

HO
$$\frac{5}{4}$$
 OH HO $\frac{5}{3}$ OH HO $\frac{5}{3}$ OH $\frac{3}{2}$ HO $\frac{3}{2}$ P-ribose β -deoxyribose

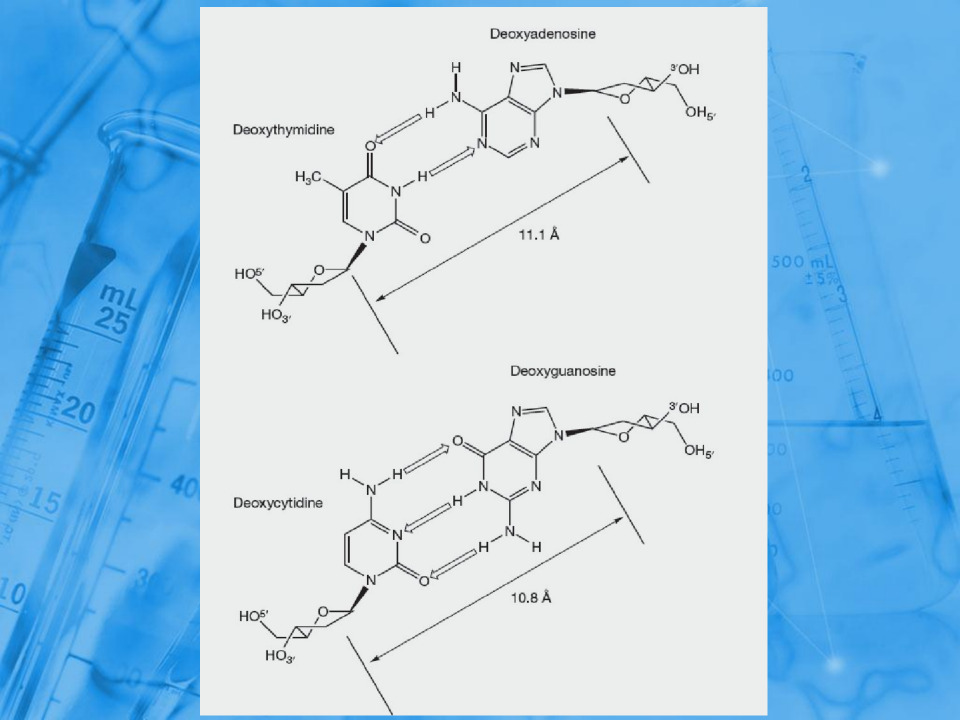
Изобразите структуру α-гликозида дезокситимидина

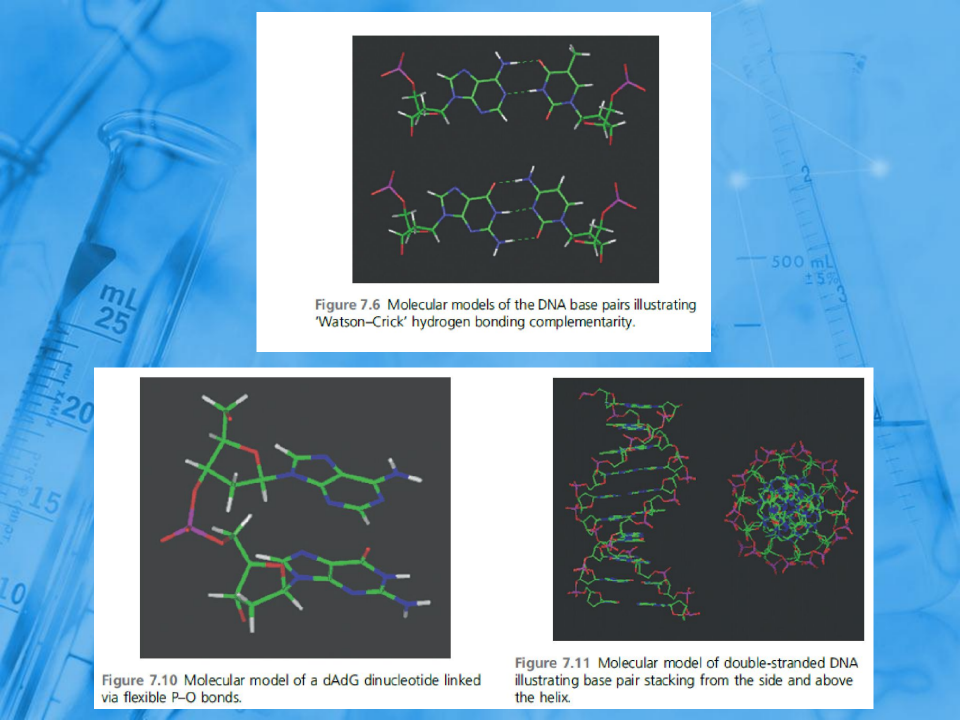
Deoxythymidine-5'-diphosphate

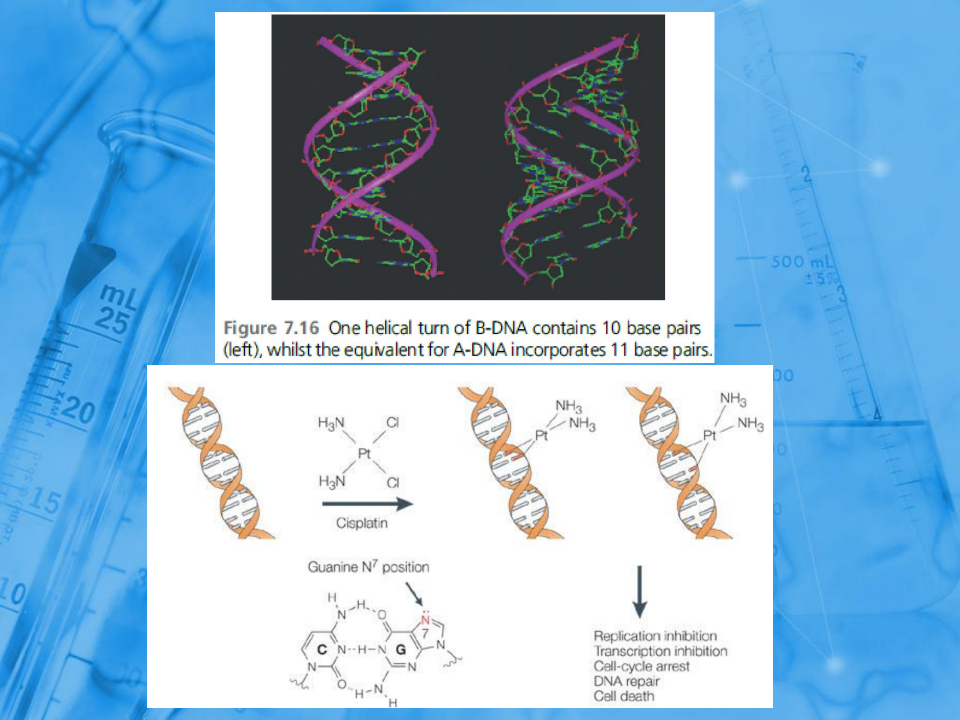
Deoxycytidine-5'-monophosphate

Deoxyadenosine-3'-monophosphate

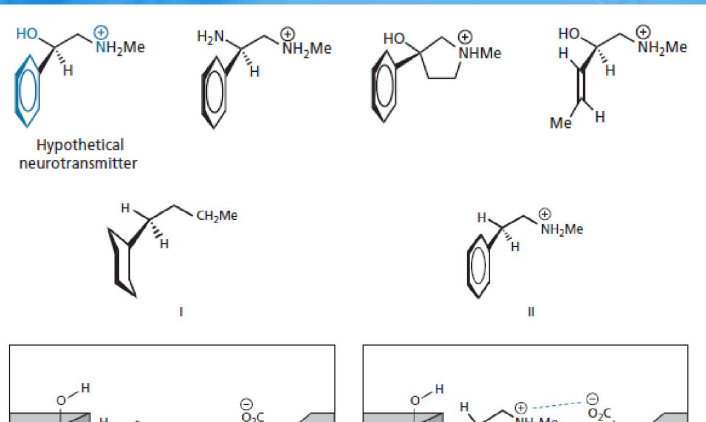
Deoxyguanosine-3',5'-biphosphate

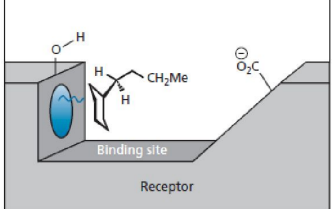






Агонисты – связывание с рецептором и его активация при низкой концентрации природного лиганда этого рецептора





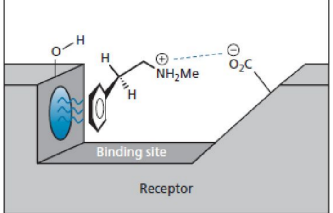
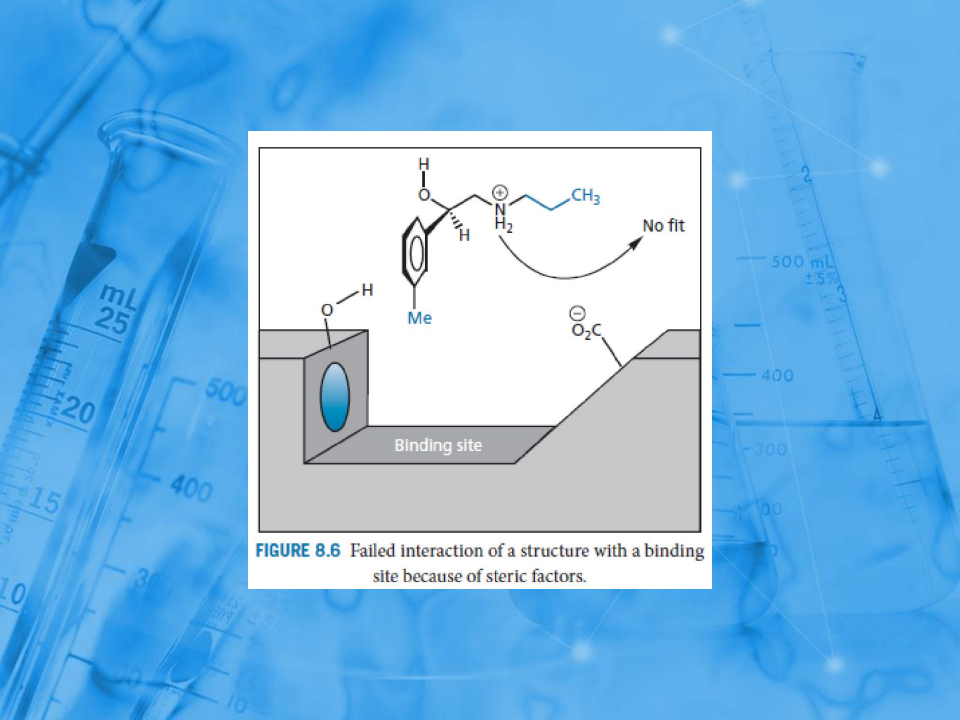
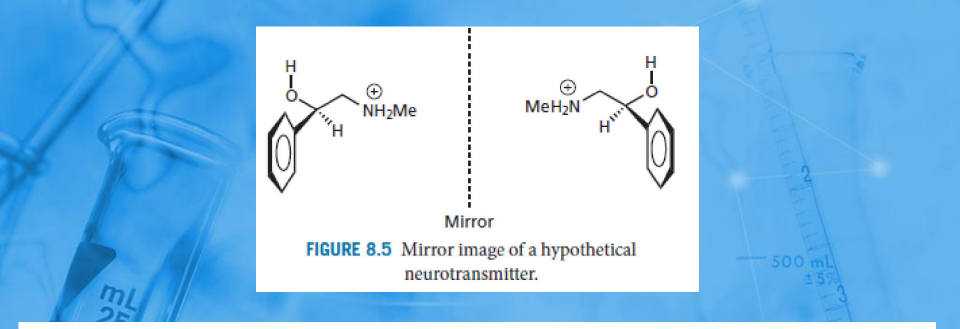
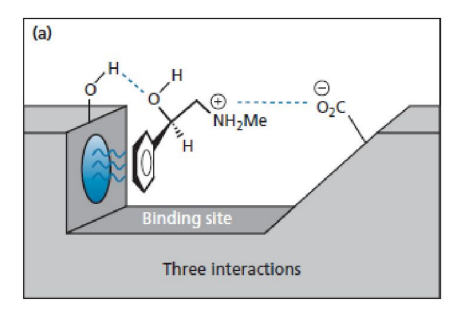


FIGURE 8.3 Weaker binding to the hypothetical receptor by structures that possess fewer than the required binding groups.







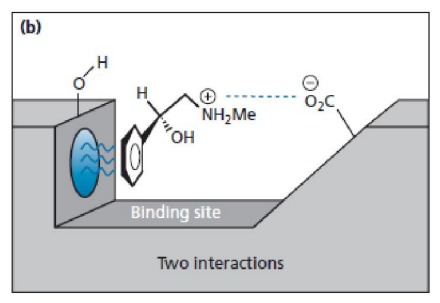


FIGURE 8.2 A comparison of interactions involving (a) the hypothetical neurotransmitter and (b) its mirror image with a hypothetical binding site.

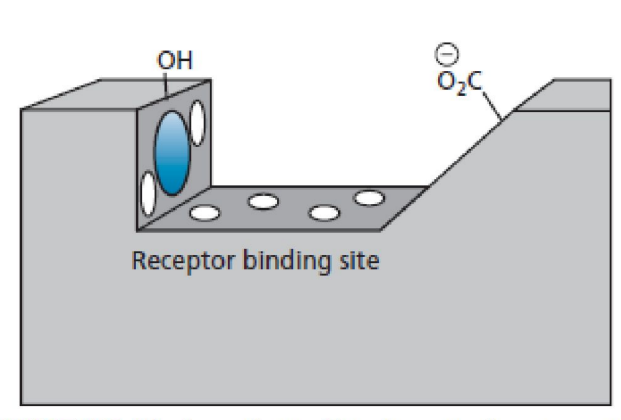


FIGURE 8.7 The hypothetical binding site showing extra binding regions (in white) that are not used by the natural chemical messenger.

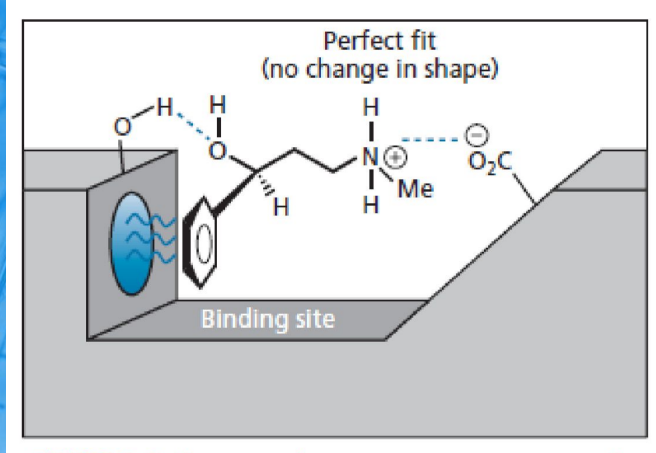
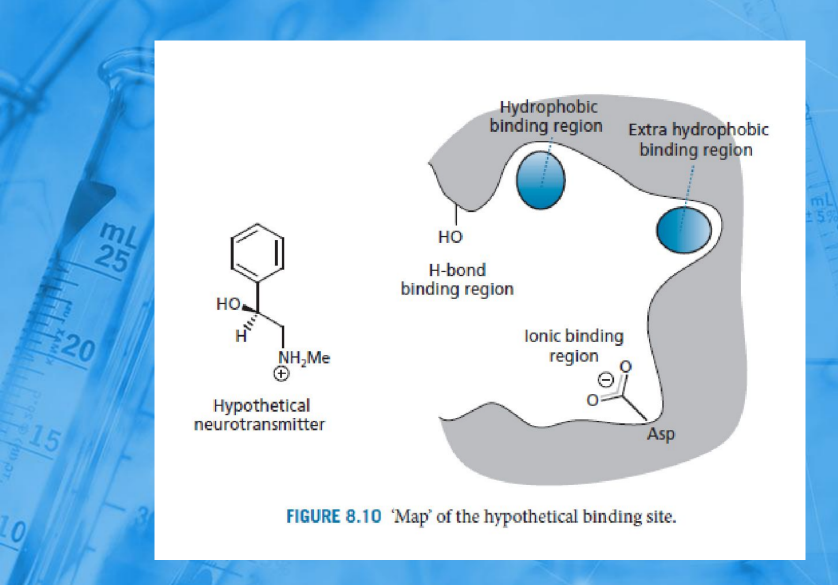


FIGURE 8.9 Compound acting as an antagonist at the binding site.



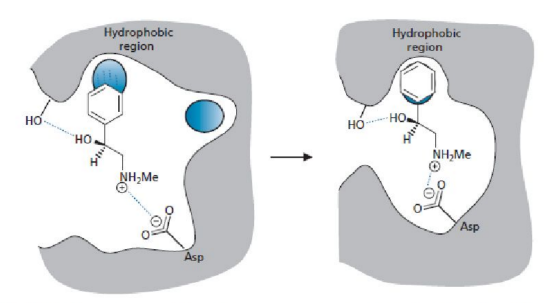


FIGURE 8.11 Binding of the natural chemical messenger resulting in an induced fit that activates the receptor.

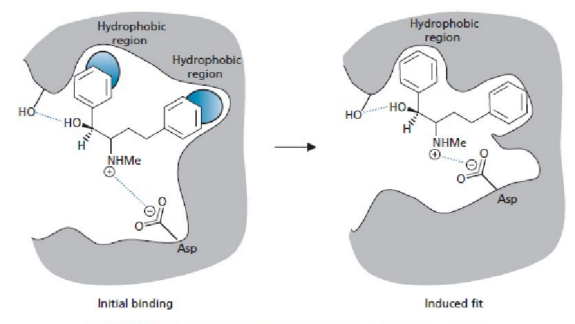


FIGURE 8.12 Binding of an antagonist leading to a different induced fit.

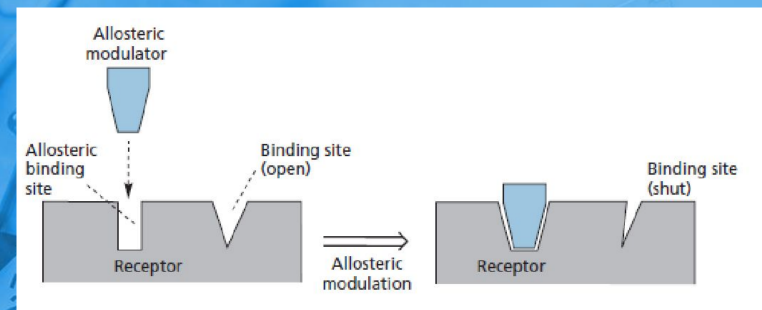


FIGURE 8.13 Principle by which an allosteric antagonist distorts a binding site.

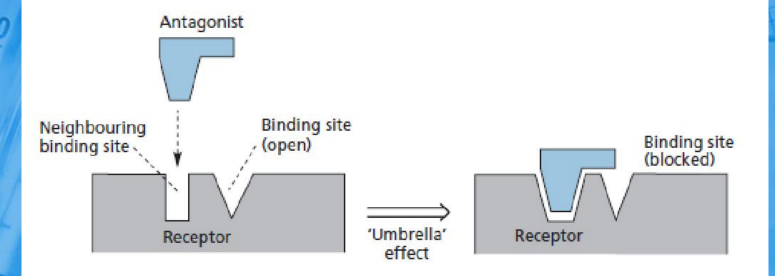


FIGURE 8.14 Antagonism by the 'umbrella effect'.



$$L + R \rightleftharpoons \underset{\substack{\text{Complex} \\ \text{complex}}}{LR}$$

$$K_{d} = \frac{[L] \times [R]}{[LR]}$$

$$[R_{tot}] = [R] + [LR]$$

This means that the number of receptors unoccupied by a ligand is

$$[R] = [R_{tot}] - [LR]$$

Substituting this into the first equation and rearranging leads to the Scatchard equation, where both [LR] and [L] are measurable:

$$\frac{[\text{Bound ligand}]}{[\text{Free ligand}]} = \frac{[\text{LR}]}{[\text{L}]} = \frac{R_{\text{tot}} - [\text{LR}]}{K_{\text{d}}}$$

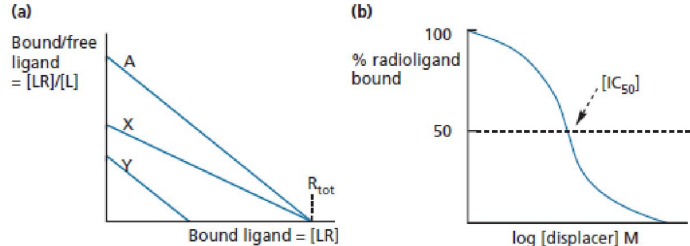
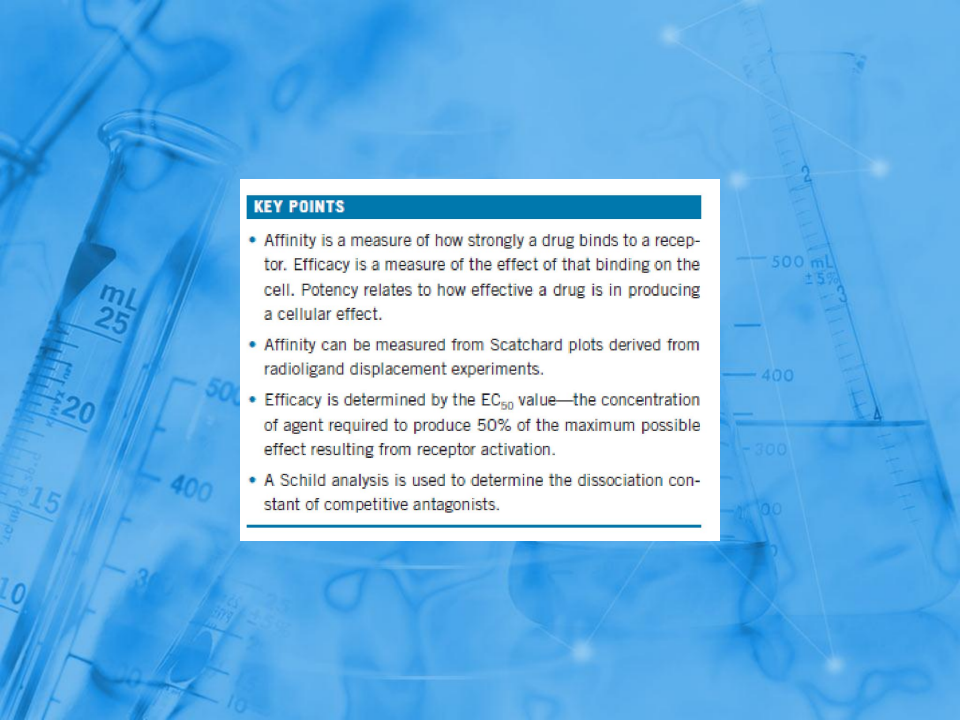


FIGURE 8.20 (a) Scatchard plot (A = radioligand only, X = radioligand + competitive ligand, Y = radioligand + noncompetitive ligand). (b) The displacement or inhibition curve.



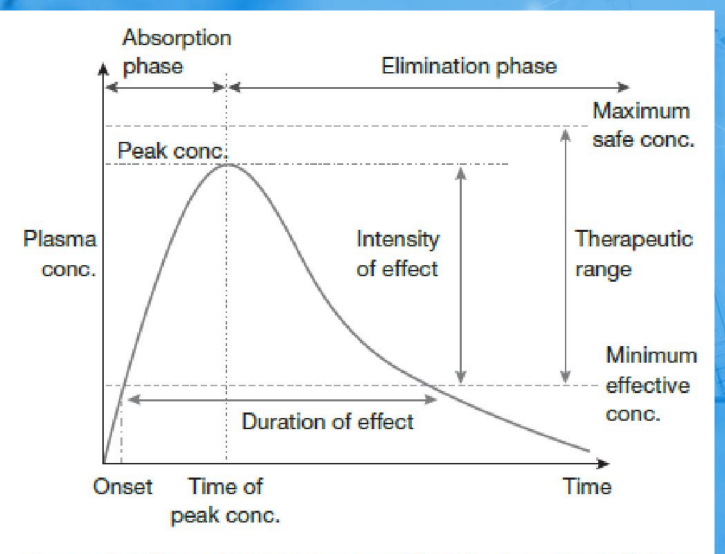
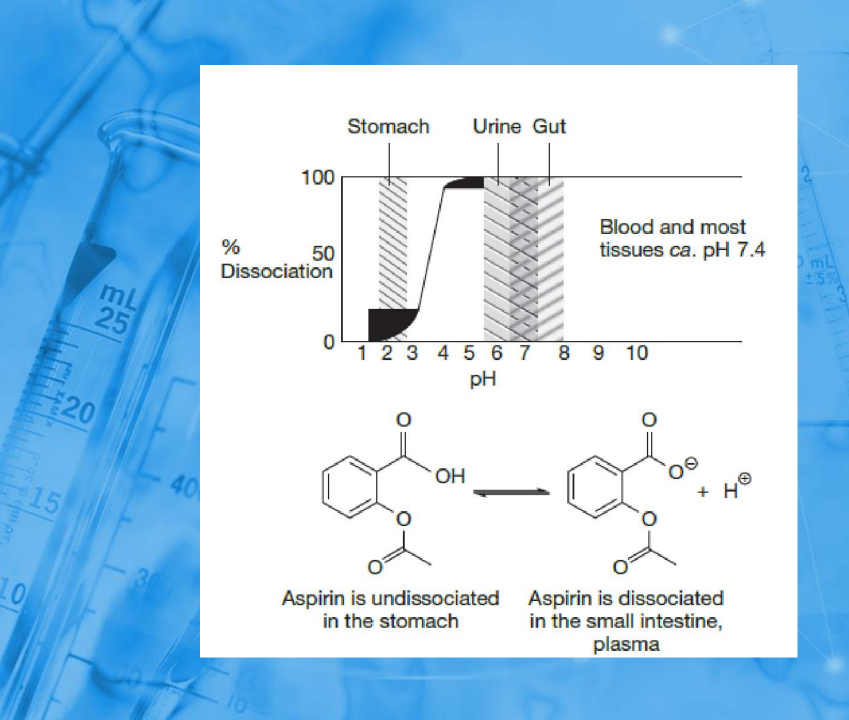


Figure 8.1 Drug dose response profile for a hypothetical drug, showing the absorption and elimination phases following administration.



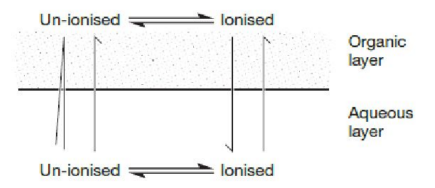


Figure 8.6 Ionised and un-ionised drug in aqueous and organic phases.

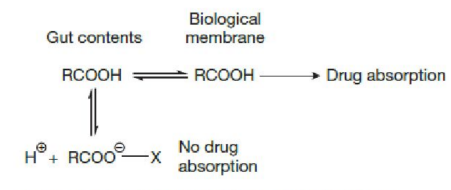


Figure 8.8 The effect of pH on the absorption of an organic acid.

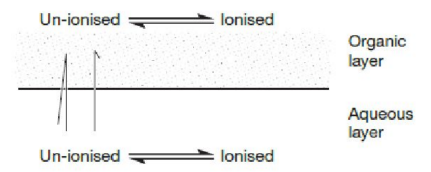


Figure 8.7 Un-ionised drug in an organic layer in equilibrium with un-ionised and ionised drug in an aqueous layer.

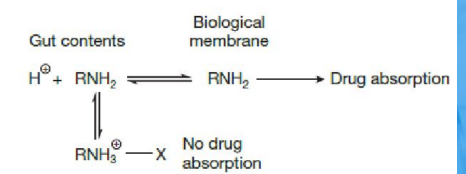
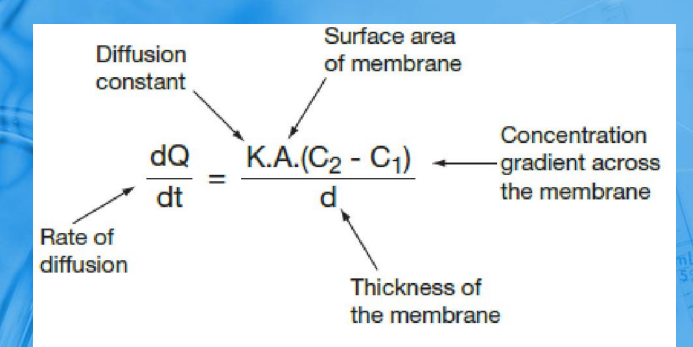


Figure 8.9 The effect of pH on the absorption of an organic base.



The diffusion constant, K, is a function of:

- aqueous solubility of the drug i.e. hydrophilic character
- lipid solubility of the drug i.e. o/w partition coefficient
- molecular size, i.e. RMM
- molecular shape
- pKa of the drug
- pH of the environment.

Рациональный подход к дизайну пролекарства:

- 1. Идентификация проблем, связанных с доставкой лекарства.
- 2. Определение физико-химических свойств, необходимых для максимального увеличения эффективности доставки.
- 3. Подбор подходящего производного, обладающего желаемыми физико-химическими свойствами, которое будет наиболее эффективно расщепляться в нужном биологическом отсеке, высвобождая при этом фармакологически активное вещество.

Several criteria have to be considered in the design of a prodrug, including:

- 1. the functional group(s) in the parent drug molecule which are amenable to chemical derivatisation
- 2. the mechanism(s) available in the body for bioactivation of the prodrug
- 3. ease of synthesis and purification of the prodrug, i.e. economic considerations
- 4. stability of the prodrug per se and its compatibility with other components of a pharmaceutical formulation
- 5. the rate and extent of regeneration of the parent drug from the prodrug in vivo, i.e. Biochemical considerations
- 6. toxicity of the prodrug and also of its transport group.

Figure 8.19 Conversion of pivampicillin into ampicillin by esterase activity followed by spontaneous decomposition.















