

بِسْمِ اللَّهِ الرَّحْمَنِ الرَّحِيمِ

Pharmacodynamics (2)

By

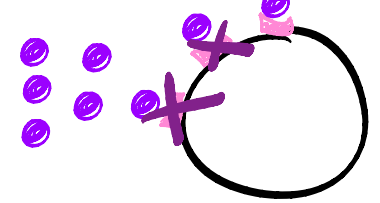
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Receptor regulation

[factor affects receptors]



1- Down-regulation: It is the decrease in number &/or sensitivity of receptors due to prolonged stimulation by agonist.
لمدة طويلة

In some receptors, **continued stimulation of the receptors** with the same dose of **agonist** results in decreased synthesis of new receptors or a state **desensitization** (also referred to as adaptation, refractoriness).

Clinically, Further administration of agonist will produce little effects (**pharmacodynamic tolerance**) in all cases of receptor down regulation.

Example of down regulation: repeated use of **β -2 adrenergic receptor agonists** as **bronchodilators** for treatment of **asthma** leads to diminished effect due to decrease number and signaling power of the beta 2 receptors.
التأثير المتناقص

الزيادة تعالج بالنقصان والنقصان يعالج بالزيادة

2- Up-regulation: It is the increase in number &/or sensitivity of receptors due to prolonged inhibition by antagonist.

Prolonged contact of receptors with an antagonist leads to synthesis of more new receptors.

This can explain the worsening of ^{تفاقم} [angina pectoris] ^{الذبحة الصدرية} in some patients following sudden withdrawal of a β -adrenergic receptor antagonist after long exposure. ^{إذا المريض رقت الدواء فجأة مسببة} [adverse effects]

On withdrawal of β -adrenergic receptor blocker; the up-regulated β -adrenergic receptors are excessively stimulated by normal concentration of circulating catecholamines (like adrenaline).

not up because no antagonist

3- The number of certain receptors is regulated by regulatory factors and **hormones** that do not bind to these receptors at all (e.g. **thyroid hormone** excess in patients suffering from hyperthyroidism causes increase the number of cardiac beta- adrenoceptors).

4- The number and function of receptors can be affected by **diseases** like **autoimmune antibodies** which destroy the receptor itself or affect the **coupling efficiency**. Examples: **diabetes**^{type II} due to destruction of insulin receptors; **myasthenia gravis** due to destruction of muscular nicotinic receptors

نعم ، السكري من النوع الأول يعتبر من أمراض المناعة الذاتية (autoimmune disease) . في هذا النوع من السكري ، يهاجم الجهاز المناعي عن طريق الخطأ خلايا بيتا المنتجة للأنسولين في البنكرياس ويدمرها ، مما يؤدي إلى نقص الأنسولين وارتفاع مستويات السكر في الدم .

أما السكري من النوع الثاني فهو ليس من أمراض المناعة الذاتية ، بل يرتبط بشكل أكبر بعوامل مثل السمنة ، والنظام الغذائي ، ونمط الحياة ، ويحدث عندما تصبح خلايا الجسم مقاومة لتأثيرات الأنسولين أو عندما لا ينتج الجسم ما يكفي من الأنسولين .

دراسة وظائف البداية ثم نقل
الفاصلية

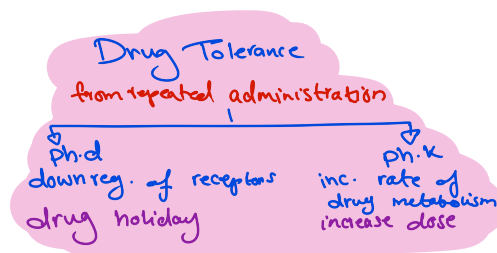
Drug Tolerance

Following the **repeated administration** of some drugs, **the intensity of response** may **decrease** during the course of therapy. Tolerance may be:

- 1- "**pharmacodynamic**" due to **downregulation of receptors** (examples include beta 2 agonists in treatment of asthma and most addicting drugs). **A drug holiday** (to stop drug administration for some time) and then to reuse it may solve the problem.
- 2- "**pharmacokinetic**" tolerance occurs due to **increased rate of drug metabolism** such as phenobarbital tolerance. In this condition, **increasing the dose** of the drug is required to get the same response.

When **tolerance happens rapidly** after the first few doses, is called "**Tachyphylaxis**"

الدكتور ما صدر نوع معين من tolerance



Cross tolerance: *two drug similar chemically or pharmacology → if tolerance happened to one of them, the second will be tolerant to.*

Development of tolerance to a chemically or pharmacologically related drugs;

Examples:

مواد افينونية

☐ *مواد افينونية* Opioids tolerance (Morphine/heroin).

☐ Benzodiazepines tolerance (Diazepam/lorazepam)

☐ Barbiturates tolerance (phenobarbitone/thiopental).

☐ CNS depressant tolerance (alcohol/anesthetics/opioids/barbiturates)

Drug intolerance

حساس جداً للجرعات الصغيرة

- It is the great susceptibility to the drug action e.g., low dose of aspirin (1/2 tablet) may be fatal in some asthmatic patients due to intolerance.

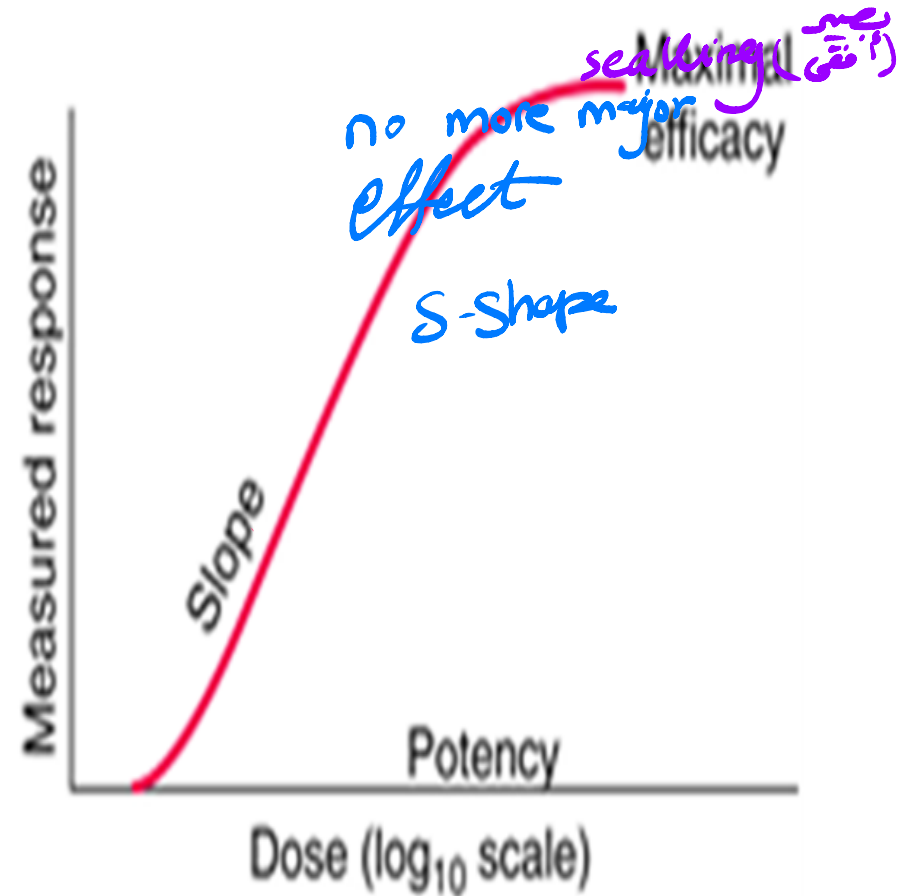
جرعة ايدرا تاكول
التي
B. conc.

Dose-response curve

dose ↑ response ↑ due to number of receptors.

It is an S-shape curve, and it shows a relation between doses (concentrations) and the corresponding responses using a logarithmic scale. The log scale allows a wider display of doses (concentrations) on the dose (concentration) axis.

Normally we plot data as response versus log [Dose].



Shape → S-shape

Scale → logarithmic

wider display of
doses
concentration

يمكن قياسها

Types of dose response curves

1- **Graded** (quantitative): measured on a continuous scale e.g. blood pressure value, blood glucose value, heart rate, etc.

2- **Quantal** (qualitative): All or non-response. Like occurrence of deaths, convulsions or arrhythmias. In this case the response is measured by the percentage of occurrence of these events (e.g., dose 1 caused 10% convulsions, dose 2 caused 22 % convulsions).

يتم قياس الاستجابة كنسبة حدوث هذه الأحداث (جرعة واحدة نسبت بـ 10٪ من التجارب)

Characteristics of graded dose-response curve

From the graded dose response curve, four characteristic variables can be identified: potency, efficacy, slope and biological variation.

بمفرق مع شركات تصنيع الأدوية. يشير هذا إلى أنها potency

(1) Potency:



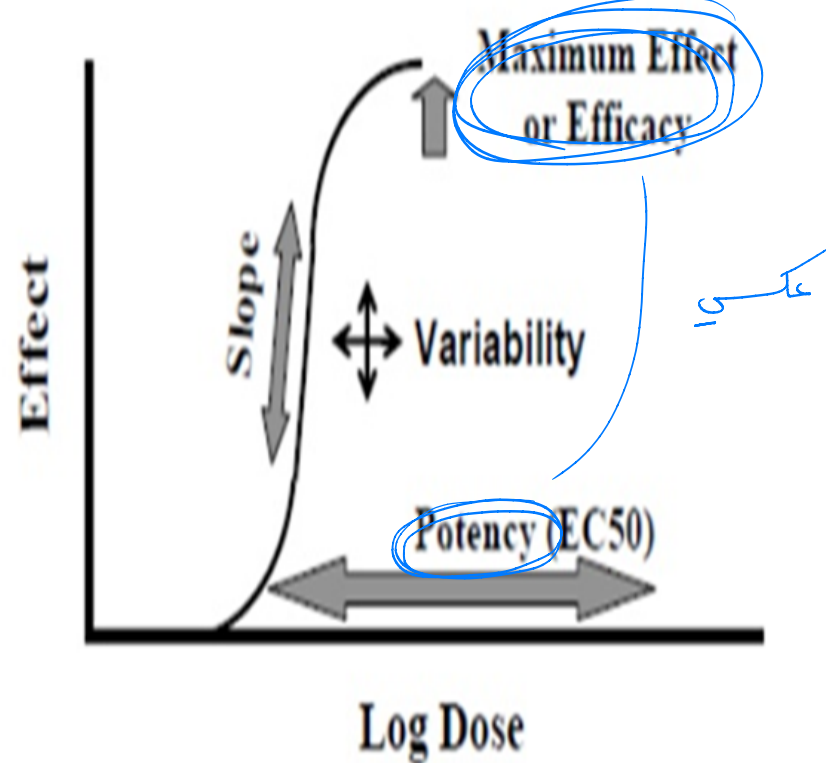
[digoxin]
from low ED₅₀
~ 3 or 4 doses
→ toxicity

It is represented by the horizontal axis of the curve (X-axis, dose axis).

- It depends on the affinity of the drug to the receptors.
- The drug is more potent when it gives response in a smaller dose.
- ED₅₀ can be used for potency comparison.

$ED_{50} \downarrow$ affinity \uparrow

Median Effective Dose "ED₅₀": in graded dose response curve, ED₅₀ is the dose which produces 50 % of maximum response

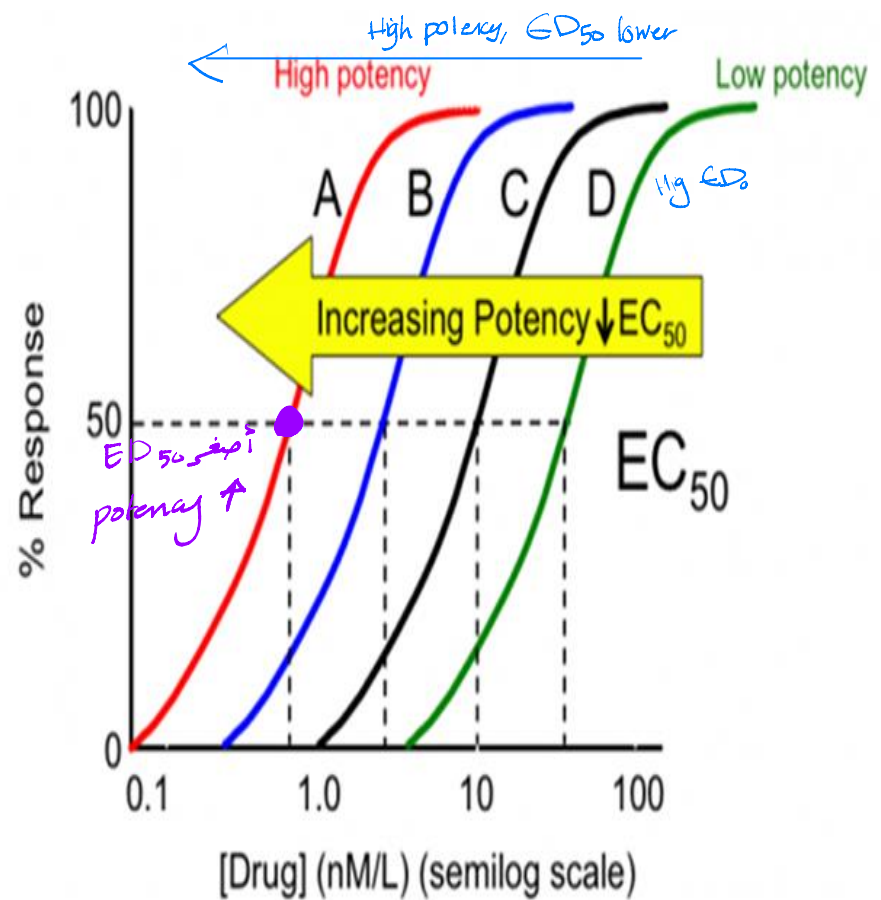


Importance of potency:

- When the potency of a drug is so low, very large dose (which may be not practical) is needed.
- When the potency of a drug is so high, any **slight increase** in the dose, can lead to toxicity.
- In pharmaceutical manufacturer, the production of **potent drugs** is considered as more economic as it consumes **less amount of the drug**.

Example:

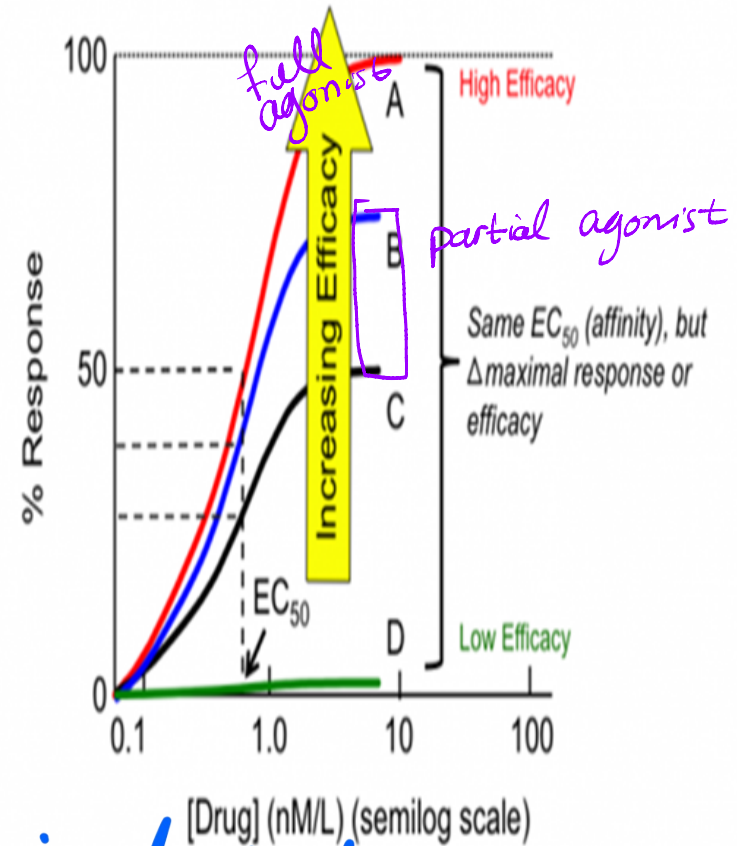
- the median effective dose (ED_{50}) or the median effective concentration (EC_{50}) is used to compare between the potency of different drugs.
- the lower the ED_{50} , the more potent drug.
- Drug A > B > C > D in potency
- EC_{50} of drug D > C > B > A



2) Efficacy:

aspirin & Morphine
من أسيل ساليسيليك و مورفين

- It is the maximal ceiling effect of the drug, after which there is no increase in the response even if the dose is increased (due to full occupancy of the receptors), but toxicity appear.
- It is the most important character of the drug clinically.
- Efficacy depends on the intrinsic activity of the drug.



no increase in R, but there is toxicity
↳ response

Examples:

i. Morphine can control any type of pain, but aspirin control only mild pain as the efficacy of morphine is higher than that of aspirin.

ii. Furosemide has efficacy higher than thiazide diuretics, so furosemide produces diuresis more than thiazide diuretics.
بوجنه مر
تبول

3) Slope:

اگه خطدار

- It is the central linear part of the curve.
- If the drug has steep slope, this indicates that the ratio between the therapeutic dose and the toxic dose of this drug is low (drugs with narrow safety margin).
- If the drug has flat slope, this indicates that the ratio between the therapeutic dose and the toxic dose of this drug is high (drugs with high safety margin).

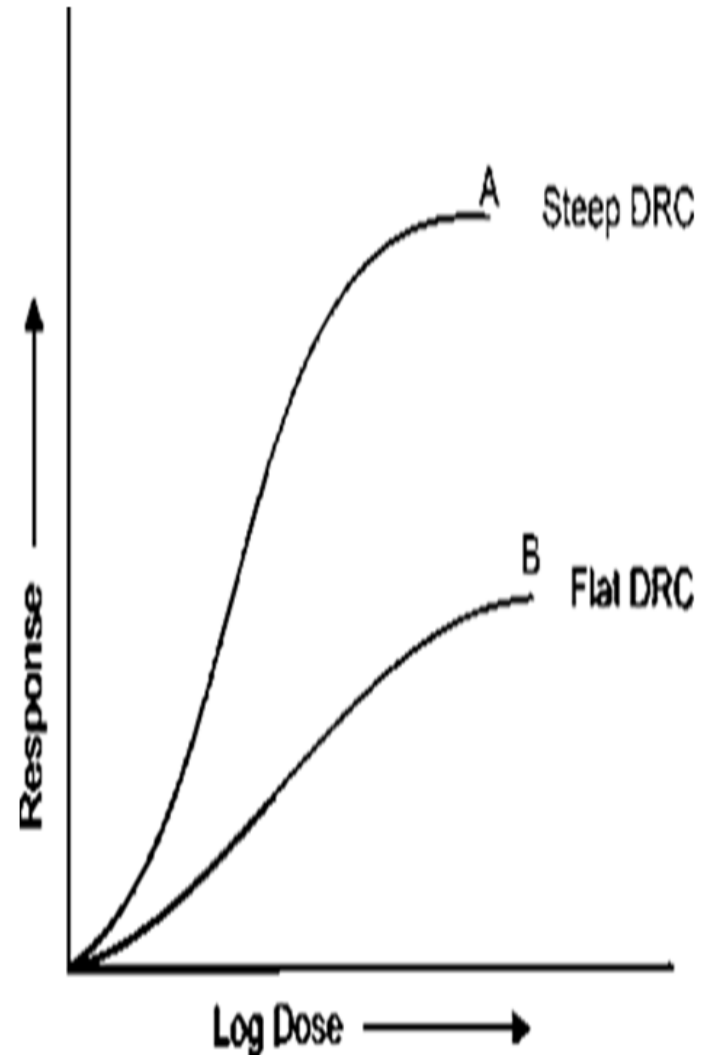
Examples:

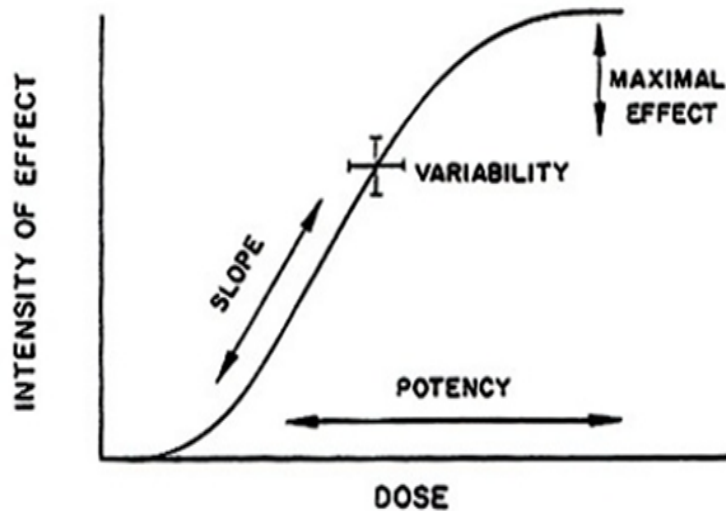
Barbiturates has steep slope while

benzodiazepine has flat slope, so

benzodiazepines are considered safer than

barbiturates.

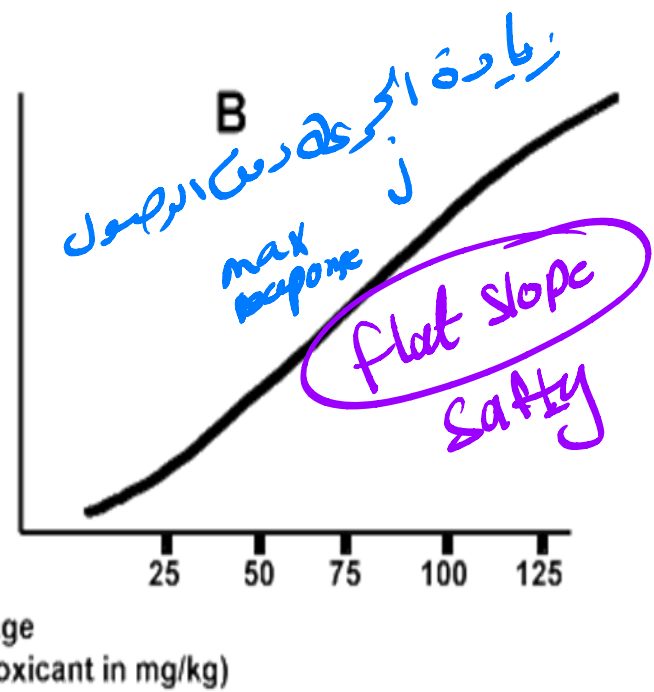
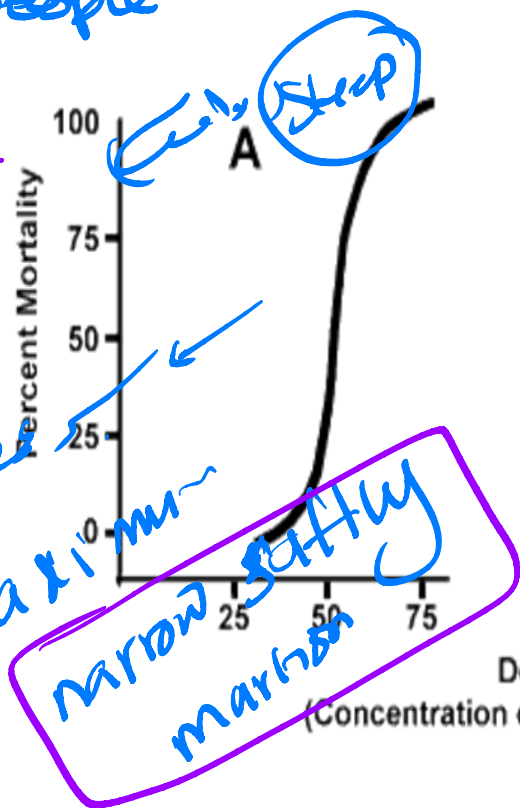




variation
 Same person
 different people

تغيير بدرجة
 maximum response
 toxicity

step
 maximum
 narrow safety margin



بين الأشخاص مختلفين
الشخص نفسه في أوقات مختلفة

4) Variability of Effects:

- Variability is another characteristic of the dose response curve, along with slope, efficacy and potency.
- A given drug at a given dose level will have different effects on different individuals and different effects on the same individual at different times.
- Causes of variability includes age, sex, race, co-medical conditions, past drug use, time of day, menstrual cycle, other drugs taken, etc. *الجزء*

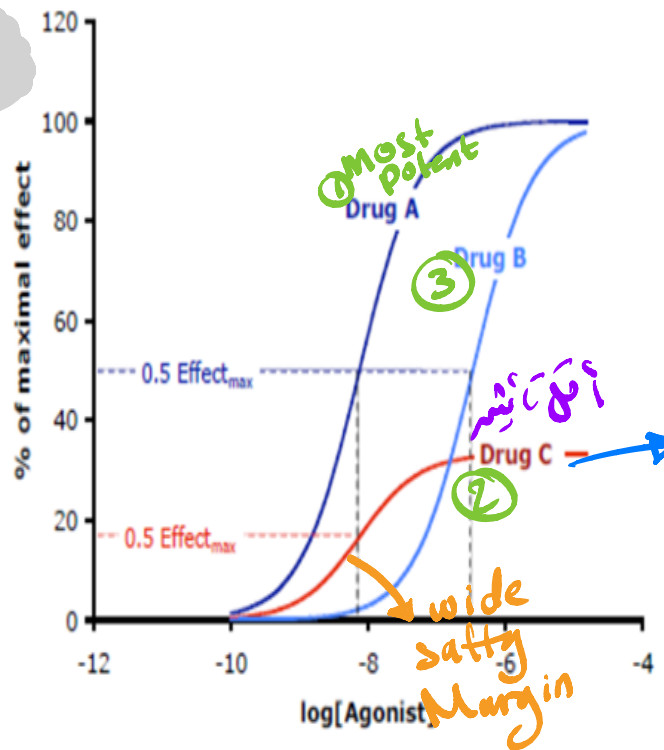
Summary of graded dose response curve

1. **Potency**: *x-axis* a comparative measure which refers to the different doses of two drugs needed to produce the same effect. For comparison of potency, ED50 is used.
2. **Efficacy**: *y-axis* is the maximum effect (Efficacy) of a drug. For comparison of efficacy, the maximum ceiling effects are compared.
3. **Slope**: *for safety* The slope of the central linear part of the curve. When the slope is steep, it means that the ratio between the therapeutic dose and toxic dose of the drug is low i.e., the drug has a narrow margin of safety.
4. **Variability of effects** among different or the same individual at different times.

(Trimethylxanthine) *أكثر دواء صو*
Caffeine

لا يوجد بينهم علاقة

Potency and efficacy



Demonstration of different potencies of drugs (A,B,C) that produce the same effect. In this figure, Drugs A & B have the same efficacy. Drug A has greater potency than B or C because the dose of B or C must be larger to produce the same effect as A. Although Drug C has lower efficacy than B, it is more potent than B at lower drug concentrations.

2- Quantal dose response curve

All or non- response are measured in percentage. Like occurrence of deaths, convulsions or arrhythmias. It is used to determine the dosage causing toxicity and lethality to experimental animals.

- Median effective dose (ED50) of quantal response = dose which produce a specific response in 50% of population.
- Median Toxic dose (TD50) = dose which produce a specific toxicity in 50% of experimental animals.
- Median lethal dose (LD50) = dose which produce death in 50% of experimental animals.

Therapeutic index (TI) → مقياس على الحيوانات

Therapeutic index (TI) is a quantitative measurement of the relative safety of drugs.

It is the ratio of TD50 or LD50 to ED50.

TI = TD50 / ED50

Therapeutic index should be greater than 1.

If TI = 1; the drug is a poison

don't use it therapeutically

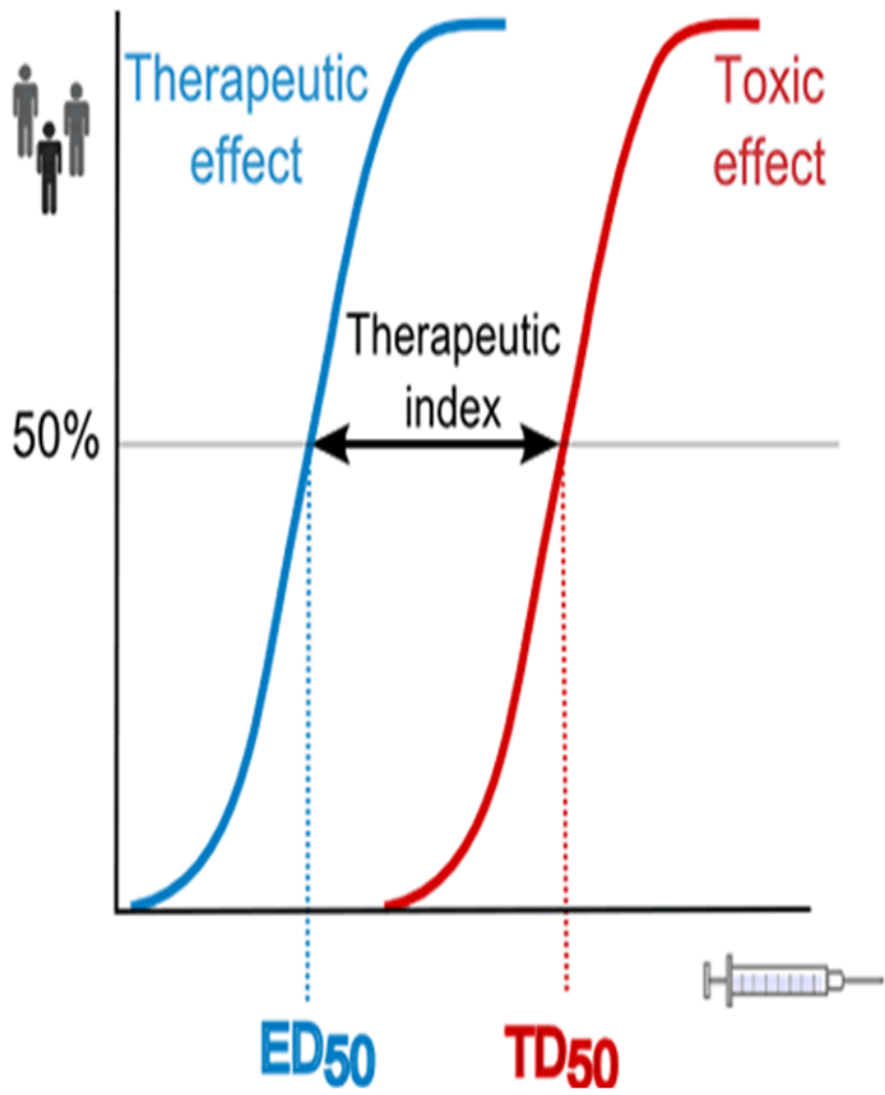
TI = 3 → not safe
TI = 4.5 → safe

بنسبتين 3 و 4.5

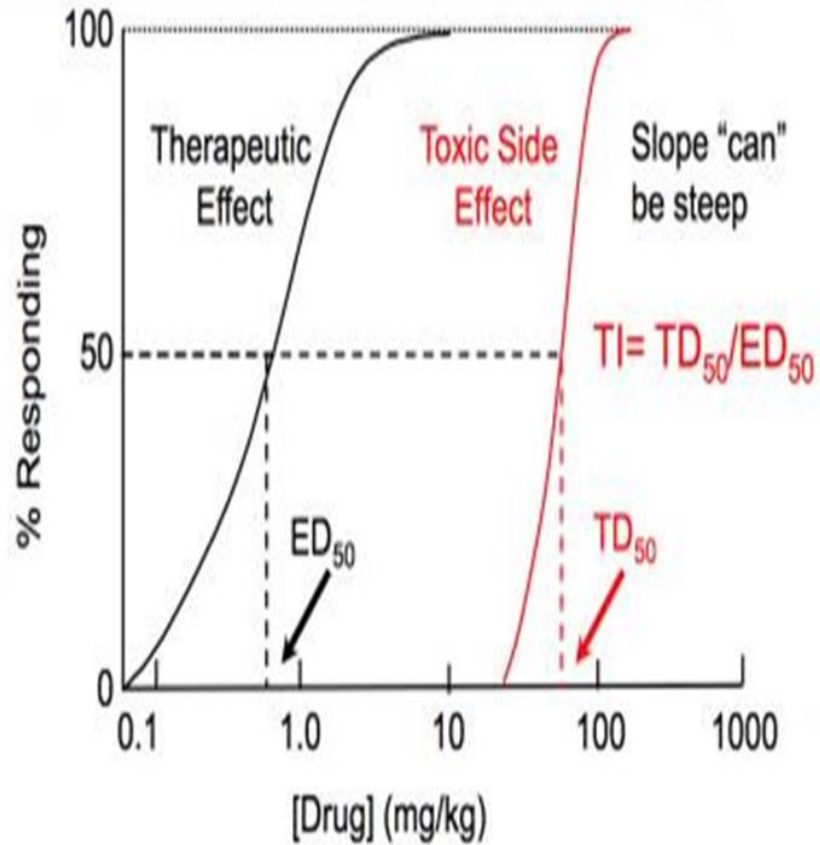
مناسب
ملائق → TI = 1.7 و 2

TI = 2 digoxin

if TI = 1 → its poison
لا يمكن استخدامه كدواء

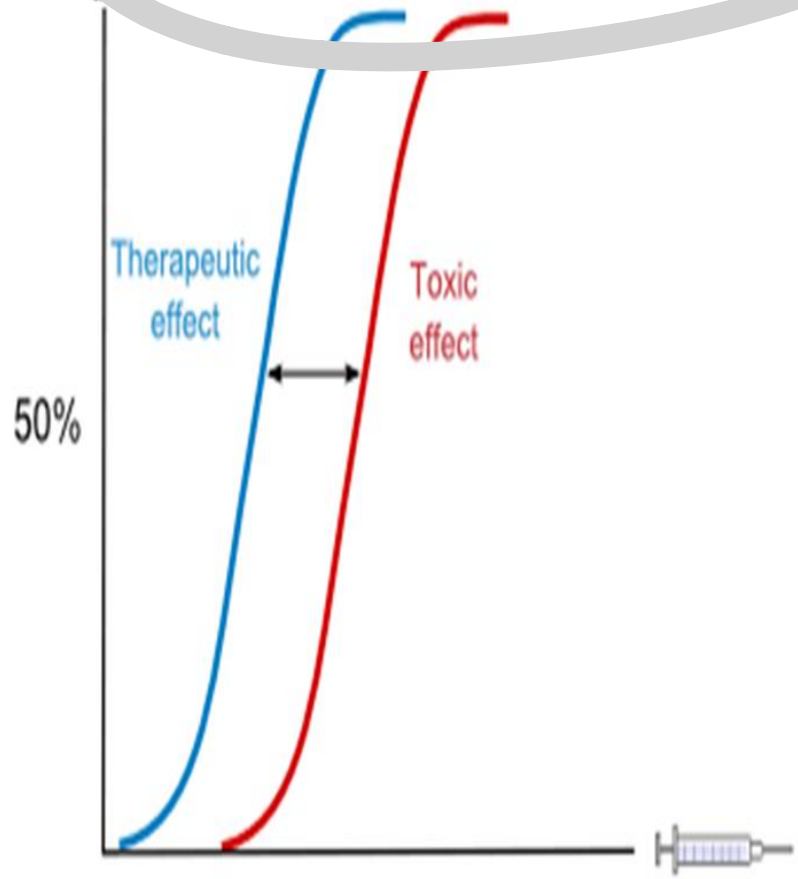


Drug Safety - Therapeutic Index



Narrow TI

Fig 10



Generally, a drug or other therapeutic agent with a narrow therapeutic range (i.e., having little difference between toxic and therapeutic doses) may have its dosage adjusted according to measurements of the actual blood levels achieved in the person taking it. This may be achieved through therapeutic drug monitoring (TDM) protocols.

[●REC]

بروتوكولات مراقبة الدواء العلاجية]

Medication with a small therapeutic window must be administered with care and control, frequently measuring blood concentration of the drug, to avoid harm. Medications with narrow therapeutic windows include digoxin, lithium, aminoglycosides, immunosuppressive drugs, antiepileptic drugs, some anticancer drugs and warfarin.

