Preface
INTRODUCTION

Many medical students’ first exposure to anesthesia happens in the hectic, often intimidating environment of the operating room. It is a challenging place to teach and learn.

“Understanding Anesthesiology: A Learner’s Guide” was created in an effort to enhance the learning experience in the clinical setting. The book introduces the reader to the fundamental concepts of anesthesia, including principles of practice both inside and outside of the operating room, at a level appropriate for the medical student or first-year (Anesthesia) resident. Residents in other programs such as Emergency Medicine or Internal Medicine, who require anesthesia experience as part of their training, will also find the guide helpful.

The guide is written at an introductory level with the aim of helping learners become oriented and functional in what might be a brief but intensive clinical experience. Those students requiring more comprehensive or detailed information should consult the standard anesthesia texts.

The author hopes that “Understanding Anesthesiology: A Learner’s Guide” succeeds not only in conveying facts but also in making our specialty approachable and appealing. I sincerely invite feedback on our efforts:

feedback@understandinganesthesiology.com

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Table of Contents
Regardless of which orientation you are in, by “pinching” the page, you will achieve the overall table of contents. In portrait mode, this appears as a typical table of contents, where sections for each chapter can be expanded and accessed from each chapter heading. In landscape mode, the table of contents appears as a series of white dots along the bottom of your ipad screen, with each dot representing a chapter. By clicking from dot to dot, you will access the specific chapter title page and see the pages for that chapter running along the bottom of the screen. The chapter title page displays the subsections of that chapter, each of which can be accessed with a touch.

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The glossary can be accessed in a second way. When you are reading the text, you will notice bolded words. The bolded words are glossary words, and touching on the word will open up its definition in a window. From that window, you can access the glossary, where you will be able to see related terms and access their definitions. Also, in the glossary, you will see the index for each term.

Index
When viewing a term in the glossary, you will find the index for that term below its definition. The index lists all the relevant references to that given term in the text, each of which you can access with a touch. Each of the index references has a definition embedded when you touch on the word where it appears in the text, even the ones that aren’t bolded.

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The figures and tables are all viewable in PDF format. Videos, interactive figures and animated slides are not viewable in PDF. The review quiz, because of its interactive element, does not appear with the PDF document. However, most of that content is available in standalone fashion on the website [www.understandinganesthesiology.com](http://www.understandinganesthesiology.com)

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ACKNOWLEDGEMENTS

Many individuals supported the production of this book.
Numerous publishers allowed the use of figures, as attributed in the text. The Wood Library-Museum of Anesthesiology provided the historic prints in Chapter 6.

Representatives from General Electric and the LMA Group of Companies were helpful in supplying the images used in the derivative figures seen in Interactive 2.1 and Figure 5 respectively.

Linda Onorato created and allowed the use of the outstanding original art seen in Figures 3 and 6, with digital mastery by Robert Barborini.

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This project would not have been possible without the efforts of Eric E. Brown, who was instrumental throughout the duration of the project, contributing to both the arduous work of formatting as well as creative visioning and problem-solving.

Karen Raymer
2012
While the contributors to this guide have made every effort to provide accurate and current drug information, readers are advised to verify the recommended dose, route and frequency of administration, and duration of action of drugs prior to administration. The details provided are of a pharmacologic nature only. They are not intended to guide the clinical aspects of how or when those drugs should be used. The treating physician, relying on knowledge and experience, determines the appropriate use and dose of a drug after careful consideration of their patient and patient’s circumstances. The creators and publisher of the guide assume no responsibility for personal injury.
THE ROLE OF THE ANESTHESIOLOGIST

Dr. Crawford Long administered the first anesthetic using an ether-saturated towel applied to his patient’s face on March 30, 1842, in the American state of Georgia. The surgical patient went on to have two small tumours successfully removed from his neck. Dr. Long received the world’s first anesthetic fee: $0.25.

Since then, the specialty of anesthesiology and the role of the anesthesiologist has grown at a rapid pace, particularly in the last several decades. In the operating room the anesthesiologist is responsible for the well-being of the patient undergoing any one of the hundreds of complex, invasive, surgical procedures being performed today. At the same time, the anesthesiologist must ensure optimal operating conditions for the surgeon. The development of new anesthetic agents (both inhaled and intravenous), regional techniques, sophisticated anesthetic machines, monitoring equipment and airway devices has made it possible to tailor the anesthetic technique to the individual patient.

Outside of the operating room, the anesthesiologist has a leading role in the management of acute pain in both surgical and obstetrical patients. As well, the anesthesiologist plays an important role in such diverse, multidisciplinary fields as chronic pain management, critical care and trauma resuscitation.
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In this chapter, you will learn about airway (anatomy, assessment and management) in order to understand the importance of the airway in the practice of anesthesiology. As well, you will develop an understanding of the fluid compartments of the body from which an approach to fluid management is developed.
Airway Management

In order to ensure adequate oxygenation and ventilation throughout the insults of anesthesia and surgery, the anesthesiologist must take active measures to maintain the *patency* of the airway as well as ensuring its protection from aspiration. A brief discussion of airway anatomy, assessment and management is given below.

**Airway Anatomy**

The *upper airway* refers to the nasal passages, oral cavity (teeth, tongue), pharynx (tonsils, uvula, *epiglottis*) and larynx. Although the *larynx* is the narrowest structure in the adult airway and a common site of obstruction, the upper airway can also become obstructed by the tongue, tonsils and epiglottis.

The *lower airway* begins below the level of the larynx. The lower airway is supported by numerous cartilaginous structures. The most prominent of these is the thyroid cartilage (Adam’s apple) which acts as a shield for the delicate laryngeal structures behind it. Below the larynx, at the level of the sixth cervical vertebra (C6), the *cricoid cartilage* forms the only complete circumferential ring in the airway. Below the cricoid, many horseshoe-shaped cartilaginous rings help maintain the rigid, pipe-like structure of the trachea. The trachea bifurcates at the level of the fourth thoracic vertebra (T4) where the *right mainstem bronchus* takes off at a much less acute angle than the left.

The airway is innervated by both sensory and motor fibres ([Table 1](#), [Figure 1](#), [Figure 2](#)). The purpose of the sensory fibres is to allow detection of foreign matter in the airway and to trigger the numerous protective responses designed to prevent aspiration. The swallowing mechanism is an example of such a response whereby the larynx moves up and under the epiglottis to ensure that the bolus of food does not enter the laryngeal inlet. The *cough reflex* is an attempt to clear the upper or lower airway of foreign matter and is also triggered by sensory input.
There are many different laryngeal muscles. Some adduct, while others abduct the cords. Some tense, while others relax the cords. With the exception of one, they are all supplied by the recurrent laryngeal nerve. The cricothyroid muscle, an adductor muscle, is supplied by the external branch of the superior laryngeal nerve.

**Table 1** Sensory innervation of the airway

<table>
<thead>
<tr>
<th>NERVE</th>
<th>AREA SUPPLIED</th>
</tr>
</thead>
<tbody>
<tr>
<td>lingual nerve</td>
<td>anterior 2/3 of tongue</td>
</tr>
<tr>
<td>glossopharyngeal nerve</td>
<td>posterior 1/3 of tongue</td>
</tr>
<tr>
<td>superior laryngeal nerve (internal branch)</td>
<td>epiglottis and larynx</td>
</tr>
<tr>
<td>recurrent laryngeal nerve</td>
<td>trachea, lower airways</td>
</tr>
</tbody>
</table>

**Figure 1** Nerve supply to the airway

This figure was published in Atlas of Regional Anesthesia, 3rd edition, David Brown, Copyright Elsevier (2006) and used with permission.

**Figure 2** Sensory innervation of the tongue

Airway Assessment

The anesthesiologist must always perform a thorough pre-operative **airway assessment**, regardless of the planned anesthetic technique. The purpose of the assessment is to identify potential difficulties with airway management and to determine the most appropriate approach. The airway is assessed by history, physical examination and occasionally, laboratory exams.

On history, one attempts to determine the presence of pathology that may affect the airway. Examples include arthritis, infection, tumors, trauma, morbid obesity, burns, congenital anomalies and previous head and neck surgery. As well, the anesthesiologist asks about symptoms suggestive of an airway disorder: dyspnea, hoarseness, stridor, sleep apnea. Finally, it is important to elicit a history of previous difficult intubation by reviewing previous anesthetic history and records.

The physical exam is focused towards the identification of anatomical features which may predict airway management difficulties. It is crucial to assess the ease of intubation. Traditional teaching maintains that exposure of the vocal cords and glottic opening by direct laryngoscopy requires the alignment of the oral, pharyngeal and laryngeal axes (Figure 3). The “**sniffing position**” optimizes the alignment of these axes and optimizes the anesthesiologist’s chance of achieving a laryngeal view.

An easy intubation can be anticipated if the patient is able to open his mouth widely, flex the lower cervical spine, extend the head at the atlanto-occipital joint and if the patient has enough anatomical space to allow a clear view. Each of these components should be assessed in every patient undergoing anesthesia.

**Figure 3** Axis alignment using the “sniffing position”

Original artwork by Linda Onorato. Digital mastery by Robert Barbortini. Used with permission of Linda Onorato.
• **Mouth opening**: Three fingerbreadths is considered adequate mouth opening. At this point in the exam, the anesthesiologist also observes the teeth for overbite, poor condition and the presence of dental prosthetics.

• **Neck motion**: The patient touches his chin to his chest and then looks up as far as possible. Normal range of motion is between 90 and 165 degrees.

• **Adequate space**: Ability to visualize the glottis is related to the size of the tongue relative to the size of the oral cavity as a large tongue can overshadow the larynx. The Mallampati classification (Table 2, Figure 4) assigns a score based on the structures visualized when the patient is sitting upright, with the head in a neutral position and the tongue protruding maximally. Class 1 corresponds well with an easy intubation. Class 4 corresponds well with a difficult intubation. Classes 2 and 3 less reliably predict ease of intubation. The **thyromental distance** is also an important indicator. The distance from the lower border of the mandible to the thyroid notch with the neck fully extended should be at least three to four fingerbreadths. A shorter distance may indicate that the oral-pharyngeal-laryngeal axis will be too acute to

**Table 2 Mallampati Classification**

<table>
<thead>
<tr>
<th>Class</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Soft palate, uvula, tonsillar pillars can be seen.</td>
</tr>
<tr>
<td>2</td>
<td>As above except tonsillar pillars not seen.</td>
</tr>
<tr>
<td>3</td>
<td>Only base of uvula is seen.</td>
</tr>
<tr>
<td>4</td>
<td>Only tongue and hard palate can be seen.</td>
</tr>
</tbody>
</table>

Figure 4 Mallampati classification

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achieve good visualization of the larynx. As well, a short thyromental distance may indicate inadequate “space” into which to displace the tongue during laryngoscopy.

Combining Mallampati classification with thyromental distance and other risk factors (morbid obesity, short, thick neck, protuberant teeth, retrognathic chin), will increase the likelihood of identifying a difficult airway. No assessment can completely rule out the possibility and so the clinician must always be prepared to manage a difficult airway.

Laboratory investigations of the airway are rarely indicated. In some specific settings, cervical spine x-rays, chest ray, flow-volume loops, computed tomography or magnetic resonance imaging may be required.

Airway Management

Airway patency and protection must be maintained at all times during anesthesia. This may be accomplished without any special maneuvers such as during regional anesthesia or conscious sedation. If the patient is deeply sedated, simple maneuvers may be required: jaw thrust, chin lift, oral airway (poorly tolerated if gag reflex is intact) or nasal airway (well tolerated but can cause epistaxis).

During general anesthesia (GA), more formal airway management is required. The three common airway techniques are:

- mask airway (airway supported manually or with oral airway)
- laryngeal mask airway (LMA)
- endotracheal intubation (nasal or oral)

The choice of airway technique depends on many factors:

- airway assessment
- risk of regurgitation and aspiration
- need for positive pressure ventilation
- surgical factors (location, duration, patient position, degree of muscle relaxation required)
A patient who is deemed to be at risk of aspiration requires that the airway be “protected” with a cuffed endotracheal tube regardless of the nature of the surgery. If the surgery requires a paralyzed patient, then in most cases the patient is intubated to allow mechanical ventilation.

**Mask Airway:** Bag mask ventilation may be used to assist or control ventilation during the initial stages of resuscitation or to pre-oxygenate a patient as a prelude to anesthetic induction and intubation. A mask airway may be used as the sole airway technique during inhalational anesthesia (with the patient breathing spontaneously) but it is only advisable for relatively short procedures as it “ties up” the anesthesiologist’s hands. It does not protect against aspiration or laryngospasm (closure of the cords in response to noxious stimuli at light planes of anesthesia). Upper airway obstruction may occur, particularly in obese patients or patients with very large tongues. In current practice, the use of a mask as a sole airway technique for anesthesia is rarely-seen although it may be used for very brief procedures in the pediatric patient.

**Laryngeal Mask Airway (LMA):** The LMA is an airway device that is a hybrid of the mask and the endotracheal tube. It is inserted blindly into the hypopharynx. When properly positioned with its cuff inflated, it sits above the larynx and seals the glottic opening. It is usually used for spontaneously breathing patients but positive pressure ventilation can be delivered through an LMA. The LMA does not protect against aspiration. Like an endotracheal tube, it frees up the anesthesiologist’s hands and allows surgical access to the head and neck area without interference. While airway obstruction due to laryngospasm is still a risk, the LMA prevents upper airway obstruction from the tongue or other soft tissues. The LMA also has a role to play in the failed intubation setting particularly when mask ventilation is difficult. The #3, #4 and #5 LMA are used in adults. Many modifications have followed the origi-
nal “classic” LMA including a design that facilitates blind endotracheal intubation through the LMA (Fastrach LMA™) and one that is specially designed for use with positive pressure ventilation with or without muscle relaxation (Proseal LMA™).

Endotracheal Intubation: There are 3 basic indications for intubation:

1. To provide a patent airway. An endotracheal tube (ETT) may be necessary to provide a patent airway as a result of either patient or surgical factors (or both). For example, an ETT is required to provide a patent airway when surgery involves the oral cavity (e.g. tonsillectomy, dental surgery). An ETT provides a patent airway when the patient must be in the prone position for spinal surgery. Airway pathology such as tumour or trauma may compromise patency, necessitating an ETT.

2. To protect the airway. Many factors predispose a patient to aspiration. ([Read more about risk of aspiration.](#)) A cuffed endotracheal tube, although not 100% reliable, is the best way to protect the airway of an anesthetized patient.

3. To facilitate positive pressure ventilation. Some surgical procedures, by their very nature, require that the patient be mechanically ventilated which is most effectively and safely achieved via an ETT. Mechanical ventilation is required when:

   - the surgery requires muscle relaxation (abdominal surgery, neurosurgery).
   - the surgery is of long duration such that respiratory muscles would become fatigued under anesthesia.
   - the surgery involves the thoracic cavity.

In rare cases, an ETT may be required to improve oxygenation in patients with critical pulmonary disease such as Acute Respiratory Distress Syndrome (ARDS), where 100% oxygen and positive end expiratory pressure (PEEP) may be needed.

While intubation is most commonly performed orally, in some settings nasotracheal intubation is preferable such as during intra-oral surgery or when long-term intubation is required. Nasotracheal intubation may be accomplished in a blind fashion (i.e. without performing laryngoscopy) in the emergency setting if the patient is breathing spontaneously.

Nasotracheal intubation is contraindicated in patients with coagulopathy, intranasal abnormalities, sinusitis, extensive facial fractures or basal skull fractures.

While there are myriad devices and techniques used to achieve intubation (oral or nasal), most often it is performed under direct vision using a laryngoscope to ex-
The blade is advanced into the **vallecula** which is the space between the base of the tongue and the epiglottis. Keeping the wrist stiff to avoid levering the blade, the laryngoscope is lifted to expose the vocal cords and glottic opening. The ETT is inserted under direct vision though the cords. A size 7.0 or 7.5 ETT is appropriate for oral intubation in the adult female and a size 8.0 or 8.5 is appropriate in the male. A full size smaller tube is used for nasal intubation.

**Movie 1.1** demonstrates the important technique to use when performing endotracheal intubation.

The view of the larynx on laryngoscopy varies greatly. A scale represented by the **“Cormack Lehane views”** allows anesthesiologists to grade and document the view that was obtained on direct laryngoscopy. Grade 1 indicates that the entire vocal aperture was visualized; grade 4 indicates that not even the epiglottis was viewed. **Figure 7** provides a realistic depiction of the range of what one might see when performing laryngoscopy.

**Movie 1.2** shows you the important anatomy to recognize on a routine intubation.

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**Movie 1.1 Intubation technique**

Video filmed and produced by Karen Raymer and Brian Colborne

pose the glottis. This technique is called **direct laryngoscopy**. The patient should first be placed in the “sniffing position” (**Figure 3**) in order to align the oral, pharyngeal and laryngeal axes. The curved **Macintosh blade** is most commonly used in adults. It is introduced into the right side of the mouth and used to sweep the tongue to the left (**Figure 6**).
Figure 6 View of upper airway on direct laryngoscopy

Original artwork by Linda Onorato, MD, FRCP(C); Digital mastery by Robert Barborini. Copyright Linda Onorato, used with permission of Linda Onorato.

Movie 1.2 Airway anatomy seen on intubation

Figure 7 Cormack Lehane views on direct laryngoscopy

Figure created by and used with permission from Kanal Medlej, M.D.; accessed from Resusroom.com
After intubation, correct placement of the ETT must be confirmed and esophageal intubation ruled out. The “gold standard” is direct visualization of the ETT situated between the vocal cords. The presence of a normal, stable end-tidal carbon dioxide (CO2) waveform on the capnograph confirms proper placement except in the cardiac arrest setting. Both sides of the chest and the epigastrium are auscultated for air entry. Vapour observed moving in and out of the ETT is supportive but not confirmative of correct tracheal placement.

If the ETT is advanced too far into the trachea, a right mainstem intubation will occur. This is detected by noting the absence of air entry on the left as well as by observing that the ETT has been advanced too far. The appropriate distance of ETT insertion, measured at the lips, is approximately 20 cm for an adult female and 22 cm for the adult male.

Complications may occur during laryngoscopy and intubation. Any of the upper airway structures may be traumatized from the laryngoscope blade or from the endotracheal tube itself. The most common complication is damage to teeth or dental prosthetics. It is imperative to perform laryngoscopy gently and not to persist with multiple attempts when difficulty is encountered. Hypertension, tachycardia, laryngospasm, raised intracranial pressure and bronchospasm may occur if airway manipulation is performed at an inadequate depth of anesthesia. Sore throat is the most common complication that presents post-extubation and is self-limited. Airway edema, sub-glottic stenosis, vocal cord paralysis, vocal cord granulomata and tracheomalacia are some of the more serious consequences that can occur and are more common after a prolonged period of intubation.
Airway Devices and Adjuncts

After performing a history and physical examination and understanding the nature of the planned procedure, the anesthesiologist decides on the anesthetic technique. If a general anesthetic is chosen, the anesthesiologist also decides whether endotracheal intubation is indicated or whether another airway device such as a LMA could be used instead.

When endotracheal intubation is planned, the technique used to achieve it depends in large part on the assessment of the patient’s airway. When intubation is expected to be routine, direct laryngoscopy is the most frequent approach. In settings where the airway management is not routine, then other techniques and adjuncts are used. Airway devices that can be used to achieve an airway (either as a primary approach or as a “rescue” method to use when direct laryngoscopy has failed) are categorized below.

- Methods for securing the upper airway only. These methods achieve what is sometimes termed a “non-invasive airway” and include the oral airway with mask; the LMA; and the King Laryngeal Tube™.
- Adjuncts for increasing the likelihood of achieving endotracheal intubation through direct laryngoscopy: alternate laryngoscope blades, endotracheal introducers (commonly referred to as gum elastic bougies), stylet.
- Methods of achieving endotracheal intubation using “indirect” visualization of the larynx: videolaryngoscope, (the Glidescope™, McGrath™); Bullard™ laryngoscope, fibreoptic bronchoscope.
- Methods of achieving endotracheal intubation in a “blind” fashion (without visualization of the larynx): blind nasal intubation, lighted stylet, retrograde intubation, Fastrach LMA™.
The Difficult Airway

Airway mismanagement is a leading cause of anesthetic morbidity and mortality and accounts for close to half of all serious complications. The best way to prevent complications of airway management is to be prepared. Anticipation of the difficult airway (or difficult intubation) and formulation of a plan to manage it when it occurs, saves lives.

Anticipated difficult intubation: The use of an alternate anesthetic technique (regional or local) may be the most practical approach. If a general anesthetic is chosen, then airway topicalization and awake intubation (with fiberoptic bronchoscope) is the preferred technique. In pediatric patients, neither a regional technique nor an awake intubation is feasible. In this case, induction of anesthesia with an inhaled agent such that the patient retains spontaneous respiration is the safest approach. Efforts are undertaken to secure the airway once the child is anesthetized.

Unanticipated difficult intubation, able to ventilate by mask: In this situation, one calls for help, repositions the patient and reattempts laryngoscopy. The guiding principle is to avoid multiple repeated attempts which can lead to airway trauma and edema resulting in the loss of the ability to ventilate the patient. During the subsequent attempts at intubation, the anesthesiologist considers using alternate airway techniques (see section on adjuncts) or awakening the patient to proceed with an awake intubation.

Unanticipated difficult intubation, unable to ventilate by mask: This is an emergency situation. One calls for help and attempts to insert an LMA which is likely to facilitate ventilation even when mask ventilation has failed. If an airway is not achievable by non-surgical means, then a surgical airway (either needle cricothyrotomy or tracheostomy) must not be delayed.

When a difficult airway is encountered, the anesthesiologist must respond quickly and decisively. As in many clinical situations which occur infrequently but are associated with high rates of morbidity and mortality, the management of the difficult airway is improved by following well-developed algorithms. The American Society of Anesthesiologists has published a “Difficult Airway Algorithm” which is widely accepted as standard of care. The algorithm is described in a lengthy document such that a full explanation is beyond the scope of this manual. The algorithm, as well as other experts’ interpretations, are readily available on the internet.
The goal of fluid management is the maintenance or restoration of adequate organ perfusion and tissue oxygenation. The ultimate consequence of inadequate fluid management is hypovolemic shock.

**Fluid Requirements**

Peri-operative fluid management must take into account the pre-operative deficit, ongoing maintenance requirements and intra-operative losses (blood loss, third space loss).

**Pre-operative Deficit**: The *pre-operative fluid deficit* equals basal fluid requirement (hourly maintenance x hours fasting) plus other losses that may have occurred during the pre-operative period.

**Maintenance fluid requirements** correlate best with lean body mass and body surface area. To calculate maintenance, use the “4/2/1 rule”:

- First 10 kilograms (i.e. 0-10 kg): 4 cc/kg/hr
- Next 10 kilograms (i.e. 11-20 kg): 2 cc/kg/hr
- All remaining kilograms over 20 kg: 1 cc/kg/hr

For example, a 60 kg woman fasting for 8 hours:

\[
\begin{align*}
10 \text{ kg} & \times 4 \text{ cc/kg/hr} = 40 \text{ cc/hr} \\
10 \text{ kg} & \times 2 \text{ cc/kg/hr} = 20 \text{ cc/hr} \\
+ & \quad 40 \text{ kg} \times 1 \text{ cc/kg/hr} = 40 \text{ cc/hr} \\
& \quad = 100 \text{ cc/hr} \times 8 \text{ hr} \\
& \quad = 800 \text{ cc}
\end{align*}
\]

Therefore, the pre-operative deficit (excluding other losses) is 800 cc.

“Other losses” (including fluid lost through sweating, vomiting, diarrhea and nasogastric drainage) are more difficult to estimate. In the febrile patient, maintenance requirements are increased by 10% per degree Celsius elevation in temperature.

As a rule, half of the deficit should be corrected prior to induction and the remainder replaced intra-operatively. However, if the pre-operative deficit is greater than 50% of the estimated blood
volume, then the surgery should be delayed, if possible, to allow for more complete resuscitation.

**Intra-operative losses**: Blood loss is usually underestimated. It is assessed by visually inspecting blood in suction bottles, on the drapes and on the floor. Sponges can be weighed (1 gram = 1 cc blood), subtracting the known dry weight of the sponge. **Third space loss** refers to the loss of plasma fluid into the interstitial space as a result of tissue trauma and can be estimated based on the nature of the surgery:

- 2-5 cc/kg/hr for minimal surgical trauma (orthopedic surgery)
- 5-10 cc/kg/hr for moderate surgical trauma (bowel resection)
- 10-15 cc/kg/hr for major surgical trauma (abdominal aortic aneurysm repair)

These are all crude estimates of fluid requirements. Adequacy of replacement is best judged by the patient’s response to therapy. Urine output greater than 1.0 cc/kg/hr is a reassuring indicator of adequate organ perfusion. Hemodynamic stability, oxygenation, pH and central venous pressures are other indicators of volume status, but may be affected by many other factors. **Figure 8** depicts the holistic approach to assessing intra-operative blood loss.

**Figure 8** Assessment of intra-operative fluid status

---

*This figure was published in “Anesthesia for Thoracic Surgery”, Jonathan Benumonf, Copyright Elsevier (1987). Used with permission of Elsevier.*
Assessment of Fluid Status

Fluid status is assessed by history, physical exam and laboratory exam. Thorough history will reveal losses of blood, urine, vomit, diarrhea and sweat. As well, the patient is questioned regarding symptoms of hypovolemia, such as thirst and dizziness.

On physical exam, vital signs, including any orthostatic changes in vital signs, are measured. A decrease in pulse pressure and decreased urine output are two of the most reliable early signs of hypovolemia. Poor capillary refill and cutaneous vasoconstriction indicate compromised tissue perfusion. Severely depleted patients may present in shock (Table 3).

Hemoglobin, sodium, urea and creatinine levels may show the concentration effect which occurs in uncorrected dehydration. When blood loss occurs, hemoglobin and hematocrit levels remain unchanged until intravascular volume has been restored with non-blood containing solutions. Therefore, only after euvolemia has been restored is the hemoglobin level a useful guide for transfusion. Lactic acidosis is a late sign of impaired tissue perfusion.

<table>
<thead>
<tr>
<th>Table 3 Classification of hemorrhagic shock in a 70 kg person</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CLASS</strong></td>
</tr>
<tr>
<td>BLOOD LOSS (cc)</td>
</tr>
<tr>
<td>BLOOD LOSS (%)</td>
</tr>
<tr>
<td>PULSE RATE</td>
</tr>
<tr>
<td>BLOOD PRESSURE</td>
</tr>
<tr>
<td>PULSE PRESSURE</td>
</tr>
<tr>
<td>RESPIRATORY RATE</td>
</tr>
<tr>
<td>URINE OUTPUT</td>
</tr>
<tr>
<td>CNS</td>
</tr>
<tr>
<td>FLUID REQUIRED</td>
</tr>
</tbody>
</table>
**Vascular Access**

Peripheral venous access is the quickest, simplest and safest method of obtaining vascular access. The upper limb is used most commonly, either at the hand or **ante-cubital fossa** (cephalic and basilic veins). The lower limb can be used if necessary, the most successful site here being the saphenous vein, located 1 cm anterior and superior to the medial malleolus.

Flow through a tube is directly proportional to the pressure drop across the tube and inversely proportional to resistance.

\[
\text{Flow } \propto \frac{\text{pressure drop}}{\text{resistance}}
\]

Resistance is directly proportional to length and inversely proportional to radius to the fourth power.

\[
\text{Resistance } \propto \frac{\text{length}}{\text{radius}^4}
\]

From these equations, we can understand how the anesthesiologist achieves rapid administration of fluids. Pressure drop is achieved by using rapid infusers that apply a squeeze to the fluid, usually with an air-filled bladder. A cannula that is of a greater radius makes a significant impact on flow; to a lesser extent, a shorter cannula allows greater flow than a longer cannula of equivalent bore.

For example, a 16 gauge cannula will allow greater flow (i.e. faster resuscitation) than a (smaller) 18 gauge cannula. Likewise, a 14 gauge peripheral IV cannula will allow greater flow than an equivalent caliber central line, which is, by necessity, significantly longer. From a practical perspective, a 16 gauge cannula is the smallest size which allows rapid administration of blood products.

Central venous access is indicated when peripheral venous access is inadequate for fluid resuscitation, or when central pressure monitoring is required. The internal jugular vein is the most common site used intraoperatively. The external jugular is also useful, but can be technically difficult in some patients due to the presence of valves. The subclavian site is associated with an increased risk of **pneumothorax**, while the femoral site is associated with an increased risk of infection, embolism and thrombosis. Multiorifaced, 6 c.m., 14 gauge catheters are the most commonly used central lines. Wide bore “introducers” (for example, the 8.5 French Arrow CV Introducer®) are also commonly used for central venous access.

There are many potential complications of central venous cannulation. They include arterial puncture, hemorrhage, pneumothorax, thoracic duct injury, neural in-
Types of Fluids

Fluids can be divided into two broad categories: crystalloids and colloids. **Crystalloids** are solutions of simple inorganic or organic salts and distribute to varying extents throughout the body water. Examples include Ringer’s Lactate (R/L), 0.9% saline (N/S) and 5% dextrose in water (D5W). Sodium chloride, a common constituent of crystalloid solutions, distributes throughout the entire extracellular space. Glucose distributes throughout the entire body water (extracellular and intracellular spaces). Whatever the active solute, water, the ubiquitous solvent, will move across membranes to maintain osmotic equilibrium.

**Colloids** are suspensions of protein or other complex organic particles. These particles cannot diffuse across capillary membranes and so remain trapped within the intravascular space. Examples of colloids are albumin (5%, 25%), hydroxyethyl starches (Pentaspan®, Voluven®, red cell concentrates, platelets, and plasma.

The partitioning throughout the body’s compartments of some of the various types of fluids for administration is summarized in Table 4 and illustrated in the animated slides, Interactive 1.1.

Normal saline or Ringer’s lactate are the preferred crystalloids for intra-operative fluid administration and resuscitation, as they provide more intravascular volume...
expansion than D5W or 2/3:1/3. Because of the partitioning in the extracellular compartment, they must be given in a 3-4:1 ratio to the estimated blood loss. Administration of large amounts of N/S results in metabolic acidosis and should be avoided. R/L contains 4 meq/L potassium, and should be avoided in patients with renal failure. Glucose-containing solutions should only be used for specific indications (such as to maintain stable glucose levels in patient with diabetes mellitus or hepatic disease), and should be based on known glucose requirements. Finally, the half-life of crystalloid redistribution is only 15-30 minutes, so it must be given at a rate that accounts for its extravasation from the intravascular space.

Colloids replace blood loss in a 1:1 ratio, assuming normal membrane permeability. The use of colloids is generally reserved for cases where greater than 20% of the blood volume needs to be replaced or when the conse-

Table 4 Partitioning of various intravenous fluid solutions

<table>
<thead>
<tr>
<th>TYPE OF FLUID</th>
<th>ECF=1/3 TBW</th>
<th>ICF = 2/3 TBW</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>IVF=1/4 ECF</td>
<td>IF=3/4 ECF</td>
</tr>
<tr>
<td>1000 cc D5W</td>
<td>83 cc</td>
<td>250 cc</td>
</tr>
<tr>
<td>1000 cc 2/3 :1/3</td>
<td>139 cc</td>
<td>417 cc</td>
</tr>
<tr>
<td>1000 cc N/S OR R/L</td>
<td>250 cc</td>
<td>750 cc</td>
</tr>
<tr>
<td>500 cc 5% ALBUMIN</td>
<td>500 cc</td>
<td>0</td>
</tr>
<tr>
<td>100 cc 25% ALBUMIN</td>
<td>500 cc</td>
<td>-400 cc</td>
</tr>
<tr>
<td>500 cc PENTASPAN</td>
<td>750 cc</td>
<td>-250 cc</td>
</tr>
<tr>
<td>1 UNIT RCC</td>
<td>400 cc</td>
<td>0</td>
</tr>
</tbody>
</table>

TBW = total body water, ECF = extracellular fluid, ICF = intracellular fluid, IVF = intravascular fluid, IF = interstitial fluid, RCC = red cell concentrates.
quences of the interstitial edema (which might occur with crystalloid administration) are serious (e.g. cerebral edema).

**Blood products** are administered for specific indications. Red cell concentrates (RCC) are given to maintain or restore oxygen carrying capacity. As hemoglobin (Hb) concentration falls, oxygen delivery is preserved by compensatory mechanisms: shifting the oxyhemoglobin dissociation curve to the right and increasing cardiac output (via an increase in heart rate and contractility). When these compensations are inadequate or detrimental, RCC should be transfused. General indications for the transfusion of blood products are outlined in Table 5.

A patient with Class 3 or 4 hemorrhagic shock (Table 3) should be transfused immediately. For the slow but steady blood loss which occurs during many types of surgery, the lowest allowable hemoglobin, the “transfusion trigger”, is determined on an individual basis. Healthy patients can tolerate Hb levels that are approximately \( \frac{1}{2} \) of normal (60-70 g/L). Compensations may be inadequate in patients with pulmonary, cardiac or cerebrovascular disease. Compensation may be harmful in patients with certain types of heart disease such as severe coronary artery disease or aortic stenosis. These patients should be transfused to relatively higher Hb levels (80-100 g/L).

Once the lowest allowable hemoglobin has been determined, then the allowable blood loss (ABL) can be calculated as follows:

\[
\frac{(Hb_{initial} - Hb_{allowable}) \times EBV}{Hb_{initial}}
\]

Estimated blood volume (EBV) is approximately 60-70 mL/kg in the adult. When blood loss approaches estimated ABL, the anesthesiologist confirms the current Hb and considers transfusing.

Transfusion of plasma, platelets or cryoprecipitate is indicated only for the correction of defective clotting and is not indicated for volume resuscitation. Impaired clotting may be observed or anticipated in a given clinical scenario. For example, after one blood volume of RCC has been transfused (6-12 units in an adult), coagulopathy is likely on a dilutional basis and transfusion of platelets and plasma will be required. Prolonged clotting times or thrombocytopenia alone, without clinical evidence of bleeding, are insufficient indications for transfusion.

Risks and benefits of transfusion should be explained to patients undergoing procedures likely to result in significant blood loss. Consent for transfusion should be obtained whenever possible.
Complications of transfusion are numerous and are generally categorized by acuity: early and late. Early complications that can occur when significant volumes of blood are transfused include hypothermia, hyperkalemia and hypocalcemia. With massive transfusion, lung injury may occur.

Transfusion reactions can occur with just a single unit of transfused blood (due to ABO incompatibility). The most common cause of transfusion reaction is clerical error, underscoring the need for careful adherence to safety procedures by all members of the healthcare team.

A more complete discussion of the indications and complications of the various blood products is beyond the scope of this manual. Many excellent reviews on the subject can be found in the current anesthesia literature.

<table>
<thead>
<tr>
<th>BLOOD PRODUCT</th>
<th>DEFICIT</th>
</tr>
</thead>
<tbody>
<tr>
<td>red cell concentrates</td>
<td>oxygen-carrying capacity</td>
</tr>
<tr>
<td>platelets</td>
<td>platelet function (quality or quantity)</td>
</tr>
<tr>
<td>fresh frozen plasma</td>
<td>clotting factor deficits</td>
</tr>
<tr>
<td>cryoprecipitate</td>
<td>fibrinogen</td>
</tr>
<tr>
<td>albumin</td>
<td>low protein or colloid volume replacement</td>
</tr>
<tr>
<td>factor concentrates</td>
<td>single clotting factor deficit (often hereditary)</td>
</tr>
</tbody>
</table>
Question 1 of 17
ALL BUT ONE of the following are predictors of a difficult intubation. Indicate the exception:

- A. Mouth opening of only 3 fingerbreadths
- B. Short, thick neck with limited mobility
- C. Mallampati Class IV
- D. Thyromental distance of 2 fingerbreadths

Check Answer
In this chapter, you will learn how the anesthesiologist assesses a patient who is scheduled to undergo anesthesia and surgery, and how the goal of risk modification is achieved through that process. As well, you will be introduced to the equipment required for the safe delivery of anesthesia: the anesthetic machine and monitors.
Pre-operative Visit

The purpose of the pre-operative assessment is three-fold:

- To review the medical and psychological status of the patient.
- To identify factors which may impact on the peri-operative course, to take measures to optimize those factors where possible, and to delay surgery if necessary. If the patient’s medical condition cannot be altered, then one can take other measures to attempt to reduce risk: substitute a lower-risk surgical procedure, modify the anesthetic technique, intensify the peri-operative monitoring or cancel the surgery altogether.
- To inform patient, alleviate anxiety and establish rapport.

This evaluation takes the form of a directed history, physical examination and laboratory exam. On history, the anesthesiologist attempts to elicit symptoms of cardiac or respiratory disease as well as a history of any other major medical illnesses, past or present. Hepatic or renal disease may impact on metabolism and excretion of anesthetic agents, fluid balance and coagulation status. The patient’s medications are reviewed including any history of adverse drug reactions. The patient’s and their relative’s previous anesthetic experience is reviewed.

The physical examination focuses on the cardiac and respiratory (including airway) systems. Recording baseline vital signs is important, as is detecting any unstable, potentially reversible conditions such as congestive heart failure or bronchospasm. The airway is assessed for ease of intubation.

Routine pre-op laboratory investigations have not been shown to improve patient outcome. Therefore, laboratory studies are ordered only as indicated, according to the medical status of the patient and the nature of the planned surgery. Studies are rarely ordered to establish a “baseline” but rather to detect abnormalities that require correction prior to surgery. The traditional “CBC and urinalysis” is no longer required in healthy patients having minor surgery. An electrocardiogram (ECG) is ordered on patients who
are known to have cardiac disease or in whom risk factors (including age) are present. Routine pre-operative chest x-rays are not required prior to most procedures.

The anesthesiologist will commonly assign an “**ASA class**” (Table 6) to the patient. The ASA (American Society of Anesthesiologists) classification was defined in the 1940’s as an attempt to identify operative risk. As the patient’s underlying health is the most important determinant of peri-operative risk, the ASA class does correlate (somewhat) to overall peri-operative risk. Though it does not lend itself to inter-rater reliability, it is an accepted method of communicating the overall physical condition of the patient and the learner should become accustomed to applying this scale to the patients he or she encounters.

<table>
<thead>
<tr>
<th>ASA CLASS</th>
<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>A normal healthy patient in need of surgery for a localized condition.</td>
</tr>
<tr>
<td>2</td>
<td>A patient with mild to moderate systemic disease; examples include controlled hypertension, mild asthma.</td>
</tr>
<tr>
<td>3</td>
<td>A patient with severe systemic disease; examples include complicated diabetes, uncontrolled hypertension, stable angina.</td>
</tr>
<tr>
<td>4</td>
<td>A patient with life-threatening systemic disease; examples include renal failure or unstable angina.</td>
</tr>
<tr>
<td>5</td>
<td>A moribund patient who is not expected to survive 24 hours with or without the operation; examples include a patient with a ruptured abdominal aortic aneurysm in profound hypovolemic shock.</td>
</tr>
</tbody>
</table>
**NPO Status**

The induction of anesthesia abolishes the normal laryngeal reflexes that prevent inhalation ("aspiration") of stomach contents. Due to gastric, biliary and pancreatic secretions (which are present even during fasting), a stomach is never “empty”. **NPO (nil per os)** indicates the restriction of oral intake for a period of time prior to surgery, minimizing the volume, acidity and solidity of stomach contents. Such measures reduce both the risk of aspiration occurring as well as the severity of pneumonitis, should an aspiration event occur.

For elective surgery, patients should not have solid food for 8 hours prior to anesthesia. Traditionally, patients were ordered to refrain from all fluids for the 8 hour pre-operative period as well. However, more recent studies have shown that the time of the last (clear) fluid intake bears little relation to the volume of gastric contents present at the induction of anesthesia. Thus, most institutions are allowing unrestricted intake of clear fluids until 2-4 hours prior to scheduled surgery. Guidelines for pediatric patients vary from institution to institution but generally are more liberal than in the adult population. For example, infants may be allowed breast milk up to 4 hours pre-operatively and formula up to 6 hours pre-operatively.

It is important to recognize that some patients remain at risk for aspiration despite strict application of NPO guidelines. Known risk factors are outlined in **Table 7**.

When it is possible to identify these patients pre-operatively, measures can be taken to reduce their risk of aspiration syndrome. Firstly, pre-medication can be given to increase gastric motility (metoclopramide) or to decrease gastric acidity (ranitidine or sodium citrate). Risk can also be reduced through careful airway management that may include the use of the **Sellick Maneuver** on its own or as part of a rapid sequence induction.

**Table 7** Risk factors for aspiration

<table>
<thead>
<tr>
<th>RISK FACTORS FOR ASPIRATION</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Gastroesophageal reflux</td>
</tr>
<tr>
<td>• Pregnancy</td>
</tr>
<tr>
<td>• Trauma</td>
</tr>
<tr>
<td>• Diabetes Mellitus</td>
</tr>
<tr>
<td>• Recent oral intake</td>
</tr>
<tr>
<td>• Bowel obstruction</td>
</tr>
<tr>
<td>• Intra-abdominal pathology</td>
</tr>
<tr>
<td>• Obesity</td>
</tr>
</tbody>
</table>
Premedication

Pre-medication can include medication that the patient takes routinely as well as medication that may be prescribed specifically for the pre-operative period. Generally speaking, patients should be given their usual medication on the morning of surgery with a sip of water. It is particularly important that patients receive their usual cardiac and antihypertensive medications pre-operatively. Discontinuation of beta-blockers, calcium channel blockers, nitrates or alpha-2 agonists (clonidine) can lead to rebound hypertension or angina. Similarly, most medications taken for chronic disease should be continued on the morning of surgery as well as throughout the peri-operative period. This is particularly important for most antidepressants, thyroid replacement and anticonvulsants.

There are certain medications that may need to be discontinued in the pre-operative period. Examples include monoamine oxidase inhibitors and anticoagulants. Patients on platelet inhibitors such as aspirin represent a special group of patients who must be considered on an individual basis such that the risk of stopping the aspirin is weighed against the risk of surgical-site bleeding. For example, a patient who is on aspirin because of the recent insertion of a coronary stent must receive their aspirin throughout the peri-operative period. On the other hand, if the patient is on aspirin for primary prevention then it is usually discontinued a full week before surgery to allow return of normal platelet function.

Some medications are ordered specifically for the pre-operative period. Examples include anxiolytics, antibiotics, bronchodilators, anti-anginal medication and anti-emetics. Beta blockers have been used to reduce the incidence of cardiac morbidity and mortality in high-risk patients undergoing high-risk procedures, although the impact of this intervention is not yet fully understood.

Currently, pre-operative sedation is used less frequently than it has been in the past as it can delay awakening at the end of anesthesia. A delayed recovery is particularly undesirable in the outpatient surgical population where a return of cognitive function is required prior to discharge home. Furthermore, a pre-operative visit has been shown to be at least as effective as pharmacologic means in allaying anxiety in surgical patients. Nonetheless, there is a role for pre-operative sedation in very anxious patients or in those for whom anxiety would be deleterious, such as the cardiac patient.

For most types of surgery, antibiotics are ordered pre-operatively to reduce the incidence of wound infection. Antibiotics may also be ordered to reduce the risk of bacterial endocarditis in at-risk patients though the
current recommendations from the American Heart Association are much more restrictive than they have been in the past.

As discussed, aspiration prophylaxis may be ordered in high risk patients. This includes agents which decrease the volume and/or acidity of gastric secretions (ranitidine, sodium citrate) as well as agents which increase gastric emptying (metoclopramide).

A history of systemic steroid use may require the delivery of a peri-operative course of steroids in order to avoid the consequences of adrenal suppression which may present as an Addisonian crisis. Adrenal suppression occurs when a patient receives long term exogenous steroids in daily dose equal to or greater than 10 mg. Once adrenal suppression has occurred, the adrenal gland takes approximately 3 months to recover function (after steroid discontinuation). Therefore, steroid supplementation is required for patients who are currently on exogenous steroids or have discontinued a long term course in the past three months. The amount and duration of supplemental steroid coverage required depends on the invasiveness of the surgery. For minor surgery, a single dose of hydrocortisone (25 mg) suffices, while for major surgery, the patient requires 100 mg of hydrocortisone daily for 2-3 days.
Anesthetic Equipment and Monitoring

In This Section
1. The Anesthetic Machine
2. Monitoring

The Anesthetic Machine
The purpose of the anesthetic machine is to deliver gases to the patient in precise, known concentrations. Although the anesthetic machine has evolved substantially over the years, the essential features have remained remarkably constant. Some of the important components of a modern anesthetic machine are depicted in Interactive 2. 1. (Tap the labels for a close-up view as well as a brief description of each component.)

Gases (oxygen, air and nitrous oxide) come from pipelines entering the operating room through the wall (Figure 9). Tanks on the back of the anesthetic machine provide an alternate source of those gases should the wall supply fail. Although 100% oxygen can be delivered to the patient, usually a mixture of oxygen (with air or nitrous oxide) is selected. The relative concentrations of the gases to be delivered are controlled by flowmeters (one flowmeter for each gas) found on the left hand side of the anesthetic machine.

The anesthetic machine also allows the delivery of a precise concentration of volatile agent. The volatile anesthetic gases, such as sevoflurane and desflurane, are contained in liquid form in the vaporizers mounted on the machine. The gas mixture from the flowmeters flows through the vaporizer and the volatile anesthetic agent is added to the mixture in gaseous form. The concentration of the volatile gas in the final mixture is determined by a dial on or near the vaporizer. For safety reasons, only one volatile agent can be delivered at a time.

The ventilator allows positive pressure ventilation of the anesthetized patient. The ventilator can be set to deliver a specific tidal volume (in which case pressure varies according to lung compliance) or to achieve a certain peak inspiratory pressure (in which case volume varies ac-
cording to lung compliance). The ventilator moves the gas mixture through the common gas outlet and into the **anesthetic circuit**, the tubing that connects to the patient’s airway. The vast majority of general anesthetics today are delivered through a circle system. The **circle circuit** has a **CO2 absorber**, a canister containing a hydroxide mixture (soda lime) that absorbs CO2. The absorption of CO2 allows the expired gas to be recycled, thus minimizing the excessive cost and pollution that would otherwise result. There are several other types of circuits which are useful in specific clinical situations or are of historical interest. The origin and pathways of gas flow that applies to most anesthetic machines is depicted in schematic form in **Figure 9**.

It is imperative that all anesthesia equipment undergo regular checks and maintenance. It is the responsibility of the anesthesiologist to ensure that the equipment is in functioning condition prior to the administration of every anesthetic. The pre-operative checklist can be found on every anesthetic machine.

**Figure 9 Pathway of gas flow in anesthetic machine**

*The shaded shapes represent (from left to right): volatile anesthetic vapourizer, ventilator and bag used for bag-mask ventilation. Image by Wikimedia user TwoOneTwo, available under the Creative Commons Attribution-Share Alike 3.0 Licence. Image modified by Emma Kolesar.*
Monitoring
The purpose of monitoring during anesthesia is to ensure the maintenance of homeostasis. The best single monitor is a vigilant anesthesiologist. The practice of anesthesia involves the use of some key monitors that are not commonly seen in other health care settings. Examples include the **pulse oximeter**, the **capnograph** and the **peripheral nerve stimulator**. The Canadian Anesthesia Society guidelines for intra-operative monitoring are listed in Table 8.

In some settings, depending on the patient status or the nature of the procedure, monitoring beyond the routine measures listed above may be deemed necessary. There are methods of invasively monitoring the cardiovascular, renal and central nervous systems in the peri-operative period. Examples include arterial catheter, **pulmonary artery catheter**, **transesophageal echocardiography**, Foley catheter and 16-channel EEG monitoring.

---

### Table 8 The Canadian Anesthesia Society guidelines for intra-operative monitoring

<table>
<thead>
<tr>
<th>Monitors which must be continuously used:</th>
</tr>
</thead>
<tbody>
<tr>
<td>• pulse oximeter</td>
</tr>
<tr>
<td>• apparatus to measure blood pressure</td>
</tr>
<tr>
<td>• electrocardiography</td>
</tr>
<tr>
<td>• capnograph when an endotracheal tube or laryngeal mask is in use</td>
</tr>
<tr>
<td>• agent-specific anesthetic gas monitor when inhaled anesthetic agents are used</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Monitors which must be exclusively available:</th>
</tr>
</thead>
<tbody>
<tr>
<td>• apparatus to measure temperature</td>
</tr>
<tr>
<td>• peripheral nerve stimulator (when neuromuscular blockers are used)</td>
</tr>
<tr>
<td>• stethoscope (precordial or esophageal)</td>
</tr>
<tr>
<td>• visualization of exposed portion of patient with adequate lighting</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Monitors which must be immediately available:</th>
</tr>
</thead>
<tbody>
<tr>
<td>• spirometer for measurement of tidal volume</td>
</tr>
</tbody>
</table>

---
Interactive 2.1 Anesthesia machine

[Interactive image of anesthesia machine with labeled parts: Monitor, Flowmeters, Scavenger, Bag, APL valve, CO₂ absorber, Ventilator, On button, Volatile agent/Vapourizer, O₂ flush button.]
Chapter 2 Review

Review 2.1 The Pre-operative Phase

Question 1 of 15
Routine preoperative hemoglobin is NOT required for:

- [ ] A. A healthy female over 50 years of age.
- [ ] B. A patient who has been crossmatched for surgery.
- [x] C. A patient on anticoagulant medication.
- [ ] D. A young female patient with menorrhagia.

Check Answer
In this chapter, you will be presented an overview of the range of techniques that can be used to provide anesthesia. Regional and general anesthesia are discussed in greater detail. The pharmacology of each of the important drugs used in the delivery of anesthesia can be found in the “Drug Finder” (Chapter 6).
Except in the most desperate of circumstances, surgical procedures are performed with the benefit of anesthesia. There are four types of anesthesia that may be employed alone or in combination:

- local
- sedation (minimal, moderate or deep)
- regional
- general

The findings on pre-operative assessment, the nature of the surgery and the patient’s preference all factor into the choice of anesthetic technique. Contrary to popular belief, studies have failed to identify one technique as superior (lower morbidity and mortality) to the others in a general patient population. Regardless of the technique employed, the anesthesiologist must ensure patient comfort, maintenance of physiologic homeostasis and provision of adequate operating conditions.

Local Anesthesia

Local anesthesia refers to the infiltration of a local anesthetic agent at the surgical site and is usually performed by the surgeon. This technique is appropriate for superficial procedures such as dental surgery, breast biopsy or carpal tunnel release. Local anesthesia may be used in an unmonitored setting. However, often it is used in combination with sedation in which case monitoring is required. While local anesthesia is inadequate for more invasive procedures such as those involving the body cavities, local infiltration is often used as an adjunct in post-operative pain management. Care must be taken to avoid intravascular injection and to avoid exceeding the toxic dose of the local anesthetic in use.
Sedation

Sedation involves the delivery of agents (usually intravenous) for the purpose of achieving a calm, relaxed patient, able to protect his own airway and support his own ventilation. The range of physiologic effects of sedation is varied and is dependent on the depth of sedation provided: minimal, moderate or deep. A patient under minimal sedation will be fully responsive to verbal commands although his or her cognitive functions and coordination would be impaired. He or she would appear calm and relaxed and would have normal cardiorespiratory function. At the other extreme, a patient receiving deep sedation would be rousable only to repeated or painful stimuli. In some instances, the patient may require assistance in maintaining a patent airway. In this case, the line between deep sedation and general anesthesia is not easily identifiable. Table 9 outlines the physiologic effects of each of the three levels of sedation as defined by the American Society of Anesthesiologists (ASA). Sedation may be used alone for minimally painful procedures such as endoscopy. Often it is used in combination with local or regional anesthesia to provide a more palatable experience for the patient. In any case, the sedated patient must be monitored due to the depressant effects of the agents used. Care must be taken to reduce the dose administered to the frail, elderly or debilitated patient, in whom depressant effects may be exaggerated. In any patient, sedation may cause disinhibition, resulting in an uncooperative, agitated patient.

Many different agents have been used for sedation. The term “neurolept anesthesia” refers to the (now historical) use of high doses of droperidol (a butyrophenone, in the same class as haloperidol) in combination with fentanyl (an opioid). Side effects were prominent. Currently, agents are chosen with specific effects in mind. Opioids, such as fentanyl or remifentanil, may be given alone if analgesia is the primary goal. The short-acting benzodiazepine, midazolam, is a popular choice because it provides amnesia as well as anxiolysis. Propofol, an anesthetic induction agent, can be infused in sub-anesthetic doses to produce a calm, euphoric patient.
| Table 9 Sedation |
|-----------------|-----------------|-----------------|-----------------|-----------------|
|                | VERBAL RESPONSE | COGNITIVE FUNCTION | AIRWAY PATENCY | RESPIRATORY FUNCTION | CARDIOVASCULAR FUNCTION |
| Light Sedation | Normal          | Conscious, but cognitive function and coordination would be impaired | Normal          | Normal          | Normal          |
| Moderate Sedation | Would respond purposefully to verbal commands | Depressed consciousness | Normal          | Normal          | Normal          |
| Deep Sedation  | Not easily aroused; verbal response only to painful or repeated physical stimuli | Depressed consciousness | Airway might require support to remain patent | Spontaneous ventilation may be inadequate | Usually maintained |
**Regional Anesthesia**

Regional anesthesia involves the blockade (with local anesthetics) of the nerve supply to the surgical site. This may be achieved by blocking peripheral nerves (e.g. ankle block) or by blocking the spinal cord and/or nerve roots (spinal, epidural block). A single nerve block may be sufficient (e.g. ulnar nerve block for repair of 5th digit) or a group of nerves may need to be blocked (e.g. brachial plexus block for forearm fasciotomy). While regional techniques are perceived to be “safer” than general anesthesia, they do carry risks of their own. The most serious “early” complication of a peripheral nerve block is local anesthetic toxicity. The most worrisome “late” complication is neuropraxis or nerve injury. The central neuraxial blocks have many potential complications, both early and late, which will be discussed in the next section.

There are some patients in whom a regional technique offers at least short term benefits over general anesthesia. For example, in those undergoing total hip arthroplasty, the use of spinal or epidural anesthesia is associated with less intra-operative blood loss, less post-operative hypoxemia and a lower risk of post-operative deep venous thrombosis formation. While it seems intuitive that physiologic homeostasis is more readily achieved when regional anesthesia is employed, the anesthesiologist must always remain vigilant: numerous case reports of sudden cardio-respiratory arrest in patients under central nerve blocks emphasize this point all too clearly. Principles of pre-operative assessment and preparation must be applied just as vigilantly to the patient undergoing regional anesthesia.
**General Anesthesia**

General anesthesia is a pharmacologically-induced, reversible state of unconsciousness. It may involve the use of intravenous agents, inhaled agents or both. The goals and techniques of general anesthesia are discussed in an upcoming chapter. General anesthesia may be used alone or in combination with local anesthesia or a regional technique. An example of such a “combined technique” would be the use of epidural and general anesthesia in a patient undergoing an abdominal aortic aneurysm repair. Such a technique allows the continued use of the epidural for post-operative pain management and may confer a lower morbidity and mortality in high risk patients.
Many different types of surgical procedures can be performed while the nerves that supply the surgical site are rendered insensate. This method is called “neural blockade” or “nerve block”. Various techniques of neural blockade comprise what is known as **regional anesthesia**. During regional anesthesia, the anesthesiologist must not only monitor and manage the patient’s physiologic status but he or she must ensure that the patient remains calm and cooperative. The anesthesiologist must be alert to the development of complications and must also be prepared to convert to a general anesthetic at any point in the procedure.

A complete description of the scope of regional anesthesia is beyond the scope of this textbook. A brief discussion of four commonly-used techniques is presented below.
**Epidural Anesthesia**

Epidural and spinal anesthesia can be used for procedures involving the abdomen, perineum or lower extremities. The two blocks differ by virtue of the anatomic space into which local anesthetic (LA) is delivered. Understanding the anatomy of the region ([Figure 10, Figure 11](#)) is crucial to understanding the blocks.

**Figure 10** Ligamentous anatomy of the spine

From “Introduction to Regional Anaesthesia” by D. Bruce Scott (1989). Used with permission from his wife, Joan and son, Nicholas B. Scott.

In **epidural anesthesia**, a tiny plastic catheter is placed into the **epidural space**, which is the anatomic space located just superficial to the dura. An epidural catheter can be placed at any point along the spinal column. Epidural catheters placed for surgical anesthesia or analgesia are most commonly used at the thoracic or lumbar regions depending on the site of the surgery. A LA solution is delivered through the catheter. From the epidural space, it is slowly absorbed into the **subarachnoid space** where it blocks the nerves of the spinal cord.
and cauda equina. The volume of anesthetic delivered and the site of the catheter determine the level or “height” of the block. The presence of an indwelling catheter allows the block to be extended in height or duration as required.

Insertion of an epidural catheter is done in a strictly sterile fashion. After local infiltration, a specially designed 17 or 18 gauge epidural needle (common trade names Tuohy® or Hustead®) is inserted into the spinous interspace. A special syringe, filled with air or saline is attached to the hub of the needle. While advancing the needle, the anesthesiologist maintains pressure on the syringe in order to sense the resistance of the tissue being traversed (Figure 12). The epidural space is a “potential space” such that when it is entered with the needle, a sudden loss of resistance is detected. The syringe is then removed so that a catheter can be threaded through the needle into the epidural space (Figure 13), after which the needle is removed. A labour epidural insertion can be viewed in Movie 3.1.

This movie demonstrates two frequent challenges faced by the anesthesiologist. The first is the difficulty of inserting an epidural in the presence of a dermal tattoo. Inserting an epidural through tattooed skin is undesirable as it may bring a plug of ink into the epidural space, the consequences of which are not known. In this case, the anesthesiologist is able to locate a small

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**Figure 12** Insertion of Tuohy needle into epidural space

Reproduced with permission from Astra Pharma Inc.

**Figure 13** Insertion of epidural catheter

Reproduced with permission from Astra Pharma Inc.
Movie 3.1 Labour epidural insertion

Procedure videotaped and presented with patient permission. Produced by Karen Raymer.
area of non-inked skin just slightly off midline, which provides satisfactory access for catheter insertion. The second challenge is performing a technical procedure in a patient who is in active labour. In this case, the anesthesiologist pauses while the patient is having contractions. The patient is able to do an excellent job of remaining still, which is quite important during this delicate procedure. As you are watching the video, watch carefully for the moment of the “loss of resistance”, when the gentle pressure on the hub of the syringe finally gives way, as the needle has entered the “potential” space that is the epidural space.

Once the catheter is in place, the anesthesiologist “tests” the catheter to ensure that neither blood nor cerebral spinal fluid (CSF) can be aspirated. While delivering LA in small (3 ml) aliquots, the anesthesiologist observes for symptoms or signs of incorrect catheter placement. If the catheter is situated in one of the veins of the epidural plexus then the patient may experience symptoms of local anesthetic toxicity: tinnitus, perioral numbness, metallic taste in the mouth and dizziness. If the catheter is situated in the intrathecal space, then the patient will develop a sensory/motor block rapidly after the administration of only a small amount of LA.

In the absence of any signs or symptoms of incorrect catheter placement, the full dose of LA is delivered over 10-20 minutes. Some LA’s, such as lidocaine, have a relatively rapid onset of effect but also have a relatively shorter duration of action. Bupivacaine, while possessing a slower onset of effect, has a longer duration of action. The dermatomal level of block is tested by pinprick or ice cube (Figure 14). It generally takes 20-30 minutes for an adequate epidural block to take effect. The required dose is determined by the range of segments which must be blocked. The higher the surgical site is, the higher the block must be. Table 10 describes the dermatomal level of block required for some of the more common surgical procedures which apply to both spinal and epidural anesthesia. Many other factors influence the amount of local anesthetic required. For example, in pregnancy, there is a significant reduction in LA effect.

<table>
<thead>
<tr>
<th>SURGICAL PROCEDURE</th>
<th>REQUIRED LEVEL OF BLOCK</th>
</tr>
</thead>
<tbody>
<tr>
<td>Caesarian section</td>
<td>T4</td>
</tr>
<tr>
<td>inguinal hernia repair</td>
<td>T10</td>
</tr>
<tr>
<td>repair fractured hip</td>
<td>L1</td>
</tr>
<tr>
<td>total knee arthroplasty (with tourniquet)</td>
<td>L2</td>
</tr>
<tr>
<td>hemorrhoidectomy</td>
<td>S4</td>
</tr>
</tbody>
</table>
tion in the required volume of LA delivered to achieve the desired block. In general, volumes in the range of 10-20 ml are required for most procedures.

Complications of epidural anesthesia can present in the early or late post-operative periods. The early complications are related either to incorrect catheter placement (LA toxicity, or “total spinal” block), to excessive volume of LA delivered (high block, with hypotension, bradycardia, respiratory compromise), or to the unavoidable blockade of sympathetic fibres (hypotension, bradycardia). Late complications are related to needle and catheter insertion, and include nerve injury, epidural abscess or hematoma, and post-dural puncture headache (if the dura is accidentally punctured).

The contraindications to epidural and spinal anesthesia are identical, and are listed in Table 11.
Spinal Anesthesia

Spinal anesthesia involves the blockade of the nerves of the spinal cord and cauda equina by injection of LA into the intrathecal space. The important anatomy is depicted in Figure 15 and Figure 16. Because the LA is injected in close proximity to its site of action, much smaller volumes are required (1-3 ml) and the onset of effect (within 5 minutes) is rapid relative to epidural an-

Table 11 Contraindications to central neural blockade

<table>
<thead>
<tr>
<th>GENERAL</th>
<th>SPECIFIC (ABSOLUTE)</th>
<th>SPECIFIC (RELATIVE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>lack of consent</td>
<td>coagulopathy</td>
<td>evolving neurological deficit</td>
</tr>
<tr>
<td>lack of resuscitative equipment</td>
<td>sepsis (systemic or at site of injection)</td>
<td>obstructive cardiac lesion (e.g. aortic stenosis)</td>
</tr>
<tr>
<td>known or suspected allergy</td>
<td>increased intracranial pressure (ICP)</td>
<td>spinal hardware</td>
</tr>
<tr>
<td>lack of familiarity with technique</td>
<td>shock</td>
<td></td>
</tr>
<tr>
<td>known or suspected allergy</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

From “Introduction to Regional Anaesthesia” by D. Bruce Scott (1989). Used with permission from his wife, Joan and son, Nicholas B. Scott.
The choice of LA used is based primarily on the anticipated length of procedure. Spinal lidocaine provides surgical anesthesia for procedures lasting up to 75 minutes however its use has been limited by the associated increased incidence of postoperative radiculopathies. Spinal bupivacaine will provide up to 3.5 hours of anesthesia and is considered to be safe. The addition of opioids (e.g. fentanyl) to the local anesthetic solution can extend the duration of block. However, if the block dissipates prior to the end of the procedure there is no way to extend the block at that point. Spinal anesthesia is performed under strict asepsis. The patient may be sitting or curled in the lateral position. A special small-bore “spinal needle” is used (22-27 gauge). The needle is inserted at a lower lumber interspace and is advanced through the dura. Because the dura is a tough membrane, a definite “pop” is often felt as the needle passes through into the intrathecal space. The stylet of the needle is removed and cerebrospinal fluid (CSF) is observed in the lumen of the needle. The local anesthetic is then injected and the needle removed. The height of the required block depends on the surgical procedure (Table 10). The height of the block achieved is determined by many factors including the mass and volume of LA administered, the position of the patient and the baricity or “heaviness” of the LA relative to CSF.

The complications of spinal anesthesia are similar to those of epidural anesthesia with a few exceptions. Because of the small dose of LA required, LA toxicity is not an issue even if intravascular injection occurs. Dural puncture is required for spinal anesthesia therefore “spinal headache” is always a risk, especially in young adult patients. The use of needles which are smaller-bore and have a “pencil-point” tip helps to decrease the incidence of post-dural puncture headache. The contraindications to spinal anesthesia are listed in Table 11.

From “Introduction to Regional Anaesthesia” by D. Bruce Scott (1989). Used with permission from his wife, Joan and son, Nicholas B. Scott.
Intravenous Regional Block (Bier Block)

An intravenous regional anesthetic (IVRA), also known as a Bier Block, is used primarily for relatively short procedures on the distal upper extremity (below the elbow). The monitors are applied and intravenous access is ensured. After cannulating a vein distal to the surgical site, the operative arm is elevated and an elastic bandage is applied to promote venous drainage. After exsanguination, an upper arm tourniquet is inflated. 40-50 ml of dilute lidocaine (without epinephrine) is then injected slowly into the cannula in the operative arm. Diffusion of the local anesthetic from the venous system to the interstitium provides surgical anesthesia within 5 minutes.

The risk of LA toxicity is high especially if the tourniquet leaks or is deflated prior to 20 minutes after injection of LA. If the surgical procedure lasts less than 20 minutes then one must wait until 20 minutes has elapsed prior to deflating the tourniquet. If less than 40 minutes (but more than 20 minutes) has elapsed, then the tourniquet should be deflated and re-inflated intermittently to avoid a sudden bolus of LA entering the systemic circulation. “Tourniquet pain” becomes prominent after 45 minutes and limits the length of time that this technique can be employed.

Brachial Plexus Block

The brachial plexus is formed from the anterior primary rami of the C5-T1 nerve roots (Figure 17) and supplies all of the motor function and most of the sensory function of the upper extremity (Figure 18). Throughout their journey to the axilla, the nerve roots merge and divide numerous times (Figure 17). The nerve roots travel through the intervertebral foramina and emerge between the anterior and middle scalene mus-
cles (Figure 19). They then travel under the clavicle and enter the axilla as three distinct cords. As they exit the axilla, the plexus divides one final time to form the axillary, radial, median, ulnar and musculocutaneous nerves. The brachial plexus block provides anesthesia for virtually any type of upper extremity surgery.

The brachial plexus can be blocked at one of its three most superficial locations: interscalene, supraclavicular or axillary. Sterile technique is used at all times and correct needle placement is ensured through the use of either ultrasound or nerve stimulator. A relatively large volume of LA is injected (30-40 ml). Up to 45 minutes may be required until adequate anesthesia is achieved, depending on how close to the nerve bundle the local anesthetic is deposited. If several hours of anesthesia are required, then bupivacaine or ropivacaine is used.

**Figure 18** Sensory innervation of upper limb

![Figure 18 Sensory innervation of upper limb](image)

Public domain image derived from Gray’s Anatomy, retrieved from Wikimedia Commons.

**Figure 19** Brachial plexus: anatomical depiction

![Figure 19 Brachial plexus: anatomical depiction](image)

Gray’s Anatomy image in public domain, retrieved from Wikimedia Commons.
A catheter can be inserted to provide surgical anesthesia or post-operative analgesia for an indefinite period of time.

There are many potential complications of a brachial plexus block. LA toxicity is a risk because of the large volumes of LA used and the proximity of the injection site to major vessels. Any block can lead to hematoma formation, nerve injury or infection. The supraclavicular and interscalene blocks pose the additional risks of pneumothorax, phrenic nerve block and recurrent laryngeal nerve block. Consequently, bilateral blocks are contraindicated. Intrathecal injection is a rare complication of interscalene block.
General Anesthesia is a pharmacologically induced reversible state of unconsciousness which is maintained despite the presence of noxious stimuli. With the continual development of new drugs, there is an ever-increasing variety of techniques used to provide general anesthesia. All techniques strive to achieve the following goals, known as the “Four A’s of Anesthesia”:

• **Lack of Awareness**: unconsciousness.
• **Amnesia**: lack of memory of the event.
• **Analgesia**: the abolition of the subconscious reactions to pain, including somatic reflexes (movement or withdrawal) and autonomic reflexes (hypertension, tachycardia, sweating and tearing).
• **Akinesia**: lack of overt movement. In some cases, the provision of muscle relaxation may be required.

In the past, general anesthesia was achieved using a single agent such as ether or chloroform. Because the above-described goals were achieved by a progressive depression of the central nervous system rather than by any direct or specific effect, relatively high concentrations of the gases were required. Consequently the associated side-effects were frequent and severe.

In current practice, we have many different agents (both intravenous and inhaled) at our disposal. The intravenous agents in particular have specific effects such as analgesia or muscle relaxation and therefore can be used to achieve the desired effect in a dose-related fashion.

The practice of using combinations of agents, each for a specific purpose, is what is termed “balanced anesthesia”. An example of a balanced technique would be the use of propofol for induction of anesthesia; the administration of desflurane and nitrous oxide for maintenance of unconsciousness; sufentanil for analgesia; and rocuronium for muscle relaxation.

The anticipated benefits of a balanced technique as compared to “ether anesthesia” of the past include:

• improved hemodynamic stability
• more effective muscle relaxation
• more rapid return of respiratory function, consciousness and airway control following the completion of the procedure
• provision of post-operative analgesia with appropriate timing and dosing of opioids administered intraperoperatively

A balanced technique is still the most common technique used for the provision of general anesthesia. However, with the development of short-acting intravenous agents such as propofol and remifentanil, the above-described goals of general anesthesia can be attained with the use of intravenous agents alone, usually by continuous infusion. This is called “Total Intravenous Anesthesia” or “TIVA”.

After the patient has been assessed, the equipment and drugs prepared and the anesthetic technique determined, one can proceed with administering the anesthetic. A general anesthetic consists of four phases: induction, maintenance, emergence and recovery.

**Induction**
The goal of the induction phase of anesthesia is to induce unconsciousness in a fashion which is pleasant, rapid and maintains hemodynamic stability. If the anesthetic plan includes control of the airway and ventilation then the induction phase also aims to achieve muscle relaxation to facilitate endotracheal intubation.

Anesthesia can be induced by having the patient breathe increasing concentrations of inhaled gases by mask. While there are settings where this is the desired technique, it tends to be slow and can be unpleasant. More commonly, anesthesia is induced with short-acting intravenous agents such as propofol, ketamine, thiopental or etomidate, followed by a muscle relaxant if indicated. In most cases, a non-depolarizing muscle relaxant (NDMR) is used. NDMR are discussed later in this chapter. Understanding the dynamics of induction requires a grasp of the essential pharmacology of these agents; the reader can do so by touching the hyperlink on each drug or by visiting Chapter 6.

**Rapid Sequence Induction**
Although regurgitation and aspiration are potential complications of any anesthetic, there are factors which place some patients at higher risk (Table 7). The obvious risk factor is recent intake of solid food. However, even a prolonged period of fasting does not guarantee an “empty stomach” if gastric emptying is delayed. Ex-
amples of conditions which impair gastric emptying in-clude diabetes, trauma, recent opioid administration and bowel obstruction. Finally, decreased integrity of the lower esophageal sphincter, as occurs in pregnancy and obesity, increases the risk of regurgitation of stomach contents. In patients deemed to be at increased risk for aspiration, the time between inducing anesthesia and securing the airway with a cuffed endotracheal tube must be minimized. Such a technique is termed a “rapid sequence induction”.

A rapid sequence induction is performed as follows:

1. Suction apparatus is checked and kept readily available.

2. Pre-oxygenation of patient with 100% oxygen for 3-5 minutes.

3. Application of cricoid pressure (Sellick’s maneuver) by assistant.

4. Induction with pre-calculated dose of induction agent followed immediately by intubating dose of depolarizing muscle relaxant (succinylcholine). A rapidly acting non-depolarizing agent (e.g. rocuronium) is commonly used in a so-called “modified” rapid sequence induction.

5. Intubation of trachea, cuff inflation and verification of proper tube position.

The purpose of pre-oxygenation is to lessen the risk of hypoxemia occurring during the apneic period after induction. Although pre-oxygenation does increase the patient’s arterial PO2 prior to induction, its most important effect is the de-nitrogenation of the functional residual capacity of the lungs. Traditionally teaching is that the Sellick maneuver provides occlusion of the esophagus between the cricoid cartilage (a complete circumferential cartilage) and the cervical vertebrae thus minimizing the risk of passive regurgitation.

Succinylcholine

Succinylcholine (Sch), a depolarizing muscle relaxant, is a very useful and very powerful drug; the anesthesiologist must understand the effects and contraindications of Sch in order to avoid causing harm or death.

Succinylcholine causes rapid, profound and brief muscular paralysis. It acts by attaching to nicotinic cholinergic receptors at the neuromuscular junction. There, it mimics the action of acetylcholine, thus depolarizing the post-junctional membrane. Neuromuscular blockade (paralysis) develops because a depolarized post-junctional membrane cannot respond to subsequent release of acetylcholine.

Succinylcholine has effects on almost every organ system, most of them being secondary to the depolarization and subsequent contraction of skeletal muscle.
Important effects include increased intracranial pressure, increased intragastric pressure and post-operative myalgia.

The most critical effects of Sch relate to its interaction with muscle cells. Sch elevates serum potassium by 0.5 mEq/L in patients with normal neuromuscular function and is therefore contraindicated in hyperkalemic patients or in patients with renal failure, for whom even a small rise in potassium could have critical implications. Sch can cause an exaggerated release of potassium, leading to fatal hyperkalemia, in those with neuromuscular or muscle disease. Examples include recent paralysis (spinal cord injury or stroke), amyotrophic lateral sclerosis, Duchenne’s muscular dystrophy and recent burn or crush injury.

Sch is a potent trigger of malignant hyperthermia (MH) and is therefore contraindicated in MH-susceptible patients. Sch is contraindicated in patients with pseudo-cholinesterase deficiency in whom the paralysis will be prolonged.

A more complete discussion of succinylcholine can be found in Chapter 6.

**Maintenance**

If no further agents were administered following the induction of anesthesia the patient would awaken within minutes. Therefore, **maintenance** of anesthesia requires the delivery of pharmacologic agents with the aim of achieving the “four A’s of anesthesia” and hemodynamic stability throughout the surgical procedure. A further consideration is the length of the procedure and the need to awaken the patient at the end of the case. The maintenance phase of anesthesia involves the use of **inhaled agents**, **opioids** and **non-depolarizing muscle relaxants** (NDMR). These agents are described later in this chapter.

The anesthesiologist must be ever vigilant. Problems related to the airway, breathing and circulation (ABC’s) are most critical and can occur during any phase of anesthesia. Several of the problems that are unique to the maintenance phase of anesthesia are discussed below.

**Awareness**

“**Awareness**” refers to a complication of anesthesia whereby a patient who has received a general anesthetic becomes conscious of his or her environment during the surgical procedure. The patient may or may not experience pain and may or may not recall the events post-operatively. The incidence of awareness with recall has been estimated at 0.2-1.0%. Some types of surgical procedures such as Caesarian section, cardiac surgery
and trauma surgery pose a higher risk of awareness because of the nature of the anesthetic given for those procedures. It may be prudent to warn such patients of the risk pre-operatively.

Intra-operatively, care should be taken to ensure delivery of adequate amounts of hypnotic drugs such as inhaled agents, propofol, benzodiazepines or ketamine. Opioids alone provide very little hypnosis and muscle relaxants provide none whatsoever! Signs of awareness should be sought. In an un-paralyzed (or partially paralyzed) paralyzed patient, this includes movement. However, a fully paralyzed patient is only able to communicate through the autonomic nervous system with signs of sympathetic hyperactivity, such as hypertension, tachycardia, sweating and tearing. Not surprisingly, the overwhelming majority of cases of awareness have been reported in paralyzed patients.

**Positioning**

The patient is positioned to facilitate surgical access. Depending on the procedure, the patient may be placed in the supine, prone, lateral, lithotomy, jack-knife, kidney or even the sitting position to name but a few. Most of the consequences of positioning involve the cardiovascular, respiratory and peripheral nervous systems.

Kinking of, or pressure on major vessels leads to decreased venous return, decreased cardiac output and hypotension. This is particularly relevant when the prone or kidney position is used. In the semi-sitting position, venous pooling in the legs has a similar effect. Very occasionally, surgery is performed in the sitting position which is associated with the risk of **venous air embolism**.

The airway may become obstructed or dislodged while the patient is in the prone position. The prone, trendelenburg and lithotomy positions may cause an upward displacement of the diaphragm due to an increase in intra-abdominal pressure. This leads to ventilation/perfusion mismatching and decreased lung compliance which may manifest as hypoxemia, hypercarbia or increased airway pressure.

Nerve injury results from compression on pressure points or stretching. Other factors such as prolonged surgery, hypothermia, hypotension, obesity and diabetes may play a role in increasing the risk of a postoperative **neuropathy**. The ulnar nerve, because of its superficiality, is at risk of compression in almost any position. Padding is commonly used but has not been shown convincingly to be helpful. Careful positioning is probably most important in this regard. The brachial plexus is at risk of stretch injury when arms are abducted in the supine position. The angle of abduction should be kept below 90 degrees and the head should be turned slightly toward the abducted arm. Many nerves including the sciatic, lateral femoral cutaneous
and common peroneal nerves are at risk of either stretch or compression injuries when the lithotomy position is used.

Other organ systems may be vulnerable in the prone position. Pressure on the orbit of the eye can lead to retinal ischemia by either arterial compression or obstruction of venous flow. The eye socket itself provides a natural protection and specially designed head rests are helpful. Constant vigilance must be maintained as patient position may shift during anesthesia. The male patient’s genitalia must be free of pressure. Finally, pressure over skin surfaces (e.g. the forehead) must be minimized as skin sloughing can result after prolonged surgery in the prone position.

**Hypothermia**

Hypothermia has deleterious effects on the cardiovascular, respiratory, central nervous, hematologic and renal systems. As well, it decreases the rate of recovery from the effects of muscle relaxants.

Heat is lost through four mechanisms:

- convection (e.g. exposure to drafts of cool air)
- conduction (e.g. contact with cold operating room table)
- evaporation (e.g. airway mucosa, prep solution, sweat)
- radiation (e.g. temperature gradient between patient and operating room environment)

Furthermore, the normal responses to hypothermia (shivering, vasoconstriction) are abolished under anesthesia. Procedures which are prolonged, involve large abdominal incisions or require administration of large volumes of intravenous fluids can be associated with particularly severe hypothermia.

Heat loss can be minimized by keeping the operating room temperature as high as tolerable (>21 C, preferably). Gas humidifiers or the use of low gas flow minimizes the heat lost through airway evaporation. Fluid warmers should be used whenever blood products or large volumes of intravenous fluids are being given. A **forced air warming system** should be used routinely except for those cases which are very short in duration.

If significant hypothermia (<35 C) results despite preventative measures then, depending on the underlying patient condition, the anesthesiologist may decide to leave the patient sedated, paralyzed and mechanically ventilated post-operatively until adequate temperature is restored. This is particularly important for patients with cardiac or respiratory disease where shivering (which increases oxygen consumption fourfold) may compromise outcome.
Drugs for Maintenance of Anesthesia
Inhaled Agents

The term “inhaled agent” refers to the volatile agents (desflurane, isoflurane, sevoflurane) and nitrous oxide (N2O). While the development of intravenous agents has largely eliminated the use of inhaled agents for induction except in the pediatric population, they continue to be the mainstay for maintenance of anesthesia. The volatile agents are so called because with vapour pressures below atmospheric pressure, they exist in equilibrium between liquid and gaseous states at room temperature. N2O is a gas at room temperature and atmospheric pressure and is therefore not a volatile agent though it is, of course, an inhaled agent.

Mechanism of Action
The mechanism of action of inhaled agents is not well understood.

Dose
While intravenous agents are given in mg (or mcg)/kg doses, the inhaled agents are given in volume percent concentrations. The volatile agents can also be termed “potent” vapours, because concentrations in the range of 0.3-6% are clinically effective while N2O (not potent) must be given in concentrations of between 30% and 70% to have any effect.

This brings us to the concept of Minimum Alveolar Concentration (MAC). The concentration of a gas in the alveoli creates an alveolar partial pressure of gas which in turn reflects its partial pressure in the active site (brain). MAC refers to the concentration of the inhaled agent in alveolar gas necessary to prevent movement of 50% of patients when a standard incision is made. This definition hints at the fact that a specific concentration of gas does not correlate to a predictable clinical effect. Many factors influence MAC, and therefore influence the concentrations required to maintain anesthesia. Some of the known factors are listed in Table 12.

The MAC values of the commonly used agents are shown in Table 13. Note that N2O, with a MAC>100%, can never be used as a sole agent to provide anesthesia.

<table>
<thead>
<tr>
<th>FACTORS WHICH DECREASE MAC</th>
<th>FACTORS WHICH INCREASE MAC</th>
</tr>
</thead>
<tbody>
<tr>
<td>advanced age</td>
<td>childhood</td>
</tr>
<tr>
<td>pregnancy</td>
<td>hyperthyroidism</td>
</tr>
<tr>
<td>hypothermia</td>
<td>hyperthermia</td>
</tr>
<tr>
<td>acute alcohol intoxication</td>
<td>chronic alcohol use</td>
</tr>
<tr>
<td>drugs: benzodiazepines, opioids, muscle relaxants, central-acting antihypertensives</td>
<td>drugs: amphetamine, cocaine</td>
</tr>
</tbody>
</table>

Table 12 Factors affecting MAC
Even the potent agents are often not administered at 1 MAC concentrations during general anesthesia. This is because other agents (such as opioids) are also being given. It is generally felt that 0.5 MAC of inhaled agent is the minimum level to provide adequate hypnosis and amnesia.

**Onset of action, duration of action and elimination**
The solubility of a gas in blood determines its rate of onset and offset of effect. An agent such as N₂O, which is relatively insoluble in blood, will build up its alveolar partial pressure (and therefore brain partial pressure) quickly and consequently will have a faster onset and offset of effect. Conversely, a soluble gas such as isoflurane will equilibrate slowly throughout body stores and therefore onset and offset lags. You may notice this theory being put into practice in the operating room: a soluble potent agent is often discontinued 5 or 10 minutes prior to the end of surgery while N₂O is delivered until moments before the desired emergence or wake-up. Table 13 summarizes these pharmacokinetic properties. The termination of effect of inhaled agents depends only on its exhalation from the lungs. The exception, of historical interest, is halothane of which up to 20% can undergo metabolism in the liver.

**Effects of the inhaled agents**
The effects of the volatile agents are quite distinct from those of nitrous oxide. Inhaled agents have effects on almost every organ system and the reader is referred to Chapter 6 for a detailed summary. Several key effects are highlighted below.

All volatile anesthetics are triggers of malignant hyperthermia while nitrous oxide is not. N₂O expands the volume of gas-containing spaces as N₂O diffuses across membranes more readily than nitrogen can diffuse out. Thus the size of a pneumothorax, an emphysematous bleb or a distended bowel loop will increase when N₂O is used.

---

**Table 13 Characteristics of inhaled agents**

<table>
<thead>
<tr>
<th></th>
<th>N₂O</th>
<th>ISOFLURANE</th>
<th>SEVOFLURANE</th>
<th>DESFLURANE</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ODOUR</strong></td>
<td>odourless</td>
<td>pungent</td>
<td>sweet</td>
<td>pungent</td>
</tr>
<tr>
<td><strong>MAC%</strong></td>
<td>104</td>
<td>1.15</td>
<td>1.8</td>
<td>6.6</td>
</tr>
<tr>
<td><strong>BLOOD/GAS PARTITIONING</strong></td>
<td>0.47</td>
<td>1.4</td>
<td>0.65</td>
<td>0.42</td>
</tr>
<tr>
<td><strong>VAPOUR PRESSURE 20°C MMHG</strong></td>
<td>38,770</td>
<td>238</td>
<td>157</td>
<td>669</td>
</tr>
</tbody>
</table>
Contraindications to the inhaled agents

The use of volatile agents is absolutely contraindicated in patients who are known or suspected to have malignant hyperthermia.

The use of nitrous oxide is contraindicated in patients with pneumothorax or bowel obstruction. As N2O raises intracranial pressure, its use is avoided in patients with intracranial pathology. Caution should be used in those patients with coronary artery disease or emphysema.

Opioids

Opioids are used intra-operatively to provide analgesia, and to reduce the requirement of other maintenance agents. The commonly used intravenous agents are the synthetic opioids fentanyl, sufentanil, remifentanil and alfentanil. They are favoured by anesthesiologists over the more familiar agents, such as morphine and meperidine. Their shorter duration of action allows finer titration to provide adequate analgesia during the variable, but intense nature of surgical stimulation, while still allowing for awakening at the end of the procedure. While there are many different opioids available for use, the discussion below is limited to the three synthetic agents which are most commonly used in anesthetic practice.

Usually, an opioid is administered in the form of a loading dose, prior to induction. Not only does this help to blunt the response to intubation, which is a very stimulating maneuver, but it establishes a plasma level of opioid which can then be supplemented as needed throughout the surgical procedure. In the case of remifentanil, which is extremely short-acting, a bolus followed by an infusion is most practical.

Mechanism of action

There are five different opioid receptors of which the most clinically relevant are the Mu and Kappa receptors. Binding to different receptors produces distinct responses. Activation of Mu receptors produces analgesia, respiratory depression, bradycardia, euphoria and decreased gastrointestinal motility. Binding to Kappa receptors produces analgesia, sedation and meiosis. The major receptors for analgesia are the Mu-1 receptor at the periaqueductal gray area of the midbrain and the Kappa receptor at the substantia gelatinosa of the spinal cord. Each opioid has its own unique profile of agonism and antagonism for each receptor. Unfortunately, an agent which possesses agonism exclusively at the analgesia receptors has not yet been developed.

Dose, onset, and duration

All opioids are relatively lipid soluble. The greater the lipid solubility, the greater the potency. As a general rule:
Onset of action is determined by lipid solubility and ionization (pKa). Duration of action is determined by both clearance and volume of distribution. Table 14 summarizes the clinically useful pharmacology of the opioids most commonly used in anesthesia.

<table>
<thead>
<tr>
<th></th>
<th>Fentanyl</th>
<th>Sufentanil</th>
<th>Remifentanil</th>
</tr>
</thead>
<tbody>
<tr>
<td>Induction dose (µg/kg)*</td>
<td>4-20</td>
<td>0.25-2</td>
<td>0.5-1</td>
</tr>
<tr>
<td>Intra-op dose for</td>
<td>2-5 µg/kg/hr</td>
<td>0.3-2.0 µg/kg/hr</td>
<td>0.1-1 µg/kg/min</td>
</tr>
<tr>
<td>maintenance*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Additional boluses*</td>
<td>25-150 µg</td>
<td>2.5-20 µg</td>
<td>0.1-1 µg/kg</td>
</tr>
<tr>
<td>Onset of action (min)</td>
<td>5-8</td>
<td>4-6</td>
<td>1-2</td>
</tr>
<tr>
<td>Duration after bolus (min)</td>
<td>45</td>
<td>45</td>
<td>1-4</td>
</tr>
</tbody>
</table>

*Loading and maintenance requirements depend greatly on the patient age and status, the doses of other anesthetic agents given, and the nature of the surgical procedure being performed.

**Elimination**

The relatively short duration of action of these agents is in part attributable to their lipid solubility. This leads to rapid redistribution away from the central nervous system to inactive tissue sites. Fentanyl and sufentanil are metabolized in the liver to (mostly) inactive metabolites which are then excreted in the urine. Remifentanil on the other hand, is susceptible to metabolism by blood and tissue esterases which accounts for its ultrashort duration of action.

**Effects**

Opioids have effects on almost every system in the body. The reader is referred to Chapter 6 for a detailed discussion.

The most important side effect of the opioids manifests on the respiratory system. Minute ventilation is reduced due to a reduction in respiratory rate. (Tidal volume actually increases.) The responsiveness to raised PCO2 is diminished such that apnea occurs until the dose-dependent “apneic threshold” of PCO2 is reached.

Opioids cause nausea and vomiting due to stimulation of the chemoreceptor trigger zone. They also cause constipation due to decreased GI motility.
**Contraindications**
Opioids must not be given to those with a known allergy. Intravenous opioids must not be given in settings where one is not able to support ventilation.

Caution should be used when administering opioids to patients with hypovolemic or cardiogenic shock, where the blunting of sympathetic tone may exacerbate hypotension.

**Non-Depolarizing Muscle Relaxants**
The decision to use non-depolarizing muscle relaxants during maintenance of anesthesia depends on both the type of surgical procedure and the type of anesthetic. Some procedures require muscle relaxation to facilitate exposure (e.g. intra-abdominal surgery). In other cases, muscle relaxation is required because patient movement would be detrimental (e.g. neurosurgery, ophthalmic surgery). In a balanced technique, the use of muscle relaxants decreases the requirements of the other agents and facilitates mechanical ventilation.

Historically, the choice of succinylcholine versus a non-depolarizing muscle relaxant (NDMR) for use during induction of anesthesia was a decision which balanced the need for rapid airway control against the side effects of succinylcholine. With the introduction of the rapidly acting non-depolarizing agent, rocuronium, the use of succinylcholine has steadily declined. In the rare circumstance where succinylcholine is used to facilitate intubation, a NDMR is given after the effects of succinylcholine have worn off. However, in the vast majority of cases NDMR is given at induction to provide relaxation for both intubation and surgery.

**Mechanism of action**
In order to appreciate how NDMR cause muscular paralysis, one must have a basic understanding of how neuromuscular transmission occurs (Figure 20). Normally a nerve impulse travels the length of the nerve to arrive at the motor nerve terminal where it causes release of acetylcholine (Ach) into the synaptic cleft. The Ach then binds to post-synaptic nicotinic Ach receptors causing a conformational change in those receptors.

This conformational change leads to a change in membrane permeability of sodium and potassium causing depolarization of the post-junctional membrane. The propagation of this action potential leads directly to muscular contraction. NDMR interfere with this process by binding to the post-synaptic Ach receptors thereby acting as a **competitive inhibitor** to Ach.

**Dose, onset, duration, elimination and effects**
The choice of which muscle relaxant to use is influenced by the speed of onset, duration of action, method of elimination and side effect profile of the various agents. Except in the case of very brief procedures, NDMR are used for relaxation during maintenance of anesthesia. Commonly-used NDMR are rocuronium.
and cis-atracurium. Table 15 presents the relevant pharmacology and includes an older NDMR, pancuronium which is a long-acting agent that is still in use. Atracurium, another short-acting NDMR, is included in Chapter 6 (Drug Finder) although it is no longer available in many jurisdictions.

**Figure 20 Neuromuscular junction**

![Neuromuscular junction diagram](image)

1. nerve terminal 2. sarcolemma 3. acetylcholine vesicles 4. acetylcholine receptors 5. mitochondrion.

Image from Wikimedia Commons, used under GNU Free Documentation License, version 1.2. Drawn by user Dake.

Many factors may exaggerate or prolong the effects of NDMR. These include:

- factors which increase the susceptibility of the neuromuscular junction to NDMR: muscular dystrophies, myasthenia gravis, hypocalcemia, hypermagnesemia, acid-base abnormalities and many drugs (aminoglycosides, lithium, diuretics, volatile anesthetics)

- factors which delay metabolism or excretion: hypothermia, renal insufficiency and liver disease

The reader is referred to Chapter 6 for a more complete discussion of the pharmacology of the non-depolarizing muscle relaxants.

During maintenance of anesthesia, the degree of muscular paralysis is best monitored using a peripheral nerve stimulator. The anesthesiologist observes the magnitude and number of twitches in response to a series of four electrical stimuli (2 per second) applied over the ulnar nerve. The importance of carefully titrating NDMR is two-fold:

- to ensure sufficient muscle relaxation during the procedure

- to ensure the ability to adequately reverse muscle relaxation at the end of the procedure
Table 15 Dose, onset, duration, elimination, and effects of NDMR

<table>
<thead>
<tr>
<th></th>
<th>PANCURONIUM</th>
<th>ROCURONIUM</th>
<th>CIS-ATRACURIUM</th>
</tr>
</thead>
<tbody>
<tr>
<td>INTUBATING DOSE (MG/KG)</td>
<td>0.08 - 0.1</td>
<td>0.6 - 1</td>
<td>0.15 - 0.25</td>
</tr>
<tr>
<td>REPEAT DOSE (MG/KG)</td>
<td>0.02</td>
<td>0.1</td>
<td>0.02</td>
</tr>
<tr>
<td>ONSET (MIN)</td>
<td>4-5</td>
<td>1.5</td>
<td>1.5-3</td>
</tr>
<tr>
<td>DURATION AFTER INTUBATING DOSE (MIN)</td>
<td>60 - 90</td>
<td>30 - 60</td>
<td>40 - 75</td>
</tr>
<tr>
<td>METABOLISM/Elimination</td>
<td>80% renal</td>
<td>&gt;70% hepatic</td>
<td>77% Hoffmann elimination</td>
</tr>
<tr>
<td>SIDE EFFECTS</td>
<td>muscarinic blocker</td>
<td>weak muscarinic blocker</td>
<td>none</td>
</tr>
</tbody>
</table>

Emergence
During the emergence phase of anesthesia, the patient begins to return to his pre-operative state of consciousness. In most cases, the anesthesiologist aims to awaken the patient at the end of the operative procedure prior to transfer to the post-anesthetic care unit (PACU). How “awake” must the patient be? Ideally the patient is conscious enough to obey commands and support his own airway. At the very least, the patient must have adequate spontaneous ventilation but may need minimal assistance to maintain patency of the airway. In between these two states lies a wide spectrum of level of consciousness. Patient factors as well as the anesthetic technique determine the rate at which emergence from general anesthesia occurs.

Spontaneous and Active Reversal
Emergence requires the offset of effect of the anesthetic agents. This is achieved by administering the anesthetic drugs in appropriate doses at the appropriate time according to the anticipated length of the procedure. The anesthesiologist relies on the normal metabolism and excretion of drugs to achieve offset of effect. Active reversal of drug effect through the administration of another drug also plays a role in emergence. The most common example of this is the reversal of muscle relaxation which is discussed in greater detail below. Aside from muscle relaxants, anesthetic agents are

Contraindications
The major contraindication to the use of any muscle relaxant is the inability to provide airway and ventilatory control. A patient who is known or suspected to be a difficult intubation or a patient who has a fixed airway obstruction should not receive muscle relaxants prior to having the airway secured.
rarely actively reversed in order to achieve emergence. There is no “antidote” to the inhaled agents; offset of effect relies on the timely discontinuation of administration followed by excretion through the lungs. While an opioid antagonist (naloxone) exists, there are several disadvantages to using it to reverse opioid effect at the end of surgery. Firstly, unless very carefully titrated, its use will lead to a startled, hyper-alert patient who complains of pain at the operative site. Hypertension, tachycardia, myocardial ischemia and pulmonary edema may result. Secondly, the duration of effect of the antagonist is shorter than that of many of the opioid agonists therefore “re-narcotization” in the PACU is a risk. Finally, naloxone is an expensive drug whose use adds unnecessarily to the cost of the anesthetic. Flumazenil is a specific benzodiazepine antagonist which may play a role in the occasional surgical patient whose decreased level of consciousness is attributed to benzodi- azepines. Like naloxone, flumazenil has a shorter duration of action than most of the benzodiazepine agonists therefore rebound sedation may occur.

**Extubation**

If an endotracheal tube is used to maintain the airway intra-operatively, it must be removed at some point during the emergence phase of anesthesia. It is important to time the extubation properly, so as to avoid the potential post-extubation complications:

- airway obstruction
- aspiration
- inadequate ventilation
- laryngospasm

If the patient meets three simple criteria, most emergence complications can be avoided. The anesthesiologist must ensure that:

- the patient has regained their drive to breathe. Sufficient offset of effect of opioids is required for the patient to resume and maintain spontaneous respiration.
- the patient has normal muscle strength. A “weak” patient will not have enough strength to keep the tongue from falling to the back of the pharynx and causing airway obstruction. Muscle strength is also required to achieve satisfactory tidal volumes. Adequate muscle strength is required for the cough reflex which protects the airway from aspiration.
- the patient is awake enough to obey commands. An adequate level of consciousness is required in order for the patient to protect his airway from aspiration and to avoid laryngospasm.

**Laryngospasm** (reflexive closure of the vocal cords) deserves special mention. Laryngospasm is the airway’s response to irritation. It can occur immediately after ex-
tubation, leading to total airway obstruction, particularly in children and young adults. Exubating the patient at a deep plane of inhalational anesthesia (when the reflex is blunted) is one way to avoid laryngospasm but is an approach that is only safely applied to the pediatric patient. In adults (and pediatric patients), performing extubation when the patient is wide awake (where consciousness abolishes the reflex) will decrease the risk of post-extubation laryngospasm. Practicing anesthesiologists understand that extubating the patient at a light plane of anesthesia (not awake, but not “asleep” either) increases the risk of post-extubation laryngospasm.

**Reversal of Muscle Relaxation**

The action of all non-depolarizing muscle relaxants must be reversed prior to emergence from anesthesia. The anticholinesterase drugs, sometimes termed “reversal agents” are edrophonium, neostigmine and pyridostigmine (Table 16), with neostigmine being most commonly used.

The anticholinesterases reverse the effects of the NDMR. However, in order for them to be completely effective, some degree of spontaneous recovery from the NDMR block must be present prior to administration of the anticholinesterase. Adequacy of reversal is assessed clinically. The peripheral nerve stimulator is used while the patient is still unconscious. Traditionally, the anesthesiologist “eyeballs” the number of twitches and presence of fade although this technique is known to result in an underestimation of the degree of residual blockade. Newer anesthetic machines are equipped to assess the same indices by measuring mechanical deflection of the thumb. The most important indicators are clinical and are measured in the awake patient. A strong hand grip and the ability to lift the head off the bed for 5 seconds reliably indicate the return of adequate muscular strength.

**Mechanism of action**

Anticholinesterases act in the synaptic cleft of the neuromuscular junction. Here, they inhibit the action of cholinesterase, thereby decreasing the rate of breakdown of acetylcholine (Ach). The increased concentration of Ach in turn displaces the NDMR from the Ach receptors and thus restores normal neuromuscular transmission.

**Dose, onset, duration and elimination**

Relevant pharmacokinetic facts are summarized in Table 16.

**Effects**

Unfortunately, the anticholinesterase drugs potentiate the action of Ach at muscarinic receptors as well as at the nicotinic receptors of the neuromuscular junction. This can lead to all of the symptoms that are associated with excessive parasympathetic tone such as bradycardia, heart block, increased airway secretions, broncho-
spasm, intestinal spasm, increased bladder tone and pupillary constriction. These effects are minimized by administering an anticholinergic (atropine or glycopyrrole) along with the anticholinesterase.

The reader is referred to Chapter 6 for a more complete discussion of the pharmacology of the anticholinesterase agents as well as the anticholinergics that must accompany their administration.

**Contraindications**
Anticholinesterases are contraindicated in patients with gastrointestinal obstruction. They should be used with caution in patients with bradycardia, asthma, seizure disorders and Parkinson’s disease. An overdose can cause a cholinergic crisis.

---

**Table 16** Dose, onset, duration, elimination of anticholinesterases

<table>
<thead>
<tr>
<th></th>
<th>EDROPHONIUM</th>
<th>NEOSTIGMINE</th>
<th>PYRIDOSTIGMINE</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>DOSE (MG/KG)</strong></td>
<td>0.5 - 1.0</td>
<td>0.025 - 0.075</td>
<td>0.1 - 0.3</td>
</tr>
<tr>
<td><strong>ONSET (MIN)</strong></td>
<td>rapid (1)</td>
<td>intermediate (5)</td>
<td>delayed (10)</td>
</tr>
<tr>
<td><strong>DURATION (MIN)</strong></td>
<td>40 - 65</td>
<td>55 - 75</td>
<td>80 - 130</td>
</tr>
<tr>
<td><strong>RENAL EXCRETION</strong></td>
<td>70%</td>
<td>50%</td>
<td>75%</td>
</tr>
</tbody>
</table>
Chapter 3 Review

Review 3.1 The Intra-operative Phase

Question 1 of 10
The abbreviation “MAC” stands for:

- A. Minimal arterial concentration
- B. Minimal alveolar concentration [Correct]
- C. Maximal arterial concentration
- D. Maximal alveolar concentration
In this chapter, you will develop an understanding of both the goals and challenges of the recovery phase of anesthesia with a special focus on postoperative pain management.
Recovery

Goals of Recovery
At the end of the operative procedure, care and monitoring of the patient is handed over from the anesthesiologist to the nurse in the post-operative care setting as the patient enters the period of recovery. For most patients, this occurs in the Post-Anesthetic Care Unit (PACU). However, some patients, such as those requiring prolonged post-operative ventilation or close hemodynamic monitoring, may instead be admitted directly to the Intensive Care Unit. Prior to transporting the patient from the operating room, the anesthesiologist must ensure the presence of the following:
- patent airway (provided either by an awake patient, oral airway or endotracheal tube)
- adequate ventilation
- stable hemodynamics
- adequate pain control

Any identified problems must be corrected before leaving the operating room to avoid transporting an unstable patient.

On arrival in the PACU, the airway patency, breathing and circulation are immediately assessed. Supplemental oxygen is provided as indicated. Routine monitors are applied. The anesthesiologist then gives the attending nurse pertinent information about the patient’s past medical history and intra-operative course. The latter includes details of the nature of the procedure, anesthetic technique, fluid balance and any intra-operative complications. Finally, instructions regarding monitoring, management of fluids, pain and nausea as well as discharge plans are given.

Monitoring in the PACU is an extension of that provided in the operating room. The patient is observed for potential complications, both surgical and anesthetic. When discharge criteria are met, the patient is transferred to their ultimate destination: the ward for inpatients or the same day surgery unit for outpatients.

Criteria for discharge are stratified into “Phase 1” (which determines when the patient is able to be transferred from the PACU to the ward) and “Phase 2” (which addresses home-readiness and only applies to those patients having “same day
surgery”). A scoring system (Aldrete score) has been developed that grades the patient’s colour, respiration, circulation, consciousness and activity on a scale of 0-2.

For Phase 1 recovery, the patient must:

- be showing no signs of respiratory depression for at least 20-30 minutes after last dose of parenteral opioid.
- be easily aroused.
- be fully oriented to person, place and time.
- be able to maintain and protect the airway on his own including evidence of a strong cough.
- have stable vital signs for at least 30 minutes.

It is also important that pain and post-operative nausea and vomiting are controlled prior to PACU discharge and that there are no ongoing surgical concerns such as surgical site bleeding. Most healthy patients undergoing routine surgery will meet the PACU discharge criteria within 60 minutes.

Prior to being discharged home (from the same day surgery unit), the patient must demonstrate the return of cognitive function, ambulation and the ability to take oral liquids.

Many types of complications can occur in the PACU. Some of them, such as airway obstruction, aspiration, post-operative hemorrhage and myocardial ischemia are life-threatening but occur relatively infrequently. With expertise in airway management and cardiovascular resuscitation, the anesthesiologist is well-positioned to detect and manage these critical events.

Hypertension is commonly seen in the PACU and has many possible underlying causes. The patient may have pre-existing essential hypertension which is poorly controlled and may be exacerbated by the omission of their usual medication on the day of surgery. Other factors can lead to hypertension in the PACU such as full bladder (which the patient may not recognize), pain and importantly, hypoxemia and hypercarbia.

Fortunately, the most common PACU complications are not usually life-threatening but are important to recognize and manage nonetheless. A brief discussion follows below.
Post-operative Nausea and Vomiting
One of the most common problems encountered in the PACU is *post-operative nausea and vomiting* (PONV). PONV is unpleasant for both patient and staff. Moreover, it places the patient at risk for aspiration of gastric contents, particularly if airway reflexes are blunted due to the residual effects of opioids, inhaled agents and muscle relaxants.

When severe PONV is encountered, it is important to rule out sinister causes such as myocardial ischemia, bowel obstruction or raised intracranial pressure. More commonly, the cause is multifactorial with patient, surgical and anesthetic factors contributing. The many risk factors for PONV are outlined in Table 17, the most important (and statistically-robust) factors highlighted in bold.

The best approach to PONV is prevention. Attention should be paid to the most emetogenic anesthetic drugs which are nitrous oxide and neostigmine. Minimizing the dose of neostigmine to less than 2 mg appears to eliminate its emetogenic effect and should be considered if neuromuscular function allows. The modification of anesthetic factors (such as avoiding the use of nitrous oxide) is at least as effective as the administration of a prophylactic *antiemetic* agent. Current guidelines recommend that prophylaxis (*ondansetron* and/or *dexamethasone*) be administered selectively to moderate or high risk patients. These agents are administered during anesthesia rather than pre-operatively. Interestingly, ondansetron is much more effective as a treatment for PONV than as a preventive, where its NNT (number needed to treat) is around 5. It is also not free of side effects such as headache, constipation and elevated liver enzymes.

In the PACU, PONV is best treated with hydration and intravenous antiemetics such as ondansetron (if not used as prophylaxis), *prochlorperazine* or *dimenhydrinate*. PONV may act as a limiting factor to the delivery of opioid analgesia; patients at high risk of PONV benefit from opioid-sparing analgesic techniques such as peripheral nerve blocks or neuraxial analgesia.

<table>
<thead>
<tr>
<th>Table 17 Risk factors for PONV</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>PATIENT</strong></td>
</tr>
<tr>
<td>female</td>
</tr>
<tr>
<td>history of PONV</td>
</tr>
<tr>
<td>history of motion sickness</td>
</tr>
<tr>
<td>non-smoking</td>
</tr>
<tr>
<td>age &lt;12 years</td>
</tr>
<tr>
<td>obesity</td>
</tr>
</tbody>
</table>
**Shivering**

Shivering is a response controlled via the hypothalamus aimed at generating endogenous heat. The cost of this heat production is a fourfold increase in oxygen consumption and carbon dioxide production. This can precipitate myocardial ischemia or respiratory failure in patients with limited coronary or ventilatory reserve. Most commonly, shivering is a direct response to hypothermia and is best treated with aggressive rewarming techniques such as forced air warming systems. In the PACU, one often observes normothermic patients who shiver. This is a poorly understood effect of residual volatile gases and often abates with the administration of small doses of intravenous meperidine.

**Pain**

Pain is a consequence of all but the most minor surgical procedures. Aside from its inherent unpleasantness, it can lead to hypertension, tachycardia, myocardial ischemia and respiratory failure. Post-operative pain is most effectively approached through preventive measures taken in the operating room. Occasionally, this is achieved with a nerve block or the administration of central neuraxial local anesthetics and/or opioids.

More commonly, however, the anesthesiologist titrates the dose and timing of intravenous opioids to anticipate an awake, comfortable patient at the end of the procedure. A brief discussion of the methods of continued post-operative pain control follows in the next chapter.
Appropriate pain control not only contributes to patient comfort, it decreases the incidence of post-operative complications such as cardiac ischemia, pulmonary atelectasis and delirium. Many factors need to be considered when determining the method of post-operative analgesia. The most important factors are the patient’s medical history and the nature of the surgical procedure. Oral analgesics (acetaminophen, codeine phosphate, non-steroidal anti-inflammatories) or intramuscular opioids may be sufficient in many cases. However, often, more sophisticated techniques are required.

**Patient-controlled analgesia**

Patient-controlled analgesia (PCA) permits the patient to administer the delivery of his own analgesic by activating a button, which then triggers the intravenous delivery of a predetermined dose of an opioid such as morphine. Limits are set on the number of doses per four-hour period and on the minimum time that must elapse between doses (lockout interval). The pharmacokinetic advantage of PCA is that by self-administering frequent, small doses, the patient is able to come closer to achieving a steady state analgesic level in the blood, avoiding the high peaks and low troughs that can be found with intermittent (intramuscular) opioid administration. Indeed, PCA has been shown to provide equivalent analgesia with less total drug dose, less sedation, fewer nocturnal disturbances and more rapid return to physical activity. In addition, patient acceptance is high since patients have a significant level of control over their pain management.

PCA analgesia is not without side effects, the most common of which is nausea and vomiting. Excessive sedation and pruritis may also be seen. Standardized orders provide “as needed” orders for medications to counteract both nausea and pruritis.

Although it does not obviate the need for close monitoring, PCA frees nursing personnel from administering analgesic medication. Since patients titrate their own therapy with PCA, they must be capable of understanding the principle, willing to participate and physically able to activate the trigger. Consequently, use is prohibited
at the extremes of age as well as in very ill or debili-
tated patients.

PCA would be appropriately used for patients recover-
ing from breast reconstruction or lumbar spine decom-
pression and fusion. Typically, the PCA modality is
used for 24-72 hours. The patient must be capable of
oral (fluid) intake prior to converting from PCA to oral
analgesics, a factor which is most relevant for those re-
covering from bowel surgery.

---

**Central Neuraxial Analgesia**

Central neuraxial analgesia involves the delivery of lo-
cal anesthetics and/or opioids to either the intrathecal
(spinal) space or the epidural space.

Because intraspinal catheters are rarely used, intrathe-
cal analgesia is usually an extension of a “one-shot” spi-
nal anesthetic used intra-operatively. Opioids added to
the (spinal) local anesthetic solution provide long-
lasting analgesia after a single injection, lasting well
into the post-operative period. The duration of effect is
directly proportional to the water-solubility of the com-
 pound, with hydrophilic compounds such as morphine
providing the longest relief.

Epidural catheters are safe and easy to insert. Contrain-
dications can be reviewed in Table 11. Epidural analge-
sia can be used to provide pain relief for days through
the infusion of a solution containing local anesthetic,
opioid or both. The infusion is usually delivered con-
tinuously. Intermittent or “bolus” doses lack titratabil-
ity and are associated with a higher incidence of side
effects such as respiratory depression. Continuous
epidural infusions provide a steady level of analgesia
while reducing the side-effects associated with bolus
administration.

Overall, epidural analgesia can provide highly effective
management of post-operative pain. It is believed to
lead to a decreased stress response to surgery, improved post-operative pulmonary function and in high risk patients, decreased cardiac morbidity. Successful management relies on proper patient selection, appropriate catheter placement (depending on the level of the surgical site), adequate post-operative monitoring and specific training of personnel to identify and treat complications (including inadequate analgesia).

Epidural analgesia is commonly used after major intra-abdominal or thoracic surgery. A common use would be following (open) abdominal aortic aneurysm repair where the catheter might be left in for 48-72 hours. Occasionally, the need for post-operative thrombosis prophylaxis triggers the removal of the catheter as catheters should not be removed or left indwelling in the anticoagulated patient.

**Peripheral Nerve Blocks**

Almost any peripheral nerve that can be reached with a needle can be blocked with local anesthetics. The brachial plexus, intercostal and femoral nerves are examples of nerves which are commonly blocked to provide post-operative analgesia. A block may be used as the sole method of post-operative analgesia or it may be useful as an adjunct to decrease the required dose of systemic opioids. Some peripheral nerve sites (e.g. the brachial plexus) lend themselves to the insertion of catheters for the continuous infusion of local anesthetics. In the absence of catheter insertion, the major drawback of this method of post-operative analgesia is that the duration of effect of a single block is limited, usually to less than 18 hours.

A typical example of the use of a peripheral nerve block for post-operative pain would be the use of a femoral/sciatic nerve block for a patient undergoing total knee arthroplasty. The block would be augmented with oral opioids and other adjuncts.
Chapter 4 Review

Review 4.1 The Post-operative Phase

Question 1 of 10
Shivering can lead to ALL BUT ONE of the following. Indicate the exception:

- A. Myocardial ischemia
- B. Increased oxygen consumption
- C. Decreased carbon dioxide production
- D. Interference with monitor signal interpretation

Check Answer
In this chapter, you will be introduced to several subsets of patients who present anesthetic concerns in addition to the ones previously discussed. You will develop an understanding of how anesthetic care is modified to accommodate these “special patients”.
Malignant Hyperthermia

Malignant Hyperthermia (MH) is a potentially life-threatening pharmacogenetic disorder characterized by the onset of a hypermetabolic crisis in response to certain triggers. Since the usual triggers are succinylcholine and volatile anesthetics, MH is known as “the anesthesiologist’s disease”.

Although MH is now known to be a genetically heterogenous disorder, up to 70% of cases involve a mutation in a gene on chromosome 19 which encodes the ryanodine receptor protein. The ryanodine receptor is located on the sarcoplasmic reticulum. This particular mutation shows autosomal dominant inheritance pattern with variable penetrance. The other known causative gene for MH is CACNA1S, which is responsible for a voltage-gated calcium channel α-subunit. Several other chromosomal loci have been linked to MH although the specific genes have not yet been identified.

The MH-associated mutations cause an abnormality in skeletal muscle metabolism whereby uncontrolled intracellular release of calcium leads to sustained muscular contraction and cellular hypermetabolism. One very rare neuromuscular disease, called central core disease, is known to be associated with MH, while other more common neuromuscular disorders, such as Duchenne Muscular Dystrophy, are possibly associated with MH. Although rare (the incidence is reported to be 1 in 126,000 general anesthetics), MH does occur in geographical clusters.

The clinical manifestations of an MH crisis reflect the hypermetabolic state and may occur intra-operatively or post-operatively. The earliest sign is tachycardia followed by evidence of increased carbon dioxide (CO2) production. Increase CO2 production manifests as tachypnea in a spontaneously-breathing patient or raised end-tidal CO2 levels in a mechanically-ventilated patient. Skeletal muscle rigidity is prominent. Hyperthermia is often delayed. Hypoxemia, acidosis, hyperkalemia, dysrhythmias and hemodynamic instability may ensue as the reaction progresses. Without treatment, the mortality rate for MH reaction is exceedingly high; even with prompt treatment, mortality may be as high as 10%.
Successful treatment of an MH crisis requires prompt recognition, discontinuation of triggering agents and administration of **dantrolene**. Dantrolene is a direct skeletal muscle relaxant which acts at the muscle cellular level. It is administered intravenously in 2.5 mg/kg doses until clinical signs show reversal of the hypermetabolic state. In most cases the symptoms will abate with a total dose of less than 20 mg/kg but the anesthesiologist must not hesitate to administer more dantrolene if the clinical indicators warrant. The remainder of treatment is supportive and involves hyperventilation with 100% oxygen, fluid administration and active cooling if temperature is elevated. One should be prepared to treat hyperkalemia and cardiac dysrhythmias. The surgical procedure should be terminated as quickly as is feasible after which the patient is transferred to the intensive care unit. Dantrolene should be continued in 1-2 mg/kg doses, every four hours for at least 24 hours. Patients should be monitored for recrudescence of the reaction as well as for complications such as myoglobinuria, renal failure and disseminated intravascular coagulation (DIC).

A very important component of care for the patient who has had an unexpected MH reaction is counseling and education for both patient and family. While the patient himself is known to be MH susceptible, his family members must be assumed to be MH susceptible until it is proven otherwise. A diagnosis of MH has implications for employment, life insurance premiums and for future anesthetic management. Specialists at “MH clinics” are best able to advise the patient and his family; it is the duty of the attending anesthesiologist to make that referral. At the MH clinic, the appropriateness of muscle biopsy will be discussed. Muscle biopsy can rule out (or in) MH susceptibility in a family member but is painful and requires an anesthetic. It is expected that in the near future, the development of a genetic blood test will obviate the need for the invasive muscle biopsy in the majority of patients.

The anesthetic management of a patient known to be MH-susceptible is straightforward. Dantrolene prophylaxis may (rarely) be given preoperatively to high risk patients. Intra-operatively, standard monitoring is used with an emphasis on end-tidal CO2, O2 saturation and temperature measurement. Triggers are avoided by using a “trigger-free” anesthetic machine which is free of vapourizers, and has been flushed clear of residual volatile gases. An anesthetic technique which does not involve the use of succinylcholine or volatile anesthetic gases is chosen. Post-operatively, the patient is usually observed in the post-anesthetic care unit for an extended period of time (e.g. 4 hours). If no suspicious signs (such as fever or unexplained tachycardia) are detected, routine post-operative care follows. The patient...
can be discharged home if instructions regarding worrisome symptoms have been given and understood, and if the patient has reasonable access to the hospital from home, should problems arise.
Physiologic Changes of Pregnancy

Physiologic and anatomic changes develop across many organ systems during pregnancy and the postpartum period. Metabolic, hormonal and physical changes all impact on anesthetic management. To the anesthesiologist, the most important changes are those that affect the respiratory and circulatory systems.

Respiratory System

There is an increased risk of difficult or failed intubation in the parturient. This is primarily due mucosal vascular engorgement which leads to airway edema and friability. Laryngoscopy can be further impeded by the presence of large breasts.

In addition, the parturient is at risk for aspiration of stomach contents. During pregnancy, the stomach is displaced cephalad and intragastric pressure increases. Gastric motility is decreased and gastric secretions increase. This, combined with a decrease in the integrity of the gastroesophageal junction predisposes to pulmonary aspiration of gastric contents. In fact, airway complications (difficult intubation, aspiration) are the most common anesthetic cause of maternal mortality. The best means of avoiding this outcome is to avoid general anesthesia (by using a regional technique) and thus maintain intact laryngeal reflexes. If a general anesthetic is required, NPO status for eight hours is preferred although not achievable in an emergency situation. Pretreatment of all parturients with a non-particulate antacid (30 cc sodium citrate p.o.) as well as with a histamine blocker (ranitidine 50 mg IV) is important. Finally, a rapid sequence induction with cricoid pressure is mandatory.

With the apnea that occurs at induction of anesthesia, the parturient becomes hypoxic much more rapidly than the non-pregnant patient. The reason for this is two-fold. Firstly, oxygen requirement has increased by 20% by term. Secondly, the functional residual capacity (FRC), which serves as an “oxygen reserve” during apnea, has decreased by 20% due to upward displacement of the diaphragm.

Adequate ventilation must be maintained during anesthesia. By term, minute ventilation has increased to 150% of baseline. This results in a de-
crease in PaCO2 (32 mmHg). The concomitant rightward shift in the oxyhemoglobin *dissociation curve* allows increased fetal transfer of O2.

**Circulatory Changes**

Blood volume increases by 40% during pregnancy in preparation for the anticipated 500-1000 cc average blood loss during vaginal or Caesarian delivery, respectively. This is significant for two reasons. Firstly, the normal signs of hypovolemia may not be seen until a relatively greater blood loss has occurred. Secondly, the expanded intravascular volume may not be tolerated by parturients with concomitant cardiovascular disease, such as mitral stenosis.

Due to the increasing uterine size, **aortocaval compression** (obstruction of the inferior vena cava and aorta) becomes relevant in the third trimester. When the pregnant patient is in the supine position, the heavy gravid uterus compresses the major vessels in the abdomen leading to maternal hypotension and fetal distress. Left lateral tilt, usually achieved with a pillow under the woman’s right hip, is an important positioning maneuver.

**Labour Analgesia**

There are many methods of relieving the pain and stress of labour. The non-invasive methods, such as **transcutaneous electrical nerve stimulation** (TENS), hypnosis and massage require a well-prepared patient who is able to accept the incomplete relief that such methods inevitably provide. Invasive methods, such as inhaled (nitrous oxide), intravenous (opioids) or regional (epidural) are associated with side effects and risks to both fetus and mother. Epidural labour analgesia will be discussed briefly in this section.

The pain of the first stage of labour is referred to the T10-L1 somatic areas. This extends to include sacral segments (S2-4) during the second stage. Thus, the principle of epidural analgesia is to administer local anesthetics (with or without opioids) into the epidural space to block the aforementioned spinal segments.

The primary advantages of epidural analgesia are its high degree of effectiveness and safety. The patient remains alert and cooperative. In the absence of complications, there are no ill effects on the fetus. Epidural analgesia can be therapeutic for patients with pre-eclampsia or cardiac disease where a high catecholamine state is detrimental. Finally, the level and intensity of an epidural block can be extended to provide anesthesia for operative delivery (Caesarian section).

As well as blocking sensory fibres, local anesthetics in the epidural space interrupt transmission along sympathetic and motor neurons. The hypotension associated with sympathetic blockade can be minimized by a one litre bolus of crystalloid prior to institution of the block, slow titration of the local anesthetic, the use of lower concentrations of local anesthetic and vigilant guarding against aortocaval compression.
There is good evidence that a labour epidural is associated with a prolongation of the second stage of labour, due to the associated motor block. Whether it also leads to an increased incidence of operative delivery remains controversial. The degree of motor block can be minimized by using lower concentrations of local anesthetics along with opioid adjunct. The use of a local anesthetic infusion (as opposed to boluses or “top-ups”) may give a more consistent level of block, lower total dose of local anesthetic, less motor block and less risk of drug toxicity.

Anesthesia for Operative Delivery

The major causes of anesthetic morbidity and mortality in the pregnant patient are those related to the respiratory system. Because of the risks of aspiration and failed intubation, and the depressant effects of anesthetic agents on the fetus, general anesthesia is avoided (where possible) in the parturient undergoing Caesarian section. Regional anesthesia is the preferred technique and can be provided by administering spinal anesthesia or by extending the depth and height of an existing epidural block.

There are two situations where a regional technique would not be chosen for Caesarian section. The first would be in the presence of an absolute contraindication to regional anesthesia (Table 11). These include coagulopathy, hypovolemia, infection, certain cardiovascular conditions and patient refusal.

The second situation where a regional technique may not be appropriate is in the setting of severe fetal distress. In this setting, general anesthesia almost always allows the most rapid delivery of the compromised fetus. If the fetal heart rate is very low and the maternal airway appears favourable, then general anesthesia will be quickly induced.

General anesthesia in the parturient is unique in several respects which reflects the many physiologic changes in this patient population. The pregnant patient has a lower anesthetic requirement (MAC) and yet, paradoxically, is at higher risk of experiencing awareness under anesthesia. Other important considerations are the risk of aspiration, rapid desaturation and the need to avoid both neonatal depression and uterine atony.
With the patient in left lateral tilt, adequate IV access is ensured and the monitors are applied. The patient is prepped and draped prior to induction. After careful pre-oxygenation, a rapid sequence induction with cricoid pressure is performed. Generally speaking, no opioids are administered until delivery of the infant in order to avoid unnecessary neonatal depression. The patient is maintained on a 50% mixture of nitrous oxide and oxygen, and a low dose of volatile agent. The volatile anesthetics, in higher doses, can decrease uterine tone, which can lead to increased blood loss. After delivery of the fetus, a moderate dose of intravenous opioid is administered. As well, oxytocin is administered to augment uterine tone. The parturient must be extubated when fully awake so that intact laryngeal reflexes will protect against aspiration.

Post-operative pain management in the post-Caesarian section patient is usually straightforward as the lower abdominal incision is relatively well-tolerated. In the instance where intrathecal morphine was administered to the patient undergoing spinal anesthesia, up to 24 hours of pain relief can be achieved.
Not just a small adult...

The principles of pre-operative assessment, anesthetic management and post-operative care described earlier apply equally well to the pediatric patient. Specific variations in management of the pediatric patient result from differences in anatomy and physiology in this patient population, as compared to adult patients. Some of these differences are discussed briefly below.

Respiratory System

The pediatric airway differs from the adult airway in several respects. The occiput is relatively prominent in infants and young children. This means that the “sniffing position” is often best achieved with the head in the neutral position, without the use of a pillow. The relatively large tongue may hinder visualization of the larynx or contribute to upper airway obstruction under anesthesia. The epiglottis is long, angled and mobile. Because of this, a Magill blade is often used (in infants and young children) to lift the epiglottis directly to expose the larynx. The larynx itself is positioned higher (C4 vs. C6 in adult) and more anteriorly. The narrowest part of the pediatric airway is the subglottic region, at the level of the cricoid cartilage. Therefore, the use of a cuffed endotracheal tube (ETT) in a child less than 10 years of age is unnecessary and undesirable, as the narrow subglottic region provides its own seal. Because the trachea is narrowed, short and easily traumatized, appropriate selection of an ETT is critical. Recommended sizes of ETT by age are indicated in Table 18. Generally, the formula below predicts the correct tube size for children over one year of age.

\[
\text{ETT size} = 4 + \frac{\text{age}}{4}
\]

The pediatric airway is relatively more prone to obstruction than the adult airway. Infants are obligate nose breathers and the nares are small and easily obstructed by edema or mucus. Due to subglottic narrowing, a small amount of edema resulting from ETT trauma or pre-existing infection (trachitis or croup) can seriously compromise airway patency. Finally, laryngospasm is common in children. This complex and potentially life-threatening phenomenon can result...
from non-specific stimuli as well as from direct irritation of the vocal cords by blood or secretions. In order to avoid laryngospasm, pediatric patients are extubated either at a deep plane of anesthesia or wide awake.

The pediatric patient is more prone to hypoxemia than most adults. Like the obstetric patient, children have a slightly smaller functional residual capacity (FRC) (Table 19). The FRC acts as a reserve tank of oxygen during apneic periods. In addition, the pediatric patient has a markedly increased oxygen consumption which is usually maintained with an increased minute ventilation. The result of both of these factors is that the pediatric patient will desaturate much more rapidly during apnea. Adequate pre-oxygenation is key to the airway management of the pediatric patient.

**Cardiovascular**

Infants and young children have a heart-rate dependent cardiac output. This means that with bradycardia, their stiff left ventricles are unable to increase stroke volume to maintain cardiac output. This explains why bradycardia is undesirable in pediatric patients. Curiously, the pediatric patient is relatively “vagotonic”. In other words, their **vagus nerve** is dominant and they are prone to developing bradycardia in response to cer-

---

**Table 18** Pediatric ETT sizes

<table>
<thead>
<tr>
<th>AGE</th>
<th>ETT SIZE</th>
</tr>
</thead>
<tbody>
<tr>
<td>TERM NEWBORN</td>
<td>3.0-3.5</td>
</tr>
<tr>
<td>0-12 MONTHS</td>
<td>3.5</td>
</tr>
<tr>
<td>1-2 YEARS</td>
<td>4.0</td>
</tr>
<tr>
<td>3 YEARS</td>
<td>4.5</td>
</tr>
<tr>
<td>4-5 YEARS</td>
<td>5.0</td>
</tr>
<tr>
<td>6-8 YEARS</td>
<td>5.5</td>
</tr>
<tr>
<td>8-9 YEARS</td>
<td>6.0</td>
</tr>
<tr>
<td>10-12 YEARS</td>
<td>6.5</td>
</tr>
</tbody>
</table>

**Table 19** Oxygen reserve, delivery and consumption

<table>
<thead>
<tr>
<th></th>
<th>CHILD</th>
<th>ADULT</th>
</tr>
</thead>
<tbody>
<tr>
<td>FRC (mL/kg)</td>
<td>30</td>
<td>35</td>
</tr>
<tr>
<td>Va (mL/kg)</td>
<td>150</td>
<td>60</td>
</tr>
<tr>
<td>Va/FRC</td>
<td>5</td>
<td>1.5</td>
</tr>
<tr>
<td>VO2 (mL/kg./min)</td>
<td>7.5</td>
<td>3.5</td>
</tr>
</tbody>
</table>

FRC=functional residual capacity, VA=minute alveolar ventilation, VO2=minute oxygen consumption
tain types of noxious stimuli. Examples include hypoxemia and laryngoscopy. It is common practice, therefore, to pre-treat infants and young children with atropine just prior to the induction of anesthesia. Bradycardia in the pediatric patient must always be assumed to be a result of hypoxemia until proven otherwise.

**Fluids and Metabolism**

Management of fluid requirements follows the same principles described in the chapter on fluid management. The “4/2/1 rule” to calculate maintenance requirements applies equally well to the pediatric patient. There are some important differences, however. The blood volume of a child is greater, relative to their weight, compared to the adult (Table 20). This becomes important when calculating estimated blood loss as a percentage of the estimated blood volume as is done to guide to transfusion therapy.

The second important issue involves the type of maintenance fluid used. Because of its glucose and sodium concentrations, 2/3 D5W-1/3 N/S is appropriate for maintenance fluid administration in adults and children. In the operating room, we routinely administer N/S or R/L for maintenance because it is the crystalloid of choice for replacing blood volume and third space losses, which make up the bulk of the fluid needs in the intra-operative period. In infants and young children, however, it is less appropriate to use N/S or R/L for maintenance especially during prolonged cases. Firstly, the immature kidney is unable to handle an excessive sodium load. Secondly, the child’s liver glycogen stores may be insufficient to maintain normal serum glucose during a more prolonged period of fasting.

### Table 20 Blood volume

<table>
<thead>
<tr>
<th>AGE</th>
<th>BLOOD VOLUME (CC/KG)</th>
</tr>
</thead>
<tbody>
<tr>
<td>preterm neonate</td>
<td>90</td>
</tr>
<tr>
<td>term neonate</td>
<td>80</td>
</tr>
<tr>
<td>infant</td>
<td>75</td>
</tr>
<tr>
<td>child</td>
<td>70</td>
</tr>
<tr>
<td>adult</td>
<td>60-70</td>
</tr>
</tbody>
</table>

**Gastrointestinal**

Children, generally speaking, present a lower risk of regurgitation and aspiration than adult patients. As well, they will become dehydrated more readily during a period of fasting. Thus, NPO guidelines for pediatric patients are more liberal than in the adult population. For
example, it is common practice to allow clear fluids from 2-4-hours pre-operatively in children under 12 years of age. Infants may be allowed breast milk up to 4 hours pre-operatively and formula up to 6 hours pre-operatively, breast milk being more readily digestible than formula.

Central Nervous System, Behaviour
Anesthetic requirements (MAC) are higher in infants and children, compared to adults, with the peak occurring at 6 months of age.

By 6 months to a year of age, infants become sufficiently aware of their surroundings to feel anxiety in the immediate pre-operative period. They are generally less inhibited about expressing their anxiety than their adult counterparts. There are many different approaches to minimizing this anxiety which must be individualized according to the needs of the patient, her parents and the anesthesiologist. Pediatric patients may be pre-medicated with benzodiazepines, opioids or anti-cholinergics. Unfortunately the administration of a pre-medication (even orally) can be distressing for these patients. Furthermore, depending on the agent chosen, recovery may be delayed. In many centers a parent is allowed in the operating room for induction to avoid separation anxiety for the child. This requires additional personnel to prepare and stay with the parent throughout. Studies have failed to show a clear benefit to the child with this technique although the parents, for the most part, seem to prefer it. Inhalation ("mask") inductions are often used in order to avoid having to insert an IV in the awake child. However, for a struggling child, a mask induction may be more traumatic than an IV induction. The use of topical tetracaine (a local anesthetic) has made awake IV starts more feasible in this patient population.
As diverse and ever-evolving as operating room practice is, anesthesiologists also have many opportunities to expand their practice outside of this realm. The ensuing discussion will look more closely at some of these challenging roles.

Anesthesiologists may provide conventional anesthetic services in locations remote from the operating room, such as radiology, burn center, endoscopy unit, lithotripsy unit, electrophysiology lab, or cardiac investigation unit. The anesthesiologist is often requested to monitor and sedate patients in order to render interventional procedures safer and more palatable to the patient. General anesthesia outside the operating room poses unique problems for the anesthesiologist and certain risks for the patient. Nonetheless, with adequate preparation and appropriate care, general anesthesia can be carried out almost anywhere.

As an expert in airway management, vascular access and fluid resuscitation, the anesthesiologist is a key member of the trauma team. The team resuscitates the patient, establishes the extent of injury and carries out appropriate investigations. These initial steps are followed by definitive therapy, when necessary. The anesthesiologist’s involvement in the care of the trauma victim does not end in the emergency room. Often, these victims arrive in the operating room urgently, where resuscitation is vigorously continued before and during the anesthetic.

Many anesthesiologists spend a proportion of their clinical time working in the intensive care setting. With the use of sophisticated physiologic monitoring, ventilatory care, pharmacologic support and acute pain management, critical care medicine is a natural extension of the anesthesiologist’s role in the operating room. Many anesthesiologists pursue specialty training in critical care after their residency training.

Most hospitals have an Acute Pain Service (APS) run by the Anesthesia departments, for postoperative surgical patients. The APS initiates and supervises various pain management therapies such as patient-controlled analgesia (PCA) and epidural analgesia. These interventions are intensive and brief; they require constant and ongoing assessment to ensure safety and effectiveness. Acute pain management requires a special set of...
skills that contributes greatly to patient recovery and satisfaction.

Large hospitals usually involve anesthesiologists in managing patients with chronic pain. Typically, the approach to chronic pain is multidisciplinary in that patients are assessed by a social worker, psychologist or psychiatrist, orthopedic surgeon and physiotherapist. Most often, these very complicated patients are referred to the clinic after seeing many other physicians. After taking a detailed history and performing a physical exam, the anesthesiologist may institute a nerve block, using local anesthetic and/or steroids. Examples of the indications for some of the more commonly performed blocks are given below:

- Complex Regional Pain Syndrome (formerly Reflex Sympathetic Dystrophy): stellate ganglion block, lumbar sympathetic block
- Chronic Back Pain: caudal or lumbar epidural steroid injection
- Occipital Headaches: occipital nerve block
- Myofascial Pain: trigger point injections
- Chronic pancreatitis: celiac plexus block

Often, these patients are followed over a period of time, giving the anesthesiologist an opportunity to establish the long-term rapport not feasible in operating room practice. Despite the notion that the anesthesiologist doesn’t talk to his patients, this patient population must be treated with a good ear and a sympathetic tone if any treatment modality is to be successful.
Question 1 of 15
All BUT ONE of the following are true regarding malignant hyperthermia (MH). Indicate the exception:

- **A.** Newer volatile anesthetics such as desflurane are not believed to trigger MH.
- **B.** Nitrous oxide is safe.
- **C.** Dantrolene is the only therapeutic drug treatment.
- **D.** Mortality is as high as 10% even with prompt treatment.
In this chapter, the drugs that are commonly used in anesthetic practice are presented in reference format, grouped together by class. The drugs can be viewed on one page, with a window for each drug opening up with a touch. Alternatively, each drug has a page devoted to it in the balance of Chapter 6 that follows.
### Opioid agonists and antagonists
- Fentanyl
- Sufentanil
- Remifentanil
- Alfentanil
- Morphine Sulfate
- Meperidine
- Naloxone

### Muscle relaxants
- Rocuronium
- Cis-Atracurium
- Pancuronium bromide
- Atracurium
- Succinylcholine chloride

### Anticholinesterases and anticholinergics
- Neostigmine
- Glycopyrrolate
- Atropine Sulfate

### Induction agents
- Propofol
- Sodium Thiopental
- Ketamine
- Etomidate

### Inhaled agents
- Desflurane
- Sevoflurane
- Isoflurane
- Nitrous Oxide

### Anxiolytics
- Midazolam

### Antiemetics
- Ondansetron
- Dimenhydrinate
- Prochlorperazine

### Vasoactive agents
- Phenylephrine
- Ephedrine sulfate
- Epinephrine

### Local anesthetics
- Bupivacaine
- Lidocaine

### Miscellaneous
- Ketorolac tromethamine
- Diphenhydramine
- Dantrolene
Opioid agonists and antagonists

Drugs

1. Fentanyl
2. Sufentanil
3. Remifentanil
4. Alfentanil
5. Morphine Sulphate
6. Meperidine
7. Naloxone

FENTANYL

Class
Synthetic opioid analgesic (intermediate-acting); adjunct to anesthesia. Fentanyl can be used as an additive to spinal and epidural anesthesia/analgesia.

Mechanism of Action
Acts at the mu-and kappa opioid receptors.

Dose
General anesthesia: 1-20 ug/kg IV according to physical status, other agents used, duration and nature of surgery.

Onset
IV 4-6 minutes

Duration
IV 30-45 minutes

Elimination
Hepatic

Effects
CNS
Potent analgesic effects; some sedative effect. Rarely causes blurred vision, seizures. All of the depressant effects of fentanyl are potentiated by concurrent use of sedatives, volatile anesthetics and nitrous oxide.

CVS
Hypotension, bradycardia. The synthetic opioids are not direct myocardial depressants but they do reduce sympathetic drive which may result in decreased cardiac output in patients who are relying on sympathetic tone to support their circulation such as those in hypovolemic or cardiogenic shock.

Respiratory
Respiratory depression which at the extreme leads to apnea.

GI
Nausea, vomiting, biliary tract spasm, constipation.

Misc
Muscle rigidity
**SUFENTANIL**

**Class**
Synthetic opioid analgesic (intermediate-acting), adjunct to anesthesia.

**Mechanism of Action**
Acts at the mu- and kappa opioid receptors.

**Dose**
General anesthesia: 0.3-1 ug/kg IV, depending on patient condition, other agents used, nature and duration of surgery.

Infusion dose: 0.3-1 ug/kg/hour

**Onset**
1-2 minutes

**Duration**
20-40 minutes

**Elimination**
Hepatic

**Effects**

**CNS**
Potent analgesic properties and some sedative effect. All of the depressant effects of sufentanil are potentiated by concurrent use of sedatives, volatile anesthetics and nitrous oxide.

**CVS**
Bradycardia, hypotension. The synthetic opioids are not direct myocardial depressants but they do reduce sympathetic drive which may result in decreased cardiac output in patients who are relying on sympathetic tone to support their circulation, such as those in hypovolemic or cardiogenic shock.

**Respiratory**
Respiratory depression, which at the extreme leads to apnea.

**GI**
Nausea, vomiting, biliary tract spasm, constipation.

**Misc.**
Muscle rigidity
REMIFENTANIL

Class
Synthetic opioid analgesic (ultra short-acting); adjunct to anestheisa.

Mechanism of Action
Acts at the mu-and kappa opioid receptors.

Dose
On induction of general anesthesia: 0.3-1 μg/kg
For maintenance of general anesthesia: 0.1-1 μg/kg/minute (by infusion)
For sedation: infusion 0.05 – 0.1 μg/kg/minute

Onset
After single bolus: 1-1.5 minutes
After initiation of infusion: 3-5 minutes

Duration
5-10 minutes; context sensitive half time 3 minutes

Elimination
Non-specific blood-tissue esterases (end-organ independent)

Effects
CNS
Potent analgesic effects, sedation. “MAC sparing” allows up to 75% reduction in dose of co-anesthetics. All of the depressant effects of remifentanil are potentiated by concurrent use of sedatives, volatile anesthetics and nitrous oxide.

CVS
Exaggerated bradycardia, hypotension (compared with other opioids). The synthetic opioids are not direct myocardial depressants but they do reduce sympathetic drive, which may result in decreased cardiac output in patients who are relying on sympathetic tone to support their circulation, such as those in hypovolemic or cardiogenic shock.

Respiratory
Profound respiratory depressant which often leads to apnea.

GI
Nausea, vomiting.

Misc.
Can cause profound muscle rigidity. Not suitable for spinal or epidural use due to glycine additive. Rapid elimination requires initiation of post-operative analgesia (usually morphine) prior to emergence.
**ALFENTANIL**

**Class**
Synthetic opioid analgesic (short-acting); adjunct to anesthesia.

**Mechanism of Action**
Acts at the mu- and kappa opioid receptors.

**Dose**
5-50 ug/kg IV, according to physical status, other agents used, nature and duration of surgery.

**Onset**
1-2 minutes

**Duration**
20 minutes

**Elimination**
Hepatic

**Effects**

**CNS**
Analgesia, sedation. All of the depressant effects of alfentanil are potentiated by concurrent use of sedatives, volatile anesthetics and nitrous oxide.

**CVS**
Bradycardia, hypotension. The synthetic opioids are not direct myocardial depressants but they do reduce sympathetic drive, which may result in decreased cardiac output in patients who are relying on sympathetic tone to support their circulation, such as those in hypovolemic or cardiogenic shock.

**Respiratory**
Potent respiratory depression which at the extreme, leads to apnea.

**GI**
Nausea, vomiting, biliary tract spasm.

**Misc.**
Muscle rigidity, pruritis.
MORPHINE SULFATE

Class
Opioid analgesic (long acting). In anesthetic practice, its main use is for postoperative analgesia. Morphine is commonly used intravenously and for spinal or epidural anesthesia/analgesia.

Mechanism of Action
Active at the mu and kappa opioid receptors.

Dose
Adults: 2.5-15 mg IV/IM/SC
Children: 0.05-0.2 mg/kg IV/IM/SC

Onset
IV 5-10 minutes
IM 15-30 minutes

Duration
2-5 hrs IV/IM/SC

Elimination
Hepatic

Effects
CNS
Reliable analgesic effects; sedation. May cause blurred vision, syncope, euphoria, dysphoria. All of the depressant effects of morphine are potentiated by concurrent use of sedatives, volatile anesthetics, nitrous oxide and alcohol. Morphine’s depressant effects are also potentiated by antihistamines, phenothiazines, butyrophenones, MAOIs and TCAs.

CVS
May cause hypotension, hypertension, bradycardia, arrhythmias.

Respiratory
Respiratory depression which at the extreme leads to apnea. May cause bronchospasm or laryngospasm.

GI
Nausea, vomiting, constipation, biliary tract spasm.

Misc.
Releases histamine. May cause pruritis, urticaria, muscle rigidity, urinary retention.
MEPERIDINE

Class
Opioid analgesic (long acting). Traditionally used for postoperative pain but currently its use is restricted (in many hospitals) to the treatment of postoperative shivering.

Mechanism of Action
Acts at the mu and kappa opioid receptors.

Dose
In adults: 25-75 mg IV/IM (0.5-2 mg/kg)

Onset
IV: 3-8 minutes
IM: 10-20 minutes

Duration
2-4 hours IV/IM

Elimination
Hepatic

Effects
CNS
Causes dose-related sedation; variable analgesic effect. Delirium in older patients is often seen. May cause seizures if used in large doses or over an extended time frame due to the accumulation of its excitatory metabolite, normeperidine. May cause euphoria and dysphoria. All of the depressant effects of meperidine are potentiated by concurrent use of sedatives, volatile anesthetics, nitrous oxide and tricyclic antidepressants.

Respiratory
Respiratory depression which at the extreme leads to apnea. May promote bronchospasm in susceptible patients (those with asthma or COPD).

GI
Nausea, vomiting, biliary tract spasm, constipation.

Misc.
Effective in the treatment of postoperative shivering. May cause muscle rigidity, urticaria, pruritis.

Contraindications:
Meperidine must not be used in patients on monoamine oxidase inhibitors in whom it can cause a fatal reaction.
**NALOXONE**

**Class**
Opioid antagonist. Used to counteract the effects of opioids.

**Mechanism of Action**
Agonist at the opioid receptors.

**Dose**
For postoperative opioid depression: 1-2 µg/kg IV in 0.5-1 µg/kg boluses, q 2-3 minutes

For neonatal opioid depression: 10 µg/kg, q 2-3 minutes IV. Infusion: 1-5 µg/kg/hr

**Onset**
1-2 minutes

**Duration**
30-60 minutes

**Elimination**
Hepatic

**Effects**

**CNS**
Rapid reversal of opioid effect can cause delirium and severe pain.

**CVS**
When opioid effect is abruptly antagonized there can be significant sympathetic activation leading to hypertension, tachycardia and in susceptible individuals, myocardial ischemia and pulmonary edema.

**Misc.**
Due to the relatively short duration of action of naloxone, “re-narcotization” can be seen when it is used to treat respiratory depression caused by long acting opioids such as morphine. In this case, close monitoring is indicated and supplemental doses may be necessary.
SECTION 2

Muscle Relaxants

Drugs

1. Rocuronium
2. Cis-atracurium
3. Pancuronium bromide
4. Atracurium
5. Succinylcholine

ROCURONIUM

Class
Non-depolarizing muscle relaxant (NDMR); short-acting

Mechanism of Action
Competitive inhibitor at the acetylcholine receptors of the post-synaptic cleft of the neuromuscular junction.

Dose
Intubation: 0.45-.9 mg/kg
Maintenance bolus 0.1-0.2 mg/kg
Not usually administered by infusion

Onset
Dose-dependent:
1-1.5 minutes (0.6 mg/kg)
0.5-1.0 minutes (0.9 mg/kg)
Higher dose is therefore suitable for rapid sequence induction.

Duration
Dose-dependent:
31 minutes (0.6 mg/kg)
60 minutes (0.9 mg/kg)

Elimination
Hepato-biliary (70%); renal (10%)

Effects
CVS
Very weak vagolytic effect.

MSK
The neuromuscular blockade effects of non-depolarizing muscle relaxants are potentiated by succinylcholine, volatile anesthetics, aminoglycosides, lithium, loop diuretics, lidocaine, magnesium, lithium, ganglionic blockers, hypothermia, hypokalemia and respiratory acidosis.

Enhanced neuromuscular blockade is seen in patients with myasthenia gravis or myopathies.

The effects of NDMR are antagonized by cholinesterase inhibitors. Increased resistance to NDMRs is seen in patients on theophylline, burn patients and those with paresis or paralysis.

Misc.
Muscle relaxants are the most common cause of anaphylactoid reactions under general anesthesia.
**CIS-ATRACURIUM**

**Class**
Non-depolarizing skeletal muscle relaxant (NDMR); intermediate-acting

**Mechanism of Action**
Competitive inhibitor at the acetylcholine receptors of the post-synaptic cleft of the neuromuscular junction.

**Dose**
Intubation: 0.15-0.2 mg/kg
Maintenance bolus: 0.03 mg/kg
Maintenance infusion: 1-2 μg/kg/minute

**Onset**
Dose-dependent:
- 2 minutes (0.15 mg/kg)
- 1.5 minutes (0.2 mg/kg)

**Duration**
Dose dependent:
- 55 minutes (0.15 mg/kg)
- 65 minutes (0.2 mg/kg)
- 20 minutes (maintenance bolus 0.03 mg/kg)

**Elimination**
Hoffman elimination (77%), renal (16%)

**Effects**

**MSK**
The neuromuscular blockade effects of non-depolarizing muscle relaxants are potentiated by succinylcholine, volatile anesthetics, aminoglycosides, lithium, loop diuretics, lidocaine, magnesium, lithium, ganglionic blockers, hypothermia, hypokalemia and respiratory acidosis.

Enhanced neuromuscular blockade is seen in patients with myasthenia gravis or myopathies.

The effects of NDMR are antagonized by cholinesterase inhibitors. Increased resistance to NDMR is seen in patients on theophylline, burn patients and those with paresis or paralysis.

**Misc.**
Histamine release may occur with rapid administration or higher dosages. Produces 5-10x less laudanosine metabolite than atracurium. Muscle relaxants are the most common cause of anaphylactoid reactions under general anesthesia.
PANCURONIUM BROMIDE

Class
Nondepolarizing skeletal muscle relaxant (NDMR); long-acting

Mechanism of Action
Competitive inhibitor at the acetylcholine receptors of the post-synaptic cleft of the neuromuscular junction.

Dose
Intubation: 0.1 mg/kg IV
Maintenance bolus: 0.01-0.03 mg/kg

Onset
4-5 minutes

Duration
45-65 minutes

Elimination
Renal (80%), hepatic (minor)

Effects
CVS
Pancuronium has a vagolytic effect and therefore causes tachycardia and hypertension. Increased risk of arrhythmias in patients receiving tricyclic antidepressants and volatile anesthetics.

Respiratory
May promote bronchospasm, salivation.

MSK
The neuromuscular blockade effects of non-depolarizing muscle relaxants are potentiated by succinylcholine, volatile anesthetics, aminoglycosides, lithium, loop diuretics, lidocaine, magnesium, lithium, ganglionic blockers, hypothermia, hypokalemia and respiratory acidosis. Enhanced neuromuscular blockade is seen in patients with myasthenia gravis or myopathies.

The effects of NDMR are antagonized by cholinesterase inhibitors. Increased resistance to NDMR is seen in patients on theophylline, burn patients and those with paresis or paralysis.

Misc.
Muscle relaxants are the most common cause of anaphylactoid reactions under general anesthesia.
ATRACURIUM

Class
Nondepolarizing skeletal muscle relaxant (NDMR); short-acting

Mechanism of Action
Competitive inhibitor at the acetylcholine receptors of the post-synaptic cleft of the neuromuscular junction.

Dose
Intubation: 0.5-0.6 mg/kg IV
Maintenance bolus: 0.1-0.3 mg/kg IV

Onset
3-4 minutes

Duration
20-35 minutes

Elimination
Hoffman elimination, ester hydrolysis

Effects
MSK
The neuromuscular blockade effects of non-depolarizing muscle relaxants are potentiated by succinylcholine, volatile anesthetics, aminoglycosides, lithium, loop diuretics, lidocaine, magnesium, lithium, ganglionic blockers, hypothermia, hypokalemia and respiratory acidosis.

Enhanced neuromuscular blockade is seen in patients with myasthenia gravis or myopathies.

The effects of NDMR are antagonized by cholinesterase inhibitors. Increased resistance to NDMR is seen in patients on theophylline, burn patients and those with paresis or paralysis.

Misc.
Histamine release may occur with rapid administration or higher dosages. Produces an excitatory metabolite called laudanosine. Muscle relaxants are the most common cause of anaphylactoid reactions under general anesthesia.
SUCCINYLCHOLINE CHLORIDE

Class
Depolarizing muscle relaxant; ultra short-acting; Used for rapid sequence induction.

Mechanism of Action
Succinylcholine (Sch) attaches to nicotinic cholinergic receptors at the neuromuscular junction. There, it mimics the action of acetylcholine thus depolarizing the post-junctional membrane. Neuromuscular blockade (paralysis) develops because a depolarized post-junctional membrane cannot respond to subsequent release of acetylcholine.

Dose
Intubation: 1-1.5 mg/kg IV or 2.5-4 mg/kg IM

Onset
30-60 seconds after IV administration
2-3 minutes after IM dose

Duration
Duration is 4-6 minutes after IV dose
10-30 minutes after IM dose

Elimination
Hydrolysis by plasma pseudocholinesterase

Effects
CNS
Raised intracranial pressure and raised intraocular pressure.

CVS
Because of cross-reactivity at the muscarinic acetylcholine receptors, Sch causes vagal cardiac dysrhythmias. Bradycardia, junctional rhythm and sinus arrest can occur particularly if a second dose is administered and particularly in children.

Respiratory
Occasionally leads to bronchospasm and excessive salivation due to muscarinic effects. Intragastric pressure is increased thereby theoretically increasing the risk of regurgitation.

Misc.
Most of the other effects are secondary to the depolarization and subsequent contraction of skeletal muscle. Sch elevates serum potassium 0.3-0.5 mEq/L in normal patients. It can cause an exaggerated release of potassium (leading to fatal hyperkalemia) in those with neuromuscular or muscle disease. Post-operative myalgia is common particularly in young adults. Succinylcholine is a potent trigger of malignant hyperthermia.

Contraindications
There is a long list of absolute and relative contraindications which can be found in any Anesthesia text. A brief summary follows:

- Malignant Hyperthermia (MH) or presence of conditions associated with MH.
- Pseudocholinesterase deficiency. Deficiency can result as a genetic defect, as a consequence of various medications or a result of liver disease. The latter two causes are usually relative while the genetic defect can produce a complete lack of pseudocholinesterase activity in homozygous individuals. The use of succinylcholine in a patient with pseudocholinesterase deficiency leads to prolonged paralysis.

- Hyperkalemia.

- Presence of neurologic or muscular condition which would predispose to hyperkalemia after Sch-induced muscle contraction. Examples include recent paralysis (spinal cord injury or stroke), amyotrophic lateral sclerosis (ALS), Duchenne’s muscular dystrophy and recent burn or crush injury. Myotonia congenita or myotonia dystrophica can manifest sustained contraction with Sch.
Anticholinesterase and Anticholinergics

Drugs

1. Neostigmine
2. Glycopyrrolate
3. Atropine sulfate

**NEOSTIGMINE**

**Class**
Anticholinesterase. In anesthesia practice, neostigmine is used for the reversal of neuromuscular blockade. Internal Medicine specialists use neostigmine (or its relative, pyridostigmine) for the treatment of myasthenia gravis.

**Mechanism of Action**
Anticholinesterases inhibit the breakdown of acetylcholine (Ach) in the synaptic cleft by inhibiting the *cholinesterase* enzyme. As a result, Ach concentrations in the synaptic cleft are increased. Ach is then better able to compete with muscle relaxants for the Ach receptors and achieve depolarization of the muscle cell.

**Dose**
For reversal of neuromuscular blockade: 0.05 mg/kg

Dose should not exceed 5 mg

Must be administered with atropine 0.015 mg/kg or glycopyrrolate 0.01 mg/kg

**Onset**
5 minutes

**Duration**
55-75 minutes

**Elimination**
Hepatic, plasma esterases

**Effects**
Most of neostigmine’s effects are related to its cholinergic action. It must be given with an anticholinergic (atropine or more commonly glycopyrrolate) in order to minimize these effects.

**CNS**
Seizures

**CVS**
Bradycardia, AV block, nodal rhythm, hypotension

**Respiratory**
Increased oral and bronchial secretions, bronchospasm

**GI/GU**
Increased peristalsis, urinary frequency

**Misc.**
Overdose may produce cholinergic crisis. Neostigmine does not antagonize succinylcholine and may prolong phase 1 block of succinylcholine.
**GLYCOPRYRROLE**

**Class**
Anticholinergic. Clinical uses in anesthesia include the treatment of bradycardia; as an **antisialogogue** for awake intubation; or (most commonly) for counteracting the muscarinic effects of the anticholinesterases used for the reversal of neuromuscular blockade.

**Mechanism of Action**
An acetylcholine receptor blocker active at the muscarinic (not nicotinic) acetylcholine receptors. Therefore, glycopyrrolate has an anti-parasympathetic effect.

**Dose**
Antisialogogue: 0.1-0.2 mg IV/IM/SC in adults or 4-6 ug/kg IV/IM/SC in children

With anticholinesterase: 0.01 mg/kg IV

**Onset**
IV: ≤1 minute

IM/SC: 30-45 minutes

**Duration**
Vagal blockade: 2-3 hrs

Antisialogogue effect: 7 hours

**Elimination**
Renal, hepatic

**Effects**
Most effects result from the anticholinergic action of glycopyrrolate.

**CNS**
Confusion is less common than with atropine, as glycopyrrolate does not cross the blood brain barrier. May cause headache, dizziness, mydriasis, blurred vision, increased intraocular pressure.

**CVS**
Causes tachycardia at high doses and may cause bradycardia at low doses.

**GU**
Urinary hesitancy, retention

**Misc.**
Must be used in caution in patients with glaucoma, gastrointestinal or genitourinary obstruction.
ATROPINE SULFATE

Class
Anticholinergic. Clinical use in anesthesia includes the treatment of bradycardia and asystole; as a antisialagogue for awake intubation; or for counteracting the muscarinic effects of the anticholinesterases used for the reversal of neuromuscular blockade.

Mechanism of Action
An acetylcholine receptor blocker active at the muscarinic (not nicotinic) acetylcholine receptors. Therefore, atropine has an anti-parasympathetic effect.

Dose
Premedication 0.4-0.6 mg IV/IM in adults, 10-20 ug/kg IV/IM in children. Reversal 0.015 mg/kg IV with neostigmine 0.05 mg/kg IV.

Onset
Immediate

Duration
1-2 hours

Elimination
Hepatic, renal

Effects
Most effects result from the anticholinergic action of atropine.

CNS
Confusion, hallucinations, mydriasis, blurred vision, increased intraocular pressure

CVS
Tachycardia (high doses), bradycardia (low doses)

GI
Gastroesophageal reflux

GU
Urinary hesitancy, retention

Misc.
Has additive anticholinergic effects with antihistamines, phenothiazines, tricyclic antidepressants, mono-amine oxidase inhibitors and benzodiazepines. Potentiates sympathomimetics. May produce central anticholinergic syndrome.

Contraindications
Contraindicated in patients with narrow-angle glaucoma, gastrointestinal or genitourinary obstruction.
Induction Agents

Drugs

1. Propofol
2. Sodium thiopental
3. Ketamine
4. Etomidate

PROPOFOL

Class
Alkylphenol intravenous anesthetic agent. Used for induction of general anesthesia. Can also be used for maintenance of anesthesia or for sedation, in each case by continuous infusion.

Mechanism of action
Not well described.

Dose
Induction: 2-2.5 mg/kg IV for adults
Induction: 3-4 mg/kg IV for children
Maintenance of anesthesia: 100-200 ug/kg/minute
Sedation: 40-100 ug/kg/minute

Onset
Within one arm-brain circulation time (approximately 20 seconds).

Duration
Approximately 5-8 minutes after single induction dose. Offset of effect is more prolonged when administered as a continuous infusion.

Elimination
Rapid redistribution away from central nervous system (CNS) into lean body compartment accounts for prompt awakening. Metabolized by liver and extra-hepatic sites then excreted by kidney.

Effects
CNS
Profound CNS depressant, potentiating the depressant effects of opioids, sedatives and volatile anesthetics. Decreases cerebral metabolic rate and intracranial pressure. Occasionally excitement, tonic-clonic movements or opisthotonus is seen on induction with propofol.

CVS
Causes direct myocardial depression and vasodilation leading to hypotension. Propofol must be used with caution in patients with poor left ventricular function or critical coronary artery insufficiency or in those who are seriously ill or debilitated.

Respiratory
Depression of respiratory centre leads to brief apnea. Propofol effectively blunts the airway’s response to manipulation thus hiccoughing and bronchospasm are rarely seen.

Misc.
Pain on injection seen in up to 20%. Mild anti-emetic properties. Patients often experience pleasant dreams under anesthesia followed by a smooth, clear-headed emergence. Strict aseptic technique must be used when handling propofol as the vehicle is capable of supporting rapid growth of micro-organisms.

Contraindications
Egg or soy allergy.
**SODIUM THIOPENTAL**

**Class**
Short-acting barbiturate. Was used as an anesthetic induction agent but has largely been replaced by propofol. It is also useful as an anticonvulsant or for the rapid reduction of elevated intracranial pressure.

**Mechanism of action**
Decreases the rate of dissociation of the inhibitory neurotransmitter GABA from its receptors resulting in depression of the **reticular activating system**.

**Dose**
- 3-5 mg/kg IV for healthy adults
- 5-6 mg/kg IV for children
- 7-8 mg/kg IV for infants

Dose must be reduced considerably in unstable or fragile patients.

**Onset**
Within one arm-brain circulation time (approximately 20 seconds).

**Duration**
Approximately 5-10 minutes after single induction dose.

**Elimination**
Rapid redistribution of drug from the central nervous system (CNS) to lean body tissue accounts for the prompt awakening. The final elimination from the body depends on hepatic metabolism and excretion by the kidneys.

**Effects**

**CNS**
- Profound CNS depressant. Decreases cerebral metabolic rate and intracranial pressure. May cause hypotension, twitching and tremors during induction. May contribute to post-operative confusion and delirium. Potentiates the depressant effects of opioids, sedatives, alcohol and volatile anesthetics.

**CVS**
- Depression of myocardial contractility and vasodilation leads to decreased cardiac output and blood pressure with a mild compensatory tachycardia. Must be used with caution in patients with poor left ventricular function or critical coronary artery insufficiency or in those who are seriously ill or debilitated.

**Respiratory**
- Depresses the rate and depth of breathing leading to brief period of apnea. Does not blunt the airway’s response to manipulation therefore coughing, hiccupping, laryngospasm and bronchospasm may be seen at light planes of anesthesia.

**GI**
- Nausea and vomiting
Misc.
Incompatible with drugs with acidic pH. For example, if given in the IV line with vecuronium (a NDMR no longer in use), precipitation would occur. Arterial or extravascular injection produces necrosis.

Contraindications
Porphyria
KETAMINE
Class
Phencyclidine derivative. Can be used as an induction agent (usually in hemodynamically-compromised patients) or for sedation during painful procedures.

Mechanism of action
Acts at numerous central nervous system receptor sites, including the N-methyl-D-aspartate (NMDA) receptor.

Dose
Induction of anesthesia: 2 mg/kg IV
Induction of anesthesia: 5 mg/kg IM

Onset
Within one arm-brain circulation time (approximately 20 seconds).

Duration
Approximately 10-15 minutes after single induction dose, with full orientation occurring after 15-30 minutes.

Elimination
Redistribution from central nervous system (CNS) to inactive tissue sites accounts for termination of unconsciousness. Ultimate clearance is via hepatic metabolism and renal excretion.

Effects
CNS:
Produces “dissociative anesthesia” with patient in a cataleptic state. Ketamine provides a state of unconsciousness and intense analgesia however the patient’s eyes may remain open and roving, and their limbs may move purposelessly. Cerebral metabolic rate and intracranial pressure are increased.

CVS
Ketamine increases sympathetic outflow from the CNS leading to increased heart rate, blood pressure and cardiac output. Because of this effect, ketamine plays an important role in the management of patients with hypovolemic shock or cardiac tamponade. However, ketamine does possess direct myocardial depressant effects which may lead to worsened hypotension in patients in a prolonged shock state.

Respiratory
Some degree of airway protection is maintained. The patient may cough or swallow. Airway secretions increase. Bronchodilatory effect is secondary to increased sympathetic tone. Apnea is rare as respiratory drive is maintained.

Misc.
Undesirable psychological reactions are common on emergence: vivid, unpleasant dreams, excitement, confusion, fear. They tend to occur in the first hour of emergence and abate within one to several hours. Pretreat-
ment with benzodiazepines may help minimize this effect.

**Contraindications**
Raised intracranial pressure, coronary ischemia, psychiatric disease, eye surgery.
ETOMIDATE

Class
Short-acting hypnotic; anesthetic induction agent. Useful in hemodynamically-compromised patients.

Mechanism of action
Potentiates the inhibitory GABA neurotransmitter resulting in depression of the reticular activating system.

Dose
Induction: 0.2-0.6 mg/kg IV

Onset
Within one arm-brain circulation time (approximately 20 seconds).

Duration
Approximately 5-10 minutes after single induction dose.

Elimination
Rapid redistribution from central nervous system (CNS) to lean body tissue accounts for brief duration of action. Ultimately metabolized by hepatic and plasma esterases to inactive products.

Effects
CNS
CNS depressant, potentiating the depressant effects of opioids, sedatives and volatile anesthetics. Decreases cerebral metabolic rate and intracranial pressure. The cerebroprotective effects of etomidate make it useful in the management of the head-injured patient. Can cause seizure-like activity.

CVS
Etomidate is notable for the lack of significant cardiovascular depression that it causes. Therefore it is commonly chosen to facilitate intubation in the trauma patient, patients with hypovolemic shock or other unstable patients.

Respiratory
Etomidate causes a brief period of apnea.

GI
Nausea and vomiting

Misc.
Etomidate suppresses corticosteroid synthesis in the adrenal cortex and can lead to primary adrenal suppression. For this reason, its use in patients with sepsis is controversial. Etomidate can result in trismus if administered too quickly.
Inhaled Agents

Drugs

1. Desflurane
2. Sevoflurane
3. Isoflurane
4. Nitrous oxide

**DESFLURANE**

**Class**
Volatile inhaled anesthetic. Used for maintenance of anesthesia.

**Mechanism of Action**
Uncertain

**Dose**
Titrated to effect; MAC (age 40) = 6.0%

**Onset**
Low solubility allows rapid uptake and equilibration. Onset of effect is hastened by using higher flows of carrier gases and by using higher concentrations of volatile agent.

**Duration**
Clinical recovery in less than 10 minutes (2-2.5 x faster washout than Isoflurane)

**Elimination**
Pulmonary (major); negligible hepatic (0.02%)

**Effects**
**CNS**
Desflurane produces an additive central nervous system (CNS) depressant effect along with other sedative/hypnotics and analgesics. Sympatho-excitation can occur with rapid increase in concentration of desflurane. Has the potential to increase intracranial pressure which can be mitigated with hyperventilation. May cause headache, agitation, dizziness.

**CVS**
Dose-related hypotension (vasodilation). Tachycardia and hypertension may be seen due to sympathetic nervous system activation.

**Respiratory**
Respiratory depression with a rapid, shallow respiratory pattern. Loss of intercostal muscle function creates a rocking boat appearance. Desflurane is irritating to the airways and can cause breath-holding, cough, laryngospasm or bronchospasm in susceptible individuals, especially if used as sole agent for induction.

**GI**

**MSK**
Potentiates neuromuscular blockade; malignant hyperthermia trigger.

**Misc.**
Significant carbon monoxide production occurs on exposure to dessicated CO₂ absorbing agents therefore must not be used with low-flow anesthesia. Rapid elimination requires initiation of post-operative analgesia prior to emergence.

**Contraindications**
Malignant hyperthermia susceptibility
SEVOFLURANE

Class
Volatile inhaled anesthetic. Used for maintenance of anesthesia. Can be used for induction of anesthesia particularly in children. Rarely may be used as a treatment for status asthmaticus.

Mechanism of Action
Uncertain

Dose
Titrated to effect; MAC (age 40) = 2.1%.

Onset
Low solubility allows rapid uptake and equilibration. Onset of effect is hastened by using higher flows of carrier gases and by using higher concentrations of volatile agent.

Duration
Clinical recovery in less than 10 minutes (usually). If given for prolonged periods, wake-up will be slower as adipose stores have been saturated and are slow to off-load.

Elimination
Pulmonary (major); hepatic (2-5%); renal (metabolites excretion only)

Effects
CNS
Sevoflurane produces an additive central nervous system (CNS)-depressant effect along with other sedative/hypnotics and analgesics. Has the potential to increase intracranial pressure which can be mitigated with hyperventilation. Delirium.

CVS
Dose-related hypotension (vasodilation).

Respiratory
Respiratory depression with a rapid, shallow respiratory pattern. Loss of intercostal muscle function creates a rocking boat appearance. Causes bronchodilation. Sevoflurane is sweet-smelling and not as irritating to the respiratory tract as desflurane.

GI
Nausea, vomiting.

MSK
Potentiates neuromuscular blockade. Malignant hyperthermia trigger.

Misc.
Potential nephrotoxicity due to Compound A which is produced through contact with soda lime. Compound A can be produced if sevoflurane is used with very low fresh gas flows or for long MAC-hours. Therefore, sevoflurane must be used with a minimum of 2 litres/minute of fresh gas flow.

Contraindications
Malignant hyperthermia susceptibility
ISOFLURANE

Class
Volatile inhaled agent. Used for maintenance of anesthesia.

Mechanism of Action
Uncertain

Dose
Titrated to effect; MAC (age 40)=1.15

Onset
Higher solubility than sevoflurane and desflurane therefore uptake is slower than the modern agents. Onset of effect is hastened by using higher flows of carrier gases and by using higher concentrations of volatile agent.

Duration
Clinical recovery in less than 15 minutes (usually). Theoretically a slower wake-up than the modern agents due to higher solubility.

Elimination
Pulmonary

Effects
CNS
Isoflurane produces an additive central nervous system (CNS)-depressant effect along with other sedative/hypnotics and analgesics. Has the potential to increase intracranial pressure which can be mitigated with hyperventilation. Delirium.

CVS
Dose-related hypotension (vasodilation).

Respiratory
Respiratory depression with rapid, shallow respiratory pattern. Loss of intercostal muscle function creates a rocking boat appearance. Isoflurane is irritating to the airways and can cause breath-holding, cough, laryngospasm or bronchospasm. Its pungent quality makes it unsuitable for use with a mask induction.

GI
Nausea, vomiting.

MSK
Potentiates neuromuscular blockade. Malignant hyperthermia trigger.

Contraindications
Malignant hyperthermia susceptibility
NITROUS OXIDE

Class
Nitrous oxide is an inhaled agent but not a volatile agent. It is used as an adjunct to general anesthesia. It has a weak effect and therefore cannot be used as the sole agent for general anesthesia and is most commonly used in combination with a volatile agent. It can be used on its own for sedation or analgesia as can be seen in the obstetric or dental setting.

Mechanism of Action
Uncertain

Dose
Delivered in concentrations of up to 70% in oxygen. Actual MAC is 104%.

Onset
Immediate due to very low solubility.

Duration
Offset of effect is rapid after discontinuation.

Elimination
Pulmonary

Effects
CNS
N₂O is a potent analgesic. It increases cerebral metabolic rate, cerebral blood flow and intracranial pressure and is therefore not a good choice for patients with decreased intracranial compliance.

CVS
N₂O has a mild sympathomimetic effect but causes direct myocardial depression. The net effect is a modest decrease in blood pressure and heart rate. Increased coronary tone may exacerbate ischemia in susceptible patients.

Respiratory
N₂O produces mild respiratory depression which is potentiated by opioids, hypnotics and volatile anesthetics. It has no bronchodilatory effect. It exacerbates pulmonary hypertension.

Misc.
N₂O expands the volume of gas-containing spaces as N₂O diffuses across membranes more readily than nitrogen can diffuse out. Thus the size of a pneumothorax, emphysematous bleb or distended bowel loop will increase when N₂O is used. Bone marrow suppression due to inhibition of methionine synthetase, can occur if N₂O is used for extended periods. N₂O enhances opioid-induced rigidity. Finally, N₂O is an operating room pollutant; N₂O levels (in parts per million) in the operating room environment are measured regularly to comply with workplace safety regulations.

Contraindications
Raised intracranial pressure, pneumothorax or bowel obstruction. Should be used with caution in patients with coronary disease or emphysema.
Anxiolytics

Drugs

1. Midazolam

**MIDAZOLAM**

**Class**
Short-acting benzodiazepine. Used for sedation or as an adjunct during general anesthesia. Midazolam has anxiolytic and sedative (but not analgesic) properties.

**Mechanism of Action**
Agonism at the inhibitory GABA receptor.

**Dose**
For sedation: 0.03-0.08 mg/kg IV

Can also be given intramuscularly, intranasally and orally.

**Onset**
Within 3-5 minutes

**Duration**
Elimination half-time is 1-4 hours, making midazolam a much shorter acting agent than diazepam.

**Elimination**
Metabolized in the liver by microsomal enzymes and excreted in the urine.

**Effects**

**CNS**

**CVS**
In larger doses, in the presence of hypovolemia or when used in combination with opioids, midazolam can lead to decreased blood pressure and increased heart rate. Cardiac output is unchanged.

**Respiratory**
Dose-related respiratory depression occurs. This response is exaggerated in the elderly, in those with COPD or when used in combination with opioids.

**Misc.**
Midazolam is water-soluble therefore the pain on injection and phlebitis that are seen with diazepam are uncommon.
Antiemetics

Drugs

1. Ondansetron
2. Dimenhydrinate
3. Prochlorperazine

**ONDANSETRON**

**Class**
Serotonin (5-HT3) antagonist. Clinical use is as an antiemetic for post-operative nausea and vomiting or for patients receiving chemotherapy.

**Mechanism of Action**
Ondansetron is a highly selective competitive antagonist of the serotonin receptor. It is believed to have its effect centrally, possibly in the area postrema of the brainstem where the chemoreceptive trigger zone is located.

**Dose**
- **Prophylaxis (adults):** 4 mg IV prior to emergence.
- **Prophylaxis (children):** 50-150 μg/kg IV
- **Treatment (adults):** 1-2 mg IV

**Onset**
Less than 30 minutes

**Duration**
9 hours

**Elimination**
Hepatic (95%)

**Effects**
- **CNS**
  - Headache
- **CVS**
  - May cause cardiac rhythm or ECG changes by prolongation of the QT interval.
- **GI**
  - Constipation, elevation of liver enzymes.
- **Misc.**
  - Elimination of ondansetron is prolonged when given with other drugs metabolized by cytochrome P450 system.
**DIMENHYDRINATE**

**Class**
Antihistamine, antiemetic. In anesthetic practice, used as a second or third line treatment of post-operative nausea and vomiting (PONV). No role in prevention of PONV.

**Mechanism of Action**
Dimenhydrinate is a competitive antagonist at the histamine H1 receptor. The antiemetic effects is related to central anticholinergic actions as well as histamine antagonism in the vestibular system in the brain.

**Dose**
50-100 mg IV q4-6h, max. 400 mg/day (adults)
1.25 mg/kg IV q6h (children)

**Onset**
5 minutes after IV administration

**Duration**
4-6 hours

**Elimination**
Hepatic

**Effects**
**CNS**
Sedation (which is additive with alcohol and sedative hypnotics), dizziness, restlessness.

**Misc.**
May cause dry mouth, blurred vision, difficult urination; more rarely causes acute glaucoma or worsening of asthma. These side effects reflect its anticholinergic activity which is additive with other anticholinergics and monoamine oxidase inhibitors (MAOI).
**PROCHLORPERAZINE**

**Class**
Although it has several uses, in anesthetic practice it is used as an antiemetic for post-operative nausea and vomiting (PONV).

**Mechanism of Action**
Central inhibition of the dopamine D2 receptors in the medullary chemoreceptor trigger zone. Prochlorperazine also inhibits the vagus nerve in the gastrointestinal tract. The anticholinergic, sedative and antihistaminic effects of prochlorperazine also contribute to its antiemetic action.

**Dose**
2.5-10 mg IV, max. 40 mg/day (adults)

**Onset**
10-20 minutes

**Duration**
3-4 hours

**Elimination**
Enterohepatic

**Effects**
Prochlorperazine has anticholinergic properties which are additive to the anticholinergic effects of other drugs. As a phenothiazine, it also has the potential to cause extrapyramidal symptoms.

**CNS**
Sedative effects which are additive to other-hypnotics. May cause extra-pyramidal syndromes (motor restlessness, oculogyric crisis, opisthotonus, dystonias), especially in young male patients.

**CVS**
Hypotension caused by \(\alpha\)-adrenergic blocking effect. May potentiate hypotensive effect of vasodilators and diuretics. Causes QT interval prolongation.

**Misc.**
Diminishes effects of anticoagulants. Possible hyperthermia in the presence of hypothalamic dysfunction. Neuroleptic malignant syndrome.
Vasoactive Agents

Drugs

1. Phenylephrine
2. Ephedrine sulfate
3. Epinephrine

Ephedra distachya. Public domain image by Prof. Dr. Otto Wilhelm Thomé Flora von Deutschland, retrieved from Wikimedia Commons.
PHENYLEPHRINE

Class
Sympathomimetic; vasopressor. Used in the treatment of hypotension.

Mechanism of Action
Direct agonist at the $\alpha$-adrenergic receptor.

Dose
Bolus dose: 50-100 $\mu$g IV (adults)
Infusion: 0.1-1.0 $\mu$g/kg/minute

Onset
<1 minute

Duration
<5 minutes

Elimination
Re-uptake by tissue, liver and gut (monoamine oxidase)

Effects
CVS
Main effect is peripheral vasoconstriction, causing an increase in blood pressure. It is most appropriately used to raise the blood pressure in patients who are peripherally vasodilated (as a result of anesthesia, for example). It has the potential to cause myocardial ischemia, and left and right ventricular failure. It routinely causes a reflex bradycardia. If used in a patient in cardiogenic or hypovolemic shock, it may lead to a further reduction in vital organ blood flow.

Misc.
The clinician may observe diminished response of phenylephrine in patients receiving $\alpha$-adrenergic blockers or drugs with $\alpha$-blocking action such as phenothiazines. On the other hand, there may be augmented response when given with other vasopressors such as vasopressin and ergonovine. Phenylephrine has prolonged action in patients using monoamine oxidase inhibitors.
EPHEDRINE SULFATE

Class
Sympathomimetic (indirect-acting); vasopressor. Used in the treatment of hypotension.

Mechanism of Action
Ephedrine causes more norepinephrine to be released from the storage vesicles in the terminal of neurons thus increasing the amount of norepinephrine in the synaptic space. Ephedrine is (mostly) an “indirect-acting” catecholamine because it doesn’t act at the post-synaptic norepinephrine receptors.

Dose
5-20 mg IV (adults)
25-50 mg IM (adults)

Onset
IV: immediate
IM: minutes

Duration
IV: 10-minutes
IM: 60 minutes

Elimination
Hepatic, renal

Effects
CNS
Increases MAC of volatile anesthetics

Respiratory
Bronchodilator

CVS
Increases heart rate, contractility and therefore cardiac output (through its β adrenergic effect). Overall effect is to increase systemic vascular resistance through its α-adrenergic effect. May cause arrhythmias especially when used with volatile anesthetics. As the mechanism of action involves the release of intracellular catecholamines, there is an unpredictable effect in patients with depleted endogenous catecholamines.

Misc.
Excessive catecholamine effects may lead to hypertension, tachycardia, arrhythmias, pulmonary edema, anxiety, tremors, hyperglycemia and transient hyperkalemia followed by hypokalemia. Skin necrosis may occur at site of injection.

Contraindications
Ephedrine should not be used in patients on monoamine oxidase inhibitors (MAOIs) or those using cocaine. In these patients, phenylephrine is a safer choice for raising blood pressure. Ephedrine should be used with caution in patients who take SSRIs (serotonin-norepinephrine re-uptake inhibitors), as it may increase the risk of “serotonin syndrome”.

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**EPINEPHRINE**

**Class**
Sympathomimetic. Epinephrine has many uses:

1) Inotropic support in the patient in cardiogenic shock.
2) Bronchodilation in status asthmaticus.
3) Treatment of allergic reactions.
4) Treatment of croup.
5) Resuscitation in cardiovascular collapse of any cause.
6) Prolongation of action of anesthetic solutions.

**Mechanism of Action**
Epinephrine is a direct-acting sympathomimetic. It stimulates α- and β-adrenergic receptors resulting in a wide range of effects attributable to the sympathetic nervous system.

**Dose**
Cardiac arrest (adults): 0.5-1.0 mg IV q5min prn
Inotropic support (adults): 0.1-1.0 µg/kg/min
Anaphylaxis/severe asthma:
Adults: 0.1-0.5 mg SC/IM or 1-5 µg/kg IV prn
Children: 0.01 mg/kg SC/IM, maximum 0.5 mg

**Onset**
IV: immediate
SC/IM: 6-15 minutes

**Duration**
IV: 5-10 minutes
SC: 1-3 hours

**Elimination**
Enzymatic degradation

**Effects**

**CNS**
Anxiety, headache, stroke

**CVS**
Hypertension, tachycardia, arrhythmias, angina, pulmonary edema. Increased risk of ventricular arrhythmias is seen when used with volatile anesthetics.

**Respiratory**
Bronchodilation

**Misc.**
Skin necrosis at site of injection, hyperglycemia, transient hyperkalemia, followed by hypokalemia.
Local Anesthetics

Drugs

1. Bupivacaine
2. Lidocaine

BUPIVACAINE

Class
Local anesthetic. Used in infiltration anesthesia, spinal and epidural anesthesia and other regional anesthesia techniques.

Mechanism of Action
Sodium channel blocker

Dose
Maximum 2mg/kg without epinephrine
Maximum 3 mg/kg with epinephrine
Safe dose depends on where and how it is being administered. For example, absorption from intercostal administration is greater than for administration in adipose tissue.

Onset
Infiltration: 2-10 minutes
Epidural: 10-30 minutes
Spinal: <5 minutes

Duration
Infiltration: 2-5 hours
Epidural and spinal: up to 3.5 hours

Elimination
hepatic, pulmonary

Effects
Local anesthetics should not have systemic effects if used appropriately. If high plasma levels are achieved due to incorrect dosing or inadvertent intravascular injection then the symptoms manifest firstly in the central nervous system and then in the cardiovascular system where hypotension, heart block and other arrhythmias may occur. Premonitory signs and symptoms are perioral numbness, metallic taste, tinnitus, restlessness, dizziness and tremors. Seizures, respiratory and circulatory depression / arrest may occur. The treatment is supportive care and the use of Intralipid. Administration of benzodiazepines will increase the seizure threshold.

High intravascular concentrations of local anesthetics may potentiate the effects of muscle relaxants (both depolarizing and non-depolarizing).
LIDOCAINE

Class
Local anesthetic. Used in infiltration anesthesia and regional anesthesia (e.g. intravenous regional anesthesia). Lidocaine is still used for epidural anesthesia, especially for Caesarian section. Lidocaine is rarely used in spinal anesthesia due to associated nerve irritation. Lidocaine is occasionally used in the treatment of ventricular arrhythmias.

Mechanism of Action
Sodium channel blocker

Dose
Anesthetic:
Maximum 4 mg/kg without epinephrine
Maximum 7 mg/kg with epinephrine

Anti-arhythmic:
1 mg/kg IV bolus followed by 0.5 mg/kg q 2-5 minutes to maximum 3 mg/kg/hr

By infusion (of 0.1% solution): 1-4 mg/min (20-50 ug/kg/min)

Onset
IV: 45-90 seconds
Infiltration: 0.5-1 minute
Epidural: 5-15 minutes
Spinal: <1 minute

Duration
IV: 10-20 minutes
Infiltration: 0.5-1 hour
Epidural and spinal: 1-3 hours

Elimination
hepatic, pulmonary

Effects
Local anesthetics should not have systemic effects if used appropriately. If high plasma levels are achieved due to incorrect dosing or inadvertent intravascular injection then the symptoms manifest firstly in the central nervous system and then in the cardiovascular system where hypotension, heart block and other arrhythmias may occur. Premonitory signs and symptoms are perioral numbness, metallic taste, tinnitus, restlessness, dizziness and tremors. Seizures, respiratory and circulatory depression / arrest may occur. The treatment is supportive care and the use of Intralipid. Administration of benzodiazepines will increase the seizure threshold.

High intravascular concentrations of local anesthetics may potentiate the effects of muscle relaxants (both depolarizing and non-depolarizing).
SECTION 10

Miscellaneous

Drugs

1. Ketorolac tromethamine
2. Diphenhydramine
3. Dantrolene

KETOROLAC TROMETHAMINE

Class
Non-steroidal anti-inflammatory analgesic. Can be used orally or intravenously.

Mechanism of Action
COX-1 inhibitor

Dose
15-30 mg IV (adults)
0.5 mg/kg, max. 15 mg/dose (children > 2 years)

IV therapy should not exceed 2 days.

Onset
<30 minutes; peak effect in 1-2 hours.

Duration
4-6 hours

Elimination
Renal (92%); enterohepatic (6%)

Effects
CNS
Analgesic with an opioid-sparing effect.

GI
Gastrointestinal bleeding, ulcer, perforation, nausea, dyspepsia.

Renal
May precipitate renal failure in patients with renal insufficiency or those using ACE inhibitors.

Hematologic
May potentiate effects of anticoagulants. Rarely may cause hemorrhage due to platelet inhibition.
**DIPHENHYDRAMINE**

**Class**
Antihistamine, antiemetic. Used in the treatment of pruritis, allergic reactions and drug-induced extrapyramidal reactions.

**Mechanism of Action**
Diphenhydramine is a competitive inhibitor at the histamine H1 receptor. The antiemetic effects is related to central anticholinergic effect as well as histamine antagonism in the vestibular system in the brain.

**Dose**
Adults: 25-50 mg PO q6-8 hours; 10-50 mg IV/IM q 6 hours; maximum daily dose 400 mg.

**Onset**
IV: 5 minutes
PO: <15 minutes

**Duration**
4-6 hours

**Elimination**
Hepatic

**Effects**
**CNS**
Sedation (which is additive with alcohol and sedative hypnotics), dizziness, restlessness.

**CVS**
Rarely causes hypotension, arrhythmias

**Misc.**
May cause dry mouth, blurred vision, difficult urination; more rarely causes acute glaucoma or worsening of asthma, and GI or GU obstruction. These side effects reflect its anticholinergic activity, which is additive with other anticholinergics and monoamine oxidase inhibitors (MAOI).
DANTROLENE

Dantrolene is used in the treatment of malignant hyperthermia. It is the only specific and direct treatment of malignant hyperthermia. It is also used in the treatment of neuroleptic malignant syndrome.

It is a direct skeletal muscle relaxant which acts at the muscle cellular level, possibly at the ryanodine receptor. It is administered intravenously in 2.5 mg/kg doses until clinical signs show reversal of the hypermetabolic state (usual total dose <20 mg/kg).

Dantrolene can cause generalized muscle weakness in higher doses. Other common side effects include sedation, dizziness and constipation.

Dantrolene is supplied as a powder that must be mixed with sterile water. Its dissolution in water is very slow, difficult and time-consuming. Special “guns” have been devised to speed the preparation of dantrolene so as to minimize any delay in administration in the urgent situation.
**Question 1 of 16**
Nitrous oxide is contraindicated in ALL BUT ONE of the following settings. Indicate the exception:

- **A.** Total hip arthroplasty in the lateral position
- **B.** Laparotomy for bowel obstruction
- **C.** Multiple trauma with pneumothorax
- **D.** Patient with history of severe post operative nausea and vomiting
Acetylcholine (Ach) is a neurotransmitter. It acts at two different types of Ach receptors: nicotinic and muscarinic.

It is released from the nerve terminal of motor neurons into the synaptic cleft of the neuromuscular junction. Here it attaches to the nicotinic receptors of the post-synaptic membrane. Ach plays a key role in neuromuscular transmission.

Ach is also the neurotransmitter of the parasympathetic nervous system where it attaches to the muscarinic Ach receptors.

Related Glossary Terms
- Anticholinergic, Atropine, Autonomic nervous system, Cholinesterase, Competitive inhibitor, Glycopyrrolate, Muscarinic, Neuromuscular junction, Nicotinic, Non-depolarizing muscle relaxants, Residual block, Succinyllcholine, Vagus nerve

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- Chapter 6 - Muscle Relaxants
- Chapter 6 - Anticholinesterase and Anticholinergics
Addisonian crisis

Addisonian crisis comprises a constellation of symptoms, including severe hypotension and coma, that results from marked adrenal insufficiency.

In the peri-operative period, Addisonian crisis can occur in a patient who has chronic adrenal suppression due to (taking) exogenous systemic corticosteroids. Even if the patient’s “normal” steroid regime is continued, it may not be adequate to meet the increased demands created by the stress of surgery. In this case, Addisonian crisis may be seen. Prevention centres on providing adequate steroid coverage perioperatively. Duration of coverage depends on the magnitude of the surgery. A single pre-operative dose is sufficient for minor surgery, while 72 hours of coverage is required for major surgery.

Related Glossary Terms
Adrenal suppression, Pre-medication
Adjunct

An adjunct is something added, but not essential. In anesthesia, we have adjunct airway devices (like stylets and bougies). As well, many pharmacologic agents are used as adjuncts to analgesia. An example would be the use of a non-steroidal anti-inflammatory for a patient receiving patient-controlled (PCA) Morphine.

Related Glossary Terms
Analgesia, Difficult airway, Fibreoptic bronchoscope, Ketorolac Tromethamine, Patient controlled analgesia, Stylet
Adrenal suppression

If a patient is receiving exogenous systemic corticosteroids for more than a week, he or she will begin to experience suppression of the hypothalamic-pituitary-adrenal axis. When this endogenous pathway shuts down, the adrenal gland atrophies and takes at least 3 months to recover its function once suppression abates.

Until adrenal function is fully recovered, the patient may experience adrenal insufficiency when exposed to the stresses of illness and surgery. Clinical guidelines exist to estimate the need for steroid replacement in patients at risk for adrenal suppression.

Related Glossary Terms
Addisonian crisis, Etomidate, Pre-medication
Airway assessment

The purpose of the airway assessment is to identify potential difficulties with airway management and to determine the most appropriate approach. The airway is assessed by history, physical examination and, occasionally, laboratory exams. Searching for past records indicating ease of intubation is also an important part of airway assessment.

The key features on physical exam are mouth opening, thyromental distance, neck range-of-motion, and Mallampati score. It is also important to assess the predicted ease of bag-mask ventilation.

It is important to understand that airway examination is imperfect in both its sensitivity and specificity for predicting ease of intubation by direct laryngoscopy.

Related Glossary Terms
Bag mask ventilation, Difficult airway, Direct laryngoscopy, Intubation, Mallampati classification, Mouth opening, Neck motion, Pre-operative assessment
Airway obstruction

Causes of airway obstruction can be categorized broadly as follows:

a) Obstruction caused by normal tissue such as the tongue, tonsils, larynx and other soft tissue. Laryngospasm is an example of airway obstruction that occurs in the anesthesia setting.

b) Obstruction caused by pathology of normal tissue such as that which occurs from infection, inflammation, tumour or trauma. Examples of infectious processes that can lead to airway obstruction include croup, epiglottitis and sublingual abscesses.

c) Obstruction caused by foreign body inhalation.

Signs of airway obstruction in the spontaneously-breathing patient include stridor, a rocking-boat appearance to the chest and tracheal indrawing.

Unresolved, airway obstruction leads to hypoxia and hypercarbia.

Related Glossary Terms
Extubation, Hypoxemia, Laryngeal mask airway (LMA), Laryngospasm, Larynx, Patency, Subglottis, Upper airway

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Aldrete score

Phase 1 of recovery must be achieved before the patient is discharged to the ward or short-stay unit. The patient must demonstrate adequacy of: ventilation and airway control; circulation; colour; level of consciousness; and activity. The Aldrete score is a detailed scoring system that rates the patient’s status from 0-2 on each of those five criteria.

When Phase 1 recovery is complete, the patient must:

- be showing no signs of respiratory depression for at least 20-30 minutes after last dose of parenteral opioids.
- be easily aroused.
- be fully oriented to person, place and time.
- be able to maintain and protect the airway on his own with evidence of a strong cough.

It is also important that pain and post-operative nausea and vomiting be controlled prior to PACU discharge and that there are no ongoing surgical concerns such as surgical site bleeding.

Related Glossary Terms
Analgesia, Opioids, Patency, Post-anesthetic care unit, Post-operative nausea and vomiting (PONV), Recovery
Alfentanil

**Class**
Synthetic opioid analgesic (short-acting); adjunct to anesthesia.

**Mechanism of Action**
Acts at the mu-and kappa opioid receptors.

**Dose**
5-50 ug/kg IV, according to physical status, other agents used, nature and duration of surgery.

**Onset**
1-2 minutes

**Duration**
20 minutes

**Elimination**
Hepatic

**Effects**
- **CNS**
  Analgesia, sedation. All of the depressant effects of alfentanil are potentiated by concurrent use of sedatives, volatile anesthetics and nitrous oxide.
- **CVS**
  Bradycardia, hypotension. The synthetic opioids are not direct myocardial depressants but they do reduce sympathetic drive, which may result in decreased cardiac output in patients who are relying on sympathetic tone to support their circulation, such as those in hypovolemic or cardiogenic shock.
- **Respiratory**
  Potent respiratory depression which at the extreme, leads to apnea.
- **GI**
  Nausea, vomiting, biliary tract spasm.
- **Misc.**
  Muscle rigidity, pruritis.

**Related Glossary Terms**
Analgesia, Apneic threshold, Balanced anesthesia, Fentanyl, Four A's of anesthesia, Morphine Sulfate, Naloxone, Opioids, Remifentanil, Sufentanil
Analgesia

Pain relief

Related Glossary Terms
Adjunct, Aldrete score, Alfentanil, Balanced anesthesia, Epidural analgesia, Fentanyl, Four A's of anesthesia, Ketorolac Tromethamine, Morphine Sulfate, Nitrous oxide (N2O), Opioids, Patient controlled analgesia, Post-anesthetic care unit, Recovery, Regional anesthesia, Remifentanil, Sufentanil

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Chapter 6 - Induction Agents
Chapter 6 - Inhaled Agents
Anesthetic circuit

The anesthetic circuit is the tubing that delivers the fresh gas (oxygen on its own or mixed with air or nitrous oxide), along with volatile gas from the anesthetic machine to the patient’s airway.

There are several types of anesthetic circuits in use. Some allow rebreathing (and must be used with a CO2 absorber) and some do not allow rebreathing. Circuits that are designed for rebreathing allow for more economical use of volatile anesthetic gases.

The most common type in use is a circle circuit (rebreathing).

Related Glossary Terms
Bag mask ventilation, Capnograph, Circle circuit, CO2 (carbon dioxide) absorber, Flowmeter, Volatile
**Antecubital fossa**

The antecubital fossa is located in the anterior (inner) surface of the arm, at the level of the elbow. It is a common space for finding peripheral intravenous access.

**Related Glossary Terms**

Drag related terms here

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Chapter 1 - Fluid Management
Anticholinergic

Anticholinergic drugs include atropine and glycopyrrolate. Anticholinergic agents act as acetylcholine receptor blockers at the muscarinic (not nicotinic) acetylcholine receptors. They have an anti-parasympathetic effect.

In anesthesia practice, anticholinergic agents are most commonly used as an accompaniment to anticholinesterase (reversal) agents. Without the addition of an anticholinergic, the reversal of NDMR might result in excessive muscarinic effect or at the extreme, a cholinergic crisis.

Anticholinergic agents are also used as antisialagogues for awake intubation.

Finally, anticholinergic agents play an important role in the treatment of clinically important bradycardias.

Related Glossary Terms

Acetylcholine, Anticholinesterase, Antisialagogue, Atropine, Autonomic nervous system, Glycopyrrolate, Muscarinic, Neostigmine, Neuromuscular junction, Nicotinic, Vagus nerve

Index
Anticholinesterase

Anticholinesterases act in the synaptic cleft of the neuromuscular junction to reverse the effect of non-depolarizing muscle relaxants (NDMR). They inhibit the action of cholinesterase thereby decreasing the rate of breakdown of acetylcholine (Ach). The increased concentration of Ach in turn displaces the NDMR from the Ach receptors and thus restores normal neuromuscular transmission.

Because anticholinesterases exert their effect at both nicotinic and muscarinic Ach receptors, their administration must be accompanied by an anticholinergic (such as atropine or glycopyrrolate) in order to avoid muscarinic effects including bradycardia, bronchospasm and excessive salivation.

Neostigmine is a commonly-used anticholinesterase.

Related Glossary Terms
Acetylcholine, Anticholinergic, Atropine, Autonomic nervous system, Cholinesterase, Extubation, Glycopyrrolate, Muscarinic, Myasthenia gravis, Neostigmine, Neuromuscular junction, Nicotinic, Non-depolarizing muscle relaxants, Peripheral nerve stimulator, Residual block, Vagus nerve

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Antiemetic agents

Antiemetic agents are those that are used to prevent or treat nausea and vomiting.

Related Glossary Terms
Chemoreceptor trigger zone, Dimenhydrinate, Ondansetron, Opioids, Post-operative nausea and vomiting (PONV), Prochlorperazine
Antisialagogue

An antisialagogue is a drug which reduces airway (including oral) secretions. An antisialagogue is often administered prior to awake intubation.

Salivation is an effect of the vagus nerve of the parasympathetic nervous system. Accordingly, the most commonly-used antisialagogues are anticholinergic agents.

Related Glossary Terms
Anticholinergic, Atropine, Autonomic nervous system, Difficult airway, Fibreoptic bronchoscope, Glycopyrrolate, Muscarinic, Vagus nerve
Aortocaval compression

In the third trimester, a pregnant woman is at risk of aortocaval compression if she lies flat on her back. In this position, the gravid uterus can compress the vena cava (compromising venous return) and/or the aorta. The result can be maternal hypotension and reduced uterine perfusion. Aortocaval compression can be prevented by having the parturient remain in the lateral position, or slightly tilted to the left.

Related Glossary Terms
Parturient
Apneic threshold

Apneic threshold is the partial pressure of carbon dioxide below which a person ceases spontaneous rhythmic breathing due to loss of central drive to breathe.

In the healthy unanesthetized patient, the apneic threshold for CO2 is 32 mmHg.

Opioids and volatile anesthetic agents raise the apneic threshold, sometimes quite considerably.

Related Glossary Terms
Alfentanil, Cough reflex, Emergence, Fentanyl, Morphine Sulfate, Naloxone, Opioids, Re-narcotization, Remifentanil, Sufentanil, Volatile
ASA class

The ASA (American Society of Anesthesiologists) classification was defined in the 1940’s as an attempt to identify operative risk. It is not an accurate predictor of peri-operative risk in current practice nor does it lend itself to inter-rater reliability. It is, however, an accepted method of indicating the overall physical condition of the patient.

<table>
<thead>
<tr>
<th>ASA CLASS</th>
<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>A normal healthy patient in need of surgery for a localized condition.</td>
</tr>
<tr>
<td>2</td>
<td>A patient with mild to moderate systemic disease; examples include controlled hypertension, mild asthma.</td>
</tr>
<tr>
<td>3</td>
<td>A patient with severe systemic disease; examples include complicated diabetes, uncontrolled hypertension, stable angina.</td>
</tr>
<tr>
<td>4</td>
<td>A patient with life-threatening systemic disease; examples include renal failure or unstable angina.</td>
</tr>
<tr>
<td>5</td>
<td>A moribund patient who is not expected to survive 24 hours with or without the operation; examples include a patient with a ruptured abdominal aortic aneurysm in profound hypovolemic shock.</td>
</tr>
</tbody>
</table>

Related Glossary Terms
Pre-operative assessment

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Chapter 2 - Pre-operative Evaluation
Asepsis is another word for sterile technique. Invasive procedures must be performed meticulously in order to minimize the risk of infection.

Sterile technique involves prepping the patient’s skin with an alcohol-based disinfectant. Once prepped, the site is isolated with sterile drapes. The operator washes his or her hands and uses sterile gown and gloves. Sterility is maintained through the careful handling of equipment and patient.

Related Glossary Terms
Regional anesthesia
Aspiration

Aspiration is the inhalation of stomach contents. Aspiration can cause pneumonia. At its most serious, aspiration can lead to acute respiratory distress syndrome (ARDS) and non-cardiogenic pulmonary edema. Patients at risk of aspiration during anesthesia include:

- Patients who are improperly fasted.
- Patients with compromised gastroesophageal sphincter (history of gastroesophageal reflux disease, pregnancy, diabetes mellitus).
- Patients with decreased gastric emptying (diabetes mellitus, pregnancy, patients receiving opioids, trauma patients, patients with bowel obstruction, patients receiving medications that slow motility).

Related Glossary Terms

Clear fluids, Cough reflex, Cricoid cartilage, Epiglottis, Intubation, Laryngeal mask airway (LMA), Laryngospasm, NPO, Pre-medication, Rapid sequence induction, Right mainstem bronchus, Sellick’s maneuver, Sodium citrate, Subglottis

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Chapter 5 - Pediatric Anesthesia
**Atracurium**

**Class**
Nondepolarizing skeletal muscle relaxant (NDMR); short-acting

**Mechanism of Action**
Competitive inhibitor at the acetylcholine receptors of the post-synaptic cleft of the neuromuscular junction.

**Dose**
Intubation: 0.5-0.6 mg/kg IV
Maintenance bolus: 0.1-0.3 mg/kg IV

**Onset**
3-4 minutes

**Duration**
20-35 minutes

**Elimination**
Hoffman elimination, ester hydrolysis

**Effects**

**MSK**
The neuromuscular blockade effects of non-depolarizing muscle relaxants are potentiated by succinylcholine, volatile anesthetics, aminoglycosides, lithium, loop diuretics, lidocaine, magnesium, lithium, ganglionic blockers, hypothermia, hypokalemia and respiratory acidosis.

Enhanced neuromuscular blockade is seen in patients with myasthenia gravis or myopathies.

The effects of NDMR are antagonized by cholinesterase inhibitors. Increased resistance to NDMR is seen in patients on theophylline, burn patients and those with paresis or paralysis.

**Misc.**
Histamine release may occur with rapid administration or higher dosages. Produces an excitatory metabolite called laudanosine. Muscle relaxants are the most common cause of anaphylactoid reactions under general anesthesia.

**Related Glossary Terms**
Anticholinesterase, Cis-Atracurium, Laudanosine, Neuromuscular junction, Non-depolarizing muscle relaxants, Pancuronium, Peripheral nerve stimulator, Residual block, Rocuronium, Urticaria

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Atropine

Class
Anticholinergic. Clinical use in anesthesia includes the treatment of bryadycardia and asystole; as an antisialagogue for awake intubation; or for countering the muscarinic effects of the anticholinesterases used for the reversal of neuromuscular blockade.

Mechanism of Action
An acetylcholine receptor blocker active at the muscarinic (not nicotinic) acetylcholine receptors. Therefore, atropine has an anti-parasympathetic effect.

Dose
Premedication 0.4-0.6 mg IV/IM in adults, 10-20 ug/kg IV/IM in children. Reversal 0.015 mg/kg IV with neostigmine 0.05 mg/kg IV.

Onset
Immediate

Duration
1-2 hours

Elimination
Hepatic, renal

Effects
Most effects result from the anticholinergic action of atropine.

CNS
Confusion, hallucinations, mydriasis, blurred vision, increased intraocular pressure

CVS
Tachycardia (high doses), bradycardia (low doses)

GI
Gastroesophageal reflux

GU
Urinary hesitancy, retention

Misc.
Has additive anticholinergic effects with antihistamines, phenothiazines, tricyclic antidepressants, mono-amine oxidase inhibitors and benzodiazepines. Potentiates sympathomimetics. May produce central anticholinergic syndrome.

Contraindications
Contraindicated in patients with narrow-angle glaucoma, gastrointestinal or genitourinary obstruction.

Related Glossary Terms
Acetylcholine, Anticholinergic, Anticholinesterase, Antisialagogue, Autonomic nervous system, Glycopyrrolate, Muscarinic, Vagus nerve
Autonomic nervous system

The autonomic nervous system is comprised of the sympathetic and parasympathetic systems.

The sympathetic system is mediated neurally through the spinal cord, exiting through thoracolumbar nerves, as well as humorally, through the adrenal gland.

The parasympathetic system is mediated through the vagus nerve and various other cranial nerves.

Related Glossary Terms
Acetylcholine, Anticholinergic, Anticholinesterase, Antisialagogue, Atropine, Cholinesterase, Ephedrine Sulfate, Epinephrine, Four A's of anesthesia, Glycopyrrolate, Shock, Sympathetic nervous system, Vagus nerve

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Chapter 3 - General Anesthesia
Awareness

Awareness is a complication of anesthesia whereby a patient who has received a general anesthetic becomes conscious of his or her environment during the surgical procedure.

Some patients with awareness have “recall” of the awareness episode while others, even most, do not. Awareness episodes are not necessarily associated with pain.

Some procedures such as Caesarian section or coronary bypass surgery are associated with a higher incidence of awareness.

Related Glossary Terms

Balanced anesthesia, Four A's of anesthesia, Non-depolarizing muscle relaxants, Reticular activating system
Bacterial endocarditis

The 2007 American Heart Association (AHA) Guidelines for the prevention of bacterial endocarditis recommend antibiotic prophylaxis only for patients with *certain* cardiac conditions undergoing *certain* surgical procedures.

The relevant procedures are listed below:

1) dental procedures involving manipulation of the gums or the roots of the teeth.
2) procedures of the respiratory tract.
3) procedures involving infected tissues.

Interestingly, procedures of the genitourinary or gastrointestinal tract do not indicate prophylaxis, assuming uninfected tissue.

The relevant cardiac conditions are listed below:

1) patients with artificial heart valves.
2) patients who have had heart repairs using prosthetic material. This does not refer to coronary stents.
3) patients with a prior history of endocarditis.
4) patients with certain un-repaired or incompletely repaired congenital heart disease.
5) patients who have transplanted hearts who now have developed a heart valve problem.

**Prophylaxis is required when a patient with one of the above conditions is undergoing one of the above-described surgical procedures.**

Updates to these recommendations are readily available online.

**Related Glossary Terms**

Pre-medication

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**Index**

Chapter 2 - Pre-operative Evaluation
Bag mask ventilation

Bag mask ventilation involves providing positive pressure ventilation to the patient through the application of a mask over the face and by manually squeezing the bag that is connected to the anesthetic circuit.

It is performed during induction of anesthesia, after the patient becomes apneic and before a definitive airway is achieved through intubation or the insertion of an LMA.

Aspiration and airway obstruction are risks during bag mask ventilation.

Related Glossary Terms
Airway assessment, Airway obstruction, Anesthetic circuit, Aspiration, Esophageal intubation, Induction, Intubation, Laryngeal mask airway (LMA), Mask Airway
Balanced anesthesia

Balanced anesthesia is the practice of using combinations of agents, each for a specific purpose. An example of a balanced technique would be the use of propofol for induction of anesthesia followed by the administration of desflurane and nitrous oxide for maintenance of unconsciousness, sufentanil for analgesia and rocuronium for muscle relaxation.

Prior to the development of the range of agents described above, ether or other volatile agents were used to achieve all the goals of anesthesia in a non-specific fashion.

Related Glossary Terms

Analgesia, Awareness, Desflurane, Four A’s of anesthesia, Induction, Inhaled agents, Isoflurane, Maintenance, Non-depolarizing muscle relaxants, Opioids, Sevoflurane, Total intravenous anesthesia
Baricity

Baricity, or heaviness, determines how a local anesthetic will distribute itself in the intrathecal space. It is expressed as a ratio of the specific gravity of the local anesthetic to that of the cerebral spinal fluid (CSF).

Related Glossary Terms
Intrathecal space, Spinal anesthesia, Subarachnoid space
Bier Block

An intravenous regional anesthetic (IVRA or “Bier block”) is a technique introduced by August Bier in 1908 and still in use today. The operative limb (arm or leg) is “exsanguinated” using gravity and an elastic bandage after which an arterial cuff is inflated proximally to maintain exsanguination. Subsequently to exsanguination, large volumes of dilute lidocaine are injected (through a small cannula that is placed prior to exsanguination) into the affected limb.

Surgical anesthesia can last for 60-90 minutes, usually limited by the pain created by the inflated cuff. It is important that the cuff pressure be maintained for at least 30 minutes to avoid a sudden bolus of local anesthetic being introduced into the systemic circulation.

Related Glossary Terms

Brachial plexus block, Intravenous regional anesthetic, Lidocaine, Local anesthetic toxicity, Regional anesthesia

Index

Chapter 3 - Regional Anesthesia
Blood products

Blood products are transfused in order to correct a deficit (in quantity or quality) of a specific blood component. Donated blood is separated into its component elements to facilitate this individualized correction.

Examples of blood products include:

Pack red blood cells (PRBC); platelets; fresh frozen plasma (FFP); cryoprecipitate; and albumin.

Related Glossary Terms

Colloids, Euvolemia, Hypothermia, Shock, Thrombocytopenia
Brachial plexus

The brachial plexus is formed from the anterior primary rami of the C5-T1 nerve roots and supplies all of the motor function and most of the sensory function of the upper extremity.

Related Glossary Terms
Brachial plexus block, Intravenous regional anesthetic

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Chapter 3 - Regional Anesthesia
Chapter 3 - General Anesthesia
Chapter 4 - Post-operative Pain Management
**Brachial plexus block**

The brachial plexus block is a technique used to provide surgical anesthesia for the upper limb. The brachial plexus is formed from the anterior primary rami of the C5-T1 nerve roots and supplies all of the motor function, and most of the sensory function of the upper extremity.

It can be accessed with a needle (and therefore blocked) at several locations: axillary, supraclavicular, interscalene. The choice of approach is determined by which nerves are most important to block (based on surgical site) and which complications are most important to avoid (based on patient co-morbidities). For example, the axillary approach will provide unreliable blockade of the more proximal arm. The supraclavicular approach and interscalene approaches carry the risk of pneumothorax and phrenic nerve block.

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**Related Glossary Terms**

Bier Block, Brachial plexus, Bupivacaine, Lidocaine, Local anesthetic toxicity, Phrenic nerve, Pneumothorax, Regional anesthesia
Bupivacaine

**Class**
Local anesthetic. Used in infiltration anesthesia, spinal and epidural anesthesia and other regional anesthesia techniques.

**Mechanism of Action**
Sodium channel blocker

**Dose**
Maximum 2mg/kg without epinephrine
Maximum 3 mg/kg with epinephrine

Safe dose depends on where and how it is being administered. For example, absorption from intercostal administration is greater than for administration in adipose tissue.

**Onset**
Infiltration: 2-10 minutes
Epidural: 10-30 minutes
Spinal: <5 minutes

**Duration**
Infiltration: 2-5 hours
Epidural and spinal: up to 3.5 hours

**Elimination**
hepatic, pulmonary

**Effects**
Local anesthetics should not have systemic effects if used appropriately. If high plasma levels are achieved due to incorrect dosing or inadvertent intravascular injection then the symptoms manifest firstly in the central nervous system and then in the cardiovascular system where hypotension, heart block and other arrhythmias may occur. Premonitory signs and symptoms are perioral numbness, metallic taste, tinnitus, restlessness, dizziness and tremors. Seizures, respiratory and circulatory depression / arrest may occur. The treatment is supportive care and the use of Intralipid. Administration of benzodiazepines will increase the seizure threshold.

High intravascular concentrations of local anesthetics may potentiate the effects of muscle relaxants (both depolarizing and non-depolarizing).

**Related Glossary Terms**
Brachial plexus block, Epidural analgesia, Epidural anesthesia, Intralipid, Lidocaine, Local anesthetic toxicity, Spinal anesthesia
Butyrophenones are a class of antipsychotic agents that includes haloperidol and droperidol.

Related Glossary Terms

Drag related terms here

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Chapter 3 - Anesthetic Techniques
Chapter 6 - Opioid agonists and antagonists
Capnograph

The capnograph is a monitor within the anesthetic machine which measures the quantity of carbon dioxide in the exhaled breath. It provides an analog waveform along with a numeric value. Various determinations can be made based on the characteristics of the waveform. For example, bronchospasm has a characteristic upslanting pattern on the capnograph that can be recognized by the astute clinician.

It is normal to have a gradient between alveolar and arterial levels of carbon dioxide, a gradient which is in the range of 4-8 cmH2O. There are conditions, however, where capnography will significantly underestimate the arterial blood levels of CO2:

1) Conditions where blood flow to the lungs is impeded, therefore carbon-dioxide is not getting delivered to the lungs. Examples include cardiac arrest and pulmonary embolus.

2) Conditions where a true alveolar gas sample is not being obtained (such that deadspace is sampled instead). Examples include the use of very small tidal volumes or any condition causing obstruction to expiration such as acute bronchospasm due to asthma exacerbation or anaphylactic reaction.

Related Glossary Terms

Anesthetic circuit, CO2 (carbon dioxide) absorber, Esophageal intubation, Intubation

Index

Chapter 1 - Airway Management
Chapter 2 - Anesthetic Equipment and Monitoring
Chemoreceptor trigger zone

The chemoreceptor trigger zone is located in the area postrema of the medulla. Its afferent input is systemic, where it can be triggered by certain drugs and hormones. Its efferent path is to the vomiting center, where it initiates vomiting.

The vomiting centre receives input from other sources, other than the chemoreceptor trigger zone, such as the gastrointestinal tract, the vestibular system and the higher cortex.

Related Glossary Terms
Antiemetic agents, Dimenhydrinate, Ondansetron, Post-operative nausea and vomiting (PONV), Prochlorperazine
Cholinesterase

Cholinesterase is a naturally-occurring enzyme in the synaptic cleft that breaks down acetylcholine (Ach). Cholinesterase reduces the activity of acetylcholine in the neuromuscular junction.

Anticholinesterase drugs inhibit cholinesterase thereby increasing the amount of acetylcholine in the neuromuscular junction. The Ach, then, is better able to compete with non-depolarizing muscle relaxant molecules for Ach receptors.

Related Glossary Terms
Acetylcholine, Anticholinesterase, Autonomic nervous system, Neostigmine, Neuromuscular junction, Nicotinic, Non-depolarizing muscle relaxants, Residual block

Chapter 3 - General Anesthesia
Chapter 6 - Anticholinesterase and Anticholinergics
Circle circuit

The circle circuit is a breathing circuit which allows rebreathing of expired gases. It allows a more economical use of gases and volatile anesthetic agents. A circle circuit requires the use of a CO2 absorber to prevent hypercarbia.

A circle system is shaped as a circle rather than a straight line. It has two one way valves (one inspiratory and one expiratory) to ensure proper direction of flow.

Related Glossary Terms
Anesthetic circuit, CO2 (carbon dioxide) absorber, Volatile
Cis-Atracurium

Class
Non-depolarizing skeletal muscle relaxant (NDMR); intermediate-acting

Mechanism of Action
Competitive inhibitor at the acetylcholine receptors of the post-synaptic cleft of the neuromuscular junction.

Dose
Intubation: 0.15-0.2 mg/kg
Maintenance bolus: 0.03 mg/kg
Maintenance infusion: 1-2 µg/kg/minute

Onset
Dose-dependent:
2 minutes (0.15 mg/kg)
1.5 minutes (0.2 mg/kg)

Duration
Dose-dependent:
55 minutes (0.15 mg/kg)
65 minutes (0.2 mg/kg)
20 minutes (maintenance bolus 0.03 mg/kg)

Elimination
Hoffman elimination (77%), renal (16%)

Effects

MSK
The neuromuscular blockade effects of non-depolarizing muscle relaxants are potentiated by succinylcholine, volatile anesthetics, aminoglycosides, lithium, loop diuretics, lidocaine, magnesium, lithium, ganglionic blockers, hypothermia, hypokalemia and respiratory acidosis.

Enhanced neuromuscular blockade is seen in patients with myasthenia gravis or myopathies.

The effects of NDMR are antagonized by cholinesterase inhibitors. Increased resistance to NDMR is seen in patients on theophylline, burn patients and those with paresis or paralysis.

Misc.
Histamine release may occur with rapid administration or higher dosages. Produces 5-10x less laudanosine metabolite than atracurium. Muscle relaxants are the most common cause of anaphylactoid reactions under general anesthesia.

Related Glossary Terms
Anticholinesterase, Atracurium, Laudanosine, Neuromuscular junction, Non-depolarizing muscle relaxants, Pancuronium, Peripheral nerve stimulator, Residual block, Rocuronium

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Chapter 3 - General Anesthesia
Chapter 6 - Drug Finder
Chapter 6 - Muscle Relaxants
Clear fluids

Clear fluids include water, apple juice, black coffee and tea.

The following liquids are NOT clear fluids: milk, orange juice, consomme (soup), coffee (or tea) with milk or cream added.

Related Glossary Terms
Aspiration, Cricoid cartilage, NPO, Pre-operative assessment, Rapid sequence induction
CO2 (carbon dioxide) absorber

The CO2 absorber is a component of the circle anesthetic circuit. Before the exhaled gas is recirculated back to the patient, it passes through the CO2 absorber where CO2 is scrubbed out. By removing the waste gas (CO2), the CO2 absorber allows “rebreathing” or recirculation of exhaled gases. Through rebreathing, air, oxygen, nitrous oxide and the volatile anesthetics can be used more economically.

The CO2 absorber contains a hydroxide called soda lime-Ca(OH)2.

The reaction takes place over three steps and can be described as follows:

\[
\text{CO}_2 + \text{H}_2\text{O} \rightarrow \text{aqueous CO}_2
\]

\[
\text{aqueous CO}_2+ \text{NaOH} \rightarrow \text{NaHCO}_3
\]

\[
\text{NaHCO}_3 + \text{Ca(OH)}_2 \rightarrow \text{CaCO}_3 + \text{H}_2\text{O} + \text{NaOH}
\]

NaOH is a catalyst and is recycled to step 2.

Each mole of CO2 (44g) reacted produces one mole of water (18g).

The overall reaction, which is a heat-producing one that requires the presence of water, is:

\[
\text{CO}_2 + \text{Ca(OH)}_2 \rightarrow \text{CaCO}_3 + \text{H}_2\text{O}
\]

The sodalime granules are manufactured with a chemical indicator such that as the soda lime becomes exhausted, it changes colour.

Related Glossary Terms
Anesthetic circuit, Capnograph, Circle circuit

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Chapter 2 - Anesthetic Equipment and Monitoring
Chapter 6 - Inhaled Agents
Colloids

Colloids are suspensions of protein or other complex organic particles. Examples of colloids are albumin (5%, 25%), hydroxyethyl starches (Pentaspan®, Voluven®), packed red blood cell concentrates (PRBC), platelets and fresh frozen plasma (FFP).

Because of their suspended proteins (or complex starches), colloids have an oncotic pressure similar to that of plasma and therefore replace blood loss in a 1:1 ratio.

Related Glossary Terms
Blood products, Crystalloid, Euvolemia, Maintenance fluid requirements, Pre-operative fluid deficit, Shock, Third space loss
Competitive inhibitor

Competitive inhibition occurs when the inhibitor (e.g. drug) binds to an active site (the receptor) thus preventing the binding of the physiologic molecule that would normally attach there to exert its effect. Non-depolarizing muscle relaxants act by competitive inhibition at the nicotinic acetylcholine receptor.

By contrast, non-competitive inhibition occurs when a drug acts by reducing the activity of an enzyme (for example) and exerts this effect whether the enzyme is attached to its substrate or not. The inhibitor, in this case, is not competing for receptor sites.

Related Glossary Terms
Acetylcholine, Neuromuscular junction, Nicotinic, Non-depolarizing muscle relaxants
Continuous Infusion

A “continuous infusion” is the intravenous delivery of a medication at a constant rate, usually requiring the use of a dedicated syringe pump.

When given by infusion, drugs reach a “steady state” in the patient’s bloodstream. The time required to reach that steady state is dependent on the half life of the drug and is usually equal to 5 half-lives. Steady state can be reached more quickly by delivering a bolus prior to commencing the continuous infusion.

Drugs can be delivered given by continuous infusion during the maintenance phase of general anesthesia when requirements are reasonably predictable. Drugs used for sedation can also be given by continuous infusion.

Propofol and remifentanil are examples of drugs that are commonly delivered by continuous infusion during anesthesia.

Related Glossary Terms

Propofol, Remifentanil, Total intravenous anesthesia
Cormack Lehane

Cormack Lehane is a grading system that describes the visualization of airway structures that is achieved on direct laryngoscopy.

Grade 1 - entire vocal folds visualized
Grade 2 - part of the vocal folds visualized
Grade 3 - epiglottis seen but none of the vocal fold structures seen
Grade 4 - none of epiglottis seen

Related Glossary Terms
Airway assessment, Difficult airway, Direct laryngoscopy, Epiglottis, Intubation, Laryngoscope, Macintosh blade, Magill blade, Mallampati classification

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Chapter 1 - Airway Management
Cough reflex

The cough reflex is an attempt to clear the upper or lower airway of foreign matter and is triggered by sensory input.

The cough reflex is an important means of protection against aspiration. The cough reflex is blunted by anything that decreases level of consciousness. Opioids, in particular, are very effective at attenuating the cough reflex and are a useful adjunct to topicalization of the airway during awake intubation.

It is important to understand that coughing is separate reflex than breathing. On emergence, patients may cough (due to the stimulation of the endotracheal tube) despite not (yet) having a drive to breath. Therefore, the anesthesiologist must resist the temptation to extubate the coughing patient until it has been established that the drive to breathe has returned.

Related Glossary Terms
Apneic threshold, Aspiration, Emergence, Fibreoptic bronchoscope, Lower airway, Opioids, Upper airway

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Chapter 1 - Airway Management
Chapter 3 - General Anesthesia
Cricoid cartilage

The cricoid is a circumferential ring of cartilage in the trachea at the level of the sixth cervical vertebra in adults. In children, it is at the level of the 4th cervical vertebra.

It is significant because it is the only tracheal ring that is complete, the other ones being open posteriorly (horseshoe-shaped). The importance of the cricoid cartilage is that when pressure is applied to it (anteriorly), its posterior aspect will occlude the esophagus.

Cricoid pressure (Sellick’s manoeuvre) is used to reduce the risk of aspiration in susceptible patients.

Related Glossary Terms
Aspiration, Clear fluids, Larynx, Needle cricothyrotomy, NPO, Rapid sequence induction, Sellick’s maneuver, Subglottis
Crystalloid

Crystalloids are solutions of simple inorganic or organic salts. Examples include normal saline, ringers lactate and “2/3 and 1/3”.

They are commonly used for maintenance fluid administration as well as replacement of fluid deficits.

Though administered into the intravascular space, crystalloids distribute throughout the entire extracellular space (not just the intravascular space) due to their low oncotic pressure.

Some types of crystalloids will distribute beyond the extracellular compartment. Hypotonic crystalloids (those that have lower osmotic pressure than extracellular fluid), such as 2/3 and 1/3, will cross into intracellular compartments.

Related Glossary Terms
Colloids, Euvolemia, Maintenance fluid requirements, Pre-operative fluid deficit, Shock, Third space loss
Dantrolene

Dantrolene is used in the treatment of malignant hyperthermia. It is the only specific and direct treatment of malignant hyperthermia. It is also used in the treatment of neuroleptic malignant syndrome.

It is a direct skeletal muscle relaxant which acts at the muscle cellular level, possibly at the ryanodine receptor. It is administered intravenously in 2.5 mg/kg doses until clinical signs show reversal of the hypermetabolic state (usual total dose <20 mg/kg).

Dantrolene can cause generalized muscle weakness in higher doses. Other common side effects include sedation, dizziness and constipation.

Dantrolene is supplied as a powder that must be mixed with sterile water. Its dissolution in water is very slow, difficult and time-consuming. Special “guns” have been devised to speed the preparation of dantrolene so as to minimize any delay in administration in the urgent situation.

Related Glossary Terms
Malignant hyperthermia, Myoglobinuria, Ryanodine

Index
Chapter 5 - Malignant Hyperthermia
Chapter 6 - Drug Finder
Chapter 6 - Miscellaneous
Desflurane

**Class**
Volatile inhaled anesthetic. Used for maintenance of anesthesia.

**Mechanism of Action**
Uncertain

**Dose**
Titrated to effect; MAC (age 40) = 6.0%

**Onset**
Low solubility allows rapid uptake and equilibration. Onset of effect is hastened by using higher flows of carrier gases and by using higher concentrations of volatile agent.

**Duration**
Clinical recovery in less than 10 minutes (2-2.5 x faster washout than Isoflurane)

**Elimination**
Pulmonary (major); negligible hepatic (0.02%)

**Effects**

**CNS**
Desflurane produces an additive central nervous system (CNS) depressant effect along with other sedative/hypnotics and analgesics. Sympatho-excitation can occur with rapid increase in concentration of desflurane. Has the potential to increase intracranial pressure which can be mitigated with hyperventilation. May cause headache, agitation, dizziness.

**CVS**
Dose-related hypotension (vasodilation). Tachycardia and hypertension may be seen due to sympathetic nervous system activation.

**Respiratory**
Respiratory depression with a rapid, shallow respiratory pattern. Loss of intercostal muscle function creates a rocking boat appearance. Desflurane is irritating to the airways and can cause breath-holding, cough, laryngospasm or bronchospasm in susceptible individuals, especially if used as sole agent for induction.

**GI**

**MSK**
Potentiates neuromuscular blockade; malignant hyperthermia trigger.

**Misc.**
Significant carbon monoxide production occurs on exposure to dessicated CO₂ absorbing agents therefore must not be used with low-flow anesthesia. Rapid elimination requires initiation of post-operative analgesia prior to emergence.

**Contraindications**
Malignant hyperthermia susceptibility

**Related Glossary Terms**
Balanced anesthesia, Four A's of anesthesia, Inhaled agents, Isoflurane, MAC, Malignant hyperthermia, Sevoflurane, Vapourizer, Volatile
Difficult airway

The American Society of Anesthesiologists Task Force has defined the difficult airway as follows: "The clinical situation in which a conventionally trained anesthesiologist experiences difficulty with mask ventilation, difficulty with tracheal intubation, or both."

The report contained some more specific definitions.

For example, difficult laryngoscopy is when no part of the larynx is visualized with direct laryngoscopy; difficult endotracheal intubation is when intubation requires more than three attempts or more than 10 minutes.

The incidence of the difficult airway has been reported to be greater in the obstetrical population (8%) than in the non-obstetrical surgical population (around 2%) although some studies have called this conventional thinking into question.

The keys to the management of the difficult airway can be summarized as follows:

a) to do a careful airway assessment in all patients in order to avoid the unanticipated difficult airway wherever possible.

b) to prioritize the establishment of ventilation (with mask or laryngeal mask) when a difficult intubation is encountered.

c) to avoid persistent, traumatic attempts at airway instrumentation.

d) to be comfortable moving to a surgical airway if neither intubation nor bag mask ventilation can be achieved.

e) to develop and maintain skills with a range of airway devices and adjuncts.


Related Glossary Terms

Adjunct, Airway assessment, Antisialagogue, Cormack Lehane, Direct laryngoscopy, Extubation, Fibreoptic bronchoscope, Hypoxemia, Intubation, Laryngeal mask airway (LMA), Laryngoscope, Macintosh blade, Magill blade, Mallampati classification, Mouth opening, Neck motion, Needle cricothyrotomy, Pre-oxygenation, Sniffing position, Stylet, Thyromental Distance, Videolaryngoscope

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Chapter 1 - Airway Management
Chapter 1 - Airway Management
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Chapter 5 - Obstetrical Anesthesia
Chapter 5 - Obstetrical Anesthesia
Dimenhydrinate

Class
Antihistamine, antiemetic. In anesthetic practice, used as a second or third line treatment of post-operative nausea and vomiting (PONV). No role in prevention of PONV.

Mechanism of Action
Dimenhydrinate is a competitive antagonist at the histamine H1 receptor. The antiemetic effects is related to central anticholinergic actions as well as histamine antagonism in the vestibular system in the brain.

Dose
50-100 mg IV q4-6h, max. 400 mg/day (adults)
1.25 mg/kg IV q6h (children)

Onset
5 minutes after IV administration

Duration
4-6 hours

Elimination
Hepatic

Effects
CNS
Sedation (which is additive with alcohol and sedative hypnotics), dizziness, restlessness.

Misc.
May cause dry mouth, blurred vision, difficult urination; more rarely causes acute glaucoma or worsening of asthma. These side effects reflect its anticholinergic activity which is additive with other anticholinergics and monoamine oxidase inhibitors (MAOI).

Related Glossary Terms
Antiemetic agents, Chemoreceptor trigger zone, Ondansetron, Post-operative nausea and vomiting (PONV), Prochlorperazine

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Chapter 4 - Recovery
Chapter 6 - Drug Finder
Chapter 6 - Antiemetics
**Diphenhydramine**

**Class**  
Antihistamine, antiemetic. Used in the treatment of pruritis, allergic reactions and drug-induced extrapyramidal reactions.

**Mechanism of Action**  
Diphenhydramine is a competitive inhibitor at the histamine H1 receptor. The antiemetic effects is related to central anticholinergic effect as well as histamine antagonism in the vestibular system in the brain.

**Dose**  
Adults: 25-50 mg PO q6-8 hours; 10-50 mg IV/IM q 6 hours; maximum daily dose 400 mg.

**Onset**  
IV: 5 minutes  
PO: <15 minutes

**Duration**  
4-6 hours

**Elimination**  
Hepatic

**Effects**  
**CNS**  
Sedation (which is additive with alcohol and sedative hypnotics), dizziness, restlessness.

**CVS**  
Rarely causes hypotension, arrhythmias

**Misc.**  
May cause dry mouth, blurred vision, difficult urination; more rarely causes acute glaucoma or worsening of asthma, and GI or GU obstruction. These side effects reflect its anticholinergic activity, which is additive with other anticholinergics and monoamine oxidase inhibitors (MAOI).

**Related Glossary Terms**  
Drag related terms here
Direct laryngoscopy

Direct laryngoscopy is the procedure where a laryngoscope is used to expose the vocal cords, usually for the purposes of endotracheal intubation.

Direct laryngoscopy is performed using a laryngoscope and is usually done while the patient is under general anesthesia.

Related Glossary Terms
Airway assessment, Cormack Lehane, Difficult airway, Fibreoptic bronchoscope, Intubation, Laryngoscope, Larynx, Macintosh blade, Magill blade, Sniffing position, Vallecula, Videolaryngoscope
Dura

The dura is the outermost (and toughest) layer of the 3 meningeal layers that cover the brain and spinal cord.

The dura is punctured during spinal anesthesia. In epidural anesthesia, one aims to avoid puncturing the dura. Instead, a catheter is placed just outside the dura, in the epidural space.

Related Glossary Terms
Epidural analgesia, Epidural anesthesia, Epidural space, Intrathecal space, Spinal anesthesia, Subarachnoid space
Emergence

During the emergence phase of anesthesia, the patient begins to return to his pre-operative state of consciousness. Emergence requires the offset of effect of the anesthetic agents.

Related Glossary Terms
Apneic threshold, Cough reflex, Extubation, Induction, Maintenance, Post-anesthetic care unit, Recovery, Residual block

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Chapter 3 - General Anesthesia
Chapter 6 - Induction Agents
Chapter 6 - Inhaled Agents
Chapter 6 - Antiemetics
**Ephedrine Sulfate**

**Class**
Sympathomimetic (indirect-acting); vasopressor. Used in the treatment of hypotension.

**Mechanism of Action**
Ephedrine causes more norepinephrine to be released from the storage vesicles in the terminal of neurons thus increasing the amount of norepinephrine in the synaptic space. Ephedrine is (mostly) an “indirect-acting” catecholamine because it doesn’t act at the post-synaptic norepinephrine receptors.

**Dose**
5-20 mg IV (adults)  
25-50 mg IM (adults)

**Onset**
IV: immediate  
IM: minutes

**Duration**
IV: 10-minutes  
IM: 60 minutes

**Elimination**
Hepatic, renal

**Effects**
**CNS**
Increases MAC of volatile anesthetics

**Respiratory**
Bronchodilator

**CVS**
Increases heart rate, contractility and therefore cardiac output (through its β-adrenergic effect). Overall effect is to increase systemic vascular resistance through its α-adrenergic effect. May cause arrhythmias especially when used with volatile anesthetics. As the mechanism of action involves the release of intracellular catecholamines, there is an unpredictable effect in patients with depleted endogenous catecholamines.

**Misc.**
Excessive catecholamine effects may lead to hypertension, tachycardia, arrhythmias, pulmonary edema, anxiety, tremors, hyperglycemia and transient hyperkalemia followed by hypokalemia. Skin necrosis may occur at site of injection.

**Contraindications**
Ephedrine should not be used in patients on monoamine oxidase inhibitors (MAOIs) or those using cocaine. In these patients, phenylephrine is a safer choice for raising blood pressure. Ephedrine should be used with caution in patients who take SSRIs (serotonin-norepinephrine re-uptake inhibitors), as it may increase the risk of “serotonin syndrome”.

**Related Glossary Terms**
Autonomic nervous system, Epinephrine, Phenylephrine, Sympathetic nervous system
Epidural analgesia

In epidural analgesia, a tiny plastic catheter is placed into the epidural space through which local anesthetics and/or opioids are delivered.

The concentration of local anesthesia used for epidural analgesia is lower than that used with epidural anesthesia. When dilute local anesthetics are used, the patient still retains some degree of motor function.

Epidural analgesia is commonly used for labouring parturients as well as for postoperative patients.

Epidural catheters placed for surgical analgesia are most commonly placed at the thoracic or lumbar regions, depending on the site of the surgery. The catheter can be left in place for days or for as long as 2 weeks, as needed.

For labour, epidural catheters are placed in the lower lumbar region. If a Caesarian section is required, the catheter can be used to establish adequate epidural anesthesia for operative delivery.

Related Glossary Terms
Analgesia, Bupivacaine, Dura, Epidural anesthesia, Epidural space, Fentanyl, Lidocaine, Local anesthetic toxicity, Morphine Sulfate, Parturient, Patient controlled analgesia, Regional anesthesia, Second stage of labour, Sympathetic nervous system
Epidural anesthesia

In epidural anesthesia, a tiny plastic catheter is placed into the epidural space through which local anesthetics and or opiates are delivered.

The presence of an indwelling catheter allows the block to be extended in height or duration, as required. For example, an epidural catheter could be used for a surgical procedure that lasted 4 hours or more, well beyond the duration of a spinal anesthetic.

An epidural catheter that was being used for labour analgesia can be used as the anesthetic technique when Caesarian section is required. An epidural used during surgical revascularization of a lower limb can be left in place for several days to a week postoperatively to provide postoperative pain. In each case, the volume and concentration of local anesthetic delivered is adjusted to achieve the specific goals.

Related Glossary Terms
Bupivacaine, Dura, Epidural analgesia, Epidural space, Fentanyl, Intrathecal space, Lidocaine, Local anesthetic toxicity, Morphine Sulfate, Parturient, Regional anesthesia, Spinal anesthesia, Sympathetic nervous system

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Epidural space

The epidural space is the anatomic space located just superficial to the dura. It is a potential space, containing fat and veins. Local anesthetics are delivered into the epidural space (through a catheter) during epidural anesthesia or analgesia.

Related Glossary Terms
Dura, Epidural analgesia, Epidural anesthesia, Intrathecal space, Spinal anesthesia
Epiglottis

The epiglottis is a cartilage located at the base of the tongue. It has an important role in preventing aspiration, as it drops over and covers the larynx during the swallowing mechanism, thus directing the food into the esophagus rather than into the trachea.

The epiglottis is the anatomic structure that one looks for when performing direct laryngoscopy. The tip of the laryngoscope is positioned in the vallecula, where the base of the tongue meets the epiglottis. When positioned properly, upward pressure on the laryngoscope blade will successfully lift the epiglottis and expose the vocal folds.

Related Glossary Terms
Aspiration, Cormack Lehane, Direct laryngoscopy, Hypopharynx, Intubation, Laryngoscope, Larynx, Macintosh blade, Magill blade, Upper airway, Vallecula

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Epinephrine

Class
Sympathomimetic. Epinephrine has many uses:
1) Inotropic support in the patient in cardiogenic shock.
2) Bronchodilation in status asthmaticus.
3) Treatment of allergic reactions.
4) Treatment of croup.
5) Resuscitation in cardiovascular collapse of any cause.
6) Prolongation of action of anesthetic solutions.

Mechanism of Action
Epinephrine is a direct-acting sympathomimetic. It stimulates α- and β-adrenergic receptors resulting in a wide range of effects attributable to the sympathetic nervous system.

Dose
Cardiac arrest (adults): 0.5-1.0 mg IV q5min prn
Inotropic support (adults): 0.1-1.0 ug/kg/min
Anaphylaxis/severe asthma:
Adults: 0.1-0.5 mg SC/IM or 1-5 ug/kg IV prn
Children: 0.01 mg/kg SC/IM, maximum 0.5 mg

Onset
IV: immediate
SC/IM: 6-15 minutes

Duration
IV: 5-10 minutes
SC: 1-3 hours

Elimination
Enzymatic degradation

Effects
CNS
Anxiety, headache, stroke
CVS
Hypertension, tachycardia, arrhythmias, angina, pulmonary edema. Increased risk of ventricular arrhythmias is seen when used with volatile anesthetics.
Respiratory
Bronchodilation
Misc.
Skin necrosis at site of injection, hyperglycemia, transient hyperkalemia, followed by hypokalemia.

Related Glossary Terms
Autonomic nervous system, Ephedrine Sulfate, Phenylephrine, Shock, Sympathetic nervous system

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Esophageal intubation

Esophageal intubation is the unintentional placement of the endotracheal tube into the esophagus rather than into the trachea.

Esophageal intubation must be recognized to avoid serious complications such as hypoxia and aspiration. Auscultation of the the chest during manual ventilation reveals the absence of air entry and the chest will not rise in the usual fashion. Auscultation of the epigastrium reveals low-pitched gurgling sounds. The capnograph usually shows no CO2 return (although it is possible to have some CO2 return if CO2 was forced into the stomach during bag mask ventilation).

When an esophageal intubation is recognized, one stops manual ventilation to avoid inflating the stomach and removes the misplaced endotracheal tube. Bag mask ventilation is resumed while preparations are made to optimize the chances of success for the next attempt at intubation.

Related Glossary Terms
Bag mask ventilation, Capnograph, Difficult airway, Direct laryngoscopy, Hypopharynx, Hypoxemia, Intubation

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Chapter 1 - Airway Management
**Etomidate**

**Class**
Short-acting hypnotic; anesthetic induction agent. Useful in hemodynamically-compromised patients.

**Mechanism of action**
Potentiates the inhibitory GABA neurotransmitter resulting in depression of the reticular activating system.

**Dose**
Induction: 0.2-0.6 mg/kg IV

**Onset**
Within one arm-brain circulation time (approximately 20 seconds).

**Duration**
Approximately 5-10 minutes after single induction dose.

**Elimination**
Rapid redistribution from central nervous system (CNS) to lean body tissue accounts for brief duration of action. Ultimately metabolized by hepatic and plasma esterases to inactive products.

**Effects**

**CNS**
CNS depressant, potentiating the depressant effects of opioids, sedatives and volatile anesthetics. Decreases cerebral metabolic rate and intracranial pressure. The cerebroprotective effects of etomidate make it useful in the management of the head-injured patient. Can cause seizure-like activity.

**CVS**
Etomidate is notable for the lack of significant cardiovascular depression that it causes. Therefore it is commonly chosen to facilitate intubation in the trauma patient, patients with hypovolemic shock or other unstable patients.

**Respiratory**
Etomidate causes a brief period of apnea.

**GI**
Nausea and vomiting

**Misc.**
Etomidate suppresses corticosteroid synthesis in the adrenal cortex and can lead to primary adrenal suppression. For this reason, its use in patients with sepsis is controversial. Etomidate can result in trismus if administered too quickly.

**Related Glossary Terms**
Addisonian crisis, Adrenal suppression, Induction, Ketamine, Propofol, Shock, Sodium Thiopental

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Euvolemia indicates that the patient has ideal intravascular volume. If a patient is volume depleted then they are “hypovolemic”; if they are volume overloaded, they are “hypervolemic”.

Related Glossary Terms
Blood products, Colloids, Crystalloid, Maintenance fluid requirements, Pre-operative fluid deficit, Shock, Third space loss
Extubation

Extubation is the removal of the endotracheal tube and is usually performed immediately upon emergence from anesthesia. Evidence suggests that significant morbidity and mortality relating to the airway are as common at emergence (at the time of extubation) as they are on induction (at the time of intubation). Readiness for extubation must be assessed in order to avoid post-extubation complications which include:

- airway obstruction
- aspiration
- inadequate ventilation
- laryngospasm

A patient who is ready to be extubated will be:

- awake enough to obey commands.
- breathing spontaneously with adequate tidal volume.
- strong (as demonstrated through hand grip or head lift); this criteria is most important when non-depolarizing muscle relaxants (NDMR) have been used.

Related Glossary Terms
Airway obstruction, Anticholinesterase, Aspiration, Difficult airway, Emergence, Laryngospasm, Non-depolarizing muscle relaxants, Patency, Residual block, Reticular activating system, Tidal volume
**Fentanyl**

**Class**
Synthetic opioid analgesic (intermediate-acting); adjunct to anesthesia. Fentanyl can be used as an additive to spinal and epidural anesthesia/analgesia.

**Mechanism of Action**
Acts at the mu-and kappa opioid receptors.

**Dose**
General anesthesia: 1-20 ug/kg IV according to physical status, other agents used, duration and nature of surgery.

**Onset**
IV 4-6 minutes

**Duration**
IV 30-45 minutes

**Elimination**
Hepatic

**Effects**

**CNS**
Potent analgesic effects; some sedative effect. Rarely causes blurred vision, seizures. All of the depressant effects of fentanyl are potentiated by concurrent use of sedatives, volatile anesthetics and nitrous oxide.

**CVS**
Hypotension, bradycardia. The synthetic opioids are not direct myocardial depressants but they do reduce sympathetic drive which may result in decreased cardiac output in patients who are relying on sympathetic tone to support their circulation such as those in hypovolemic or cardiogenic shock.

**Respiratory**
Respiratory depression which at the extreme leads to apnea.

**GI**
Nausea, vomiting, biliary tract spasm, constipation.

**Misc**
Muscle rigidity

**Related Glossary Terms**
Alfentanil, Analgesia, Apneic threshold, Balanced anesthesia, Epidural analgesia, Epidural anesthesia, Four A's of anesthesia, Morphine Sulfate, Naloxone, Opioids, Remifentanil, Sufentanil

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Fibreoptic bronchoscope

Fibreoptic bronchoscopy is the approach that is used when performing an awake intubation. After applying topical anesthetic to the airway, the anesthesiologist uses a fibreoptic bronchoscope (loaded with an endotracheal tube) to identify and pass through the larynx into the trachea. Once in the trachea, the bronchoscope serves as a guide over which the endotracheal tube is passed.

Fibreoptic bronchoscopy is used in the management of patients with airways that are known or suspected to be “difficult” from the perspective of direct laryngoscopy. Fibreoptic intubations are also indicated in patients with unstable cervical spines because they can be performed with the neck in the neutral position rather than in the sniffing position. It is also used in the occasional patient with very limited mouth opening as it can be passed though the smallest of openings.

Fibreoptic bronchoscopic intubations are usually performed on the awake patient. Opioids and/or anxiolytics are used judiciously as adjuncts to blunt the cough reflex as well as to make the experience more palatable for the patient.

Fibreoptic bronchoscopy is not easily performed on the anesthetized patient. After the tube is placed, anesthesia is then induced.

The fibreoptic bronchoscope, passed through an existing endotracheal tube, can be also used by the anesthesiologist to examine the lower airways and remove secretions or other matter.

Related Glossary Terms
Adjunct, Airway assessment, Antisialagogue, Cough reflex, Difficult airway, Direct laryngoscopy, Intubation, Laryngoscope, Larynx, Lower airway, Mallampati classification, Midazolam, Mouth opening, Neck motion, Opioids, Sniffing position

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Chapter 1 - Airway Management
Flowmeter

The anesthetic flowmeter is a component of the anesthetic delivery unit (anesthetic machine) that simultaneously controls and displays the rate of delivery of a given medical gas. Flowmeters are used for the delivery of air, oxygen and nitrous oxide. (Volatile anesthetic gases are delivered via a vapourizer, not a flowmeter.)

The flowmeter allows delivery of one or two (but not three) gases at once.

Many different designs of flowmeters have been used over the years. There are numerous sophisticated safety features built into the design of the modern anesthetic flowmeter.

Related Glossary Terms
Anesthetic circuit, Nitrous oxide (N2O), Volatile
Flumazenil

Flumazenil is a benzodiazepine antagonist. It can be used to reverse the effect of a benzodiazepine like midazolam.

Related Glossary Terms
Midazolam

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Forced air warming system

A forced air warming system is a device that will blow warmed air over the patient. Two different types of blankets (upper body and lower body) can be used to appropriately exclude the surgical site. Forced air warming should be applied to patients undergoing all but the briefest of surgical procedures.

Of note, the device can also be used to cool patients who are hyperthermic (such as those experiencing a malignant hyperthermic reaction), a much less commonly-indicated application.

Related Glossary Terms
Hypothermia, Malignant hyperthermia
Four A’s of anesthesia

Lack of Awareness: unconsciousness.

Amnesia: lack of memory of the event.

Analgesia: implies the abolition of the subconscious reactions to pain. This includes somatic reflexes (such as movement or withdrawal) as well as the autonomic reflexes (such as hypertension, tachycardia, sweating and tearing).

Akinesia: implies lack of overt movement as well as the provision of adequate degree of muscle relaxation according to the procedure.

Related Glossary Terms
Analgesia, Autonomic nervous system, Awareness, Balanced anesthesia, Desflurane, Isoflurane, MAC, Maintenance, Nitrous oxide (N2O), Non-depolarizing muscle relaxants, Opioids, Reticular activating system, Sevoflurane, Sympathetic nervous system, Total intravenous anesthesia, Volatile

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Functional residual capacity

Functional residual capacity (FRC) is the volume of air left in the lungs at the end of exhalation during quiet respiration. It represents a balance between two opposing forces: elastic recoil of the lung and the tendency of the ribcage to spring outwards.

FRC is reduced in pregnant patients, pediatric patients and those patients who are obese. A reduced FRC can lead to rapid desaturation on induction of anesthesia. Careful pre-oxygenation is even more important in these patients. (Access glossary index for diagram)
Glycogen

Glycogen is a polysaccharide that serves as a form of long-term energy storage. Glycogen is primarily located in the liver and to a lesser extent, in muscle. Glycogen is broken down to glucose and used to maintain serum glucose levels during fasting.

Glycogen synthesis and breakdown are controlled by insulin and glucagon, each hormone counterbalancing the effect of the other.

Related Glossary Terms
Drag related terms here

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Chapter 5 - Pediatric Anesthesia
**Glycopyrrolate**

**Class**
Anticholinergic. Clinical uses in anesthesia include the treatment of bradycardia; as an antisialogogue for awake intubation; or (most commonly) for counteracting the muscarinic effects of the anticholinesterases used for the reversal of neuromuscular blockade.

**Mechanism of Action**
An acetylcholine receptor blocker active at the muscarinic (not nicotinic) acetylcholine receptors. Therefore, glycopyrrolate has an anti-parasympathetic effect.

**Dose**
Antisialogogue: 0.1-0.2 mg IV/IM/SC in adults or 4-6 ug/kg IV/IM/SC in children

With anticholinesterase: 0.01 mg/kg IV

**Onset**
IV: <1 minute
IM/SC: 30-45 minutes

**Duration**
Vagal blockade: 2-3 hrs
Antisialogogue effect: 7 hours

**Elimination**
Renal, hepatic

**Effects**
Most effects result from the anticholinergic action of glycopyrrolate.

**CNS**
Confusion is less common than with atropine, as glycopyrrolate does not cross the blood brain barrier. May cause headache, dizziness, mydriasis, blurred vision, increased intraocular pressure.

**CVS**
Causes tachycardia at high doses and may cause bradycardia at low doses.

**GU**
Urinary hesitancy, retention

**Misc.**
Must be used in caution in patients with glaucoma, gastrointestinal or genitourinary obstruction.

**Related Glossary Terms**
Acetylcholine, Anticholinergic, Anticholinesterase, Antisialogogue, Atropine, Autonomic nervous system, Muscarinic, Vagus nerve

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Hypopharynx

The hypopharynx is the lower part of the pharynx. Just below the level of the hypopharynx, the upper airway splits into the esophagus and the larynx.

Related Glossary Terms
Epiglottis, Esophageal intubation, Laryngeal mask airway (LMA), Larynx, Upper airway

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Chapter 1 - Airway Management
Hypothermia indicates a low core body temperature. Intra-operative loss of body heat can occur through four mechanisms:

- Convection (e.g. exposure to drafts of cool air)
- Conduction (e.g. contact with cold operating room table)
- Evaporation (e.g. airway mucosa, prep solution, sweat)
- Radiation (e.g. temperature gradient between patient and operating room environment)

Hypothermia has effects on almost every organ system. These include decreased wound healing, increased wound infection, decreased platelet function and increased systemic vascular resistance.

**Related Glossary Terms**
Forced air warming system
Hypoxemia

Hypoxemia is low oxygen tension in the blood. It can quickly lead to inadequate oxygen delivery to vital organs, especially the brain, heart, kidney and liver.

Related Glossary Terms
Difficult airway, Esophageal intubation, Functional residual capacity, Intubation, Laryngospasm, Mechanical ventilation, Oxygen saturation, Oxyhemoglobin dissociation curve, Positive pressure ventilation, Pre-oxygenation, Pulse oximeter
Induction

Induction is the phase of anesthesia where unconsciousness is first achieved, usually through the administration of a short-acting hypnotic agent. Induction can also be achieved with inhaled agents as is most commonly seen in the care of pediatric patients.

During the induction period, the patient usually becomes apneic and ventilation is maintained with bag-mask ventilation until (in most cases) a definitive airway is achieved.

The period of induction is a treacherous time for the patient. Cardiovascular instability can result due to the effects of the induction agent. The airway is not yet secured, so aspiration is a risk. The anesthesiologist must maintain a patent airway during bag mask ventilation and be prepared for an unexpected difficult intubation.

Related Glossary Terms
Aspiration, Balanced anesthesia, Difficult airway, Emergence, Etomidate, Functional residual capacity, Intubation, Ketamine, Maintenance, Mask Airway, Pre-oxygenation, Propofol, Rapid sequence induction, Sodium Thiopental

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Inhaled agents

The term “inhaled agent” refers to the volatile agents (commonly desflurane, isoflurane, sevoflurane) and nitrous oxide (N₂O).

Related Glossary Terms
Desflurane, Isoflurane, MAC, Nitrous oxide (N₂O), Sevoflurane, Volatile

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Intralipid

Intralipid is a fat emulsion. It is used most commonly in total parenteral nutrition (TPN). It is also present in propofol. Anecdotal reports suggest that intralipid is a useful treatment of local anesthetic toxicity. The exact mechanism is unknown but it is suggested that it acts as a “lipid sink”, binding and therefore sequestering local anesthetic.

The appropriate dose is unknown. Suggested dose (of a 20% intralipid solution) is 1.5 mg/kg bolus which can be repeated once or twice as needed.

Related Glossary Terms
Bupivacaine, Lidocaine, Local anesthetic toxicity, Propofol
Intrathecal space

The intrathecal space is another term for the subarachnoid space. It refers to the fluid-filled space between the arachnoid and the pia mater of the brain and spinal cord. It is the space into which local anesthetic is injected during spinal anesthesia.

Related Glossary Terms
Baricity, Dura, Epidural anesthesia, Epidural space, Pencil-point, Spinal anesthesia, Subarachnoid space

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Intravenous regional anesthetic

An intravenous regional anesthetic (IVRA or “Bier block”) is a technique introduced by August Bier in 1908 and still in use today. The operative limb (arm or leg) is “exsanguinated” using gravity and an elastic bandage after which an arterial cuff is inflated proximally to maintain exsanguination. Subsequently to exsanguination, large volumes of dilute lidocaine are injected (through a small cannula that is placed prior to exsanguination) into the affected limb.

Surgical anesthesia can last for 60-90 minutes, usually limited by the pain created by the inflated cuff. It is important that the cuff pressure be maintained for at least 30 minutes to avoid a sudden bolus of local anesthetic being introduced into the systemic circulation.
**Intubation**

Endotracheal intubation is the process in which a breathing tube is placed through the larynx into the trachea. Intubation can be achieved through a variety of techniques. It is usually performed to achieve one or more of the following goals:

a) maintain airway patency

b) protect the airway from aspiration

c) facilitate the delivery of mechanical ventilation

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**Related Glossary Terms**

Airway assessment, Aspiration, Bag mask ventilation, Capnograph, Cormack Lehane, Difficult airway, Esophageal intubation, Fibreoptic bronchoscope, Hypoxemia, Induction, Laryngoscope, Larynx, Lower airway, Macintosh blade, Magill blade, Mallampati classification, Mechanical ventilation, Mouth opening, Nasotracheal intubation, Needle cricothyrotomy, Rapid sequence induction, Right mainstem bronchus, Sniffing position, Stylet, Vallecula, Videolaryngoscope

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Isoflurane

Class
Volatile inhaled agent. Used for maintenance of anesthesia.

Mechanism of Action
Uncertain

Dose
Titrated to effect; MAC (age 40)=1.15

Onset
Higher solubility than sevoflurane and desflurane therefore uptake is slower than the modern agents. Onset of effect is hastened by using higher flows of carrier gases and by using higher concentrations of volatile agent.

Duration
Clinical recovery in less than 15 minutes (usually). Theoretically a slower wake-up than the modern agents due to higher solubility.

Elimination
Pulmonary

Effects
CNS
Isoflurane produces an additive central nervous system (CNS)-depressant effect along with other sedative/hypnotics and analgesics. Has the potential to increase intracranial pressure which can be mitigated with hyperventilation. Delirium.

CVS
Dose-related hypotension (vasodilation).

Respiratory
Respiratory depression with rapid, shallow respiratory pattern. Loss of intercostal muscle function creates a rocking boat appearance. Isoflurane is irritating to the airways and can cause breath-holding, cough, laryngospasm or bronchospasm. Its pungent quality makes it unsuitable for use with a mask induction.

GI
Nausea, vomiting.

MSK
Potentiates neuromuscular blockade. Malignant hyperthermia trigger.

Contraindications
Malignant hyperthermia susceptibility.

Related Glossary Terms
Balanced anesthesia, Desflurane, Four A's of anesthesia, Inhaled agents, MAC, Malignant hyperthermia, Sevoflurane, Vapourizer, Volatile
Ketamine

**Class**
Phencyclidine derivative. Can be used as an induction agent (usually in hemodynamically-compromised patients) or for sedation during painful procedures.

**Mechanism of action**
Acts at numerous central nervous system receptor sites, including the N-methyl-D-aspartate (NMDA) receptor.

**Dose**
Induction of anesthesia: 2 mg/kg IV
Induction of anesthesia: 5 mg/kg IM

**Onset**
Within one arm-brain circulation time (approximately 20 seconds).

**Duration**
Approximately 10-15 minutes after single induction dose, with full orientation occurring after 15-30 minutes.

**Elimination**
Redistribution from central nervous system (CNS) to inactive tissue sites accounts for termination of unconsciousness. Ultimate clearance is via hepatic metabolism and renal excretion.

**Effects**

**CNS**
Produces "dissociative anesthesia" with patient in a cataleptic state. Ketamine provides a state of unconsciousness and intense analgesia however the patient’s eyes may remain open and roving, and their limbs may move purposelessly. Cerebral metabolic rate and intracranial pressure are increased.

**CVS**
Ketamine increases sympathetic outflow from the CNS leading to increased heart rate, blood pressure and cardiac output. Because of this effect, ketamine plays an important role in the management of patients with hypovolemic shock or cardiac tamponade. However, ketamine does possess direct myocardial depressant effects which may lead to worsened hypotension in patients in a prolonged shock state.

**Respiratory**
Some degree of airway protection is maintained. The patient may cough or swallow. Airway secretions increase. Bronchodilatory effect is secondary to increased sympathetic tone. Apnea is rare as respiratory drive is maintained.

**Misc.**
Undesirable psychological reactions are common on emergence: vivid, unpleasant dreams, excitement, confusion, fear. They tend to occur in the first hour of emergence and abate within one to several hours. Pretreatment with benzodiazepines may help minimize this effect.

**Contraindications**
Raised intracranial pressure, coronary ischemia, psychiatric disease, eye surgery.

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**Related Glossary Terms**
Etomidate, Induction, Propofol, Shock, Sodium Thiopental

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Ketorolac Tromethamine

Class
Non-steroidal anti-inflammatory analgesic. Can be used orally or intravenously.

Mechanism of Action
COX-1 inhibitor

Dose
15-30 mg IV (adults)
0.5 mg/kg, max. 15 mg/dose (children > 2 years)
IV therapy should not exceed 2 days.

Onset
<30 minutes; peak effect in 1-2 hours.

Duration
4-6 hours

Elimination
Renal (92%); enterohepatic (6%)

Effects
CNS
Analgesic with an opioid-sparing effect.

GI
Gastrointestinal bleeding, ulcer, perforation, nausea, dyspepsia.

Renal
May precipitate renal failure in patients with renal insufficiency or those using ACE inhibitors.

Hematologic
May potentiate effects of anticoagulants. Rarely may cause hemorrhage due to platelet inhibition.

Related Glossary Terms
Adjunct, Analgesia
Laryngeal mask airway (LMA)

The LMA is an airway device that is a hybrid of the mask and the endotracheal tube. It is inserted blindly into the hypopharynx. When properly positioned, with its cuff inflated, it sits above the larynx and seals the glottic opening. It is usually used for spontaneously breathing patients but positive pressure ventilation can be delivered through an LMA. The LMA does not protect against aspiration.

Like an endotracheal tube, the LMA frees up the anesthesiologist’s hands and allows surgical access to the head and neck area without interference. While airway obstruction due to laryngospasm is still a risk, the LMA prevents upper airway obstruction from the tongue or other soft tissues.

The LMA also has a role to play in the failed intubation setting, particularly when mask ventilation is difficult.

Related Glossary Terms
Airway obstruction, Aspiration, Bag mask ventilation, Difficult airway, Hypopharynx, Laryngospasm, Larynx, Mask Airway, Upper airway
Laryngoscope

A laryngoscope is the device used to expose the vocal cords in order to pass an endotracheal tube. It is most commonly used under general anesthesia. It is used in most routine intubations.

The most common laryngoscope blade is the Macintosh blade. It is positioned in the vallecula such that upward pressure on the tongue will expose the larynx.

The traditional laryngoscope allows direct laryngoscopy. Newer devices, such as the video-laryngoscope, allow for indirect laryngoscopy where the larynx is viewed on a screen and a direct line of sight is not achieved.

Related Glossary Terms
Cormack Lehane, Difficult airway, Direct laryngoscopy, Fibreoptic bronchoscope, Intubation, Larynx, Macintosh blade, Magill blade, Vallecula, Videolaryngoscope
Laryngospasm

Laryngospasm is the reflexive closure of the glottis. Laryngospasm is the airway’s response to irritation. It can occur immediately after extubation, leading to total airway obstruction, particularly in children and young adults.

Extubating the patient at a deep plane of inhalational anesthesia, when the reflex is blunted, is one way to reduce the risk of laryngospasm but is an approach that is only safely applied to the pediatric patient.

In adults (and pediatric patients), performing extubation when the patient is wide awake, (such that consciousness abolishes the reflex), will decrease the risk of post-extubation laryngospasm. Performing extubation when the patient is at a light plane of anesthesia (not awake, but not “asleep” either) is generally thought to increase the risk of post-extubation laryngospasm.

Laryngospasm is more likely to occur in patients with inadequate reversal of muscle relaxation. It is more common in smokers. Laryngospasm may be an indication of aspiration.

Treatment of laryngospasm must be individualized to the patient according to the presumed underlying cause and the severity of obstruction. Suctioning the airway clear of mucous or other irritants is important. Ensuring full reversal of muscle relaxation is also key. Simple maneuvers such as airway support and gentle positive pressure ventilation will often “break” the spasm. Very occasionally, administration of anesthetic agents (possibly including a very small dose of muscle relaxant) will be required to relieve the airway obstruction.

It is important that the anesthesiologist remain alert to the airway after extubation to recognize laryngospasm (which can be silent when obstruction is complete) so that deep levels of hypoxia are not reached.
Larynx

The larynx is that part of the airway that connects the pharynx and hypopharynx to the trachea. It serves several functions: breathing, airway protection and phonation. The vocal cords are just part of the laryngeal structure.

The larynx is located from the 3rd to 6th cervical vertebrae in adults.

Related Glossary Terms
Airway obstruction, Direct laryngoscopy, Epiglottis, Fibreoptic bronchoscope, Hypopharynx, Intubation, Laryngeal mask airway (LMA), Laryngoscope, Laryngospasm, Lower airway, Recurrent laryngeal nerve, Subglottis, Superior laryngeal nerve, Upper airway, Vallecula
Laudanosine

Laudanosine is a metabolite of atracurium and (to a lesser extent) cis-atracurium. Laudanosine can lower the seizure threshold (and therefore cause seizures) in susceptible individuals.

Related Glossary Terms
Atracurium, Cis-Atracurium, Non-depolarizing muscle relaxants
Lidocaine

**Class**
Local anesthetic. Used in infiltration anesthesia and regional anesthesia (e.g., intravenous regional anesthesia). Lidocaine is still used for epidural anesthesia, especially for Caesarian section. Lidocaine is rarely used in spinal anesthesia due to associated nerve irritation. Lidocaine is occasionally used in the treatment of ventricular arrhythmias.

**Mechanism of Action**
Sodium channel blocker

**Dose**
Anesthetic:
- Maximum 4 mg/kg without epinephrine
- Maximum 7 mg/kg with epinephrine

Anti-arrhythmic:
- 1 mg/kg IV bolus followed by 0.5 mg/kg q 2-5 minutes to maximum 3 mg/kg/hr
- By infusion (of 0.1% solution): 1-4 mg/min (20-50 ug/kg/min)

**Onset**
- IV: 45-90 seconds
- Infiltration: 0.5-1 minute
- Epidural: 5-15 minutes
- Spinal: <1 minute

**Duration**
- IV: 10-20 minutes
- Infiltration: 0.5-1 hour
- Epidural and spinal: 1-3 hours

**Elimination**
hepatic, pulmonary

**Effects**
Local anesthetics should not have systemic effects if used appropriately. If high plasma levels are achieved due to incorrect dosing or inadvertent intravascular injection then the symptoms manifest firstly in the central nervous system and then in the cardiovascular system where hypotension, heart block and other arrhythmias may occur. Premonitory signs and symptoms are perioral numbness, metallic taste, tinnitus, restlessness, dizziness and tremors. Seizures, respiratory and circulatory depression/arrêt may occur. The treatment is supportive care and the use of Intralipid. Administration of benzodiazepines will increase the seizure threshold.

High intravascular concentrations of local anesthetics may potentiate the effects of muscle relaxants (both depolarizing and non-depolarizing).

**Related Glossary Terms**
Bier Block, Brachial plexus block, Bupivacaine, Epidural analgesia, Epidural anesthesia, Intralipid, Intravenous regional anesthetic, Local anesthetic toxicity, Spinal anesthesia

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Chapter 3 - Regional Anesthesia
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Chapter 3 - Regional Anesthesia
Chapter 6 - Drug Finder
Chapter 6 - Local Anesthetics
**Lithotomy**

The lithotomy position is the position which sees the patient on his or her back, hips and flexed and abducted. It is used for many types of gynecologic, urologic and ano-rectal procedures.

**Related Glossary Terms**

Neuropathy, Prone, Supine, Trendelenburg
Local anesthetic toxicity

Local anesthetic toxicity causes symptoms of tinnitus, perioral numbness, metallic taste in the mouth and dizziness. Ultimately, it will lead to seizures, arrhythmias and cardiovascular collapse. It is a risk of any regional technique that uses significant doses of local anesthetic.

Related Glossary Terms
Bier Block, Brachial plexus block, Bupivacaine, Epidural analgesia, Epidural anesthesia, In-tralipid, Intravenous regional anesthetic, Lidocaine

Index
Chapter 3 - Anesthetic Techniques
Chapter 3 - Regional Anesthesia
Lower airway

The lower airway begins below the level of the larynx and includes the subglottic region, the trachea, mainstem bronchi and bronchioles.

Related Glossary Terms
Cough reflex, Fibreoptic bronchoscope, Intubation, Larynx, Right mainstem bronchus, Subglottis, Upper airway

Index
Chapter 1 - Airway Management
Chapter 1 - Airway Management
MAC

MAC refers to the concentration inhaled agent in alveolar gas necessary to prevent movement of 50% of patients when a standard incision is made. The factors that affect the MAC for a given patient are outlined below.

<table>
<thead>
<tr>
<th>FACTORS WHICH DECREASE MAC</th>
<th>FACTORS WHICH INCREASE MAC</th>
</tr>
</thead>
<tbody>
<tr>
<td>advanced age</td>
<td>childhood</td>
</tr>
<tr>
<td>pregnancy</td>
<td>hyperthyroidism</td>
</tr>
<tr>
<td>hypothermia</td>
<td>hyperthermia</td>
</tr>
<tr>
<td>acute alcohol intoxication</td>
<td>chronic alcohol use</td>
</tr>
<tr>
<td>drugs: benzodiazepines, opiates, muscle relaxants, central-acting antihypertensives</td>
<td>drugs: amphetamine, cocaine</td>
</tr>
</tbody>
</table>

Related Glossary Terms
Desflurane, Four A’s of anesthesia, Inhaled agents, Isoflurane, Nitrous oxide (N2O), Sevoflurane, Vapourizer, Volatile
Macintosh blade

The Macintosh blade is a curved blade used in direct laryngoscopy and is the one in most common use today. Size 3 is the standard size for use on an average-sized adult. Smaller sizes are used in children; a size 4 is available for very large adults.

The tip of the Macintosh blade is placed in the vallecula such that upward pressure on the tongue will lift the epiglottis, thus exposing the larynx.

Related Glossary Terms
Direct laryngoscopy, Epiglottis, Intubation, Laryngoscope, Magill blade, Vallecula

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Chapter 1 - Airway Management
Magill blade

A Magill blade, used in direct laryngoscopy, is straight (not curved). It is designed to lift the epiglottis directly. It is useful in the pediatric patient due to the long angled epiglottis seen in this patient population.

Related Glossary Terms
Cormack Lehane, Direct laryngoscopy, Epiglottis, Intubation, Laryngoscope, Macintosh blade

Index
Chapter 5 - Pediatric Anesthesia
Maintenance

Maintenance of anesthesia requires the delivery of pharmacologic agents with the aim of achieving the “four A’s of anesthesia” and hemodynamic stability throughout the surgical procedure.

Related Glossary Terms
Balanced anesthesia, Emergence, Four A’s of anesthesia, Induction, Total intravenous anesthesia

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Chapter 3 - General Anesthesia
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Chapter 3 - General Anesthesia
Chapter 3 - General Anesthesia
Chapter 3 - General Anesthesia
Maintenance fluid requirements correlate best with lean body mass and body surface area.

To calculate maintenance, use the “4/2/1 rule”:

- First 10 kilograms (i.e. 0-10 kg): 4 cc/kg/hr
- Next 10 kilograms (i.e. 11-20 kg): 2 cc/kg/hr
- All remaining kilograms over 20 kg: 1 cc/kg/hr

For example, a 60 kg woman fasting for 8 hours:

\[
\begin{align*}
10 \text{ kg} \times 4 \text{ cc/kg/hr} & = 40 \text{ cc/hr} \\
10 \text{ kg} \times 2 \text{ cc/kg/hr} & = 20 \text{ cc/hr} \\
40 \text{ kg} \times 1 \text{ cc/kg/hr} & = 40 \text{ cc/hr} \\
\text{Total} & = 100 \text{ cc/hr} \times 8 \text{ hr} \\
& = 800 \text{ cc}
\end{align*}
\]

Therefore, the pre-operative deficit (excluding other losses) is 800 cc.

Related Glossary Terms
- Colloids
- Crystalloid
- Euvolemia
- Pre-operative fluid deficit
- Shock
- Third space loss

Chapter 1 - Fluid Management
Chapter 5 - Pediatric Anesthesia
Malignant hyperthermia

Malignant hyperthermia (MH) is a potentially life-threatening pharmacogenetic disorder characterized by the onset of a hypermetabolic crisis in response to certain triggers. Since the usual triggers are succinylcholine and volatile anesthetics, MH is known as “the anesthesiologist’s disease”.

Immediate discontinuation of triggering agents is key if an MH reaction is suspected. Dantrolene, a direct skeletal muscle relaxant, is the only specific treatment for an MH reaction. Active cooling of the hyperthermic patient may also be required.

Related Glossary Terms
Dantrolene, Desflurane, Forced air warming system, Isoflurane, Myoglobinuria, Ryanodine, Sevoflurane, Succinylcholine

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Chapter 3 - General Anesthesia
Chapter 5 - Malignant Hyperthermia
Chapter 6 - Muscle Relaxants
Chapter 6 - Inhaled Agents
Chapter 6 - Miscellaneous
Mallampati classification

The Mallampati classification assigns a score, based on the structures visualized when the patient is sitting upright, with the head in a neutral position and the tongue protruding maximally. Class 1 corresponds well with an easy intubation. Class 4 corresponds well with a difficult intubation. Classes 2 and 3 less reliably predict ease of intubation.

<table>
<thead>
<tr>
<th>Class 1</th>
<th>Soft palate, uvula, tonsillar pillars can be seen.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Class 2</td>
<td>As above except tonsillar pillars not seen.</td>
</tr>
<tr>
<td>Class 3</td>
<td>Only base of uvula is seen.</td>
</tr>
<tr>
<td>Class 4</td>
<td>Only tongue and hard palate can be seen.</td>
</tr>
</tbody>
</table>

Related Glossary Terms
Airway assessment, Cormack Lehane, Difficult airway, Direct laryngoscopy, Esophageal intubation, Intubation
**Mask Airway**

A mask airway can be achieved with the correct placement of a mask over the patient’s face along with appropriate jaw lift (with or without an oral airway) in order to create a patent airway. Positive pressure can then be performed to achieve adequate ventilation and oxygenation.

Bag mask ventilation is performed immediately after induction of anesthesia prior to intubation or the insertion of a laryngeal mask. Occasionally, the mask airway is the airway of choice for the duration of the anesthetic, as can be seen in very short procedures.

Effective bag mask ventilation is key during failed intubation attempts in order to ensure adequate oxygenation of the patient.

**Related Glossary Terms**

Bag mask ventilation, Induction, Intubation, Laryngeal mask airway (LMA), Positive pressure ventilation

**Index**

Chapter 1 - Airway Management
Mechanical ventilation

Mechanical ventilation is the process of achieving tidal volume through the application of positive pressure to the patient’s airway. It is most easily achieved through an endotracheal tube or tracheostomy tube.

Mechanical ventilation is required in patients receiving muscle relaxation.

Related Glossary Terms
Hypoxemia, Intubation, Positive pressure ventilation, Tidal volume, Ventilator
Meperidine

Class
Opioid analgesic (long acting). Traditionally used for postoperative pain but currently its use is restricted (in many hospitals) to the treatment of postoperative shivering.

Mechanism of Action
Acts at the mu and kappa opioid receptors.

Dose
In adults: 25-75 mg IV/IM (0.5-2 mg/kg)

Onset
IV: 3-8 minutes
IM: 10-20 minutes

Duration
2-4 hours IV/IM

Elimination
Hepatic

Effects
CNS
Causes dose-related sedation; variable analgesic effect. Delirium in older patients is often seen. May cause seizures if used in large doses or over an extended time frame due to the accumulation of its excitatory metabolite, normeperidine. May cause euphoria and dysphoria. All of the depressant effects of meperidine are potentiated by concurrent use of sedatives, volatile anesthetics, nitrous oxide and tricyclic antidepressants.

Respiratory
Respiratory depression which at the extreme leads to apnea. May promote bronchospasm in susceptible patients (those with asthma or COPD).

GI
Nausea, vomiting, biliary tract spasm, constipation.

Misc.
Effective in the treatment of postoperative shivering. May cause muscle rigidity, urticaria, pruritis.

Contraindications:
Meperidine must not be used in patients on monoamine oxidase inhibitors in whom it can cause a fatal reaction.

Related Glossary Terms
Naloxone, Opioids
Metabolic acidosis

Metabolic acidosis occurs when the blood concentration of bicarbonate is lower than normal. It can occur as a primary defect (as a result of one of a long list of causes) or as a compensatory response to a respiratory alkalosis. Patients with metabolic acidosis can be acidemic, alkalemic or have a normal blood pH.

Related Glossary Terms

Shock
Midazolam

**Class**
Short-acting benzodiazepine. Used for sedation or as an adjunct during general anesthesia. Midazolam has anxiolytic and sedative (but not analgesic) properties.

**Mechanism of Action**
Agonism at the inhibitory GABA receptor.

**Dose**
For sedation: 0.03-0.08 mg/kg IV
Can also be given intramuscularly, intranasally and orally.

**Onset**
Within 3-5 minutes

**Duration**
Elimination half-time is 1-4 hours, making midazolam a much shorter acting agent than diazepam.

**Elimination**
Metabolized in the liver by microsomal enzymes and excreted in the urine.

**Effects**

**CNS**

**CVS**
In larger doses, in the presence of hypovolemia or when used in combination with opioids, midazolam can lead to decreased blood pressure and increased heart rate. Cardiac output is unchanged.

**Respiratory**
Dose-related respiratory depression occurs. This response is exaggerated in the elderly, in those with COPD or when used in combination with opioids.

**Misc.**
Midazolam is water-soluble therefore the pain on injection and phlebitis that are seen with diazepam are uncommon.

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**Related Glossary Terms**
Fibreoptic bronchoscope, Flumazenil

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**Index**
Fibreoptic bronchoscope, Flumazenil
Morphine Sulfate

**Class**
Opioid analgesic (long acting). In anesthetic practice, its main use is for postoperative analgesia. Morphine is commonly used intravenously and for spinal or epidural anesthesia/analgesia.

**Mechanism of Action**
Active at the mu and kappa opioid receptors.

**Dose**
Adults: 2.5-15 mg IV/IM/SC
Children: 0.05-0.2 mg/kg IV/IM/SC

**Onset**
IV 5-10 minutes
IM 15-30 minutes

**Duration**
2-5 hrs IV/IM/SC

**Elimination**
Hepatic

**Effects**

**CNS**
Reliable analgesic effects; sedation. May cause blurred vision, syncope, euphoria, dysphoria. All of the depressant effects of morphine are potentiated by concurrent use of sedatives, volatile anesthetics, nitrous oxide and alcohol. Morphine’s depressant effects are also potentiated by antihistamines, phenothiazines, butyrophenones, MAOIs and TCAs.

**CVS**
May cause hypotension, hypertension, bradycardia, arrhythmias.

**Respiratory**
Respiratory depression which at the extreme leads to apnea. May cause bronchospasm or laryngospasm.

**GI**
Nausea, vomiting, constipation, biliary tract spasm.

**Misc.**
Releases histamine. May cause pruritis, urticaria, muscle rigidity, urinary retention.

**Related Glossary Terms**
Alfentanil, Analgesia, Apneic threshold, Epidural analgesia, Epidural anesthesia, Fentanyl, Naloxone, Opioids, Patient controlled analgesia, Re-narcotization, Remifentanil, Spinal anesthesia, Sufentanil

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Chapter 4 - Post-operative Pain Management
Chapter 4 - Post-operative Pain Management
Chapter 5 - Obstetrical Anesthesia
Chapter 6 - Drug Finder
Chapter 6 - Opioid agonists and antagonists
**Mouth opening**

Three fingerbreadths is considered adequate mouth opening for the purpose of airway manipulation.

A patient with very limited mouth opening may require a fibreoptic bronchoscopic (awake) intubation.

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**Related Glossary Terms**

Airway assessment, Difficult airway, Fibreoptic bronchoscope, Intubation
Muscarinic acetylcholine (Ach) receptors are the type of Ach receptors that are found on the nerves of the parasympathetic nervous system. The vagus nerve, for example, has muscarinic Ach receptors.

The other type of Ach receptors are the nictoinic variety which are found at the neuromuscular junction.

Related Glossary Terms
Acetylcholine, Anticholinergic, Anticholinesterase, Antisialagogue, Atropine, Glycopyrrolate, Neuromuscular junction, Nicotinic, Vagus nerve
Myasthenia gravis

Myasthenia gravis is a neuromuscular disease of immune origin such that antibodies attack the acetylcholine receptor of the neuromuscular junction.

Because the pathology of the disease is analogous to the pharmacologic effects of NDMR, patients with myasthenia gravis are exquisitely sensitive to the effects of NDMR, which should be avoided in these patients wherever possible.

Anticholinesterases (NDMR reversal agents) are used in the treatment of myasthenia gravis.

Related Glossary Terms
Anticholinesterase, Neuromuscular junction, Non-depolarizing muscle relaxants

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Chapter 3 - General Anesthesia
Chapter 6 - Muscle Relaxants
Chapter 6 - Muscle Relaxants
Chapter 6 - Muscle Relaxants
Chapter 6 - Muscle Relaxants
Chapter 6 - Anticholinesterase and Anticholinergics
Myoglobinuria

When pathological muscle injury or breakdown occurs, myoglobin is released into the bloodstream and can be filtered by the kidney, causing acute tubular necrosis and acute renal failure.

Some causes of myoglobinuria are malignant hyperthermia, compartment syndrome, limb ischemia and traumatic crush injury.

Myoglobinuria is treated by addressing the underlying cause, where possible. Volume resuscitation is key. Alkalization of the urine (by the administration of an intravenous bicarbonate infusion) is also used.

Related Glossary Terms
Dantrolene, Malignant hyperthermia, Ryanodine
Naloxone

Class
Opioid antagonist. Used to counteract the effects of opioids.

Mechanism of Action
Agonist at the opioid receptors.

Dose
For postoperative opioid depression: 1-2 μg/kg IV in 0.5-1 μg/kg boluses, q 2-3 minutes
For neonatal opioid depression: 10 μg/kg, q 2-3 minutes IV. Infusion: 1-5 μg/kg/hr

Onset
1-2 minutes

Duration
30-60 minutes

Elimination
Hepatic

Effects

CNS
Rapid reversal of opioid effect can cause delirium and severe pain.

CVS
When opioid effect is abruptly antagonized there can be significant sympathetic activation leading to hypertension, tachycardia and in susceptible individuals, myocardial ischemia and pulmonary edema.

Misc.
Due to the relatively short duration of action of naloxone, “re-narcotization” can be seen when it is used to treat respiratory depression caused by long acting opioids such as morphine. In this case, close monitoring is indicated and supplemental doses may be necessary.

Related Glossary Terms
Alfentanil, Apneic threshold, Fentanyl, Meperidine, Morphine Sulfate, Neonatal depression, Neonatal opioid depression, Opioids, Re-narcotization, Remifentanil, Sufentanil

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Chapter 6 - Opioid agonists and antagonists
Nasotracheal intubation

Nasotracheal intubation is the procedure of placing a breathing tube (endotracheal tube) into the trachea through the nose rather than through the mouth. Nasotracheal intubation is done to facilitate intra-oral surgery amongst other indications. It carries additional complications, most notably, epistaxis.

Related Glossary Terms
Intubation
Neck motion

Normal range of motion is between 90 and 165 degrees. Restricted range of motion may indicate that direct laryngoscopy will be difficult. Adequate neck mobility is required to achieve the desired “sniffing position” that facilitates visualization of the larynx on direct laryngoscopy.

Related Glossary Terms
Airway assessment, Difficult airway, Direct laryngoscopy, Fibreoptic bronchoscope, Intubation, Sniffing position
Needle cricothyrotomy

A needle cricothyrotomy is an emergency procedure where an airway is achieved percutaneously through the cricothyroid membrane. A cricothyrotomy kit should be on every difficult airway cart.

A cricothyrotomy is usually performed as a last-ditch emergency measure after failed attempts at conventional intubation. It is a temporary, stop-gap measure. A formal tracheostomy will usually be required as soon as is feasible.

Studies of anesthetic morbidity and mortality related to the airway underscore the importance of not hesitating to perform a cricothyrotomy when other measures have failed and oxygenation of the patient is not being maintained.

Related Glossary Terms
Cricoid cartilage, Difficult airway, Hypoxemia, Intubation, Subglottis
**Neonatal depression**

Neonatal depression refers to insufficient respiratory function in the newborn. It can result from administration of opioids to the mother prior to delivery but is also a result of fetal asphyxia.

**Related Glossary Terms**

Naloxone, Neonatal opioid depression

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Chapter 5 - Obstetrical Anesthesia
Neonatal opioid depression

The respiratory system of the neonate can be depressed by opioids delivered to the mother intravenously, intramuscularly and less commonly, epidurally. The effects are reversed with naloxone.
Neostigmine

**Class**
Anticholinesterase. In anesthesia practice, neostigmine is used for the reversal of neuromuscular blockade. Internal Medicine specialists use neostigmine for the treatment of myasthenia gravis.

**Mechanism of Action**
Anticholinesterases inhibit the breakdown of acetylcholine (Ach) in the synaptic cleft by inhibiting the cholinesterase enzyme. As a result, Ach concentrations in the synaptic cleft are increased. Ach is then better able to compete with muscle relaxants for the Ach receptors and achieve depolarization of the muscle cell.

**Dose**
For reversal of neuromuscular blockade: 0.05 mg/kg
Dose should not exceed 5 mg
Must be administered with atropine 0.015 mg/kg or glycopyrrolate 0.01 mg/kg

**Onset**
5 minutes

**Duration**
55-75 minutes

**Elimination**
Hepatic, plasma esterases

**Effects**
Most of neostigmine’s effects are related to its cholinergic action. It must be given with an anticholinergic (atropine or more commonly glycopyrrolate) in order to minimize these effects.

**CNS**
Seizures

**CVS**
Bradycardia, AV block, nodal rhythm, hypotension

**Respiratory**
Increased oral and bronchial secretions, bronchospasm

**GI/GU**
Increased peristalsis, urinary frequency

**Misc.**
Overdose may produce cholinergic crisis. Neostigmine does not antagonize succinylcholine and may prolong phase 1 block of succinylcholine.

**Related Glossary Terms**
Anticholinergic, Anticholinesterase, Cholinesterase, Neuromuscular junction, Non-depolarizing muscle relaxants

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Chapter 4 - Recovery
Neuromuscular junction

The neuromuscular junction is located where the nerve terminal meets the motor end-plate. The motor end-plate is a specialized area of the muscle cell that is responsible for the initiation of an action potential which then propagates through the entire muscle cell.

A stimulated nerve releases acetylcholine from nerve terminal into the synaptic cleft, where it attaches to acetylcholine (Ach) receptors on the motor end-plate, triggering a muscular contraction.

Muscle relaxants exert their action at the neuromuscular junction.

Related Glossary Terms
Acetylcholine, Anticholinergic, Cholinesterase, Competitive inhibitor, Muscarinic, Myasthenia gravis, Neostigmine, Nicotinic, Non-depolarizing muscle relaxants, Peripheral nerve stimulator

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Chapter 3 - General Anesthesia
Chapter 6 - Muscle Relaxants
Neuropathy

A neuropathy occurs as a result of damage to a nerve, often from compression or irritation, which can result in motor and or sensory deficits. It can also result in pain.

In anesthetic practice, neuropathies can be seen due to the effects of patient positioning and as a complication of regional anesthetic techniques.

Related Glossary Terms
Lithotomy, Prone, Radiculopathies, Regional anesthesia
Nicotinic

Nicotinic receptors are the type of acetylcholine (Ach) receptors that are found at the neuromuscular junction. When bound by Ach, the nicotinic receptors act to initiate a muscle contraction. Non depolarizing muscle relaxants act as competitive inhibitors at the nicotinic Ach receptors.

The other type of Ach receptor are the muscarinic receptors which are found in the parasympathetic nervous system.

Both types of receptors are activated by acetylcholine.

Anticholinesterases used to increase the effects of Ach at the nicotinic receptors of the neuromuscular junction also increase the Ach effect at muscarinic receptors. For this reason, the anticholinesterases are always administered with an anticholinergic drug such as glycopyrrolate.

Related Glossary Terms
Acetylcholine, Anticholinergic, Anticholinesterase, Cholinesterase, Competitive inhibitor, Muscarinic, Neuromuscular junction, Non-depolarizing muscle relaxants
Nitrous oxide (N2O)

Class
Nitrous oxide is an inhaled agent but not a volatile agent. It is used as an adjunct to general anesthesia. It has a weak effect and therefore cannot be used as the sole agent for general anesthesia and is most commonly used in combination with a volatile agent. It can be used on its own for sedation or analgesia as can be seen in the obstetric or dental setting.

Mechanism of Action
Uncertain

Dose
Delivered in concentrations of up to 70% in oxygen. Actual MAC is 104%.

Onset
Immediate due to very low solubility.

Duration
Offset of effect is rapid after discontinuation.

Elimination
Pulmonary

Effects
CNS
N2O is a potent analgesic. It increases cerebral metabolic rate, cerebral blood flow and intracranial pressure and is therefore not a good choice for patients with decreased intracranial compliance.

CVS
N2O has a mild sympathomimetic effect but causes direct myocardial depression. The net effect is a modest decrease in blood pressure and heart rate. Increased coronary tone may exacerbate ischemia in susceptible patients.

Respiratory
N2O produces mild respiratory depression which is potentiated by opioids, hypnotics and volatile anesthetics. It has no bronchodilatory effect. It exacerbates pulmonary hypertension.

Misc.
N2O expands the volume of gas-containing spaces as N2O diffuses across membranes more readily than nitrogen can diffuse out. Thus the size of a pneumothorax, emphysematous bleb and distended bowel loop will increase when N2O is used. Bone marrow suppression due to inhibition of methionine synthetase, can occur if N2O is used for extended periods.

N2O enhances opioid-induced rigidity. Finally, N2O is an operating room pollutant; N2O levels (in parts per million) in the operating room environment are measured regularly to comply with workplace safety regulations.

Related Glossary Terms
Analgesia, Flowmeter, Four A's of anesthesia, Inhaled agents, MAC, Volatile
Non-depolarizing muscle relaxants

The non-depolarizing muscle relaxants (NDMR) are those which achieve muscle relaxation through competitive-inhibition of the acetylcholine receptor at the neuromuscular junction. Unlike the depolarizing muscle relaxants (such as succinylcholine), the non-depolarizers do not cause a contraction of the muscle prior to achieving relaxation. Examples include rocuronium, cis-atracurium, pancuronium and atracurium.

The use of NDMR must be monitored carefully using a peripheral nerve stimulator, to avoid excessive paralysis and inadequate return of strength at the end of the procedure, so-called “residual block”. The effect of NDMR is antagonized using anticholinesterase agents.

Related Glossary Terms
Acetylcholine, Anticholinesterase, Atracurium, Awareness, Cholinesterase, Cis-Atracurium, Competitive inhibitor, Extubation, Four A’s of anesthesia, Laudanosine, Myasthenia gravis, Neostigmine, Neuromuscular junction, Nicotinic, Pancuronium, Peripheral nerve stimulator, Residual block, Rocuronium
NPO stands for “nil per os” and indicates that the patient has no oral intake of fluids or solids.

Each department of anesthesia has their own policy dictating how long a patient must be “NPO” prior to surgery and anesthesia. Commonly-used guidelines are detailed below:

For elective surgery, patients should not have solid food for 8 hours prior to anesthesia. Unrestricted intake of clear fluids is permitted until 2-4 hours prior to scheduled surgery.

Guidelines for pediatric patients are more liberal than in the adult population. For example, infants may be allowed breast milk up to 4 hours pre-operatively and formula up to 6 hours pre-operatively.

For emergency surgery, the risk of aspiration is weighed against the risk of delaying surgery to achieve a longer “NPO” interval. It is important to realize that most patients undergoing emergency surgery will have a “full stomach” regardless of how long they have been NPO as the effects of trauma, pain, illness, opiates and other medications all serve to delay gastric emptying.

Related Glossary Terms
Aspiration, Clear fluids, Cricoid cartilage, Rapid sequence induction, Sellick’s maneuver

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Chapter 2 - Pre-operative Evaluation
Chapter 5 - Obstetrical Anesthesia
Chapter 5 - Pediatric Anesthesia
Ondansetron

**Class**
Serotonin (5-HT₃) antagonist. Clinical use is as an antiemetic for post-operative nausea and vomiting or for patients receiving chemotherapy.

**Mechanism of Action**
Ondansetron is a highly selective competitive antagonist of the serotonin receptor. It is believed to have its effect centrally, possibly in the area postrema of the brainstem where the chemoreceptive trigger zone is located.

**Dose**
Prophylaxis (adults): 4 mg IV prior to emergence.
Prophylaxis (children): 50-150 μg/kg IV
Treatment (adults): 1-2 mg IV

**Onset**
Less than 30 minutes

**Duration**
9 hours

**Elimination**
Hepatic (95%)

**Effects**

**CNS**
Headache

**CVS**
May cause cardiac rhythm or ECG changes by prolongation of the QT interval.

**GI**
Constipation, elevation of liver enzymes.

**Misc.**
Elimination of ondansetron is prolonged when given with other drugs metabolized by cytochrome P450 system.

**Related Glossary Terms**
Antiemetic agents, Chemoreceptor trigger zone, Dimenhydrinate, Opioids, Post-operative nausea and vomiting (PONV), Prochlorperazine

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Chapter 4 - Recovery
Chapter 6 - Drug Finder
Chapter 6 - Antiemetics
Opioids

Opioids are used to provide analgesia and act through agonism at the mu opiate receptor.

Opioids can be administered by almost any route. The common routes of administration in anesthetic practice are intravenous, epidural and intrathecal. In addition, oral opiates are often used as adjuncts along with patient-controlled analgesia (PCA).

For intravenous use, the synthetic opioids fentanyl, sufentanil and remifentanil are commonly used in modern anesthesia care. Morphine and hydromorphone are also used, most commonly for post-operative analgesia.

The most important side effect of opioids is their effect on the respiratory centre, the medulla oblongata. Opioids depress the drive to breathe by raising the apneic threshold for carbon dioxide. The effects of opioids can be reversed with naloxone.

Opioids are a major cause of post-operative nausea and vomiting.

Related Glossary Terms

Aldrete score, Alfentanil, Analgesia, Antiemetic agents, Apneic threshold, Balanced anesthesia, Cough reflex, Fentanyl, Fibreoptic bronchoscope, Four A's of anesthesia, Meperidine, Morphine Sulfate, Naloxone, Neonatal depression, Neonatal opioid depression, Ondansetron, Patient controlled analgesia, Re-narcotization, Remifentanil, Sufentanil
**Oxygen saturation**

The oxygen saturation is the percentage of hemoglobin that is bound with oxygen. Each hemoglobin molecule can carry four oxygen molecules. Oxygen saturation is determined by but not synonymous with PO2 (partial pressure of oxygen). The relationship between PO2 and oxygen saturation is described by the oxyhemoglobin dissociation curve.

Normal oxygen saturation is between 95-100%.

Oxygen saturation can be measured non-invasively with a pulse oximeter, while PO2 is measured by analyzing an arterial blood sample.

**Related Glossary Terms**

Hypoxemia, Oxyhemoglobin dissociation curve, Pulse oximeter
**Oxyhemoglobin dissociation curve**

The oxyhemoglobin dissociation curve describes the affinity of hemoglobin for oxygen as a function of partial pressure of oxygen. It is a sigmoidal curve. The curve can shift to the right with hyperthermia, hypercarbia, hypoxemia and elevated levels of 2,3 DPG. When shifted to the right, hemoglobin has less affinity for oxygen and therefore offloads oxygen more readily to the tissue.

**Related Glossary Terms**
- Hypoxemia
- Oxygen saturation
- Pre-oxygenation
- Pulse oximeter

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- Chapter 1 - Fluid Management
- Chapter 5 - Obstetrical Anesthesia
Oxytocin

Oxytocin is used in obstetrical care to augment uterine tone. It is used to stimulate labour. After delivery, it is used to minimize uterine atony and in so doing, decrease bleeding. It can cause hypotension if a bolus dose is delivered too quickly.

Related Glossary Terms

Uterine tone

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Chapter 5 - Obstetrical Anesthesia
Pancuronium

Class
Nondepolarizing skeletal muscle relaxant (NDMR); long-acting

Mechanism of Action
Competitive inhibitor at the acetylcholine receptors of the post-synaptic cleft of the neuromuscular junction.

Dose
Intubation: 0.1 mg/kg IV
Maintenance bolus: 0.01-0.03 mg/kg

Onset
4-5 minutes

Duration
45-65 minutes

Elimination
Renal (80%), hepatic (minor)

Effects
CVS
Pancuronium has a vagolytic effect and therefore causes tachycardia and hypertension. Increased risk of arrhythmias in patients receiving tricyclic antidepressants and volatile anesthetics.

Respiratory
May promote bronchospasm, salivation.

MSK
The neuromuscular blockade effects of non-depolarizing muscle relaxants are potentiated by succinylcholine, volatile anesthetics, aminoglycosides, lithium, loop diuretics, lidocaine, magnesium, lithium, ganglionic blockers, hypothermia, hypokalemia and respiratory acidosis. Enhanced neuromuscular blockade is seen in patients with myasthenia gravis or myopathies.

The effects of NDMR are antagonized by cholinesterase inhibitors. Increased resistance to NDMR is seen in patients on theophylline, burn patients and those with paresis or paralysis.

Misc.
Muscle relaxants are the most common cause of anaphylactoid reactions under general anesthesia.

Related Glossary Terms
Anticholinesterase, Atracurium, Cis-Atracurium, Neuromuscular junction, Non-depolarizing muscle relaxants, Peripheral nerve stimulator, Residual block, Rocuronium, Vagolytic
Parturient

The parturient is the pregnant patient who is in labour or will otherwise soon be delivering the baby.

Related Glossary Terms
Aortocaval compression, Epidural analgesia, Epidural anesthesia, Second stage of labour
Patency

Airway patency indicates that the airway is clear and free from obstruction. Causes of airway obstruction can be categorized broadly as follows:

a) Obstruction caused by normal tissue such as the tongue, tonsils, larynx and other soft tissue. Laryngospasm is an example of airway obstruction that occurs in the anesthesia setting.

b) Obstruction caused by pathology of normal tissue, such as that which occurs from infection, inflammation, tumour or trauma. Examples of infectious processes that can lead to airway obstruction include croup, epiglottitis and sublingual abscesses.

c) Obstruction caused by foreign body inhalation.

Related Glossary Terms
Airway obstruction, Aldrete score, Extubation, Laryngospasm, Upper airway

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Chapter 1 - Airway Management
Chapter 3 - General Anesthesia
Chapter 4 - Recovery
Chapter 5 - Pediatric Anesthesia
Patient controlled analgesia

Patient-controlled analgesia (PCA) permits the patient to control his own analgesia by activating a button, which then triggers the intravenous delivery of a predetermined dose of an opioid such as morphine or hydromorphone. Limits are set on the number of boluses per unit time as well as on the minimum time that must elapse between doses (lockout interval).

Complications of PCA include respiratory depression, nausea and vomiting, excessive sedation and inadequate analgesia.

Related Glossary Terms
Adjunct, Analgesia, Epidural analgesia, Morphine Sulfate

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Chapter 4 - Post-operative Pain Management
Chapter 5 - Anesthesia Outside the Operating Room
Pencil-point

Pencil point spinal needles have a blunt tip that is designed to spread rather than cut through the dural fibres. Their use is associated with a lower incidence of post-dural puncture headaches. Brand names include Pencan ® and Whitacre ®.

Related Glossary Terms
Intrathecal space, Spinal anesthesia, Subarachnoid space

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Find Term
Chapter 3 - Regional Anesthesia
Peripheral nerve stimulator

A peripheral nerve stimulator is used to assess neuromuscular transmission when neuromuscular blocking agents (NMBA) are used. The careful assessment of the depth of neuromuscular blockade allows the anesthesiologist to provide appropriate relaxation to facilitate surgery (when required) and to time the delivery of NMBA and their reversal agents to ensure the adequate return of muscle strength on emergence.

Typically, the peripheral nerve stimulator is used to deliver 4 successive electrical stimuli (2 per second) to a superficial peripheral nerve such as the median or facial nerve. The anesthesiologist will observe the number of twitches that the muscle produces and whether there is “fade” associated with those twitches. Fade is the successive decrease in amplitude of each twitch. The presence of less than four twitches or the presence of four twitches with fade provide evidence of residual neuromuscular blockade.

A patient could have 70% of his acetylcholine (Ach) receptors blocked and still register four equal twitches! It is for this reason that the limits of the peripheral nerve stimulator must be understood: it is specific but not sensitive for the presence of neuromuscular blockade. The most reliable indicators of the return of muscle strength are clinical ones: grip strength, head lift, ability to generate an adequate tidal volume and strong cough.

Related Glossary Terms
Anticholinesterase, Neuromuscular junction, Non-depolarizing muscle relaxants, Residual block

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Phenylephrine

Class
Sympathomimetic; vasopressor. Used in the treatment of hypotension.

Mechanism of Action
Direct agonist at the α-adrenergic receptor.

Dose
Bolus dose: 50-100 µg IV (adults)
Infusion: 0.1-1.0 µg/kg/minute

Onset
<1 minute

Duration
<5 minutes

Elimination
Re-uptake by tissue, liver and gut (monoamine oxidase)

Effects
CVS
Main effect is peripheral vasoconstriction, causing an increase in blood pressure. It is most appropriately used to raise the blood pressure in patients who are peripherally vasodilated (as a result of anesthesia, for example). It has the potential to cause myocardial ischemia, and left and right ventricular failure. It routinely causes a reflex bradycardia. If used in a patient in cardiogenic or hypovolemic shock, it may lead to a further reduction in vital organ blood flow.

Misc.
The clinician may observe diminished response of phenylephrine in patients receiving α-adrenergic blockers or drugs with α-blocking action such as phenothiazines. On the other hand, there may be augmented response when given with other vasopressors such as vasopressin and ergonovine. Phenylephrine has prolonged action in patients using monoamine oxidase inhibitors.

Related Glossary Terms
Ephedrine Sulfate, Epinephrine

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Chapter 6 - Drug Finder
Chapter 6 - Vasoactive Agents
Phrenic nerve

The phrenic nerve, arising from the third-fifth cervical nerve roots, controls the diaphragm. It is possible to block the phrenic nerve with a brachial plexus block when the supraclavicular or interscalene approach is used. For this reason (amongst others) bilateral brachial plexus blocks are contraindicated.

Related Glossary Terms
Brachial plexus block

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Chapter 3 - Regional Anesthesia
Pneumothorax

Pneumothorax indicates that air has entered the pleural space. If a pneumothorax is large enough, it may lead to a collapsed lung, as the negative intrapleural pressure that maintains lung distension is lost, allowing the lung to succumb to its natural elastic recoil.

Pneumothoraces have many causes. They can occur spontaneously or as a result of lung disease or trauma. They can arise as a complication of central line insertion or brachial plexus block if the needle punctures the pleural cavity.

Related Glossary Terms
Brachial plexus block

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Chapter 1 - Fluid Management
Chapter 3 - Regional Anesthesia
Chapter 3 - General Anesthesia
Chapter 6 - Inhaled Agents
Positive pressure ventilation

Positive pressure ventilation, often referred to as mechanical ventilation, is the process of achieving tidal volume through the application of positive pressure to the patient’s airway. It is most easily achieved through an endotracheal tube or tracheostomy tube.

Positive pressure ventilation can be achieved through bag mask ventilation or with a ventilator.

Most (but not all) patients under anesthesia receive positive pressure ventilation.

Related Glossary Terms

Bag mask ventilation, Hypoxemia, Mechanical ventilation, Tidal volume, Ventilator
Post-anesthetic care unit

The post-anesthetic care unit (PACU) is the unit to which the patient is transferred in the immediate post-operative period. Here, direct care is passed on from anesthesiologist to PACU nurse, although the anesthesiologist maintains responsibility for the patient. The PACU nurse is trained to monitor the patient and to recognize and manage the common post-operative problems and complications.

Related Glossary Terms
Aldrete score, Analgesia, Emergence, Post-operative nausea and vomiting (PONV), Recovery

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Chapter 3 - General Anesthesia
Chapter 4 - Recovery
Post-operative nausea and vomiting (PONV)

PONV is the most common post-operative complication.

The most important risk factors are female gender, non-smoking, past history of PONV or motion sickness, and the requirement of perioperative opioids. For each risk factor present, the risk of PONV goes up by 20%, that is 20% with one risk factor, 80% with four. Those with no risk factors have a risk of PONV of 10%.

Prevention involves includes modifying the anesthetic given to minimize the administration of emetogenic agents such as nitrous oxide and neostigmine. Prevention also includes the administration of prophylactic agents which should be directed towards the higher risk patients only. Commonly-used anti-emetics are ondansetron and dexamethasone. Ondansetron is much more effective as a treatment of PONV than as a preventive where its NNT (number needed to treat) appears to be around 5.

Related Glossary Terms
Aldrete score, Antiemetic agents, Chemoreceptor trigger zone, Dimenhydrinate, Ondansetron, Post-anesthetic care unit, Prochlorperazine, Recovery
Pre-medication refers to medication that the patient receives prior to surgery and can include medication that the patient takes routinely, as well as medication that may be prescribed specifically for the pre-operative period.

Examples of the latter category include:

a) antibiotics for prevention of wound infection or prevention of bacterial endocarditis

b) anxiolytics

c) aspiration prophylaxis (sodium citrate for the parturient)

d) corticosteroids for the adrenal-suppressed patient

Related Glossary Terms
Addisonian crisis, Adrenal suppression, Aspiration, Bacterial endocarditis, Pre-operative assessment, Sodium citrate

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Chapter 2 - Pre-operative Evaluation
Chapter 2 - Pre-operative Evaluation
Chapter 5 - Pediatric Anesthesia
Pre-operative assessment

Every patient undergoing anesthesia requires a careful pre-operative assessment. If the surgery is scheduled in advance and the patient is not (currently) a hospital in-patient, the assessment will occur in the pre-operative clinic. For patients who are already admitted to hospital, which includes those patients undergoing emergency surgery, assessment will take place at the bedside, occasionally (but not ideally) just prior to surgery. In any case, the goals of the assessment remain constant:

1) To review the medical and psychological status of the patient.

2) To identify factors which may impact on the peri-operative course, to take measures to optimize those factors and to delay surgery if necessary.

3) To inform patient, establish rapport, alleviate anxiety and obtain consent.

Related Glossary Terms

Airway assessment, ASA class, Clear fluids, Pre-medication, Pre-operative fluid deficit
Pre-operative fluid deficit

The pre-operative fluid deficit is determined by calculating the basal fluid requirement over the period of time that the patient has been fasting, and adding an estimation of other losses that may have occurred during the pre-operative period. Causes of losses would include bleeding, vomiting, diarrhea and nasogastric drainage.

Related Glossary Terms

Colloids, Crystalloid, Euvolemia, Maintenance fluid requirements, Pre-operative assessment, Shock, Third space loss
Pre-oxygenation

The purpose of pre-oxygenation is to lessen the risk of hypoxia occurring during the apneic period after induction. Proper and thorough pre-oxygenation is particularly important in patients who have a decreased functional residual capacity, such as the obese patient, the pediatric patient and the obstetric patient.

Pre-oxygenation works through denitrogenation of the functional residual capacity, creating a “reserve tank” of oxygen that will allow for gas exchange during apnea.

Pre-oxygenation is also key when a difficult intubation and/or difficult bag mask ventilation is encountered. In these situations, pre-oxygenation ensures maintenance of oxygenation for a longer period of time while the attempts at securing the airway are made.

Related Glossary Terms
Difficult airway, Functional residual capacity, Hypoxemia, Induction, Oxyhemoglobin dissociation curve

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Chapter 5 - Obstetrical Anesthesia
Chapter 5 - Pediatric Anesthesia
Prochlorperazine

Class
Although it has several uses, in anesthetic practice it is used as an antiemetic for post-operative nausea and vomiting (PONV).

Mechanism of Action
Central inhibition of the dopamine D₂ receptors in the medullary chemoreceptor trigger zone. Prochlorperazine also inhibits the vagus nerve in the gastrointestinal tract. The anticholinergic, sedative and antihistaminic effects of prochlorperazine also contribute to its antiemetic action.

Dose
2.5-10 mg IV, max. 40 mg/day (adults)

Onset
10-20 minutes

Duration
3-4 hours

Elimination
Enterohepatic

Effects
Prochlorperazine has anticholinergic properties which are additive to the anticholinergic effects of other drugs. As a phenothiazine, it also has the potential to cause extrapyramidal symptoms.

CNS
Sedative effects which are additive to other-hypnotics> May cause extra-pyramidal syndromes (motor restlessness, oculogyric crisis, opisthotonus, dystonias), especially in young male patients.

CVS

Misc.
Diminishes effects of anticoagulants. Possible hyperthermia in the presence of hypothalamic dysfunction. Neuroleptic malignant syndrome.

Related Glossary Terms
Antiemetic agents, Chemoreceptor trigger zone, Dimenhydrinate, Ondansetron, Post-operative nausea and vomiting (PONV)
Prone

In the prone position, the patient is lying on his or her front. It is used for spinal surgery, posterior fossa neurosurgical surgery and for surgery for pilonidal abscesses amongst others.

Special operating room tables and bolsters are used to increase the safety of the prone position, but in all cases, care must be taken to avoid its many potential complications. The prone position can compromise respiratory function, venous return as well as causing ocular complications.

Related Glossary Terms

Lithotomy, Neuropathy, Supine

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Chapter 1 - Airway Management
Chapter 3 - General Anesthesia
Propofol

**Class**
Alkylphenol intravenous anesthetic agent. Used for induction of general anesthesia. Can also be used for maintenance of anesthesia or for sedation, in each case by continuous infusion.

**Mechanism of action**
Not well described.

**Dose**
- Induction: 2-2.5 mg/kg IV for adults
- Induction: 3-4 mg/kg IV for children
- Maintenance of anesthesia: 100-200 ug/kg/minute
- Sedation: 40-100 ug/kg/minute

**Onset**
Within one arm-brain circulation time (approximately 20 seconds).

**Duration**
Approximately 5-8 minutes after single induction dose.

**Elimination**
Rapid redistribution away from central nervous system (CNS) into lean body compartment accounts for prompt awakening. Metabolized by liver and extra-hepatic sites then excreted by kidney.

**Effects**

**CNS**
Profound CNS depressant, potentiating the depressant effects of opioids, sedatives and volatile anesthetics. Decreases cerebral metabolic rate and intracranial pressure. Occasionally excitement, tonic-clonic movements or opisthotonus is seen on induction with propofol.

**CVS**
Causes direct myocardial depression and vasodilation leading to hypotension. Propofol must be used with caution in patients with poor left ventricular function or critical coronary artery insufficiency or in those who are seriously ill or debilitated.

**Respiratory**
Depression of respiratory centre leads to brief apnea. Propofol effectively blunts the airway’s response to manipulation thus hiccoughing and bronchospasm are rarely seen.

**Misc.**
Pain on injection seen in up to 20%. Mild anti-emetic properties. Patients often experience pleasant dreams under anesthesia followed by a smooth, clear-headed emergence. Strict aseptic technique must be used when handling propofol as the vehicle is capable of supporting rapid growth of micro-organisms.

**Contraindications**
Egg or soy allergy.

**Related Glossary Terms**
Continuous Infusion, Etomidate, Induction, Intralipid, Ketamine, Sodium Thiopental, Total intravenous anesthesia
Pseudocholinesterase deficiency

Pseudocholinesterase deficiency is an inheritable enzyme deficiency of the enzyme that breaks down succinylcholine. Those affected will demonstrate prolonged effect of succinylcholine.

Deficiency can result as a genetic defect, as a consequence of various medications or a result of liver disease. The latter two causes are usually relative while the genetic defect can produce a complete lack of pseudocholinesterase function in homozygous individuals. Individuals who are heterozygous have reduced (but still measurable) pseudocholinesterase activity.

Related Glossary Terms
Succinylcholine
Pulmonary artery catheter

A pulmonary artery catheter is a catheter that is fed from the internal jugular vein (or subclavian vein) through the right side of the heart into the pulmonary artery. It is used to estimate the preload of the left side of the heart. It can also be used to measure cardiac output by the thermodilution technique. It can be useful in determining the cause of shock in critically ill patients. Unfortunately, studies have failed to show an improvement in outcomes as a result of this diagnostic tool.
Pulse oximeter

The oxygen saturation monitor non-invasively measures the oxygen saturation of hemoglobin in the patient’s arterial blood. It is also known as a pulse oximeter.

The pulse-ox is placed on a finger, toe or earlobe. Two different wavelengths of light (one red, one infrared) are passed through the patient’s tissue from one side of the pulse-oximeter to a photodetector on the other side. Absorption of light at these wavelengths differs significantly between oxyhemoglobin and its deoxygenated form. Accordingly, the percentage of oxygenated hemoglobin can be calculated from the ratio of the absorption of these two wavelengths of light. By searching for a pulsatile signal, the device can measure the saturation in arterial (not venous) blood.

There are many factors that can affect the ability of the pulse-oximeter to give an accurate reading. Good peripheral blood flow is required which can be impeded in a cold or shocked patient. Measurements can be erroneous when nail polish is used as well as in the presence of carbon monoxide, cyanide and methemoglobin.

Related Glossary Terms
Hypoxemia, Oxygen saturation, Oxyhemoglobin dissociation curve, Shock

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Chapter 2 - Anesthetic Equipment and Monitoring
Pulse pressure

Pulse pressure is the difference between systolic and diastolic blood pressure. It is affected by several factors, including intravascular volume and systolic vascular resistance (SVR).

For example, a high SVR will narrow the difference between systolic and diastolic blood pressure, and therefore lower the pulse pressure. A low SVR will lower the diastolic blood pressure (disproportionately to the systolic) and will cause a widened pulse pressure.

Imagine two unfortunate patients in the emergency room, each involved in a motor vehicle accident, each with a systolic blood pressure of 110 mmHg. Patient A, who has a ruptured spleen and is in the early stages of hypovolemic shock, will have a narrow pulse pressure (BP 110/75), indicating that the sympathetic nervous system has been activated and that his “normal” blood pressure is maintained due to high systemic vascular resistance.

Patient B has suffered a cervical spine injury and has a BP of 110/35, his wide pulse pressure betraying the loss of sympathetic tone that has resulted from disruption of sympathetic outflow, which exits the spinal column from T1-L2 to form the sympathetic chain.

Related Glossary Terms

Shock, Sympathetic nervous system
Radiculopathies

Radiculopathy describes the consequence of nerve root damage or irritation, of any cause. It can comprise both sensory and motor abnormalities.

Related Glossary Terms
Neuropathy

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Chapter 3 - Regional Anesthesia
Rapid sequence induction

In patients deemed to be at increased risk for aspiration, the time between inducing anesthesia and securing the airway with a cuffed endotracheal tube must be minimized. Such a technique is termed a “rapid sequence induction”.

A rapid sequence induction is performed as follows:

1. Suction apparatus is checked and kept readily available.
2. Pre-oxygenation of patient with 100% oxygen for 3-5 minutes.
3. Application of cricoid pressure (Sellick’s maneuver) by assistant.
4. Induction with pre-calculated dose of induction agent followed immediately by intubating dose of depolarizing muscle relaxant (succinylcholine). A rapidly acting non-depolarizing agent (e.g. rocuronium) is commonly substituted in a so-called “modified” rapid sequence induction.
5. Intubation of trachea, cuff inflation and verification of proper tube position.

Related Glossary Terms

Aspiration, Clear fluids, Cricoid cartilage, Induction, Intubation, Rocuronium, Sellick’s maneuver, Stylet, Succinylcholine
**Re-narcotization**

A narcotized patient is one who is exhibiting the respiratory-depressant effects of opioids. When a naloxone has been used to reverse the effects of opioids, the patient may become “re-narcotized” after an initial period of response. The reason for this “re-narcotization” is that the duration of effect of naloxone is shorter than that of many of the opioids in use. Therefore, one must continue to monitor the patient who has received naloxone rather than be falsely reassured by the initial (immediate) response.

**Related Glossary Terms**

Morphine Sulfate, Naloxone, Neonatal opioid depression, Opioids
Recovery

Phase 1 of recovery must be achieved before the patient is discharged from the post-anesthetic care unit (PACU) to the ward or short-stay unit. The patient must demonstrate adequate airway control, ventilation, circulation, colour, level of consciousness and activity. The Aldrete score is a detailed scoring system that rates the patient’s status from 0-2 on each of those five criteria.

For Phase 1 recovery to be complete, the patient must:

- be showing no signs of respiratory depression for at least 20-30 minutes after last dose of parenteral opioids
- be easily aroused
- be fully oriented to person, place and time
- be able to maintain and protect the airway on his own, with evidence of a strong cough
- have stable vital signs for at least 30 minutes

It is also important that pain, nausea and vomiting are controlled prior to PACU discharge and that there are no ongoing surgical concerns, such as surgical site bleeding.

Phase 2 recovery focuses on the necessary criteria that must be met before the patient is returned home and requires the return of cognitive function, ambulation and the ability to take oral liquids, to name a few.

Related Glossary Terms
Aldrete score, Analgesia, Emergence, Post-anesthetic care unit, Post-operative nausea and vomiting (PONV)

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Chapter 2 - Pre-operative Evaluation
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Chapter 4 - Recovery
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Recurrent laryngeal nerve

The recurrent laryngeal nerve is a branch of the vagus nerve (cranial nerve X). It supplies all the laryngeal muscles, with the exception of one. The exception is the cricothyroid muscle, an adductor muscle, which is supplied by the external branch of the superior laryngeal nerve.

Related Glossary Terms
Larynx, Superior laryngeal nerve, Vagus nerve

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Chapter 1 - Airway Management
Chapter 3 - Regional Anesthesia
Regional anesthesia

Regional anesthesia is the anesthetic technique which relies on the blockade of a nerve or group of nerves to render the surgical field insensate. Regional anesthesia can be used on its own or in combination with sedation or general anesthesia.

Related Glossary Terms
Analgesia, Asepsis, Bier Block, Brachial plexus block, Epidural anesthesia, Neuropathy, Spinal anesthesia

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Chapter 3 - Anesthetic Techniques
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Chapter 5 - Obstetrical Anesthesia
Chapter 6 - Local Anesthetics
Chapter 6 - Local Anesthetics
Remifentanil

**Class**
Synthetic opioid analgesic (ultra short-acting); adjunct to anesthesia.

**Mechanism of Action**
Acts at the mu-and kappa opioid receptors.

**Dose**
On induction of general anesthesia: 0.3-1 μg/kg
For maintenance of general anesthesia: 0.1-1 μg/kg/minute (by infusion)
For sedation: infusion 0.05 – 0.1 μg/kg/minute

**Onset**
After single bolus: 1-1.5 minutes
After initiation of infusion: 3-5 minutes

**Duration**
5-10 minutes; context sensitive half time 3 minutes

**Elimination**
Non-specific blood-tissue esterases (end-organ independent)

**Effects**

**CNS**
Potent analgesic effects, sedation. “MAC sparing” allows up to 75% reduction in dose of co-anesthetics. All of the depressant effects of remifentanil are potentiated by concurrent use of sedatives, volatile anesthetics and nitrous oxide.

**CVS**
Exaggerated bradycardia, hypotension (compared with other opioids). The synthetic opioids are not direct myocardial depressants but they do reduce sympathetic drive, which may result in decreased cardiac output in patients who are relying on sympathetic tone to support their circulation, such as those in hypovolemic or cardiogenic shock.

**Respiratory**
Profound respiratory depressant which often leads to apnea.

**GI**
Nausea, vomiting.

**Misc.**
Can cause profound muscle rigidity. Not suitable for spinal or epidural use due to glycine additive. Rapid elimination requires initiation of post-operative analgesia (usually morphine) prior to emergence.

**Related Glossary Terms**
Alfentanil, Analgesia, Apneic threshold, Balanced anesthesia, Continuous Infusion, Fentanyl, Four A's of anesthesia, Morphine Sulfate, Naloxone, Opioids, Sufentanil, Total intravenous anesthesia
Residual block

Residual block (or residual paralysis) refers to the patient who displays the ongoing effects of non-depolarizing muscle relaxants (NDMR) on or after emergence. It can occur when excessive doses of NDMR are given intraoperatively, or if appropriate antagonism (with anticholinesterases) has not occurred.

Some factors predispose to residual block and include an unexpectedly short duration of surgery or inadequate renal or hepatic function required to metabolize the NDMR.

The use of a peripheral nerve stimulator as well as clinical examination of the patient on emergence should identify a residual block. If possible, an additional dose of anticholinesterase can be given. If the residual block is significant and/or the maximum dose of anticholinesterase has been given, then the patient should remain intubated, with appropriate sedation, until normal muscle strength returns.

Extubating a patient with a residual block puts the patient at risk of airway obstruction, laryngospasm, aspiration and inadequate ventilation.

Related Glossary Terms
Airway obstruction, Anticholinesterase, Cholinesterase, Emergence, Extubation, Laryngospasm, Non-depolarizing muscle relaxants, Peripheral nerve stimulator
Reticular activating system

The reticular activating system is the part of the brain that governs consciousness.

Related Glossary Terms
Awareness, Extubation, Four A's of anesthesia

Index
Chapter 6 - Induction Agents
Right mainstem bronchus

The trachea splits into right and left mainstem bronchi. The right mainstem bronchus has a more vertical orientation which explains why aspirated material most commonly affects the right lung.

If the endotracheal tube is advanced too far on intubation, it will pass into the right mainstem bronchus and only one lung (the right) will be ventilated. Careful auscultation will reveal the absence of breath sounds on the left.

Related Glossary Terms
Aspiration, Intubation, Lower airway
Rocuronium

**Class**
Non-depolarizing muscle relaxant (NDMR); short-acting

**Mechanism of Action**
Competitive inhibitor at the acetylcholine receptors of the post-synaptic cleft of the neuromuscular junction.

**Dose**
Intubation: 0.45-0.9 mg/kg
Maintenance bolus 0.1-0.2 mg/kg

**Onset**
Dose-dependent:
1-1.5 minutes (0.6 mg/kg)
0.5-1.0 minutes (0.9 mg/kg)
Higher dose is therefore suitable for rapid sequence induction.

**Duration**
Dose-dependent:
31 minutes (0.6 mg/kg)
60 minutes (0.9 mg/kg)

**Elimination**
Hepato-biliary (70%); renal (10%)

**Effect**
CVS
Very weak vagolytic effect.

MSK
The neuromuscular blockade effects of non-depolarizing muscle relaxants are potentiated by succinylcholine, volatile anesthetics, aminoglycosides, lithium, loop diuretics, lidocaine, magnesium, lithium, ganglionic blockers, hypothermia, hypokalemia and respiratory acidosis.

Enhanced neuromuscular blockade is seen in patients with myasthenia gravis or myopathies.

The effects of NDMR are antagonized by cholinesterase inhibitors. Increased resistance to NDMRs is seen in patients on theophylline, burn patients and those with paresis or paralysis.

Misc.
Muscle relaxants are the most common cause of anaphylactoid reactions under general anesthesia.

**Related Glossary Terms**
Anticholinesterase, Atracurium, Cis-Atracurium, Neuromuscular junction, Non-depolarizing muscle relaxants, Pancuronium, Peripheral nerve stimulator, Rapid sequence induction, Residual block
Ryanodine

Up to 70% of Malignant Hyperthermia cases involve a mutation in the gene encoding the ryanodine receptor protein on chromosome 19. This particular mutation shows autosomal dominant inheritance pattern, with variable penetrance.

Related Glossary Terms
Dantrolene, Malignant hyperthermia, Myoglobinuria, Succinylcholine
Second stage of labour

The second stage of labour is the stage where the cervix is fully dilated and the woman is pushing. It ends when the baby is out. The duration of the second stage can be just a few minutes or up to several hours. It is longer in the primiparous female and also will generally be longer if epidural analgesia is being used.

Related Glossary Terms

Epidural analgesia, Parturient

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Chapter 5 - Obstetrical Anesthesia
Sellick's maneuver

Sellick's maneuver is the application of cricoid pressure during the induction of a patient who is at risk of aspiration. The maneuver aims to use the cricoid cartilage, the only circumferential cartilage of the airway, to temporarily occlude the esophagus to prevent passive regurgitation of stomach contents while the airway is unprotected.

Related Glossary Terms
Aspiration, Cricoid cartilage, Intubation, Rapid sequence induction, Subglottis

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Chapter 3 - General Anesthesia
Serotonin syndrome

Serotonin syndrome is a serious, potential fatal condition caused by excessive action of the serotonin neurotransmitter in the central nervous system usually by increased action of serotonin re-uptake inhibitors (SSRIs), a common class of anti-depressants.

Augmented effects of SSRI’s can occur through decreased clearance of the drug, drug interactions or through increased serum levels of precursors such as tryptophan.

Symptoms and signs of serotonin syndrome include confusion, hallucinations, agitation, coma, rigidity, tremors, hyper-reflexia, autonomic instability and hyperthermia.

Related Glossary Terms
Drag related terms here

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Chapter 6 - Vasoactive Agents
Sevoflurane

Class
Volatile inhaled anesthetic. Used for maintenance of anesthesia. Can be used for induction of anesthesia particularly in children. Rarely may be used as a treatment for status asthmaticus.

Mechanism of Action
Uncertain

Dose
Titrated to effect; MAC (age 40) = 2.1%.

Onset
Low solubility allows rapid uptake and equilibration. Onset of effect is hastened by using higher flows of carrier gases and by using higher concentrations of volatile agent.

Duration
Clinical recovery in less than 10 minutes (usually). If given for prolonged periods, wake-up will be slower as adipose stores have been saturated and are slow to off-load.

Elimination
Pulmonary (major); hepatic (2-5%); renal (metabolites excretion only)

Effects
CNS
Sevoflurane produces an additive central nervous system (CNS)-depressant effect along with other sedative/hypnotics and analgesics. Has the potential to increase intracranial pressure which can be mitigated with hyperventilation. Delirium.

CVS
Dose-related hypotension (vasodilation).

Respiratory
Respiratory depression with a rapid, shallow respiratory pattern. Loss of intercostal muscle function creates a rocking boat appearance. Causes bronchodilation. Sevoflurane is sweet-smelling and not as irritating to the respiratory tract as desflurane.

GI
Nausea, vomiting.

MSK
Potentiates neuromuscular blockade. Malignant hyperthermia trigger.

Misc.
Potential nephrotoxicity due to Compound A which is produced through contact with sodalime. Compound A can be produced if sevoflurane is used with very low fresh gas flows or for long MAC-hours. Therefore, sevoflurane must be used with a minimum of 2 litres/minute of fresh gas flow.

Related Glossary Terms
Balanced anesthesia, Desflurane, Four A's of anesthesia, Inhaled agents, Isoflurane, MAC, Malignant hyperthermia, Vapourizer, Volatile
Shock

Shock occurs when perfusion of oxygenated blood to the tissues is inadequate to support vital organ function. Categories of shock are hypovolemic, cardiogenic, neurogenic and distributive.

Hypovolemic shock is inadequate organ perfusion that results from decreased circulatory volume, which can result from dehydration or blood loss.

Related Glossary Terms
Autonomic nervous system, Blood products, Colloids, Crystalloid, Epinephrine, Etomidate, Euvolemia, Ketamine, Maintenance fluid requirements, Metabolic acidosis, Pre-operative fluid deficit, Pulmonary artery catheter, Pulse oximeter, Pulse pressure, Sympathetic nervous system, Third space loss, Venous air embolism
Sniffing position

The sniffing position is achieved when the lower cervical spine is flexed and the upper cervical spine is extended. It is believed to achieve the best alignment of the tracheal, laryngeal and oropharyngeal axes. Simply put, the sniffing position facilitates the anesthesiologist’s ability to visualize the larynx by direct laryngoscopy. Careful use of pillows (or of a specially-designed “Troop pillow®”) will aid in achieving a proper sniffing position.

Related Glossary Terms
Difficult airway, Direct laryngoscopy, Fibreoptic bronchoscope, Intubation, Neck motion
Sodium citrate

Sodium citrate is a non-particulate antacid commonly used to reduce the consequences (but not the occurrence) of aspiration in the pregnant patient.

Related Glossary Terms
Aspiration, Pre-medication
Sodium Thiopental

**Class**
Short-acting barbiturate. Was used as an anesthetic induction agent but has largely been replaced by propofol. It is also useful as an anticonvulsant or for the rapid reduction of elevated intracranial pressure.

**Mechanism of action**
Decreases the rate of dissociation of the inhibitory neurotransmitter GABA from its receptors resulting in depression of the reticular activating system.

**Dose**
- 3-5 mg/kg IV for healthy adults
- 5-6 mg/kg IV for children
- 7-8 mg/kg IV for infants

Dose must be reduced considerably in unstable or fragile patients.

**Onset**
Within one arm-brain circulation time (approximately 20 seconds).

**Duration**
Approximately 5-10 minutes after single induction dose.

**Elimination**
Rapid redistribution of drug from the central nervous system (CNS) to lean body tissue accounts for the prompt awakening. The final elimination from the body depends on hepatic metabolism and excretion by the kidneys.

**Effects**

**CNS**
Profound CNS depressant. Decreases cerebral metabolic rate and intracranial pressure. May cause hypertonus, twitching and tremors during induction. May contribute to postoperative confusion and delirium. Potentiates the depressant effects of opioids, sedatives, alcohol and volatile anesthetics.

**CVS**
Depression of myocardial contractility and vasodilation leads to decreased cardiac output and blood pressure with a mild compensatory tachycardia. Must be used with caution in patients with poor left ventricular function or critical coronary artery insufficiency or in those who are seriously ill or debilitated.

**Respiratory**
Depresses the rate and depth of breathing leading to brief period of apnea. Does not blunt the airway’s response to manipulation therefore coughing, hiccoughing, laryngospasm and bronchospasm may be seen at light planes of anesthesia.

**GI**
Nausea and vomiting

**Misc.**
Incompatible with drugs with acidic pH. For example, if given in the IV line with vecuronium (a NDMR no longer in use), precipitation would occur. Arterial or extravascular injection produces necrosis.

**Contraindications**
Porphyria

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**Related Glossary Terms**
Etomidate, Induction, Ketamine, Propofol

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Spinal anesthesia

Spinal anesthesia is the technique where a small-calibre needle is placed in the intrathecal space, through which a small volume of local anesthetic is delivered. Spinal anesthesia provides up to 3 1/2 hours of surgical anesthesia and is most commonly used for lower limb surgery, inguinal hernia repair, genitourinary procedures, Caesarian section and some gynecological procedures.

Any procedure for which pain sensation is restricted to T4 (nipple line) or lower can be performed under spinal anesthesia. Any intra-peritoneal procedure requires a T4 block, even if the incision is much lower, because the peritoneal innervation is transmitted at that level. Blocking the spinal cord above that level is undesirable, as the neural control of the heart and muscles of respiration begin to be compromised. From a practical perspective, abdominal procedures are rarely performed under spinal anesthesia because the precise level of block achieved by a given dose is difficult to predict. The notable exception is Caesarian sections, for which spinal anesthesia is the preferred technique.

Related Glossary Terms
Baricity, Bupivacaine, Dura, Epidural anesthesia, Epidural space, Intrathecal space, Lido-caine, Morphine Sulfate, Pencil-point, Regional anesthesia, Subarachnoid space, Sympathetic nervous system
A stylet is a firm, bendable device that is placed inside the lumen of endotracheal tube prior to intubation. It allows the anesthesiologist to make a bend in the endotracheal tube to facilitate intubation.

Just after the tube is passed through the laryngeal inlet, the stylet is pulled out (by an assistant) whereupon the anesthesiologist advances the tube into its final position in the trachea. The anesthesiologist must hold the tube firmly as the stylet is pulled out, to avoid the displacement of the endotracheal tube.

The stylet has the potential to cause trauma to the airway and must never be positioned beyond the tip of the endotracheal tube. Furthermore, the stylet should be removed as soon as the tip of the tube has passed the glottis. One is not meant to pass the styletted tube into the mid-trachea.

A stylet is not required for most intubations; the use of a stylet should not be a compensation for improper intubation technique.

Related Glossary Terms
Adjunct, Difficult airway, Direct laryngoscopy, Intubation, Laryngoscope, Macintosh blade, Magill blade, Rapid sequence induction, Sniffing position
Subarachnoid space

The subarachnoid space contains the spinal cord, cauda equina and cerebrospinal fluid. Spinal anesthesia is performed by injecting a small volume of local anesthetic into the subarachnoid (or “intrathecal” space).

Related Glossary Terms
Baricity, Dura, Intrathecal space, Pencil-point, Spinal anesthesia
Subglottis

The subglottis is the region just below the larynx. It is located at the level of the cricoid cartilage. In adults, this is at the level of the 6th cervical vertebrae; in children, it is higher, at C4. The subglottis is the narrowest region of the pediatric airway. The pediatric patient, therefore, is susceptible to airway obstruction at this level, when the subglottic region is swollen either through viral inflammation (croup) or traumatic airway maneuvers.

Related Glossary Terms
Airway obstruction, Aspiration, Cricoid cartilage, Larynx, Lower airway, Needle cricothyrotomy, Sellick’s maneuver, Upper airway

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Chapter 5 - Pediatric Anesthesia
**Succinylcholine**

**Class**
Depolarizing muscle relaxant; ultra short-acting; Used for rapid sequence induction.

**Mechanism of Action**
Succinylcholine (Sch) attaches to nicotinic cholinergic receptors at the neuromuscular junction. There, it mimics the action of acetylcholine thus depolarizing the post-junctional membrane. Neuromuscular blockade (paralysis) develops because a depolarized post-junctional membrane cannot respond to subsequent release of acetylcholine.

**Dose**
- Intubation: 1.0-1.5 mg/kg IV or 2.5-4 mg/kg IM
- Onset: 30-60 seconds after IV administration
- Duration: 2-3 minutes after IM dose

**Onset**
- Duration: is 4-6 minutes after IV dose
- Duration: 10-30 minutes after IM dose

**Elimination**
Hydrolysis by plasma pseudocholinesterase

**Dose**
Intubation: 1-1.5 mg/kg IV or 2.5-4 mg/kg IM

**Onset**
30-60 seconds after IV administration
2-3 minutes after IM dose

**Duration**
Duration: 4-6 minutes after IV dose
Duration: 10-30 minutes after IM dose

**Effects**

**CNS**
Raised intracranial pressure and raised intraocular pressure.

**CVS**
Because of cross-reactivity at the muscarinic acetylcholine receptors, Sch causes vagal cardiac dysrhythmias. Bradycardia, junctional rhythm and sinus arrest can occur particularly if a second dose is administered and particularly in children.

**Respiratory**
Occasionally leads to bronchospasm and excessive salivation due to muscarinic effects. Intragastardic pressure is increased thereby theoretically increasing the risk of regurgitation.

**Mac.**
Most of the other effects are secondary to the depolarization and subsequent contraction of skeletal muscle. Sch elevates serum potassium 0.3-0.5 mEq/L in normal patients. It can cause an exaggerated release of potassium (leading to fatal hyperkalemia) in those with neuromuscular or muscle disease. Post-operative myalgia is common particularly in young adults. Succinylcholine is a potent trigger of malignant hyperthermia.

**Contraindications**
There is a long list of absolute and relative contraindications which can be found in any Anesthesia text. A brief summary follows:

- Malignant Hyperthermia (MH) or presence of conditions associated with MH.
- Pseudocholinesterase deficiency: Deficiency can result as a genetic defect, as a consequence of various medications or as a result of liver disease. The latter two causes are usually relative while the genetic defect can produce a complete lack of pseudocholinesterase activity in homozygous individuals. The use of succinylcholine in a patient with pseudocholinesterase deficiency leads to prolonged paralysis.
- Hyperkalemia.
- Presence of neurologic or muscular condition which would predispose to hyperkalemia after Sch-induced muscle contraction. Examples include recent paralysis (spinal cord injury or stroke), amyotrophic lateral sclerosis (ALS), Duchenne’s muscular dystrophy and recent burns or crush injury. Myotonia congenita or myotonia dystrophica can manifest sustained contraction with Sch.

**Related Glossary Terms**
Acetylcholine, Dantrolene, Malignant hyperthermia, Myoglobinuria, Neuromuscular junction, Non-depolarizing muscle relaxants, Pseudocholinesterase deficiency, Rapid sequence induction, Ryanodine
Sufentanil

Class
Synthetic opioid analgesic (intermediate-acting), adjunct to anesthesia.

Mechanism of Action
Acts at the mu- and kappa opioid receptors.

Dose
General anesthesia: 0.3-1 ug/kg IV, depending on patient condition, other agents used, nature and duration of surgery.
Infusion dose: 0.3-1 ug/kg/hour

Onset
1-2 minutes

Duration
20-40 minutes

Elimination
Hepatic

Effects
CNS
Potent analgesic properties and some sedative effect. All of the depressant effects of sufentanil are potentiated by concurrent use of sedatives, volatile anesthetics and nitrous oxide.

CVS
Bradycardia, hypotension. The synthetic opioids are not direct myocardial depressants but they do reduce sympathetic drive which may result in decreased cardiac output in patients who are relying on sympathetic tone to support their circulation, such as those in hypovolemic or cardiogenic shock.

Respiratory
Respiratory depression, which at the extreme leads to apnea.

GI
Nausea, vomiting, biliary tract spasm, constipation.

Misc.
Muscle rigidity

Related Glossary Terms
Analgesia, Apneic threshold, Balanced anesthesia, Fentanyl, Four A’s of anesthesia, Morphine Sulfate, Naloxone, Opioids, Remifentanil

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Superior laryngeal nerve

The superior laryngeal nerve supplies the only muscle of the larynx which is not supplied by the recurrent laryngeal nerve: the cricothyroid muscle, an adductor muscle of the larynx.

Like the recurrent laryngeal nerve, the superior laryngeal nerve is a branch of the vagus nerve.

Related Glossary Terms
Larynx, Recurrent laryngeal nerve, Vagus nerve

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Chapter 1 - Airway Management
**Supine**

Supine describes the position where the patient is lying on his or her back.

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**Related Glossary Terms**

Lithotomy, Prone, Trendelenburg

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Chapter 3 - General Anesthesia
Chapter 5 - Obstetrical Anesthesia
Sympathetic nervous system

The sympathetic system is mediated neurally, through the sympathetic chain, and hormonally, through the adrenal gland. Epinephrine and norepinephrine are the neurotransmitters of the sympathetic nervous system in contrast to Ach which is the messenger of the parasympathetic nervous system.

From the neural perspective, the sympathetic chain is comprised of presynaptic nerves exiting the spinal cord through T1-L2; therefore, epidural or spinal blockade that involves any or all of those levels will reduce sympathetic outflow. The sympathetic outflow to the heart is at the level of T1-T4 and causes positive inotropy and chronotropy. The heart is not the only way the sympathetic nervous system effects the cardiovascular system: sympathetic innervation along the entire distribution (T1-L2) controls the vascular tone of the blood vessels.

From the humoral perspective, the adrenal gland, when stimulated, exerts its effect by releasing epinephrine and norepinephrine into the bloodstream.

Related Glossary Terms
Autonomic nervous system, Ephedrine Sulfate, Epidural analgesia, Epidural anesthesia, Epinephrine, Four A's of anesthesia, Pulse pressure, Shock, Spinal anesthesia, Vagus nerve

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Tachypnea

Increased respiratory rate

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Drag related terms here

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Chapter 5 - Malignant Hyperthermia
Theophylline

Theophylline is a medication that is used occasionally in the treatment of asthma or chronic obstructive pulmonary disease (COPD). Theophylline can be administered orally or intravenously. Patients on theophylline are resistant to the effects of non-depolarizing muscle relaxants.

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Third space loss

Third space loss refers to the loss of plasma fluid into the interstitial space as a result of tissue trauma and can be estimated based on the nature of the surgery:

- 2-5 cc/kg/hr for minimal surgical trauma (orthopedic surgery)
- 5-10 cc/kg/hr for moderate surgical trauma (bowel resection)
- 10-15 cc/kg/hr for major surgical trauma (abdominal aortic aneurysm repair)

Related Glossary Terms

Colloids, Crystalloid, Euvolemia, Maintenance fluid requirements, Pre-operative fluid deficit, Shock

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Chapter 5 - Pediatric Anesthesia
Thrombocytopenia

Low platelet count

Related Glossary Terms

Blood products

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Chapter 1 - Fluid Management
Thyromental Distance

The distance from the lower border of the mandible to the thyroid notch with the neck fully extended should be at least three to four fingerbreadths. A shorter distance may indicate that the oral-pharyngeal-laryngeal axis will be too acute to achieve good visualization of the larynx with direct laryngoscopy.

Related Glossary Terms
Airway assessment, Difficult airway, Direct laryngoscopy, Intubation

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Chapter 1 - Airway Management
Tidal volume

Tidal volume is the volume of a normal breath. In the context of a patient who is being mechanically ventilated, the tidal volume is set by the anesthesiologist.

Related Glossary Terms
Extubation, Functional residual capacity, Mechanical ventilation, Positive pressure ventilation

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Chapter 3 - General Anesthesia
Total intravenous anesthesia

Total intravenous anesthesia (TIVA) is a technique whereby the goals of general anesthesia are attained with the use of intravenous agents alone, usually by continuous infusion. Propofol and remifentanil are commonly-used during TIVA.

The patient would be ventilated with air and oxygen, but no inhalational agents.

Related Glossary Terms
Balanced anesthesia, Continuous Infusion, Four A's of anesthesia, Maintenance, Propofol, Remifentanil
Transcutaneous electrical nerve stimulation

Transcutaneous electrical nerve stimulation (TENS) is the use of an electrical stimulus, applied to the skin, for therapeutic purposes. It can be used to provide analgesia during labour.

Related Glossary Terms
Drag related terms here

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Chapter 5 - Obstetrical Anesthesia
Transesophageal echocardiography

Transesophageal echocardiography (TEE) is performed by the anesthesiologist during certain types of cardiac surgery. It can also be used to help determine the cause of cardiovascular collapse in other (non-cardiac) surgical patients.

Related Glossary Terms

Drag related terms here

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Chapter 2 - Anesthetic Equipment and Monitoring
**Trendelenburg**

The trendelenburg position indicates a head-down tilt. It can be used when the patient is in the supine or lithotomy positions to facilitate surgical exposure. It is commonly used during gynecologic surgery.

The trendelenburg position increases venous return to the heart. As well, it decreases lung compliance due to upwards pressure (from the abdominal contents) on the diaphragm.

**Related Glossary Terms**

Lithotomy, Supine

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Chapter 3 - General Anesthesia
Upper airway

The upper airway ends at the level of the larynx or glottic opening. Therefore, it includes the mouth and nose, the tongue, the pharynx, epiglottis and laryngeal structures. The lower airway begins below the level of the larynx.

Related Glossary Terms
Airway obstruction, Cough reflex, Epiglottis, Hypopharynx, Laryngeal mask airway (LMA), Larynx, Lower airway, Patency, Subglottis, Vallecula
Urticaria

Hives. Urticaria is caused by histamine release which can result from the direct effect of a histamine-releasing medication (such as morphine) or by an allergic reaction.

Muscle relaxants are the most common anesthetic cause of allergic reaction.

Related Glossary Terms
Non-depolarizing muscle relaxants

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Uterine tone

Inhaled volatile anesthetic agents decrease uterine tone. Uterine hypotonicity can lead to excessive bleeding. As a result, the concentration of volatile agent delivered during Caesarian section must be limited. Oxytocin is given after all deliveries to ensure adequate uterine tone.

Related Glossary Terms

Oxytocin, Volatile

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Chapter 5 - Obstetrical Anesthesia
**Vagolytic**

Drugs with a vagolytic effect counteract or inhibit the action of the vagus nerve.

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**Related Glossary Terms**
Pancuronium, Vagus nerve

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Chapter 6 - Muscle Relaxants
Chapter 6 - Muscle Relaxants
**Vagus nerve**

The vagus nerve is the tenth cranial nerve. It carries the parasympathetic outflow to the heart and other visceral organs. The most important clinical effect of the vagus nerve is bradycardia.

The vagus nerve can be stimulated by many different types of noxious stimuli, including laryngoscopy, uterine or cervical traction and ophthalmic pressure. It can also be stimulated by emotional triggers. If you have fainted at the sight of blood, then you have experienced the impact of your own vagus nerve!

**Related Glossary Terms**

Acetylcholine, Anticholinergic, Anticholinesterase, Antisialagogue, Atropine, Autonomic nervous system, Glycopyrrolate, Muscarinic, Recurrent laryngeal nerve, Superior laryngeal nerve, Sympathetic nervous system, Vagolytic
Vallecula

The vallecula is the space between the base of the tongue and the epiglottis. During direct laryngoscopy, one aims to position the tip of the laryngoscope in the vallecula. Correct positioning ensures that the lifting maneuver will successfully displace the epiglottis upwards to expose the larynx.

Related Glossary Terms
Direct laryngoscopy, Intubation, Laryngoscope, Larynx, Upper airway

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Chapter 1 - Airway Management
Vapourizer

An anesthetic vapourizer is a device that is attached to the anesthetic machine and through which the fresh gas from the flowmeters (oxygen, usually mixed with air or nitrous oxide) flows. The vaporizer is able to deliver a precise concentration of volatile anesthetic agent into that fresh gas mixture, according to the concentration selected by the anesthesiologist.

The vaporizer is a sophisticated device which is designed to adjust to variations in ambient temperature, fresh gas flow and the vapour pressure of the volatile agent in use. There are three different types in use in modern anesthesia care: the plenum, the drawover and the dual-circuit gas–vapour blender. The latter type is required for the use of desflurane.

A given vapourizer is dedicated to the use of one particular volatile agent. For safety reasons, they are colour coded. As well, they have unique nozzle designs that prevents the wrong type of volatile anesthetic from filling the vapourizer. They are also designed such that only one volatile cannister can be “on” at once. In other words, you can not deliver more than one type of volatile agent at a time.

Related Glossary Terms
Desflurane, Inhaled agents, Isoflurane, MAC, Sevoflurane, Volatile

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Chapter 5 - Malignant Hyperthermia
Venous air embolism (VAE) is a serious condition where air enters the venous circulation and travels to the right side of the heart and (possibly) the lungs. If large enough, VAE can cause cardiovascular collapse. VAE can occur iatrogenically, particularly during procedures where veins are “opened” while situated at a level higher than that of the right atrium.

The sitting position, rarely used in modern practice, places the patient at high risk of VAE.
## Ventilator

A ventilator is a machine which delivers an inspiratory breath to the patient using positive pressure and allows the passive expiration of that breath. Modern ventilators have many different modes that allow it to accommodate the pathophysiology of a variety of pulmonary and systemic conditions as well as the specific goals of a given anesthetic technique.

The most basic distinctions in modes are:

i) Volume control, where the tidal volume is set and the pressure generated is variable, a function of pulmonary compliance.

ii) Pressure control, where the pressure delivered is set and the tidal volume is variable, (also) a function of pulmonary compliance.

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**Related Glossary Terms**

Mechanical ventilation, Positive pressure ventilation, Tidal volume

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Chapter 2 - Anesthetic Equipment and Monitoring
Videolaryngoscope

The videolaryngoscope is an airway device that is used, much like a direct laryngoscope, to achieve endotracheal intubation. The difference is that rather than having a light at the tip (like a direct laryngoscope), the videolaryngoscope has a tiny fibreoptic camera at its tip.

The camera gives the operator a view of the anatomy that is at the tip of the scope, and that view is projected on a small screen. Therefore the operator uses this “indirect” view of the laryngeal anatomy to guide the placement of the endotracheal tube.

The advantage of the videolaryngoscope is that it improves the view of the structures that would otherwise be seen through direct laryngoscopy, usually by 1-2 grades.

Although the videolaryngoscope provides a great view of the cords (in most cases), it requires some skill and practice to place the tube, as one has not necessarily achieved a “straight line” from mouth to larynx as must happen in the case of direct laryngoscopy.

A videolaryngoscope is useful in the management of the difficult airway. It is usually used on the anesthetized patient.

The Glidescope™ and the McGrath™ are two types of videolaryngoscopes.

Related Glossary Terms
Difficult airway, Direct laryngoscopy, Esophageal intubation, Intubation, Laryngoscope, Macintosh blade, Magill blade

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Chapter 1 - Airway Management
Volatile agents are a subset of inhaled agents and include desflurane, sevoflurane and isoflurane. They are “volatile” because at room temperature and 1 atmosphere, they are in equilibrium between liquid and gaseous states. They are used for maintenance of anesthesia. Sevoflurane is also used for (mask) induction of anesthesia.

Nitrous oxide is not a volatile agent, though it is an inhaled agent.

Related Glossary Terms
Anesthetic circuit, Apneic threshold, Desflurane, Flowmeter, Four A's of anesthesia, Inhaled agents, Isoflurane, Nitrous oxide (N2O), Sevoflurane, Uterine tone, Vapourizer

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