Anaesthetics

A- General Anaesthesia

General anaesthesia is a reversible state of central nervous system (CNS) depression, causing loss of response to and perception of stimuli. For patients undergoing surgical or medical procedures, anaesthesia provides five important benefits:

- Sedation and reduced anxiety
- Lack of awareness and amnesia
- Skeletal muscle relaxation
- Suppression of undesirable reflexes
- Analgesia

❖ Because no single agent provides all desirable properties, several categories of drugs are combined to produce optimal anaesthesia (Figure 1).
❖ Preanesthetics help calm patients, relieve pain, and prevent side effects of subsequently administered anesthetics or the procedure itself.
❖ In addition, the neuromuscular blockers facilitate tracheal intubation and surgery.
❖ Potent general anesthetics usually are delivered via inhalation and/or intravenous (IV) injection.

Patient factors in selection of anaesthesia

Drugs are chosen to provide safe and efficient anaesthesia based on the type of procedure and patient characteristics such as organ function, medical conditions and concurrent medications.

A. Status of organ systems

   So, they should be used with caution in patients with coronary artery disease, heart failure, dysrhythmias, and other cardiovascular disorders.
   To treat the hypotension, which may develop during anaesthesia and lead to reduce perfusion pressure and ischemic injury to tissues, vasoactive agents may be used.
   Some anaesthetics, such as halothane, sensitize the heart to arrhythmogenic effects of sympathomimetic agents.

Figure 1: Actions of anesthesia adjunct drugs.
2. **Respiratory system:** Respiratory function must be considered for all anesthetics. Asthma and ventilation or perfusion abnormalities complicate control of inhalation anesthetics. Inhaled agents depress respiration but also act as bronchodilators. IV anesthetics and opioids suppress respiration. These effects may influence the ability to provide adequate ventilation and oxygenation during and after surgery.

3. **Liver and kidney:** The liver and kidneys influence long-term distribution and clearance of drugs and are also target organs for toxic effects.

4. **Nervous system:** The presence of neurologic disorders (for example, epilepsy, myasthenia gravis, neuromuscular disease, compromised cerebral circulation) influences the selection of anesthetic.

5. **Pregnancy:** Special precautions should be observed when anesthetics and adjunctive agents are administered during pregnancy. Effects on fetal organogenesis are a major concern in early pregnancy. Transient use of nitrous oxide may cause aplastic anemia in the fetus. Oral clefts have occurred in fetuses when mothers received benzodiazepines in early pregnancy. Benzodiazepines should not be used during labor because of resultant temporary hypotonia and altered thermoregulation in the new born.

**B- Concomitant use of drugs**

1. **Multiple adjunct agents:** Premedications play an important role in anesthesia as they can facilitate the smooth induction of anesthesia and lower required anesthetic doses. However, they can also enhance undesirable anesthetic effects (hypoventilation) and, when coadministered, may produce negative effects not observed when given individually.

2. **Concomitant use of other drugs:** Patients may take medications for underlying diseases or abuse drugs that alter response to anesthetics. For example, **alcoholics** have elevated levels of liver enzymes that metabolize anesthetics, and **drug abusers** may be tolerant to opioids.

**STAGES AND DEPTH OF ANESTHESIA**

General anesthesia has three stages: induction, maintenance, and recovery. **Induction** is the time from administration of a potent anesthetic to development of effective anesthesia. Maintenance provides sustained anesthesia. Recovery is the time from discontinuation of anesthetic until consciousness and protective reflexes return. Induction of anesthesia depends on how fast effective concentrations of anesthetic reach the brain. Recovery is essentially the reverse of induction and depends on how fast the anesthetic diffuses from the brain. Depth of anesthesia is the degree to which the CNS is depressed.
**Induction**

General anesthesia in adults is normally induced with an IV agent like propofol, producing unconsciousness in 30 to 40 seconds. Additional inhalation and/or IV drugs may be given to produce the desired depth of anesthesia. [Note: This often includes an IV neuromuscular blocker such as rocuronium, vecuronium, or succinylcholine to facilitate tracheal intubation and muscle relaxation.]

**B. Maintenance of anesthesia**

After administering the anesthetic, vital signs and response to stimuli are monitored continuously to balance the amount of drug inhaled and/or infused with the depth of anesthesia. Maintenance is commonly provided with volatile anesthetics, which offer good control over the depth of anesthesia. Opioids such as fentanyl are used for analgesia along with inhalation agents, because the latter are not good analgesics. IV infusions of various drugs may be used during the maintenance phase.

**C. Recovery**

Postoperatively, the anesthetic admixture is withdrawn, and the patient is monitored for return of consciousness. For most anesthetic agents, recovery is the reverse of induction. Redistribution from the site of action (rather than metabolism of the drug) underlies recovery.

If neuromuscular blockers have not been fully metabolised, reversal agents may be used. The patient is monitored to assure full recovery, with normal physiologic functions (spontaneous respiration, acceptable blood pressure and heart rate, intact reflexes, and no delayed reactions such as respiratory depression).

**THE DEPTH OF ANESTHESIA**

The depth of anesthesia has four sequential stages characterized by increasing CNS depression as the anesthetic accumulates in the brain. [Note: These stages were defined for the original anesthetic ether, which produces a slow onset of anesthesia. With modern anesthetics, the stages merge because of the rapid onset of stage III.]

1. **Stage I—Analgesia:** Loss of pain sensation results from interference with sensory transmission in the spinothalamic tract. The patient progresses from conscious and conversational to drowsy. Amnesia and reduced awareness of pain occur as stage II is approached.

2. **Stage II—Excitement:** The patient displays delirium and possibly combative behavior. A rise and irregularity in blood pressure and respiration occur, as well as a risk of laryngospasm. To shorten or eliminate this stage, rapid-acting IV agents are given before inhalation anesthesia is administered.

3. **Stage III—Surgical anesthesia:** There is gradual loss of muscle tone and reflexes as the CNS is further depressed. Regular respiration and relaxation of skeletal muscles with eventual loss of spontaneous movement occur. This is the ideal stage for surgery. Careful monitoring is needed to prevent undesired progression to stage IV.
4. **Stage IV—Medullary paralysis:** Severe depression of the respiratory and vasomotor centers occurs. Ventilation and/or circulation must be supported to prevent death.

A brief summary about depth of anaesthesia and its stages are illustrated in Figure 2.

![Figure 2: Stages of anesthesia.](image)

**Figure 2: Stages of anesthesia.**

There are 3 different routes of administration of the anaesthetics, which are:

A- **INHALATION ANESTHETICS**
- Inhaled gases are used primarily for maintenance of anesthesia after administration of an IV agent.
- Depth of anesthesia can be rapidly altered by changing the inhaled concentration. The inhaled anaesthetics are summarised in Figure 3.

![Figure 3: Inhaled anaesthetics](image)

**Common features of inhalation anesthetics**
- Modern inhalation anesthetics are non-flammable, nonexplosive agents.
- They include *nitrous oxide* and volatile, halogenated hydrocarbons.
- Movement of these agents from the lungs to various body compartments depends upon
  - Their solubility in blood and tissues, as well as on blood flow.
  - The above-mentioned factors play a role in induction and recovery.
Mechanism of action of inhaled anesthetics

❖ Generally, no specific receptor has been identified as the locus of general anesthetic action. The fact that chemically unrelated compounds produce anesthesia argues against the existence of a single receptor.

❖ At clinically effective concentrations, general anesthetics increase the sensitivity of the γ-aminobutyric acid (GABAA) receptors to the inhibitory neurotransmitter GABA. This increases chloride ion influx and hyperpolarization of neurons. Postsynaptic neuronal excitability and, thus, CNS activity are diminished.

❖ Unlike other anesthetics, nitrous oxide and ketamine do not have actions on GABAA receptors. Their effects are likely mediated via inhibition of the N-methyl-d-aspartate (NMDA) receptors. [Note: The NMDA receptor is a glutamate receptor. Glutamate is the body’s main excitatory neurotransmitter.]

Halothane

❖ Halothane is the prototype to which newer inhalation anesthetics are compared. When halothane was introduced, its rapid induction and quick recovery made it an anesthetic of choice. Due to adverse effects and the availability of other anesthetics with fewer complications, halothane has been replaced in most countries.

❖ Therapeutic uses

✓ Halothane is a potent anesthetic but a relatively weak analgesic. Thus, it is usually coadministered with nitrous oxide, opioids, or local anesthetics.

✓ Halothane relaxes both skeletal and uterine muscles and can be used in obstetrics when uterine relaxation is indicated.

✓ Halothane is not hepatotoxic in children (unlike its potential effect on adults).

✓ Combined with its pleasant odor, it is suitable in pediatrics for inhalation induction, although sevoflurane is now the agent of choice.

✓ Adverse effects such as cardiac arrhythmias and produce concentration-dependent hypotension, which can be best treated with a direct-acting vasoconstrictor, such as phenylephrine. Moreover, it can cause Malignant hyperthermia (MH).

Nitrous oxide (NO)

• NO (“laughing gas”) is a non-irritating potent analgesic but a weak general anesthetic. It is frequently used at concentrations of 30 to 50% in combination with oxygen for analgesia, particularly in dentistry.

• NO alone cannot produce surgical anesthesia, but it is commonly combined with other more potent agents.
• **NO** is poorly soluble in blood and other tissues, allowing it to move very rapidly in and out of the body.

• Within closed body compartments, *nitrous oxide* can increase the volume (for example, causing a pneumothorax) or pressure (for example, in the sinuses), because it replaces nitrogen in various air spaces faster than the nitrogen leaves. Its speed of movement allows *nitrous oxide* to retard oxygen uptake during recovery, thereby causing “diffusion hypoxia,” which can be overcome by significant concentrations of inspired oxygen during recovery.

• **NO** does not depress respiration and does not produce muscle relaxation.

• When coadministered with other anesthetics, it has moderate to no effect on the cardiovascular system or on increasing cerebral blood flow, and it is the least hepatotoxic of the inhalation agents.

• Therefore, it is probably the safest of these anesthetics, provided that sufficient oxygen is administered simultaneously.

**INTRAVENOUS ANESTHETICS**

• IV anesthetics cause rapid induction often occurring within one “arm–brain circulation time,” or the time it takes to travel from the site of injection (usually the arm) to the brain, where it has its effect.

• Anesthesia may then be maintained with an inhalation agent.

• IV anesthetics may be used as sole agents for short procedures or administered as infusions to help maintain anesthesia during longer cases.

• In lower doses, they may be used for sedation.

**Propofol**

• *Propofol* is an IV sedative/hypnotic used for induction and/or maintenance of anesthesia.

• It is widely used and has replaced *thiopental* as the first choice for induction of general anesthesia and sedation.

• Because *propofol* is poorly water soluble, it is supplied as an emulsion containing soybean oil and egg phospholipid, giving it a milk-like appearance.

• Induction is smooth and occurs 30 to 40 seconds after administration.

• *Propofol* is commonly infused in lower doses to provide sedation.

• The incidence of postoperative nausea and vomiting is very low, as this agent has some antiemetic effects.

**Barbiturates**

• *Thiopental* is an ultra–short-acting barbiturate with high lipid solubility. It is a potent anesthetic but a weak analgesic.

• Barbiturates require supplementary analgesic administration during anesthesia.

• When given IV, agents such as *thiopental* and *methohexital* quickly enter the CNS and depress function, often in less than 1 minute.

• These drugs may remain in the body for relatively long periods, because only about 15% of a dose entering the circulation is metabolized by the liver per hour.

• *Thiopental* has minor effects on the normal cardiovascular system but may contribute to severe hypotension in patients with hypovolemia or shock.
Benzodiazepines
- The benzodiazepines are used in conjunction with anesthetics for sedation.
- The most commonly used is midazolam.
- Diazepam and lorazepam are alternatives. All three facilitate amnesia while causing sedation, enhancing the inhibitory effects of various neurotransmitters, particularly GABA.
- Benzodiazepines can induce a temporary form of anterograde amnesia in which the patient retains memory of past events, but new information is not transferred into long-term memory. Therefore, important treatment information should be repeated to the patient after the effects of the drug have worn off.

Opioids
- Because of their analgesic property, opioids are commonly combined with other anesthetics. The choice of opioid is based primarily on the duration of action needed. The most commonly used opioids are fentanyl, sufentanil, and remifentanil, because they induce analgesia more rapidly than morphine.
- They may be administered intravenously, epidurally, or intrathecally (into the cerebrospinal fluid).
- Opioids are not good amnesics, and they can all cause hypotension, respiratory depression, and muscle rigidity, as well as postanesthetic nausea and vomiting.
- Opioid effects can be antagonized by naloxone.
- In addition to the above-mentioned drugs, there are Etomidate, Ketamine, Neuromuscular blockers and Dexmedetomidine. Their important properties are summarised in figure 4.

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**Figure 4: Therapeutic disadvantages and advantages of some anesthetic agents.**
(MEMORISABLE)
LOCAL ANESTHETICS

- Local anesthetics block nerve conduction of sensory impulses and, in higher concentrations, motor impulses from the periphery to the CNS.
- Na⁺ ion channels are blocked to prevent the transient increase in permeability of the nerve membrane to Na⁺ that is required for an action potential.
- When propagation of action potentials is prevented, sensation cannot be transmitted from the source of stimulation to the brain.
- Different delivery techniques can be used such as topical administration, infiltration, peripheral nerve blocks.
- Structurally, local anesthetics all include a lipophilic group joined by an amide or ester linkage to a carbon chain, which, in turn, is joined to a hydrophilic group (Figure 5).
- The most widely used local anesthetics are bupivacaine, lidocaine, mepivacaine, procaine, ropivacaine, and tetracaine. Bupivacaine is noted for cardiotoxicity if inadvertently injected IV.
- Bupivacaine liposome injectable suspension may provide postsurgical analgesia lasting 24 hours or longer after injection into the surgical site.
- Non-bupivacaine local anesthetics may cause an immediate release of bupivacaine from the liposomal suspension if administered together locally.
- Mepivacaine should not be used in obstetric anesthesia due to its increased toxicity to the neonate.

![Ester and Amide Anesthetics](image)

**Figure 5:** Representative structures of ester and amide anesthetics.

Metabolism

- Biotransformation of amides occurs primarily in the liver. Prilocaine, a dental anesthetic, is also metabolized in the plasma and kidney, and one of its metabolites may lead to methemoglobinemia.
- Esters are biotransformed by plasma cholinesterase (pseudocholinesterase). Patients with pseudocholinesterase deficiency may metabolize ester local anesthetics more slowly. At normal doses, this has little clinical effect.
Onset and duration of action
The onset and duration of action of local anesthetics are influenced by several factors including tissue pH, nerve morphology, concentration, pKa, and lipid solubility of the drug. Of these, the pH of the tissue and pKa are the most important. At physiologic pH, these compounds are charged. The ionized form interacts with the protein receptor of the Na+ channel to inhibit its function and achieve local anesthesia. The pH may drop in infected sites, causing onset to be delayed or even prevented. Within limits, higher concentration and greater lipid solubility improve onset somewhat. Duration of action depends on the length of time the drug can stay near the nerve to block sodium channels.

Actions
- Local anesthetics cause vasodilation, leading to rapid diffusion away from the site of action and shorter duration when these drugs are administered alone. By adding the vasoconstrictor epinephrine, the rate of local anesthetic absorption and diffusion is decreased. This minimizes systemic toxicity and increases the duration of action.
- Hepatic function does not affect the duration of action of local anesthesia, which is determined by redistribution and not biotransformation.
- Some local anesthetics have other therapeutic uses (for example, lidocaine is an IV antiarrhythmic).

Allergic reactions
- Patient reports of allergic reactions to local anesthetics are fairly common, but often times reported “allergies” are actually side effects from epinephrine added to the local anesthetic. Psychogenic reactions to injections may be misdiagnosed as allergic reactions and may also mimic them with signs such as urticaria, edema, and bronchospasm. True allergy to an amide local anesthetic is exceedingly rare, whereas the ester procaine is somewhat more allergenic. Allergy to one ester rules out use of another ester, because the allergenic component is the metabolite para-aminobenzoic acid, produced by all esters.
- In contrast, allergy to one amide does not rule out the use of another amide. A patient may be allergic to other compounds in the local anesthetic, such as preservatives in multidose vials.

Administration to children and the elderly
- Before administering local anesthetic to a child, the maximum dose based on weight should be calculated to prevent accidental overdose.
- There are no significant differences in response to local anesthetics between younger and older adults.
- It is prudent to stay well below maximum recommended doses in elderly patients who often have some compromise in liver function.
Because some degree of cardiovascular compromise may be expected in elderly patients, reducing the dose of epinephrine may be prudent.

Local anesthetics are safe for patients who are susceptible to malignant hyperthermia.

**Systemic local anesthetic toxicity**

- Toxic blood levels of the drug may be due to repeated injections or could result from a single inadvertent IV injection.
- Aspiration before every injection is imperative. The signs, symptoms, and timing of local anesthetic systemic toxicity are unpredictable. One must consider the diagnosis in any patient with altered mental status or cardiovascular instability following injection of local anesthetic. CNS symptoms may be apparent but may also be nonspecific or absent.
- Treatment for systemic local anesthetic toxicity includes airway management, support of breathing and circulation, seizure suppression and, if needed, cardiopulmonary resuscitation. Administering a 20% lipid emulsion infusion (lipid rescue therapy) is a valuable asset.

Figure 6 summarizes pharmacologic properties of some local anesthetics.

<table>
<thead>
<tr>
<th>CHARACTERISTIC</th>
<th>ESTERS</th>
<th>AMIDES</th>
<th>POTENCY</th>
<th>ONSET</th>
<th>DURATION</th>
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<tbody>
<tr>
<td>Metabolism</td>
<td>Proacne</td>
<td>Lidocaine</td>
<td>Low</td>
<td>Rapid</td>
<td>Short</td>
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<td></td>
<td>Chloroprocaine</td>
<td>Bupivacaine</td>
<td>Low</td>
<td>Rapid</td>
<td>Short</td>
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<td></td>
<td>Tetracaine</td>
<td>Prilocaine</td>
<td>High</td>
<td>Slow</td>
<td>Long (spinal)</td>
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<td></td>
<td>Cocaine</td>
<td>Ropivaclene</td>
<td>Low</td>
<td>Moderate</td>
<td>Intermediate</td>
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<tr>
<td>Systemic toxicity</td>
<td>Less likely</td>
<td>More likely</td>
<td></td>
<td></td>
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<tr>
<td>Allergic reaction</td>
<td>Possible</td>
<td>PABA derivatives form</td>
<td>Very rare</td>
<td></td>
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<tr>
<td>Stability in solution</td>
<td>Breaks down in ampules (heat, sun)</td>
<td>Very stable chemically</td>
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<tr>
<td>Onset of action</td>
<td>Slow as a general rule</td>
<td>Moderate to fast</td>
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<td>pH₄'s</td>
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<td>Close to physiologic pH (7.6–8.1)</td>
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Figure 6: Summary of pharmacologic properties of some local anesthetics. PABA = para-aminobenzoic acid. (MEMORISABLE)