

ILOS

Determine quantitative aspects of drug receptor binding.

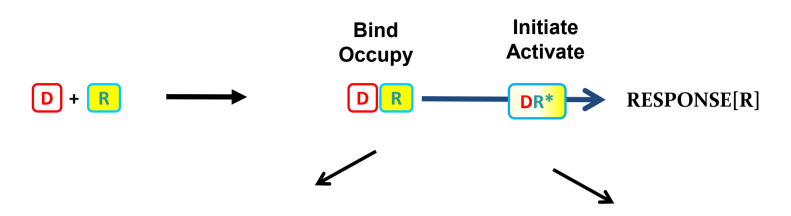
**Recognize concentration binding curves.** 

Identify dose response curves and the therapeutic utility of these curves.



**Classify different types of antagonism** 

#### QUANTIFY ASPECTS OF DRUG ACTION



Relate concentration [C] of D used (x- axis) Relate concentration [C] of D used (xto the binding capacity at receptors (y-axis) axis) to response produced (y-axis)

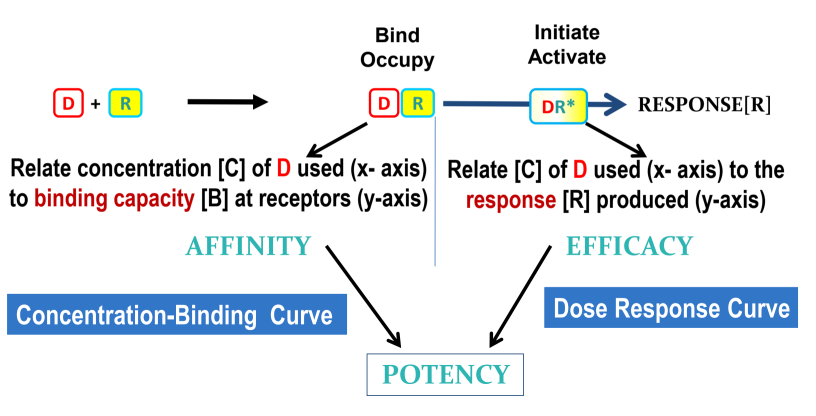
Concentration-Binding Curve
Dose Response Curves

AFFINITY
EFFICACY
POTENCY

The tendency of a drug to bind to the receptors is governed by itsAFFINITYaffinity.

The ability for it, once bound, to activate the receptor is denoted by its efficacy.

**EFFICACY** 



- Is a correlation between drug concentration
- [C] used (x- axis) and drug binding capacity at receptors [B] (y-axis).
- = is relation between concentration & drug binding
- = i.e. Affinity

**Concentration-Binding curves are used to determine:** 

## **OB**<sub>max</sub> (the binding capacity)

is the total density of receptors in the tissues.

# **K**<sub>D50</sub>

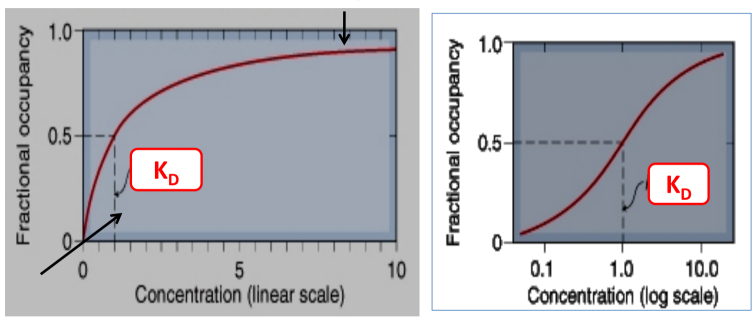
is the concentration of the drug required to occupy 50% of receptors at equilibrium.

## The affinity of drug for receptor

The higher the affinity of D for receptor, the lower is the  $K_D$ 

i.e. inverse relation ( Binding Potential = Bmax /K<sub>D</sub> )

#### **Concentration-Binding Curve**



(**B**<sub>max</sub>): Total density of receptors in the tissue

(k<sub>D</sub>)= [C] of D required to occupy 50% of receptors at equilibrium



#### DOSE -RESPONSE CURVES

- Is a correlation between <u>drug concentration [D]</u>
   used (x- axis) and <u>drug response [R] (y-axis)</u>.
- Used to study how response varies with the concentration of the drug or dose.
- i.e. relation between concentration & Response

### TYPES OF DOSE -RESPONSE CURVES

✓ Graded dose-response curve

✓ Quantal dose-response curve (all or none)

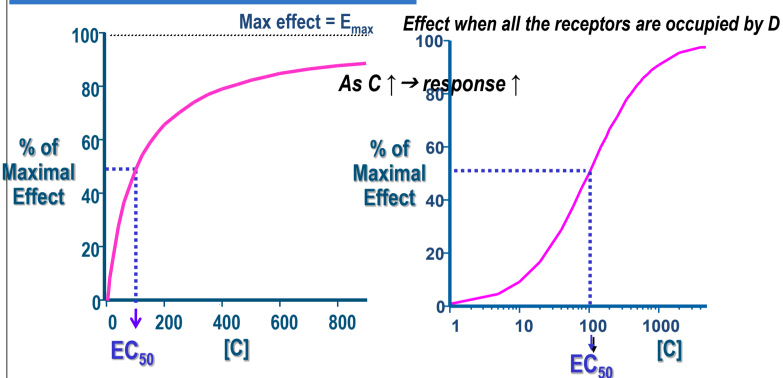
## **Graded Dose-Response Curve**

- Relate drug concentration to response.
- Response is gradual.
- Continuous response: increase in response by increasing the dose
- o e.g. ↓blood pressure, heart rate, blood glucose level, cholesterol,...
- Curve is usually sigmoid in shape

### **Graded dose-response curves are used to determine:**

- E<sub>max</sub>
- EC<sub>50</sub>
- Potency
- Efficacy

#### **GRADED DOSE RESPONSE CURVE**



**EC**<sub>50</sub> that gives half the maximal effect

**Graded dose-response curves are used to determine:** 

#### Maximum Efficacy (Emax):

is the maximal biological response produced by a drug.

### **Median Effective concentration (EC50):**

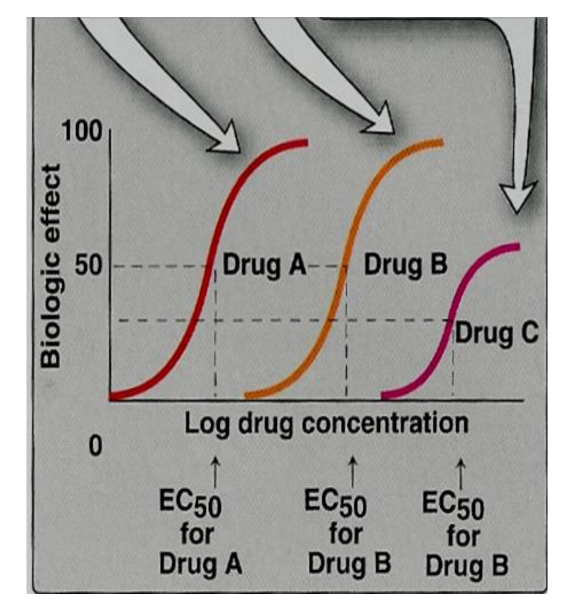
is the concentration of the drug that produces 50% of the maximal response (Emax)

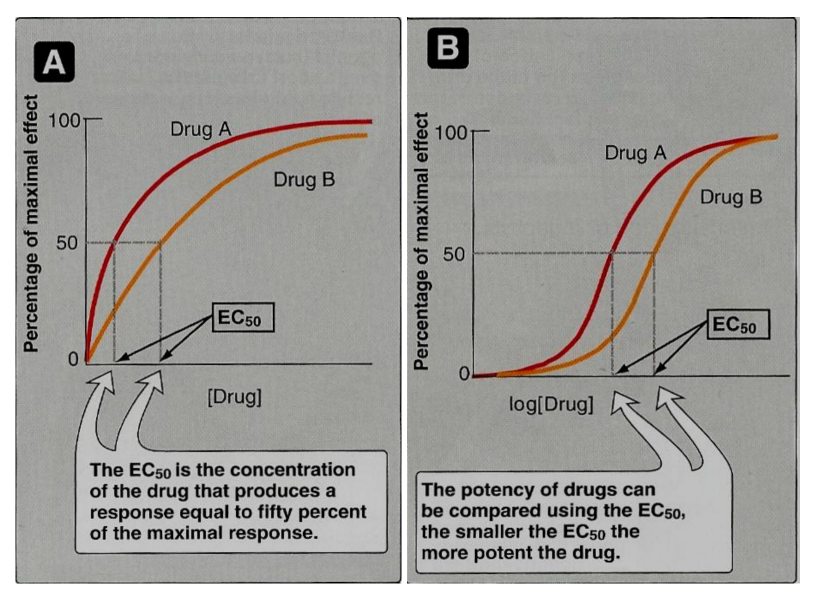
**Graded dose-response curves are used to determine:** 

**Potency:** the concentration of the drug required to produce a specified response (**50% of the maximal response = EC\_{50}**).

**Potency** of drugs can be compared using  $EC_{50}$ , The smaller the  $EC_{50}$ , the more potent the drug.

Potency is inversely proportional to EC<sub>50</sub>

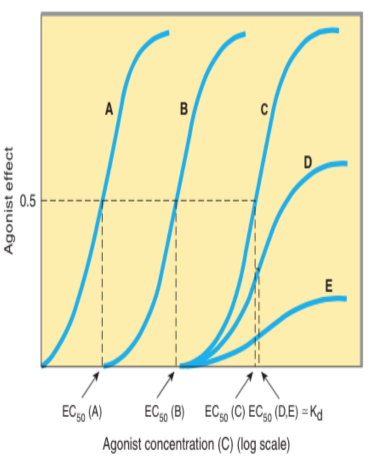




### **Graded Dose-Response Curve**

Which of the following curves represent the least potent drugs ?

Which of the following drugs have the lowest efficacy ?

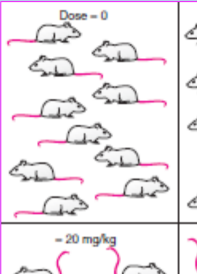


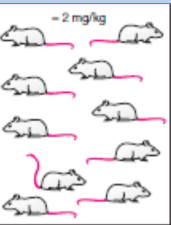
## **Quantal Dose-response Curve**

- Relate drug concentration to % percentage of patients responding (all or none response).
- The response may be therapeutic response, adverse effect or lethal effect.
- o e.g. prevention of convulsion, arrhythmias or death.
- Used to determine
  - ED<sub>50</sub>
  - TD<sub>50</sub> & LD<sub>50</sub>
  - $\circ$  Therapeutic index (**TI**).

#### QANTAL DOSE RESPONSE CURVE

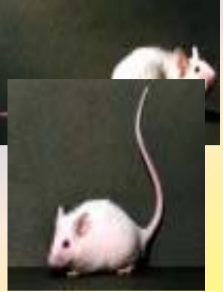
#### **All-non respor**

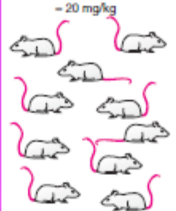


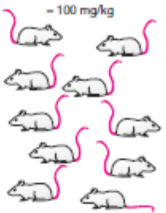




100









**Dose-frequency relationship** 

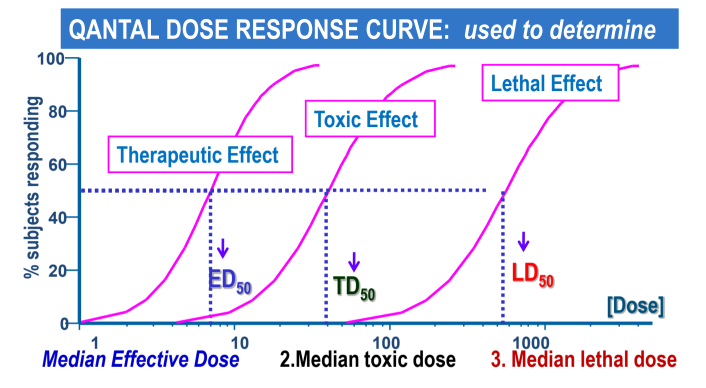
## **Median Effective Dose (ED**<sub>50</sub>):

is a dose of the drug required to produce a therapeutic effect in 50% of individuals.

## Median Toxic Dose (TD<sub>50</sub>):

is the dose of a drug required to produce toxic effects in 50 % of individuals.

**Median Lethal Dose (LD**<sub>50</sub>): is the dose of a drug required to produce death in 50 % of individuals.



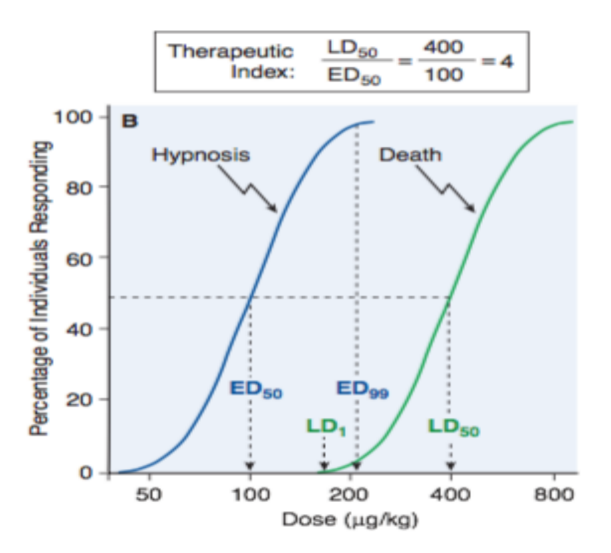
 $ED_{50}$  = 50% of individuals exhibit the specified therapeutic response

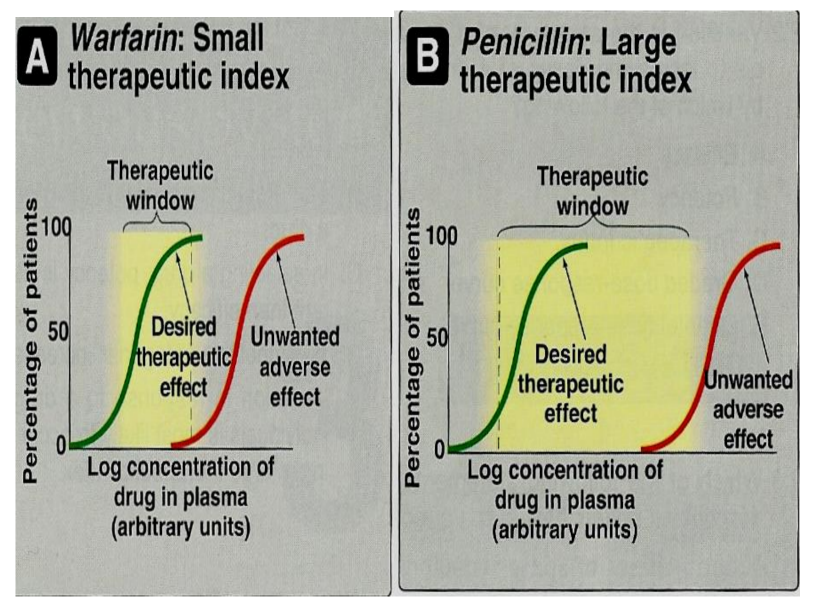
- $TD_{50}$  = 50% of individuals exhibit toxic effects
- LD<sub>50</sub> = 50% of individuals exhibit death

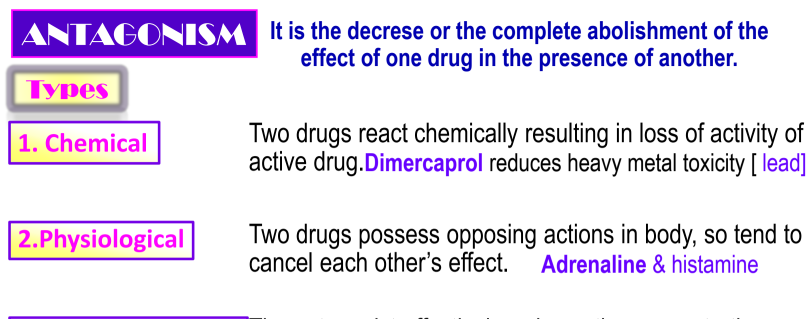
## **Therapeutic Index (TI)**

- Therapeutic index =  $TD_{50}/ED_{50}$  or  $LD_{50}/ED_{50}$ 
  - $TD_{50}$  is the dose that produces a toxic effect in 50% of the population.
  - LD<sub>50</sub> is the dose that is lethal in 50% of the population
  - ED<sub>50</sub> is the dose that produces therapeutic response in 50% of the population
- Is a measure of safety profile
- **High value** = drug with wide margin of safety e.g diazepam, penicillin
- **Small value** = a narrow margin of safety e.g. digoxin, warfarin

# Therapeutic Index







**3. Pharmacokinetic** The antagonist effectively reduces the concentration of the active drug at the site of action **Phenobarbitone** accelerates hepatic metabolism warfarin



## Antagonism

It is the decrease or the complete loss of the effect of one drug by the co-administration (concurrent administration) or combination with another drug.

# **Types of Antagonism**

- ✓ Chemical antagonism.
- ✓ Physiological antagonism.
- Pharmacokinetic antagonism
- Pharmacodynamic antagonism (receptorblockade antagonism)
  - Competitive
    - ✓ Reversible
    - ✓ Irreversible
  - Non-competitive

## Antagonist

### Types

- Physiological antagonist.
- Chemical antagonist.
- Pharmacokinetic antagonist.
- Pharmacodynamic antagonist.

### **Chemical Antagonism**

- Simple chemical reaction between two drugs resulting into loss of activity.
- $\,\circ\,$  No receptors are involved.
- e.g. Dimercaprol used as antidote to reduce heavy metal toxicity (lead toxicity).

#### **Pharmacokinetic Antagonism**

- The antagonist effectively reduces the concentration of the active drug at the site of action.
- e.g. Phenobarbitone accelerates hepatic metabolism of warfarin

## **Physiological Antagonism**

Two drugs act on different receptors to produce opposite physiological effects.

#### e.g. histamine and adrenaline

## **Histamine** $\rightarrow$ vasodilatation ( $\downarrow$ BP) & bronchoconstriction Adrenaline $\rightarrow$ vasoconstriction ( $\uparrow$ BP) & bronchodilation.

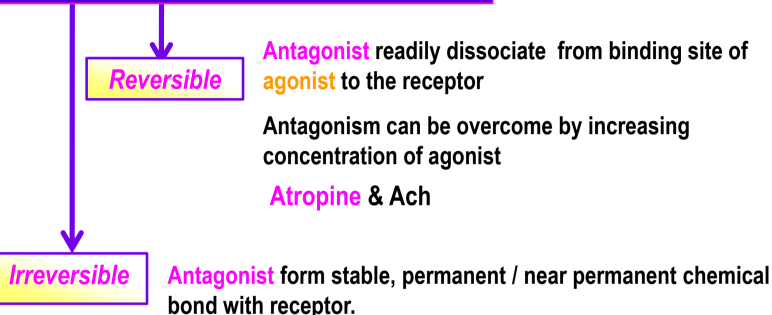
## **\*\***Adrenaline is used in anaphylactic shock

**Pharmacodynamic antagonism** (Receptor-blockade antagonism)

## Types

- Competitive
  - Reversible
  - Irreversible
- Non-Competitive

#### COMPETATIVE ANTAGONISM

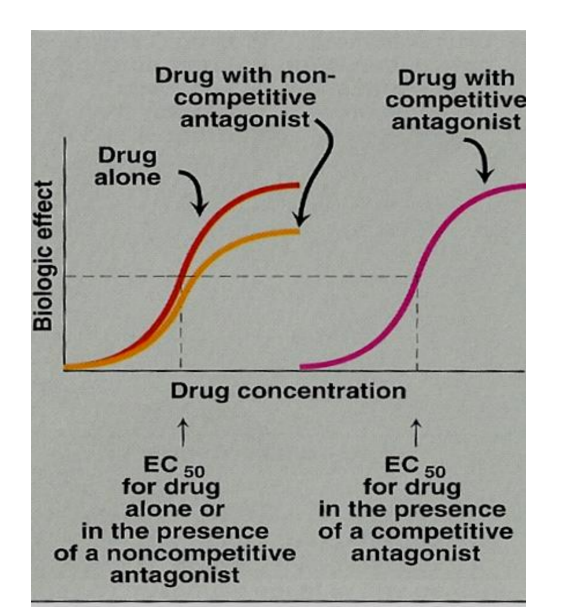


Inactivation lasts for duration of receptor turnover or its denovo synthesis  $\rightarrow$  explains its longevity of action

**Phenoxybenzamine & Noradrenaline** 

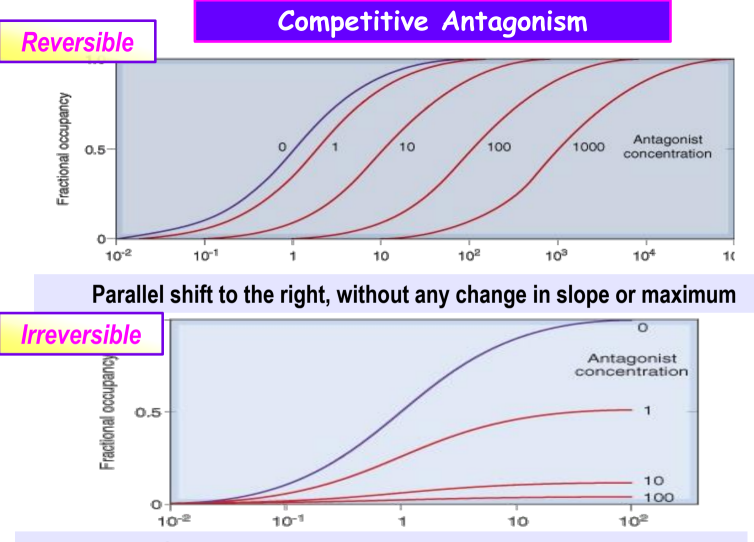
# Pharmacodynamic antagonism Competitive (reversible)

- Two drugs compete for the same receptor (only one is bound).
- The antagonist partially or completely prevents the pharmacological effect of agonist.
- Antagonist dissociate rapidly from receptor.
- Antagonism can be overcome by increasing the concentration of the agonist.
- Parallel shift of the D-R curve to the right, without any change in slope or maximum.
- e.g. acetylcholine and atropine.



# Pharmacodynamic antagonism Competitive (irreversible)

- Two drugs compete for the same receptor.
- Antagonist forms stable, permanent chemical bond with receptor.
- The original response <u>can not be overcome</u> even by increasing the dose of the agonist.
- No parallel shift of D-R curve
- a decrease in slope and a reduced maximal response are obtained.
  - e.g. phenoxybenzamine and noradrenaline.



No parallel shift

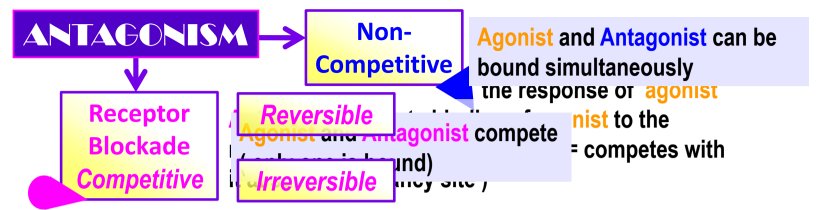
But both a decrease in slope and a reduced maximum are obtained.

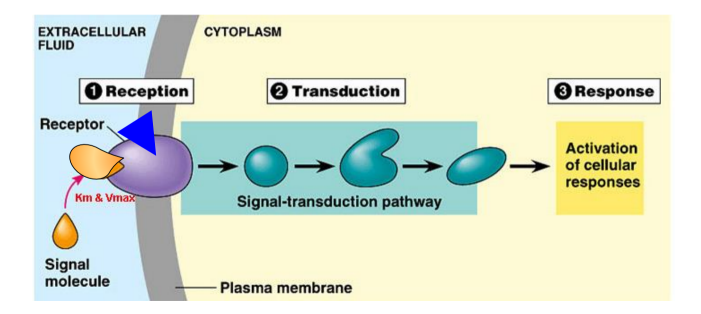
#### **Competitive (reversible)**

- Two drugs compete for the same receptor.
- Antagonist dissociate rapidly from receptor.
- The original response <u>can be</u> <u>obtained by</u> increasing the concentration of the agonist.
- Parallel shift to the right
- No change in slope
- No change in maximum effect
- e.g. acetylcholine and atropine.

#### **Competitive (irreversible)**

- Two drugs compete for the same receptor.
- Antagonist forms stable, permanent chemical bond with receptor.
- The original response <u>can not</u> <u>be obtained</u> even by increasing the dose of the agonist.
- No parallel shift
- A decrease in slope
- Decrease in maximum effect e.g. phenoxybenzamine and noradrenaline.





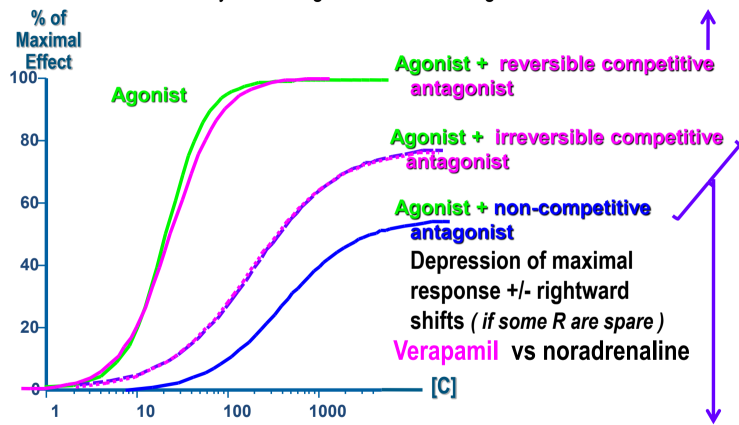
# Pharmacodynamic antagonism Non-competitive

- Agonist and Antagonist can bound simultaneously.
- Antagonist block at some point the chain of events that stimulate the response of agonist.
- Antagonism <u>cannot be overcome</u> by increasing concentration of agonist.

e.g. verapamil and noradrenaline.

#### Competitive vs Noncompetative Antagonism

Antagonism can be overcome by increasing concentration of agonist = **SURMOUNTABLE** 



Antagonism cannot be overcome by increasing concentration of agonist = **NON-SURMOUNTABLE** 



PHARMACOLOGY