibiting gas exchange

to impaired gas exchange. Dead space ventilation can lead to both hypoxia and hypercapnia, due to CO_2 retention. Table 3.1 provides clinical examples of shunts as compared to dead space.

There are several other mechanisms of hypoxemia. The next most common mechanism is alveolar hypoventilation. If a patient is not breathing adequately to facilitate gas exchange, such as with an opioid overdose or splinting due to rib fractures, they can become hypoxemic (Fig. 3.8).

TABLE 3.1 Etiologies of hypoxemia from shunts or dead space

Shunts	Dead space
Atelectasis	Pulmonary embolus
Pneumonia	Low cardiac output
Pulmonary edema	Hyperinflation
ARDS	
Pneumothorax/hemothorax	
Hyperinflation	

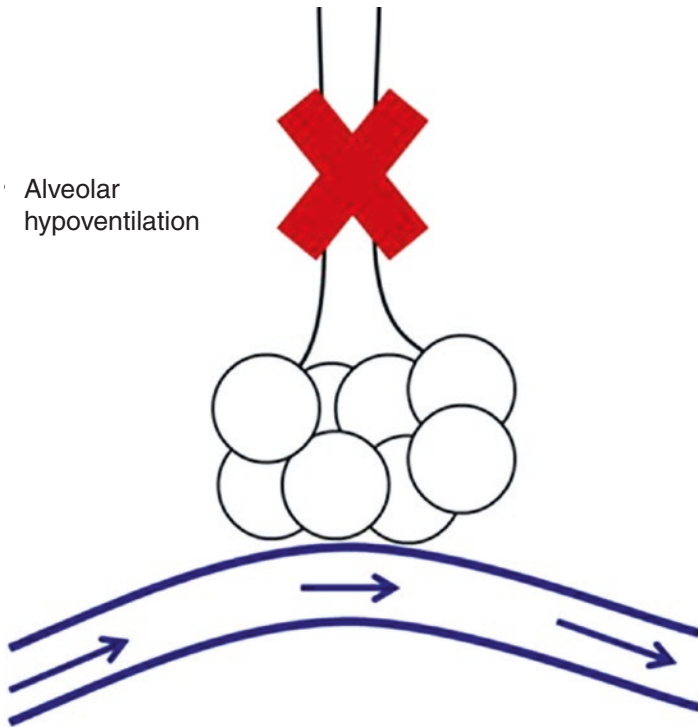


FIGURE 3.8 Decreased airflow to alveoli inhibiting gas exchange

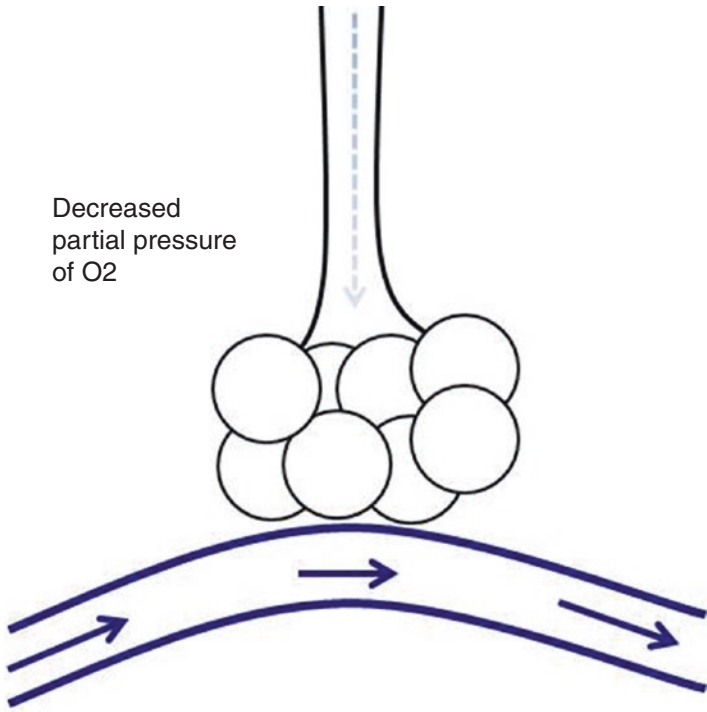


FIGURE 3.9 Decreased partial pressure of oxygen inhibiting oxygenation

Occasionally, patients can demonstrate hypoxemia from a decreased partial pressure of oxygen. While this can occur at altitude, it is less commonly seen in the ED (Fig. 3.9).

Patients may be hypoxemic due to decreased diffusion. Decreased diffusion can occur with increased interstitial thickness, as occurs in interstitial lung diseases (Fig. 3.10), but probably even more commonly, diffusion is decreased due to a loss of surface area, as occurs with emphysema (Fig. 3.11).

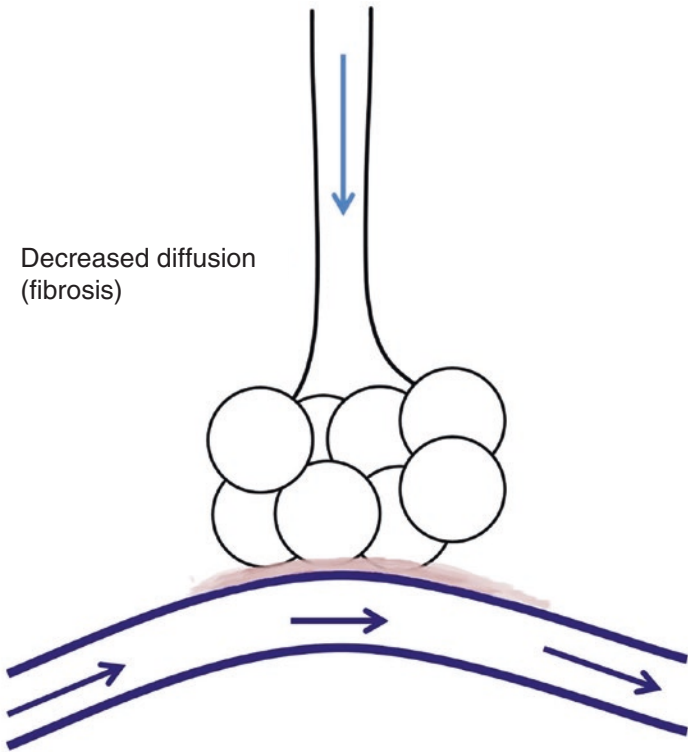


FIGURE 3.10 Increased interstitial thickness inhibiting gas exchange

Hypoxic Vasoconstriction

When an area of the lung is hypoxic, or there is impairment in the oxygen delivery, the lung tries to optimize ventilation and perfusion ration (V/Q matching) by means of *hypoxic vasoconstriction*. In this schematic below, the cluster of alveoli is not receiving oxygen. Therefore, the arterioles leading to the alveoli constrict, diverting blood away from this under-ventilated area, in an effort to improve oxygenation (Fig. 3.12).

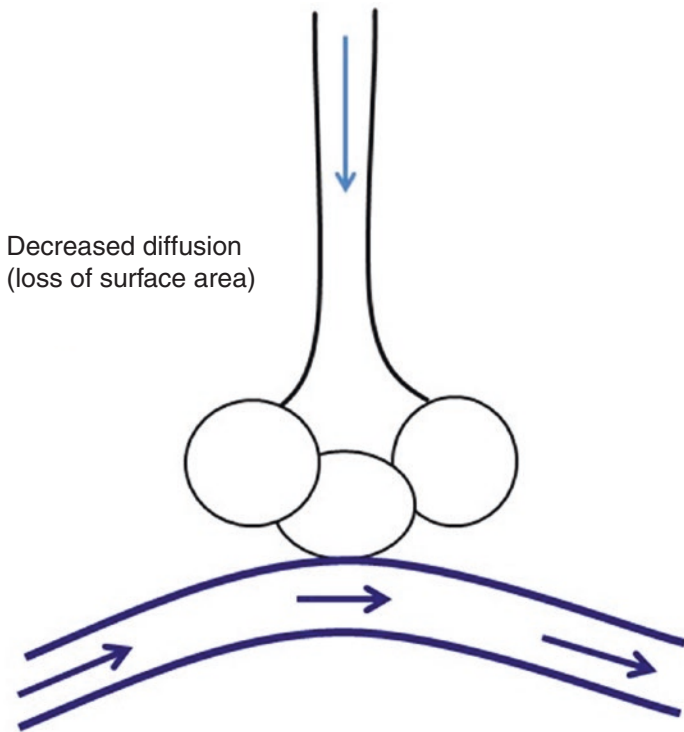


FIGURE 3.11 Loss of surface area inhibiting gas exchange

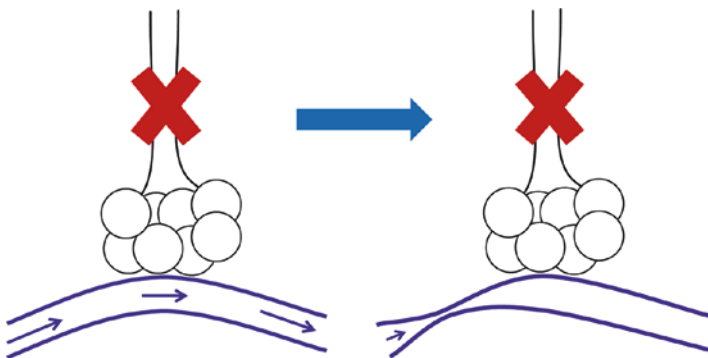


FIGURE 3.12 Hypoxic vasoconstriction leads to decreased perfusion of ineffective lung units

Atelectasis and Derecruitment

Maximizing V/Q matching, by preventing atelectasis, is a key principle in management of respiratory failure. Alveolar derecruitment, or atelectasis, leads to the creation of shunts. Such shunts are created when lying supine to sleep. However, they are compounded by excessive lung weight (such as with pulmonary edema), chest wall weight (as with morbid obesity), abdominal contents and distention (as with small bowel obstructions), and even cardiac compresses (as with pericardial effusion). The addition of sedation and paralysis to positive pressure ventilation can further augment this derecruitment. The diagram in Fig. 3.13 reflects the pressures leading to compression of the lungs when lying a patient supine – the weight of the heart, the weight of the chest wall, the weight of the abdominal contents, and the weight of the lungs themselves.

Issues with Ventilation

Many of the same issues that lead to issues with oxygenation can lead to problems with ventilation, clinically manifested as hypercapnia, as well. Patients in respiratory failure may present with predominantly hypoxemia, predominantly hypercapnia, or both.

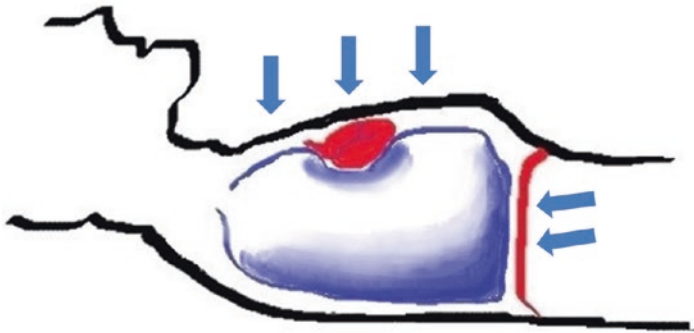


FIGURE 3.13 Collapse of many lung units, or atelectasis on a large scale, is derecruitment

Some of the variability in hypoxemia and hypercapnia arises from the differential transport of oxygen and carbon dioxide as described above. Three of the major etiologies of hypoxemia, dead space, alveolar hypoventilation, and decreased diffusion, lead to hypercapnia. While a patient may have disproportionate hypercapnia, a patient having a completely normal oxygenation with clinically important hypercapnia is unlikely to occur, as oxygen transport is more involved and therefore more susceptible to physiologic derangements.

The arterial alveolar gradient (A-a gradient) is useful to determine if the patient has a combined oxygenation-ventilation problem or simply an oxygenation problem. Although not necessary for many patients who present in the ED with a clear etiology of respiratory failure (an obvious pneumonia, for example), checking an A-a gradient for patients with hypoxemia of uncertain etiology may help narrow the differential diagnosis.

The A-a gradient is the difference between the alveolar pressure of oxygen (PAO_2) and the pressure of the oxygen in the arterial blood (PaO_2). This measurement requires an ABG.

The PAO_2 is calculated using the alveolar gas equation, or:

$$PAO_2 = PiO_2 - PaCO_2 / 0.8$$

Where the PiO_2 is the pressure of the inspired oxygen.

A normal A-a gradient is <15 mmHg for most patients (Table 3.2).

TABLE 3.2 Normal and increased A-a gradients

Normal A-a gradient	Increased A-a gradient
Low partial pressure O_2	V/Q mismatch
Alveolar hypoventilation	Cardiac or pulmonary shunt
	Decreased diffusion

Compliance and Resistance

Two other important physiologic concepts to review are *compliance* and *resistance*.

Resistance is the impedance of flow in the tubing and airways and therefore can only occur when there is airflow. According to Ohm's law:

$$\begin{aligned} \text{Resistance } (R) &= \Delta \text{ pressure} / \Delta \text{ volume} \\ R &= (\text{Peak inspiratory pressure} - \text{Plateau pressure}) / \text{Tidal volume} \\ R &= (\text{PIP} - \text{Pplat}) / (\text{TV}) \end{aligned}$$

Assuming a constant tidal volume, the resistance equation can be simplified to:

$$R \approx (\text{PIP} - \text{Pplat})$$

Normal airway resistance should be ≤ 5 cmH₂O. Resistance is a factor in ventilating all patients but can become particularly important when ventilating patients with COPD or asthma. The resistance in a system increases with decreasing diameter. While common examples include a very small endotracheal tube (ETT) or bronchospasm leading to narrowing of the airways, recall that a "decrease in the diameter" can also occur at just one point, such as with kinking or biting of the ETT, or a mucous plug in a large airway.

Compliance refers to the distensibility of the system and is the inverse of elastance. In other words, it is a measure of the lung's ability to stretch and expand. The more elastic a system, or higher the "recoil," the lower the compliance. A common analogy to understand the concepts of elastance is to analyze the recoil of springs. Imagine a very tightly wound and stiff spring. This spring is difficult to stretch and wants to stay in the coiled position. This spring would have

a high elastance and a low compliance. Envision a second, loosely coiled spring. Very little force is required to stretch out this spring, and therefore, it has low elastance but high compliance.

Although compliance commonly is used to describe the lung parenchyma, remember that compliance actually involves all components of the system. In other words, a patient with pulmonary edema may have low compliance due to an issue with the lung parenchyma, but another patient may have a similarly low compliance due to severe chest wall stiffness after a third-degree burn. Clinically, knowing the exact cause of decreased compliance in a given patient can be challenging. Physicians should not, therefore, always assume that it is always related to “stiff lungs.”

In the schematic shown in Fig. 3.14, the top “lungs” are healthy. The lungs on the left have a resistance problem or impairment in air flow. The lungs on the right have a compliance problem or impairment in stretch and recoil. In this diagram, both figures could have elevated peak inspiratory pressures (PIP), due to the excess pressure generated in the

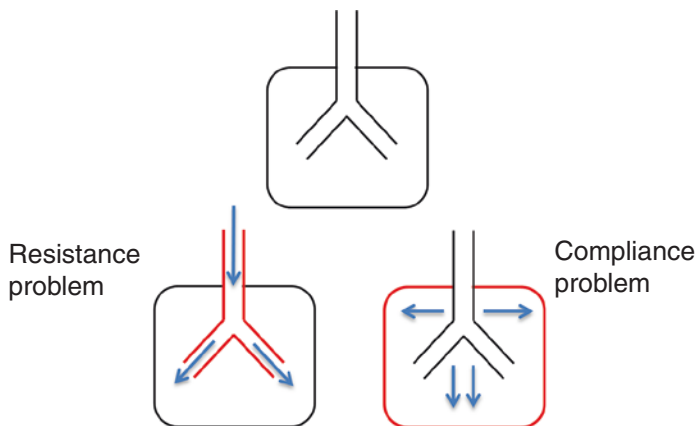


FIGURE 3.14 Resistance to flow in airways vs. decreased distensibility of the entire respiratory system

system. However, only the right-hand side figure would have an elevated plateau pressure (Pplat), since this process occurs when there is absence of air flow.

$$\text{Compliance}(C) = \Delta \text{volume} / \Delta \text{pressure}$$

$$C = \text{Tidal volume} / \text{Plateau pressure} - \text{Peak inspiratory pressure}$$

$$C = (\text{TV}) / (\text{Pplat} - \text{PEEP})$$

Therefore, when trouble shooting high pressure ventilator alarms, two values are needed. The peak airway pressure (PIP) should be displayed on the ventilator screen, while the plateau pressure (Pplat) is obtained by holding the “inspiratory hold” or “inspiratory pause” button on the ventilator. An elevated PIP and normal Pplat is indicative of increased airway resistance. An elevated PIP and elevated Pplat is indicative of an abnormal compliance. Determining whether the patient has a resistance problem or a compliance problem can assist in the differential diagnosis of respiratory failure in the ED, as outlined in Table 3.3.

Atelectasis, or collapse of alveoli and decruitment, is another key physiologic concept in mechanical ventilation. Atelectasis has multiple detrimental effects in ventilated patients. First, atelectasis decreases the surface area for gas exchange. Atelectasis also worsens compliance. Consider blowing up a small party balloon. To start to open the balloon, a large amount of pressure is required. Once the balloon starts to inflate, blowing it up further is easy, until it reaches the point of overdistention. Atelectasis leads to shunts and can cause impaired oxygenation.

Air trapping, also referred to as *breath-stacking*, can lead to the development of *auto-PEEP*, or *intrinsic PEEP* (*iPEEP*). These pressures should be differentiated from applied PEEP, or *extrinsic PEEP* (*ePEEP*). *ePEEP* refers to the additional end-expiratory positive pressure set during mechanical ventilation to prevent alveolar collapse and recruitment. In contrast, *auto-PEEP*, or *iPEEP*, is a pathophysiological process that can occur when the ventilator initiates the next breath

TABLE 3.3 Characteristics of high resistance and abnormal compliance

High resistance	Abnormal compliance
High PIP, Low/Normal P_{plat}	High PIP, High P_{plat}
Kinked/obstructed ETT	Mainstem intubation
Mucus plugging	Atelectasis
Bronchospasm	Pulmonary edema
ETT too narrow (small)	ARDS
Coughing	Hemo/pneumothorax
Obstructive lung disease	Pneumonia
	Bronchospasm (obstructive lung disease)
	Pulmonary fibrosis (restrictive lung disease)
	Obesity
	Abdominal compartment syndrome
	Circumferential burns of the chest
	Scoliosis
	Supine position

prior to complete exhalation. While this is most common in patients with prolonged expiratory phases, such as asthma or COPD, it can also occur in patients who have a fast respiratory rate or those who are being ventilated with large tidal volumes. The amount of auto-PEEP can be measured by pressing the “expiratory hold” or “expiratory pause” button on the ventilator. When this button is pressed, the ventilator will display the total PEEP. The auto-PEEP is the difference between the total PEEP and the set PEEP.

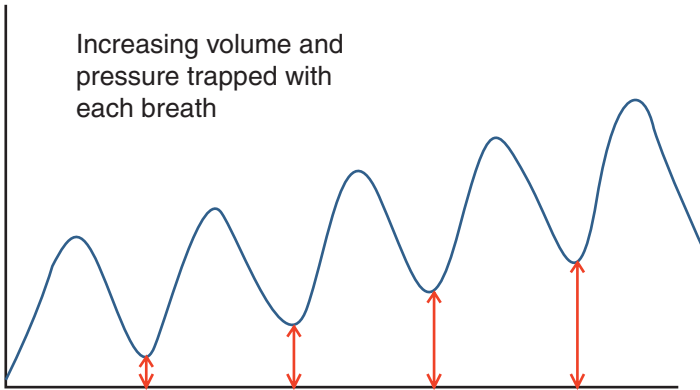


FIGURE 3.15 Conceptual illustration of air trapping

$$\text{Auto - PEEP (iPEEP)} = \text{Total PEEP} - \text{ePEEP}$$

The schematic in Fig. 3.15 represents the effects of air trapping. Please note that this diagram is for illustration purposes only and does not represent the expected tracings on actual ventilator screens.

Air trapping, or autoPEEP, can lead to significant adverse cardiopulmonary effects. The increased intrathoracic pressure from autoPEEP can decrease venous return and lead to hemodynamic instability, even cardiac arrest in severe cases. The increased pressures may also result in a pneumothorax or pneumomediastinum. Additionally, air trapping can lead to ineffective ventilation due to collapse of the capillaries responsible for gas exchange, with worsening hypercarbia and hypoxemia. While this may seem like a paradox, as one may assume that increasing the minute ventilation, or moving more air, will improve ventilation, there is a limit to the beneficial effects. Once the lungs are overdistended, gas exchange is ineffective. In these circumstances, allowing the patient sufficient time to exhale can decrease CO_2 retention.

Suggested Reading

1. Dantzker DR. Physiology and pathophysiology of pulmonary gas exchange. *Hosp Pract.* 1986;21(1):135–9.
2. Henig NR, Pierson DJ. Mechanisms of hypoxemia. *Respir Care Clin N Am.* 2000;6(4):501–21.
3. Levy BD, Kitch B, Fanta CH. Medical and ventilatory management of status asthmaticus. *Intensive Care Med.* 1998;24(2):105–17.
4. West JB. New advances in pulmonary gas exchange. *Anesth Analg.* 1975;54(4):409–18.

Chapter 4

Noninvasive Respiratory Support



The clinician should first assess whether the patient has an oxygenation problem or a ventilation problem. Many patients will have both simultaneously. Determining which problem needs addressing will help determine the next appropriate steps in support. Please note that patients with airway compromise, profoundly altered mental status, or severe shock should, in most circumstances, be intubated rather than maintained with noninvasive means.

Oxygen Support

Many patients who present with hypoxemia can be well-supported by supplemental oxygen. Patients should be given only the minimal support they need to maintain their desired oxygen level, as hyperoxia, or too much oxygen, is increasingly appreciated as a risk factor for poor outcomes [1].

High Flow Nasal Cannula

High flow nasal cannula (HFNC) is an excellent means of supporting hypoxemic patients [2]. As illustrated in Fig. 4.1, a typical nasal cannula can provide up to 6 L/min of supplemental oxygen. Each additional L/min provides about % extra oxygen. HFNC, conversely, can provide about 45–60 L/

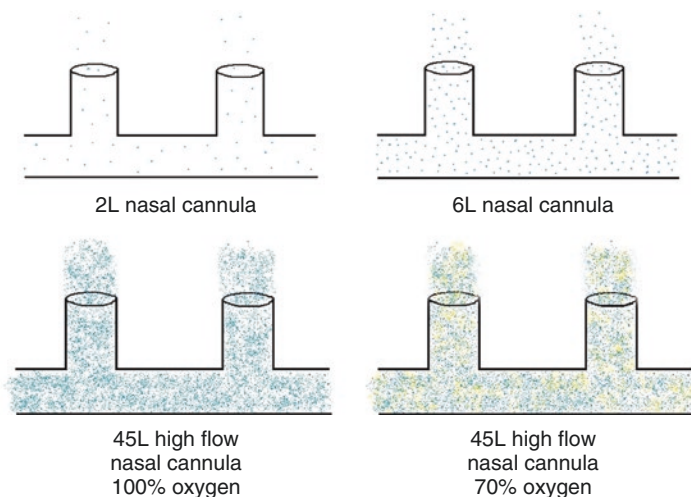


FIGURE 4.1 In this illustration, the blue dots represent theoretical oxygen delivery. A small amount of oxygen is delivered and mixed with ambient air in the typical nasal cannula, in the top two figures. The bottom two depict HFNC, showing the increased flow as well as the ability to blend oxygen and air at desired concentrations

min, depending upon the variations of the setup. Whereas the typical nasal cannula provide additional oxygen blended with the ambient air, the HFNC has a blender attached to the apparatus. This means that HFNC has two components, the L/min delivered, as well as the percentage of oxygen delivered. Figure 4.1 illustrates these different mechanisms for oxygen delivery, as well as the differences between flow and percent oxygen.

HFNC not only provides the option of delivering a high concentration of oxygen (90–100%), but it provides a small level of positive pressure, given the high flows. This positive pressure and associated CO_2 washout also seem to be helpful to some degree in hypercapnic respiratory failure, making HFNC an excellent initial option for respiratory support. Table 4.1 lists contraindications for HFNC.

TABLE 4.1 Contraindications to HFNC

Airway compromise
Facial trauma
Other indication for intubation
Altered mental status
Severe shock
Primarily hypercapnic respiratory failure

Noninvasive Positive Pressure Ventilation

Noninvasive positive pressure ventilation (NIPPV) is one of the most important advances in emergency and critical care of patients in respiratory failure. Numerous studies have demonstrated improved outcomes for patients with respiratory failure from COPD and congestive heart failure (CHF) using noninvasive ventilation [3–5].

As opposed to invasive ventilation after placement of an endotracheal tube (ETT), NIPPV is delivered via a tight-fitting face mask or nasal prongs. There are several indications for NIPPV, as it is a terrific method to oxygenate and ventilate many patients. However, there are a few key contraindications. Patients must be awake and able to protect their airway, as this is not a definitive airway. If the patient is too obtunded to remove the mask, should they vomit or have any other threat to the airway, they should not be placed on NIPPV. Additionally, nausea and vomiting are contraindications, due to the risk of aspiration. Facial trauma, precluding the tight-fitting mask, is a contraindication, as is a recent GI surgery (such as a partial gastrectomy) that would not tolerate pressure on suture lines. These contraindications are outlined in Table 4.2.

There are two forms of NIPPV: continuous positive pressure ventilation and bilevel positive airway pressure.

TABLE 4.2 Contraindications to noninvasive ventilation

Obtundation, inability to remove mask
GI pathology with vomiting or high risk for vomiting
Recent ENT or GI surgery
Airway compromise
Facial trauma
Other indication for intubation
Altered mental status
Severe shock
Severe hypoxemic respiratory failure

Continuous positive pressure ventilation (CPAP) is a continuous positive pressure that is delivered throughout the respiratory cycle, and along with the FiO_2 , assists with oxygenation by recruiting alveoli, preventing alveolar collapse, and decreasing the work of breathing. In function, CPAP is analogous to positive end-expiratory pressure (PEEP) for an intubated patient. The difference between CPAP and PEEP is one of nomenclature, as the PEEP is only measurable at the end of expiration.

In patients with congestive heart failure (CHF), CPAP can increase the intrathoracic pressure to decrease the venous return and therefore reduce lung congestion. In addition, this positive pressure can also decrease the afterload on the left ventricle, leading to increased stroke volume and cardiac output. CPAP is primarily used in the treatment algorithm of patients with hypoxemic respiratory failure, or those who need the additional positive pressure to assist with alveolar recruitment.

Bilevel positive airway pressure (BPAP or BiPAP) is another mode of NIPPV, which provides two different levels of pressure throughout the respiratory cycle. The high pressure, or

the inspiratory peak airway pressure (IPAP), is analogous to the PIP of invasive ventilation. A second lower pressure, the expiratory peak airway pressure (EPAP), is similar to the CPAP described above or PEEP applied in invasive mechanical ventilation. Providing these pressures, in addition to the FiO_2 , assists in improving the patient's oxygenation. The difference between the IPAP and EPAP serves as the driving pressure and assists with ventilation. In contrast to CPAP, which is beneficial in hypoxemia, BPAP is useful in patients with hypoxemic and hypercapnic respiratory failure. Figure 4.2 illustrates a typical BPAP ventilator screen.

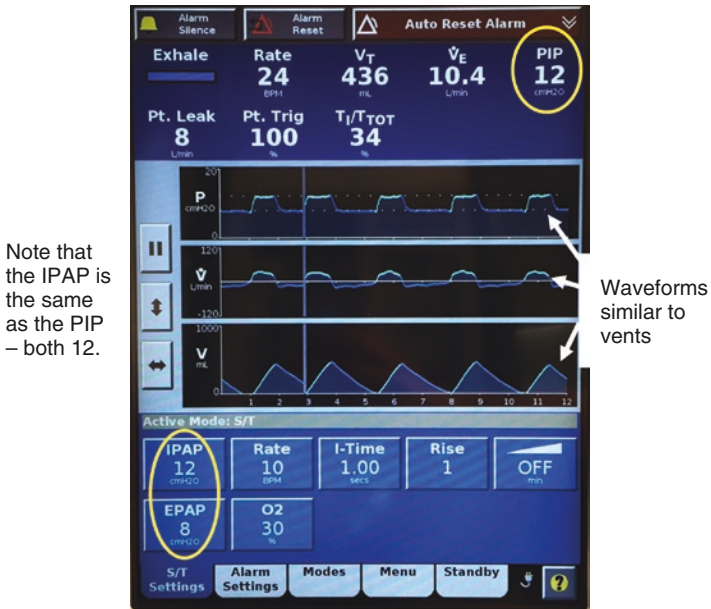


FIGURE 4.2 Typical screen for BPAP, highlighting the IPAP, EPAP, and the peak inspiratory pressure, the PIP. By convention, with non-invasive ventilation, the IPAP and the PIP are the same. The waveforms are similar to those of invasive mechanical ventilation. Please refer to Fig. 2.5 and 6.1 for additional examples

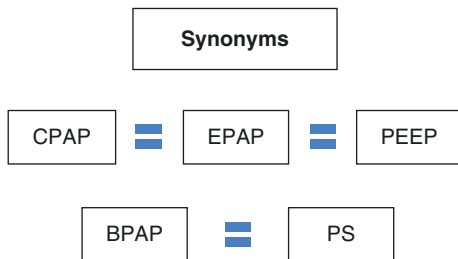


FIGURE 4.3 Although several terms are used for the same principles, the concepts are simple. Continuous positive airway pressure (CPAP), expiratory positive airway pressure (EPAP), and positive end expiratory pressure (PEEP) all refer to a baseline positive pressure, over which the patient breathes. Bilevel positive airway pressure (BPAP) and pressure support (PS) are both modes of ventilation in which a patient receives an additional pressure over the baseline pressure to support their ventilation. By convention, BPAP refers to this pressure being provided via a mask, and PS refers to this pressure being provided via an endotracheal tube

Where bilevel positive airway pressure differs from CPAP is that once a patient triggers a breath, the machine will provide additional, supportive pressure, or the inspiratory positive airway pressure (IPAP.) By assisting the patients with IPAP, BPAP is a great tool for patients with poor ventilation, such as COPD patients. The clinician can set both the IPAP and the EPAP with BPAP, based upon the patient's needs. In this way, BPAP is very similar to pressure support, discussed in detail in Chap. 5. Figure 4.3 demonstrates the multiple synonyms that represent the same concepts.

References

1. Page D, Ablordeppey E, Wessman BT, Mohr NM, Trzeciak S, Kollef MH, Roberts BW, Fuller BM. Emergency department hyperoxia is associated with increased mortality in mechanically ventilated patients: a cohort study. *Crit Care*. 2018;22(1):9.
2. Frat JP, Thille AW, Mercat A, Girault C, Ragot S, Perbet S, et al; FLORALI Study Group; REVA Network. High-flow oxygen

- through nasal cannula in acute hypoxemic respiratory failure. *N Engl J Med.* 2015;372(23):2185–96.
3. Rose L, Gerdtz MF. Review of non-invasive ventilation in the emergency department: Clinical considerations and management priorities. *J Clin Nurs.* 2009;18:3216–24.
 4. Cabrini L, Landoni G, Oriani A, et al. Noninvasive ventilation and survival in acute care settings: A comprehensive systematic review and meta-analysis of randomized controlled trials. *Crit Care Med.* 2015;43:880–8.
 5. Archambault PM, t-Onge M. Invasive and noninvasive ventilation in the emergency department. *Emerg Med Clin North Am.* 2012;30(2):421–49. ix.

Chapter 5

Modes of Invasive Mechanical Ventilation



Modes of Invasive Ventilation

As already illustrated, the terminology used for mechanical ventilation can be confusing, as many clinicians use various terms for the same settings. The “*mode*” of ventilation simply refers to the way the ventilator is set to interact with the patient. A key distinguishing factor among modes is whether the patient can alter the breath they receive, or whether the ventilator will administer the same breath each time, regardless of the patient’s effort.

Assist control (AC) is one of the most commonly used modes of ventilation. AC can be set to target (control) either a pressure or a volume, as described in further detail below. In assist control, the clinician the independent variable (tidal volume or pressure), the respiratory rate, and the FiO₂. If the patient exerts no respiratory effort, she/he will receive an identical breath each time. Also, if the patient starts to breathe, or “triggers” the ventilator, the ventilator will give that identical breath. This allows the patient to “overbreathe” but she/he cannot alter the other clinician-set properties of the breath. For example, if a patient is set to receive 400 mL per breath in AC volume-controlled ventilation, at a flow of 60 L/min, with a respiratory rate of 12 breaths per minute, this is what the patient will receive if they make no efforts to breathe. If the patient is then less sedated and starts to make

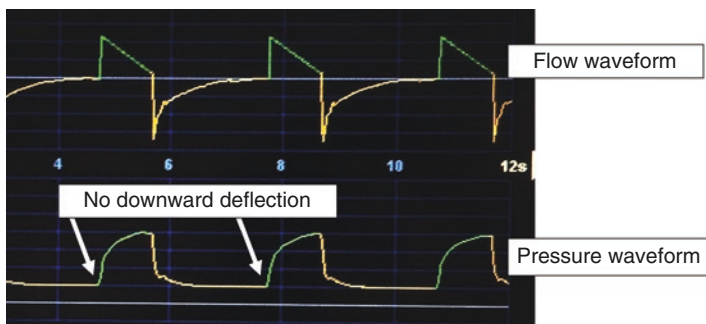


FIGURE 5.1 Illustration of typical volume control waveforms for pressure and flow

respiratory efforts, he or she can increase the respiratory rate, and each breath will still have approximately 400 mL delivered at a rate of 60 mL/min.

In Fig. 5.1, the flow curve is on the top line, and the pressure curve on the bottom line. Note that every waveform is identical. Also note that there is no downward deflection at the initiation of each breath, indicating that these are machine-triggered breaths.

As we continue to review ventilator screen shots, it is important to start recognizing patterns, as the placement of the volume, flow, and pressure curves can vary based on clinician preference and do not reflect patient physiology.

Synchronized intermittent mandatory ventilation (SIMV) involves components of both AC and PS. A respiratory rate is set in SIMV, but it is usually a low rate, such as 8–10 breaths per minute. The patient will receive those “mandatory” breaths, and they will receive the set breath parameters, with a set volume or pressure, rate, and flow or inspiratory time, as determined by the clinician, just as in AC. However, in between those mandatory breaths, the patient can take additional spontaneous breathe with pressure support, allowing them to vary their breathing pattern. This mode was previously used as a weaning mode, but studies have shown that it offers no benefit over other modes.

Pressure support (PS or PSV) is a partially supported, or spontaneous, pressure-controlled mode of ventilation. In this mode, there is no set respiratory rate or tidal volume, and the patient must be awake enough to trigger each breath. The patient receives a set baseline pressure, the PEEP, and, with the triggering of the breath, an additional, supportive pressure above that baseline to help overcome the resistive airway forces and decrease the work of breathing. The clinician sets the PEEP and the supporting pressure.

The other significant difference is that in pressure support, the ventilator can sense when the patient stops exerting effort for the breath. Once the flow drops to a preset limit (usually 25%), the ventilator stops providing the additional pressure support for that breath. In this way, the patient has more control over the breathing pattern.

Figure 5.2 is a ventilator screen shot of a patient breathing with PSV. Note the downward deflection at the initiation of each breath, indicating that the patient triggered the breath. Also note that in contrast to the last diagram of a patient breathing on AC ventilation, the patient on PSV generated flow waveforms that have subtle variety in shape, size, and rhythm, because the patient determines each breath. However, the pressure waveforms on the top line are constant across these five breaths, because the ventilator is providing that maximum pressure, as dialed in by the clinician. Lastly, note

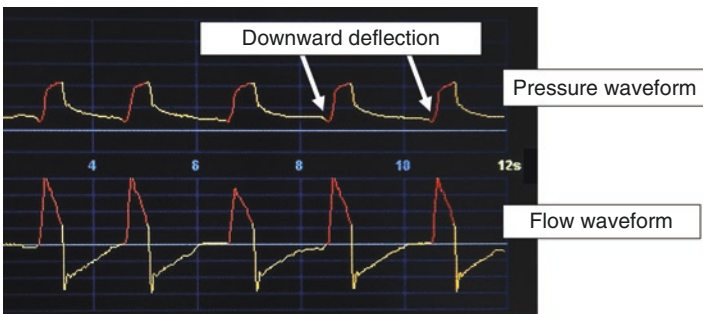


FIGURE 5.2 Illustration of typical pressure support waveforms

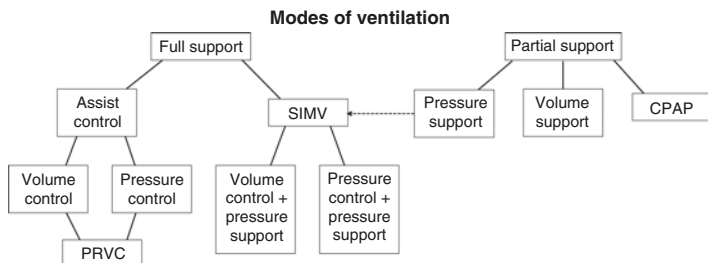


FIGURE 5.3 Relationship among the commonly used modes of mechanical ventilation. Note that SIMV usually incorporates aspects of both assist control ventilation and pressure support ventilation. Pressure regulated volume control is a volume-targeted mode that has a maximum pressure allowed to reach that volume

that in this figure, the pressure waveform is now on top and the flow is on the bottom. Again, this is a matter of preference, and reflects nothing about the patient’s physiology.

Figure 5.3 demonstrates the relationship among the commonly used modes of ventilation, separating them as full support or partial support modes.

The modes used most commonly will vary hospital to hospital. By and large, as long as the patient is getting the level of support appropriate for their condition (a critically ill patient with severe respiratory failure, requiring heavy sedation receiving full support, or a patient intubated for airway swelling only requiring partial support, as examples), the mode has not really been shown to make a significant difference in outcome [1].

Each mode, assist control, SIMV, or partial support modes, can be set to be volume-targeted (such as volume control, or VC) or pressure-targeted (pressure control, PC). When the volume is set (“volume control” or “volume targeted” ventilation), the patient’s resistance and compliance will determine the pressures. When the pressure is set (“pressure control” or “pressure targeted ventilation”), the resistance and compliance will determine the volume.

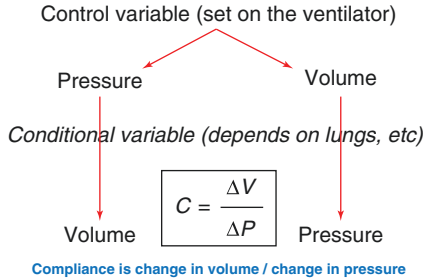


FIGURE 5.4 Compliance is the relationship between the change in pressure and the change in volume. For any ventilator setting, the clinician can only set the pressure or the volume. The compliance of the respiratory system will determine the other value

Understanding this relationship is important for clinicians to monitor the ventilated patient. This relationship is illustrated in Fig. 5.4.

Beyond the mode, clinicians should understand other basic ventilator settings and their relationships. The following examples illustrate the settings of the ventilator.

In volume control AC (AC/VC), the provider sets a predetermined tidal volume (e.g., 500 mL), flow rate (e.g., 60 mL/min), and respiratory rate (e.g., 12 breaths/min). In this mode of ventilation, the inspiratory to expiratory (I:E) ratio is indirectly determined by the RR and flow rate, as demonstrated below, since this mode of ventilation is not determined by a set time, otherwise known as “time-cycled.”

For the VC settings:

- TV = 500 mL
- Flow = 60 L/min = 1 L/s
- RR = 20 breaths/min

The resulting calculations demonstrate the I:E ratio:

$$\text{Total cycle time (TCT)} = (60 \text{ s/min}) / (20 \text{ breaths/min}) = 3 \text{ s/breath cycle}$$

$$\begin{aligned} \text{Inspiratory time (iTime)} &= (500 \text{ mL}) / (1 \text{ L/s}) = 0.5 \text{ s} \\ \text{Expiratory time (eTime)} &= \text{TCT} - \text{iTime} = 3 \text{ s} - 0.5 \text{ s} = 2.5 \text{ s} \\ \text{I:E ratio} &= 1:5 \end{aligned}$$

In contrast, in pressure control AC (AC/PC), the ventilator is set to give the desired pressure for a set time. For example, the clinician can set the ventilator for peak pressure such as 15 cmH₂O, and the inspiratory time, such as 1 s. Therefore, one can set the I:E ratio directly, since PC is time-cycled or, in other words, gives the selected pressure for a set time.

For the PC settings:

$$\begin{aligned} \text{Set pressure} &= 15 \text{ cmH}_2\text{O} \\ \text{RR} &= 20 \text{ breaths/min} \\ \text{Inspiratory time} &= 0.5 \text{ s} \end{aligned}$$

The resulting calculation demonstrates the I:E ratio

$$\begin{aligned} \text{Total cycle time (TCT)} &= (60 \text{ s/min}) / (20 \text{ breaths/min}) = 3 \text{ s} \\ &\text{breath cycle} \\ \text{Inspiratory time (iTime)} &= 0.5 \text{ s} \\ \text{Expiratory time (eTime)} &= \text{TCT} - \text{iTime} = 3 \text{ s} - 0.5 \text{ s} = 2.5 \text{ s} \\ \text{I:E ratio} &= 1:5 \end{aligned}$$

Pressure regulated volume control, or PRVC, is another mode of mechanical ventilation that blends the best aspects of both volume and pressure targeted ventilation. It is an assist-control (AC) mode that is largely volume targeted, in that the clinician selects a desired tidal volume. However, the ventilator strives to administer the tidal volume at the lowest possible pressure, based on the peak pressure limit set by the clinician. If the peak inspiratory pressure reaches the limit set by the clinician, the ventilator will then cycle to expiration phase to protect the lungs from barotrauma before the set tidal volume is achieved. The clinician will then be alerted to the high pressures, allowing for an intervention to assist in reaching the desired tidal volume.

Pressures on the Ventilator

Modern mechanical ventilators all deliver positive pressure ventilation, as opposed to the negative pressure ventilation used in normal physiologic breathing. This pressure, which allows for both oxygenation and ventilation, can be potentially detrimental to the patient in excess. Therefore, the goal is to use the minimum pressure required to oxygenate and ventilate adequately, while minimizing the risks of barotrauma and volutrauma.

The *peak inspiratory pressure (PIP)* represents pressures in the entire airway system and is a measure of both the resistance and compliance. The PIP is displayed on the vent screen with each breath.

The *plateau pressure (Pplat)*, which is measured when there is an absence of airflow during the plateau phase of mechanical breathing, is the reflection of the pressure delivered to the alveoli and the compliance of the system. Therefore, to prevent alveolar injury, the Pplat should be maintained <30 cmH₂O. The Pplat is not directly displayed on the ventilator but can be calculated by pressing the inspiratory pause button, briefly allowing all pressures to equilibrate when there is an absence of airflow. The machine will then display this calculated value.

In Fig. 5.5, the pressure waveform is on the top, and the flow is on the bottom. The PIP is a little over 50 cmH₂O, looking at the scale to the left of the screen. The Pplat is 38 cmH₂O, as noted on the scale on the left in the first breath on this image, as well as calculated value noted on the upper right-hand corner of the ventilator, after pressing the inspiratory pause button. This indicates that there is a compliance problem. The difference between the PIP and Pplat is greater than 5 cmH₂O, indicating there is also a resistance problem. This tracing was taken from the ventilator of a patient with end-stage COPD who developed pneumonia.



FIGURE 5.5 Ventilator screen showing the relationship between the peak inspiratory pressure (PIP) and the plateau pressure (Pplat.) The Pplat is only seen with an inspiratory hold maneuver

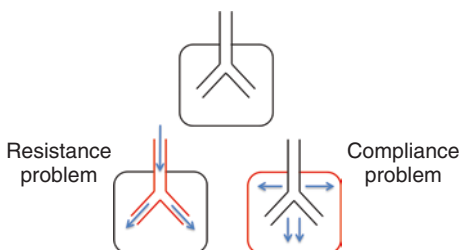


FIGURE 5.6 Resistance versus compliance

To return to our diagram of resistance and compliance in Fig. 5.6, we can imagine that the patient on the left side may have a very high PIP, given the resistance in the system. But, with healthy lungs and a normal compliance, the Pplat would be low or normal. Therefore, there may be a large gap between the PIP and Pplat, indicating an issue with airway resistance. In the lungs on the right, the PIP could still be elevated, since there is a lot of excess pressure being delivered to this system, and PIP includes measures of both resistance and compliance. In addition, the Pplat would also be elevated in this diagram because there is excess pressure being delivered to the alveoli. However, if the difference between the elevated PIP and Pplat is $<5 \text{ cmH}_2\text{O}$, this would indicate a compliance problem alone.

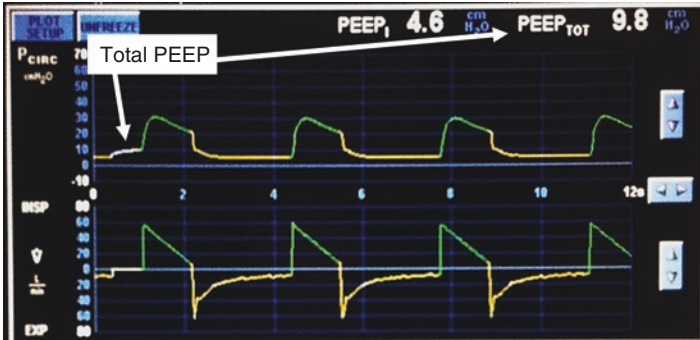


FIGURE 5.7 Ventilator screen demonstrating an expiratory hold maneuver. The total PEEP is 9.8, for an intrinsic PEEP, also known as autoPEEP, of 4.6

Another important pressure measure on the ventilator is that of the *autoPEEP* or *intrinsic PEEP* (*iPEEP*). When air is trapped in the alveoli at the end of exhalation, it exerts a pressure above and beyond the set PEEP. This pressure can actually be quantified on the ventilator by pressing the expiratory pause button, allowing the ventilator to briefly equilibrate the pressure at the end of expiration.

In Fig. 5.7, the clinician has performed an expiratory hold maneuver, as noted in the first breath on this diagram. The expiratory hold calculates the total PEEP ($PEEP_{TOT}$) in the system. Assuming the clinician set a PEEP of 5 cmH₂O, we can determine the intrinsic PEEP as follows:

$PEEP_{TOT} = ePEEP + iPEEP$. Therefore, *iPEEP*, as noted in the top of this figure, is approximately 4.6 cmH₂O. In other words, this patient is not completely exhaling at the end of each breath, leaving some additional pressure in the alveoli. This can also be noted in the flow tracings on the bottom of this figure, as the value at the end of each breath does not return to baseline. With such excess pressure at the end of exhalation, as can be seen in patients with COPD, the work of breathing can dramatically increase, leading to problems with ventilation.

Reference

1. Chacko B, Peter JV, Tharyan P, et al. Pressure-controlled versus volume-controlled ventilation for acute respiratory failure due to acute lung injury (ALI) or acute respiratory distress syndrome (ARDS). *Cochrane Database Syst Rev.* 2015;1:CD008807.

Suggested Reading

1. Singer BD, Corbridge TC. Basic invasive mechanical ventilation. *South Med J.* 2009 Dec;102(12):1238–45.
2. Wood S, Winters ME. Care of the intubated emergency department patient. *J Emerg Med.* 2011 Apr;40(4):419–27.
3. Mosier JM, Hypes C, Joshi R, et al. Ventilator strategies and rescue therapies for management of acute respiratory failure in the emergency department. *Ann Emerg Med.* 2015;66:529–41.
4. Archambault PM, St-Onge M. Invasive and noninvasive ventilation in the emergency department. *Emerg Med Clin North Am.* 2012;30(2):421–49. ix.

Chapter 6

Understanding the Ventilator Screen



Ventilators at times seem intimidating as there are numerous waveforms and values on the screen. Additionally, the data are presented slightly differently on the screens of each mechanical ventilator brand, increasing confusion. However, using the terms we have just reviewed, close inspection of ventilator screens will show that most of the waves and data are actually simple, given a little familiarity. To increase clinicians' comfort with ventilator screens, we have deliberately selected screenshots from a few different types of machines and modes of ventilation. Additionally, we have changed the colors of the backgrounds to demonstrate that the presentation is less important than the data provided.

Key concepts for evaluating ventilator screens are as follows:

1. The values set by clinicians are found on the bottom of the screen. The patient's response is located at the top of the screen.
2. Data are provided in both numerical and graphical contexts on the screen.
3. Much like studying EKGs, interpreting flow lines simply comes with experience. Unlike EKGs, however, there are very few variations to know! Ventilators provide three types of tracings: flow, pressure, and volume. Some mechanical ventilators show all three, while other brands allow the

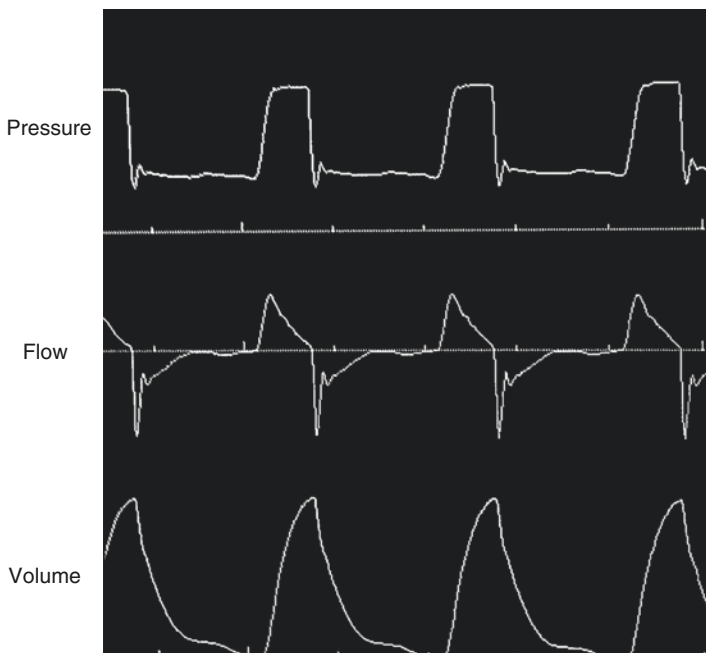


FIGURE 6.1 Typical waveforms for pressure, flow, and volume are illustrated

clinician to choose two tracings to display on the screen. Fortunately, all are labeled directly on the ventilator screens.

Figure 6.1 illustrates typical pressure, flow, and volume waveforms on a ventilator screen. Please also refer the theoretical illustration of Fig. 2.5, highlighting the relationship between the volume, flow, and pressure.

Examine the image of the mechanical ventilator screen in Fig. 6.2 closely and try to answer the following questions:

1. What is the PEEP?
2. What is the respiratory rate? (Hint: it is also known as the “frequency.”) Is the patient overbreathing? How could you tell?

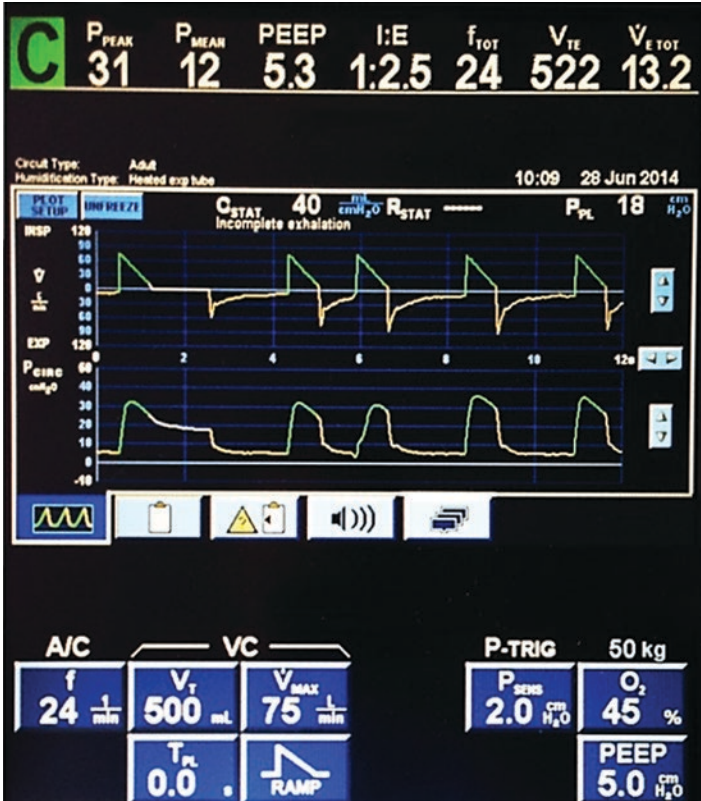


FIGURE 6.2 Example ventilator screen from an ICU patient

3. What is the set tidal volume? What is the tidal volume the patient is actually receiving?
4. What is the peak inspiratory pressure? What is the Pplat?
5. What is the I:E ratio? Is this set directly or indirectly on this particular patient?
6. What is represented by the top tracing? What is represented by the bottom tracing? What value (pressure, flow, or volume) is not shown here?
7. What is the minute ventilation?

Answers for Fig. 6.2:

1. PEEP is 5cmH₂O
2. The set RR is 24 (the frequency, or f). This patient is not overbreathing, as the rate up top is also 24.
3. The set tidal volume (VT) is 500, but the patient is receiving 522. This small variation is to be expected from breath to breath.
4. The peak inspiratory pressure (PIP) is 31. The Pplat is 18.
5. The I:E ratio is 1:2.5. Looking at the bottom of the screen, the max inspiratory flow is set at 77 L/min. We do not see any setting for the specific I:E ratio. Therefore, the I:E is set indirectly. Please refer to Chap. 5 for a discussion of setting the I:E indirectly.
6. The top tracing is the flow. Note that it is labeled at the left side of the screen. The bottom is pressure. Volume is not pictured.
7. The minute ventilation (VE) is 13.2.

Figure 6.3 is another ventilator screen from a different mechanical ventilator brand. Again, practice looking for certain values.

1. What is the set tidal volume?
2. What is the PEEP?
3. What is the set respiratory rate? Is the patient overbreathing?
4. What is the PIP? What is the Pplat?
5. What is the inspiratory time?
6. What do the 50 cmH₂O, 100 L/min, and 500 mL signify to the left side of the screen?

Answers for Fig. 6.3:

1. The set tidal volume is 350
2. The PEEP is 20
3. The set rate is 24, but the patient is overbreathing, as the actual rate is 26
4. The PIP is 33, and the Pplat is 31
5. The inspiratory time is 0.9 s
6. These values are labels on the Y axis for the pressure, flow, and volume tracings, respectively

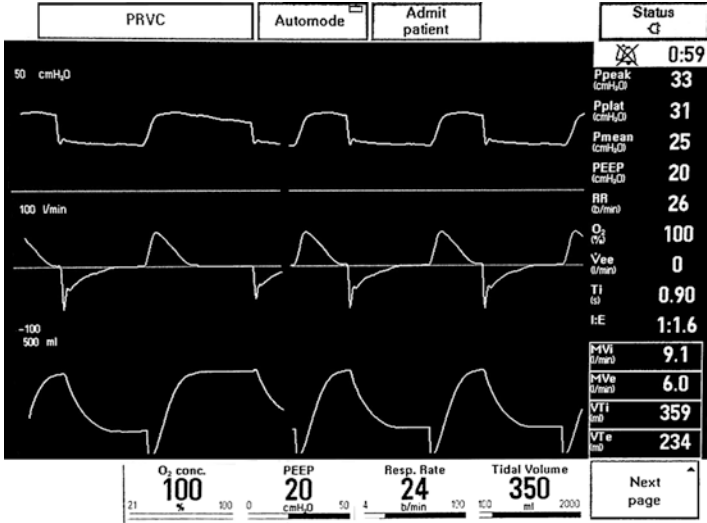


FIGURE 6.3 Example ventilator screen. Note that although the design differs slightly from Fig. 6.2, the general formatting is consistent. The variables set by the clinician are at the bottom, and resultant values and graphical information are on the top of the screen

As a final example of the variability, yet similarity, among ventilator interfaces, Fig. 6.4 is a screenshot from yet another style of ventilator.

1. What is the mode?
2. What tidal volume is the patient receiving? Bonus: What does this tell you about the patient’s compliance?
3. What is the respiratory rate (or frequency)?
4. What is the PIP? Bonus: Is it possible to check a Pplat?
5. What is the minute ventilation?

Answers for Fig. 6.4:

1. Pressure support. There are several clues. The “S” in the upper left-hand corner indicates “Support.” Review Fig. 6.2. There is a “C” in that corner, indicating that the most recent breath delivered was a “Controlled” breath. Many ventilators will also show an “A” for “Assist” when a patient is in Assist Control mode and triggers a breath.



FIGURE 6.4 Example ventilator screen, highlighting yet another style, yet the same basic data are all provided

2. The other clues are that there is no set respiratory rate, and the settings at the bottom of the screen feature pressures.
3. The patient is receiving 959 mL of tidal volume. This is a very high volume and may need to be intervened upon!
4. There is no set respiratory rate, but the patient is averaging 8.3 breaths per minute.
5. The PIP is 17. This should be, and is, very close to the set pressure support of 5cmH₂O + the PEEP of 10cmH₂O. It is not uncommon to have small variances in numbers between the set and delivered pressures and volumes.
6. The minute ventilation is 8.52. This is intuitive, as the patient is taking about a liter per breath and is breathing just over 8 times a minute, leading to 8.5 L/min of airflow.

Suggested Reading

1. Singer BD, Corbridge TC. Basic invasive mechanical ventilation. *South Med J.* 2009;102(12):1238–45.
2. Mosier JM, Hypes C, Joshi R, et al. Ventilator strategies and rescue therapies for management of acute respiratory failure in the emergency department. *Ann Emerg Med.* 2015;66:529–41.

Chapter 7

Placing the Patient on the Ventilator



Anticipating Physiologic Changes

Critically ill patients are at high risk of deterioration with intubation and initiation of mechanical ventilation. Much of this chapter is devoted to reviewing the effects positive pressure ventilation (PPV) can have on pulmonary physiology. However, mechanical ventilation can also have extrapulmonary effects that warrant review. Specifically, PPV can lead to an increase in the intrathoracic pressure, which leads to decreased venous return and decreased preload. While we use this principle to care for those with congestive heart failure (CHF), in excess, this phenomenon can lead to a decrease in the cardiac output and hypotension, especially in the intravascularly depleted patient, those with shock physiology, or with air trapping. Additionally, PPV leads to decrease the left ventricular afterload. Again, using the patient with an acute CHF exacerbation as an example, this principle can lead to an increase in the stroke volume and cardiac output.

When intubating and placing the patient on the ventilator, the emergency medicine clinician should anticipate these effects. A volume depleted patient, such as a patient with a GI bleed, may have hemodynamic collapse with initiation of positive pressure ventilation.

When initiating mechanical ventilation in the ED, the practitioner must be conscientious to ensure adequate gas

exchange to meet the metabolic demands of the patient. For example, a patient with severe metabolic acidosis with respiratory compensation might be very tachypneic. One must be cognizant to increase the respiratory rate on the ventilator to help meet the patient's metabolic demands. Failure to do so can be detrimental for the patient and lead to rapid decompensation.

Along the same lines, the practitioner must be careful to set and then adjust the ventilator settings to prevent further decompensation or injury. For example, excessive volumes ventilator can lead to volutrauma and impaired gas exchange. Excess pressure can lead to hemodynamic instability or barotrauma.

Setting the Ventilator

The goal of reviewing the terms, physiology, and concepts behind mechanical ventilation is to be able to put the pieces together and improve our care of mechanically ventilated patients in the ED. Also, please remember that ventilator settings may require adjustment as the patient's disease evolves or resolves. Therefore, once the initial settings are placed, the clinician must assess the patient, and continuously adjust best to meet the patient's metabolic demands, while trying to reduce harm.

To that end, let us practice selecting ventilator settings. Imagine that you just intubated a patient who presented to your ED after an overdose of an unknown medication, leading to apnea and a GCS of 3. How would you select ventilator settings for this patient?

Mode To start, select a mode. Most patients in the ED, especially shortly after intubation, should be ventilated in assist control (AC). Assist control would be appropriate for our hypothetical patient, as she is making no respiratory efforts. The next decision involves selecting a volume-targeted or a pressure-targeted mode. In the clear majority of cases,

this decision is one of personal preference and local customs. Numerous studies have found no differences for patients ventilated with one or the other. Most clinicians chose volume-targeted assist control, or volume control.

Tidal Volume (TV) The appropriate tidal volume is based upon the patient's height and biological sex, as these parameters determine predicted body weight and lung size. Take care to use predicted body weight, and not actual body weight, as using actual body weight can greatly overestimate the appropriate tidal volume. In contrast to older practices, which used "high" tidal volumes of 10–12 mL/kg, current practice based on several trials suggests that patient should be ventilated with "lower" tidal volumes of 6–8 mL/kg.

Respiratory Rate (RR) A reasonable approach is to consider the desired minute ventilation and chose a respiratory rate to approximate this value. Assuming there are no acid-base derangements, targeting relatively normal minute ventilation is appropriate. If we selected a tidal volume of 400 based on her height, a respiratory rate of 15 breaths per minute will lead to a minute ventilation of 6 L/min.

Conversely, if there is an acid-base disturbance, such as can occur with the ingestion of a toxin like ethylene glycol or in sepsis, the patient will need larger minute ventilation to correct the acidosis. Setting her rate at 24 breaths per minute will give a minute ventilation of 9.6 L/min. Regardless, about 20–30 min after selecting initial settings, clinicians should check an arterial blood gas (ABG) to assess acid/base status and oxygenation and make ventilator changes as needed. Also, as the disease process improves, the respiratory rate may need to be adjusted.

PEEP PEEP should always be set at least 5 cmH₂O, to reduce atelectasis. The conditions that will require a higher PEEP are those leading to worsening hypoxemia, wherein more atelectasis or derecruitment would be detrimental. Additionally, patients with large abdominal or chest walls may

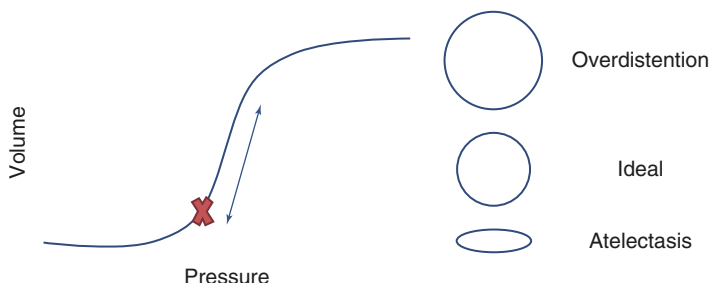


FIGURE 7.1 Theoretical representation of ideal PEEP. The PEEP should be high enough to prevent atelectasis with exhalation, but low enough that inhalation does not result in overdistention. The red “x” in this diagram shows this ideal spot for the relationship between the volume and pressure for this hypothetical patient. The double-ended arrow represents the changes in inspiration and exhalation

require a higher PEEP to prevent compression from abdominal contents. The concept behind the ideal PEEP is illustrated in Fig. 7.1. Every patient will have a relationship between the change in pressure and the change in volume with each breath. The PEEP should be set above the threshold for atelectasis, but such that the breath will not lead to overdistention.

Using our hypothetical patient with intubated for a GCS of 3, if she has a small to average habitus, a PEEP of 5 is likely appropriate to start. If she is heavier or has a larger abdomen or chest wall, she may be more prone to atelectasis. This would make a higher initial PEEP of 7–10 cmH_2O reasonable.

Inspiratory flow and I:E ratio The inspiratory flow and I:E ratios are commonly set at 60 L/min and 1:1.5 to 1:2, respectively. Common inspiratory times are 0.75–1 s. In certain circumstances, such as in airway obstruction with asthma, allowing more time for exhalation is beneficial. In these cases, one can increase the inspiratory flow or decrease the I:E ratio, to 1:3 or 1:4. Reexamine the ventilator screen shown in Chap. 6, as Fig. 7.2.

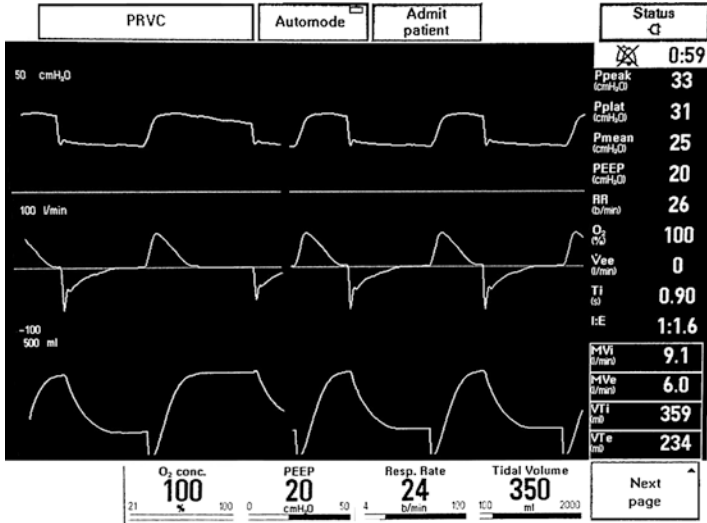


FIGURE 7.2 Ventilator screen demonstrating the relationship between respiratory rate, inspiratory time, and I:E ratio

In this example, the respiratory rate is 26, meaning that each breath is allotted 2.3 s ($60 \text{ s}/26 \text{ breaths} = 2.3 \text{ s/breath}$).

The inspiratory time is 0.9 s. This means that expiratory time is 1.4 s ($2.3 - 0.9 \text{ s}$).

The ratio of inspiratory time to expiratory time is therefore 0.9:1.4 – or approximately 1:1.6.

At the bedside, ventilators will provide this information, just as illustrated in Fig. 7.2. The clinician does not have to perform the calculations, but understanding the concepts is important for setting and adjusting the ventilator. To return to the example of our patient intubated for the overdose, we could consider what changes we would make if she had bronchospasm. In addition to treating with bronchodilators, we would give more time for exhalation, it is important to understand that this would mean either decreasing the respiratory rate or decreasing the inspiratory time.

After Initial Settings

Mechanical ventilation is a dynamic intervention, and once a patient is intubated and mechanically ventilated, the clinician must continuously reassess the patient and determine the best settings to help meet the metabolic and oxygen demands while avoiding any additional injuries. All intubated patients should have an arterial blood gas (ABG) checked 20–30 min after intubation. While venous blood gases (VBG) are excellent in the ED and are useful for evaluating a patient's pH and ventilation, a VBG cannot provide any data regarding oxygenation. Most patients are intubated and started on a FiO_2 of 100%, although this can be titrated down, to reduce the risks of oxygen toxicity, a condition increasingly appreciated in relation to numerous causes of critical illness.

To report these ventilator settings, such as when speaking to the intensivist, one would say, “The patient is on Assist Control/volume control, tidal volume 400mL, rate of 15 breaths per minute, PEEP 5 cmH₂O, and an FiO_2 of 100%. She is occasionally overbreathing to a rate of 18 breaths per minute. Her initial ABG after 30 min on these settings showed...”

Patients also remain at risk for hemodynamic perturbations after the initiation of ventilation or with changes in ventilation, due to changes in physiology along with the fluctuations in preload and afterload. Therefore, clinicians must continue to be mindful of the patient's intravascular status in ventilated patients and resuscitate these patients as needed.

Suggested Reading

1. The acute respiratory distress syndrome network. Ventilation with lower tidal volumes as compared with traditional tidal volumes for acute lung injury and the acute respiratory distress syndrome. *N Engl J Med.* 2000;342:1301–8.
2. Chacko B, Peter JV, Tharyan P, et al. Pressure-controlled versus volume-controlled ventilation for acute respiratory failure due to

- acute lung injury (ALI) or acute respiratory distress syndrome (ARDS). *Cochrane Database Syst Rev.* 2015;1:CD008807.
3. Mosier JM, Hypes C, Joshi R, et al. Ventilator strategies and rescue therapies for management of acute respiratory failure in the emergency department. *Ann Emerg Med.* 2015;66:529–41.
 4. Slutsky AS, Ranieri VM. Ventilator-induced lung injury. *N Engl J Med.* 2013;369:2126–36.

Chapter 8

Specific Circumstances: Acute Respiratory Distress Syndrome (ARDS)



Acute respiratory distress syndrome (ARDS) is a condition of diffuse alveolar damage and inflammation, secondary to any number of possible processes. ARDS is defined by four criteria [1]:

1. The condition must be acute (<7 days).
2. The findings are not solely explained by cardiogenic pulmonary edema.
3. The chest X-ray must have bilateral opacities, as shown in Fig. 8.1.

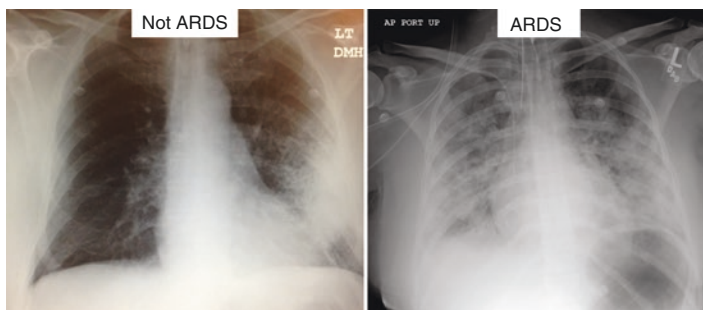


FIGURE 8.1 Chest X-rays illustrating the difference between ARDS and pneumonia. Note that both patients can be severely hypoxemic, but ARDS has bilateral, diffuse infiltrates

4. While on at least 5 cmH₂O of positive pressure ventilation, the ratio of PaO₂ to FiO₂ (expressed as a decimal, such as 0.7) must be <300.
 - (a) Mild ARDS is a PaO₂/FiO₂ ratio of 200–300.
 - (b) Moderate ARDS is 100–199.
 - (c) Severe ARDS is <100.

Although patients rarely present to the ED in fulminant ARDS, as it usually develops later in the critically ill, ARDS can be seen in the ED. Of all the interventions in critical care, few have been as reproducibly beneficial to patients as low tidal volume ventilation [2]. Positive pressure ventilation, especially with large tidal volumes or high pressures, has been shown to cause injury in both patients with ARDS as well as patients who do not yet have ARDS. Prevention of ARDS, or prevention of exacerbating ARDS with ventilator-induced lung injury, is a key benefit of active ventilator management in the ED.

Many of the maneuvers used in severe hypoxemia to improve oxygenation and ventilation can be deleterious in the long term. Increasing the mean airway pressure (MAP) is one of the major goals of positive pressure ventilation, and higher MAPs are often associated with improved oxygenation. However, higher pressures in the alveoli are also associated with worse outcomes. Therefore, the clinician must balance the risk of increasing the MAP with using evidence-based ventilator management, shown in Fig. 8.2.

As described above, tidal volumes are best represented in both mLs and mLs/kg of predicted body weight. The predicted body weight is a surrogate for the patient's anticipated lung volume. Lung volumes depend upon a patient's height and biological sex. As many patients weigh more than their predicted body weight, the actual body weight should never be used as a replacement for the predicted body weight.

Once the correct tidal volume is selected, the pressures should be assessed. In ARDS, as well as other patients, maintaining a Pplat <30cmH₂O is key to preventing ventilator-induced lung injury [3]. Using an inspiratory hold, the Pplat

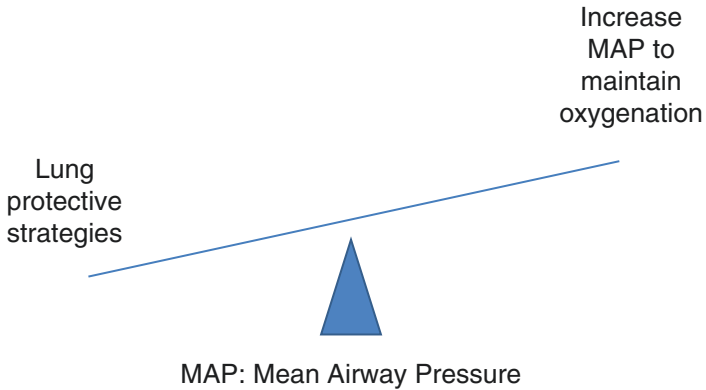


FIGURE 8.2 Although the clinician should do what they must to stabilize a hypoxemic patient, the principles of evidence-based ventilator management should hold weight. For example, large tidal volumes may lead to a rapid improvement in hypoxemia, but their use trades short-term benefit for long-term harm

should be confirmed to be less than 30 cmH₂O. If Pplat is >30 cmH₂O, a lower tidal volume can be considered, even down to 4 mL/kg. Figure 8.3 shows an example of a Pplat.

Patients being ventilated with low tidal volumes will require a higher rate to maintain minute ventilation. Most patients with ARDS will require RR of 20 breaths per minute or greater.

PEEP is the next setting to address. Clearly, oxygenation is a critical factor for these patients. Most modest increases in PEEP do not substantially recruit collapsed alveoli, but PEEP can prevent further derecruitment. Recall from Chap. 7 that the goal of PEEP is to prevent atelectasis with exhalation. Stiff or edematous lungs are more prone to atelectasis, mandating a higher PEEP. Many of these patients will need higher PEEPs of 10–16 cmH₂O, and at times, even over 20 cmH₂O! The PEEP will contribute to the Pplat, and therefore, the Pplat should be checked with any PEEP change, just as with any TV change. In addition to minimizing D recruitment, PEEP may offer the benefit of minimizing “atelecta-

At times, patients may have refractory, severe hypoxemic respiratory failure. After checking all ventilator setting as described above, the clinician should employ additional evidence-based maneuvers.

Recruitment Maneuvers

In well-sedated and possibly chemically relaxed patients, the first maneuver is to provide a recruitment maneuver. Recalling that derecruitment is a common cause of hypoxemia, gently recruiting alveoli can improve oxygenation. The concept behind a recruitment maneuver is simple: the application of a sustained pressure, for 20–40 s, to open up collapsed alveoli [7]. However, there are two potential downsides.

First, the damage to lungs is heterogeneous. Some areas are atelectatic, some are fluid-filled, some are already overdistended, and some are even normal. The goal of the recruitment maneuver is to reopen the atelectatic areas, as illustrated in Fig. 8.4.

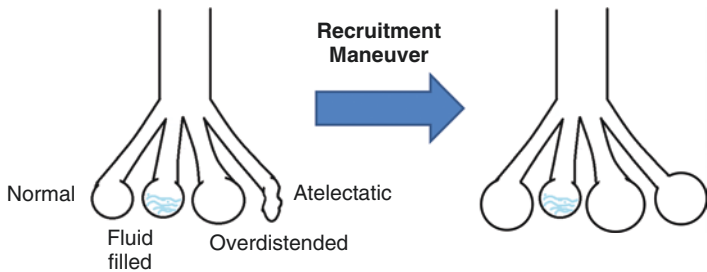


FIGURE 8.4 ARDS is a heterogeneous condition. The “alveoli” here represent areas of lung, with various lung units being normal, fluid-filled, overdistended, or atelectatic. The recruitment maneuver may transiently overdistend the normal and already overdistended lung units, but the expectation is that recruiting the atelectatic areas will overall improve oxygenation after the maneuver is completed

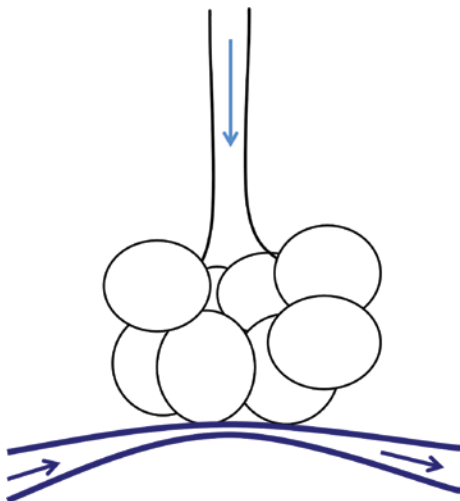


FIGURE 8.5 A lung unit that becomes overdistended can surpass the pressure in the capillaries, transiently reducing blood flow and gas exchange to that portion of the lung. This is what is responsible for the temporary desaturation during a recruitment maneuver

However, note that the normal and overdistended areas may also become even more overdistended during the recruitment maneuver. This overdistention within the previously “good” parts of the lung can lead to decreased gas exchange during the recruitment, causing desaturation, as seen in Fig. 8.5. This effect should be temporary and improve after the maneuver.

The second effect is that the patient can become hemodynamically unstable, due to a significant increase in the intra-thoracic pressure and resultant decrease in preload. Again, this should be temporary with reduction in the pressure, but in very unstable or preload dependent patients, this can precipitate hemodynamic collapse.

There are many methods of performing recruitment maneuvers. One of the methods least likely to cause hemodynamic perturbations is to serially increase PEEP in small increments [7]. For example, increasing the PEEP to a final maximum of 20–30 cmH₂O in increments of 2 cmH₂O,

each held for 10–20 s, while keeping the PIP <45 cmH₂O, can be effective in many patients. Recruitment maneuvers should never be performed without a respiratory therapist and nurse present. All clinicians should be aware of the risks of transient hypoxemia and hypotension.

Neuromuscular Blockade

Once the patient is adequately sedated, neuromuscular blockade can be considered in patients with ARDS and a P/F ratio <120 . For example, a continuous infusion of cisatracurium, when administered within the first 48 h of severe ARDS, for 48 additional hours, improves 90-day mortality and decreases ventilator days [8, 9].

The next maneuver is proning the patient, or placing them in the prone position, to improve oxygenation to the posterior lungs. Proning the patient improves V/Q matching and allows the patient to have gas exchange along the posterior aspects of the lungs, illustrated in Fig. 8.6. Proning has been shown to improve mortality in severe ARDS in a large multicenter study [10]. However, this maneuver requires specialized expertise and a coordinated effort among providers to avoid dislodging the endotracheal tube and patient harm. If a patient has such severe hypoxemia in the ED that the clinicians are considering proning, expert consultation should be sought.

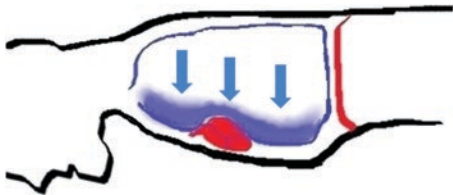


FIGURE 8.6 The posterior aspect of the lungs holds a large surface area for gas exchange. Additionally, moving the heart anteriorly and off the lungs helps reduce atelectasis behind the heart. Coupled with mechanical changes in the chest wall, proning can significantly improve oxygenation

Another consideration is the administration of inhaled pulmonary vasodilators, such as inhaled nitric oxide (not to be confused with nitrous oxide, the anesthetic agent) or prostacyclins, such as epoprostenol. Hypoxemic patients generally have heterogeneous lung pathology, with some damaged areas, not participating in oxygenation and ventilation, as well as some relatively unharmed areas that are doing the bulk of gas exchange. Inhaled pulmonary vasodilators will vasodilate the areas that are participating in gas exchange, effectively increasing blood flow to the good areas of the lung and allowing the ineffective areas to continue to have hypoxic vasoconstriction. This principle is illustrated in Fig. 8.7.

Finally, patients with severe, refractory hypoxemia should be referred to an ECMO center for consideration of ECMO support. A discussion of ECMO is beyond the scope of this chapter; however, transfer for ECMO has been shown to improve survival in patients with severe ARDS [11].

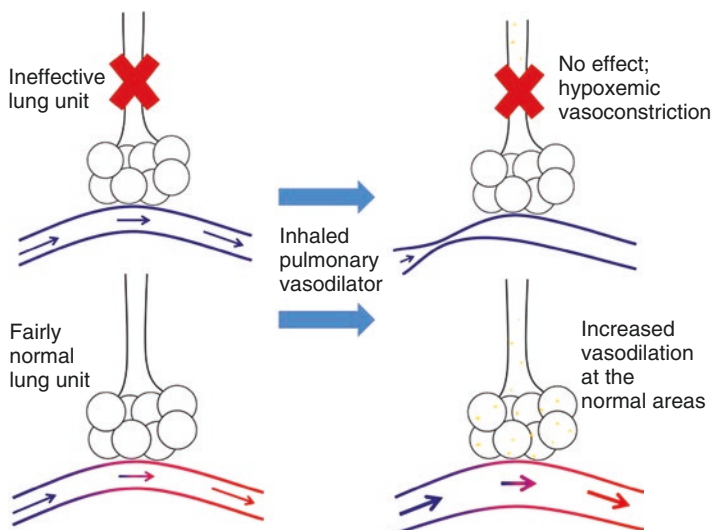


FIGURE 8.7 Inhaled pulmonary vasodilators only reach the alveoli of lung units participating in gas exchange. They dilate the capillaries for these “good” lung units and thereby direct more blood flow to the areas participating in gas exchange

References

1. ARDS Definition Task Force, Ranieri VM, Rubenfeld GD, et al. Acute respiratory distress syndrome: the berlin definition. *JAMA*. 2012;307:2526–33.
2. The Acute Respiratory Distress Syndrome Network. Ventilation with lower tidal volumes as compared with traditional tidal volumes for acute lung injury and the acute respiratory distress syndrome. *N Engl J Med*. 2000;342(18):1301–8.
3. Slutsky AS, Ranieri VM. Ventilator-induced lung injury. *N Engl J Med*. 2013;369:2126–36.
4. Damiani E, Adrario E, Girardis M, et al. Arterial hyperoxia and mortality in critically ill patients: a systematic review and meta-analysis. *Crit Care*. 2014;18:711.
5. Helmerhorst HJ, Roos-Blom MJ, van Westerloo DJ, et al. Association between arterial hyperoxia and outcome in subsets of critical illness: a systematic review, meta-analysis, and meta-regression of cohort studies. *Crit Care Med*. 2015;43:1508–19.
6. Page D, Ablordeppey E, Wessman BT, Mohr NM, Trzeciak S, Kollef MH, Roberts BW, Fuller BM. Emergency department hyperoxia is associated with increased mortality in mechanically ventilated patients: a cohort study. *Crit Care*. 2018;22(1):9.
7. Keenan JC, Formenti P, Marini JJ. Lung recruitment in acute respiratory distress syndrome: what is the best strategy? *Curr Opin Crit Care*. 2014;20:63–8.
8. Papazian L, Forel JM, Gacouin A, et al. Neuromuscular blockers in early acute respiratory distress syndrome. *N Engl J Med*. 2010;363:1107–16.
9. Murray MJ, DeBlock H, Erstad B, et al. Clinical practice guidelines for sustained neuromuscular blockade in the adult critically ill patient. *Crit Care Med*. 2016;44:2079–103.
10. Guerin C, Reignier J, Richard JC, et al. Prone positioning in severe acute respiratory distress syndrome. *N Engl J Med*. 2013;368:2159–68.
11. Peek GJ, Mugford M, Tiruvoipati R, et al. Efficacy and economic assessment of conventional ventilatory support versus extracorporeal membrane oxygenation for severe adult respiratory failure (CESAR): a multicentre randomised controlled trial. *Lancet*. 2009;374:1351–63.

Chapter 9

Specific Circumstances: Asthma and COPD



In asthma, the patient has constriction of the bronchial smooth muscles in the airways, leading to reversible air trapping. This is indicated in the schematic of Fig. 9.1. Note that the bronchial muscles do not extend into the small airways.

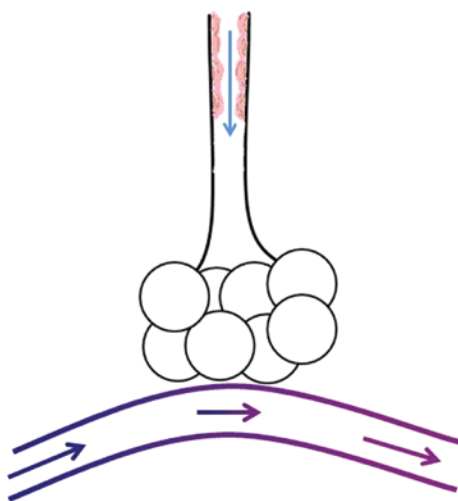


FIGURE 9.1 In asthma, the patient has intermittent constriction of the smooth muscles of the bronchi, thereby limiting airflow

Intubation of an asthmatic in the ED is a dreaded complication of this illness, as asthmatics can deteriorate rapidly on the ventilator without close monitoring and active management. The goal with a ventilated asthmatic is to prevent breath-stacking or autoPEEP, and the hemodynamic instability that can result.

Before discussing the ventilator management of asthma, clinicians should note that intubation of an asthmatic should trigger even more active management with medications, rather than less. Intubated asthmatic patients should continue to receive aggressive treatment with bronchodilators, steroids, magnesium, as well as deep sedation and possibly even neuromuscular blockade in the initial hours after intubation, in an effort to relax the chest wall musculature and gain control of the situation. Please note that neuromuscular blockade only works on skeletal muscle and therefore will not bronchodilate smooth muscle in the airways. In addition, it is very critical to be aware of the patient's intravascular volume status, as the excess positive pressure can lead to hemodynamic collapse. Moreover, the excess pressure, including the auto-PEEP, can result in barotrauma, such as the development of a pneumothorax very quickly in this patient population.

Four ventilator maneuvers increase expiratory time, namely, decreasing the respiratory rate, decreasing the I:E ratio, decreasing the inspiratory time, or increasing the inspiratory flow. Of these, decreasing the respiratory rate is the most effective means to allow more time to exhale.

Figure 9.2 shows a schematic of 30 s with two patients, set with the same I:E ratio of 1:2. The first patient has a rate of 10 breaths per minute, allowing 6 s per breath cycle. The second patient has only 3 s per breath cycle, given the respiratory rate of 20. The blue represents inspiration, the red the time for exhalation. Note that even with the same I:E, the lower rate offers a substantially longer time to exhale.

In looking further at this diagram, one can imagine the effects of changing the I:E ratio, the inspiratory flow, or the

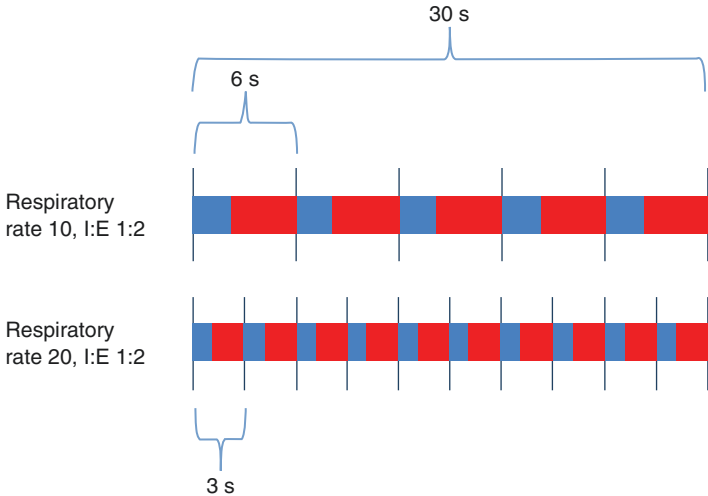


FIGURE 9.2 In this figure, the red represents time to exhale. Note the dramatic effect of decreasing the respiratory rate from 20 to 10, even while all other parameters are held equal

I time. Figure 9.3 shows a hypothetical example of the effects of these changes in a patient on volume control. In a given patient, the exact values will vary, but the purpose of the illustration is to show the relationship among the parameters of I:E, inspiratory time, and inspiratory flow.

In addition to a slow respiratory rate, a low I:E ratio, a short inspiratory time, and/or a fast inspiratory flow rate, asthmatics should also be ventilated with low tidal volumes. Considering that the larger the tidal volume, the more the patient has to exhale, this is fairly intuitive.

In monitoring an intubated asthmatic, looking for air trapping is key. In the vent tracing in Fig. 9.4, note that the flow tracing, in the middle, does not return to the baseline before the next breath (Red arrows). This represents that the patient is still exhaling when the next breath is given, leading

Expressed in I:E ratio	Expressed in inspiratory time	Expressed in inspiratory flow
1:3	0.75 s	90 L/min
1:2	1 s	70 L/min
1:1	1.5 s	50 L/min

FIGURE 9.3 This figure illustrates the relationship between the I:E ratio, the inspiratory time, and the inspiratory flow. Decreasing the I:E ratio, decreasing the inspiratory time, and increasing the inspiratory flow all provide more time for exhalation



FIGURE 9.4 In this image of a ventilator screen, the flow does not return to the baseline before the next breath is given, indicating that the patient is still exhaling when being forced to inhale. This creates air trapping. Note that the quantity of air trapping cannot be determined by the flow waveform; this is qualitative data only

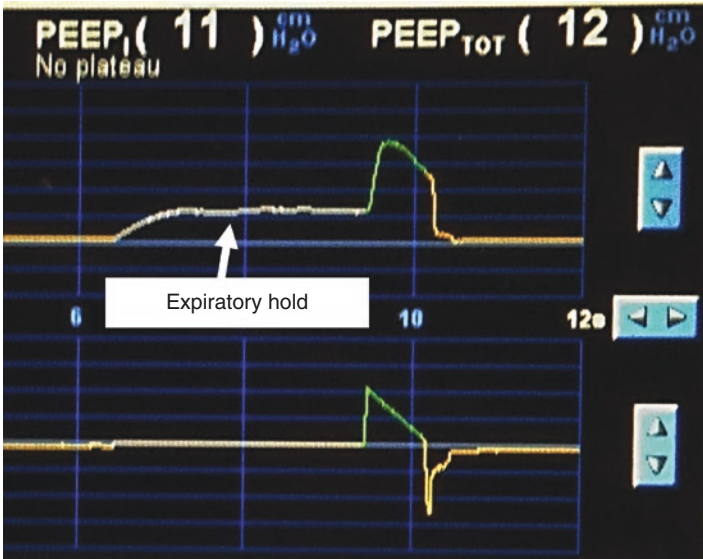


FIGURE 9.5 The amount of air trapping can be quantified by an expiratory hold. In this figure, the PEEP was turned to 1 cmH_2O (for illustration purposes only, not recommended for clinical practice). The total PEEP is 12 cmH_2O , and the autoPEEP, or intrinsic PEEP, is therefore 11 cmH_2O

to air trapping. Seeing this pattern on the ventilator can be an early clue that the patient is air trapping. If you were caring for this patient, how would you address this air trapping? (Fig. 9.5).

In this patient, you could first decrease the respiratory rate, or increase sedation if the patient is overbreathing. The I:E ratio is only 1:2, so changing the I time to make a ratio of 1:3 or 1:4 is also appropriate. Also continued treatment with bronchodilators to decrease the bronchospasm associated with this disease will also mitigate the excess auto-PEEP.

Recall that to quantify the pressure exerted by air trapping, one should check for autoPEEP by checking an expiratory hold button on the mechanical ventilator. In this tracing, what is the autoPEEP, or the intrinsic PEEP? What is the total PEEP?

The intrinsic PEEP is 11, and the total PEEP is 12. This indicates that the patient was only set on 1 of PEEP (an unusual setting, but used in this circumstance for demonstration purposes only).

Thus, to set the ventilator for an asthmatic, select a low tidal volume of 6–8 mL/kg of predicted body weight. The respiratory rate should be low, less than 20 breaths per minute, and often around 10. The I:E ratio should be changed to 1:3 or less. PEEP should be set at 5 cmH₂O. The FiO₂ should be down-titrated as tolerated. These patients continue to receive heavy sedation, possibly NMB if required, continuous bronchodilators, and close monitoring for breath stacking and autoPEEP. AutoPEEP should be monitored periodically or after any ventilator change with an expiratory hold. Arterial blood gases (ABGs) should be checked to ensure that the patient is being adequately ventilated.

COPD

There are two types of obstructive lung disease falling under the umbrella of COPD, namely, chronic bronchitis and emphysema (Fig. 9.6).

Chronic bronchitis can resemble the asthmatic schematic above, with the notable exception that muscles hypertrophy and are not entirely reversible. Additionally, chronic bronchitis is associated with increased mucous production (Fig. 9.7).

Emphysema is a disease of parenchymal destruction. Not only is there loss of alveoli, resulting in decreased surface area, or decreased diffusion area (leading to an increased DLCO), but the small airways can become floppy due to the loss of other tissues holding them open (Fig. 9.8).

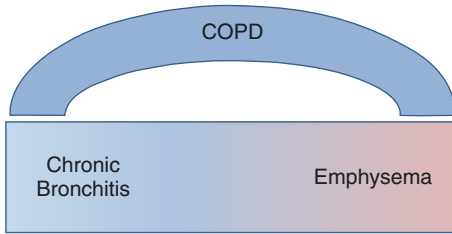


FIGURE 9.6 Both chronic bronchitis and emphysema fall under the umbrella term of COPD. Most patients with COPD will have aspects of both

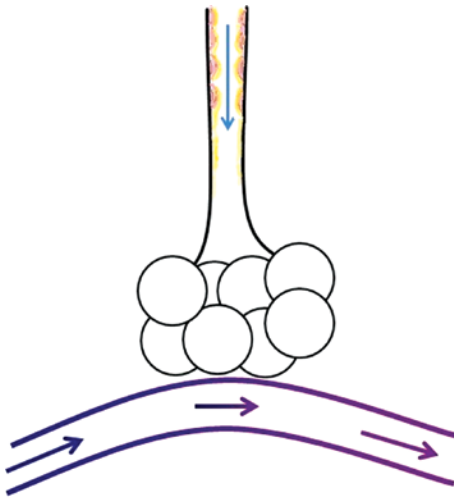


FIGURE 9.7 In chronic bronchitis, the patient has narrowing of the airways with nonreversible hypertrophy and increased mucous production

Understanding the pathophysiology of COPD is important for considering how to best ventilate these patients. It should be noted, however, that most patients with COPD have some mixing of elements of chronic bronchitis and emphysema. These conditions exist on a spectrum rather than a dichotomy.

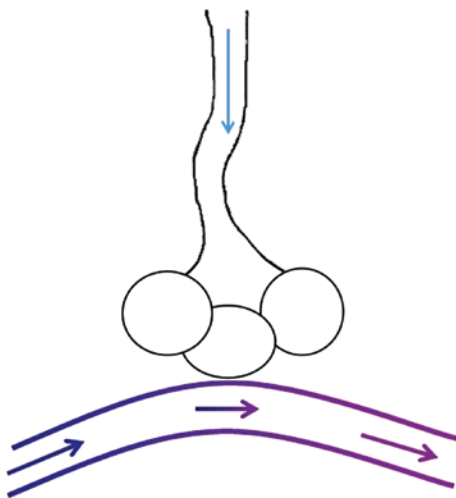


FIGURE 9.8 In emphysema, there is parenchymal destruction, with loss of alveoli and collapsibility of the small airways

Most patients with COPD are now managed with BPAP, with improved outcomes over intubation. However, on occasion, a patient with COPD is not a candidate for BPAP or fails to improve with a trial of BPAP, mandating intubation and invasive mechanical ventilation. Many of the principles that apply in mechanical ventilation for asthma also apply in COPD. Both are obstructive diseases, and in both processes, the patients require adequate time to exhale. Therefore, low tidal volumes, low rates, and low I:E ratios are appropriate. However, a key difference involves the role of PEEP.

Patients with COPD are at high risk of developing autoPEEP. Due to their obstructive disease, they require additional time to exhale. However, the mechanism of obstruction can differ between asthma and COPD, especially COPD with emphysematous changes as illustrated above. With the destruction of parenchyma, the small airways can collapse with exhalation, trapping air behind. In this circumstance, this

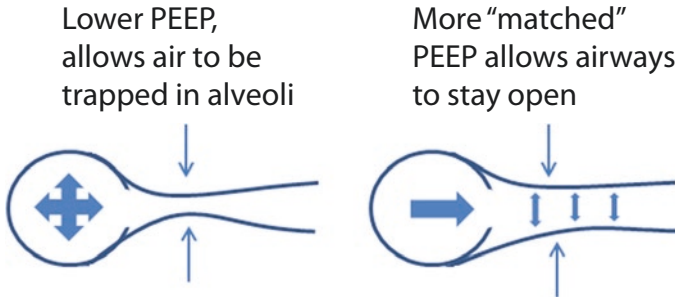


FIGURE 9.9 This figure illustrates how increasing PEEP to match autoPEEP can help reduce air trapping. By maintaining the patency of the small airways, the patient is better able to exhale

trapped air leads to autoPEEP. Increasing the set PEEP, to match the autoPEEP, is not necessarily an intuitive solution. However, as illustrated by the diagram below, increasing the PEEP to prevent collapse of these small airways can allow the patient to exhale more fully (Fig. 9.9).

Reexamine the tracing of Fig. 9.5 from the Asthma section, imagining that this patient has COPD. If this patient has 11 of autoPEEP, or intrinsic PEEP, what PEEP would you select? (Fig. 9.10).

To match the autoPEEP, 11 cmH₂O would be an appropriate PEEP selection.

Lastly, patients with COPD are often chronically hypoxemic. Indications of chronic hypoxemia physical exam findings of chronic hypoxemia can be demonstrated with nail clubbing. Additionally, an elevated hemoglobin level on the CBC can be an indication of a patient's compensation for chronic lung disease. Because these patients are baseline hypoxemic, and ventilation is often a relatively greater issue for them than hypoxemia, the oxygen saturation for a patient with COPD should be targeted at 88–92% in most circumstances. This is increasingly important as more data demonstrating the risks of hyperoxia continue to accumulate.



FIGURE 9.10 Illustration of an autoPEEP, or intrinsic PEEP, of 11

Suggested Reading

1. Archambault PM, St-Onge M. Invasive and noninvasive ventilation in the emergency department. *Emerg Med Clin North Am.* 2012;30(2):421–49, ix.
2. Levy BD, Kitch B, Fanta CH. Medical and ventilatory management of status asthmaticus. *Intensive Care Med.* 1998;24(2):105–17.
3. Medoff BD. Invasive and noninvasive ventilation in patients with asthma. *Respir Care.* 2008;53(6):740–8; discussion 749–50.

Chapter 10

Specific Circumstances: Neurologic Injury



Patients with neurologic injury requiring mechanical ventilation are an especially fragile patient population in the ED. Although EM clinicians cannot prevent the primary injury, as this has already occurred by the time the patient presents, preventing secondary injury is key. Patients with neurologic injury have specific needs for oxygenation, ventilation, and blood pressure management. In essence, the most important concept is to keep parameters normal—neither too high nor too low. These patients require very careful monitoring as the brain not only has no physiologic reserve, but also these patients are at risk of becoming unstable.

Traumatic Brain Injury

Management of oxygenation and ventilation in patients with traumatic brain injury has been well studied. Numerous studies have shown that early proper management of patients with TBI can improve outcomes, or at least, reduce the risk of secondary injury [1–3]. Studies of pre-hospital intubation and ventilation have shown worse outcomes for patients with TBI, due in large part to the hyperventilation that can occur in a setting with less monitoring [3, 4].

TBI patients are at risk for increased intracranial pressure. This hyperventilation leads to hypocapnia. With decreased CO_2 , blood vessels in the brain vasoconstrict, leading to

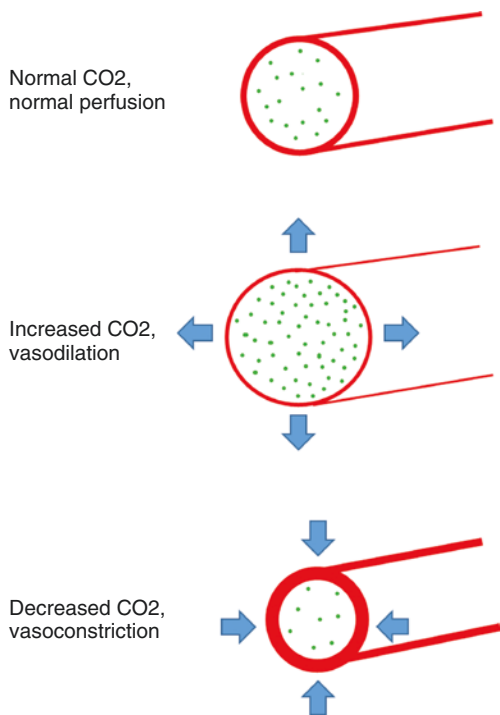


FIGURE 10.1 This figure illustrates the effects of PaCO_2 on the cerebral vasculature. In the top image, the PaCO_2 is normal, around 40 mmHg. In the second, the concentration is much increased, as may be seen with hypoventilation. This leads to vasodilation and can increase intracranial pressure. The bottom illustration demonstrates a low PaCO_2 , as would be seen with hyperventilation. This causes vasoconstriction and can lead to worsening ischemia in vulnerable areas of the brain

decreased blood flow as illustrated in Fig. 10.1. This vasoconstriction risks secondary injury from hypoperfusion and ischemia. Although in decades past, clinicians used to recommend hyperventilation of patients at risk for increased ICP, this is no longer recommended due to worsening outcomes with this hypoperfusion. Guidelines recommend targeting a normal PaCO_2 of 35–40.

Upon intubating a patient with TBI, bag-ventilating the patient should be minimized, to reduce the risk of unintentional hyperventilation. A useful tip for those bagging the patient is to only squeeze the bag until you see chest rise and only give the patient a breath when you take a breath. The patient should then be placed on the mechanical ventilator as soon as possible, targeting a reasonable starting minute ventilation of 7–8 liters/min. The trauma patient is likely to be hypermetabolic, so a slightly higher than normal MV is appropriate to start.

All patients with neurologic injury are at risk for ARDS. In other patient populations, permissive hypercapnia is accepted to maintain low tidal volumes and allow for adequate time to exhale. However, permissive hypercapnia is not appropriate in neurologically injured patients, and the ventilator should be adjusted accordingly.

Capnography can be very useful in this population. As the capnography is started, checking an ABG is beneficial to correlate the partial pressure of carbon dioxide (PaCO_2) with the end-tidal CO_2 (ETCO_2). Some patients, especially those with chest trauma or underlying lung disease, may have substantial dead space, leading to a larger than anticipated discrepancy between the two. Normally, the difference should be about 5. Once the relationship is established, the ETCO_2 can be followed for trends, assuming no significant changes in the pulmonary status.

It is intuitive that hypoxemia in TBI is also associated with worsened outcomes and secondary injury. However, clinicians may not realize that numerous studies have shown that patients are frequently HYPERoxygenated in emergency scenarios, right after intubation. This hyperoxia is also deleterious, with a supposed mechanism of worsening reperfusion injury and free radical production. Therefore, normoxia is the goal, and this is another reason an ABG should be checked 15–20 min after intubation. The FiO_2 should be lowered, targeting a PaO_2 75–100, for a corresponding O_2 saturation of 95–99%, depending upon the individual oxygen-hemoglobin dissociation curve.

TBI patients are also very susceptible to further injury from hypotension and hypertension. With intubation and initiation of ventilation, the clinicians should take care to manage hemodynamics aggressively. Volume depleted patients should be resuscitated with fluids or blood, as indicated by the circumstance, and any patient with hypotension or at risk for hypotension should receive vasopressors to maintain cerebral perfusion pressure. Similarly, laryngeal stimulation may at times lead to hypertension, and BPs over 140 systolic should be avoided or at least treated with sedation, opioids, and on rare occasions, antihypertensives as needed.

Ischemic Stroke

Patients with ischemic stroke may require intubation and mechanical ventilation for a variety of reasons, including the need for airway protection, respiratory failure after aspiration, or need for invasive procedures. While the need for intubation in ischemic stroke portends a poor prognosis, it is imperative that the emergency clinician prevents secondary injury to the vulnerable area of the brain, the penumbra, to the greatest extent possible.

Just as with TBI, patients with ischemic stroke are at risk for hypocapnia-induced vasoconstriction, leading to secondary ischemia. This vasoconstriction can then worsen outcomes by decreasing perfusion to the penumbra. Similarly, hypercapnia should be avoided to reduce vasodilation and the consequent increase in intracranial pressure. Therefore, clinicians should target normal PaCO₂ parameters of 35–45 mmHg [5]. The risk of increased ICP in an acute ischemic stroke is lower than in TBI, allowing a more liberal PaCO₂ target. Low tidal volume ventilation should be initiated, with a goal of 6–8 mL/Kg of predicted body weight. The MV should be targeted to 5–6 L/min, as these patients are less likely to be hypermetabolic than the TBI patients.

Both hypoxia and hyperoxia can be damaging to the ischemic stroke patient. Current guidelines recommend maintaining oxygen saturation greater than 94% but do not provide a recommended upper limit [6]. Because hyperoxia has been associated with increased mortality in ischemic stroke patients [7], the minimum FiO_2 required to maintain an O_2 saturation of 95% or above should be used. The value of ABGs in the neurologic injury population is not just to evaluate hypoxemia but to evaluate for hyperoxia, which is not readily detected with pulse oximetry.

Hypotension must also be avoided in the ischemic stroke patient. These patients are at risk of dehydration, and volume repletion before intubation, if possible, is advisable. Maintaining a systolic blood pressure of at least 140 mmHg until vessel recanalization is performed (if this is planned.) However, hypertension increases the risks of hemorrhage, especially in cases of intravenous thrombolysis, and the target should be less than 180/105.

Intracranial Hemorrhage

The principles of management for patients with intracranial hemorrhage are similar to patients with TBI and ischemic stroke. These patients are similarly susceptible to the ischemic effects of hypocapnia from hyperventilation [8]. Because the risk of increased intracranial pressure is higher in ICH than ischemic stroke, targeting a PaCO_2 of 35–40 mmHg is reasonable. Hyperoxia has been associated with increased mortality in this population as well. Patients with intracranial hemorrhage are at risk of developing ARDS and, as such, should also be ventilated with low tidal volumes of 6–8 mL/Kg predicted body weight. Close monitoring with ABGs and capnography is mandatory.

As with other patients with neurologic injury, the blood pressure can vary widely, and maintaining normal perfusion is key. The clinicians should be aware of the risks of hemodynamic lability with intubation and initiation of ventilation and be prepared to treat hypertension or hypotension rapidly.

TABLE 10.1 Goals for mechanical ventilation in neurologic injury

Traumatic brain injury	Ischemic stroke	Intracranial hemorrhage	Status epilepticus
PaCO ₂ 35–40 mmHg	PaCO ₂ 35–45 mmHg	PaCO ₂ 35–40 mmHg	PaCO ₂ 35–45 mmHg
MV 7–8 L/min	MV 5–6 L/min	MV 6–7 L/min	MV 8–10 L/min
PaO ₂ 75–100 mmHg	PaO ₂ 75–100 mmHg	PaO ₂ 75–100 mmHg	PaO ₂ 75–100 mmHg
O ₂ saturation 95–99%	O ₂ saturation 95–99%	O ₂ saturation 95–99%	O ₂ saturation 95–99%
SBP 120–140	SBP 140–180	SBP <140	Varies by etiology

PaCO₂ partial pressure of oxygen, *MV* minute ventilation, *PaO₂* partial pressure of oxygen, *SBP* systolic blood pressure

Status Epilepticus

Patients in status epilepticus requiring intubation have a few unique challenges. If possible, only short acting paralytics should be used with intubation to minimize obscuring the exam. Recall that these patients will be hypermetabolic, with a lactic acidosis, and the MV should be increased accordingly, likely starting at least at 8–10 L/min. Acid-base status should be followed closely to reduce the risk of secondary injury from additional metabolic insults. The core concepts are highlighted in Table 10.1.

References

1. Davis DP, Idris AH, Sise MJ, et al. Early ventilation and outcome in patients with moderate to severe traumatic brain injury. *Crit Care Med.* 2006;34:1202–8.
2. Warner KJ, Cuschieri J, Copass MK, et al. Emergency department ventilation effects outcome in severe traumatic brain injury. *J Trauma.* 2008;64:341–7.

3. Davis DP, Douglas DJ, Koenig W, et al. Hyperventilation following aero-medical rapid sequence intubation may be a deliberate response to hypoxemia. *Resuscitation*. 2007;73:354–61.
4. von Elm E, Schoettker P, Henzi I, et al. Pre-hospital tracheal intubation in patients with traumatic brain injury: systematic review of current evidence. *Br J Anaesth*. 2009;103:371–86.
5. Lahiri S, Schlick K, Kavi T, Song S, Moheet AM, Yusufali T, Rosengart A, Alexander MJ, Lyden PD. Optimizing outcomes for mechanically ventilated patients in an era of endovascular acute ischemic stroke therapy. *J Intensive Care Med*. 2017;32(8):467–72.
6. Jauch EC, Saver JL, Adams HP Jr, et al. American Heart Association Stroke Council; Council on Cardiovascular Nursing; Council on Peripheral Vascular Disease; Council on Clinical Cardiology. Guidelines for the early management of patients with acute ischemic stroke: a guideline for healthcare professionals from the American Heart Association/American Stroke Association. *Stroke*. 2013;44(3):870–947.
7. Rincon F, Kang J, Maltenfort M, et al. Association between hyperoxia and mortality after stroke: a multicenter cohort study. *Crit Care Med*. 2014;42(2):387–96.
8. Badenes R, Bilotta F. Neurocritical care for intracranial haemorrhage: a systematic review of recent studies. *Br J Anaesth*. 2015;115(Suppl 2):ii68–74.

Chapter 11

Troubleshooting the Ventilated Patient



Patients in the emergency department are at high risk for deterioration after intubation. Understanding the common ways in which a patient can deteriorate, and having a systematic approach, is key to acting in these high-stress situations.

When the ventilator loses pressure, the *low pressure alarm* will go off. The differential for the low pressure alarm includes a break in the circuitry, anywhere from the ventilator to the lungs. Causes can include:

- A disconnection of the ventilator tubing and endotracheal tube (ETT).
- Increased patient effort—air hunger, “sucking in” during breaths.
- ETT displacement or extubation—the ETT tip may be right at the level of the vocal cords, and therefore, the patient is not yet desaturating.
- The ETT cuff may have a leak, allowing air to escape.
- Although less likely, a hole may also be present somewhere in the circuit.

The *high pressure alarm* similarly can arise from an issue anywhere from the patient to the ventilator.

- If the patient is fighting or “bucking” while on the mechanical ventilator, the peak airway pressure can rise.
- Elevation in autoPEEP.

- Any impedance to airflow, such as a mainstem intubation, bronchospasm, a pneumothorax, or a mucous/bloody plug in the ETT or airways, can also lead to elevated pressures.
- A disruption in the ETT, such as kinking or biting of the tube may be responsible.

An alarm on the vent should NEVER be ignored, and these alarms should be thought of as a pre-code situation. Ventilated patients are among the highest risk for deterioration in the ED, and they must be attended to promptly for both high pressure and low pressure alarms. It is important in such situations to assemble your team to fully assess the patient, including the assistance of a respirator therapist (RT) if available.

To quickly assess for issues, clinicians should recall the DOPES mnemonic, outlined in Fig. 11.1. Running quickly through this mnemonic will remind clinicians to consider common causes of deterioration while on the ventilator.

A separate, but related mnemonic, DOTTS, reminds clinicians how to respond to a ventilator alarm. Although the type of alarm (low pressure or high pressure) will influence the differential, the immediate actions are all the same. The first

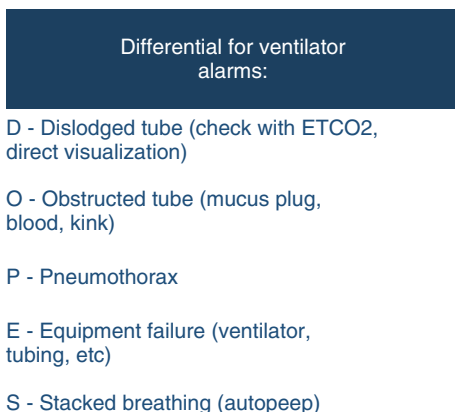


FIGURE 11.1 DOPES mnemonic for assessing cause of alarm or deterioration for an intubated patient

Approach to ventilator alarms:

D - Disconnect the patient from the ventilator, provide gentle chest pressure

O - Oxygen 100% by manual bagging, check compliance by squeezing the bag.

T - Tube position/function (pass a suction catheter)

T - Tweak the vent

S - Sonography (PTX; Mainstem intubation; plugging)

FIGURE 11.2 DOTTTS mnemonic for steps in assessing and treating the deterioration of an intubated patient

maneuver, disconnecting the patient from the ventilator immediately, takes much of the equipment out of the equation. The patient should then be connected to 100% oxygen via a bag, and ventilation provided by bagging. This can be diagnostic, allowing the clinician to assess if there is increased pressure or decreased pressure. The next maneuver is to pass a flexible suction catheter down the ETT, checking for resistance and patency of the tube. Suctioning, in the case of a mucous plug, may also be therapeutic. From here, information regarding the differential diagnosis should now be available. As indicated, the clinician can tweak the ventilator if autoPEEP is thought to be an issue, and perform a bedside US to assess for lung sliding if pneumothorax is suspected (Fig. 11.2).

Suggested Reading

1. Archambault PM, St-Onge M. Invasive and noninvasive ventilation in the emergency department. *Emerg Med Clin North Am.* 2012;30(2):421–49, ix.
2. Wood S, Winters ME. Care of the intubated emergency department patient. *J Emerg Med.* 2011;40(4):419–27.

Chapter 12

Case Studies in Mechanical Ventilation



Case 1

Thirty-four-year-old female presents to the ED with 1 day of feeling unwell, with myalgias, headache (HA), nausea, and fevers. The patient is in respiratory distress. Her vitals are: Temp: 36.6C (98F), HR 121, BP 89/45, SpO₂ 82% on room air.

1. What are the options for respiratory support for this patient? What are the risks and benefits of each?
2. You elect to intubate the patient. How would you set this patient's ventilator? What other information would you need to know?
3. What are your goals for ventilating this patient?
4. Does this patient have ARDS? How can you tell?
5. The ABG is 7.14/54/89 on 100%. How do you interpret this ABG?
6. Fig. 12.1 shows the patient's ventilator screen.
 - 6a. What is the TV?
 - 6b. What is the PEEP?
 - 6c. What is the PIP?
 - 6d. What is the Pplat?

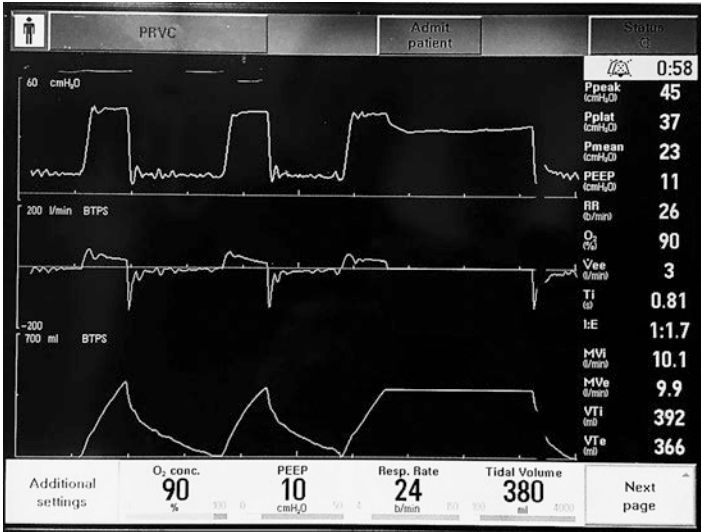


FIGURE 12.1 Ventilator screen for Case 1

7. What changes would you make to this ventilator?
8. While awaiting her bed in the ICU, the patient starts to desaturate. How do you assess this situation?

Case 2

Fifty-six-year-old male, long h/o 2 PPD smoking, suspected COPD, presents with wheezing, shortness of breath (SOB), and chest tightness. EMS gave him bronchodilators, but he arrives in the ED in extremis. Temp: 36.6C (98F), HR 115, BP 160/82, SpO₂ 87% while receiving an albuterol neb with oxygen.

1. What are the options for respiratory support for this patient? What are the risks and benefits of each? What are absolute and relative contraindications to these types of support?
2. How would you select ventilation settings for this scenario? What are your goals of respiratory support?

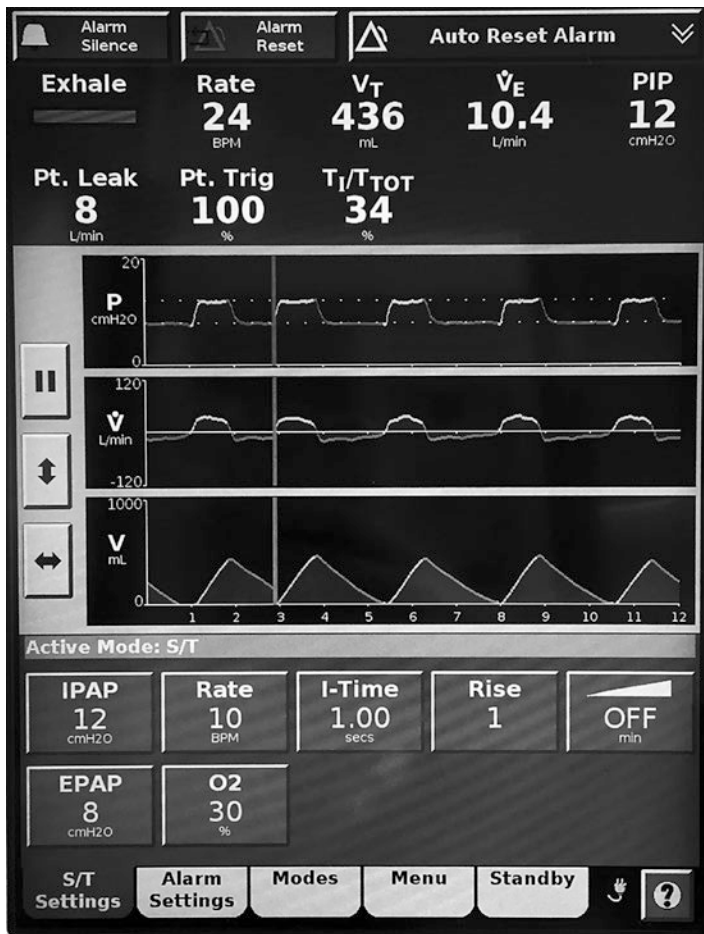


FIGURE 12.2 Ventilator screen for Case 2

3. How do you assess the adequacy of ventilation on noninvasive ventilation?
4. Figure 12.2 shows the patient's ventilator screen.
 - 4a. What is the IPAP?
 - 4b. What is the EPAP?

- 4c. What do each of those mean?
- 4d. What is the tidal volume?
- 4e. What is the minute ventilation?
5. The patient has an ABG of 7.25/75/68 on the BPAP. How would you interpret this ABG in this scenario?
6. The patient is awaiting transport to the ICU when the alarms in the room start to go off. The repeat vitals are: BP 85/45, HR 130, RR 30, SpO₂ 85%. How would you approach this emergency?

Case 3

Twenty-four-year-old male, long h/o poorly controlled asthma, presents to the ED with increasing SOB after an upper respiratory infection (URI). You use all the appropriate pharmacologic measures, but the patient is beginning to look fatigued and remains in distress.

1. What are the options for respiratory support for this patient? What are the risks and benefits of each?
2. You elect to intubate the patient. How would you set this patient's ventilator? What other information would you need to know?
3. Figure 12.3 shows the patient's ventilator screen.
 - 3a. What is the TV?
 - 3b. What is the PEEP?
 - 3c. What is the PIP?
 - 3d. What is the I:E ratio?
4. Do you think this patient has a compliance problem or a resistance problem? How could you tell?
5. Can you quantify air trapping?
6. What changes would you make to these ventilator settings?
7. While awaiting his ICU bed, the low-pressure alarm starts to go off on the ventilator. How do you address this?

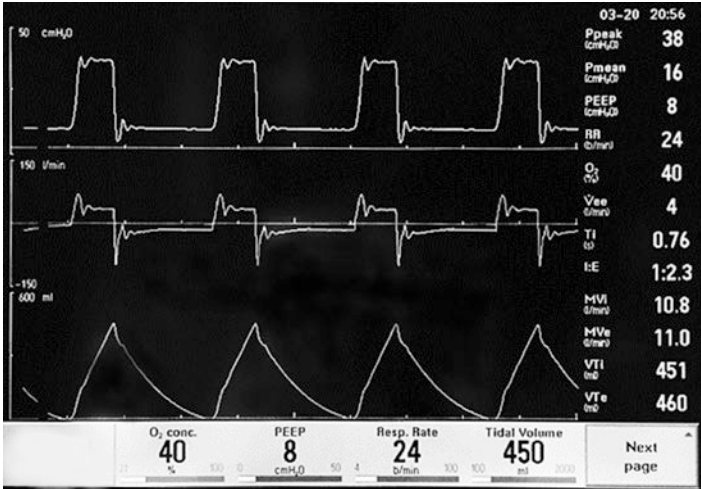


FIGURE 12.3 Ventilator screen for Case 3

Case 4

A twenty-eight-year-old male presents after a single-passenger MVC. He is obtunded, with gurgling respirations, GCS 5. He is intubated promptly upon arrival in the ED.

- The respiratory therapist asks you if she should put him on the ventilator now or if she should keep bagging the patient, since he is probably going to CT scan soon. What do you say? Why?
- How would you set the patient's ventilator when you do put him on the vent? What additional data would you like to know?
- Fig. 12.4 shows the ventilator screen.
 - What is the TV?
 - What is minute ventilation?
 - What is the PIP?
- His ABG returns as 7.54/28/225. What changes would you make to these vent settings?

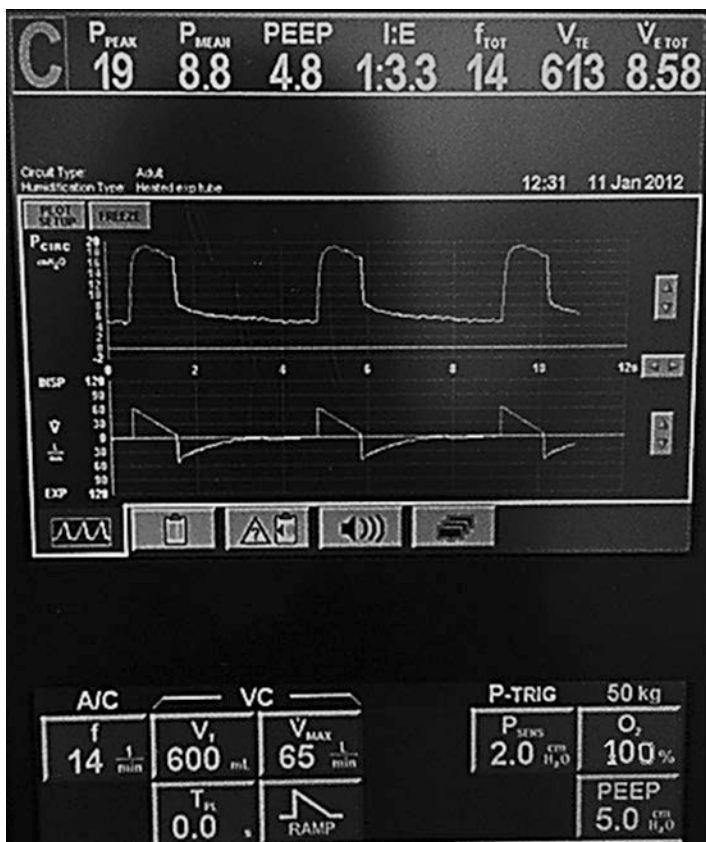


FIGURE 12.4 Ventilator screen for Case 4

- The patient's head CT shows a large subdural hematoma. Would that change your approach to your ventilator settings? Would you lower the PEEP?
- If this patient developed a hemothorax, what changes would you expect to see on the ventilator?

Case Study Answers

Case 1

1. There are three options for respiratory support for this patient.
 1. The patient could be trialed on high flow nasal cannula. The benefit of this is that it provides excellent noninvasive support for oxygenation. The downside is that if the patient develops shock, or any other form of instability, high flow nasal cannula will not be sufficient.
 2. Noninvasive positive pressure ventilation. This is an excellent way to oxygenate and ventilate a patient as well. However, if the patient has any alterations mental status or develops shock, the patient will need to be intubated.
 3. Intubation and mechanical ventilation. While this method has the downside of being the most invasive, for a patient in septic shock, this may be the most reasonable option.
2. This patient should be set on low tidal volume ventilation, with a goal of 6–8 mL/kg of predicted body weight. Her major issue is that she is hypoxemic, and therefore, she should have adequate PEEP support. The data point that is important is to know her height so that her predicted body weight can be calculated.
3. The goals for ventilating this patient are to maintain her on low tidal volume ventilation, keeping her plateau pressure less than 30. The PEEP should be set to maintain adequate oxygenation, trying to minimize the D recruitment that happens with sedation in a division. The FiO_2 should be decreased as soon as possible, targeting an oxygen saturation of 88–92%. This patient is appropriate for permissive hypercapnia.

4. The question stem does not provide enough data to determine if the patient has ARDS. To determine if the patient has ARDS, we would need to know the results of her chest x-ray to evaluate for bilateral infiltrates. Additionally, presumably based upon the question stem, this is not cardiogenic in nature; however, that cannot be known for certain without requiring more history. However, if this patient were to meet these requirements for ARDS, she would fall into the severe ARDS category.
5. The patient has poor oxygenation as indicated by her P/F ratio of 89 (89/1.0). She also has a combined metabolic and respiratory acidosis given that she has a mild hypercapnia at 54 mmHg but has a substantial acidemia at 7.14.
 - 6a. The TV is set at 380 mL. Her last inspiration was 392 mL, and her last exhalation was 366 mL.
 - 6b. 10
 - 6c. 45
 - 6d. 37
7. This is a challenging question because the patient is already presumably on a low tidal volume at 380. Her peak inspiratory pressure and plateau pressure are both very elevated. Increasing PEEP may help improve her compliance. If the patient has large areas of derecruited lung, performing a recruitment maneuver and increasing her PEEP may open previously atelectatic lung units and thereby improve compliance.

Case 2

1. For patient with COPD, the target oxygen levels are often 88–92%. Therefore, while this patient is certainly hypoxic, his increased work of breathing is a greater concern than his relatively mild to moderate hypoxemia. Although high flow nasal cannula may help with management of mild hypercapnia, and improving oxygenation can sometimes decrease the work of breathing, with the patient's

suspected COPD, noninvasive ventilatory support is likely a better option. Noninvasive ventilatory support has been shown to improve outcomes in patients with COPD. This patient could be intubated and mechanically ventilated, however, unless the patient has a contraindication, most patients with COPD should be trialed on bilevel positive pressure ventilation first. The absolute contraindications to high flow nasal cannula include airway compromise. The absolute contraindications to noninvasive ventilatory support are airway compromise, severely altered mental status, recent ENT/upper GI surgeries, small bowel obstruction, or other pathology that will put the patient at high risk for vomiting.

2. When initiating bilevel noninvasive support, one will often begin with relatively low settings of 10/5 cmH_2O . The patient's resultant tidal volume, respiratory rate, and overall comfort can then be reassessed. A blood gas should be checked approximately 15–30 min after initiation of support to ensure that the patient is trending toward improvement.
3. The noninvasive ventilator will provide a tidal volume and a minute ventilation just as with the patient on invasive mechanical ventilation. In addition to monitoring these values, monitoring the patient clinically, looking at the oxygen saturation, the respiratory rate, the work of breathing, the accessory muscle use, as well as checking blood gases are important to ensure the adequacy of ventilation.
- 4a. 12
- 4b. 8
- 4c. In BPAP, on noninvasive ventilation, the inhaled positive airway pressure (IPAP) is equivalent to the PIP in invasive ventilation. The expired positive airway pressure (EPAP) is equivalent to PEEP or CPAP. In this example, the EPAP of 8 is the baseline pressure. With every breath, the patient receives an additional 4 cmH_2O of support, for a total of 12 cmH_2O .
- 4d. 10.4 L/min.

5. The patient has a chronic respiratory acidosis with an acute respiratory acidosis superimposed. The patient also has some hypoxemia with a partial pressure of oxygen of 68 mmHg on 30% FiO₂.
6. The DOPES and DOTTS mnemonic devices are designed for patients who are intubated. However, similar concepts can apply to the patient on positive pressure ventilation. DOPES begins with displacement, which is not relevant in this scenario. However, obstruction, pneumothorax, equipment failure, and stacking provide a reasonable start for building a differential diagnosis. Similarly, the patient does not necessarily have to be disconnected and bagged with 100% oxygen; however, taking the patient briefly off the noninvasive ventilation, talking to the patient, assessing the patency of the airway, considering obstruction, and listening for bilateral breath sounds for a possible pneumothorax are all reasonable steps to complete within the first 30 s of assessing the patient.

Case 3

1. This case with the patient presenting in respiratory failure with severe asthma. The patient has reactive airways disease leading to an obstructive process. As oxygenation is not his primary issue, high flow nasal cannula is unlikely to be the best option. It is reasonable to try the patient on noninvasive positive pressure ventilation, with an understanding that if the patient does not respond promptly, he will require intubation. The downsides of noninvasive ventilation with asthma are that the patient is at risk for air trapping and cannot be heavily sedated with noninvasive. Additionally, if the patient is starting to fatigue or developing altered mental status, noninvasive ventilation is not the appropriate means of supporting the patient. The patient can be intubated and mechanically ventilated; however, this is also an area from with risk for a patient with asthma. Patient's heart high risk of air trapping with mechanical ventilation and must be aggressively treated and monitored.

2. The key principles of this patient's ventilator involved in maintaining a low respiratory rate, a low I:E ratio, and potentially a high flow rate. Of all the interventions, keeping a low respiratory rate is the most effective and giving the patient adequate time to excrete. Additionally, patients with asthma should be ventilated with low tidal volume ventilation to minimize the amount of gas that needs to be exhaled. Permissive hypercapnia or allowing the patient to have a mild to moderate respiratory acidosis is acceptable in a patient with asthma.
 - 3a. 450 mL
 - 3b. 8 cmH₂O
 - 3c. 38 cmH₂O
 - 3d. 1:2.3.
4. This patient likely has a resistance problem. The way to be certain is to check his pulmonary mechanics. By checking a plateau pressure, or checking an inspiratory hold, the clinician can determine the difference between his peak inspiratory pressure and his plateau pressure. If this difference is 5 cm of water or less, the patient has minimal issues with resistance. Conversely, if the patient has a high peak inspiratory pressure but a low plateau pressure, this indicates a significant issue with resistance. The inspiratory hold maneuver is not shown in Fig. 12.3; however, when it was checked, the patient had a plateau pressure of only 24. This indicates that the patient's issue was with resistance, not compliance.
5. Air trapping can be readily quantified at the bedside by performing an expiratory hold maneuver. This will give the autoPEEP. Although the expiratory hold maneuver is also not shown in Fig. 12.3, the autoPEEP in this scenario was 9 cmH₂O. This indicates that the patient has 9 cmH₂O of pressure trapped in his lungs due to inability to fully exhale. Looking closely at the patient's monitor, one can see that the patient is not fully exhaling as the wave form for flow never reaches the baseline before the next breath occurs. Recall that evaluating the wave form is a qualitative measure only and does not quantify the amount of air trapping.

6. This patient has elevated peak inspiratory pressure. In addition to providing aggressive care with continuous bronchodilators, steroids, magnesium, and any other medically appropriate interventions, the patient's ventilator should be adjusted. His respiratory rate is 24, which is likely far too high for a patient with asthma. It would be reasonable to drop the respiratory rate to 14 breaths per minute and reassess. Additionally, clinician could decrease the tidal volume to minimize the volume of gas the patient must exhale. Lastly, although some PEEP is always appropriate, the patient's PEEP could be decreased from 8 to 5 cmH₂O.
7. Although this patient is high risk for breath stacking, which would lead to a high-pressure alarm, the question stem indicates the patient has a low-pressure alarm. The mnemonic DOPES addresses etiologies leading to both low and high pressure alarms. Displacement and equipment failure are the two most likely causes of low-pressure alarm. In this scenario, the patient had coughed vigorously, and he had partially self-extubated leading to the low-pressure alarm sounding.

Case 4

1. Although the patient is likely to travel to the CT scanner soon, a patient with neurologic injury should be placed on a mechanical ventilator as soon as possible to minimize the risk of unintentional secondary injury. Use of a mechanical ventilators, including portable or transport ventilators for travel to radiology or interfacility transport, is important to ensure a consistent ventilation by means of providing consistent tidal volumes and respiratory rate. This monitoring and consistency minimize risk of inadvertent hyper- or hypoventilation.
2. This is a trauma patient with an apparent neurologic injury. It is appropriate to place the patient on low tidal volume

ventilation, with a tidal volume of 6–8 mL/kg of predicted body weight, and set the respiratory rate such that the patient has a starting minute ventilation of at least 7–8 L/m. Therefore, it is also important to know how tall the patient is such that his predicted body weight can be calculated. The patient should be given at least 5 of PEEP and the FiO_2 should be weaned as rapidly as possible, targeting an oxygen saturation of 95–99%. An ABG should be checked within 15–30 min after intubation to ensure a PaCO_2 of 35–40 and a PaO_2 of 80–100.

- 3a. The tidal volume is set for 600 mL and the patient is receiving 613 mLs.
- 3b. 8.58 L/min
- 3c. 19 cmH_2O .
4. This ABG indicates that the patient is being hyperventilated. His PaCO_2 is in the 20s, below the target of 35–40. Therefore, his respiratory rate or tidal volume should be decreased to decrease his overall minute ventilation. Additionally, the patient has hyperoxia with a PaO_2 of 225. This level is too high and could lead to secondary brain injury. The FiO_2 should be dropped substantially, likely to 60%, monitoring the SPO_2 to ensure that the patient does not become hypoxemic.
5. Even with a subdural hematoma, keeping the patient on a low amount of PEEP is appropriate. Patients with trauma and other neurologic injuries are at high risk for development of ARDS. PEEP is thought to help prevent ARDS to the extent that it prevents “atelectatrauma,” or the injury to alveoli that occur with repeated opening and snapping shut. A total of 5 cmH_2O of PEEP is an appropriate minimum for all patients.
6. If the patient developed a hemothorax, the patient’s compliance should go down. This would be manifested as an increase in peak inspiratory pressure and the plateau pressure.

Suggested Reading

1. Archambault PM, St-Onge M. Invasive and noninvasive ventilation in the emergency department. *Emerg Med Clin North Am.* 2012;30(2):421–49. ix
2. Spiegel R, Mallema H. Emergency Department Treatment of the Mechanically Ventilated Patient. *Emerg Med Clin North Am.* 2016;34(1):63–75.
3. Wright BJ. Lung-protective ventilation strategies and adjunctive treatments for the emergency medicine patient with acute respiratory failure. *Emerg Med Clin North Am.* 2014;32(4):871–87.
4. Mosier JM, Hypes C, Joshi R, et al. Ventilator strategies and rescue therapies for management of acute respiratory failure in the emergency department. *Ann Emerg Med.* 2015;66:529–41.

Chapter 13

Conclusions and Key Concepts



In summary, management of mechanical ventilation is an important procedure performed by emergency medicine clinicians to assist with oxygenation and ventilation, decrease the work of breathing, and help the patient meet their metabolic demands while critically ill. It is also important to recognize that mechanical ventilation can lead to several complications, which must be considered and minimized in all intubated patients. While no course can replace care from respiratory therapists and intensivists, having a shared vocabulary and understanding will allow for improved collaboration and care of these patients.

As a reminder, the goals of this text are to:

1. *Familiarize ED clinicians with common terms in mechanical ventilation.*
 - Many terms are used interchangeably in mechanical ventilation, and this leads to confusion. Select appropriate terms, and use them consistently.
 - Key concepts include tidal volume, respiratory rate, minute ventilation, PEEP, resistance, compliance, peak inspiratory pressure, plateau pressure, autoPEEP, and derecruitment.
 - Modes of ventilation are assist control (including volume control and pressure control, as well as pressure-regulated volume control), pressure support, and synchronized intermittent mandatory ventilation.

2. *Review the fundamental principles of pulmonary physiology relevant to mechanical ventilation.*

- There are two types of V/Q mismatch: Shunt is perfusion without ventilation, and dead space is ventilation without perfusion. The body tries to optimize V/Q matching by hypoxemic vasoconstriction.
- Resistance involves flow, but compliance is the distensibility of the entire system. The peak inspiratory pressure includes factors of resistance and compliance; however, plateau pressure only involves compliance.

3. *Discuss the basic principles of selecting ventilator settings.*

- Ventilator screens provide much data, but in general, the settings selected by the clinician appear along the bottom, and the patient's response appears along the top.
- Tidal volume should be selected for 6–8 mL/kg of predicted body weight, based upon height and sex. The respiratory rate should be selected to target a reasonable minute ventilation.
- PEEP should be set at minimum of 5 cmH₂O, and titrated higher as needed to correct for hypoxemia and derecruitment.
- Once the ventilator settings are selected the patient must be continuously reassessed, settings such be titrated based on ABG results, and peak inspiratory pressures and plateau pressures monitored to reduce harm.

4. *Develop strategies for caring for the ventilated ED patients with ARDS, asthma, COPD, and neurologic injury.*

- ARDS: The most important concepts in management of ARDS patients are low tidal volume ventilation while targeting a plateau pressure <30 cmH₂O. These patients might also require high levels of PEEP for improved oxygenation. Following ABGs on these patients is critical to determine the PaO₂ to FiO₂ ratios, and titrate their FiO₂ requirements. They may develop severe

hypoxemia, even after intubation, requiring recruitment maneuvers and neuromuscular blockade to assist with oxygenation. Additional techniques can be considered including proning, use of inhaled pulmonary vasodilators, and even ECMO.

- **Asthma:** These patients are at high risk of breath stacking, leading to autoPEEP. They should be ventilated with similarly low tidal volumes, a low respiratory, and low I:E ratios. They must be monitored for air trapping and autoPEEP checked with an expiratory hold.
- **COPD:** COPD patients often respond well to BPAP. If they require intubation, they should be treated very similarly to asthmatic patients. One difference is that patients with COPD may require higher levels of PEEP to match autoPEEP from collapsible distal airways.
- **Neurologic Injury:** These patients are at risk for secondary injury during and after intubation from hypoxemia as well as hypocapnia. Therefore, efforts should be made not to hyperventilate these patients, instead targeting normoxia and eucapnia.

5. *Assess and respond to emergencies during mechanical ventilation.*

- The differential diagnosis for ventilator alarms is DOPES: Displacement, Obstruction, Pneumothorax, Equipment Failure, and Stacked Breathing.
- The mnemonic for action is DOTTS: Disconnect, Oxygen (bagging), Tube Position, Tweak the Vent, Sonography.

Index

A

- Acute respiratory distress syndrome (ARDS), 2, 18, 19, 116, 117
 - definition, 69
 - MAPs, 70, 71
 - neuromuscular blockade, 75, 76
 - P/F ratio, 72
 - PEEP, 71
 - Pplat, 70, 72
 - predicted body weight, 70
 - recruitment maneuver, 73–75
- Air trapping, 31–33
- Airway resistance, 6
- Arterial alveolar gradient (A-a gradient), 28
- Arterial blood gas (ABG), 63, 84
- Assist control (AC), 11, 43, 44, 48, 62
- Asthma, 2, 80, 82
 - air trapping, 83
 - autoPEEP, 84
 - bronchial smooth muscles, 79
 - I:E ratio, 80, 82
 - intrinsic PEEP, 84
 - intubation, 80
 - neuromuscular blockade, 80
 - respiratory rate, 80, 81
 - total PEEP, 84
 - ventilator screen, 81, 82
- Atelectasis, 31
- AutoPEEP, 9, 19, 33, 51, 84

B

- Bilevel positive airway pressure (BPAP/BiPAP), 13, 38–40
- Breath-stacking, 31–33
- Bronchospasm, 29

C

- Chronic bronchitis, 84, 85
- Chronic obstructive pulmonary disease (COPD), 2, 21, 117
 - autoPEEP, 86–88
 - BPAP, 86
 - chronic bronchitis, 84, 85
 - emphysema, 84, 86
- Congestive heart failure (CHF), 38, 61
- Continuous positive airway pressure (CPAP), 12, 38

D

- Dead space, 20
- Derecruitment, 6
- Deterioration, 97
- DOPEs, 98, 110
- DOTTS, 98, 99, 110
- Driving pressure (ΔP), 9

E

Emphysema, 84, 86
 Endotracheal tube (ETT),
 29, 37, 97
 Expiratory peak airway pressure
 (EPAP), 39
 Expiratory time (eTime), 9, 65
 Expired positive airway pressure
 (EPAP), 109
 Extrinsic PEEP (ePEEP), 9

F

Fraction of inspired oxygen
 (FiO₂), 11

G

Gas exchange, 15–18

H

Hemothorax, 19
 High flow nasal cannula
 (HFNC), 35–37
 High pressure alarm, 97, 98
 Hypercapnia, 92
 Hyperinflation, 19
 Hyperoxia, 93
 Hypocapnia, 89, 93
 Hypoperfusion, 90
 Hypotension, 93
 Hypoxia, 93

I

Inhaled positive airway pressure
 (IPAP), 109
 Inhaled pulmonary
 vasodilators, 76
 Inspiratory pressure (iPAP), 13
 Inspiratory time (iTime), 9, 65
 Inspiratory to expiratory (I:E)
 ratio, 9, 47, 48, 93
 Intrinsic PEEP (iPEEP), 9, 51
 Invasive ventilation, 49
 assist control, 43, 44

compliance, 47
 pressure control, 48
 pressure support, 45
 pressure ventilation (*see*
 Pressure ventilation)
 PRVC, 48
 SIMV, 44, 46
 volume control, 47
 Ischemic stroke, 92, 93

L

Low pressure alarm, 97
 Lung compliance, 6

M

Mean airway pressure (MAP), 70
 Mechanical ventilation, 62–64,
 101–113, 116
 ABG, 66
 air trapping, 31–33
 airway resistance, 6
 ARDS, 116, 117
 assessment and response, 2,
 117
 asthma, 117
 atelectasis, 31
 blood gas, 3
 clinicians, 115
 compliance, 29–32
 conditional variables, 5
 control/target variables, 5
 conventional modes, 11–13
 COPD, 117
 critical care and respiratory
 therapy, 3
 critical illness, 66
 derecruitment, 6
 educational principle, 2
 emergent procedures, 1
 exhalation, 8
 experienced clinicians, 2
 hemodynamic perturbations, 66
 indications, 1
 initiation phase, 6, 7
 inspiratory phase, 7

- intubation and initiation, 1
 - issues, 28
 - lung compliance, 6
 - materials, 2
 - medicine practice, 1
 - neurologic injury, 117
 - patient history
 - COPD, wheezing and chest tightness, 102–104, 108–110
 - gurgling respirations, GCS 5, 105, 106, 112, 113
 - poorly controlled asthma, 104, 105, 110–112
 - respiratory distress, 101–102, 107, 108
 - PBW, 6
 - physiologic changes, 61, 62
 - plateau phase, 7
 - pulmonary physiology, 2, 116
 - recruitment, 6
 - resistance, 29, 30, 32
 - settings, 8–11
 - strategies for caring, 2
 - trigger, 5
 - unconventional modes, 13
 - VBG, 66
 - ventilator management, 1, 2
 - ventilator settings, 2, 64, 65
 - initial settings, 62
 - inspiratory flow and I:E ratio, 64, 65
 - mode, 62
 - PEEP, 63, 64
 - principles, 116
 - respiratory rate (RR), 63
 - tidal volume (TV), 63
 - volume, pressure/time cycled, 5
 - Minute ventilation, 11, 63
- N**
- Neurologic injury
 - intracranial hemorrhage, 93
 - ischemic stroke, 92, 93
 - status epilepticus, 94
 - traumatic brain injury, 89–92
 - Neuromuscular blockade, 75, 76
 - Noninvasive positive pressure ventilation (NIPPV), 12
 - BPAP, 38–40
 - contraindications, 37
 - CPAP, 38
 - definition, 37
 - Noninvasive respiratory support, 37–40
 - HFNC, 35, 36
 - NIPPV
 - BPAP, 38–40
 - CPAP, 38
 - definition, 37
 - oxygen therapy, 35
- O**
- Oxygenation
 - atelectasis, 27
 - hypoxemia, 17–26
 - hypoxic vasoconstriction, 25, 26
- P**
- PaO₂ to FiO₂ (P/F) ratio, 72
 - Partial support modes, 46
 - Patent foramen ovals (PFOs), 20
 - Peak airway pressure (PIP), 30, 31
 - Peak inspiratory flow, 9
 - Peak inspiratory pressure (PIP/P_{peak}), 8, 48, 49
 - Plateau pressure (P_{plat}), 8, 31, 49, 50, 70, 72
 - Pneumonia, 18
 - Pneumothorax, 19
 - Positive end-expiratory pressure (PEEP), 8, 38
 - Positive pressure ventilation (PPV), 61
 - Predicted body weight (PBW), 6
 - Pressure control (PC), 48

Pressure of oxygen (PAO_2), 28
Pressure regulated volume
 control (PRVC), 12, 48
Pressure support (PS), 12, 45
Pressure ventilation
 autoPEEP/iPEEP, 51
 PIP, 49
 Pplat, 49, 50

R

Recruitment maneuver, 6, 73–75
Resistance, 29
Respirator therapist (RT), 98
Respiratory rate (RR/f), 10

S

Shunts, 18–21
Status epilepticus, 94
Synchronized intermittent
 mandatory ventilation
 (SIMV), 11, 44

T

Tidal volume
 (TV/V_T), 9, 10
Traumatic brain injury,
 2, 89–92

V

Venous blood gases
 (VBG), 66
Ventilation and perfusion ration
 (V/Q matching), 25
Ventilator-induced lung
 injury, 70
Ventilator screens
 clinicians' comfort, 53
 evaluation, 53
 example, 54–58
 pressure, flow, and volume
 waveforms, 54
Ventilator tubing, 97
Volume control
 (VC), 47