

DOSE-EFFECT RELATIONSHIP

The intensity and duration of a drug's effects are a function of the drug dose and drug concentration at the effect site

Frank M. Balis, M.D.

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Monitoring Dose-Effect

- * **Level**
 - **Molecular (e.g, enzyme inhibition)**
 - **Cellular (*in vitro* tissue culture, blood cells)**
 - **Tissue or organ (*in vitro* or *in vivo*)**
 - **Organism**
- * **Endpoint used to measure effect may be different at each level**
- * **Overall effect = sum of multiple drug effects and physiological response to drug effects**

Dose-Effect Endpoints

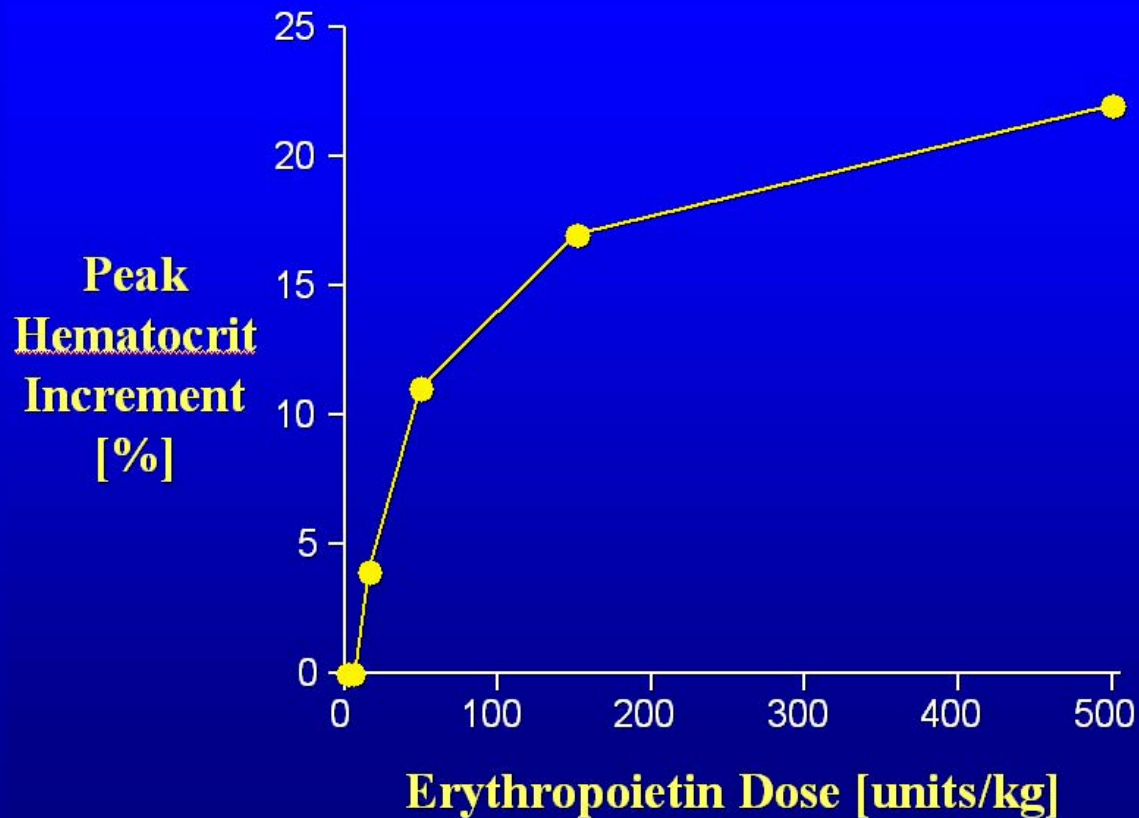
Graded

- * Continuous scale (\uparrow dose \rightarrow \uparrow effect)
- * Measured in a single biologic unit
- * Relates dose to intensity of effect

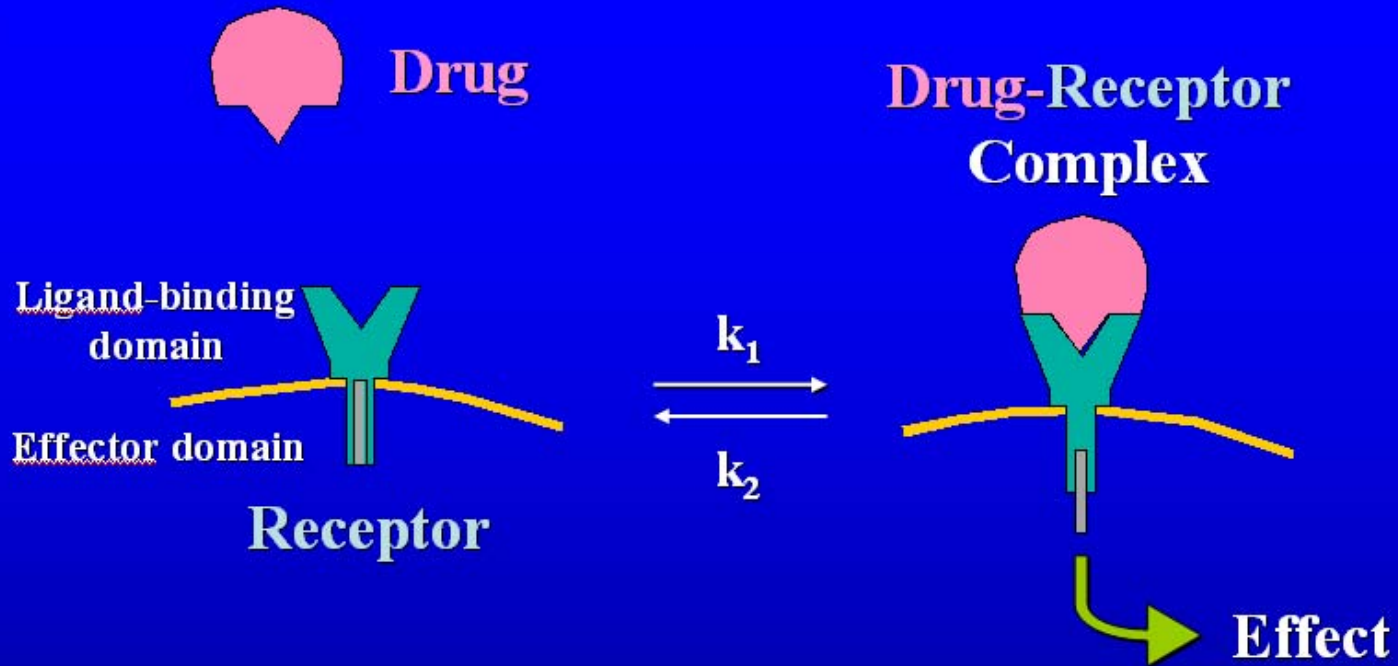
Quantal

- * All-or-none pharmacologic effect
- * Population studies
- * Relates dose to frequency of effect

Erythropoietin and Anemia



Drug-Receptor Interactions



$$\text{Effect} = \frac{\text{Maximal effect} \cdot [\text{Drug}]}{K_D + [\text{Drug}]}$$

$$(K_D = k_2/k_1)$$

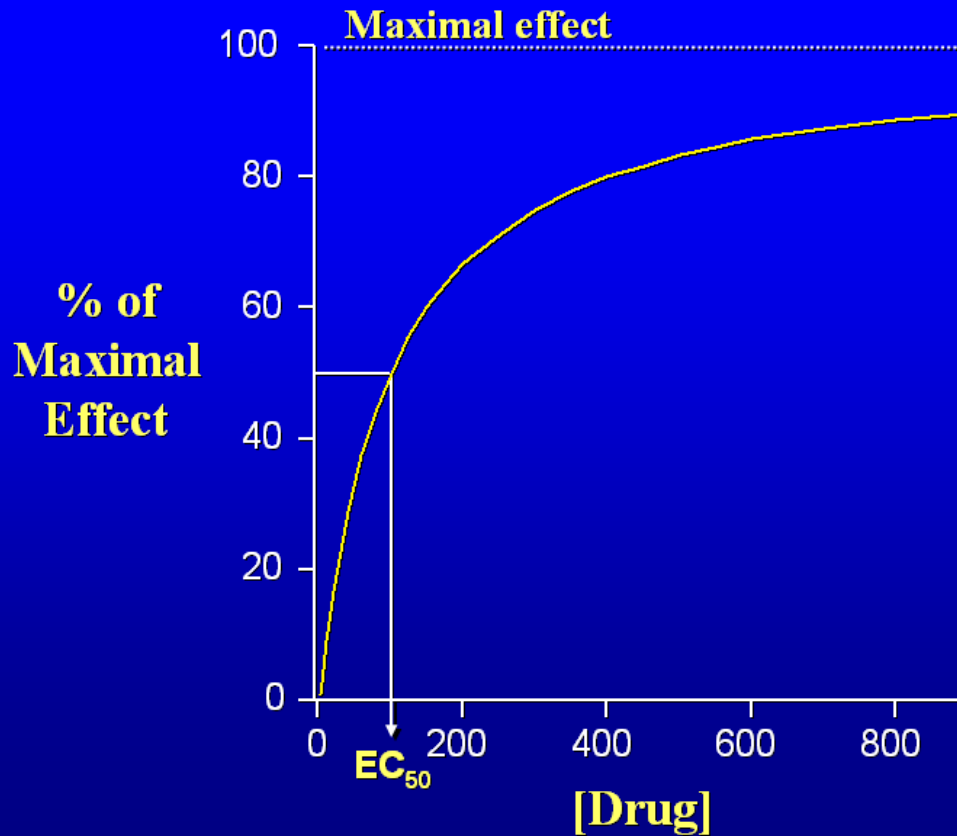
Dose-Effect Relationship

$$\text{Effect} = \frac{\text{Maximal effect} \cdot [\text{Drug}]}{K_D + [\text{Drug}]}$$

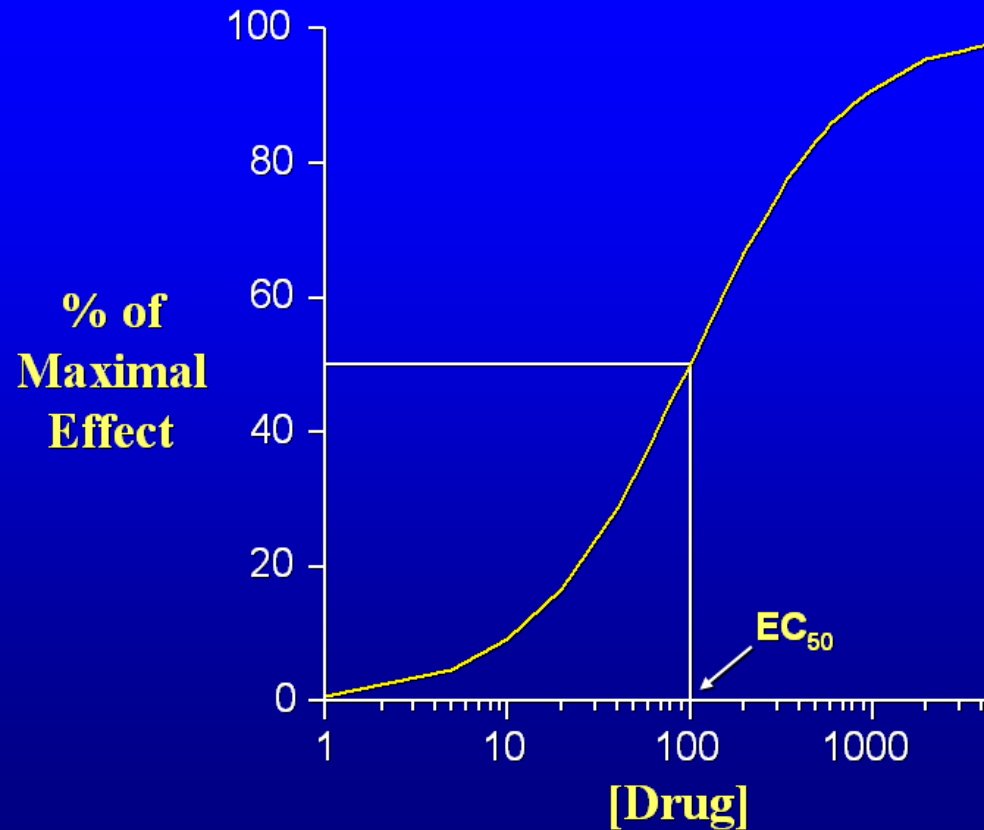
$$\text{Effect} = \text{Maximal effect} \frac{[\text{Drug}]}{K_D + [\text{Drug}]}$$

$$\text{Effect} = \text{Maximal effect} \quad \text{if } [\text{Drug}] \gg K_D$$

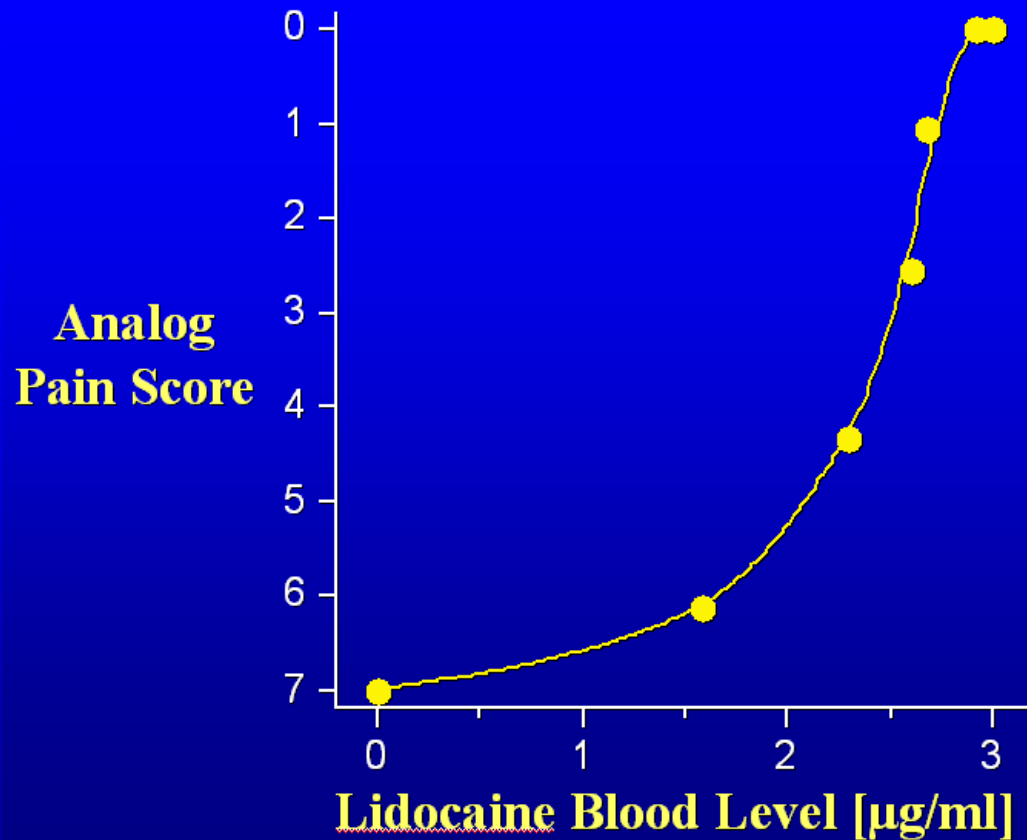
Graded Dose-Effect Curve



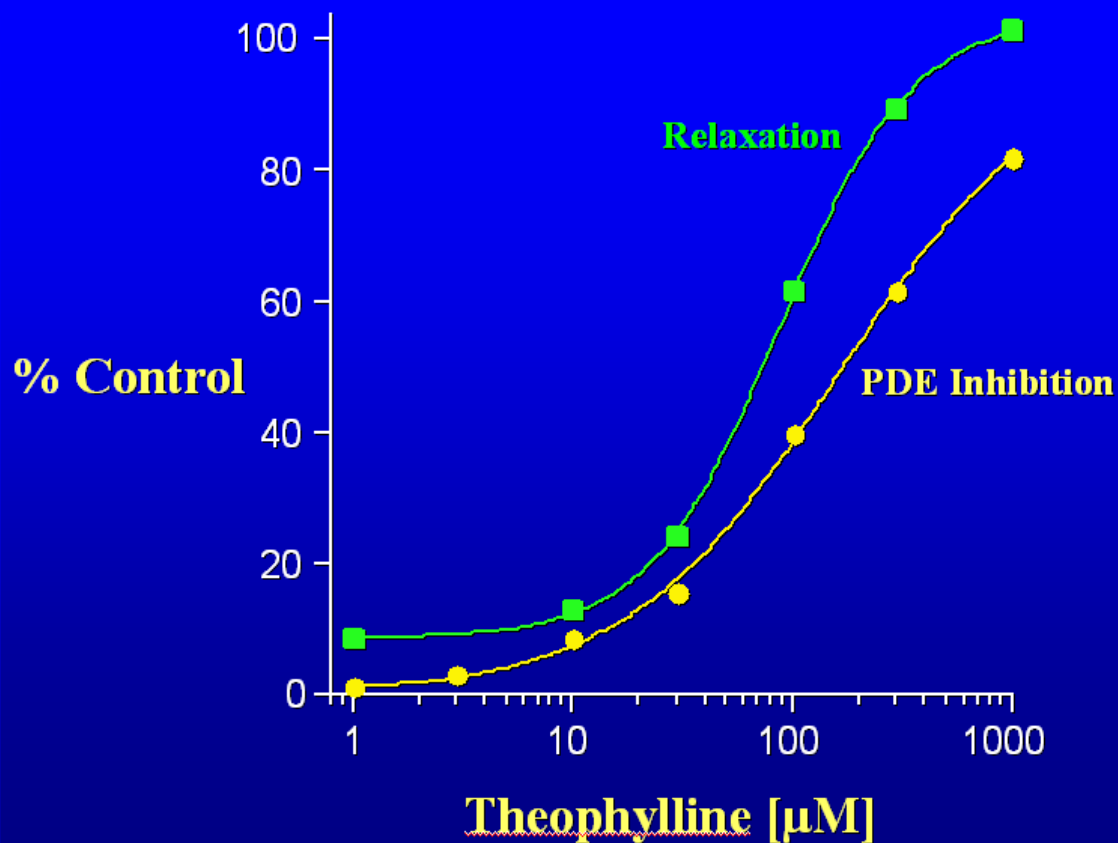
Log Dose-Effect Curve



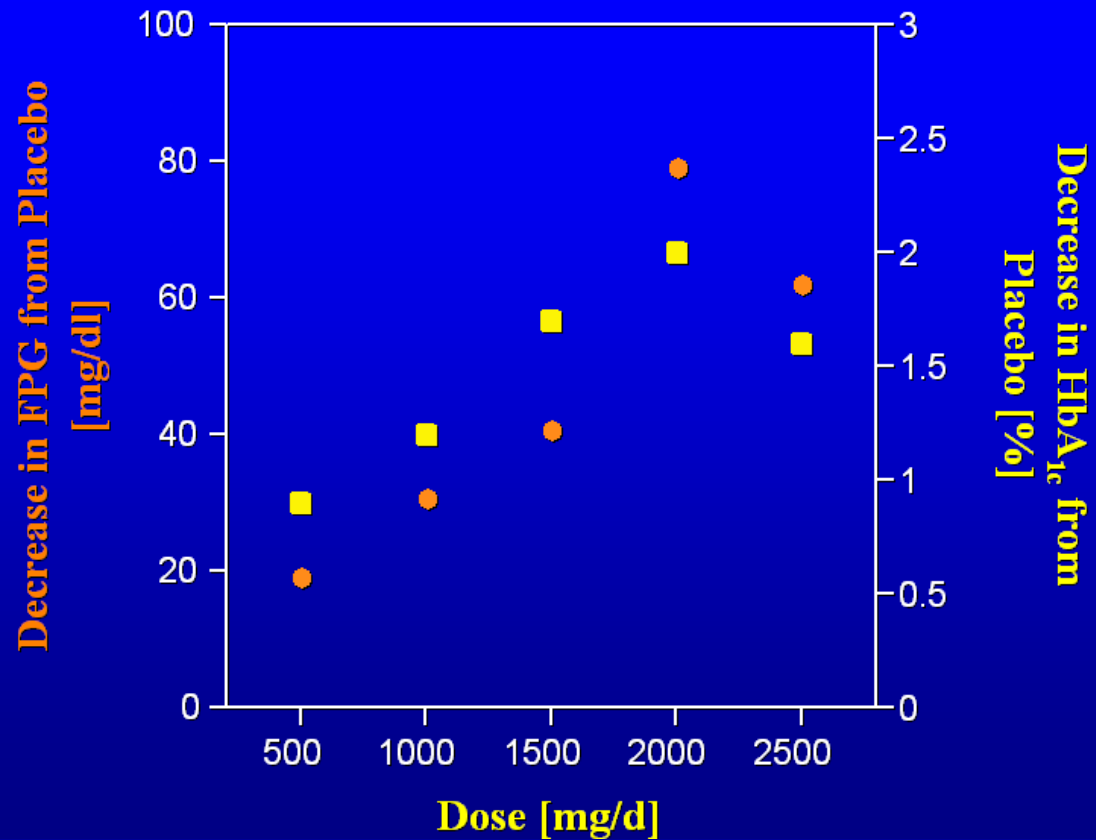
Lidocaine Graded Dose-Effect



Theophylline Dose-Effect



Metformin Dose-Response

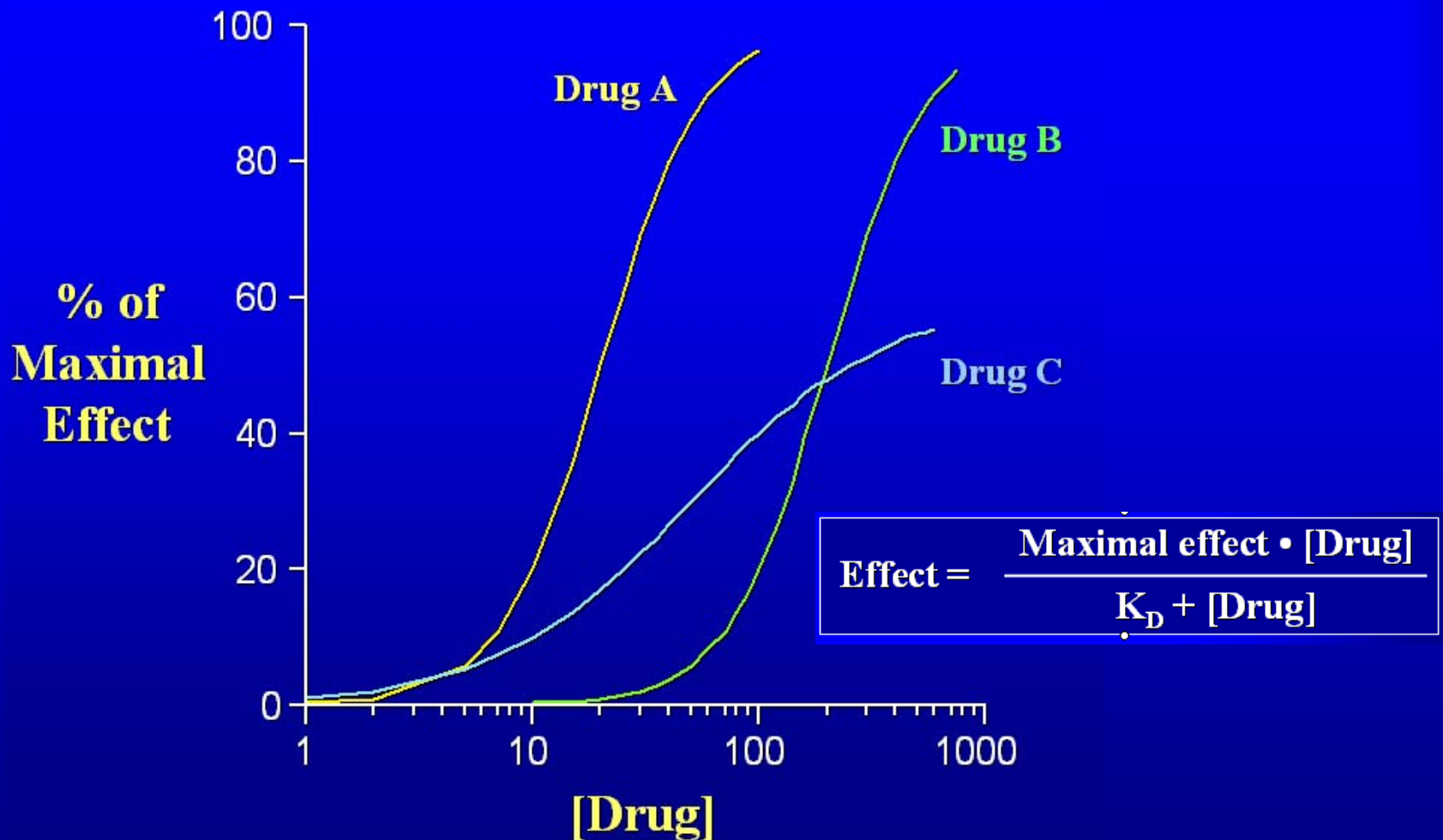


Dose-Effect Parameters

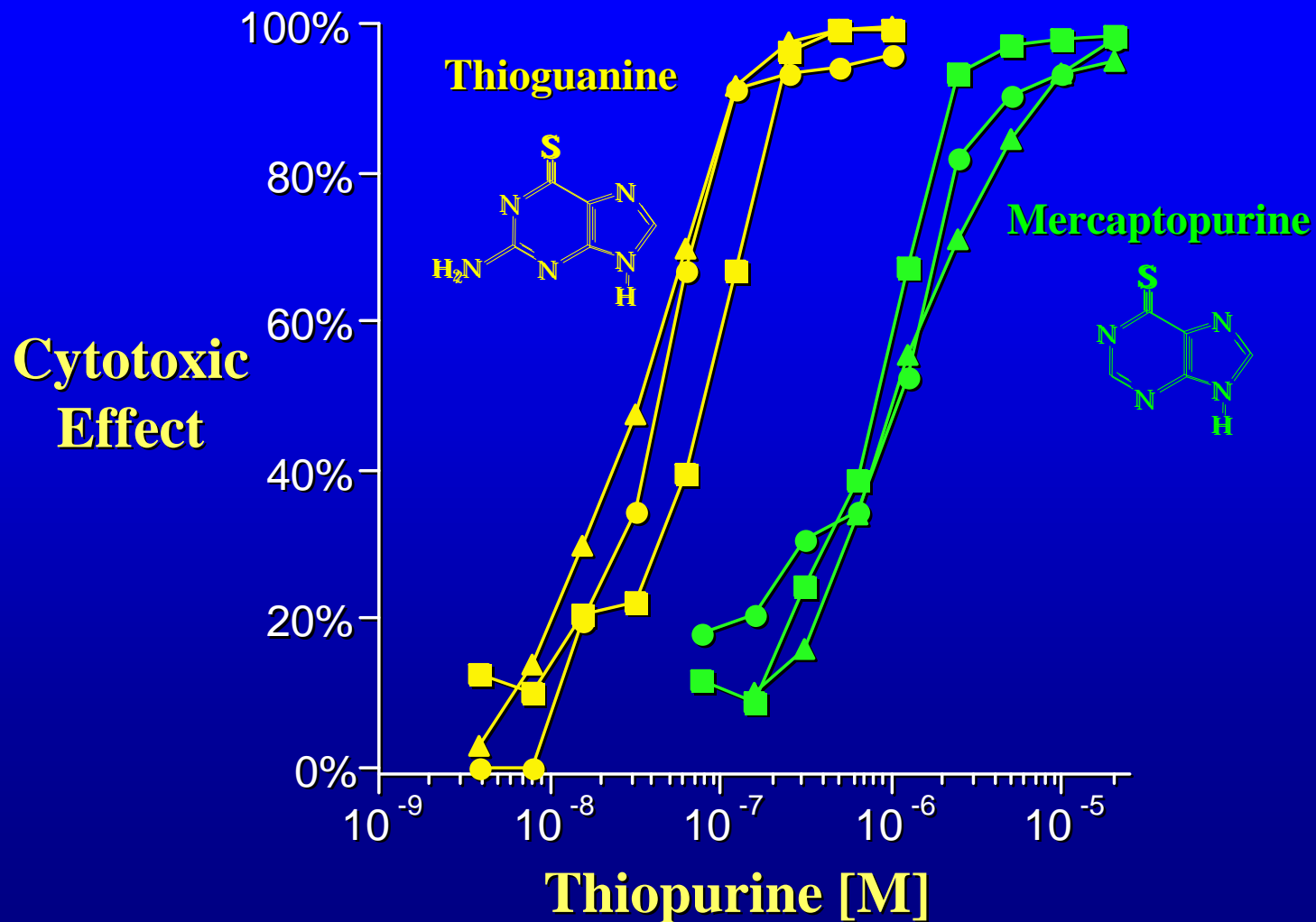
POTENCY: The sensitivity of an organ or tissue to the drug

EFFICACY: The maximum effect

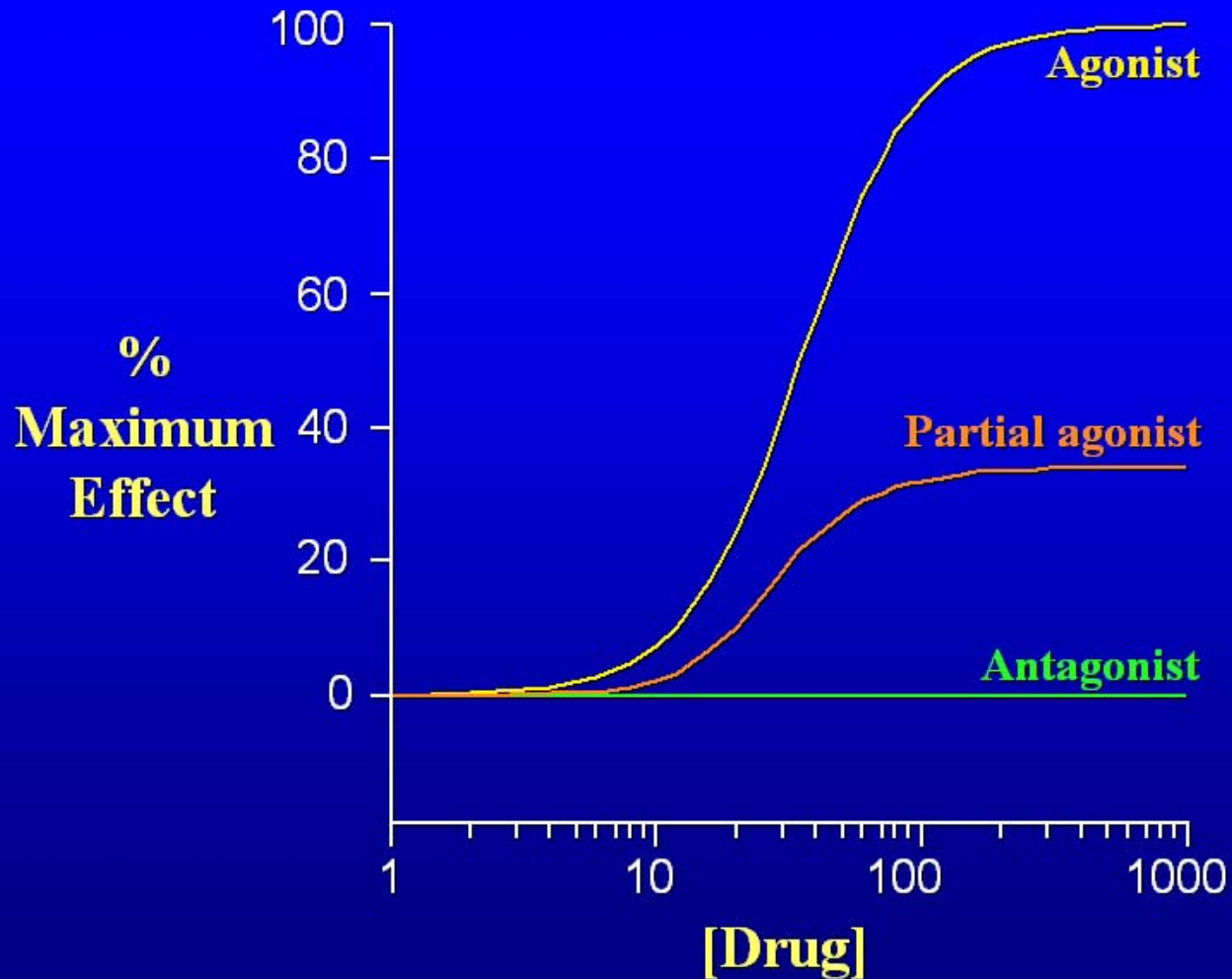
Comparing Dose-Effect Curves



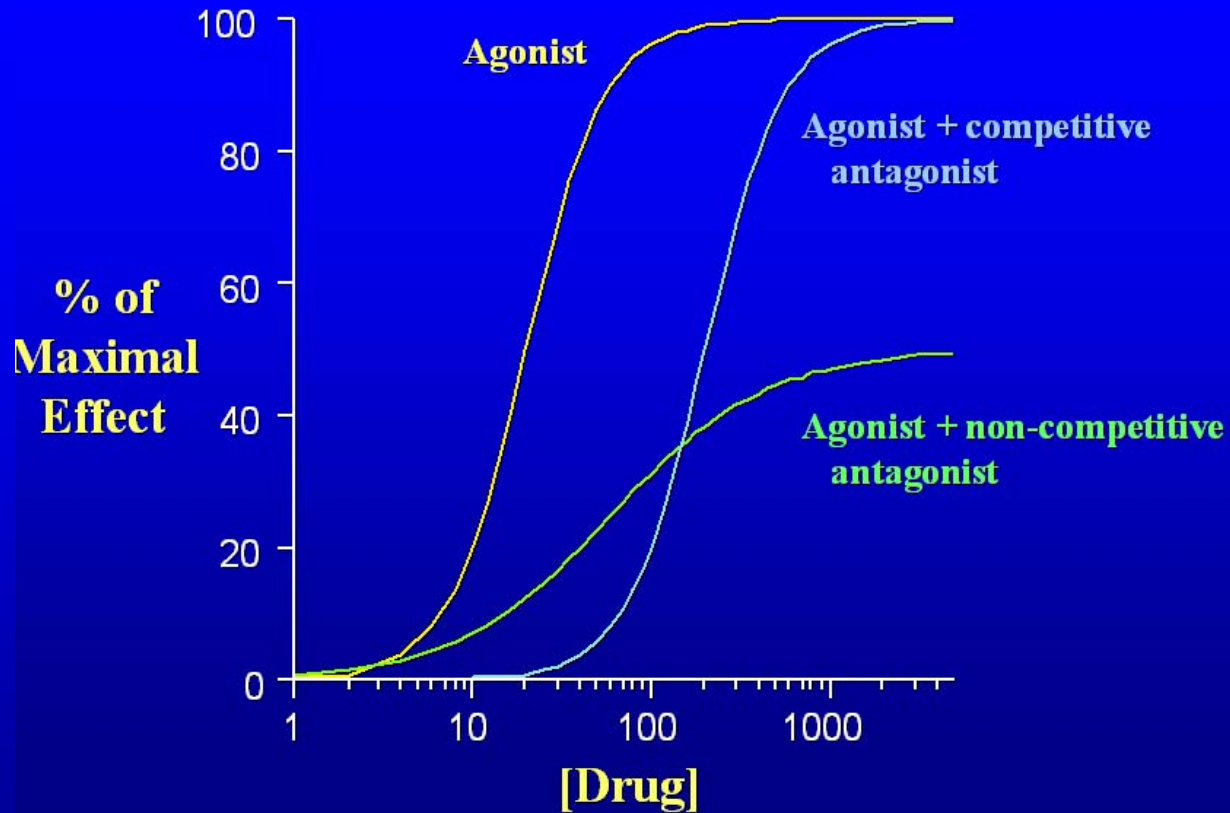
Thiopurine Cytotoxicity



Receptor-Mediated Effects



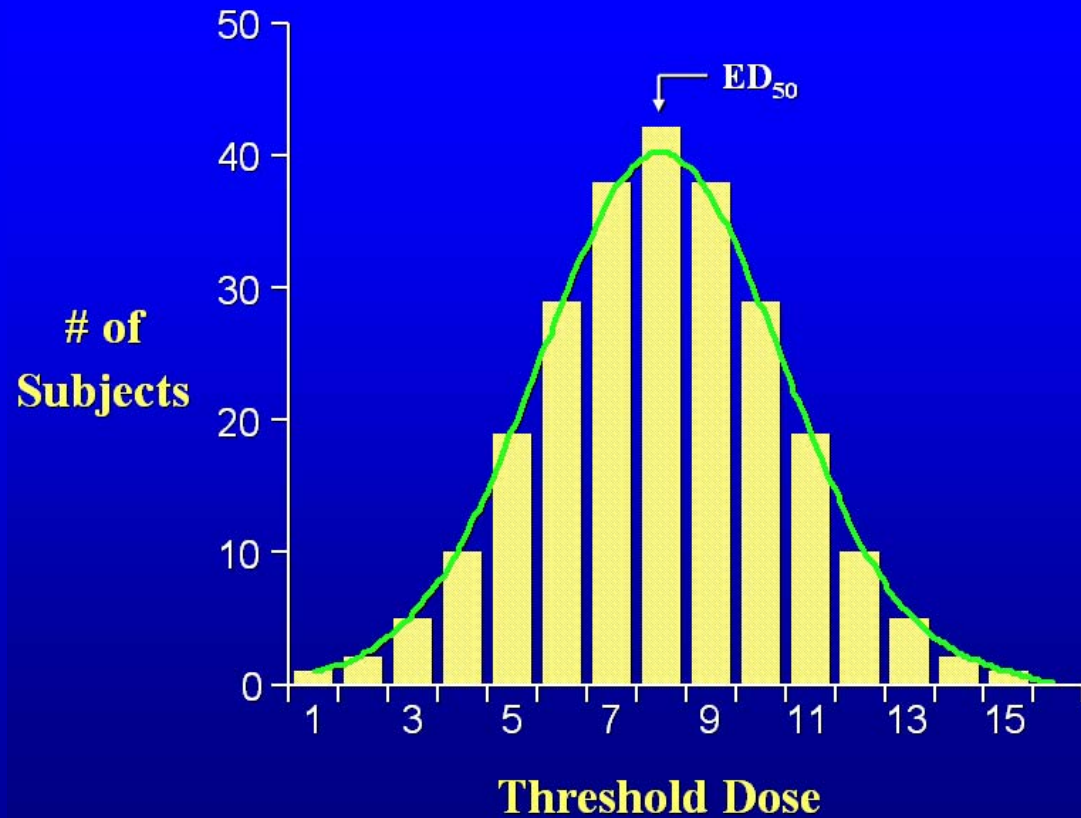
Drug Interactions



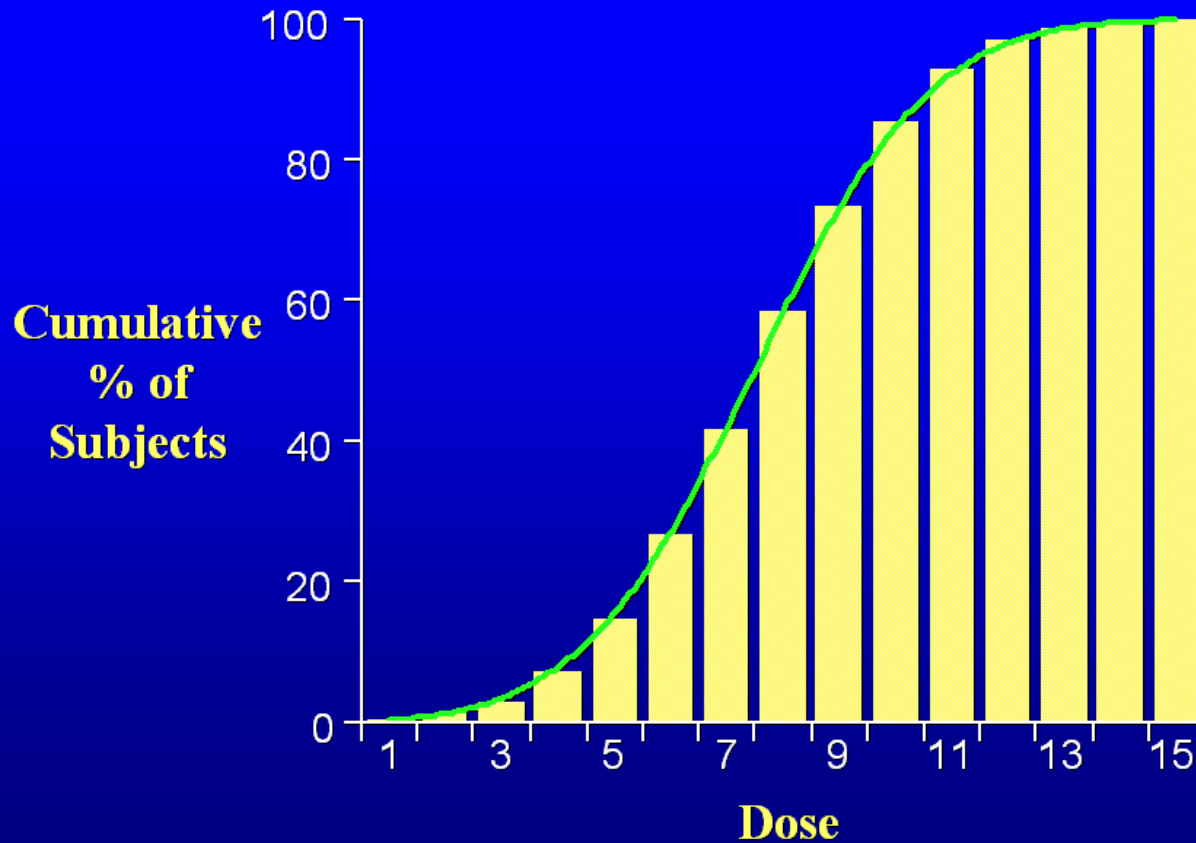
Graded Dose-Effect Analysis

- * Identify the therapeutic dose/concentration**
- * Define site of drug action (receptor)**
- * Classify effect produced by drug-receptor interaction (agonist, antagonist)**
- * Compare the relative potency and efficacy of drugs that produce the same effect**
- * Assess mechanism of drug interactions**

Quantal Dose-Effect Distribution



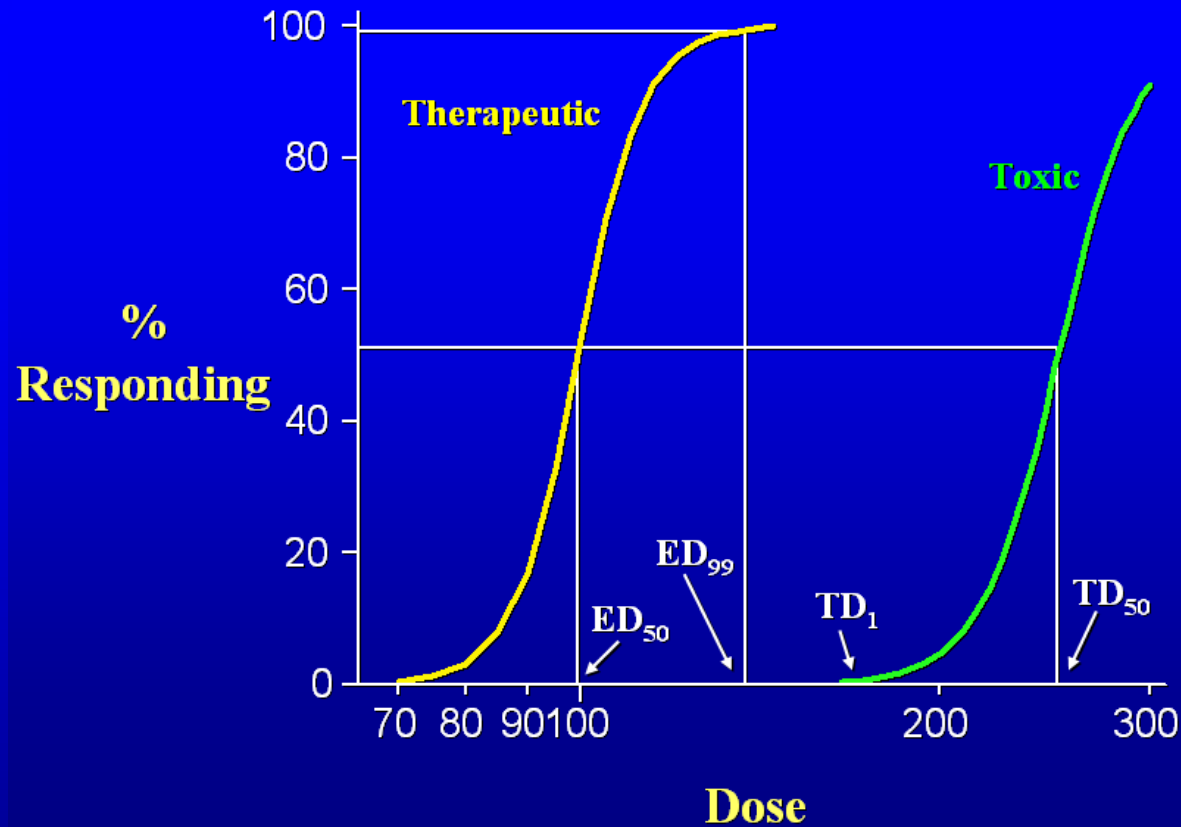
Cumulative Dose-Effect Curve



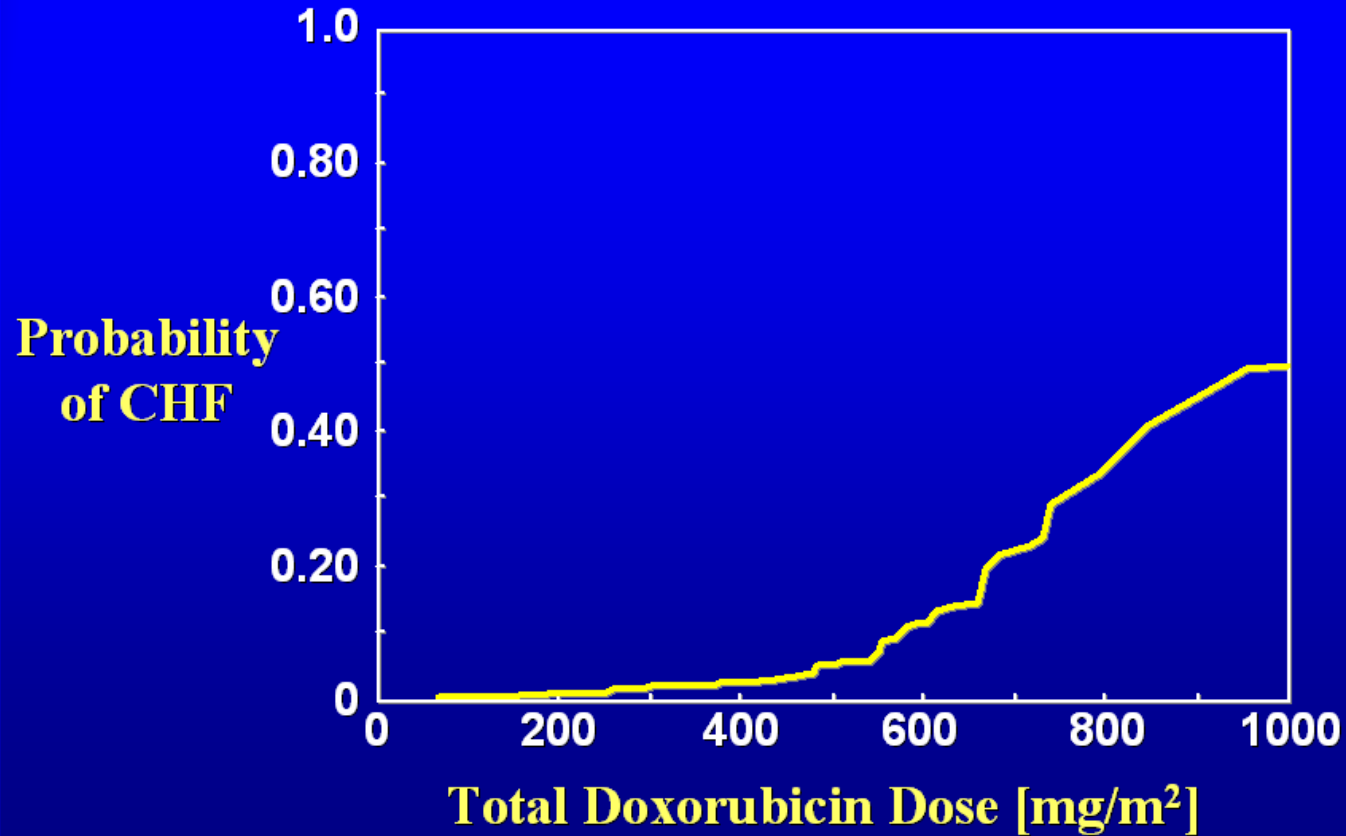
Cumulative Dose-Effect Study

Dose Level	No. of Subjects	No. Responding	% Response
1	10	0	0
2	10	1	10
3	10	3	30
4	10	5	50
5	10	7	70
6	10	8	80
7	10	9	90
8	10	10	100

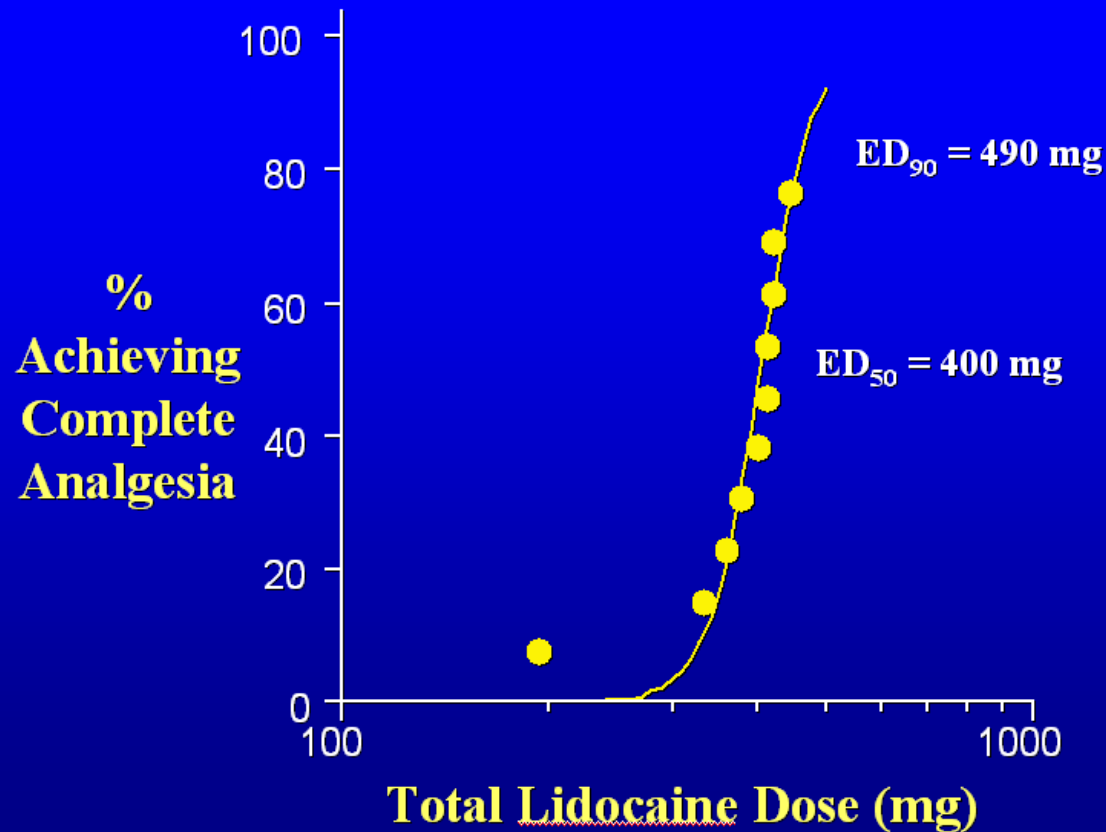
Therapeutic and Toxic Effects



Doxorubicin Cardiotoxicity



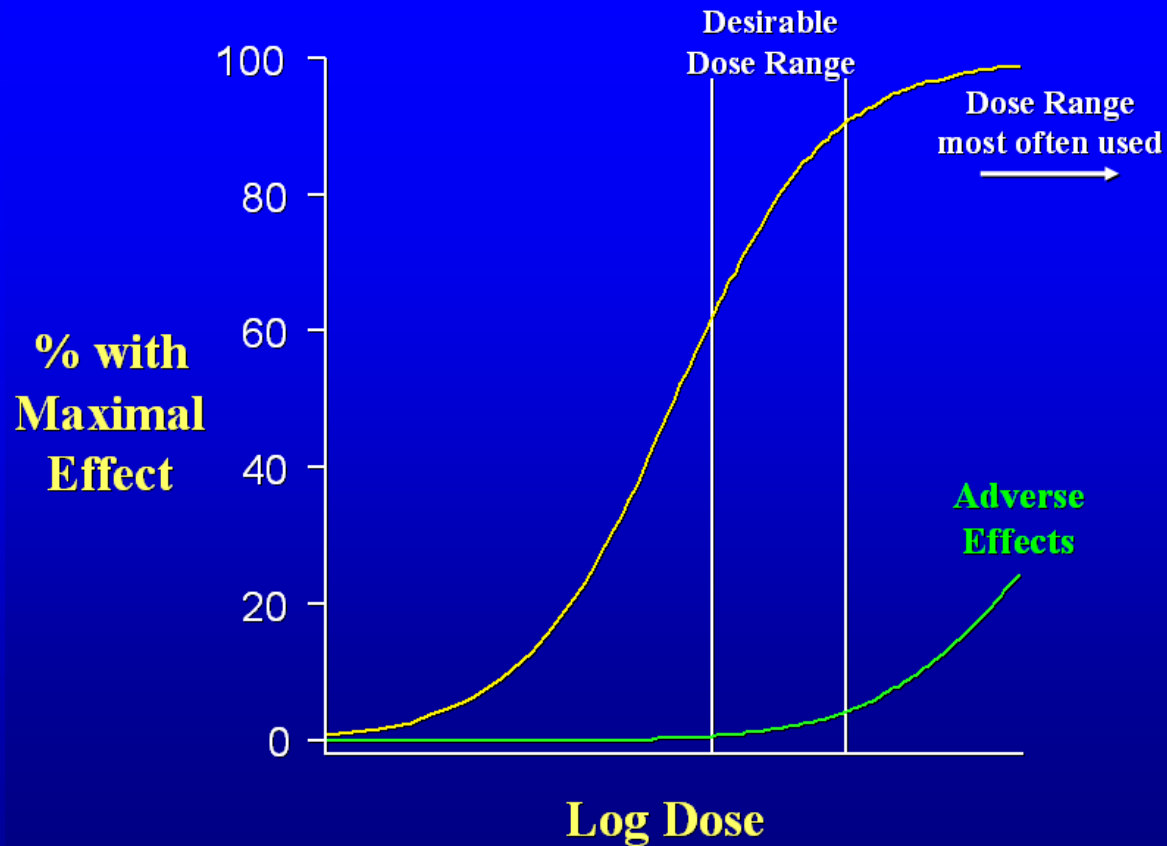
Lidocaine Quantal Dose-Effect



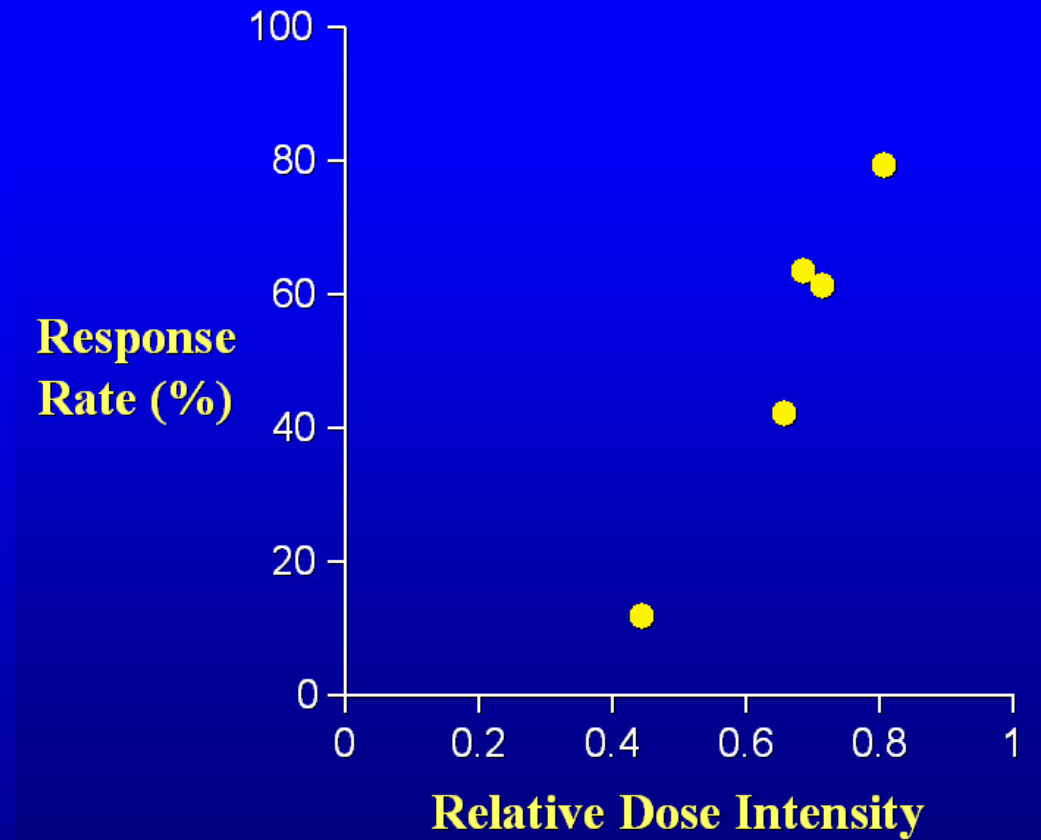
Antihypertensive Dose-Effect

Drug	Dose Range [mg]		Lowest Effective Dose [mg]
	Early Studies	Present Dose	
<u>Propranolol</u>	160-5000	160-320	80
<u>Atenolol</u>	100-2000	50-100	25
<u>Hydrochlorthiazide</u>	50-400	25-50	12.5
<u>Captopril</u>	75-1000	50-150	37.5
<u>Methyldopa</u>	500-6000	500-3000	750

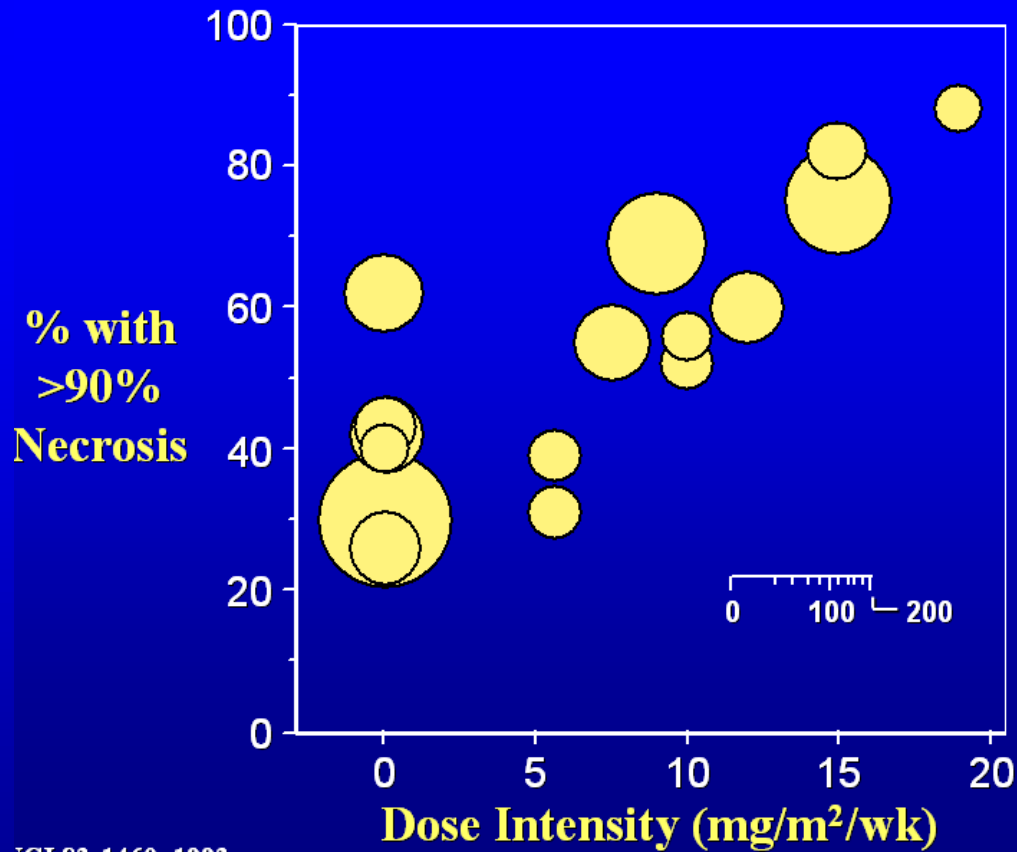
Antihypertensive Drugs



Dose Intensity in Breast Cancer



Doxorubicin Dose in Osteosarcoma



Relating Dose to Effect *In Vivo*



Age

Absorption

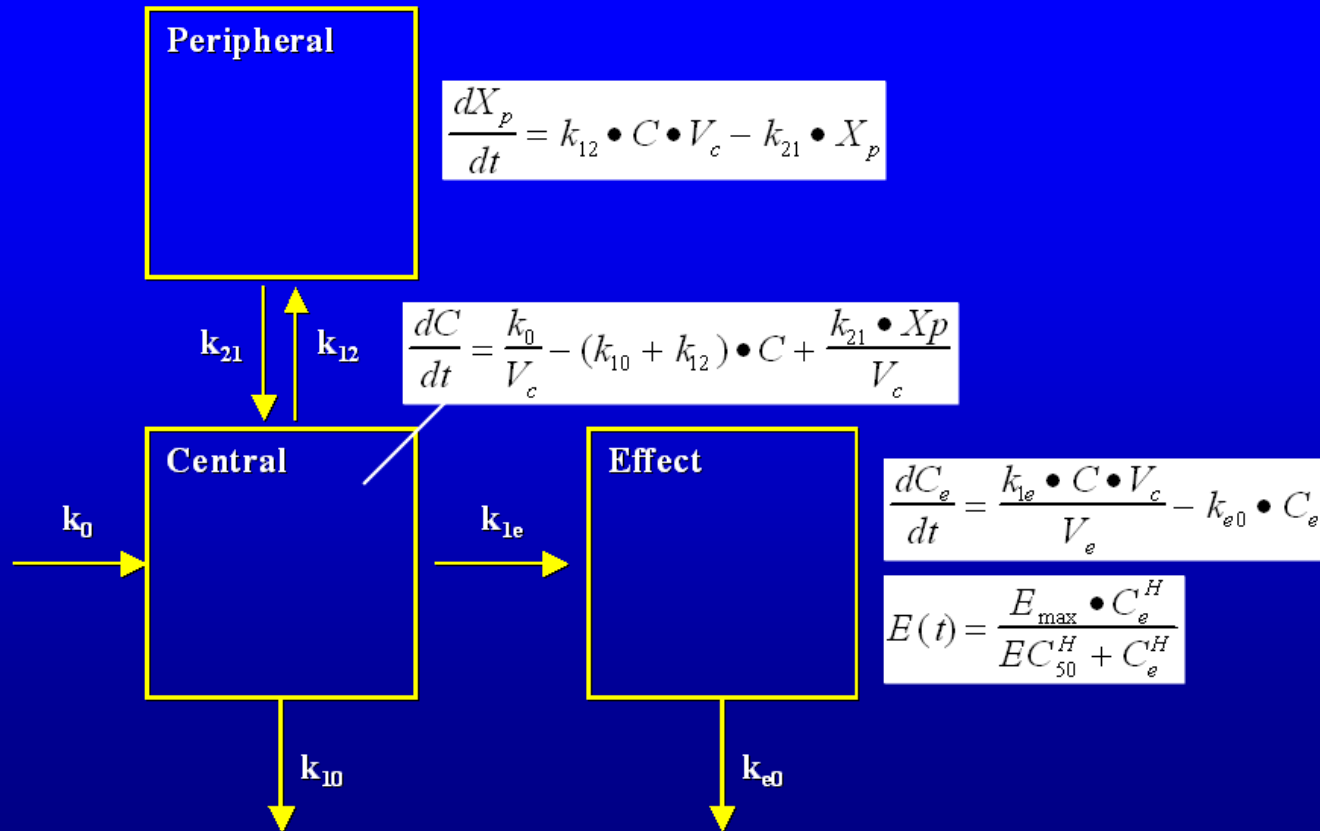
Distribution

Elimination

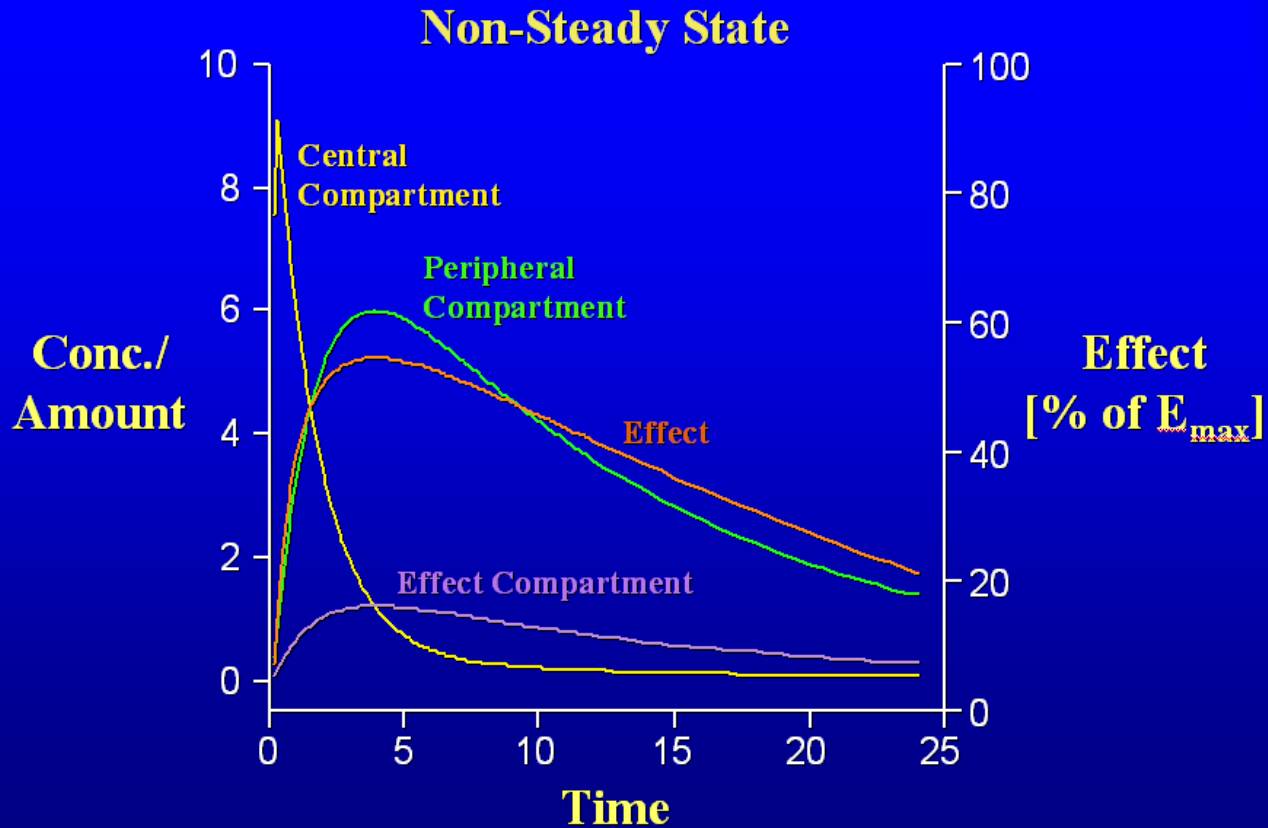
Drug interactions

**Tissue/organ sensitivity
(receptor status)**

Effect Compartment (PK/PD Model)



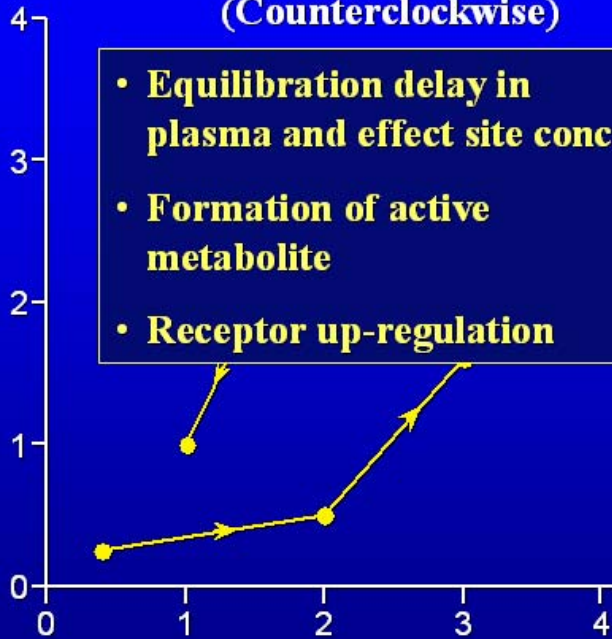
Concentration and Effect vs. Time



Hysteresis and Proteresis Loops

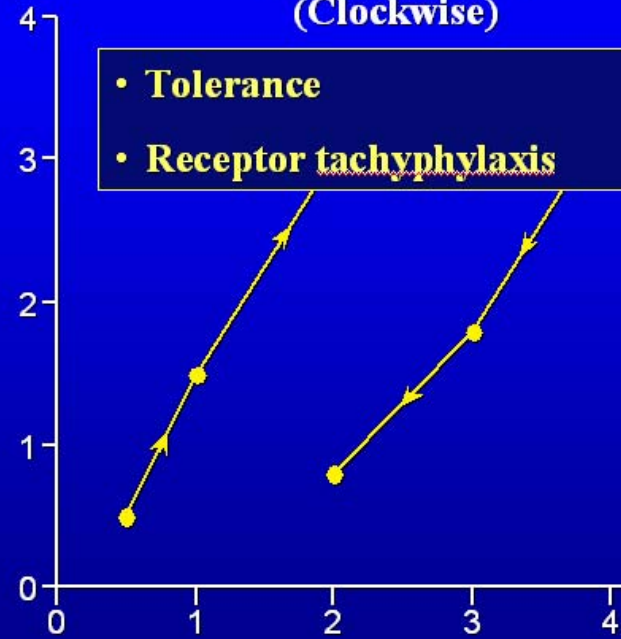
Intensity of Drug Effect

Hysteresis Loop
(Counterclockwise)



Intensity of Drug Effect

Proteresis Loop
(Clockwise)



Plasma Drug Concentration

Role of Dose-Effect Studies

*** Drug development**

- Site of action**
- Selection of dose and schedule**
- Potency, efficacy and safety**
- Drug interactions**

*** Patient management**

- Therapeutic drug monitoring**
- Risk-benefit (therapeutic indices)**

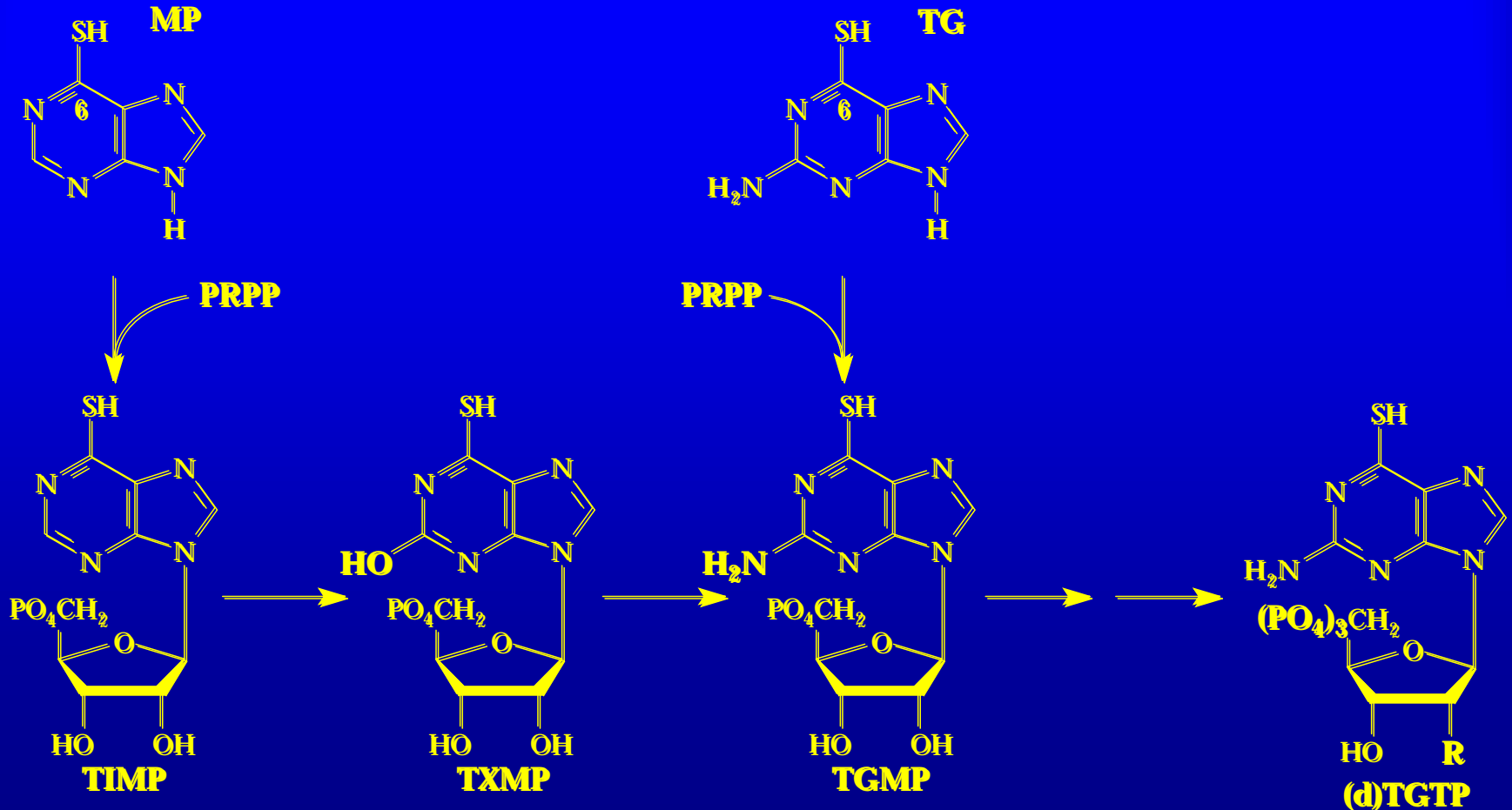
THE END

Endpoints to Monitor Drug Effect

Farnesyltransferase Inhibitors for Cancer

LEVEL	ENDPOINT
Molecular	<u>Farnesyltransferase inhibition</u>
Cellular	Proliferation rate, apoptosis
Tumor	Response (change in tumor size)
Organism	Survival, quality of life

Thiopurine Metabolic Activation



Therapeutic Indices

$$\text{Therapeutic Ratio} = \frac{\text{TD}_{50}}{\text{ED}_{50}} = 2.5$$

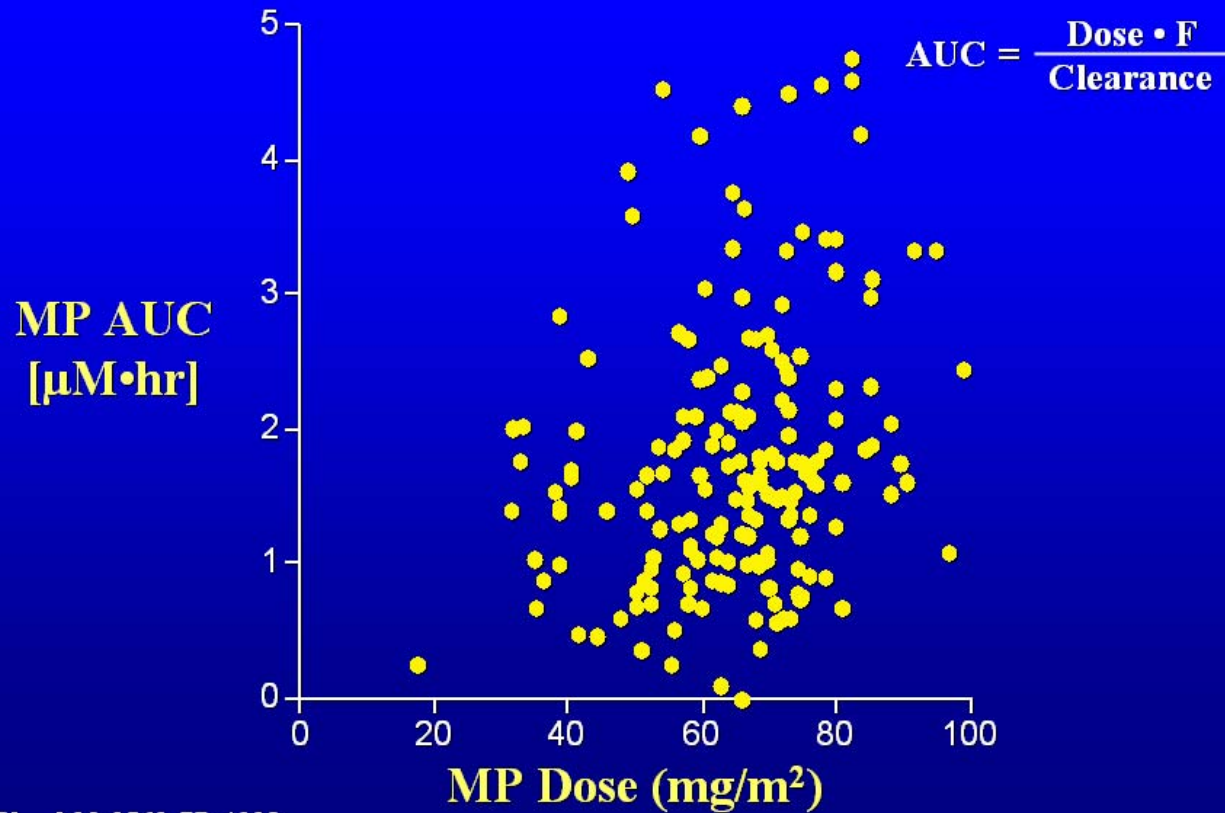
$$\text{Certain Safety Factor} = \frac{\text{TD}_1}{\text{ED}_{99}} = 1.3$$

$$\text{Standard Safety Margin} = \frac{\text{TD}_1 - \text{ED}_{99}}{\text{ED}_{99}} \times 100 = 31\%$$

Relative Dose Intensity

Regimen	Drugs	Dose Rate mg/m ² /wk	R.D.I.	
			Drugs	Regimen
CAF-1	Cyclo	350	1	1
	Doxo	15	1	
	FU	250	1	
CAF-2	Cyclo	125	0.36	0.56
	Doxo	12.5	0.83	
	FU	125	0.50	

Oral Mercaptopurine



Pharmacodynamic Models

* Fixed effect model

$$Effect = E_0 + S \cdot [Drug]$$

* Linear model

$$Effect = I + S \cdot \text{Log}([Drug])$$

* Log-linear model

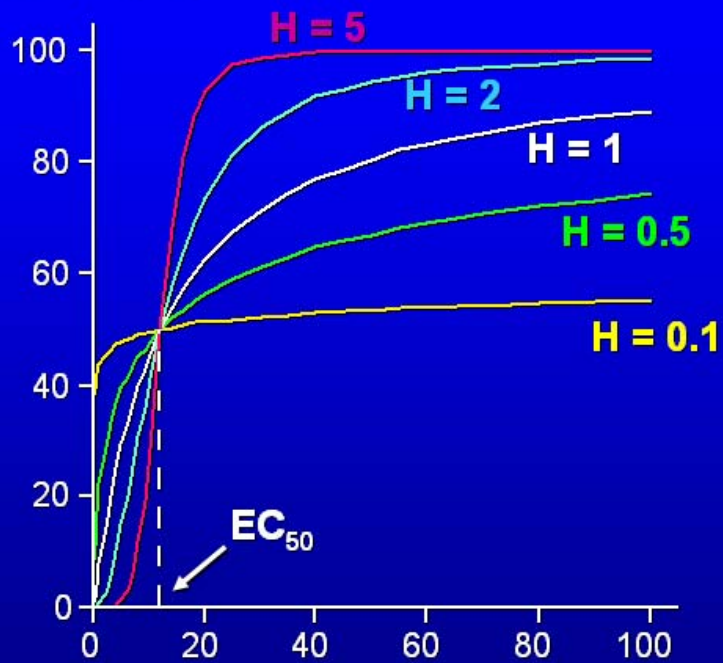
$$Effect = \frac{E_{max} \cdot [Drug]^H}{EC_{50}^H + [Drug]^H}$$

* E_{max} model

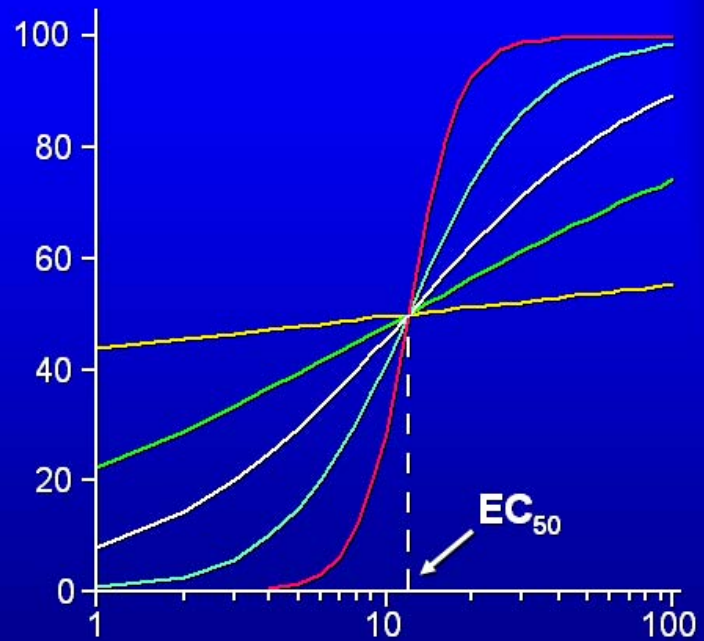
* Sigmoid E_{max} model

Sigmoid E_{\max} PD Model

Effect (%)

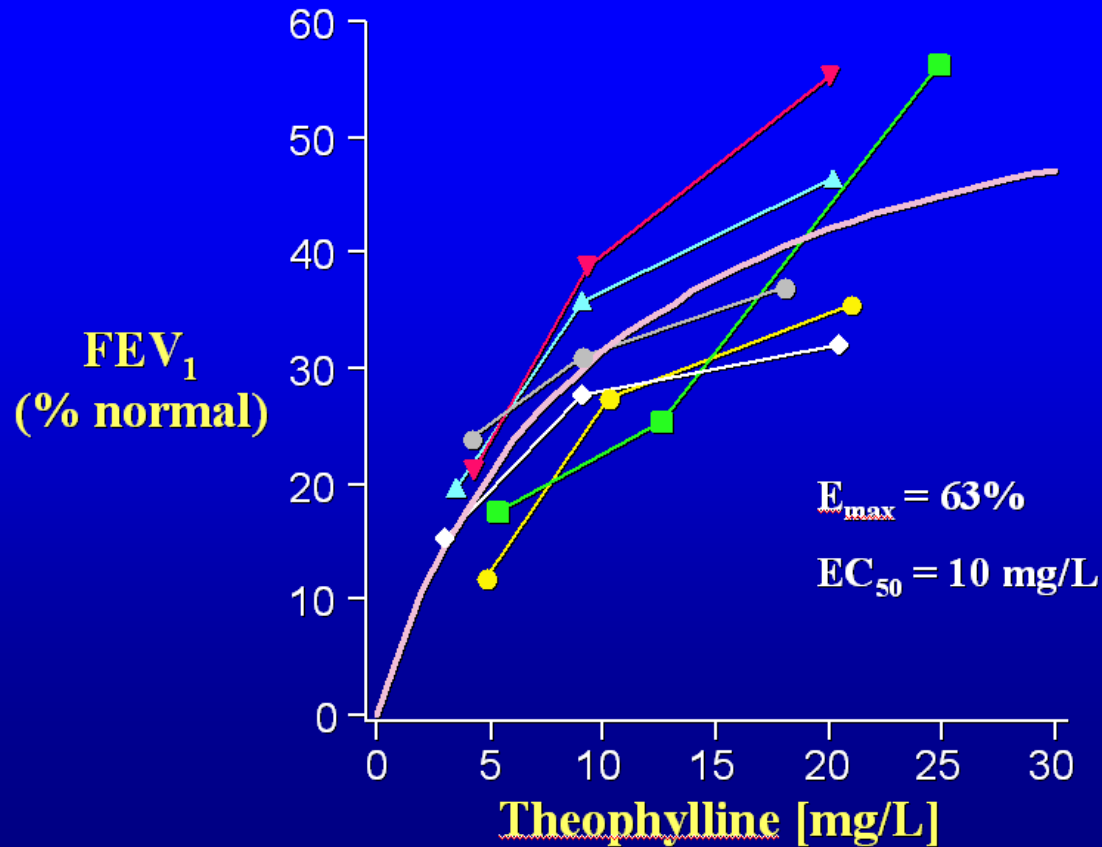


Effect (%)



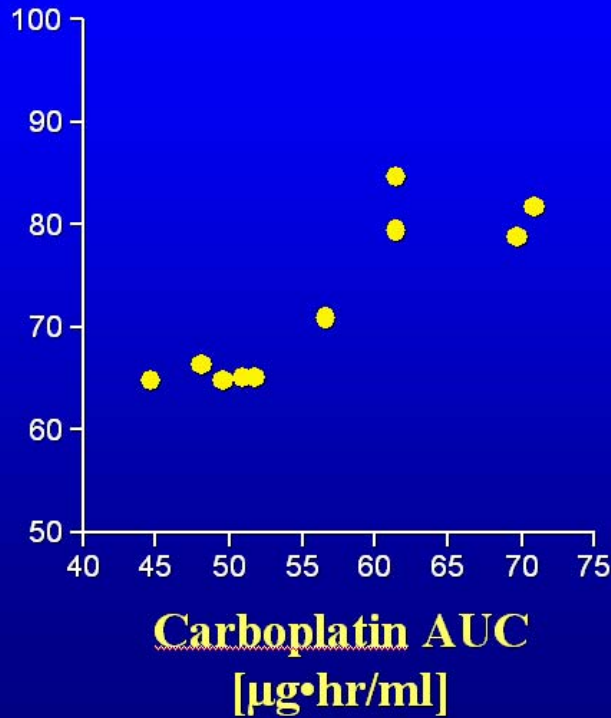
[Drug]

Theophylline Pharmacodynamics

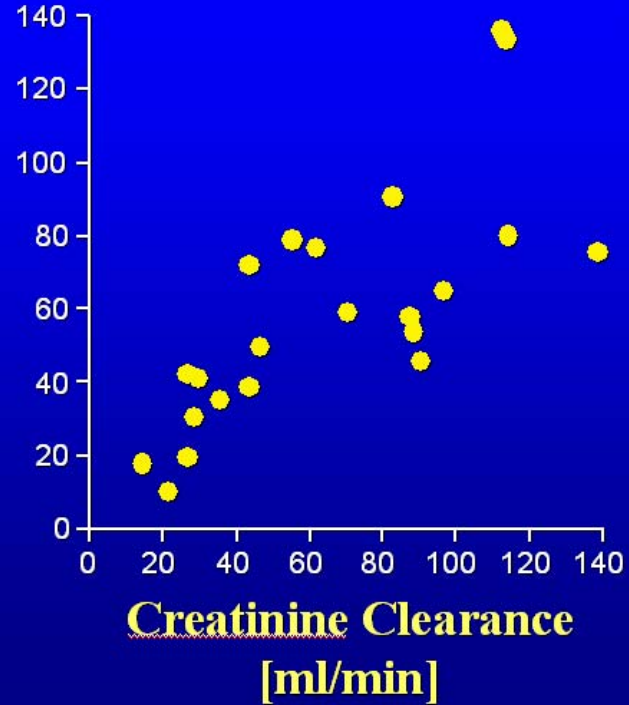


Carboplatin PK/PD

**% Decrease
Platelet**



**Carboplatin
 Cl_{TB} [ml/min]**



Carboplatin Adaptive Dosing

ADULTS

$$D[\text{mg} / \text{m}^2] = 0.091 \times CL_{CR}[\text{ml} / \text{min} / \text{m}^2] \times \left(\frac{\text{prePlt} - \text{trgtPlt}}{\text{prePlt}} \times 100 - \text{priorRx} \right) + 86$$

$$D[\text{mg}] = \text{trgtAUC}[\text{mg} \cdot \text{min} / \text{ml}] \times (\text{GFR}[\text{ml} / \text{min}] + 25)$$

CHILDREN

$$D[\text{mg} / \text{m}^2] = \text{trgtAUC}[\text{mg} \cdot \text{min} / \text{ml}] \times (0.93 \times \text{GFR}[\text{ml} / \text{min} / \text{m}^2] + 15)$$

$$D[\text{mg}] = \text{trgtAUC}[\text{mg} \cdot \text{min} / \text{ml}] \times (\text{GFR}[\text{ml} / \text{min}] + (0.36 \times \text{BW}[\text{kg}]))$$