Neurotransmitters in the Brain: from rapid information transfer to long term synaptic plasticity



Molecular basis of memory







STAGE 2

NEUROTRANSMITTERS

• Provide enormous diversity in variety of transmitters and their receptors.

•Amino acids (e.g. glutamate, gamma amino butyric acid [GABA], glycine [gly]), amines (e.g. noradrenaline [NA] and dopamine [DA]) and neuropeptides (e.g. opioid peptides).

- May mediate rapid (µs ms) or slower effects (ms)
- •Vary in abundance from mM to nM in CNS tissue •Neurones receive multiple transmitter influences

which are integrated to produce diverse responses





Protein complex formation between vesicle, membrane and cytoplasmic proteins enables both vesicle docking and a rapid response to Ca entry leading to membrane fusion and exocutorsis.







Neurotransmitter action is defined by receptor kinetics



Glutamate, gamma amino butyric acid (GABA), glycine nicotinic receptors



muscarinic receptors, dopamine (DA), noradrenaline (NA), 5-hydroxytryptamine (5HT) and neuropeptides



GABA_A receptors mediate most fast INHIBITORY responses

 Activate chloride ion conductance Somatic location - profound effect •Subunits encoded by 17 genes: α (6), β (4), $\gamma(4)$, delta(1) and rho(2) subtypes Each subunit contributes a unique property and exhibits a distinct pattern of distribution: a1 is most abundant, a3 forebrain, a6 cerebellum

The most common native receptor contains $\alpha 1$, $\beta 2$ and $\gamma 2$

GABA_A receptor subunits possess different properties

- GABA affinity ($\mu M \text{ to } mM$) defined by the α subunit
- Each subunit confers a different response to BZ, Barbiturates, ethanol, steroids

 BZ modulation requires β1 Ethanol modulation most evident in: $\alpha 6 \beta 2 \gamma 2_1$ receptors which are localised in cerebellum (motor incoordination)



GLUTAMATE RECEPTORS

 Excitatory transmission in primarily mediated by Ionotropic glutamate receptors (iGLURs)

 iGLURs mediate basic information processing and underlie changes in synaptic efficacy

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•e.g. learning and memory, developing and maintaining cellular connections,pain perception

Receptors are composed of 4 subunits, each with 3 TMs and a loop



There are 3 main glutamate receptors, each having a unique role

AMPA - main receptor mediating fast CNS transmission (GluR1,2,3,4) NMDA - coincidence detection and synaptic adaptation (NR1, NR2A, B, C, D, NR3)

Kainate - modulatory role at pre and post synaptic sites (GluR5,6,7, KA1,2). Agonists are potent convulsants and environmental neurotoxins.

AMPA = (S)- -Amino-3-hydroxy-5-methyl-4isoxazole propionic acid, NMDA = N-methyl-D-aspartate





Blocked by extracellular Mg2+ in a

strongly voltage dependent manner

The native **AMPA** receptor is a tetrameric (dimer of dimers) complex formed from GLUR1, GLUR2, GLUR3 and GLUR4

 Most receptors contain GLUR2 which confers Ca²⁺ impermeability (typical of AMPA receptors in vivo)
 Expression of GLUR2 with either GLUR1 or GLUR3 yields a receptor with little divalent Ca²⁺ permeability (Equivalent to AMPA receptor response in situ in pyramidal cells).

Expression of GLUR1 or GLUR3 alone or in combination yields a Ca²⁺ permeable channel (unlike the native receptor !)

GLUR2 has functional dominance



Na⁺ Ca²⁺ Na⁺ In GluR2 there is a glutamine residue (Q) present in the primary genomic sequence which is edited to arginine R. This confers the change in property from calcium permeable to calcium impermeable.

GluR2 editing is ~100% in adult mammalian brain

Editing requires:

RNA EDITING

formation of double stranded RNA around the critical region with adjacent pairing in the intronic region
Double stranded RNA adenosine deaminase (CAG to CIG)

GluR2 determines calcium permeability



Q/R editing is essential in the adult for mediating physiological excitation

Transgenic mice in which the GluR2 editing site has been mutated

Calcium permeability increased Severe epileptic seizures Cell death in hippocampus Mice die at 3 weeks

Rapid excitatory transmission: NMDA receptors

- Strong afferent stimulation is needed to produce sufficient depolarisation to remove the Mg²⁺ block and activate N-methyl-D-aspartate receptors causing Ca²⁺ and Na⁺ influx.
- NMDA receptors are critical for the induction of certain forms of synaptic plasticity
- Receptors associate with other signalling molecules e.g. kinases to mediate their postsynaptic response.

NMDA receptors (NR1 + NR2A-D/NR3)



Receptors with NR2B subunits instead of NR2A stay open longer - functional consequences? NR2B-receptors have a longer phase of memory activation increasing the window during which coincidence detection can occur. Animals with increased expression of the NR2B subunit have an enhanced ability to learn (see later).

The modulatory NR2 subunits have profound effects on receptor kinetics

Cull-Candy & Leszkiewicz (2004)

Ionotropic Glutamate receptors are formed from subunits which are encoded by 18 genes which gives structural DIVERSITY

- due to multiple subunit combinations
- **RNA** editina
- multiple splice variants
- post-translational modifications

which translates into functional DIVERSITY

- ion channel properties
- regional specificity
- developmental specificity
- adaptation to synaptic activity

STAGE 3 Integration of Signals

The NMDA synapse provides a "molecular model of associative memory"

If two neurones are excited at the same time then the active synapse between them is strengthened (Hebb's rule).

COINCIDENCE DETECTION

•A change in the efficiency of synaptic transmission ("Long term potentiation"- LTP.)

•Occurs during development, learning new skills,

responding to environmental changes (harmless/noxious)

Long-term potentiation (LTP)

Bliss and Lømo (1973) J. Physiol. 232: 331-356

Long lasting effects require protein synthesis

Evoked

in the

potentials

hippocampus



Spatial navigation learning in rats depends on a hippocampal "map of spatial relations"

The hippocampus is a major site of learning:

Morris water maze

•NMDA R antagonists impair learning Several forms of ASSOCIATIVE memory are mediated by GLUTAMATE at the NMDA receptor and these are widely used models

Effects are long lasting and depend on protein synthesis.



The rat finds a platform

NMDA receptor-mediated Long Term Potentiation in the hippocampus provides a model of learning

Dendritic spine changes associated with hippocampal long-term synaptic plasticity Florian Engert & Tobias Bonhoeffer











High frequency stimulation leads to Ca²⁺ gating through the NMDA receptor

- CaMKII (calcium and calmodulin dependent kinase) is activated by Ca²⁺
- CaMKII is autophosphorylated (Thr286) and translocates to the subsynaptic region. Point mutation inhibits LTP and memory formation
- Transient Ca²⁺ signal prolongs kinase activity until dephosphorylated by protein phosphatase

What is the molecular process ?



CaMKIV-dominant negative mutants have normal E-LTP but impaired L-LTP and impaired memory consolidation Nuclear calmodulin inhibition impairs LTP and LTM













G-protein-coupled receptors mediate the action of many neurotransmitters

Amines: Dopamine, noradrenaline, 5-hydroxy-tryptamine (5-HT), muscarinic cholinergic mACh

Amino acids: GABA_B metabotropic glutamate mGLUR Neuropeptides: somatostatin, substance P, enkephalin

•~1000 G-protein coupled 2. receptors in the genome (many are 'orphan') - All have 7 alpha helices

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RECEPTOR

• G-proteins have 3 subunits 16 alpha subunits define transduction system, 5 beta, 11 gamma subunits

 Effectors: Adenyl cyclase, Phospholipase C (PLC), cGMP phosphodiesterase, Phopholipase A2, K⁺ or Ca²⁺ channels



Many GPCR actions are nervous system specific



Targets for phosphorylation

- lon channels * cell excitability, increases (activation) or decreases (inactivates) open probability
- Enzymes affects transmitter availability
- Receptors facilitate or desensitise transmitter response
- Transcription factors (e.g. CREB) cause long term effects on protein synthesis

* Direct gating of ion channels by G-proteins also occurs.

Dopamine pathways in the brain



5-hydroxytryptamine pathways in the brain



DA receptors:

D1 and D5 (D1-type) stimulate cAMP

D2, D3 and D5 (D2-type) inhibit cAMP/ increase IP3



less well characterised.

Noradrenergic pathways in the brain



Adrenoceptors:

Beta receptors stimulate cAMP (G_S) Alpha₂ inhibit cAMP (G_{i/o}) Alpha_ increase IP3 (Gq)