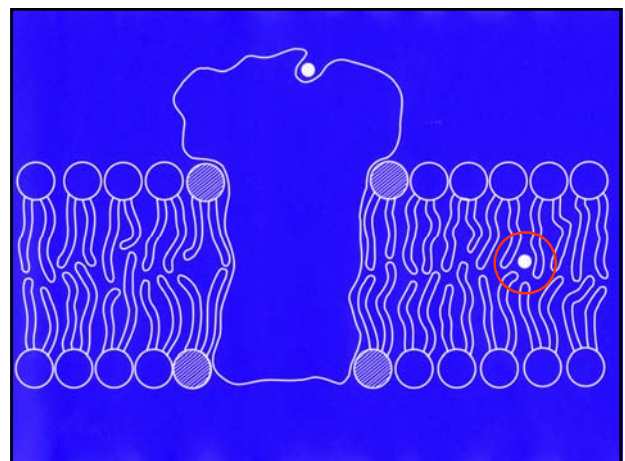
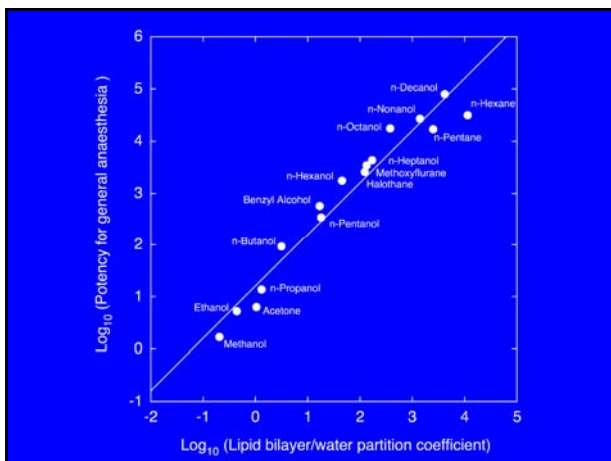
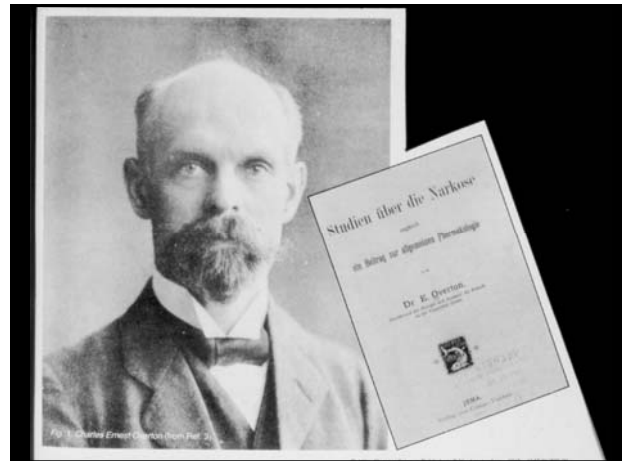
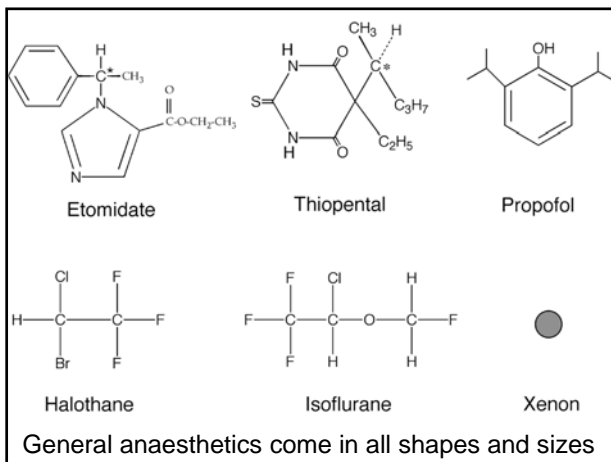


Molecular Targets of General Anaesthetics

Dr Robert Dickinson
 Anaesthetics, Pain Medicine &
 Intensive Care Section
 Imperial College
 Biophysics Group
 Blackett Laboratory
 South Kensington Campus
 r.dickinson@imperial.ac.uk

Molecular Targets of General Anaesthetics

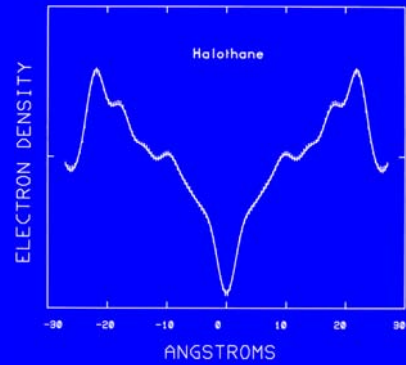
- Meyer-Overton correlation
- Theories of general anaesthesia
 - Lipid theories
 - Protein theories
- Molecular interactions with proteins
- Ion channel targets
 - Criteria for putative targets



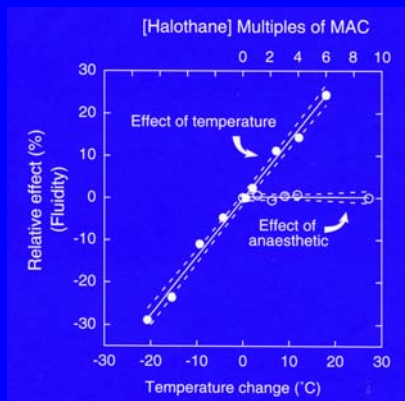
Lipid theories of anaesthesia

- Unitary hypothesis
- Diversity of theories
 - membrane expansion
 - membrane fluidity
 - membrane phase transitions

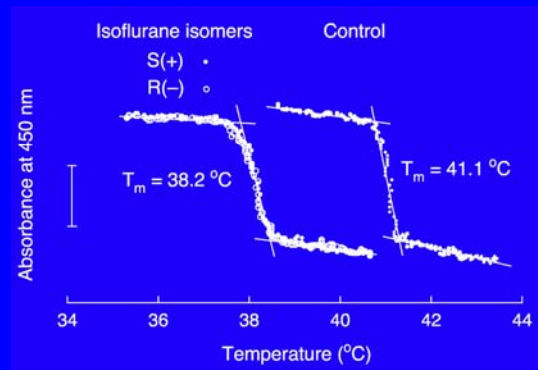
General anaesthetics do not change lipid bilayer dimensions



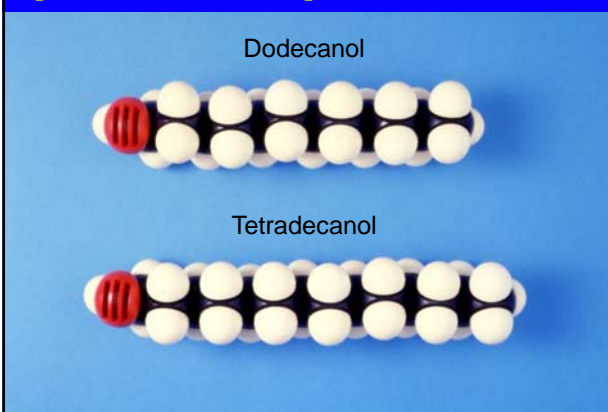
Anaesthetics do not affect lipid bilayer fluidity



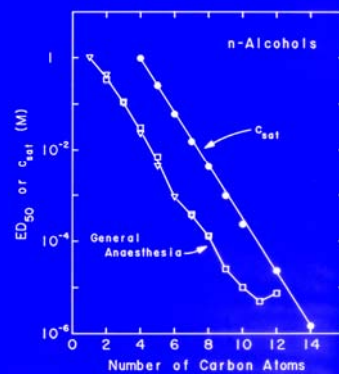
Effect of anaesthetics on lipid phase transitions is not stereoselective



Lipid theories cannot explain the “cutoff effect”

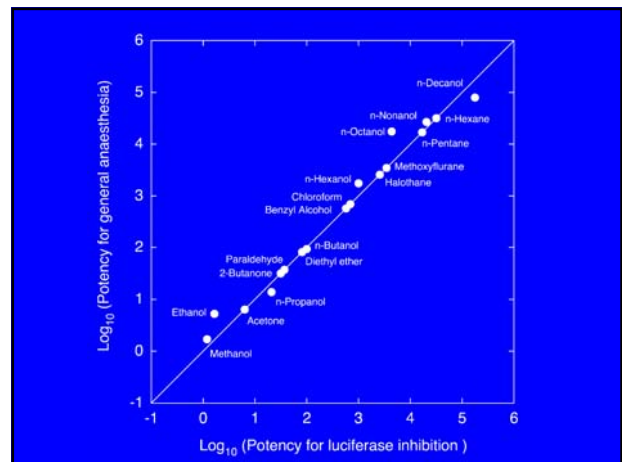


Lipid theories cannot explain the “cutoff effect”

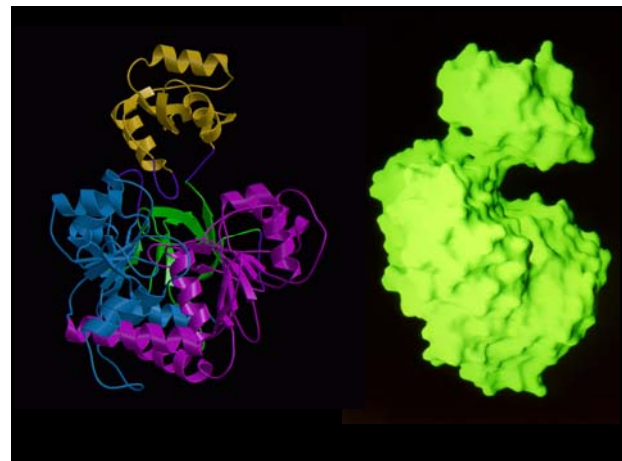
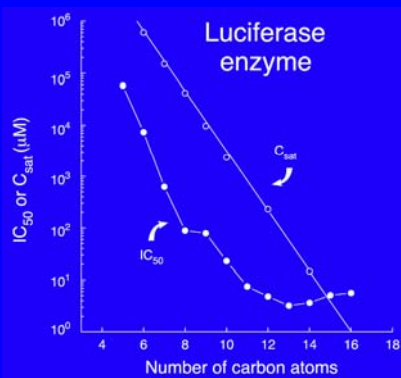


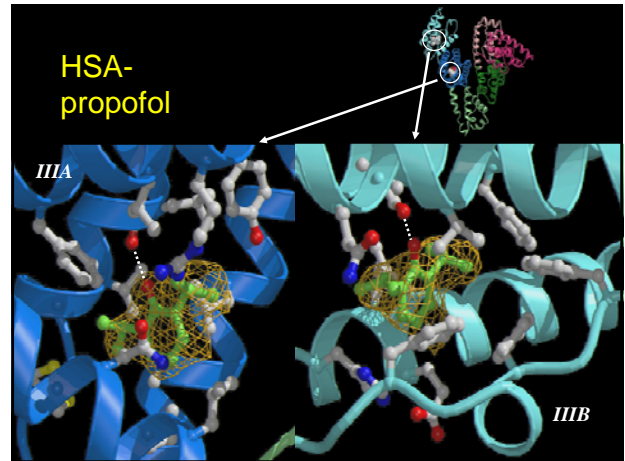
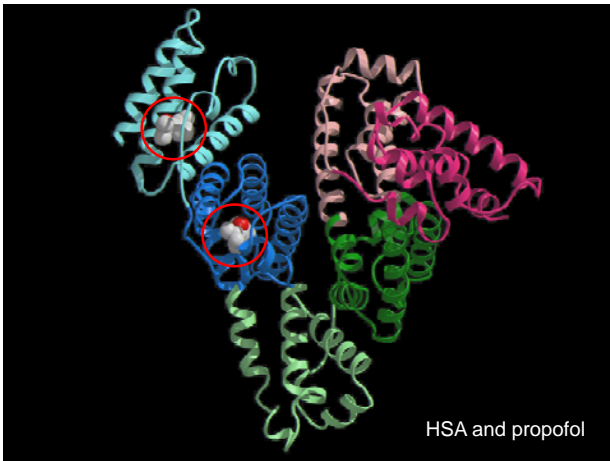
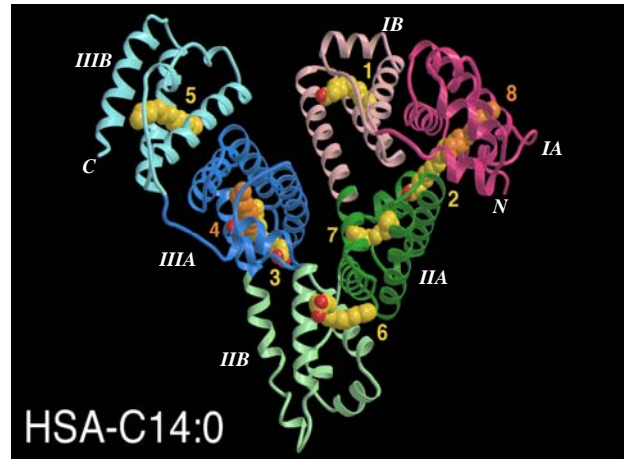
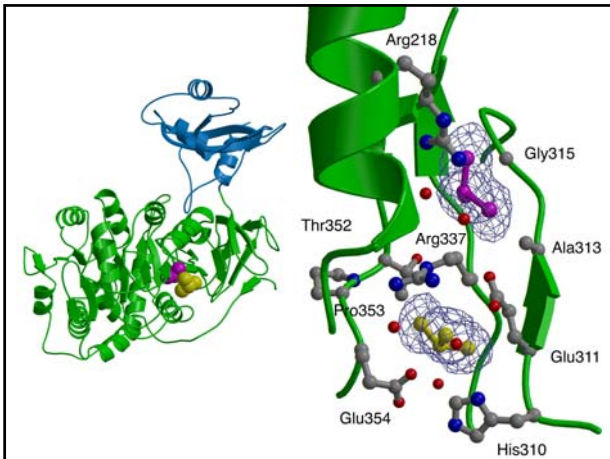
Anaesthetics do not act by disrupting lipid bilayers

Do anaesthetics act by binding to proteins?



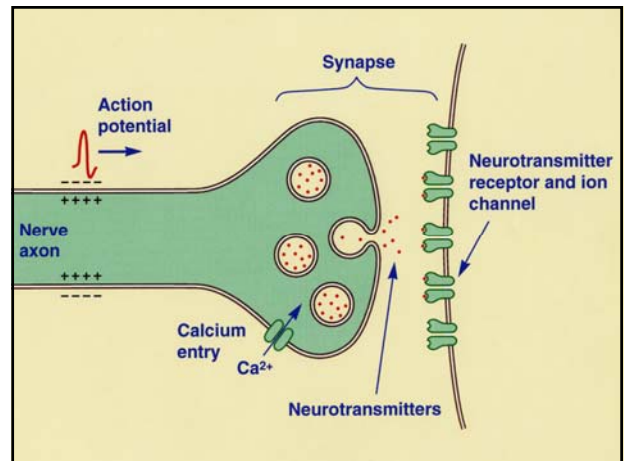
Firefly luciferase exhibits a "cutoff" effect





Anaesthetics act by binding directly to sensitive protein targets in pre-formed cavities or clefts

.....but which proteins are relevant?



Criteria for putative targets

- Plausibility
- Sensitivity
- Stereoselectivity

Anaesthetic endpoints & free aqueous concentrations for thiopental

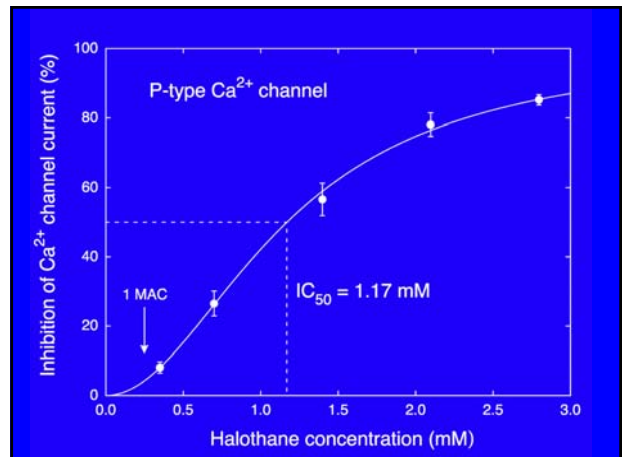
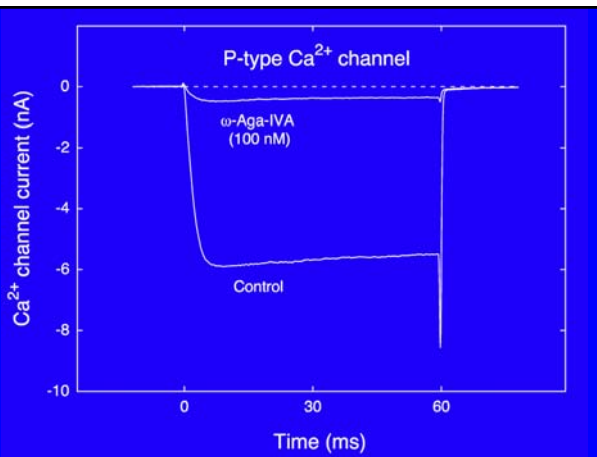
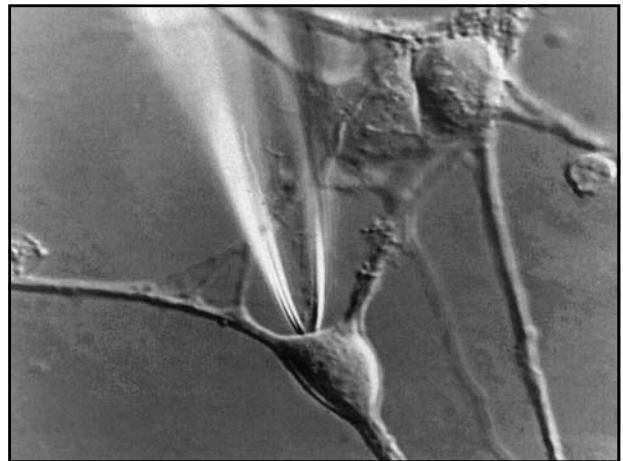
| Rat | | Human | |
|---|------------------------------|--|------------------------------|
| | Thiopental (μM) | | Thiopental (μM) |
| Righting reflex | 9 | Response to verbal command | 9 |
| Response to painful stimulus (Tail clamp) | 22 | Response to painful stimulus (Surgical incision) | 23 |
| Tracheal intubation | 39 | Tracheal intubation | 46 |

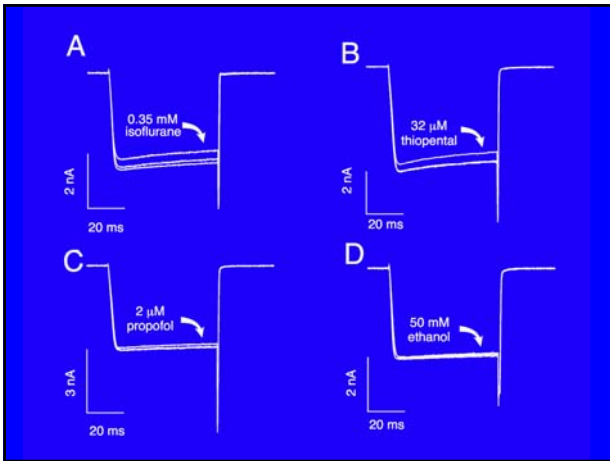
Data from Becker (1978) *Anesthesiology* 49, 192-196, Hung et al. (1992) *Anesthesiology* 77, 237-244 and Gustafsson et al. (1996) *Anesthesiology* 84, 415-427.

General anaesthetic EC_{50} concentrations for mammals

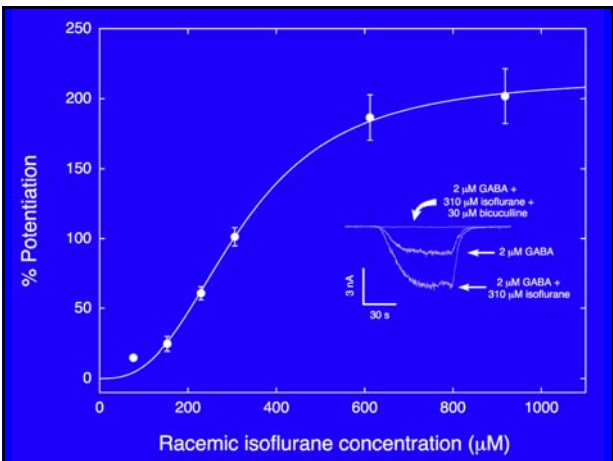
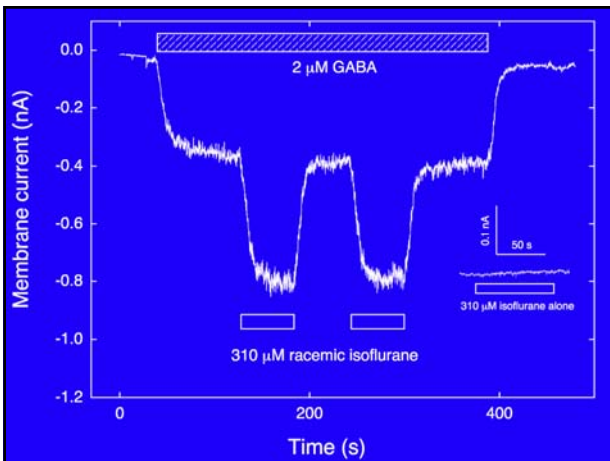
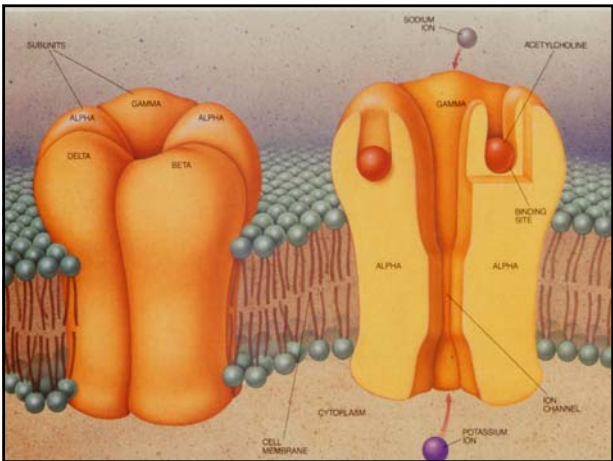
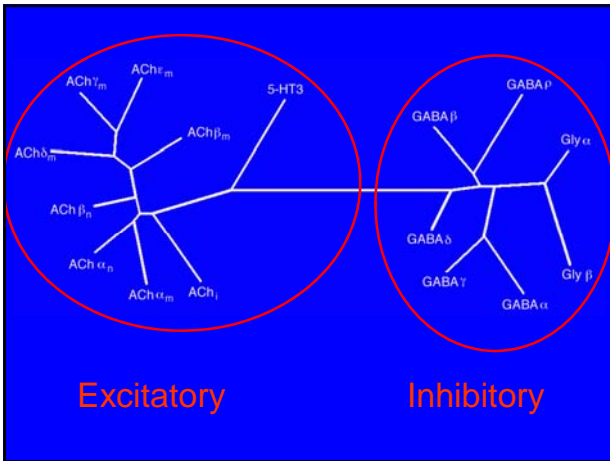
(inhibition of a response to a painful stimulus)

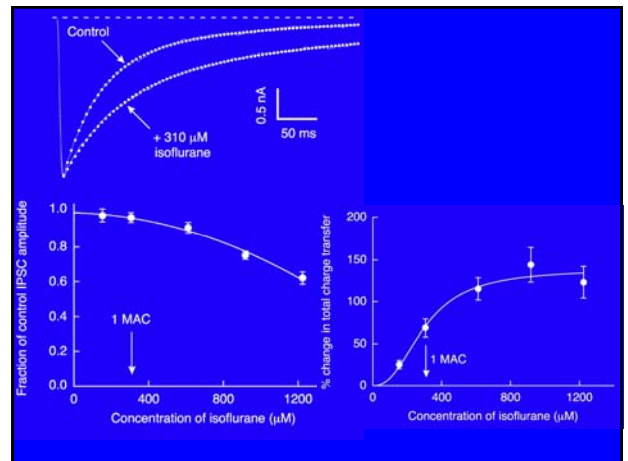
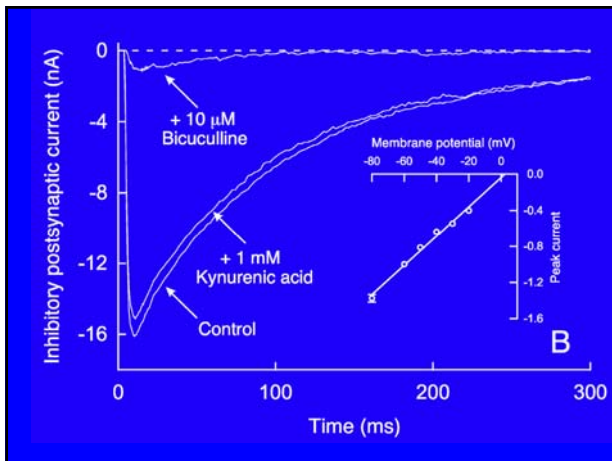
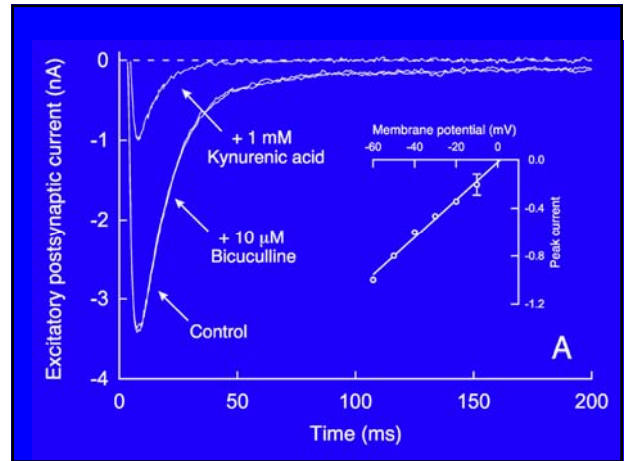
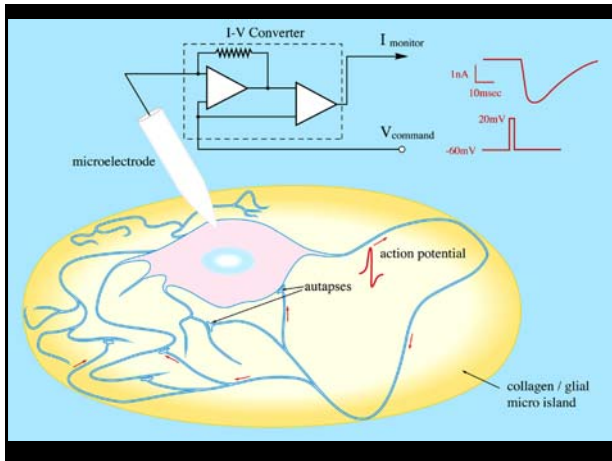
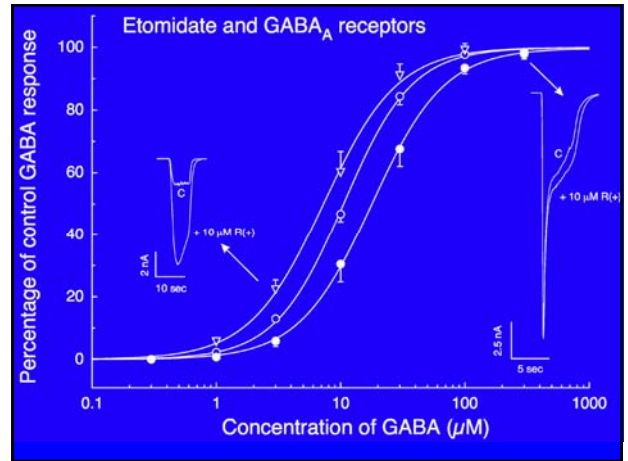
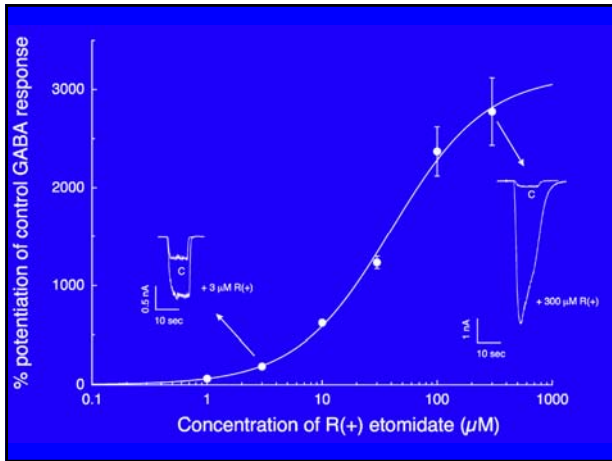
| | |
|------------|-------------------|
| Halothane | 230 μM |
| Isoflurane | 280 μM |
| Thiopental | 25 μM |
| Propofol | 1.5 μM |

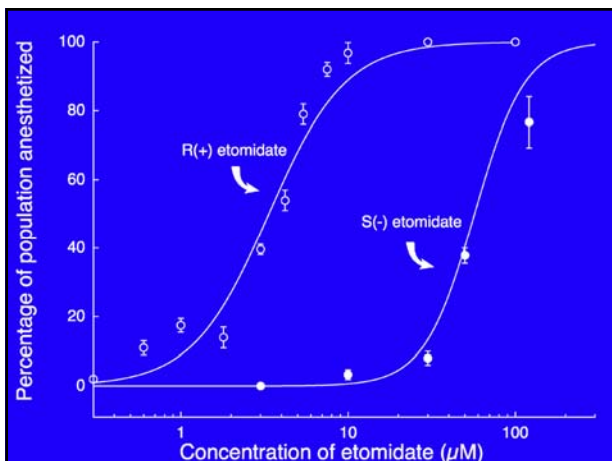
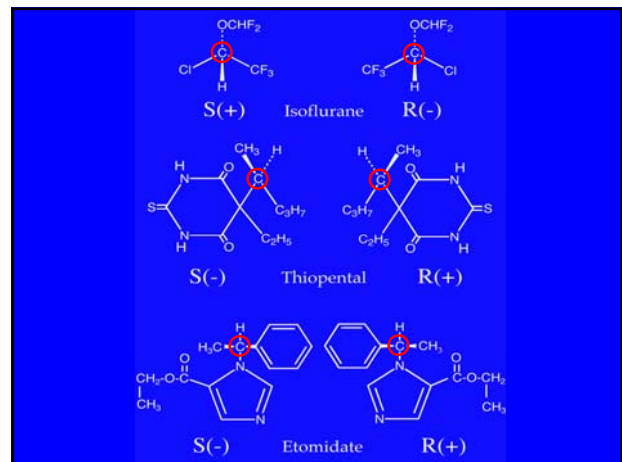
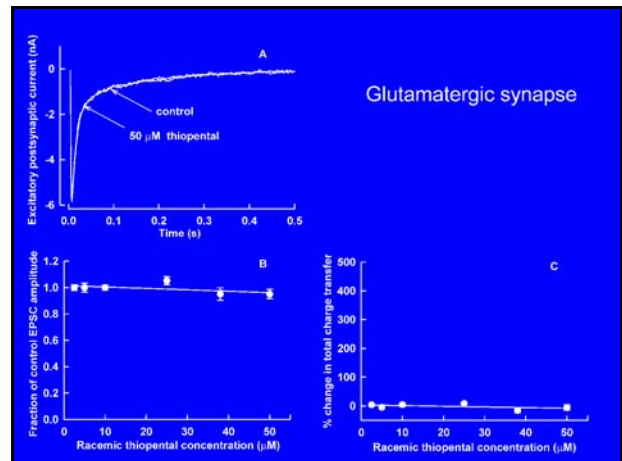
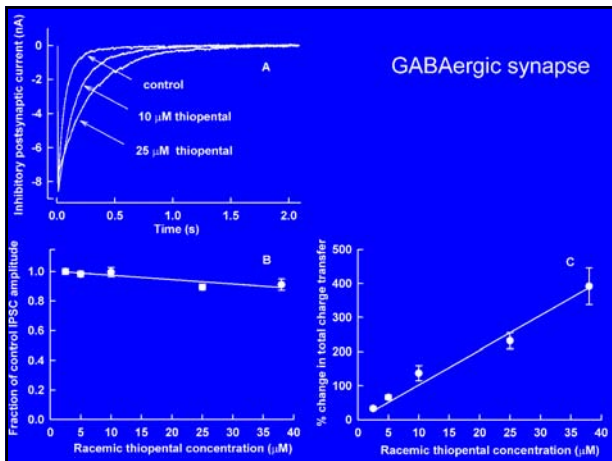




Most voltage-gated ion channels are insensitive to anaesthetics

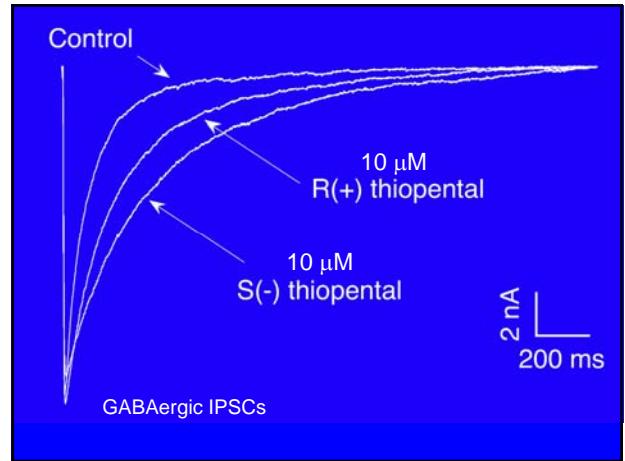
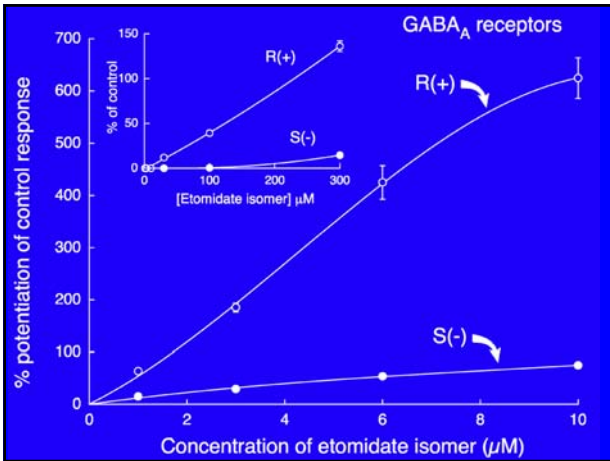




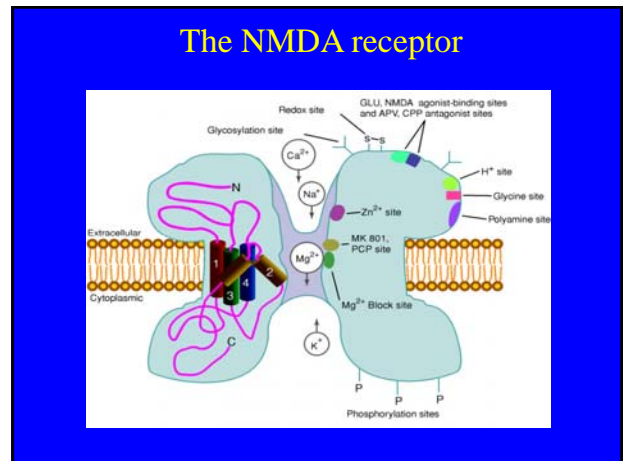
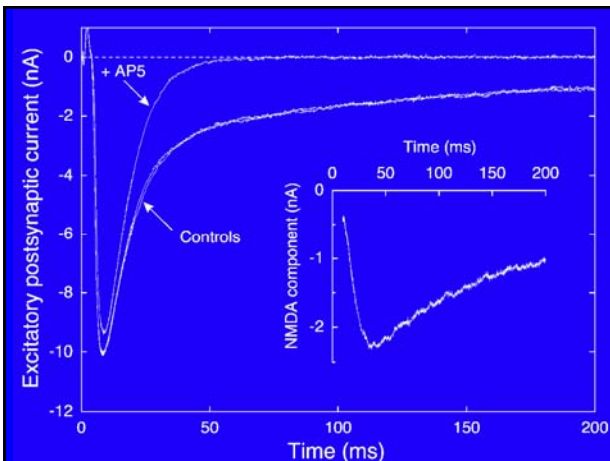
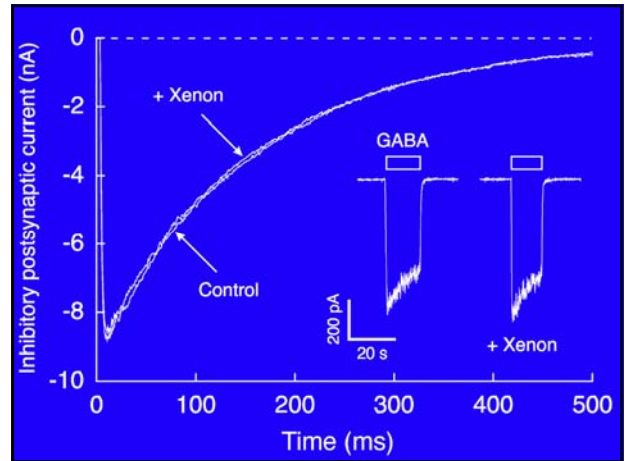


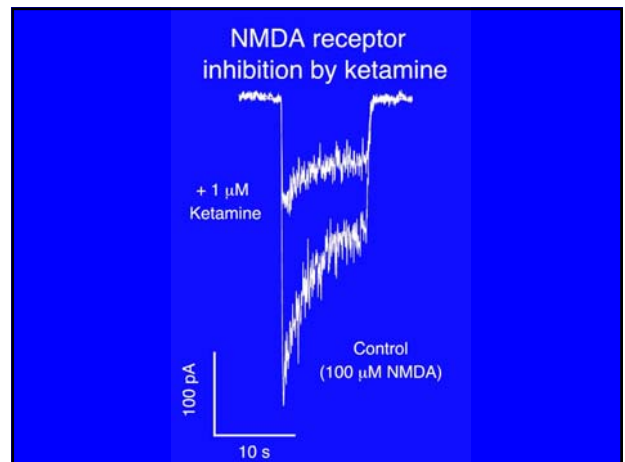
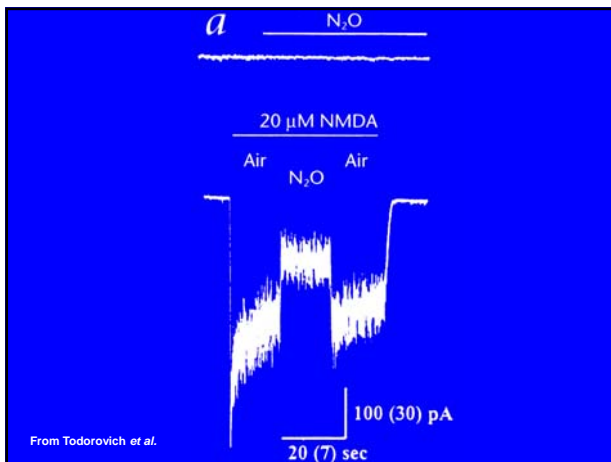
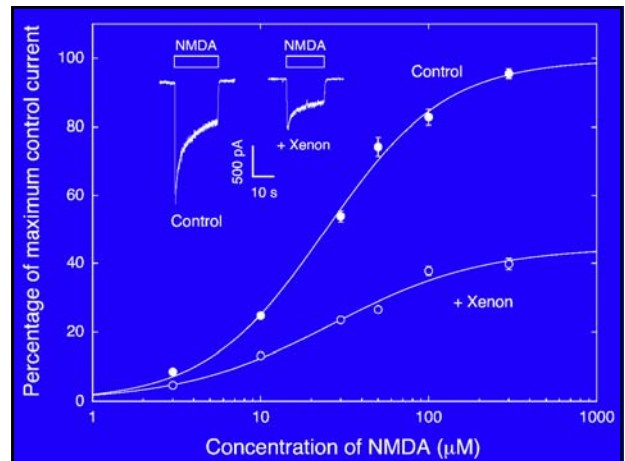
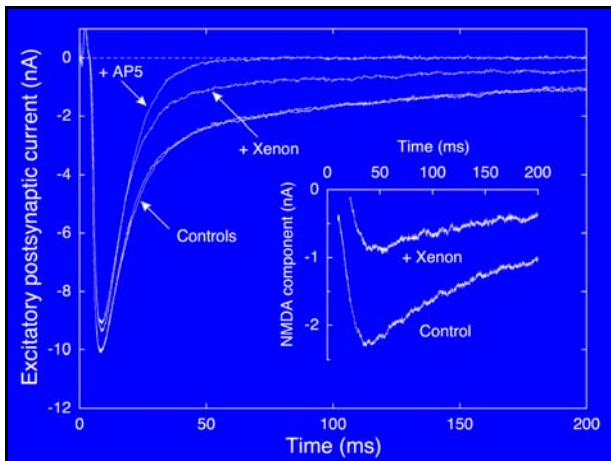
Stereoselectivity for general anaesthesia

| | |
|---|------|
| Isoflurane (Dickinson et al., 2000) | ~1.5 |
| Barbiturates (Andrews & Mark, 1982) | 2-4 |
| Ketamine (White et al., 1985) | 2-4 |
| Etomidate (Tomlin et al., 1998) | >10 |
| Neurosteroids (Wittmer et al., 1996) | >10 |



Not all anaesthetics act on GABA_A receptors





Where does xenon act on NMDA receptors?

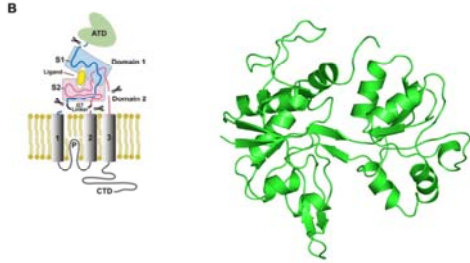
- Can molecular modelling combined with electrophysiology provide the answer?

Molecular Modelling

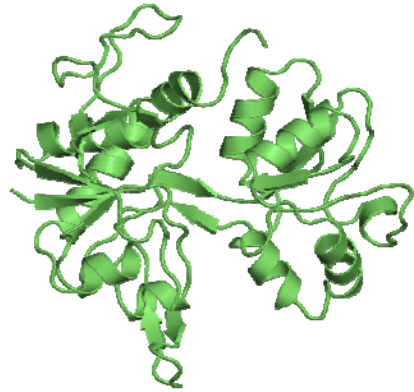
- Why might molecular modelling work for Xe binding?
- Xe is simple “noble” gas with only two relevant force/energy terms
 - van de Waals
 - charge-induced dipole
- Use GCMC simulations

Mechanisms of activation, inhibition and specificity: crystal structures of the NMDA receptor NR1 ligand-binding core

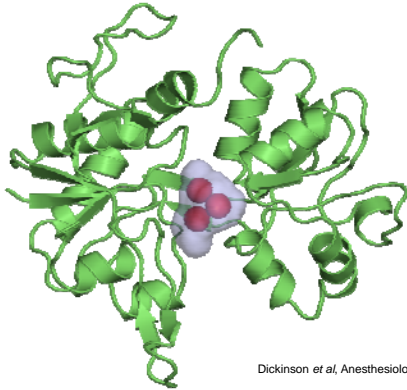
Hiroyasu Furukawa and Eric Gouaux¹
Department of Biochemistry and Molecular Biophysics and Howard Hughes Medical Institute, Columbia University, 650 West 168th Street, New York, NY 10032, USA



Structure of NMDA receptor NR1 subunit

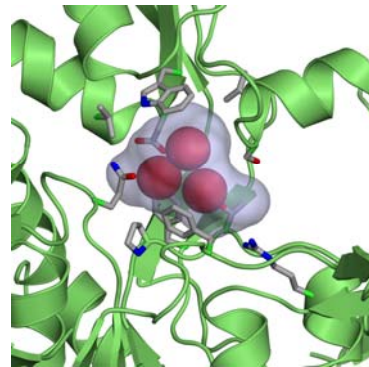


Modelling predicts xenon binds at glycine site



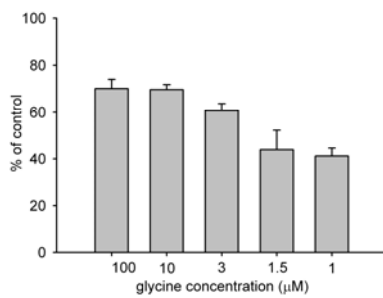
Dickinson et al, Anesthesiology, Nov 2007

Modelling predicts xenon binds at glycine site



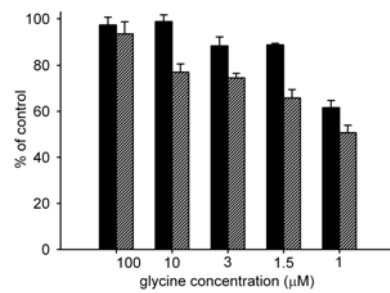
Dickinson et al, Anesthesiology, Nov 2007

Xenon inhibition increases at low glycine concentration



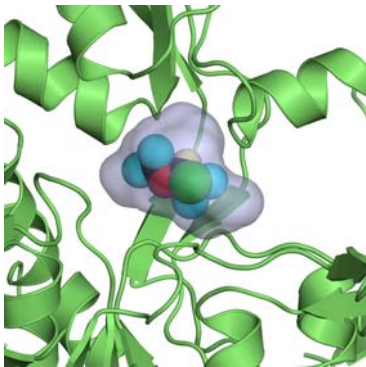
Dickinson et al, Anesthesiology, Nov 2007

Isoflurane inhibition increases at low glycine concentration



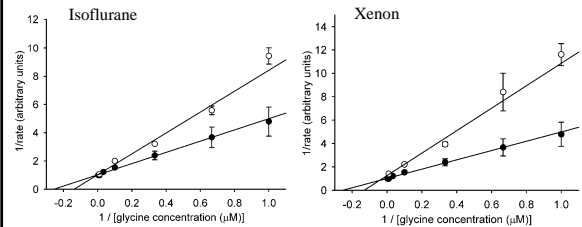
Dickinson et al, Anesthesiology, Nov 2007

Isoflurane fits in same site as xenon



Dickinson et al, Anesthesiology, Nov 2007

Lineweaver-Burk plots: quantitative analysis of competitive inhibition

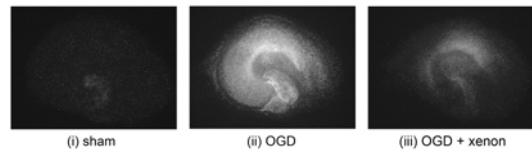


Dickinson et al, Anesthesiology, Nov 2007

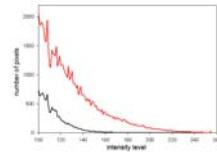
Clinical implications/Neuroprotection

- NMDA receptors critical in signalling pathways involved in cell death & neuronal injury in stroke, neonatal asphyxia & head trauma.
- NMDA receptor glycine site antagonists (e.g gavestinel) well tolerated & devoid of psychotomimetic side effects.

In-vitro neuroprotection studies

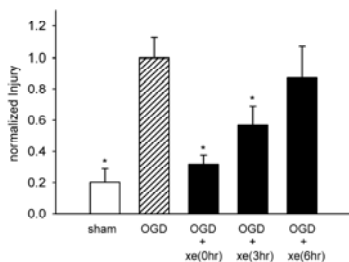


- inflict ischemic injury by OGD
- measure cell death by quantitative propidium iodide (PI) fluorescence



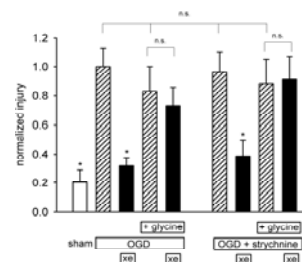
Banks, Franks & Dickinson, Anesthesiology, 2010

50% xenon protects up to 3hr post insult



Banks, Franks & Dickinson, Anesthesiology, 2010

xenon neuroprotection is reversed by adding glycine

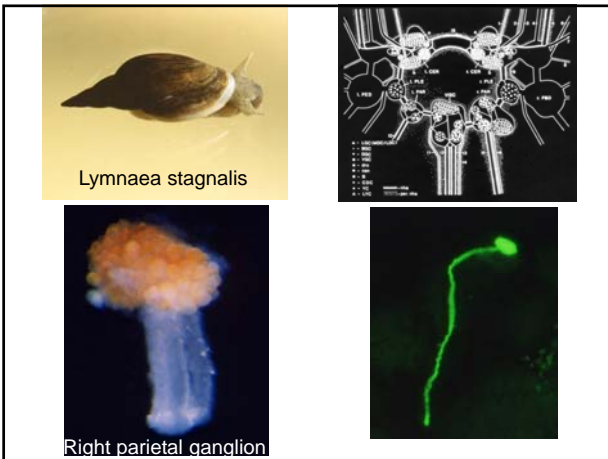


Banks, Franks & Dickinson, Anesthesiology, 2010

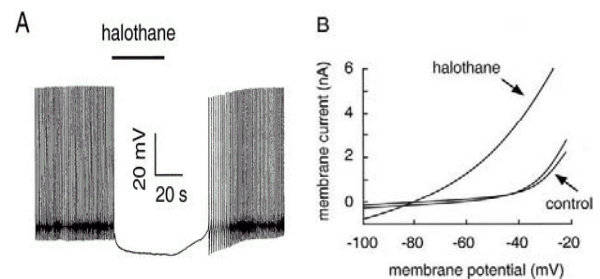
- Xenon neuroprotection against ischemia mediated by glycine-site inhibition
- Identifies NMDA receptor as target for xenon neuroprotection against ischemia
- Clinical implications
 - Glycine-site antagonists well tolerated in patients
 - Low blood/gas coefficient \Rightarrow rapid onset
 - Therapeutic window up to 3hrs post-insult
 - Neonatal asphyxia, perioperative stroke, cardiac arrest

Ion channels sensitive to general anaesthetics

- GABA_A receptor
- NMDA receptor
- 2 pore K⁺ channels
- glycine receptor – spinal chord (immobility)
- neuronal nACh receptor – function unclear (amnesia?)



Anaesthetic actions on neuronal properties.



Determinants of the Anesthetic Sensitivity of Two-pore Domain Acid-sensitive Potassium Channels

MOLECULAR CLONING OF AN ANESTHETIC-ACTIVATED POTASSIUM CHANNEL FROM *LYMNAEA STAGNALIS*
Received for publication November 17, 2006, and in revised form April 13, 2007. Published, JBC Papers in Press, June 4, 2007; DOI: 10.1074/jbc.M610692.000

Isabelle Andres-Enguix^{1,2}, Alex Caley^{1,2}, Raquel Yustos¹, Mark A. Schumacher^{1,3}, Pietro D. Spanu¹, Robert Dickinson^{1,2}, Mervyn Maze^{1,2}, and Nicholas P. Franks^{1,2}

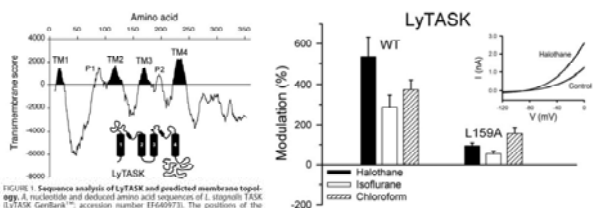


FIGURE 1. Sequence analysis of LyTASK and predicted membrane topology. A, nucleotide and deduced amino acid sequences of *L. stagnalis* LyTASK (LyTASK, GenBank™ accession number EF649973). The positions of the

Lymnaea – the movie

Further reading:

see reference list on handout