

Drug Receptor Interactions

What would you do?

- thoracotomy
- premed:
buprenorphine
(partial agonist)
- intra-op: fentanyl
(full agonist)
- recovery: naloxone
(antagonist)
- post op analgesia?



agonist

- A drug which interacts with a specific receptor to produce a response
 - ie, it has efficacy

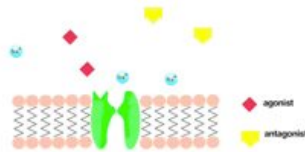
efficacy

- The ability to produce a response after binding

antagonist

- A drug which occupies a receptor stopping an agonist getting in
- it produces no effect on its own
 - ie, it has no efficacy

competitive antagonist



inverse agonist

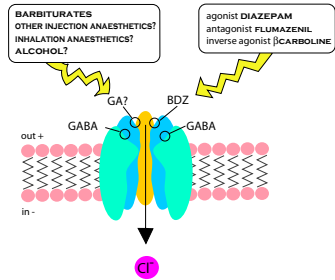
- A drug which occupies a receptor to produce the opposite effect to an agonist
 - ie, it has negative efficacy
- it is also blocked by an antagonist
- constitutive activation required

partial agonist

- a drug which occupies a receptor and produces a response which is smaller than that of a full agonist
 - ie it has low efficacy

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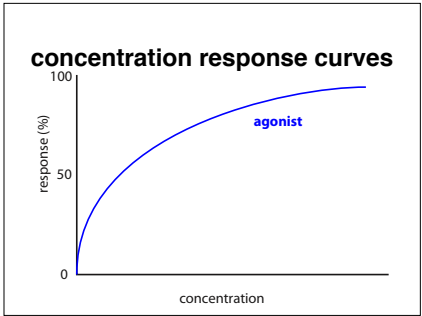
affinity

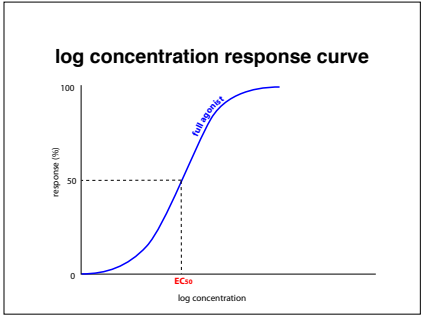
- The tendency of a drug to bind to receptors

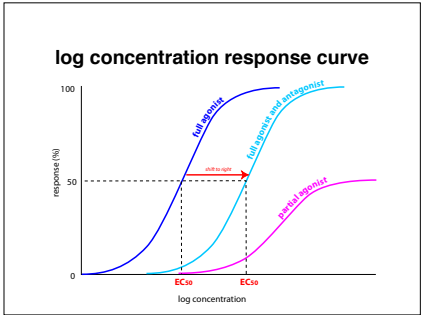
$$KA = 1 \\ KD$$

affinity

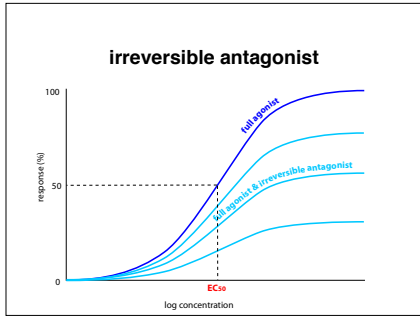
- high affinity drug
 - high occupancy at low concentration
- low affinity drug
 - high occupancy at high concentration



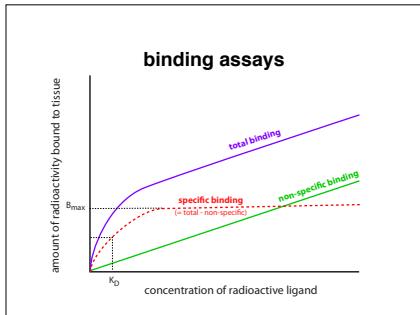


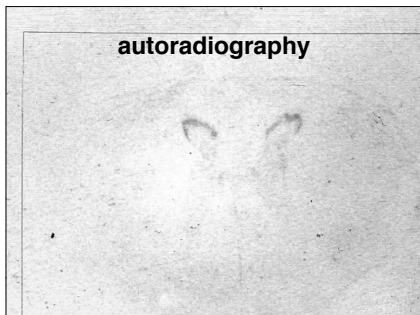


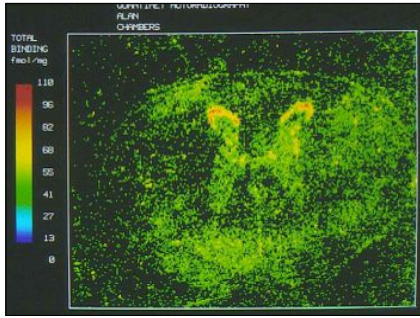
- antagonism**
- **competitive**
 - reversible
 - irreversible
 - **non-competitive**
 - usually channel blockers
 - **physiological**
 - **chemical**
 - **pharmacokinetic**

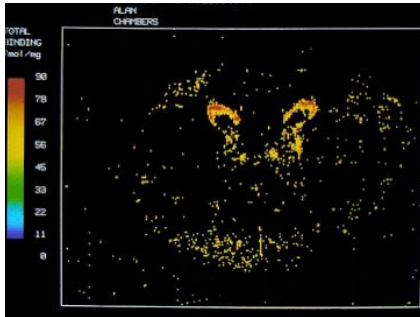


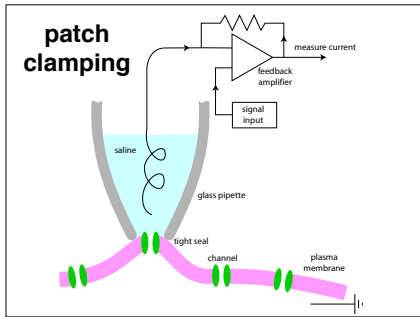
- ### binding assays
- tissues homogenised
 - cell membranes collected
 - incubated with radioligand
 - recovered by filtration & washed
 - radioactivity measured
 - KD and Bmax calculated

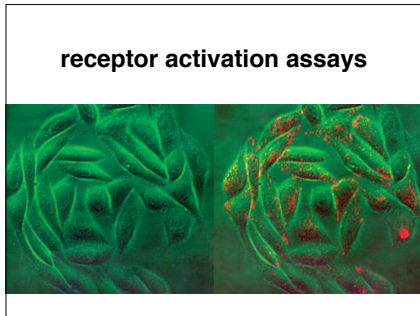












receptor numbers

- change with use
- up and down regulation

receptor reserve

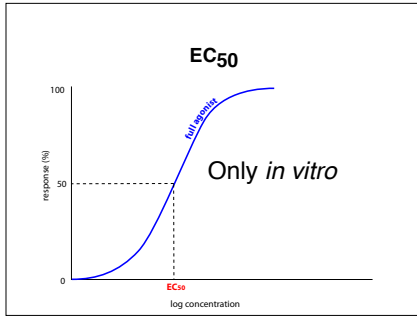
- = spare receptors
- more receptors in tissue than required for full response
- partial agonists may produce a full response in a tissue with many spare receptors
- common in smooth muscle

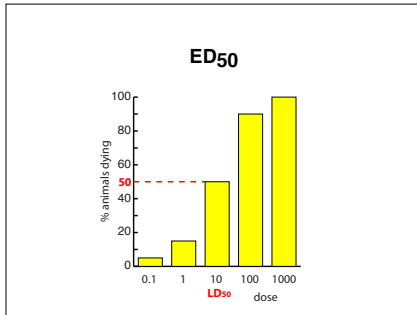
desensitisation / tachyphylaxis (receptors)

- receptor down regulation
- conformation changes
- transducer changes
- mediator depletion

tolerance (animals)

- increased metabolism
- adaptation
 - progression of disease
- drugs pumped out





therapeutic ratio

- an index of a drug's safety

$$= \frac{LD_{50}}{ED_{50}}$$

therapeutic ratio

- difference between effective dose and dose which produces side effects is clinically important
- LD₅₀ ethically unacceptable

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drug receptor interactions

- agonists produce an effect
- competitive antagonists block the effect but the blockade can be overcome by increasing the agonist concentration
- drugs can be compared using EC_{50} values in vitro and ED_{50} values in vivo
- therapeutic index is a measure of how safe a drug is
