

Medical Pharmacology: The science of substances which are used for the prevention, diagnosis and treatment of diseases.

Pharmacy: It is the science that deals with the preparing & dispensing the drugs

Drug: It is the chemical molecules which alter biochemistry & physiology of the living system when introduced inside the body
The Physical Nature of Drugs
Drugs may be
Solid at room temperature (e.g. aspirin, atropine)
Liquid (e.g. nicotine, ethanol)
Gaseous (e.g. nitrous oxide)
These factors often determine the best route of administration.

Routes of administration, bioavailability, and general characteristics

Route	Biovailability (%)	Characteristics
Intravenous (IV)	100 (by definition)	Most rapid onset
intramuscular (IM)	75 to ≤ 100	Large volumes often feasible; may be painful
Subcutaneous (SC)	75 to ≤ 100	Smaller volumes than IM; may be painful
Oral (PO)	5 to < 100	Most convenient; first- pass effect may be important
Rectal (PR)	30 to < 100	Less first-pass effect than oral
Inhalation	5 to < 100	Often very rapid onset
Transdermal	80 to ≤ 100	Usually very slow absorption; used for lack of first-pass effect; prolonged duration of action

Drugs may be: Synthesized within the body (e.g. hormones) Chemicals *not* synthesized in the body, ie, xenobiotics (from the Greek *xenos*, meaning stranger). Poisons are drugs that have almost exclusively harmful effects. Toxins are poisons of biologic origin, ie, synthesized by plants or animals, in contrast to inorganic poisons such as lead.

Drugs may be

- Organic compounds (carbohydrates, proteins, lipids, and their constituents). organic drugs are weak acids or bases. This fact has important implications for the way they are handled by the body, because pH differences in the various compartments of the body may alter the degree of ionization of such drugs
- Inorganic elements, e.g. iron

Drug Size

- The molecular size of drugs varies from very small (MW 7) to very large (MW 59,050).
- most drugs have molecular weights between 100 and 1000.
- Drugs much larger than MW 1000 do not diffuse readily between compartments of the body (Therefore, very large drugs (usually proteins) must often be administered directly into the compartment where they have their effect. In the case of alteplase, a clot-dissolving enzyme, the drug is administered directly into the vascular compartment by intravenous or intraarterial infusion

Drug Reactivity & Drug-Receptor Bonds

Drugs interact with receptors by means of chemical forces or bonds. There are three major types

- 1. Covalent
- 2. Electrostatic
- 3. Hydrophobic

Covalent Bonds

Very strong(not broken) and in many cases not reversible under biologic conditions. EXAMPLE: the covalent bond formed between the acetyl group of aspirin and its enzyme target in platelets cyclooxygenase **Electrostatic Bonds** Weaker than covalent bonds. **Hydrophobic Bonds** Quite weak and are probably important in the interactions of highly lipid-soluble drugs with the lipids of cell membranes

Drug Shape

The shape of a drug molecule must be permit binding to its receptor site via the bonds, the drug's shape is complementary to that of the receptor site in the same way that a key is complementary to a lock.

Drug-body Interactions

The interactions between a drug and the body are divided into two classes.

Pharmacodynamic

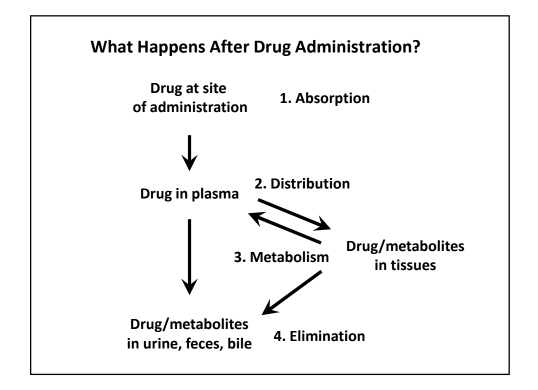
What a drug does to the body, such as mechanism of action and therapeutic and toxic effects.

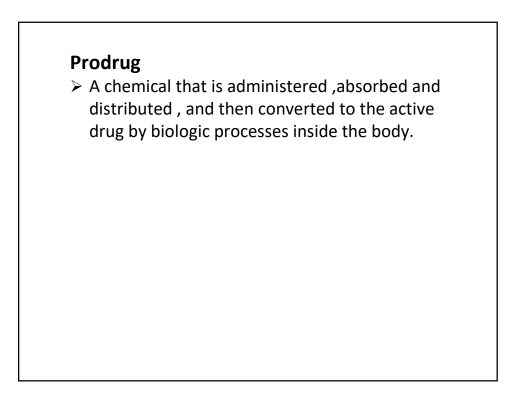
Pharmacokinetic

What the body does to the drug or how the body deals with the drug or the fate of the drug in the body

Pharmacokinetics processes are

Absorption, Distribution, Metabolism and Excretion





Drug receptors Coupling

The overall transduction process that links drug occupancy of receptors and pharmacologic response is called **coupling**

Agonist

Drug or endogenous regulatory molecule that occupies a receptor& make conformational changes &produce a pharmacologic response.

Agonists can be divided into

- Full agonists produce maximal pharmacologic response
- Partial agonists produce a lower response, at full receptor occupancy, than do full agonists.

NOTE: Partial agonists produce concentration-effect curves that resemble those observed with full agonists in the presence of an antagonist that irreversibly blocks some of the receptor sites

Receptor antagonists

Bind to receptors but do not activate them; the primary action of antagonists is to reduce the effects of agonists

Antagonist drugs

- > Competitively
- > Noncompetitively

Note :Some drugs mimic agonist drugs by inhibiting the molecules responsible for terminating the action of an endogenous agonist **Example:**

acetylcholinesterase inhibitors, Inhibit acetylcholinesterase enzyme(the enzyme responsible for the destruction of endogenous acetylcholine, cause cholinomimetic effects(resemble the actions of cholinoceptor agonist)

Antagonist drugs

> Competitively

> Noncompetitively

Pharmacologic antagonist

- Drugs activate the agonist binding site, by binding to a receptor, compete with and prevent binding by other molecules.
- Example
- Atropine(acetylcholine receptor blockers) are antagonists because they prevent access of acetylcholine to the acetylcholine receptor site and they stabilize the receptor in its inactive state (or some state other than the acetylcholineactivated state).

Allosteric modulators

Drugs that bind to the same receptor but do not prevent binding of the agonist , may enhance or inhibit action of the agonist molecule.

Allosteric inhibitors (Non competetive antagonists)(negative allosteric modulators)bind very tightly to the receptor site in an irreversible or pseudoirreversible fashion and cannot be displaced by increasing the agonist concentration(by increasing the dose of agonist)

Example of irreversible α -adrenoceptor antagonist

Phenoxybenzamine, an irreversible α -adrenoceptor antagonist, is used to control the hypertension caused by catecholamines released from pheochromocytoma, a tumor of the adrenal medulla. If administration of phenoxybenzamine lowers blood pressure, blockade will be maintained even when the tumor releases very large amounts of catecholamine.

Allosteric activators (positive allosteric modulators) may increase the efficacy of the agonist or its binding

Example

Benzodiazepines bind noncompetitively to ion channels activated by the neurotransmitter γ -aminobutyric acid (GABA), thereby enhancing the net activating effect of GABA on channel conductance

Note:

Allosteric antagonists do not bind to the agonist receptor site; they bind to some other region of the receptor molecule that results in inhibition of the response to agonists ,they do not prevent binding of the agonist.

In contrast, pharmacologic antagonists bind to the agonist site and prevent access of the agonist, High concentrations of agonist displace or prevent

the binding of a pharmacologic antagonist but not an allosteric antagonist.

Antagonism by receptor block(Receptor Antagonist) ≻Competitive: They compete for the binding site Reversible

1. Antagonist binds with the same receptor as the agonist

2-Antagonist resembles chemically with agonist

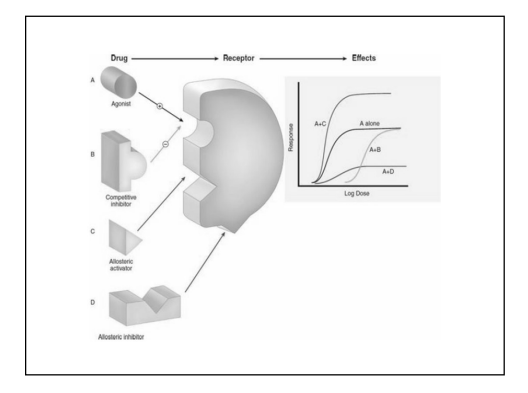
3-Antagonist reduces affinity of agonist

4-Intensity depend on conc. Of both agonist and antagonist

> Non-competitve

- 1.Antagonist binds to another binding site of the recepter
- 2- Not resembles chemically with agonist
- 3-Antaginist reduces efficacy of agonist
- 4-depend only on concentration of antagonist

Some antagonists exhibit "**inverse agonist**" activity ,because they also reduce receptor activity below basal levels observed in the absence of any agonist at all(activate the recepter to produce an effect in the opposite direction to that of the agonist



Constitutive activity(Basal Signal):

The receptor is able to have 2 states

Ri & Ra

- Ri state, it is inactive and produces no effect (nonfunctional), even when combined with a drug (D) molecule.
- > Ra state, it activates its effectors and

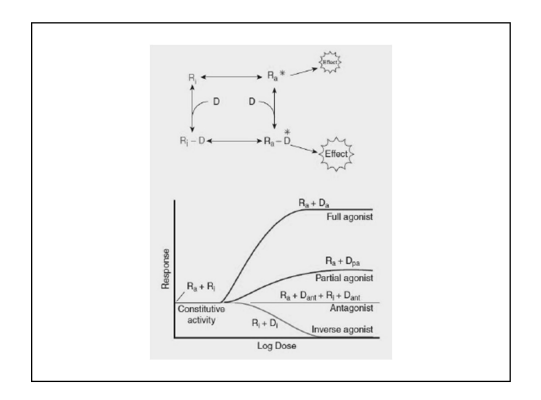
an effect is recorded, even in the absence of ligand.

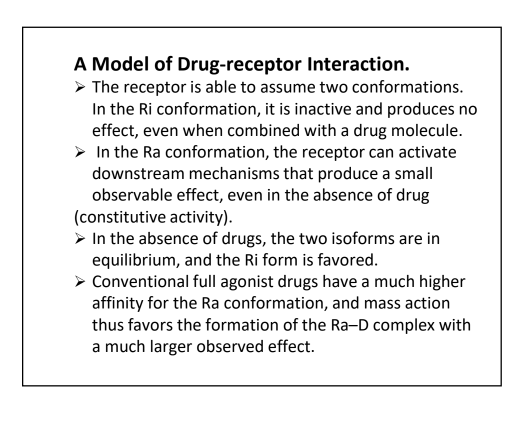
- In the absence of drug, the equilibrium between Ri and Ra determines the degree of constitutive activity.
- Thermodynamic considerations indicate that even in the absence of any agonist, some of the receptor pool must exist in the Ra form some of the time and may produce the same physiologic effect as agonist-induced activity

- A full agonist drug (Da) has a much higher affinity for the Ra than for the Ri receptor conformation, and a maximal effect is produced at sufficiently high drug concentration.
- A partial agonist drug (Dpa) has somewhat greater affinity for the Ra than for the Ri &produces less effect, even at saturating concentrations.
- A neutral antagonist (Dant) binds with equal affinity to both receptor conformations and prevents binding of agonist.
- > An inverse agonist (Di) binds much more to the

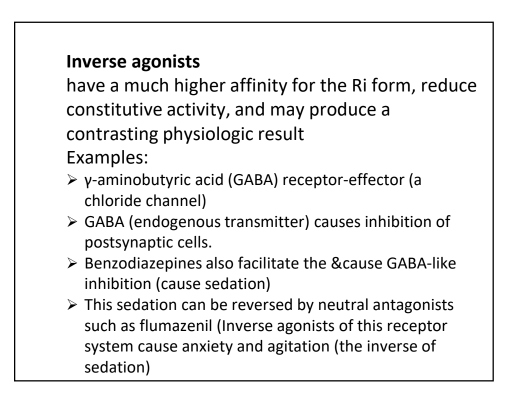
Ri receptor conformation, prevents conversion to the Ra state, and reduces constitutive activity

- The receptor is exist in the inactive, nonfunctional form (Ri) and in the activated form (Ra).
- Thermodynamic considerations indicate that even in the absence of any agonist, some of the receptor pool must exist in the Ra form some of the time and may produce the same physiologic effect as agonistinduced activity.
- This effect, occurring in the absence of agonist, is termed constitutive activity.





- Partial agonists have an intermediate affinity for both Ri and Ra forms.
- Conventional antagonists, according to this hypothesis, have equal affinity for both receptor forms and maintain the same level of constitutive activity.



Properties of an Ideal Drug

Effectiveness

Safety

Selectivity

Reversible action

Ease of administration

Freedom from drug Interactions

Low cost

Chemical Stability

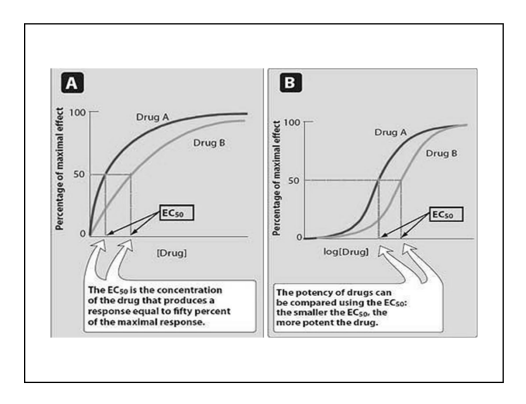
Efficacy

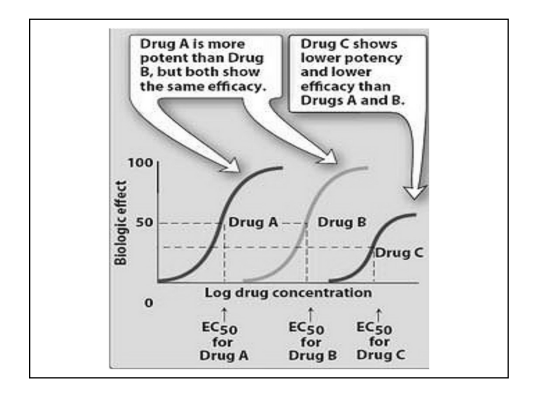
is the capacity of a drug to produce a specific effect **Potency**

Refers to the amount of drug (in weight) that give an effect

Example

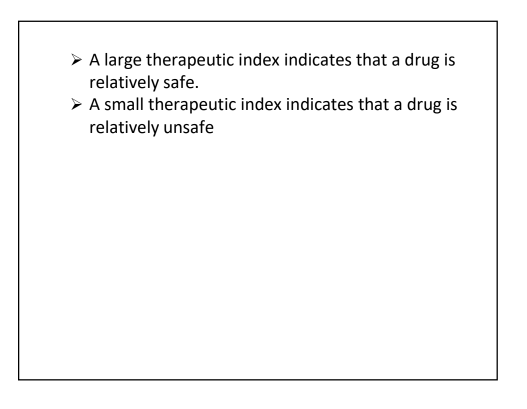
Morphine 10mg can relief severe pain, meperidine 100 mg can relief the same degree of pain relived by morphine, so morphine is more potent than meperidine.

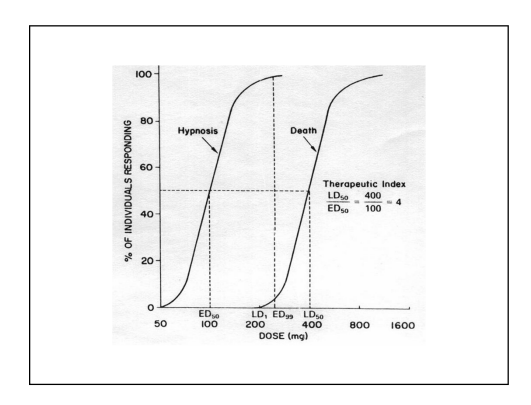


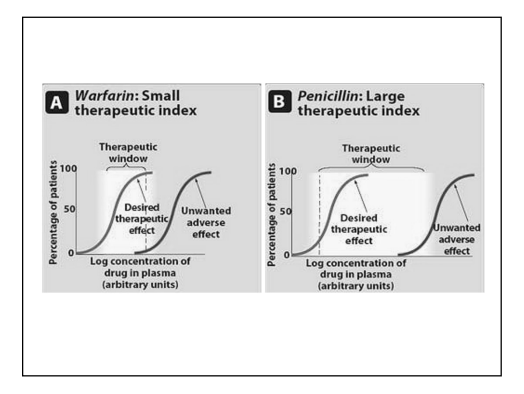


Toxicology

It is part of pharmacology which deals with the undesired or toxic effect of drugs. The therapeutic index The ED 50 and LD50 are used to calculate an important value in pharmacology, the ratio of a drug's LD 50 and its ED 50 ---- the higher the value, the safer the medication **TI=LD50/ED50 Median lethal dose (LD 50)** The dose of a chemical that causes death in 50% of the animals







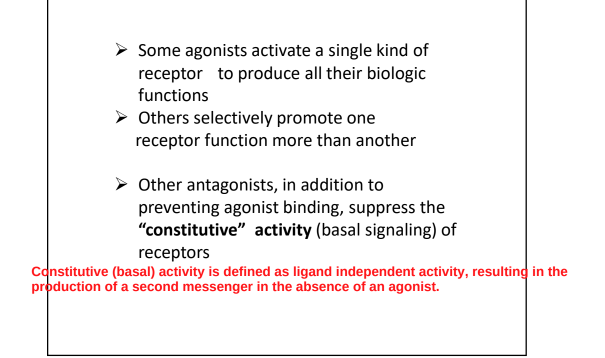
Selectivity of drug

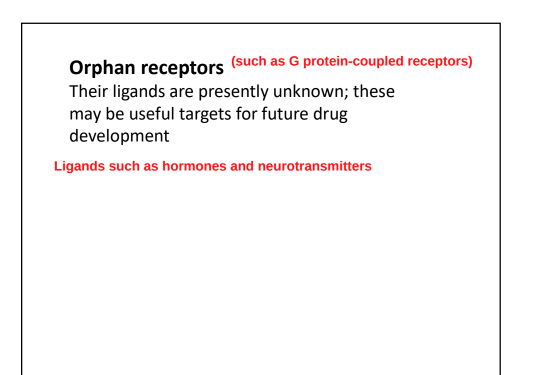
Receptors are responsible for selectivity of drug action.

The molecular size, shape, and electrical charge of a drug determine , changes in the chemical structure of a drug can dramatically increase or decrease a new drug's affinities for different classes of receptors, with resulting alterations in therapeutic and toxic effects.

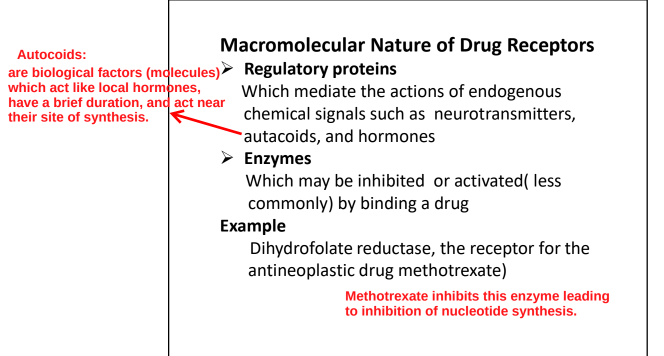
The sensitivity of a cell or tissue to a particular concentration of agonist depends on:

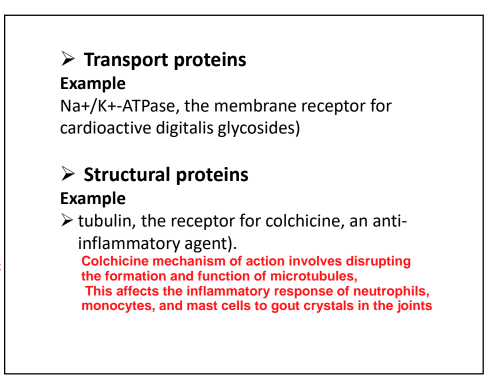
- The affinity of the receptor for binding the agonist
- The degree of spareness (the total number of receptors present compared with the number actually needed to elicit a maximal biologic response





24



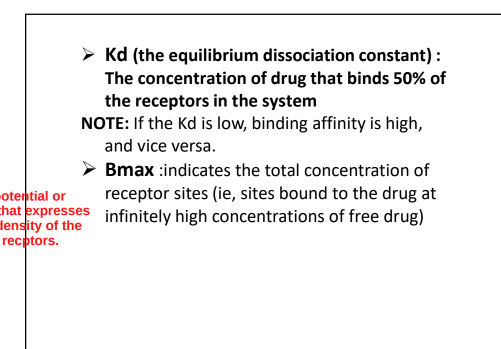


Building blocks of microtubules-- anti-cancerous effect

25

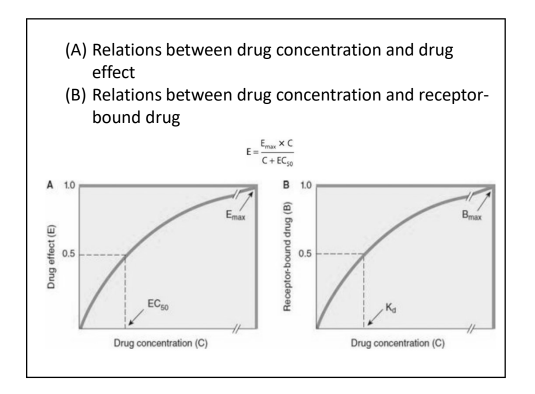
Concentration-Effect Curves & Receptor Binding of Agonists

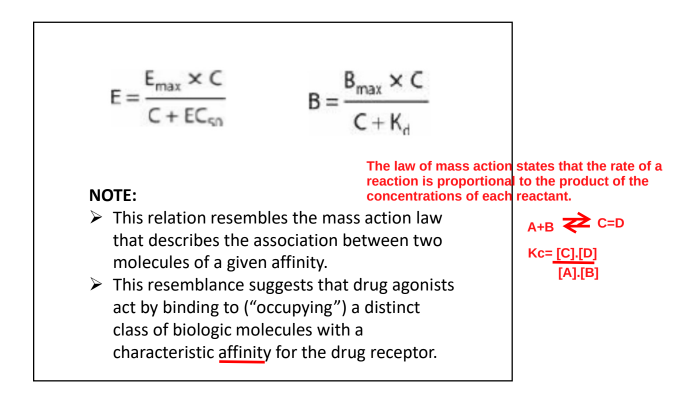
- E is the effect observed at concentration C
- > E max: is the maximal response that can be produced by the drug.
- EC50: is the concentration of drug that produces 50% of maximal effect.



B max:

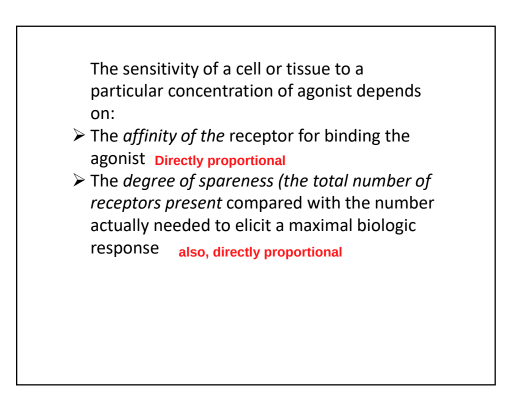
Binding potential or capacity that expresses the total density of the occupied recptors.





The Kd of the agonist-receptor interaction determines what fraction (B/Bmax) of total receptors will be occupied at a given free concentration (C) of agonist regardless of the receptor concentration

$$\frac{B}{B_{max}} = \frac{C}{C + K_d}$$



Spare Receptors

If it is possible to elicit a maximal biologic response at a concentration of agonist that does not result in occupancy of the full complement of available receptors.

Example

The same maximal inotropic response of heart muscle to catecholamines can be elicited even when 90% of the β adrenoceptors are occupied by a irreversible antagonist,,this mean myocardial cells contain a large proportion of spare β adrenoceptors.

Spare receptors

Experimental demonstration of spare receptors, using different concentrations of an irreversible antagonist.

- Curve A shows agonist response in the absence of antagonist.
- Curve B After treatment with a low concentration of antagonist, the curve is shifted to the right. Maximal responsiveness is preserved, however, because the remaining available receptors are still in excess of the number required.

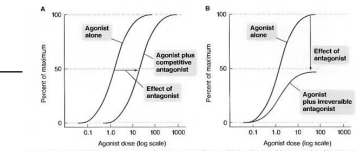
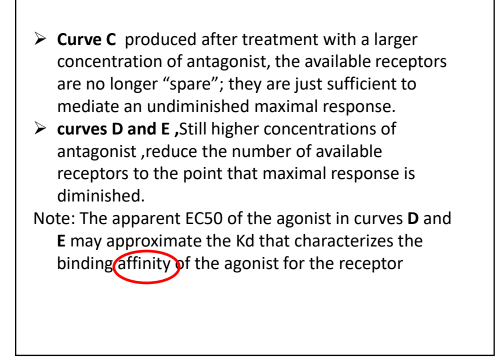
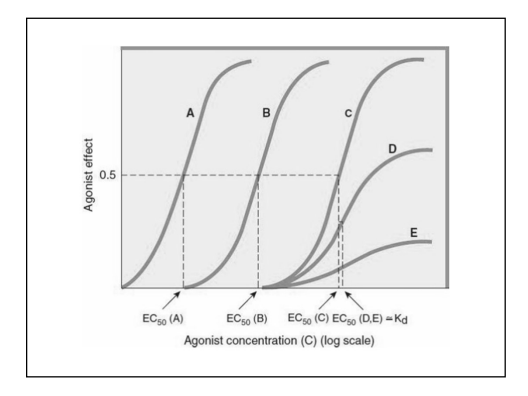


Figure 2–5. Agonist dose-response curves in the presence of competitive and inverse instead and accession in the presence of competitive and inverse instead and accession is not the the use of a logarithmic scale for drug concentration. A A competitive antagonist has an effect illustrated by the shift of the agonist curve to the right. B. A noncompetitive antagonist shifts the agonist curve downward.

This refers to the condition in a tissue whereby the agonist needs to activate only a small fraction of the existing receptor population to produce the maximal system response. The magnitude of the reserve depends upon the sensit vity of the tissue and the efficacy of the agonist.





some antagonists exhibit "inverse agonist" activity ,because they also reduce receptor activity below basal levels observed in the absence of any agonist at all.

Chemical antagonist i.e there is a chemical reaction between the agonist and the antagonist. some types of antagonism do not involve a receptor Example Protamine, a protein that is positively charged at physiologic pH, can be used clinically to counteract the effects of heparin, an anticoagulant that is negatively charged. In this case, one drug acts as a **chemical antagonist** of the other simply by ionic binding that makes the other drug unavailable for interactions with proteins involved in blood clotting. It negates the charge needed for interaction

Physiologic antagonism

mediated by different receptors Example

Elevated glucocorticoid by endogenous synthesis (a tumor of the adrenal cortex) or as a result of glucocorticoid therapy, several catabolic actions of the glucocorticoid hormones lead to increased blood sugar, an effect that is physiologically opposed by insulin.

The clinician administer insulin to oppose the hyperglycemic effects of a glucocorticoid hormone

NOTE:

physiologic antagonist produces effects that are less specific and less easy to control than are the effects of a receptor-specific antagonist. Example

To treat bradycardia caused by increased release of

acetylcholine from vagus nerve endings, the physician could use isoproterenol, a β -adrenoceptor agonist that increases heart rate by mimicking sympathetic stimulation of the heart.

- Use of this physiologic more dangerous than use of a receptor-specific antagonist such as atropine (a competitive
- antagonist at the receptors at which acetylcholine

