AOP Title

Binding of antagonist to N-methyl-D-aspartate receptors (NMDARs) during brain development induces impairment of learning and memory abilities

Short name: Binding of antagonist to NMDARs impairs cognition

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OECD Project 1.22: The Adverse Outcome Pathway from Binding of Antagonists to NMDAR During Brain Development (Synaptogenesis) Induces Impairment of learning and Memory Abilities

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Abstract

It is well documented and accepted that learning and memory processes rely on physiological functioning of the glutamate receptor N-methyl-D-aspartate (NMDAR). Both animal and human studies investigating NMDA itself, experiments with NMDAR antagonists and mutant mice lacking NMDAR subunits strongly support this statement (Rezvani, 2006). Activation of NMDARs results in long-term potentiation (LTP), which is related to increased synaptic strength, plasticity and memory formation in the hippocampus (Johnston et al., 2009). LTP induced by activation of NMDA receptors has been found to be elevated in the developing rodent brain compared to the mature brain, partially due to ‘developmental switch’ of the NMDAR 2A and 2B subunits (Johnston et al., 2009). Activation of the NMDAR also enhances brain derived neurotrophic factor (BDNF) release, which promotes neuronal survival, differentiation and synaptogenesis (Tyler et al., 2002; Johnston et al., 2009). Consequently, the blockage of NMDAR by chemical substances during synaptogenesis disrupts neuronal network formation resulting in the impairment of learning and memory processes (Toscano and Guilarte, 2005). This AOP is relevant to developmental neurotoxicity (DNT). The molecular initiating event (MIE) is described as the binding of antagonist to NMDAR in neurons during synaptogenesis (development) in hippocampus (one of the critical brain structures for learning and memory formation). Chemicals that block NMDAR are known developmental neurotoxicants such as lead (Pb\textsuperscript{2+}), toluene and ethanol. Additionally there are some specific pharmaceuticals such as MK-801, ketamine, dextromethorphan, pethidine, methadone, dextropropoxyphene, tramadol and ketobemidone that are selective antagonists of NMDAR too.
Summary of the AOP

Molecular Initiating Event

**Molecular Initiating Event**

**NMDARs, Binding of antagonist**

**Short name:** NMDA receptors, Binding of antagonist

**How this Key Event works**

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**Biological state:** L-glutamate (Glu) is a neurotransmitter with important role in the regulation of brain development and maturation processes. Two major classes of Glu receptors, ionotropic and metabotropic, have been identified. Due to its physiological and pharmacological properties, Glu activates three classes of ionotropic receptors named: α-amino-3-hydroxy-5-methyl-4-isoxazolopyridine (AMPA receptors), 2-carboxy-3-carboxymethyl-4-isopropylpyrrolidine (kainate receptors) and N-methyl-D-aspartate (NMDA receptors, NMDARs), which transduce the postsynaptic signal. Ionotropic glutamate receptors are integral membrane proteins formed by four large subunits that compose a central ion channel pore. In case of NMDA receptors, two NR1 subunits are combined with either two NR2 (NR2A, NR2B, NR2C, NR2D) subunits and less commonly are assembled together with a combination of NR2 and NR3 (A, B) subunits (reviewed in Traynelis et al., 2010). To be activated NMDA receptors require simultaneous binding of both glutamate to NR2 subunits and of glycine to either NR1 or NR3 subunits that provide the specific binding sites named extracellular ligand-binding domains (LBDs). Apart from LBDs, NMDA receptor subunits contain three more domains that are considered semiautonomous: 1) the extracellular amino-terminal domain that plays important role in assembly and trafficking of these receptors; 2) the transmembrane domain that is linked with LBD and contributes to the formation of the core of the ion channel and 3) the intracellular carboxyl-terminal domain that influences membrane targeting, stabilization, degradation and post-translation modifications.

**Biological compartments:** The genes of the NMDAR subunits are expressed in various tissues and are not only restricted to the nervous system. The level of expression of these receptors in neuronal and non-neuronal cells depends on: transcription, chromatin remodelling, mRNA levels, translation, stabilization of the protein, receptor assembly and trafficking, energy metabolism and numerous environmental stimuli (reviewed in Traynelis et al., 2010).

In hippocampus region of the brain, NR2A and NR2B are the most abundant NR2 family subunits. NR2A-containing NMDARs are mostly expressed synaptically, while NR2B-containing NMDARs are found both synaptically and extrasynaptically (Neal and Guilarte, 2010).

**General role in biology:** NMDA receptors, when compared to the other Glu receptors, are characterized by higher affinity for Glu, slower activation and desensitisation kinetics, higher permeability for calcium (Ca2+) and susceptibility to potential-dependent blockade by magnesium ions (Mg2+). NMDA receptors are involved in fast excitatory synaptic transmission and neuronal plasticity in the central nervous system (CNS). Functions of NMDA receptors:

1. They are involved in cell signalling events converting environmental stimuli to genetic changes by regulating gene transcription and epigenetic modifications in neuronal cells (Cohen and Greenberg, 2008).
2. In NMDA receptors, the ion channel is blocked by extracellular Mg2+ and Zn2+ ions, allowing the flow of...
Na+ and Ca2+ ions into the cell and K+ out of the cell which is voltage-dependent. Ca2+ flux through the NMDA receptor is considered to play a critical role in pre- and post-synaptic plasticity, a cellular mechanism important for learning and memory (Barria and Malinow, 2002).

3. The NMDA receptors have been shown to play an essential role in the strengthening of synapses and neuronal differentiation, through long-term potentiation (LTP), and the weakening of synapses, through long-term depression (LTD). All these processes are implicated in the memory and learning function (Barria and Malinow, 2002).

How it is Measured or Detected

Methods that have been previously reviewed and approved by a recognized authority should be included in the Overview section above. All other methods, including those well established in the published literature, should be described here. Consider the following criteria when describing each method: 1. Is the assay fit for purpose? 2. Is the assay directly or indirectly (i.e. a surrogate) related to a key event relevant to the final adverse effect in question? 3. Is the assay repeatable? 4. Is the assay reproducible? There is no OECD advised method for measuring NMDA receptor binding of antagonists. However, there are methods described in the scientific literature that allow measuring:

1. **Ex vivo:** The most common assay used is the NMDA receptor (MK801 site) radioligand competition binding assays (Reynolds, 2001; Gao et al., 2013; http://pdsp.med.unc.edu/UNC-CH%20Protocol%20Book.pdf; http://www.currentprotocols.com/WileyCDA/CPUnit/refId-ph0120.html). This assay is based on the use of the most potent and specific antagonist of this receptor, MK801 that is used to detect and differentiate agonists and antagonists that bind to this specific site of the receptor.

2. **In silico:** The prediction of NMDA receptor targeting is achievable by combining database mining, molecular docking, structure-based pharmacophore searching, and chemical similarity searching methods together (Koutsoukos et al., 2011; Gao et al., 2013).

Evidence Supporting Taxonomic Applicability

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The evolution of NMDAR subunits (NR1, NR2, NR3) is well-conserved throughout different species from lower organism to mammals, including humans (Ewald and Cline, 2009; Tikhonov and Magazanik, 2009; Koo and Hampson, 2010; Teng et al., 2010; Flores-soto et al., 2012).

Many of the binding sites for the noncompetitive or competitive antagonists e.g. for binding of dizocilpine (MK-801), phencyclidine, D-2-amino-5-phosphonopentanoate (AP5) and 3-((R)-2-carboxypiperazin-4-yl)-propyl-1-phosphonic acid (R-CPP) are also conserved in Drosophila (reviewed in Xia and Chiang, 2009).

Cellular membranes can be prepared from different brain areas of distinct species. Using [3H]MK-801, high affinity binding sites for MK-801 were detected in membranes of the rat brain (Woodruff et al., 1987). The same binding assay has been used in preparations from human brains mostly by patients with neurodegenerative disorders (Slater et al., 1993) as well as from different marine, avian species (Scheuhammer et al., 2008) and insects (Eldefrawi et al., 1993).

Evidence for Chemical Initiation of this Molecular Initiating Event

Glu and glycine are endogenous agonists that bind to LBD of specific NMDA receptor subunits. In this binding site numerous competitive exogenous antagonists have been identified to cause closure of binding
site and inhibition of NMDA receptor (reviewed in Traynelis et al., 2010). Here, are listed some known competitive antagonists for NMDA receptor, some of them are specific to NR1 subunit and some to NR2 subunit: α-AA, α-aminoadipate;

5,7-DCKA, 5,7-dichlorokynurenic acid;

7-CKA, 7-chlorokynurenic acid;

ACEA-1011, 5-chloro-7-trifluoromethyl-1,4-dihydro-2,3-quinoxalinedione;

ACEA-1021, licostinel;

AP5, 2-amino-5-phosphonopentanoate;

AP7, 2-amino-7-phosphonopentanoate;

CGP-61594, (±)-trans-4-[2-(4-azidophenyl)acetylamino]-5,7-dichloro-1,2,3,4-tetrahydroquinoline-2-carboxylic acid;

CGP-40116, d-(E)-2-amino-4-methyl-5-phosphono-3-pentenoic acid;

CGP-43487, d-(E)-2-amino-4-methyl-5-phosphono-3-pentenoic acid methyl ester;

CGP-58411, 7-chloro-4-hydroxy-3-phenyl-1H-quinolin-2-one;

CGS-19755, (2R,4S)-4-(phosphonomethyl)piperidine-2-carboxylic acid;

CPP, 4-(3-phosphonopropyl) pizervazine-2-carboxylic acid;

GV150,526A, gavestinel;

GV196,771A, (E)-4,6-dichloro-3-[(2-oxo-1-phenyl-3-pyrrolidinylidene)methyl]-1H-indole-2-carboxylic acid;

L-689,560, 4-trans-2-carboxy-5,7-dichloro-4-phenylaminocarbonylamino-1,2,3,4-tetrahydroquinoline;

L-701,324, 7-chloro-4-hydroxy-3-(3-phenoxy)phenyl-2(1H)-quinolone;

MDL105,519, (E)-3-(2-phenyl-2-carboxyethyl)-4, 6-dichloro-1H-indole-2-carboxylic acid;

PBPD, (2S,3R)-1-(biphenyl-4-carbonyl)piperazine-2,3-dicarboxylic acid;

PMPA, (R,S)-4-(phosphonomethyl)-piperazine-2-carboxylic acid;

PPDA, (2S,3R)-1-(phenanthren-2-carbonyl)piperazine-2,3-dicarboxylic acid

Besides competitive antagonists, noncompetitive antagonists have also been designed like phenylethanolamine ifenprodil that interacts with the NR2B extracellular amino-terminal domain. It has been suggested that they act by stabilizing an agonist-bound state in which the receptor has a low open probability. Other more potent derivatives of ifenprodil are: α-(4-hydroxyphenyl)-β-methyl-4-(phenylmethyl)-1-piperidine propanol (Ro 25-6981), 1-[2-(4-hydroxy-phenoxy)-ethyl]-4-(4-methyl-benzyl)-piperidin-4-ol (Ro 63-1908), besonprodil (CI-1041), and traxoprodil mesylate (CP-101,606). Ethanol has been proposed to be a noncompetitive antagonist of NMDA receptors, binding to NR2 subunit (Nagy, 2008). Inhibition of NMDA receptor function by ethanol and interactions between EtOH and the noncompetitive NMDA receptor antagonist ifenprodil have been examined in neocortical neurons from rat and human embryonic kidney (HEK) 293 cells expressing recombinant NMDA receptors (Lovinger, 1995). Recently, a structural model has been suggested that predicts the presence of four sites of ethanol action on the NMDA receptor, each containing four pairs of positions in the NR1/NR2 subunits (reviewed in Chandrasekar, 2013). Some other antagonists can become trapped in the pore of the NMDA receptor after channel closure and
these antagonists are called uncompetitive or trapping blockers. The most well studied NMDA receptor uncompetitive antagonists are Mg2+, polyamines, phencyclidine, ketamine, MK-801, memantine, amantadine, pentamidine, 9-tetrahydroaminoacridine, dextromethorphan, and its metabolite dextrorphan. MK-801 has been shown to prevent toluene-induced alterations in pattern-elicited visual-evoked potentials in vivo, suggesting the possibility that the binding site of toluene might be common with the one of MK-801 (Bale et al., 2007). However, another study suggests that toluene interference with the NMDA receptor might not be exclusively because of the binding to the channel pore (Smothers and Woodward, 2007) but it may involve some other binding sites. Pb2+ is considered a voltage independent antagonist of NMDA receptors and it is believed that possibly shares the same binding site with Zn2+ (reviewed in Neal and Guilarte, 2010; Traynelis et al., 2010). However, studies done in recombinant NR2A- and NR2B- containing NMDA receptors with mutated Zn2+ binding sites exhibit that additional structural elements, different from those important for Zn2+ binding are involved in Pb2+ binding site (reviewed in Neal and Guilarte, 2010). Similarly, there are contradicting experimental evidence and disagreement about Pb2+’s role as competitive or non-competitive antagonist (Neal and Guilarte, 2010).

References


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**Key Events**

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<td>Release of BDNF, Reduced</td>
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<td>Dendritic morphology, Aberrant</td>
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<td>Synaptogenesis, Decreased</td>
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<td>Neuronal network function, Decreased</td>
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<td>Presynaptic release of glutamate, Reduced</td>
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<td>Cell death, N/A</td>
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**Calcium influx, Decreased**

**Short name: Calcium influx, Decreased**

**How this Key Event works**
**Level of Biological Organization**

**Cellular**

**Biological state:** Under physiological resting conditions of the cell, the free intracellular Ca²⁺ reaches around 100 nM, whereas the extracellular Ca²⁺ can be found at higher concentrations of 1.2 mM that under certain stimulus may invade the cell (Berridge et al, 2000). Six to seven oxygen atoms surround Ca²⁺, whereas the protein chelator of Ca²⁺ is the EF motif that is present in many proteins such as calmodulin (Clapham, 2007). The EF-hand is a helix-loop-helix calcium-binding motif in which two helices pack together at an angle of approximately 90 degrees (Lewit-Bentley and Réty, 2000). The two helices are separated by a loop region where calcium actually binds. The EF notation for the motif is derived from the notation applied to the structure of parvalbumin, in which the E and F helices were originally identified as forming this calcium-binding motif.

**Biological compartments:** Ca²⁺ ions accumulate in the cytoplasm, cellular organelles (e.g. mitochondria and endoplasmic reticulum) and nucleus in response to diverse classes of stimuli.

**General role in biology:** In order to adapt to altered stimulus from exposure to different environmental factors, cells require signal transmission. However, signalling needs messengers whose concentration is modified upon stimulus (Clapham, 2007). Ca²⁺ ions act as an important intracellular messenger playing the role of ubiquitous signalling molecules and consequently regulate many different cellular functions (Berridge, 2012; Hagenston and Bading, 2011). Given its important role in processes that are fundamental to all cell types, Ca²⁺ homeostasis is tightly regulated by intracellular and extracellular mechanisms (Barhoumi et al., 2010). Intracellular Ca²⁺ concentration is regulated by opening or closing channels in the plasma membrane. Additionally, the Ca²⁺-ions can be released from intracellular stores of the endoplasmic reticulum (ER) through ryanodine receptors (RYRs) or inositol 1,4,5-trisphosphate receptors (InsP3Rs). Ca²⁺ homeostasis is also regulated by the mechanisms that remove Ca²⁺ from the cytosol, for example pumps in both cell membrane and ER membrane. In addition, cytosolic Ca²⁺ regulation involves accumulation of Ca²⁺ in mitochondria that have the capacity to buffer the excess of cytoplasmic Ca²⁺-ions. In neurons, Ca²⁺ ions regulate many critical functions. Firstly, they contribute to dendritic electrical signalling, producing postsynaptic depolarization by the current carried by Ca²⁺-ions. Secondly, Ca²⁺ activates Ca²⁺-sensitive proteins such as different kinases, calcineurin and calpain, triggering signalling pathways critical for cell physiology. Modification of the gene transcription is the final outcome of the Ca²⁺-ions impact on long-term modifications affecting neurotransmitters release (reviewed in Neher and Sakaba, 2008), neuronal differentiation, synapse function and cell viability (Clapham, 2007; Higley and Sabatini, 2012). Thus, the Ca²⁺ that enters and accumulates in cytoplasm and nucleus is a central signalling molecule that regulates synapse and neuronal cell function, including learning and memory processes (Berridge, 2012; Hagenston and Bading, 2011).

**How it is Measured or Detected**

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No OECD method is available to measure intracellular Ca²⁺.

In vitro, well-established flow cytometric or high content imaging analysis with specific fluorescent dyes (Ca²⁺-sensitive fluorophores) such as Fura-2, Oregon Green-BAPTA, Fluo-4 and X-Rhod exist for determination of intracellular Ca²⁺ concentration. The use of different fluorometric calcium indicators in neuroscience and neurotoxicology have been recently reviewed by Grienberger and Konnerth (2012).

Barhoumi et al. 2010 summarised all the methods to measure cytosolic Ca²⁺ alterations due to exposure to neurotoxic compounds, including steady state, short-term kinetic measurements of stimulated Ca²⁺
transients and dynamic measurements. This paper further discusses the strengths and weaknesses of each approach in intracellular Ca2+ measurements and its applicability in high throughput screening.

For quantitative estimation of Ca2+ in dendritic spines, besides of Ca2+-sensitive fluorophores the use of two-photon released caged neurotransmitters has been suggested as it allows direct stimulation of visualized spines (Higley and Sabatini, 2012). In Higley and Sabatini 2012 further technical information can be found in relation to study Ca2+ in dendritic spines.

Furthermore, there are three methods for measuring Ca2+ influx in NMDA receptors that involve the measurement of 1) relative Ca2+ permeability, 2) channel blockage by Ca2+, and 3) fractional Ca2+ currents from whole-cell currents determined in the presence of high concentrations of intracellular Fura-2 (Traynelis et al., 2010).

In vivo, two-photon Ca2+ imaging using Ca2+-sensitive fluorescent indicators that measure changes in intracellular Ca2+ concentration as a readout for suprathreshold and subthreshold neuronal activity has also been used to study learning and memory in live rodents (Chen et al., 2013) The last two decades the neuronal function of the larval and adult zebrafish has been extensively studied using Ca2+ imaging methods. By applying simple Ca2+ indicators such as dextran or acetoxy methyl esters to more powerful genetically encoded Ca2+ indicators, zebrafish provides a transparent model where live Ca2+ imaging can be successfully achieved (Kettunen, 2012).

Evidence Supporting Taxonomic Applicability

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<td>Mus gratus</td>
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<td>zebrafish</td>
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Ca2+ homeostatic system is known to be highly conserved throughout evolution and is present from humans to invertebrates (Case et al., 2007).

References


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**NMDARs, Inhibition**

Short name: NMDARs, Inhibition

**How this Key Event works**

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**Biological state:** Please see MIE NMDARs, Binding of antagonist

**Biological compartments:** Please see MIE NMDARs, Binding of antagonist

**General role in biology:** Please see MIE NMDARs, Binding of antagonist

**How it is Measured or Detected**

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No OECD methods are available to measure the activation state of NMDA receptors.

The measurement of the activation or the inhibition of NMDA receptors is done indirectly by recording the individual ion channels that are selective to Na+, K+ and Ca+2 by the patch clamp technique. This method relies on lack of measurable ion flux when NMDA ion channel is closed, whereas constant channel specific conductance is recorded at the open state of the receptor (Blanke and VanDongen, 2009). Furthermore, this method is based on the prediction that activation or inhibition of an ion channel results from an increase in the probability of being in the open or close state, respectively.

The whole-cell patch clamp recording techniques have also been used to study synaptically-evoked NMDA receptor-mediated excitatory or inhibitory postsynaptic currents (EPSCs and IPSCs, respectively) in brain
slices and neuronal cells, allowing the evaluation of the activated or inhibited state of the receptor.

Microelectrode array (MEA) recordings are also used to measure electrical activity in cultured neurons in response to NMDA receptor activation or inactivation (Keefer et al., 2001, Gramowski et al., 2000 and Gopal, 2003; Johnstone et al., 2010). MEAs can also be applied in higher throughput platforms to facilitate screening of numerous chemical compounds based on electrical activity measurements (McConnell et al., 2012).

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The cellular expression of the NMDAR subunits has been studied in both adult human cortex and hippocampus (Scherzer et al., 1998) as well as during the development of the human hippocampal formation (Law et al., 2003). The whole-cell patch clamp recording techniques have been used in NMDA receptors expressed in human TsA cells (derivative of the human embryonic kidney cell line HEK-293) (Ludolph et al., 2010). Cell-attached single-channel recordings of NMDA channels has been carried out in human dentate gyrus granule cells acutely dissociated from slices prepared from hippocampi surgically removed from human patients (Lieberman and Mody, 1999).

It is important to note that in invertebrates the glutamatergic synaptic transmission has inhibitory and not excitatory role like in vertebrates. This type of neurotransmission is mediated by glutamate-gated chloride channels that are members of the ‘cys-loop’ ligand-gated anion channel superfamily found only in invertebrates. The subunits of glutamate-activated chloride channel have been isolated from C. elegans and from Drosophila (Blanke and VanDongen, 2009).

**References**


Release of BDNF, Reduced
Release of BDNF, Reduced

How this Key Event works

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**Biological state:** BDNF belongs to a family of closely related neurotrophic factors named neurotrophins and is widely expressed in the developing and mature CNS. In the rodent cortex, postnatal BDNF expression is initially low but slowly increases to reach high levels around weaning. Therefore, BDNF expression peaks at a time when both structural and functional maturation of cortical circuitry occurs. During postnatal development, BDNF levels are dynamically regulated, in part by neuronal activity dependent mechanisms (Waterhouse and Xu, 2009). Glutamate has been shown to increase the transcription and release of BDNF. Indeed, BDNF is synthesized, stored and released from glutamatergic neurons (Lessmann et al., 2003).

**Biological compartments:** BDNF initially is synthesized as precursor proteins (proBDNF), which is processed intracellularly to be transformed in its mature form (mBDNF) after proteolytically cleaved in the synaptic cleft by plasmin which is a protease activated by tissue plasminogen activator (tPA) (Cohen-Cory et al., 2010). proBDNF is constantly secreted while tPA release and mBDNF production depends on neuronal excitation (Head et al., 2009). Storage and activity-dependent release of BDNF has been demonstrated in both dendrites and axon terminals (Waterhouse and Xu, 2009). More specifically, in hippocampus, BDNF appears to be stored in dendritic processes of neurons (Balkowiec and Katz, 2002). BDNF is abundant in cerebellum and cortex and has also been measured in cerebrospinal fluid (CSF) (Zhang et al., 2013), whole blood, plasma, serum (plasma without clotting factors) and platelets (Trajkovska et al., 2007). BDNF can has been found be produced by astrocytes under both physiological and pathological conditions (Endo, 2005; Coco et al., 2013; Nelson and Alkon, 2014).

In humans, mBDNF is sequestered in platelets, consequently BDNF can reach all tissues and organs. Lymphocytic cells have been shown to express BDNF in vitro similarly to eosinophils, dendritic cells, and endothelial cells. The visceral and airway epithelium are also significant sources of BDNF. Female reproductive system including ovaries, placenta and uterus also express BDNF (Wessels et al., 2014).

**General role in biology:** The biological functions of mBDNF are mediated by binding to tyrosine kinase B (TrkB) receptor that leads to the activation of three major intracellular signalling pathways, including MAPK, PI3K and PLCγ1 (Soulé et al., 2006). TrkB-mediated signaling regulates gene transcription in the nucleus through the activation of several transcription factors. These genes are involved in neurite outgrowth, synaptogenesis, synapse maturation and stabilization (Pang et al., 2004; Lu et al., 2005; Nelson and Alkon, 2014).

On the other hand, proBDNF binds to the p75 neurotrophin receptor (p75NTR) and activates RhoA, a small GTPase that regulates actin cytoskeleton polymerization leading to inhibition of axonal elongation, growth
cone collapse, and apoptosis (Dubreuil et al., 2003; Yamauchi et al., 2004; Head et al., 2009).

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No OECD methods are available to measure BDNF protein and mRNA levels. Depending on the tissue or fluid measurements distinct methods are used.

Brain tissue: BDNF protein levels can be measured by commercial available antibody sandwich ELISA kits, Western blotting, immunohistochemistry and immunofluorescence. BDNF primers for different exons are available to determine mRNA levels by RT-PCR. The Bdnf gene consists of multiple alternative exons (ten in human, eight in rodents and six in lower vertebrates), and a single exon coding for the entire pro-BDNF protein (Cohen-Cory et al., 2010).

Cerebro-spinal fluid (CSF): There are available commercial antibody sandwich ELISA kits (Trajkovska et al., 2007) and immunobead-based multiplex assays for high throughput screening (Zhang et al., 2013).

Whole blood, serum, plasma and platelets: There are several commercial double antibody sandwich ELISA kits that can be used for identification of BDNF levels in biological fluids (Trajkovska et al., 2007).

Methodological considerations that have to be taken into account during sample preparation and measurement of BDNF by ELISA have been recently reviewed in Elfving et al. 2010. A study measuring BDNF by a commercially available ELISA kit in various tissues and biological liquids derived from distinct species revealed that BDNF is undetectable in mouse blood, pig plasma BDNF levels were comparable to levels reported in humans and that there is positive correlations between blood BDNF levels and hippocampal BDNF levels in rats and pigs (Klein et al., 2011).

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Bdnf gene is well conserved among species, from avian species to fish and mammals (Heinrich and Paktakan, 2004; Aid et al., 2007; Pruunsild et al., 2007, Brenowitz, 2013).

Klein et al. (2011) examined blood, serum, plasma and brain-tissue and measured BDNF levels in three different mammalian species: rat, pig, and mouse, using an ELISA method, whereas Trajkovska et al. 2007 determined BDNF levels in human blood.

References


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Dendritic morphology, Aberrant

How this Key Event works

Level of Biological Organization
Cellular

Biological state: After becoming post-mitotic and during the differentiation process, neuronal cells, apart from interneurons, undergo lengthening, branching, dendrite and dendritic spine formation (Scott and Luo, 2001). In human, dendrites appear as early as 13.5 weeks gestation in the subplate neurons while arborization begins only after 26 weeks (Mrzljak et al., 1988 and Mrzljak et al., 1990). In rodents, during the first postnatal week, both pyramidal and nonpyramidal neurons go through extensive and fast dendrite growth, branching, and elaboration. Dendrite arbor’s capacity and complexity continue to increase in the second and third postnatal week, however, much slower. During the same developmental window, dendritic spines begin to appear as a group. The first spines look like filopodia (Dailey and Smith, 1996; Fiala et al., 1998). Filopodia can grow and retract within seconds to minutes, permitting them to explore and identify appropriate presynaptic targets (Dailey and Smith, 1996). As dendrite spines mature, these long and thin structures change and the spines shorten and acquire a bulbous ending or ‘head’ (Dailey and Smith, 1996). At this final stage of dendrite growth, a neuron possesses a dynamic dendrite tree, which has a greater potential for connectivity and synapse creation because of dendritic spine formation.

Biological compartments: Dendritic morphology determines many aspects of neuronal function, including
action potential propagation and information processing. Postsynaptic density-95 (PSD-95), a protein involved in dendritic spine maturation and clustering of synaptic signalling proteins, plays a critical role in regulating dendrite outgrowth and branching, independent of its synaptic functions. In immature neurons, over-expression of PSD-95 decreases the proportion of primary dendrites that undergo additional branching, resulting in a marked reduction of secondary dendrite number. Conversely, knocking down PSD-95 protein in immature neurons increases secondary dendrite number. Binding of cypin to PSD-95 (that regulates PSD-95 location) correlates with formation of stable dendrite branches. Finally, overexpression of PSD-95 in COS-7 cells disrupts microtubule organization, indicating that PSD-95 may modulate microtubules to regulate dendritic branching. Proteins primarily involved in synaptic functions can also play developmental roles in shaping how a neuron patterns its dendrite branches (Komau et al., 1995). New spines containing PSDs are formed by conversion of dynamic filopodia-like spine precursors in which PSDs appeared de novo, or by direct extension of spines or spine precursors carrying preformed PSDs from the shaft. PSDs are therefore highly dynamic structures that can undergo rapid structural alteration within dendrite shafts, spines and spine precursors, permitting rapid formation and re-modelling of synaptic connections in developing CNS tissues.

Dendritic spines are important sites of excitatory synaptic transmission and changes in the strength of these synapses are likely to underlie important higher brain functions such as learning and memory. Spines form biochemical compartments for isolating reactions that occur at one synapse from those at other synapses thereby providing a possible way to ensure the specificity of connections between neurons in the brain.

The stages of dendrite development have been clearly described in neurons located in the developing rodent cortex and hippocampus (Dailey and Smith, 1996; Fiala et al., 1998; Redmond, 2008) and human prefrontal cortex (Mrzljak et al., 1988; Mrzljak et al., 1990).

General role in biology: Functionally, dendrites serve as post-synaptic part of a synapse, playing a critical role in the processing of information transmitted through synapses. They receive the majority of synaptic inputs comparing to the soma or the axon. Consequently, it is not surprising that postsynaptic activity is closely related to the properties of the dendritic arbor itself, implying that the dendrites strongly influence and control synaptic transmission and plasticity (Sjöström et al., 2008).

How it is Measured or Detected

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Elaboration of dendritic processes is measured from electron and fluorescent micrographs. These processes are identified primarily by the presence of microtubule associated protein 2 (MAP-2) and the absence of components characteristic of axons and glia (e.g. small vesicles, myelin, glial filaments). These measurements can also be carried out by automated imaging systems in cells prepared for immunohistochemistry with specific antibodies that recognise MAP-2 (Harrill and Mundy, 2011).

Two-photon time-lapse images can also be used to visualise dendrites in GFP-transfected neurons, whereas Golgi Stain Kit is used to measure both dendrites and dendritic spines. A combination of Golgi-Cox and immunofluorescence using confocal microscopy has also been suggested in brain slices from rodents and non-human primates (Levine et al., 2013).

The morphological analysis of neurons, include the use of fluorescent markers, such as Dil (1,1'-dioctadecyl-3,3,3',3'-tetramethylindocarbocyanine perchlorate) that permits not only the visualisation of detailed dendritic arborizations and spines in cell culture and tissue sections but also permits the quantitative analysis of dendritic spines.

Fluorescent labelling for MARCM (mosaic analysis with a repressible cell marker) system can also be used but only in case of transparent larval body wall found in Drosophila.
Evidence Supporting Taxonomic Applicability

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*Drosophila* is one of the best-studied models that allow examining how diverse dendrite morphologies are formed during development (Grueber et al., 2002). The chick embryo (*Gallus domesticus*) is another important model in vertebrate developmental neurobiology where the dendritic arbor development has been extensively studied (Rubel and Fritzsch, 2002). Different methods have also been used to study dendritic arborization and spine formation in brain sections and cell cultures derived by rodents (Stansfield et al., 2012) and primates (Khazipov et al., 2001).

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Synaptogenesis, Decreased
Synaptogenesis, Decreased

How this Key Event works

**Level of Biological Organization**

**Biological state:** Synaptogenesis is a multi-step process that is crucial for brain development and involves the formation of synapses. It follows axonal migration, at which stage presynaptic and postsynaptic differentiation occurs (Garner et al., 2002). "Synaptic assembly" that refers to the gathering of the appropriate components and "synaptic formation" that is defined by the mechanisms involved in recruitment of molecules required for differentiation, stabilization and maturation of synapse, are the main phases that characterise synaptogenesis (Colón-Ramos, 2009). Elimination is a physiological step involved in synaptogenesis regarding the synapses that fail to get stabilised and mature.

The first step is the recognition of synaptic post synaptic part followed by the establishment of contact between an axon and a dendritic spine in which pre- and postsynaptic neurons play important role. The presynaptic differentiation occurs followed by excretion of neurotransmitters that bind to appropriate receptors located on the target spine. However, a postsynaptic neuron does not passively receive guidance from a presynaptic axon but are the same dendritic filopodia that gradually are transformed into spines that select and engage their presynaptic neurons. The transformation of dendritic filopodia into dendritic spines that involves the expression of the whole postsynaptic machinery such as postsynaptic density (PSD), receptor subunits, scaffolding proteins and actin cytoskeleton, is the first step to give nascent synapses. However, to become functional and mature these synapses need an important number of cell-cell interactions, including stimulation from glutamatergic synapses as well as the influence of neurotrophic factors (Munno and Syed, 2003).

However, all this is true for glutamatergic synapses because GABAergic synapses do not appear in dendritic spines, but rather form on dendritic shafts, nerve cell somata and axon initial segments. These inhibitory synapses besides their distinct location are also structurally different compared to excitatory synapses (reviewed in Gatto and Broadie, 2010).

**Biological compartments:** Synaptogenesis is spaciously and temporally strictly controlled process. It does not happen in a uniform way in all brain regions and there important differences between the times of appearance of the main two types of synapses (reviewed in Erecinska et al., 2004). For example, in rat hippocampus excitatory synapses are well established or fully mature within the two first postnatal weeks, whereas inhibitory synapses cannot be found prior to PND 18, after which it increases steadily to reach adult levels at PND 28. In addition, in rat neostriatal neurons the excitatory responses to both cortical and thalamic stimuli can be observed by PND 6, but the long-lasting hyperpolarization and late depolarization is never seen before PND 12.
Structural remodelling of synapses and formation of new synaptic contacts has been postulated as a possible mechanism underlying the late phase of long-term potentiation (LTP), a form of plasticity which is involved in learning and memory. LTP induction results in a sequence of morphological changes consisting of a transient remodelling of the postsynaptic membrane followed by a marked increase in the proportion of axon terminals contacting two or more dendritic spines. Three-dimensional reconstruction revealed that these spines arose from the same dendrite. As pharmacological blockade of LTP prevented these morphological changes, it is suggested that LTP is associated with the formation of new, mature and probably functional synapses contacting the same presynaptic terminal and thereby duplicating activated synapses (Erik et al., 2006).

In human, synaptogenesis does not happen at the same time in all brain regions, as the prefrontal cortex lags behind in terms of synapse formation compared to the auditory and visual cortices. In contrast, synaptogenesis appears to proceed concurrently in different brain areas for rhesus monkey.

**General role in biology:** The period of rapid synaptogenesis or the so-called brain growth spurt is considered one of the most important processes that take place during brain development (Garner et al., 2002). This process is crucial not only in neurodevelopment but also plays a vital role in synaptic plasticity, learning and memory and adaptation throughout life. Without this process no complex brain network can be established as synapse is the fundamental unit of connectivity and communication between neurons (Tau and Peterson, 2010). Cell adhesion represents the most direct way of coordinating synaptic connectivity in the brain. Recent evidence highlights the importance of a trans-synaptic interaction between postsynaptic neuroligins and presynaptic neurexins. These transmembrane molecules bind each other extracellularly to promote adhesion between dendrites and axons, facilitating synapse establishment (Dean and Dresbach, 2006). Furthermore, the number of excitatory versus inhibitory synapses created at single neuron dictates neuronal excitability and function (Schummers et al., 2002).

**How it is Measured or Detected**

Methods that have been previously reviewed and approved by a recognized authority should be included in the Overview section above. All other methods, including those well established in the published literature, should be described here. Consider the following criteria when describing each method: 1. Is the assay fit for purpose? 2. Is the assay directly or indirectly (i.e. a surrogate) related to a key event relevant to the final adverse effect in question? 3. Is the assay repeatable? 4. Is the assay reproducible?

There is no OECD advised method for measuring synaptogenesis.

Anatomical methods can be used to structurally estimate the number of excitatory or inhibitory synapses. Immunostaining can be employed with specific antibodies that recognize vesicular glutamate transporters (VGLUTs) and the postsynaptic density protein-95 kDa (PSD-95) that are characteristic of excitatory synapses, while inhibitory synapses are identified by the presence of the vesicular GABA (VGAT) and vesicular inhibitory amino acid (VIAAT) transporters and the postsynaptic adaptor protein gephyrin (Gatto and Broadie, 2010). There are commercial available synaptogenesis assay kits that rely on the immunostaining of cells with MAP-2, PSD-95 and synaptophysin. Some other presynaptic (Bassoon) and postsynaptic (ProSAP1/Shank2) markers have been suggested and showed to correlate well with the ultrastructural studies in cultured hippocampus primary cells (Grabrucker et al., 2005). Electron microscopy can also be applied to assess the prevalence of excitatory and inhibitory synapses amongst convergent contacts (Megias et al., 2001). Recently, a high content image analysis based on RNAi screening protocols has been suggested as a useful tool to create imaging algorithm for use in both in vitro and in vivo synaptic punctae analysis (Nieland et al., 2014).

**Evidence Supporting Taxonomic Applicability**

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The mechanisms governing synapse formation is considered conserved among both vertebrates and invertebrates (Munno and Syed, 2003). Invertebrates have served as simple animal models to study synapse formation. Indeed, Colón-Ramos (2009) has recently reviewed the early developmental events that take place in the process of synaptogenesis pointing out the importance of this process in neural network formation and function. The experimental evaluation of synaptogenesis has been performed using invertebrates and in particular C. elegans and Drosophila as well as vertebrates (Colón-Ramos, 2009).

This vulnerable period of synaptogenesis appears to happen in different developmental stages across species. For example, in rodents primarily synaptogenesis occurs during the first two weeks after birth (Bai et al., 2013). For rhesus monkeys, this period ranges from approximately 115-day gestation up to PND 60 (Bai et al., 2013). In humans, it starts from the third trimester of pregnancy and continues 2-3 years following birth (Bai et al., 2013).

References


Biological state: There are striking differences in neuronal network formation and function among the developing and mature brain. The developing brain shows a slow maturation and a transient passage from spontaneous, long-duration action potentials to synaptically-triggered, short-duration action potentials.

Furthermore, at this precise developmental stage the neuronal network is characterised by "hyperexcitability", which is related to the increased number of local circuit recurrent excitatory synapses and the lack of γ-amino-butyric acid A (GABAA)-mediated inhibitory function that appears much later. This "hyperexcitability" disappears with maturation when pairing of the pre- and postsynaptic partners occurs and synapses are formed generating population of postsynaptic potentials and population of spikes followed by developmental GABA switch. Glutamatergic neurotransmission is dominant at early stages of development and NMDA receptor-mediated synaptic currents are far more times longer than those in maturation, allowing more calcium to enter the neurons. The processes that are involved in increased calcium influx and the subsequent intracellular events seem to play a critical role in establishment of wiring of neural circuits and strengthening of synaptic connections during development (reviewed in Erecinska et al., 2004). Neurons that do not receive glutaminergic stimulation are undergoing developmental apoptosis.

During the neonatal period, the brain is subject to profound alterations in neuronal circuitry due to high levels of synaptogenesis and gliogenesis. For example, in neuroendocrine regions such as the preoptic area-anterior hypothalamus (POA-AH), the site of gonadotropin-releasing hormone (GnRH) system is developmentally regulated by glutamatergic neurons. The changes in the expression of the N-methyl-D-aspartate (NMDA) receptor subunits NR1 and NR2B system begin early in postnatal development, before the onset of puberty, thereby playing a role in establishing the appropriate environment for the subsequent maturation of GnRH neurons (Adams et al., 1999).

Biological compartments: Neural network formation and function happen in all brain regions but it appears to onset at different time points of development (reviewed in Erecinska et al., 2004). Glutamatergic neurotransmission in hippocampus is poorly developed at birth. Initially, NMDA receptors play important role but the vast majority of these premature glutamatergic synapses are “silent” possibly due to delayed development of hippocampal AMPA receptors. In contrast, in the cerebral cortex the maturation of excitatory glutamatergic neurotransmission happens much earlier. The “silent” synapses disappear by PND 7-8 in both brain regions mentioned above.

There is strong evidence suggesting that NMDA receptor subunit composition controls synaptogenesis and synapse stabilization (Gambrill and Barria, 2011). It is established fat that during early postnatal development in the rat hippocampus, synaptogenesis occurs in parallel with a developmental switch in the subunit composition of NMDA receptors from NR2B to NR2A. It is suggested that early expression of NR2A in organotypic hippocampal slices reduces the number of synapses and the volume and dynamics of spines. In contrast, overexpression of NR2B does not affect the normal number and growth of synapses. However, it does increase spine motility, adding and retracting spines at a higher rate. The C terminus of NR2B, and specifically its ability to bind CaMKII, is sufficient to allow proper synapse formation and maturation.
Conversely, the C terminus of NR2A was sufficient to stop the development of synapse number and spine growth. These results indicate that the ratio of synaptic NR2B over NR2A controls spine motility and synaptogenesis, and suggest a structural role for the intracellular C terminus of NR2 in recruiting the signalling and scaffolding molecules necessary for proper synaptogenesis. Interestingly, it was found that genetic deletion of NR3A accelerates glutamatergic synaptic transmission, as measured by AMPAR-mediated postsynaptic currents recorded in hippocampal CA1. Consistent, the deletion of NR3A accelerates the expression of the glutamate receptor subunits NR1, NR2A, and GluR1 sugesting that glutamatergic synapse maturation is critically dependent upon activation of NMDA-type glutamate receptors (Henson et al., 2012).

**General role in biology:** The development of neuronal networks can be distinguished into two phases: an early ‘establishment’ phase of neuronal connections, where activity-dependent and independent mechanisms could operate, and a later ‘maintenance’ phase, which appears to be controlled by neuronal activity (Yuste and Sur, 1999). These neuronal networks facilitate information flow that is necessary to produce complex behaviors, including learning and memory (Mayford et al., 2012).

**How it is Measured or Detected**

*Methods that have been previously reviewed and approved by a recognized authority should be included in the Overview section above. All other methods, including those well established in the published literature, should be described here. Consider the following criteria when describing each method: 1. Is the assay fit for purpose? 2. Is the assay directly or indirectly (i.e. a surrogate) related to a key event relevant to the final adverse effect in question? 3. Is the assay repeatable? 4. Is the assay reproducible?*

**In vivo:** The recording of brain activity by using electroencephalography (EEG), electrocorticography (ECoG) and local field potentials (LFP) assists towards the collection of signals generated by multiple neuronal cell networks. Advances in computer technology have allowed quantification of the EEG and expansion of quantitative EEG (qEEG) analysis providing a sensitive tool for time-course studies of different compounds acting on neuronal networks' function (Binienda et al., 2011). The number of excitatory or inhibitory synapses can be functionally studied at an electrophysiological level by examining the contribution of glutamatergic and GABAergic synaptic inputs. The number of them can be determined by variably clamping the membrane potential and recording excitatory and inhibitory postsynaptic currents (EPSCs or IPSCs) (Liu, 2004).

**In vitro:** Microelectrode array (MEA) recordings are also used to measure electrical activity in cultured neurons (Keefter et al., 2001, Gramowski et al., 2000; Gopal, 2003; Johnstone et al., 2010). MEAs can be applied in high throughput platforms to facilitate screening of numerous chemical compounds (McConnell et al., 2012). Using selective agonists and antagonists of different classes of receptors their response can be evaluated in a quantitative manner (Novellino et al., 2011; Hogberg et al., 2011).

**Evidence Supporting Taxonomic Applicability**

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In vitro studies in brain slices applying electrophysiological techniques showed significant variability among species related to synaptic latency, duration, amplitude and efficacy in spike initiation (reviewed in Erecinska et al., 2004). These brain tissues had derived from immature rats, rabbits and kittens (Erecinska et al., 2004).

**References**


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Presynaptic release of glutamate, Reduced
Presynaptic release of glutamate, Reduced

How this Key Event works
**Level of Biological Organization**

**Molecular**

**Biological state:** Glutamate is an amino acid with neurotransmitter function that is stored in presynaptic vesicles by the action of vesicular glutamate transporters (VGLUTs) and under physiological conditions is found at a concentration of 100 mmol/L per vesicle. Different mechanisms are involved in the release of glutamate (reviewed in Meldrum, 2000). Glutamate is mainly released from the vesicles in a Ca2+-dependent mechanism that involves N- and P/Q-type voltage-dependent Ca2+ channels, closely linked to vesicle docking sites. However, glutamate can also be released by reverse operation during reduction of Na+ and K+ gradient across the membrane like for example during cerebral ischemia. Interestingly, the synaptic release of glutamate is controlled by a wide range of presynaptic receptors that are not only glutamatergic like Group II and Group III of glutamate metabotropic receptors but also cholinergic such as nicotinic and muscarinic, adenosine (A1), kappa opioid, γ-aminobutyric acid (GABA)B, cholecystokinin and neuropeptide Y (Y2) receptors.

The synaptic effects of glutamate are rapidly terminated by action of glutamate transporters (excitatory amino acid transporters [EAATs]) located on the plasma membrane of astrocytes and neurons. Therefore, pre-synaptically released glutamate is mostly re-uptaken by astrocytes but also transported from the synaptic cleft into the presynaptic terminals (Rothstein et al., 1994; Blanke and VanDongen, 2009).

Following its release, glutamate exerts its effects via inotropic and metabotropic receptors. Although glutamate is available for binding to receptors for a short time, NMDA receptors show high affinity for this specific neurotransmitter that causes their activation compared to other receptors.

**Biological compartments:** Glutamate is the most abundant amino acid in the diet, consequently is found at higher levels in plasma compared to cerebrospinal fluid. The blood brain barrier prevents the entry of glutamate, meaning that the glutamate present in CNS is derived from de novo synthesis of this neurotransmitter relying on the recycling of the main resources. Glutamine and α-ketoglutarate are the major precursors of glutamate. Glutamine is converted via phosphate-activated glutaminase to glutamate and ammonia, whereas α-ketoglutarate is transaminated into glutamate (Platt, 2007). In glial cells, the glutamate is metabolised via glutamine synthase into glutamine or metabolised into α-ketoglutarate. These products are actively transported out of the glial cells and back into the pre-synaptic terminals for subsequent re-synthesis and storage of glutamate.

Five transporters of glutamate have been identified in the CNS. Two are expressed predominantly in glia and three in neurons (reviewed in Meldrum, 2000). The presence of glutamate has also been demonstrated in other tissues and organs as glutamate receptors have been found to be expressed in pancreatic β-cells, osteoblasts and osteoclasts of bones (Nedergaard et al., 2002).

**General role in biology:** In mature nervous system, glutamate is known to play important role in synaptic plasticity. Similarly important is this neurotransmitter during development where it regulates neurogenesis, neurite outgrowth, synaptogenesis and apoptosis (reviewed in Mattson, 1996; Meldrum, 2000; Mattson, 2008).

The proper functioning of the central nervous system relays on the physiological homeostasis between glutamate and GABA, creating the opposite excitatory/inhibitory forces in the brain. Together, these two neurotransmitters constitute more than 90% of all neurotransmission. If this homeostasis is disturbed it could lead to anxiety disorders (Wieronska et al., 2015).

**How it is Measured or Detected**

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No OECD methods are available to measure glutamate release.

There are radioactive assays like [3H]glutamate release assay and spectrophotometric commercially available kits to measure glutamate in cell culture medium (release) or intracellular (cell lysate) using LC-MS. Furthermore, neurotransmitters including glutamate can be measured by HPLC with fluorescence detector.

**Evidence Supporting Taxonomic Applicability**

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Whereas glutamatergic transmission in vertebrates is excitatory, mediated by glutamate-gated cation channels, glutamate serves as both an excitatory and an inhibitory transmitter in invertebrates (Cleland, 1996).

**References**


**Cell injury/ death, N/ A**

**Short name: Cell injury/ death, N/ A**

**How this Key Event works**
The pathogenesis of drug- or toxin-induced cell injury usually involves toxic metabolites that either elicit an immune response or directly affect the biochemistry of the cell. Intracellular stress can lead to apoptotic or necrotic cell death, depending on the extent of mitochondrial involvement (ATP and ROS production, cytochrome c and apoptosis inducing factor (AIF) release, opening of permeability transition pore (PTP) and the balance of factors that activate and inhibit the Bcl2 family of proteins and the caspases. Chemicals and their metabolites can trigger oxidative stress causing depletion of reduced glutathione with consequent proteins, lipids, and DNA oxidation. These chemical consequences can directly affect organelles such as mitochondria, cytoskeleton, endoplasmic reticulum, microtubules, or nucleus or indirectly influence these organelles through activation or inhibition of signalling kinases, transcription factors, and gene and protein expression. The outcome may be either triggering of the necrotic or apoptotic process or sensitization to the lethal action of cytokines of the immune system.

Two types of cell death can be distinguished by morphological features, although it is likely that these are two ends of a spectrum with possible intermediate forms. Apoptosis involves shrinkage, nuclear disassembly, and fragmentation of the cell into discrete bodies with intact plasma membranes. These are rapidly phagocytosed by neighbouring cells. An important feature of apoptosis is the requirement for adenosine triphosphate (ATP) to initiate the execution phase. In contrast, necrotic cell death is characterized by cell swelling and lysis. This is usually a consequence of profound loss of mitochondrial function and resultant ATP depletion, leading to loss of ion homeostasis, including volume regulation, and increased Ca2+. The latter activates a number of nonspecific hydrolases (i.e., proteases, nucleases, and phospholipases) as well as calcium dependent kinesis. Activation of calpain I, the Ca2+-dependent cysteine protease cleaves the death-promoting Bcl-2 family members Bid and Bax which translocate to mitochondrial membranes, resulting in release of truncated apoptosis-inducing factor (tAIF), cytochrome c (cyt c) and endonuclease G (endoG) in the case of Bid and cyt c in the case of Bax. tAIF translocates to cell nuclei, and together with cyclophilin A and phosphorylated histone H2AX (γH2AX) is responsible for DNA cleavage, a feature of programmed necrosis. Activated calpain I has also been shown to cleave the plasma membrane Na+-Ca2+ exchanger, which leads to buildup of intracellular Ca2+, so this is a source of additional increased intracellular Ca2+. Cytochrome c (cyt c) in cellular apoptosis is a component of the apoptosome.

DNA damage activates nuclear poly(ADP-ribose) polymerase-1(PARP-1), a DNA repair enzyme. PARP-1 forms poly(ADP-ribose) polymers, to repair DNA, but when DNA damage is extensive, PAR accumulates, exits neuronal nuclei and travels to mitochondrial membranes, where it, like calpain I, is involved in AIF release from mitochondria. A fundamental distinction between necrosis and apoptosis is the loss of plasma membrane integrity; this is integral to the former but not the latter. As a consequence, lytic release of cellular constituents promotes a local inflammatory reaction, whereas the rapid removal of apoptotic bodies minimizes such a reaction. The distinction between the two modes of death is easily accomplished in vitro but not in vivo. Thus, although claims that certain drugs induce apoptosis have been made, these are relatively unconvincing. DNA fragmentation can occur in necrosis, leading to positive TUNEL staining. Conversely, when apoptosis is massive, it can exceed the capacity for rapid phagocytosis, resulting in the eventual appearance of secondary necrosis.

Two alternative pathways - either extrinsic (receptor-mediated) or intrinsic (mitochondria-mediated) - lead to apoptotic cell death. The initiation of cell death begins either at the plasma membrane with the binding of TNF or FasL to their cognate receptors or within the cell. The latter is due to the occurrence of intracellular stress in the form of biochemical events such as oxidative stress, redox changes, covalent binding, lipid peroxidation, and consequent functional effects on mitochondria, endoplasmic reticulum, microtubules, cytoskeleton, or DNA. The intrinsic mitochondrial pathway involves the initiator, caspase-9, which, when activated, forms an “apoptosome” in the cytosol, together with cytochrome c, which translocates from mitochondria, Apaf-1 and dATP. The apoptosome activates caspase-3, the central effector caspase, which in turn activates downstream factors that are responsible for the apoptotic death of a cell (review by Fujikawa, 2015). Intracellular stress either directly affects mitochondria or can lead to effects on other organelles, which then send signals to the mitochondria to recruit participation in the death process.
Constitutively expressed nitric oxide synthase (nNOS) is a Ca2+-dependent cytosolic enzyme that forms nitric oxide (NO) from L-arginine, and NO reacts with the free radical such as superoxide (O2−) to form the very toxic free radical peroxynitrite (ONOO−). Free radicals such as ONOO−, O2− and hydroxyl radical (OH−) damage cellular membranes and intracellular proteins, enzymes and DNA (review by Fujikawa, 2015).

**How it is Measured or Detected**

Necrosis: LDH is a soluble cytoplasmic enzyme that is present in almost all cells and is released into extracellular space when the plasma membrane is damaged. To detect the leakage of LDH into cell culture medium, a tetrazolium salt is used in this assay. In the first step, LDH produces reduced nicotinamide adenine dinucleotide (NADH) when it catalyzes the oxidation of lactate to pyruvate. In the second step, a tetrazolium salt is converted to a colored formazan product using newly synthesized NADH in the presence of an electron acceptor. The amount of formazan product can be colorimetrically quantified by standard spectroscopy. Because of the linearity of the assay, it can be used to enumerate the percentage of necrotic cells in a sample.

The MTT assay is a colorimetric assay for assessing cell viability. NAD(P)H-dependent cellular oxidoreductase enzymes may reflect the number of viable cells present. These enzymes are capable of reducing the tetrazolium dye MTT 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide to its insoluble formazan, which has a purple color. Other closely related tetrazolium dyes including XTT, MTS and the WSTs. Tetrazolium dye assays can also be used to measure cytotoxicity (loss of viable cells) or cytostatic activity (shift from proliferation to quiescence) of potential medicinal agents and toxic materials. MTT assays are usually done in the dark since the MTT reagent is sensitive to light (Berridge et al., 2005)

Propidium iodide (PI) is an intercalating agent and a fluorescent molecule used to stain necrotic cells. It is cell membrane impermeant so it stains only those cells that cell membrane is destroyed. When PI is bound to nucleic acids, the fluorescence excitation maximum is 535 nm and the emission maximum is 617 nm. (Moore et al., 1998).

Apoptosis: TUNEL is a common method for detecting DNA fragmentation that results from apoptotic signaling cascades. The assay relies on the presence of nicks in the DNA which can be identified by terminal deoxynucleotidyl transferase or TdT, an enzyme that will catalyze the addition of dUTPs that are secondarily labeled with a marker. It may also label cells that have suffered severe DNA damage.

Caspases activity assays measured by fluorescence. During apoptosis, mainly caspase-3 and -7 cleave PARP to yield an 85 kDa and a 25 kDa fragment. PARP cleavage is considered to be one of the classical characteristics of apoptosis. Antibodies to the 85 kDa fragment of cleaved PARP or to caspase-3 both serve as markers for apoptotic cells that can be monitored using immunofluorescence (Li et al., 2004)

Hoechst 33342 staining: Hoechst dyes are cell-permeable and bind to DNA in live or fixed cells. Therefore, these stains are often called supravital, which means that cells survive a treatment with these compounds. The stained, condensed or fragmented DNA is a marker of apoptosis (Kubbies et al., 1983).

**Evidence Supporting Taxonomic Applicability**

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<td>human</td>
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<td>rodents</td>
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<td>human and other cells in culture</td>
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</table>
Cell death is an universal event occurring in cells of any species.

References

1. ↑ Malhi, H. et al., (2010), Hepatocyte death: a clear and present danger. Physiol Rev. 90, 1165-1194


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Adverse Outcome

**Learning and memory, Impairment**

**Short name:** Learning and memory, Impairment

**How this Key Event works**

**Level of Biological Organization**

Learning can be defined as the process by which new information is acquired to establish knowledge by systematic study or by trial and error (Ono, 2009). Two types of learning are known in neurobehavioral studies: a) associative learning and b) non-associative learning.

The associative learning is the learning by doing associations between different events. In associative learning, a subject learns the relationship among two different stimuli or between the stimulus and the subject's behavior. Classical conditioning, operant conditioning and category learning are some examples of the associative learning.

On the other hand, non-associative learning can be defined as an alteration in the behavioral response that occurs over time in response to a single type of stimulus. Habituation and sensitization are some examples of non-associative learning.

The memory to be formed requires acquisition, retention and retrieval of information in the brain, which is characterised by the nonconscious recall of information (Ono, 2009). Memory is considered very important as it allows the subjects to access the past, to form experience and consequently to acquire skills for surviving purposes. There are three main categories of memory, including sensory memory, short-term or working memory (up to a few hours) and long-term memory (up to several days or even much longer). At the cellular level the storage of long-term memory is associated with increased gene expression and protein
synthesis as well as formation of novel synaptic connections (Lynch et al., 2014).

Amnesia is defined as the impairment or loss of memory. Depending on the cause amnesia can be characterised as functional, organic amnesia or infantile amnesia. Dementia is the intellectual impairment observed mainly in elderly people due to the progress of a neudegenerative disease, whereas in younger people this type of impairment is known as presenile dementia.

It is appropriate to state that while much emphasis has been given on the key role of the hippocampus in memory, it would probably be simplistic to attribute memory deficits solely to hippocampal damage (Barker and Warburton, 2011). There is substantial evidence that fundamental memory functions are not mediated by hippocampus alone but require a network that includes, in addition to the hippocampus, anterior thalamic nuclei, mammillary bodies and cortex (Aggleton and Brown, 1999; Mitchell et al., 2002). Each of these brain structures can be potentially damaged leading to more or less severe impairment of learning and memory.

**How it is Measured or Detected**

**In humans:** Initially, neuropsychological tests have been used to identify mainly risks from occupational exposure to chemicals. Later neurosensory assessment has been incorporated, whereas recently there has been a shift from occupational exposure towards identification of altered neurobehaviors in vulnerable populations such as children (Rohlman et al., 2008). Intelligence tests, perceptual motor tests, planning tests, and logical, spatial, short term, long term, and working memory tasks can be used in neurobehavioral studies to assess learning and memory in humans.

**In laboratory animals:** Cognitive function including learning and memory is an important endpoint required by the OECD Developmental Neurotoxicity study TG 426 and OECD Neurotoxicity Study in Rodents TG 424. The methods applied to assess learning and memory have been recently reviewed in the OECD Series on testing and assessment number 20, Guidance document for Neurotoxicity Testing (2004). This document is considered an essential supplement to a substantial number of already existing OECD Test Guidelines relevant for neurotoxicity testing.

**Evidence Supporting Taxonomic Applicability**

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Learning and memory have been studied in invertebrates such as gastropod mollusks and drosophila and vertebrates such as rodents and primates. Recently, larval zebrafish has also been suggested as model for the study of learning and memory (Roberts et al., 2013).

**Regulatory Examples Using This Adverse Outcome**

Impairment of learning and memory is considered a chemical induced adverse outcome that is used for risk assessment and management purposes. Neurotoxicity testing guidelines (OECD TG 424 and 426) are implemented in a number of occasions where the neurotoxic properties of a compound have to be assessed in order to comply with relevant EU regulations. These regulations are as follows: REACH regulation (EC, No 1907/2006), Plant protection products regulation (EC, No 1107/2009), Biocidal products regulation (EC, No 528/2012), Test methods regulation (EC, No 440/2008), Classification, labelling and packaging of substances and mixtures (EC, No 1272/2008) and Maximum residue levels of pesticides in or on food and feed of plant and animal origin regulation (EC, No 396/2005).

**References**


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### Scientific evidence supporting the linkages in the AOP

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<td>Synaptogenesis, Decreased</td>
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### NMDARs, Binding of antagonist Directly Leads to NMDARs, Inhibition

How Does This Key Event Relationship Work
The binding of antagonist to the NMDA receptor causes inhibition of the NMDAR function.

**Weight of Evidence**

**Biological Plausibility**

There is structural mechanistic understanding supporting the relationship between MIE (NMDARs, binding of antagonists) and KE1 (NMDARs, inhibition). Crystal structure studies are used to study the binding of antagonists/agonists to NMDA receptors. In case of NMDAR antagonists, the binding to the receptor causes LBD opening and promotion of channel closure leading to reduced Ca\(^{2+}\) influx (Blanke and VanDongen, 2009). This lack of measurable ion flux is applied as an indication of NMDAR inhibition.

**Empirical Support for Linkage**

*Include consideration of temporal concordance here*

In slices of cerebellum derived from postnatal days 6-30 (PND 6-30) Sprague Dawley rats, 10 \(\mu M\) MK-801 completely blocked evoke NMDA excitatory postsynaptic currents (EPSCs) as it has been demonstrated by patch clamp technique (Rumbaugh and Vicini, 1999). The same technique has been employed in cortical slices from C57BL/6 mice of both genders and different age groups (P8-12, P21-28 or P45-90), showing that 1 \(\mu M\) APV and 50 nM NVP-AAM077 antagonise a similar amount of NMDA receptor current independently of the age (de Marchena et al., 2008).

Pb\(^{2+}\) has potent inhibitory effects on the NMDA receptor (Alkondon et al., 1990; Guilarte and Miceli, 1992; Guilarte, 1997; Gavazzo et al., 2001). In rat hippocampal neurons, Pb\(^{2+}\) (2.5-50 \(\mu M\)) inhibits NMDA-induced whole-cell and single-channel currents in a concentration-dependent manner, suggesting that Pb\(^{2+}\) can decrease the frequency of NMDA-induced channel activation (Alkondon et al., 1990). In the same study, they have examined the effect of Pb\(^{2+}\) on the binding of \([3H]\)MK-801 to the rat brain hippocampal membranes and showed that Pb\(^{2+}\) inhibits the binding of \([3H]\)MK-801 in a concentration dependent manner with an IC\(_{50}\) value close to 700 \(\mu M\) (Alkondon et al., 1990). These inhibitory effects of Pb\(^{2+}\) on NMDA receptors activation appear to be age and brain region specific (Guilarte, 1997; Guilarte and Miceli, 1992). The Pb\(^{2+}\) IC\(_{50}\) is significantly lower in cortical membranes prepared from neonatal than from adult rats, whereas the hippocampus is more sensitive than the cerebral cortex since the Pb\(^{2+}\) IC\(_{50}\) is significantly lower in the hippocampus (Guilarte and Miceli, 1992). The number of \([3H]\)MK-801 binding sites associated with the high and low affinity sites of Pb\(^{2+}\) inhibition in the hippocampus of rats is increased as a function of age, peaking at PND 28 and 21 (Guilarte, 1997). High and low affinity Pb\((2+)\)-sensitive \([3H]\)MK-801 binding sites have also been measured in the cerebral cortex during early development, but that has not be possible to be evaluated after PND 14.

The developing brain is more sensitive than the adult brain to Pb\(^{2+}\)-induced effects mediated through the NMDA receptor. Moreover, the hippocampus appears to be particularly vulnerable as in this brain structure NMDA receptors undergo subunit specific changes during developmental Pb\(^{2+}\) exposure (Guilarte and McGlothan, 1998). Exposure to Pb\(^{2+}\) during synaptogenesis causes decreased expression of hippocampal NR2A-subunit of NMDARs at synapses and increased targeting of NR2B-NMDARs to dendritic spines (without increased NR2B-NMDARs expression) (Nihei and Guilarte, 1999; Neal et al., 2011; Zhang et al., 2002).

**Uncertainties or Inconsistencies**

Pb\(^{2+}\) has been found to produce either potentiation or inhibition depending on: a) the subunit composition of NMDA receptors, b) endogenous glutamate concentration and c) Pb\(^{2+}\) dosage. In case that the NMDA receptors are saturated by agonist, Pb\(^{2+}\) at low concentrations (<1 \(\mu M\)) acts as a positive modulator of agonist action at NR1b-2AC and NR1a-2AB subunit complexes, whereas at higher concentrations, Pb\(^{2+}\) it behaved as a potent inhibitor of all recombinant NMDA receptors tested and was least potent at NR1b-2AC (Omelchenko et al., 1996; 1997), meaning that Pb\(^{2+}\) is not always acting as NMDAR inhibitor but it can also behave as NMDAR activator under certain conditions.
Quantitative Understanding of the Linkage

Is it known how much change in the first event is needed to impact the second? Are there known modulators of the response-response relationships? Are there models or extrapolation approaches that help describe those relationships?

To predict how potent an antagonist can be, it is usually measured the half maximal inhibition concentrations (IC50) for the inhibition of glutamate/glycine and the half maximal effective concentration (EC50) induced currents in NMDA receptors from brain slices and cells or in recombinantly expressed receptors. Traynelis et al. 2010 summarised the IC50 values for competitive, noncompetitive and uncompetitive antagonists in different subunits of NMDA receptors. The inhibitory effect (efficacy) of antagonists on NMDA receptors has been found to be dependent on:

- the type of subunits that form the NMDA receptors depending on the developmental stage -the chemical structure of the antagonists
- how tightly an antagonists binds to the receptor (affinity)

\[ \text{Pb2}^+ \]: Although the NR2 subunits have different Zn2+ binding sites i.e. the NR2A-NMDAR binds Zn2+ at a high-affinity site (nM affinity) while the NR2B-NMDAR binds Zn2+ with lower affinity (µM range), the Pb2+ IC50 for wild type NR2A-NMDARs was reported to be 1.3 µM, while the Pb2+ IC50 of wild type NR2B-NMDARs was 1.2 µM (Gavazzo et al., 2008). Similar finding were published by Lasley and Gilbert (1999) using cortical neurons from adult rats. The IC50 for Pb2+ ranged from 1.52 to 4.86 µM, with the rank of Pb2+ potency in inhibition of NMDA receptor subunits to be NR1b-2A>NR1b-2C>NR1b-2D>NR1b-2AC after experiments that have been conducted in Xenopus oocytes injected with cRNAs for different combinations of NMDA receptor subunits (Omelchenko et al., 1997).

Ketamine: Ketamine IC50 values were not significantly different between recombinant human and rat NR1/NR2A and NR1/NR2C NMDA receptor subunits expressed in Xenopus oocytes (Hedegaard et al., 2012). However, ketamine IC50 values were 1.6- and 1.7-fold lower in rat NR1/NR2B and NR1/NR2D, respectively, compared to the corresponding human subtypes. Memantine IC50 values were not significantly different between human and rat NR1/NR2A, NR1/NR2B, and NR1/NR2D, but were 1.3-fold higher in rat NR1/NR2C compared to human NR1/NR2C. The rank order of potencies at both human and rat subtypes was the same for both antagonists, ketamine and memantine, (NR1/NR2A > NR1/NR2B > NR1/NR2D > NR1/NR2C), and similar findings have been reported as well for recombinant rat NMDA receptors (Dravid et al., 2007). Recently, NMDAR occupancy time course following ketamine administration to rats, nonhuman primates and humans has been carried out to allow direct interspecies comparisons of specific ketamine-mediated pharmacodynamics through normalization (Shaffer et al., 2014). Total plasma concentration -time profiles of ketamine has been generated from rats and nonhuman primates following a single, memory-impairing dose of ketamine, whereas neuropharmacokinetics have been determined in rats and [(3)H]MK-801-displacement studies in rats (Shaffer et al., 2014).

Evidence Supporting Taxonomic Applicability

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The biophysical properties of rat and human receptors have been mostly assessed through recombinant studies, whereas the pharmacological properties of rat and human NMDA receptors have not been fully explored and compared yet (Hedegaard et al., 2012). Mean channel open times for human NMDA receptor subtypes in recombinant protein studies are similar to those of the corresponding rat NMDA receptor subtypes. However, mean single-channel conductances for human NMDA receptor subtypes appear lower than those of the corresponding rat NMDA receptor subtypes. Regarding pharmacological properties of the receptors, the differences were less than 2-fold and were not observed at the same subtypes for all the
antagonists testing, suggesting that the molecular pharmacology of NMDA receptor is conserved between human and rat, although some inter-species differences are seen in IC50 values using two-electrode voltage-clamp recordings (Hedegaard et al., 2012).

References


How Does This Key Event Relationship Work

The NMDA receptor is distinct in two ways: firstly, it is both ligand-gated and voltage-dependent and secondly, it requires co-activation by two ligands: glutamate and either D-serine or glycine.

NMDA receptor activation allows the influx of Ca2+ only when the receptor is occupied by L-glutamate or other agonists (and removal of Mg++ block) resulting in the postsynaptic membrane depolarization. In contrast, binding of antagonist to NMDA receptor decreases or eliminates Ca2+ influx and consequently dramatically decreases intracellular influx of Ca2+ levels (reviewed in Higley and Sabatini, 2012).

Weight of Evidence

Biological Plausibility

The relationship between KE1 (NMDARs, Inhibition) and KE2 (Calcium influx, Decreased) is plausible as the function evaluation of NMDA receptors is commonly carried out by measurement of intracellular influx of Ca2+ upon NMDA receptor stimulation by agonist. Calcium imaging techniques have been extensively utilized to investigate the relationship between these two KEs. Almost 15% of the current through NMDA receptors is mediated by Ca2+ under physiological conditions (Higley and Sabatini, 2012).

It has been shown that less than five and, occasionally only a single NMDA receptor opens under physiological conditions, causing a total Ca2+ influx of about 6000 ions into a dendritic spine head reaching a concentration of 10 µM (Higley and Sabatini, 2012). However, the majority of the ions are rapidly eliminated by binding Ca2+ proteins, reaching 1 µM of free Ca2+ concentration (Higley and Sabatini, 2012).

In rat primary forebrain cultures the intracellular Ca2+ increases after activation of the NMDA receptor and this increase is blocked when the cells are cultured under Ca2+ free conditions, demonstrating that the NMDA-evoked increase in intracellular Ca2+ derives from extracellular and not intracellular sources (Liu et al., 2013).

Neurons in brain slices from wild-type (GluRe2+/+) mice showed increase of intracellular Ca2+ in the presence of 100 µM NMDA that was completely inhibited after exposure to 100 mM APV. In contrast, the NMDA-mediated increase in Ca2+ was absent in brain slices from GluRe2−/− mice that do not possess any functional NMDA receptors in the developing neocortex (Okada et al., 2004).

Empirical Support for Linkage

Include consideration of temporal concordance here

Pb2+: There are a few studies examining the effect of Pb2+ exposure on the changes in intracellular Ca2+. Incubation of rat synaptosomes with Pb2+ stimulates the activity of calmodulin reaching the higher effect at 30 µM, whereas higher concentrations of Pb2+ causes inhibition (Sandhir and Gill, 1994). Pb2+ exposure increases the activity of calmodulin by 45% in animal models. The IC50 values for inhibition of Ca2+ ATPase by Pb2+ has been found to be 13.34 and 16.69 µM in calmodulin-rich and calmodulin-
depleted synaptic plasma membranes, respectively. Exposure of rats to Pb2+ has also inhibitory effect on Ca2+ ATPase activity, causing increase in intrasynaptosomal Ca2+ (Sandhir and Gill, 1994). In embryonic rat hippocampal neurons, exposure to 100 nM Pb2+ for periods from 1 hour to 2 days showes decrease of Ca2+ (Ferguson et al., 2000).

There is evidence that Pb2+ exposure affects Ca2+ homeostasis causing alterations in the phosphorylation state of different kinases. For example, Pb2+ has been shown to interfere with MAPK signaling as it increases the phosphorylation of both ERK1/2 and p38(MAPK) (Cordova et al., 2004). However, the findings regarding calcium/calmodulin kinase II (CamKII) activity are not clear (Toscano et al., 2005). On one hand, Pb2+ has been found to cause reduction of CREB phosphorylation in the hippocampus of rats exposed during brain development (Toscano et al., 2003; Toscano et al., 2002). One the other hand, the levels of phosphorylation of CamKII have not been explored but only the mRNA expression levels have been studied in rat pups on PND 25 that received Pb2+ (180 and 375-ppm lead acetate in food for 30 days) and reached blood Pb2+ levels 5.8 to 10.3 μg/dl on PND 55 (Schneider et al., 2012). More specifically, CamKIIα gene expression has been found to be very sensitive to Pb2+ exposure in the frontal cortex but not in the hippocampus, whereas CamKIIβ gene expression in both brain structures remained unchanged (Schneider et al., 2012).

Acute Pb2+ (10μM) exposure impairs LTP (125.8% reduction of baseline) in CA1 region of hippocampus derived from Sprague-Dawley rats (15-18 PND) as it has been recorded by whole cell patch-clamp technique (Li et al., 2006). In the same study, through calcium imaging, it has been shown in the 10mM caffeine-perfused cultured hippocampal neurons that 10μM Pb2+ reduces intracellular Fluo-4 fluorescence ratio to 0.44 (Li et al., 2006).

Pb2+ chronically or acutely applied, significantly reduces LTP in CA1 region of hippocampus from Wistar or Sprague-Dawley rats (30 and 60 PND) (Carpenter et al., 2002). These animals were exposed to Pb2+ via the mother’s drinking water either through gestation and lactation (upto day 21) (perinatal), only by lactation through the mother’s drinking water and then in the pup’s drinking water (post) or from gestation (pre and post). The concentrations of Pb2+ used in the drinking water were 0.1 and 0.2%. In CA1, LTP has been reduced at both ages and Pb2+ concentrations or duration of exposure. In CA3, there have been no differences with time of exposure, but there was a dramatic difference in response as the age of aniams increased. At 30 days LTP was significantly reduced, but at 60 days LTP was increased by about 30% (Carpenter et al., 2002). In the same brain structure and area (CA3) the effects of Pb2+ on LTP have been different in 30 PND and 60 PND rats after either acute perfusion of Pb2+ or from slices derived from rats after chronic developmental exposure to Pb2+, as inhibition of LTP has been recorded in 30 PND CA3, whereas potentiation has been measured in 60 PND CA3 with either exposure paradigm that have been attributed to possible involvement of protein kinase C (Hussain et al., 2000).

**Ketamine:** Ketamine has been suggested to inhibit the NMDA receptor by two distinct mechanisms: (1) by blocking the open channel and thereby reducing channel mean open time, and (2) by decreasing the frequency of channel opening through an allosteric mechanism (Orser et al., 1997).

Recently, the effects of ketamine, MK-801 and NMDA on Ca2+ oscillations (measured with Fluo-4-loaded scanning microscope) of hippocampal neurons from 24-h postnatal Sprague-Dawley rats have been investigated (Huang et al., 2013). In this cellular model, 100 µM NMDA causes an increase of Ca2+ oscillation frequency (25% compared to control) that is not statistically significant, whereas MK-801, at 40 µM significantly decreases the amplitude and frequency of the Ca2+ oscillations both by 25% (Huang et al., 2013). However, Sinner et al. (2005) have shown that the same dose of MK-801 causes complete inhibition of the Ca2+ oscillations in the same cellular model at 17-18 DIV and not at the 5 DIV, suggesting that different effects are triggered depending on the developmental window.

In primary cortical neurons derived from CD1 mice (E15-16), the addition of 10 µM of MK-801 completely suppresses Ca2+ influx. When MK-801 is removed, neurons show no response in terms of Ca2+ influx to a subsequent exposure to glutamate (Jiang et al., 2010). On the other hand, memantine at 15 µM concentration has been effective in inhibiting Ca2+ influx (approximately 40%) in cortical neurons. However, this inhibition is reversible as after washing out, neurons are able to respond immediately to the presence of glutamate showing an increase in Ca2+ influx in cortical neurons. These results suggest that
memantine can inhibit glutamate-mediated Ca2+ influx in cortical neurons in a transient and reversible manner (Jiang et al., 2010).

Ketamine has been tested in a large range of concentrations but only above 300 µM inhibits significantly the frequency of the Ca2+ oscillations (25%) and at 3,000 µM completely abolishes the neuronal Ca2+ oscillations (Huang et al., 2013). Lower concentrations (1-10 µM) cause small increase in Ca2+ oscillations similar to the studies of Sinner et al. 2005 that showed increase of the Ca2+ oscillation frequency at 3 µM of ketamine. In cell-attached patches, 1 µM ketamine decreased both the frequency of channel opening and the mean open time (44% and 68% of control values, respectively) (Orser et al., 1997).

Uncertainties or Inconsistencies

The structural diversity of NMDA subunits can influence the functionality of the receptors and their permeability to Ca2+. For example, NR2B subunits show higher affinity for glutamate binding and higher Ca2+ permeability (reviewed in Higley and Sabatini, 2012). But NMDA receptor subunit composition is not the only parameter that influences Ca2+ entrance in the cytosol. Membrane potential due to pore blockade by extracellular Mg2+ and receptor phosphorylation are two additional regulator of Ca2+ influx through NMDA receptors (reviewed in Higley and Sabatini, 2012).

Entrance of Ca2+ into neuronal cell can also happen through KA and AMPA receptors but to a smaller extend compared to NMDA receptors (reviewed in Higley and Sabatini, 2012). However, recent findings suggest that AMPA receptors may also contribute to Ca2+ signalling during CNS development (reviewed in Cohen and Greenberg, 2008). Early in development cortical pyramidal neurons express calcium-permeable, GluR2 subunit-lacking AMPA receptors. During postnatal development these neurons undergo a switch in the subunit composition of AMPA receptors, expressing instead GluR2-containing, calcium-impermeable AMPA receptor suggesting that the main point entrance of Ca2+ at this developmental stage are NMDA receptors.

Furthermore, Ca2+ entry occurs through L-type voltage-dependent Ca2+channels (L-VDCCs) (Berridge, 1998; Felix, 2005) that are encountered in neurons, suggesting that there are more possible entrance sites for Ca2+ to get into the cytosol rather than only through NMDA receptors.

Ketamine has been shown to cause increase of Ca2+ in cytosol although it is an antagonist of NMDA receptors. The reason behind is that ketamine has been found together with D-AP5, GGS-19755 and MK-801 to upregulate NMDA receptors subunits that is considered to be associated with the excessive entry of Ca2+ inside the cell after exposure to these specific antagonists (reviewed in Wang, 2013). Ethanol has also been found to increase intracellular Ca2+ in cultured septo-hippocampal neurons (derived from 0 PND) in a dose dependent manner (Webb et al., 1997).

Interestingly, Pb2+ has the ability to mimic or even compete with Ca2+ in the CNS (Flora et al., 2006). Indeed, Pb2+ is accumulated in the same mitochondrial compartment as Ca2+ and it has been linked to disruptions in intracellular calcium metabolism (Bressler and Goldstain, 1991). So, it can be that the reduced levels of Ca2+ after Pb2+ exposure may not be attributed to NMDA receptor inhibition but also to the ability of this heavy metal to compete with Ca2+. To make things more complicated, recent findings suggest that BDNF can also acutely elicit an increase in intracellular Ca2+ concentration, which is attributed not only to the influx of extracellular Ca2+ but also to Ca2+ mobilization from intracellular calcium stores (Numakawa et al., 2002; He et al., 2005). These finding derive from primary cultures of cortical neurons (E18 or 2-3 PND), where BDNF-evoked Ca2+ signals have not been altered neither by tetrodotoxin nor by a cocktail of glutamate receptor blockers (CNQX and APV), pointing out the importance of BDNF in Ca2+ homeostasis (Numakawa et al., 2002; He et al., 2005).

Quantitative Understanding of the Linkage

Is it known how much change in the first event is needed to impact the second? Are there known modulators of the response-response relationships? Are there models or extrapolation approaches that help describe those relationships?
Evidence Supporting Taxonomic Applicability

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Besides the above studies described in rodents, intracellular Ca2+ regulation has been studied at the neuromuscular junction of larval Drosophila exposed to 0, 100 μM or 250 μM Pb2+ (He et al., 2009).

References


Calcium influx, Decreased Indirectly Leads to Release of BDNF, Reduced

How Does This Key Event Relationship Work

Mainly, NMDA receptor activation initiates Ca2+-dependent signaling events that regulate the expression of genes involved in regulation of neuronal function including bdnf (reviewed in Cohen and Greenberg, 2008). Inhibition of NMDA receptors results in low levels of Ca2+ and decreased transcription of BDNF and consequently to low level of BDNF protein production and release.
Weight of Evidence

Biological Plausibility

BDNF transcription is induced by Ca2+ entering through either L type voltage gated calcium channel (L-VGCC) (Tao et al., 1998) or NMDA receptor (Tabuchi et al., 2000; Zheng et al., 2011) that can last up to 6 h. BDNF IV that is the most studied among its different exons has been shown to bind three Ca2+ elements within the regulatory region (reviewed in Zheng et al., 2012). One of these Ca2+ elements binds to CREB facilitating transcription. However, more transcription factors rather than only CREB are implicated in the transcription process of BDNF such as NFAT (nuclear factor of activated T cell), MEF2 (myocyte enhancer factor 2) and NFκB (nuclear factor kB) (reviewed in Zheng et al., 2012). The activation of the relevant transcription factor is triggered by the initial activation of CaM kinase, cAMP/PKA and Ras/ERK1/2 pathways mediated by the elevated intracellular Ca2+. Inhibitory studies targeting different elements of these pathways show that can cause decrease at mRNA BDNF levels (reviewed in Zheng et al., 2012).

In particular, exon IV BDNF mRNA transcription is regulated by a transcriptional silencer, methyl-CpG binding protein 2 (MeCP2), demonstrating that epigenetic alterations can also regulate BDNF transcription. Increase of intracellular Ca2+ levels phosphorylates MeCP2, which inactivates its repressor function and permits the transcription of BDNF exon IV (Chen et al., 2003; Greer and Greenberg, 2008; Tao et al., 2009; Zhou et al., 2006). Indeed, NMDA receptor activation has been shown to upregulate BDNF transcripts containing exon IV not only via Ca2+-dependent CREB but also through Ca2+ activation of MeCP2 transcription (Metsis et al., 1993; Shieh et al., 1998; Tao et al., 1998; Tabuchi et al., 2000; Chen et al., 2003; Jiang et al., 2005; Zheng et al., 2011), whereas NMDAR antagonists decrease BDNF exon IV expression (Zafra et al., 1991; Stansfield et al., 2012). Furthermore, BDNF mRNA is also targeted in different locations within the cell during the process of translation, depending on the promoter used (reviewed in Tongiorgi et al., 2006).

Empirical Support for Linkage

Include consideration of temporal concordance here

There is no direct evidence linking reduced levels of Ca2+ to decreased BDNF levels as they have not been ever measured both in the same study after exposure to stressors. However, there are findings that strongly link the different elements of Ca2+-dependent signalling events to transcription of BDNF.

Pb2+: Pb2+ decreases the ratio of phosphorylated versus total MeCP2 and consequently MeCP2 maintains its repressor function and prevents BDNF exon IV transcription (Stansfield et al., 2012). MeCP2 gene expression in the frontal cortex is very sensitive to Pb2+ exposure while in the hippocampus, the same gene is affected only at the higher exposure group in rat pups with blood Pb2+ levels 5.8 to 10.3 μg/dl on PND 55 (Schneider et al., 2012). In two different in vivo studies from the same research group, the use of doses of Pb2+ that result in learning and LTP deficits in rats causes decrease in phosphorylation of CREB in cerebral cortex at 14 PND and the same reduction in phosphorylation state of CREB in both cortex and hippocampus at 50 PND (Toscano et al., 2002; 2003). Interestingly, under similar experimental conditions no alteration at the phosphorylation state of CAMKII has been recorded (Toscano et al., 2005).

MK-801: Administration of MK-801 to 7 PND rats has been found to cause sustained depletion of BDNF mRNA levels (Hansen et al., 2004). The same effect on BDNF mRNA levels has been recorded in immature neocortical neurons after treatment with MK801 (5 μM). In primary hippocampal neurons exposed to 1 μM Pb2+ for 5 days during the period of synaptogenesis (DIV7–DIV12), both the cellular proBDNF protein and extracellular levels of mBDNF decrease with the latter to smaller extend (Neal et al., 2010). In the same in vitro model Pb2+ also decreases dendritic proBDNF levels throughout the length of the dendrites and caused impairment of BDNF vesicle transport to sites of release in dendritic spines (Stansfield et al., 2012). Furthermore, Pb2+ treatment resulted in a specific reduction of Bdnf exon IV and IX mRNA transcripts causing no alteration in the expression of exons I and II (Stansfield et al., 2012). Rat pups on PND 25 exposed to Pb2+ (180 and 375-ppm lead acetate in food for 30 days) demonstrated blood Pb2+ levels 5.8 to 10.3 μg/dl on PND 55 and show no change at gene levels of BDNF (Schneider et al., 2012). In mouse
embryonic stem cells (ESCs), Bdnf exon IV has been found to be down-regulated in cells treated with 0.1 µM Pb, whereas Bdnf exon IX has been found up-regulated (Sánchez-Martín et al., 2013).

**Uncertainties or Inconsistencies**

Autry and his colleagues (2010) demonstrated that ketamine and some other NMDA receptor antagonists produce fast-acting behavioral antidepressant-like effects in mouse models, and that these effects depend on the rapid synthesis of BDNF. Increased BDNF levels in prefrontal cortex have been found on PND 15 after treatment of rats with 0.25 mg/kg MK-801 twice daily from PND 5 to PND 14 but went back to normal control baseline on PND 42 and 77. In contrast, BDNF expression in the hippocampus has not been influenced by neonatal MK-801 administration on PND 15 but there has been recorded significant late increase of BDNF expression on PND 42 and 77 (Guo et al., 2010). Acute administration of memantine at high dose (20 mg/kg), but not imipramine, increases BDNF protein levels in the rat hippocampus (Réus et al., 2010). Combined treatment of rats with ketamine (5 and 10 mg/kg) and imipramine (10 and 20 mg/kg) produces significant increase of CREB and BDNF protein levels in the prefrontal cortex, hippocampus and amygdala as well as PKA phosphorylation in the hippocampus and amygdala and PKC phosphorylation in prefrontal cortex (Réus et al., 2011).

Whereas the majority of the studies investigating the effect of ethanol in different areas of developing brain show decrease of BDNF, few reports fail to do so (reviewed in Davis, 2008). For example, Heaton et al. 1999 have found no differences in BDNF protein levels in cerebellum with numerous exposure paradigms around this age of interest. Few years later the same research group reported an increase of BDNF immediately after exposure to ethanol on PND 4 or 7 that dropped to normal levels by 2 and 12 hours, respectively (Heaton et al., 2003). It is believed that the discrepancy in these data is not likely due to timing of exposure since conflicting results have also obtained with gestational exposure, but may be due to the age of the animals that measurement has been carried out (reviewed in Davis, 2008).

In a gene expression study, where gene analysis has been performed in the hippocampus derived from male or female rats fed with 1500 ppm Pb2+-containing chow for 30 days beginning at weaning, two molecular networks have been identified that were different between male and female treated rats. In these networks, CREB was the highly connected node, common for both networks (Schneider et al., 2011). However, no change has been reported in the expression of bdnf gene neither in male nor in female rats treated with Pb2+ (Schneider et al., 2011).

**Quantitative Understanding of the Linkage**

*Is it known how much change in the first event is needed to impact the second? Are there known modulators of the response-response relationships? Are there models or extrapolation approaches that help describe those relationships?*

**Evidence Supporting Taxonomic Applicability**

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**References**


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**Release of BDNF, Reduced Indirectly Leads to Dendritic morphology, Aberrant**

**How Does This Key Event Relationship Work**

The dendritically synthesized BDNF when secreted activates tyrosine kinase B (TrkB) receptors that induce the synthesis of a number of proteins involved in the development of proper dendritic spine morphology.

**Weight of Evidence**

**Biological Plausibility**

After activation of tyrosine kinase B (TrkB) receptors by BDNF proteins such as Arc, Homer2, LIMK1 (Kang and Schuman, 1996, Schratt et al., 2004 and Yin et al., 2002) that are known to promote actin polymerization and consequently enlargement of spine heads (Sala et al., 2001) are released. Recently, it has been shown that BDNF promotes dendritic spine formation by interacting with Wnt signaling. Indeed, Wnt signaling inhibition in cultured cortical neurons caused disruption in dendritic spine development, reduction in dendritic arbor size and complexity and blockage of BDNF-induced dendritic spine formation and maturation (Hiester et al., 2013).

In addition, it has been shown that the inhibition of BDNF synthesis reduces the size of spine heads and impairs LTP (An et al., 2008; Waterhouse and Xu, 2009). BDNF has been characterized as a critical factor in
promoting dendritic morphogenesis in various types of neurons (reviewed in Jan and Jan, 2010; Park and Poo, 2013).

BDNF that is synthesized in dendrites is known to regulate the morphology of spines (Tyler and Pozzo-Miller, 2003; An et al., 2008). For example, spines in the absence of spontaneous electrical activity are significantly smaller than normal (Harvey et al., 2005). On the other hand, simultaneous electrical activity and glutamate release increase the size of the spine head, which has been shown to be dependent on BDNF (Tanaka et al., 2008).

Mice harboring the Val66Met mutation of Bdnf gene show dendritic arborization defects in the hippocampus. Interestingly, human subjects with the Val66Met SNP demonstrate similar anatomical features (reviewed in Cohen and Greenberg, 2008).

More targeted studies have shown that, within the physiological range of expression, dendritic spine density is tightly regulated by BDNF in the dentate gyrus but not in CA1 pyramidal cells (Alexis and Stranahan, 2011).

**Empirical Support for Linkage**

*Include consideration of temporal concordance here*

Exposure of rat hippocampal neurons in culture to BDNF causes increase in cypin mRNA and protein levels, which is a known guanine deaminase that activates dendritic arborisation. This increase of cypin induced by BDNF appears after 72 h but not at earlier time points (Kwon et al., 2011), meaning that BDNF has to act first in order to stimulate dendritic arbor formation.

**Pb2+:** The first hint for involvement of Pb2+ in dendritic morphology was described by Alfano and Petit. 1982. They have demonstrated reduction in the length of dendritic processes and the number of dendritic branches in hippocampal dentate granule cells after developmental Pb2+ exposure of Long-Evans hooded rat pups (Alfano and Petit, 1982). More recently, it has been shown that the chronic exposure of rats to environmentally relevant levels (Pb2+ blood levels 25.8 ± 1.28 μg/dL) during early life alters cell morphology in the dentate gyrus as immature granule cells immunolabeled with doublecortin display aberrant dendritic morphology (Verina et al., 2007).

Exposure of rats to Pb2+ that initiated at embryonic phase and terminated at PND 21 have revealed that at PND 14 (Pb2+ concentration in the hippocampus 0.249±0.06 μg/g) and PND 21 (Pb2+ concentration in the hippocampus 0.471±0.11 μg/g) the number of dendritic spine on hippocampal CA1 area decreases by 32.83% and 24.11%, respectively (Hu et al., 2014). The length-density of the doublecortin-positive apical dendrites in the outer portion of the dentate gyrus molecular layer has been found significantly decreased up to 36% in chronically exposed rats to environmentally relevant levels of Pb2+ (Pb2+ blood levels 25.8 ± 1.28 μg/dL) (Verina et al., 2007). In another in vivo study, lower blood levels of Pb2+ (10 ± 1.28 μg/dL) in similar age of rats has led to significant decrease of BDNF levels (39% in forebrain cortex and 29% in hippocampus) (Baranowska-Bosiacka et al., 2013).

In cultured rat hippocampal neurons, low levels of Pb2+ (0.1 and 1 μM) cause reduction of dendritic spine density in a dose-dependent manner (Hu et al., 2014). In the same in vitro model, exposure to 1 μM Pb2+ for 5 days during the period of synaptogenesis (DIV7–DIV12), significantly reduces proBDNF protein and extracellular levels of mBDNF (Neal et al., 2010). When mouse embryonic stem cells are differentiated into neurons, exposure to lead (II) acetate causes reduction in the percentage of microtubule-associated protein 2 (MAP-2)-positive cells and in the mRNA levels of MAP-2 in a dose-dependent manner (Baek et al., 2011).

**Ketamine:** Female pregnant rats on gestational day 14 (n = 3) received ketamine with an initial intramuscular injection (40 mg/kg) followed by continuous intravenous infusion by pump via tail vein at a rate of 40–60 mg/kg/h for 2 h. The pyramidal dendrites in the ketamine treated pups (PND 0 and PND 30) have been found to be less branched, whereas the total branch length has been detected to be shorter than in controls only in the CA3 and not in the CA1 region of hippocampus (Zhao et al., 2014). Additionally, spine density has been observed to be significantly decreased in the ketamine treated offspring compared
MK-801: After a 20-min preincubation of a synaptoneurosomal preparation with MK-801, BDNF-induced increase of Arc synthesis has been significantly reduced (Yin et al., 2002), demonstrating that MK-801 can alter the morphology of dendritic spines through BDNF regulation (Yin et al., 2002).

Uncertainties or Inconsistencies

Various molecular mechanisms have been identified as regulators of dendritic arborisation patterns and dendritic spine formation (Jan and Jan, 2010). More specific, transcription factors, growth factors, receptor-ligand interactions, various signalling pathways, local translational machinery, cytoskeletal elements, Golgi outposts and endosomes have been identified as contributors to the organization of dendrites of individual neurons and the contribution of these dendrites in the neuronal circuitry (Jan and Jan, 2010). This study suggests that more parameters rather than only BDNF may be involved in dendritic arbor and spine formation during development.

Female pregnant rats on gestational day 14 (n = 3) received ketamine with an initial intramuscular injection (40 mg/kg) followed by continuous intravenous infusion by pump via tail vein at a rate of 40–60 mg/kg/h for 2 h. Pyramidal dendrites in the ketamine treated pups PND 0 and PND 30 were less branched and total branch length was shorter than in controls in the CA3, but not in the CA1 region (Zhao et al., 2014). Additionally, spine density was significantly decreased in the ketamine treated offspring compared with that of the controls in the CA3 region (Zhao et al., 2014). Previously, Ibla et al. 2009 showed that treatment with ketamine (20 mg/kg) at 90-minute intervals over 9 hours to PND 7 rat pups increase BDNF cDNA products and protein levels. However, the day of ketamine treatment and the age of animals among these two studies are not similar and consequently do not allow us to evaluate the strength of this KE relationship for this specific stressor.

Quantitative Understanding of the Linkage

Is it known how much change in the first event is needed to impact the second? Are there known modulators of the response-response relationships? Are there models or extrapolation approaches that help describe those relationships?

Evidence Supporting Taxonomic Applicability

In organotypic slice cultures derived from the ferret visual cortex application of exogenous BDNF increased the length and complexity especially of Layer IV pyramidal neurons (McAllister et al., 1995) that was also activity-dependent (McAllister et al., 1996). Several studies conducted in rodents further support that the in vitro treatment of hippocampal cultures with exogenous BDNF increases dendritic growth in developing neurons (reviewed in Zagrebelsky and Korte, 2014).

References


Release of BDNF, Reduced Indirectly Leads to Presynaptic release of glutamate, Reduced

How Does This Key Event Relationship Work

BDNF, acting via its specific presynaptic receptor TrkB, has been shown to increase excitatory synaptic transmission by triggering presynaptic glutamate release in hippocampal cultures as well as in hippocampal and cortical slices (Lessmann et al., 1994; Kang and Schuman, 1995; Carmignoto et al., 1997; Mohajerani et al., 2007).

Weight of Evidence

Biological Plausibility

Experimentally, it has been shown that presynaptically, BDNF enhances glutamate release and increases the frequency of mEPSCs in hippocampal neurons of rat (Lessmann and Heumann, 1998; Takei et al., 1998; Minichiello, 2009). It has been reported that BDNF rapidly induces glutamate transporter-mediated glutamate release via phospholipase C-γ (PLC-γ)/Ca2+ signaling and that antidepressants enhance PLC-γ/Ca2+ signaling leading to reduced levels of BDNF that cause decreased glutamate release (Numakawa et al., 2002; Yagasaki et al., 2006).

Empirical Support for Linkage

Include consideration of temporal concordance here

- In cortical cultured neurons obtained from PND 2-3 rat pups, BDNF fails to induce glutamate release at DIV 3 and 4. However, after 5 days in vitro culture or more (DIV 6-9), BDNF (100 ng/ml) induces significant glutamate release (2-2.8 fold) within 1 min after exogenous application (Numakawa et al., 2002).
- It has been shown that there is a dose-dependent effect of BDNF on the glutamate release. The glutamate release is initially observed at 5 ng/ml BDNF and reaches a plateau at 100 ng/ml (Numakawa et al., 2002).
- No studies have been found in the literature measuring both KEs after exposure to the stressors. Interestingly, proton magnetic resonance spectroscopy in adults with childhood lead exposure shows decrease in a composite of glutamate and glutamine in vermis and in parietal white matter of the brain (Cecil et al., 2011).

Uncertainties or Inconsistencies
A dose-response study using microdialysis techniques in conscious rats shows that low doses of ketamine (10, 20, and 30 mg/kg) increase glutamate outflow in the prefrontal cortex (Moghaddam et al., 1997). A higher dose close to the anaesthetic dose of ketamine (200 mg/kg) decreases glutamate levels, whereas an intermediate dose of 50 mg/kg has no effect (Moghaddam et al., 1997). In another microdialysis study, NMDA (250 and 500 µM) causes dose-dependent increase in extracellular concentrations of glutamate in the striatum of rats (Bustos et al., 1992). This neurotransmitter increase has been reduced in a dose-related way by prior perfusion with 75 µM MK-801. MK-801, at the same dose causes no changes on basal levels of glutamate. However, 100 µM MK-801 did increase glutamate extracellular concentrations (Bustos et al., 1992).

In mice, quantitative RT-PCR analysis has shown that BDNF mRNA expression is unaltered by ketamine (3.0 mg/kg) or MK-801 at either 30 min or 24 h after treatment (Autry et al., 2011). In contrast, Western blot and ELISA analyses have shown that BDNF protein levels increase at 30 min but not at 24 h after ketamine or MK-801 treatment (Autry et al., 2011).

Recently, in heterozygous BDNF-knockout (BDNF+/−) mice it has been demonstrated that the reduced BDNF levels did not affect presynaptic glutamate release (Meis et al., 2012).

**Quantitative Understanding of the Linkage**

*Is it known how much change in the first event is needed to impact the second? Are there known modulators of the response-response relationships? Are there models or extrapolation approaches that help describe those relationships?*

**Evidence Supporting Taxonomic Applicability**

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**References**


Meis S, Endres T, Lessmann V. (2012) Postsynaptic BDNF signalling regulates long-term potentiation at
Cell death, N/A Indirectly Leads to Synaptogenesis, Decreased

How Does This Key Event Relationship Work

Under physiological conditions, in the developing nervous system, apoptosis occurs during the process of synaptogenesis, where competition leads to the loss of excess neurons and to the connection of the appropriate neurons (Buss et al., 2006; Mennerick and Zorumski, 2000; Oppenheim, 1991). However, when a stressor increases the number of apoptotic cells this KE has a negative effect on synaptogenesis as the reduced number of neurons (besides the ones that have been already eliminated through the physiological process of apoptosis) provides limited dendritic fields for receiving synaptic inputs from incoming axons. At the same time the loss of neurons also means that there are less axons to establish synaptic contacts (Onley, 2014), leading to reduced synaptogenesis and neuronal networking.

Weight of Evidence

Biological Plausibility

Recently, Dekkers et al. 2013 have reviewed how under physiological conditions components of the apoptotic machinery in developing brain regulate synapse formation and neuronal connectivity. For example, caspase activation is known to be required for axon pruning during development to generate neuronal network (reviewed in Dekkers et al., 2013). Experimental work carried out in Drosophila melanogaster and in mammalian neurons shows that components of apoptotic machinery are involved in axonal degeneration that can consequently interfere with synapse formation (reviewed in Dekkers et al., 2013). Furthermore, Bax mutant mice studies indicate that the lack of this pro-apoptotic protein BAX leads to disruption of intrinsically photosensitive retinal ganglion cells spacing and dendritic stratification that affects synapse localization and function (Chen et al., 2013).

Empirical Support for Linkage

Include consideration of temporal concordance here
Synaptogenesis and refinement of the cortical network precedes the programmed cell death of neurons during development (Innocenti and Price, 2005).

**Pb2+:** Elevated blood Pb2+ concentrations that have been evident in new-born rats prenatally exposed to 30 or 200 mg/l Pb2+ caused postnatally delay in synaptogenesis (McCauley et al., 1982). In this study, Pb2+ treatment depresses synaptic counts in pups of PND 11 to 15 but not in older pups (McCauley et al., 1982). In rat hippocampal primary cultures, Pb2+ treatment has no effect on PSD95 puncta density nor has any effect on Synapsin Ia/b total gray value, puncta density, and integrated intensity but only reduces the phosphorylation of Synapsin Ia/b (Stansfield et al., 2012). Pb2+ exposure also represses the expression of presynaptic vesicular proteins implicated in neurotransmitter release, such as synaptobrevin (VAMP1) and synaptophysin (SYN) (Neal et al., 2010).

**NMDA receptor antagonists:** Developmental treatment of rats (PND 15 up to PND 30) with the competitive NMDAR antagonist 2-amino-5-phosphonovaleric acid (APV) or the noncompetitive antagonist phencyclidine (PCP) causes decrease in the total number of synapses (Brooks et al., 1997). Withdrawal from NMDA antagonist initially affects similarly the synapses, whereas by PND 36 there has been full recovery in synapse number compared to the control (Brooks et al., 1997).

**Various anaesthetics:** The detrimental effect of general anaesthetics on synaptogenesis has been shown to be age-dependent and most severe if the exposure occurs at the peak of synaptogenesis (Yon et al., 2005). Exposure of rats to anaesthesia mixture (isoflurane, nitrous oxide, and midazolam) during the peak of synaptogenesis, causes long-lasting injury in subiculum characterised by mitochondria degeneration, decrease in the number of neurons possessing multiple synaptic boutons and reduction of synapse volumetric densities (Lunardi et al., 2010; 2011). The same research group has previously reported that this type of anaesthesia causes neuronal loss (30-40% compared to controls) in the same brain area in both rats and guinea pigs (Rizzi et al. 2008; Nikizad et al. 2007) and has concluded that this non selective synaptic loss after anaesthesia is attributed to the overall neuronal loss.

In a recent study, it has been shown that exposure of mice to general anaesthesia at the peak of synaptogenesis causes significant reductions in dendritic filopodial spines and synapse formation in hippocampal neurons (Head et al., 2009). These results indicate strong age-dependent vulnerability of synaptic spines and boutons during early stages of brain development.

**Uncertainties or Inconsistencies**

In adult nervous system, the role of apoptotic machinery in axon pruning and synapse elimination, which are necessary to refine mature neuronal network has been extensively studied (reviewed in Hyman and Yuan, 2012), whereas in developing nervous system the regulatory importance of apoptotic machinery in synapse formation and function is less clear (reviewed in Dekkers et al., 2013).

**Pb2+:** In mouse ESCs cultured in 3D aggregates, the treatment with 0.1 µM Pb2+ causes around 25% of cell loss that is not attributed to apoptosis as no change in caspase 3 levels has been detected (Sánchez-Martín et al., 2013). In the same study but in an in vivo model, Pb2+ causes downregulation of Syn1 gene expression in the hippocampus of male offspring (PND 60) derived from female mice exposed to 0 or 3 ppm of lead acetate in drinking water from 8 weeks prior to mating, through gestation and until postnatal day PND 10 (Sánchez-Martín et al., 2013).

**Ketamine:** The expression of PSD-95 is differently regulated between males and females and it has been found to be decreased in males but significantly increased in females exposed to ketamine (50 mg/kg) during gestation day 15 (GD 15) to GD 20 (Aligny et al., 2014). In the same study, apoptosis has been measured but not under the same experimental conditions (Aligny et al., 2014).

**Memantine:** Memantine treatment of cultured cerebellar granule cells has a stimulatory effect on dendritic spine maturation and excitatory synapse formation (Wei et al., 2012).

**Quantitative Understanding of the Linkage**
Is it known how much change in the first event is needed to impact the second? Are there known modulators of the response-response relationships? Are there models or extrapolation approaches that help describe those relationships?

Evidence Supporting Taxonomic Applicability

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| Experiments have been conducted both in Drosophila melanogaster and in mammals in order to elucidate the relationship between components of apoptotic machinery and synaptogenesis (reviewed in Dekkers et al., 2013).

References


Dendritic morphology, Aberrant Indirectly Leads to Synaptogenesis, Decreased

How Does This Key Event Relationship Work

It is well-established that loss of dendritic spine density and dendrite branch complexity leads to loss of synapse formation. Indeed, huge amount of research has been performed on dendrite arbour, dendritic spines and the molecular components of these structures that led to the elucidation of their role in higher order brain functions, including learning and memory (reviewed in Sjöström et al., 2008).

Weight of Evidence

Biological Plausibility

It has been proved that the appearance of extensive dendritic arbor and new spines coincides with synapse formation (Zito et al., 2004). Zhang and Benson (2001) have investigated the role of actin (the main component of dendritic spines) during the early stages of neuronal development by introducing an actin depolymerization protein named latrunculin A and conducting fluorescent imaging of synapse formation. At the early stages of neuronal development, it has been reported that the depolymerisation of filamentous actin (F-actin) significantly reduces the number of stable synapses and the presence of postsynaptic...
proteins (PSD-95, neuroligins, and Bassoon). Most importantly, pre- and postsynaptic vesicles needed for synaptogenesis have not been found at contact sites as a result of depolymerisation of F-actin (Zhang and Benson, 2001). Furthermore, synapsin I-deficient neurons have been shown to be unable to form synapses during the first week in culture even after establishing axon-dendritic contacts (Ferreira et al., 1996).

Empirical Support for Linkage

Include consideration of temporal concordance here

Many studies have indicated that synaptogenesis and spine formation happen in any order, meaning that not always synaptogenesis follows the spine formation but it can also happen the other way around (Bhatt et al., 2009; McAllister, 2007; Okabe et al., 2001).

**Pb2+:** Newborn rats exposed to 10 mg/ml of lead acetate from PND 2 up to PND 20 and 56 have showed significant decrease in the spine density as shown in Golgi staining of hippocampal pyramidal neurons of the CA1 region (Kiraly and Jones, 1982).

**Ketamine:** Peripheral administration of the N-methyl-D-aspartate receptor antagonist dizocilpine maleate, DL-2-amino-5-phosphonovalerate or ketamine on PND 20 has revealed a 30% decrease in asymmetric synapses in the dorsolateral striatum of treated rats (Butler et al., 1999).

**Ethanol:** Prenatal alcohol exposure causes significant decrease in the number of dendritic spines of pyramidal neurons in the visual cortex when mice pups have been assessed from PND 0 up to PND 30. The changes have been dose dependent and persisted until PND 30. At the same time point decreased numbers of synaptic vesicles have been recorded that also persisted up to PND 30 (Cui et al., 2010). In embryonic rat hippocampal neurons, 6 days of ethanol (200, 400 or 600 mg/dl) treatment, causes decrease in total dendritic length per cell, dendrite number per cell, length of individual dendrites and synapse number per innervated dendrite but has no effect on cell survival (Yanni and Lindsley, 2000). This reduction in synapse number has been associated with dendrite length, suggesting that ethanol’s effects on synapse number are secondary to its effects on dendritic morphology (Yanni and Lindsley, 2000).

Uncertainties or Inconsistencies

**Ketamine:** Administration of either ketamine-xylazine or isoflurane in 1-month-old mice rapidly alters dendritic filopodial dynamics but has no significant effects on spine dynamics (Yang et al., 2011). Ketamine-xylazine increases filopodial formation, whereas isoflurane decreases filopodial elimination during 4 h of anesthesia (Yang et al., 2011). Both effects have been transient and disappeared within a day after the animals woke up.

**Ethanol:** Rats (PND 19) that have been pair-fed with 6.6% (v/v) ethanol-containing liquid diets on a chronic basis prior to parturition show no change at the density of synapses in layer I of the motor cortex compared to controls (Druse et al., 1986). However, this study has not excluded that synaptogenesis is possibly affected by ethanol but at another age or in other regions of the cerebral cortex.

Quantitative Understanding of the Linkage

Is it known how much change in the first event is needed to impact the second? Are there known modulators of the response-response relationships? Are there models or extrapolation approaches that help describe those relationships?

Evidence Supporting Taxonomic Applicability

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Synaptogenesis, Decreased Directly Leads to Neuronal network function, Decreased

How Does This Key Event Relationship Work

The ability of a neuron to communicate is based on neural network formation that relies on functional synapse establishment (Colón-Ramos, 2009). The main roles of synapses are the regulation of intercellular communication in the nervous system, and the information flow within neural networks. The connectivity and functionality of neural networks depends on where and when synapses are formed. Therefore, the decreased synapse formation during the process of synaptogenesis is critical and leads to decrease of neural network formation and function in the adult brain.
Weight of Evidence

Biological Plausibility

Neuronal connections are established via the process of synaptogenesis. The developmental period of synaptogenesis is critical for the formation of the basic circuitry of the nervous system, although neurons are able to form new synapses throughout life (Rodier, 1995). The brain electrical activity dependence on synapse formation is critical for proper neuronal communication. Alterations in synaptic connectivity lead to refinement of neuronal networks during development (Cline and Haas, 2008). Indeed, knockdown of PSD-95 arrests the functional and morphological development of glutamatergic synapses (Ehrlich et al., 2007).

Empirical Support for Linkage

Include consideration of temporal concordance here

Pb2+: At low Pb2+ levels (less than 30 micrograms/dl), slow wave voltage tending have been observed that are positive in children under five years old but negative in children over five years. However, reverse age-related polarity shift has been observed in children with higher Pb2+ levels (Otto and Reiter, 1984).

In experiments carried out in Wistar rats that have been fed with lead acetate (400 micrograms lead/g body weight/day) from PND 2 until PND 60, EEG findings show statistically significant reduction in the delta, theta, alpha and beta band of EEG spectral power in motor cortex and hippocampus with the exception of the delta and beta bands power of motor cortex in wakeful state (Kumar and Desiraju, 1992).

Male Sprague-Dawley rats have been exposed to Pb2+ from parturition to weaning though their dams' milk that received drinking water containing 1.0, 2.5, or 5.0 mg/ml lead acetate (McCarren and Eccles, 1983). Beginning from 15 weeks of age, the characteristics of the electrically elicited hippocampal afterdischarge (AD) and its alteration by phenytoin (PHT) showed significant increase in primary AD duration only in the animals exposed to the higher dose of Pb2+, whereas all groups responded to PHT with increases in primary AD duration (McCarren and Eccles, 1983).

Ketamine: In primary rat cortical neurons (12-22 DIV), ketamine (50 µM) has been reported to reduce mean firing rate (MFR) by around 90%, whereas Pb2+ at the same concentration slightly increases MFR as measured by MEA technology (McConnell et al., 2012).

Whole-cell patch-clamp recordings in an in vitro preparation of forebrain slices from immature and mature rats have revealed that ketamine exposure (1, 5 and 10 µM) inhibits evoked NMDAR-mediated excitatory postsynaptic currents (eEPSCs) in a dose-dependent manner in both immature and mature neurons. However, more extensive inhibition of eEPSCs has been noted in neonatal neurons (75%) rather than in adult neurons (55%) after application of ketamine at higher concentration (10 µM) (Jin et al., 2013).

Ethanol: The recording in hippocampal slices derived from rats (PND 50-70) have showed that ethanol exposure during the GD 1-10 or PND 1-14 produces only minor changes in synaptic plasticity of offsprings after measuring theta-burst stimulation (TBS) and high-frequency stimulation (HFS). However, ethanol exposure during GD 11-22 causes pronounced reduction (approximately 54%) of long-term potentiation (LTP), suggesting that the timing of exposure influences the severity of neuronal network dysfunction (Helfer et al., 2012). A previous study has also shown that prenatal treatment with 5 g/kg/day ethanol causes significantly higher theta score compared to controls after recording with bipolar electrode that has been surgically implanted in rats of 90 days of age (Cortese et al., 1997).

Male Sprague-Dawley rats have been exposed to ethanol (6.0 g/kg/day) between PND 4-9 and recording electrodes have been implanted into the brain of each rat at the age of 3.5-4 months. The EEG activity derived from these animals reveals that ethanol exposure increases the peak frequency in the frontal cortical and parietal cortical 16-32 Hz frequency bands, whereas measurement of auditory event-related potentials (ERPs) shows that parietal cortical N1 amplitude is reduced (Slawecki et al., 2004).
Uncertainties or Inconsistencies

There has been a case report in which exposure to alcohol during the second half of twins pregnancy caused slowing down of the background activity on EEG and delay in both motor and cognitive function during the first year of life but only in one of the twins that were born (Riikonen, 1994).

Neonatal rat pups and their mothers have been exposed to ethanol for 4 h/d between PND 2 and PND 12 (pup serum ethanol concentration, 0.16 g/dl) and the function of climbing fibers-Purkinje cells synapses have been assessed by using patch-clamp electrophysiological techniques in slices from the cerebellar vermis. This study revealed that the baseline characteristics of the complex spike remains unaffected by ethanol exposure (Zamudio-Bulcock et al., 2014).

Quantitative Understanding of the Linkage

*Is it known how much change in the first event is needed to impact the second? Are there known modulators of the response-response relationships? Are there models or extrapolation approaches that help describe those relationships?*

Evidence Supporting Taxonomic Applicability

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Neuronal network function, Decreased Indirectly Leads to Learning and memory, Impairment

How Does This Key Event Relationship Work

Mammalian learning and memory is one of the outcomes of the functional expression of neurons and neural networks. Damage or destruction of neurons by chemical compounds during development when they are in the process of synapses formation, integration and formation of neural networks, will derange the organization and function of these networks, thereby setting the stage for subsequent impairment of learning and memory. Exposure to the potential developmental toxicants during neuronal differentiation and synaptogenesis will increase risk of functional neuronal network damage leading to learning and memory impairment.

Weight of Evidence

Biological Plausibility

Long-term potentiation (LTP) is a long-lasting increase in synaptic efficacy after high-frequency stimulation of afferent fibers, and its discovery potentiated the idea that individual synapses possess the properties expected for learning and memory (reviewed in Lynch et al., 2014). Moreover, LTP is intimately related to the theta rhythm, an oscillation long associated with learning. Learning-induced enhancement in neuronal excitability, a measurement of neural network function, has also been shown in hippocampal neurons following classical conditioning in several experimental approaches (reviewed in Saar and Barkai, 2003).

On the other hand, memory requires the increase in magnitude of EPSCs to be developed quickly and to be persistent for few weeks at least without disturbing already potentiated contacts. Once again, a substantial body of evidence have demonstrated that tight connection between LTP and diverse instances of memory exist (reviewed in Lynch et al., 2014).

Empirical Support for Linkage

Include consideration of temporal concordance here

A series of important findings support that the biochemical changes that happen after induction of LTP also occur during memory acquisition, showing temporality between the two KEs (reviewed in Lynch et al.,
Furthermore, a review on Morris water maze (MWM) as a tool to investigate spatial learning and memory in laboratory rats also pointed out that the disconnection between neuronal networks rather than the brain damage of certain regions is responsible for the impairment of MWM performance (D’Hooge and De Deyn, 2001). Functional integrated neural networks that involve the coordination action of different brain regions are consequently important for spatial learning and MWM performance.

Morris et al. 1986 found that blocking the NMDA receptor with AP5 inhibits spatial learning in rats. Most importantly, in the same study they measured brain electrical activity and recorded that this agent also inhibits LTP, however, they have not proved that spatial learning and LTP inhibition are causally related (Morris et al., 1986). Since then a number of NMDA receptor antagonists have been studied towards their ability to induce impairment of learning and memory. It is worth mentioning that similar findings have been found in human subjects, where by combining behavioural and electrophysiological data from patients with temporal lobe epilepsy exposed to ketamine, they demonstrated involvement of NMDA receptors in human memory processes (Grunwald et al., 1999).

Pb2+: Exposure to low levels of Pb2+, during early development, has been implicated in long-lasting behavioural abnormalities and cognitive deficits in children (Bellinger et al., 1991; Baghurst et al., 1992; Finkelstein et al., 1998; Lanphear et al., 2000; Canfield et al., 2003; Bellinger 2004; Lanphear et al., 2005; Surkan et al., 2007) and experimental animals (Brockel et al., 1998; Murphy and Regan, 1999; Moreira et al., 2001). Multiple lines of evidence suggest that Pb2+ can impair hippocampus-mediated learning in animal models (reviewed in Toscano and Guilarte, 2005).

The majority of the studies addressing the effects of Pb2+ on hippocampal-associated spatial learning and memory processes have been carried out mainly in male rats (Cao et al., 2008, Gilbert et al., 2005); only a few studies have examined both sexes simultaneously (Jett et al., 1997, Xu et al., 2009). Female rats exposed to Pb2+ through gestation and lactation have shown more severe impairment of memory than male rats with similar Pb2+ exposures (Jett et al., 1997). De Souza Lisboa et al. (2005) have reported that exposure to Pb2+ during both pregnancy and lactation causes depressive-like behaviour (detected in the forced swimming test) in female but not male rats. A more recent study has investigated the neurobehavioral outcomes in Pb2+-exposed rats (250, 750 and 1500 ppm Pb2+ acetate in food) during gestation and through weaning and demonstrated that these outcomes are very much influenced by sex and rearing environment (Anderson et al., 2012). In females, Pb2+ exposure lessens some of the benefits of enriched environment on learning, whereas, in males, enrichment does help to overcome detrimental effects of Pb2+ on learning. Regarding reference memory, environmental enrichment has not been beneficial in females when exposure to Pb2+ occurs, in contrast to males (Anderson et al., 2012).

Wistar rat pups exposed to 0.2% Pb2+ via their dams' drinking water from PND 1 to PND 21 and directly via drinking water from weaning until PND 30 (Jaako-Movits et al., 2005). At PND 60 and 80, the neurobehavioural assessment has revealed that developmental Pb2+ exposure induces persistent increase in the level of anxiety and inhibition of contextual fear conditioning (Jaako-Movits et al., 2005). The same behavioural syndrome in rats has been described in Salinas and Höff, 2002 and is in agreement with observations on humans as children exposed to low levels of Pb2+ display attention deficit, increased emotional reactivity and impaired memory and learning (Finkelstein et al., 1998).

In experiments carried out in Wistar rats that were fed with lead acetate (400 micrograms lead/g body weight/day) from PND 2 until PND 60, EEG findings show statistically significant reduction in the delta, theta, alpha and beta band EEG spectral power in motor cortex and hippocampus but not in delta and beta bands power of motor cortex in wakeful state (Kumar and Desiraju, 1992). After 40 days of recovery, animals have been assessed for their neurobehaviour and revealed that Pb2+ treated animals show more time and sessions in attaining criterion of learning than controls (Kumar and Desiraju, 1992). Further data obtained using animal behavioral techniques demonstrate that NMDA mediated synaptic transmission is decreased by Pb2+ exposure (Cory-Slechta, 1995; Cohn and Cory-Slechta, 1993; 1994).

Ketamine: A number of retrospective clinical studies have shown that there is an association between anaesthesia early in life and late-onset learning impairment (Hansen et al., 2011; Ing et al., 2012; Kalkman...
et al., 2009). Most of the experimental approaches have looked into the effects of general anaesthesia on neurodevelopmental consequences following postnatal exposure both in rodents (Kong et al., 2012; Palanisamy et al., 2011; Zheng et al., 2013) and primates (Slikker et al., 2007; Paule et al., 2011). For example, rhesus monkeys (PND 5 or 6) exposed to a dose of ketamine sufficient to produce a light surgical plane of anaesthesia for 24 h have been tested for neurobehavioural alterations. By using an operant test battery to evaluate the cognitive function, ketamine-treated monkeys have shown lower training scores for at least 10 months after the administration of ketamine compared to controls (Paule et al., 2011).

The Morris water maze (MWM) is a test for learning and spatial memory in rodents that relies on distal cues to navigate from the start locations around the perimeter of an open swimming arena to locate a submerged escape platform (D'Hooge and De Deyn, 2001). Extensive evidence supports its validity as a measure of hippocampal dependent spatial learning and memory. In case of ketamine, rat offspring treated with this anaesthetic have needed more time to escape onto the platform compared to the controls (Zhao et al., 2014).

**MK-801:** Rats treated with MK-801 (0.25mg/kg twice daily) at PND 5-14 showed moderate working memory impairments in adolescence (PND 35) but a pronounced deficit in adulthood (PND 63) (Su et al., 2014).

**Uncertainties or Inconsistencies**

One of the most difficult issues for neuroscientists is to link neuronal network function to cognition, including learning and memory. It is still unclear what exactly modifications in neuronal circuits need to happen in order to alter motor behaviour as it is recorded in a learning and memory test (Mayford et al., 2012), meaning that there is no clear understanding about the how these two KEs are connected.

Several epidemiological studies where Pb2+ exposure levels have been studied in relation to neurobehavioural alterations in children have been reviewed in Koller et al. 2004. This review has concluded that in some occasions there is negative correlation between Pb2+ dose and cognitive deficits of the subjects (Koller et al. 2004), meaning that not always Pb2+ exposure is positive associated with learning and memory impairment in children.

**Quantitative Understanding of the Linkage**

*Is it known how much change in the first event is needed to impact the second? Are there known modulators of the response-response relationships? Are there models or extrapolation approaches that help describe those relationships?*

**Evidence Supporting Taxonomic Applicability**

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<td>mouse</td>
<td>Mus sp.</td>
<td>Strong</td>
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Release of BDNF, Reduced Indirectly Leads to Cell death, N/A

How Does This Key Event Relationship Work

BDNF influences the apoptosis occurring in developing neurons through two distinct mechanisms (Bernd, 2008). mBDNF can trigger prosurvival signaling after binding to TrkB receptor through inactivation of components of the cell death machinery and also through activation of the transcription factor cAMP-response element binding protein (CREB), which drives expression of the pro-survival gene Bcl-2 (West et al., 2001). On the other hand, proBDNF binds to the p75 neurotrophin receptor (p75NTR) and activates RhoA that regulates actin cytoskeleton polymerization resulting in apoptosis (Lee et al., 2001; Miller and Kaplan, 2001; Murray and Holmes, 2011). It is proved that reduced levels of BDNF can severely interfere with the survival of neurons in different brain regions, leading to cell death (Lee et al., 2001; Miller and Kaplan, 2001; Murray and Holmes, 2011).

Weight of Evidence

Biological Plausibility

BDNF mRNA levels dramatically increase between embryonic days 11 to 13 during rat development, playing important role in neuronal differentiation and survival (reviewed in Murray and Holmes, 2011). The latter has been supported by transgenic experiments where BDNF−/− mice demonstrated a dramatic increase in cell death among developing granule cells leading to impaired development of the layers of the cerebellar cortex (Schwartz et al., 1997). BDNF has also been shown to provide neuroprotection after hypoxic-ischemic brain injury in neonates (P7) but not in older (P21) animals (Cheng et al., 1997; Han et al., 2000). The neuroprotective role of BDNF has been further supported by the observed correlation between elevated BDNF protein levels and resistance to ischemic damage in hippocampus in vivo (Kokaia et al., 1996) and K+ rich medium-induced apoptosis in vitro (Kubo et al., 1995).

Empirical Support for Linkage

Include consideration of temporal concordance here

Several in vitro and in vivo studies on cortical neurons have demonstrated that the survival of developing neurons is closely related with the activation of the NMDA receptors and subsequent BDNF synthesis/release that fully support the BDNF neurotrophic theory (Ikonomidou et al., 1999; Yoon et al. 2003; Hansen et al. 2004).
**Pb2+:** Neonatal mice exposed to Pb2+ (350 mg/kg lead twice every 4 h) and sacrificed after 8-24 h have shown increased apoptotic neurodegeneration above that seen in normal controls. This effect has been recorded only in animals treated with Pb2+ at PND 7, but not on PND 14 (Dribben et al., 2011), confirming the importance of the time of exposure during development in order for Pb2+ to induce apoptosis. Two to four week old rats treated for 7 days with 15 mg/kg daily dose of lead acetate showed increased apoptosis in hippocampus (Sharifi et al., 2002). In rats (30 PND), it has also been shown that Pb2+ (2, 20 and 200 mg/kg/d) can induce apoptosis (Liu et al., 2010). However, in contrast to the first two in vivo studies, the animals in this experimental approach were old enough to evaluate the most sensitive window of vulnerability of developing neurons to Pb2+ exposure (Liu et al., 2010), confirming that only Pb2+ treatment during synaptogenesis can lead to neuronal cell apoptosis. In vitro evidence of apoptosis induced by Pb2+ also derive from PC12 cells exposed to Pb2+ (0.1, 1, 10 μM) that have shown increased activation of caspase-3 (Xu et al., 2006). Besides PC12 cells (Xu et al., 2006; Sharifi and Mousavi, 2008), lead-induced apoptosis has also been studied in cultured rat cerebellar neurons (Oberto et al., 1996), hippocampal neurons (Niu et al., 2002) and retinal rod cells (He et al., 2000). In primary rat hippocampal neurons exposed to 1 μM Pb2+ for 5 days during the period of synaptogenesis (DIV7–DIV12), decreased cellular proBDNF protein (40% compared to control) and extracellular levels of mBDNF (25% compared to control) have been recorded (Neal et al., 2010). Significant reductions specifically in dendritic proBDNF levels throughout the length of the dendrites have also been described by Stansfield et al. (2012) after exposure to the same concentration of Pb2+ using this in vitro model. In an in vivo study, mice at PND 7 with mean Pb2+ blood levels of 8.10 μg/mL have showed increased apoptosis in the cortex, hippocampus, caudate-putamen, and thalamus compared to controls with F (1,14) = 19.5, 8.40, 4.15, 4.53, respectively (Dribben et al., 2011). These Pb2+ levels in blood (Dribben et al., 2011) were a bit higher than the levels determined in Guilarte et al. 2003 (3.90 μg/dl) that served as the base to calculate in vitro doses in Neal et al. 2010 and Stansfield et al. 2012.

**MK-801:** Systemic administration of three doses of 0.5 mg/kg MK-801 every eight hours apart to rat pups on PND 7 causes widespread apoptosis only in neurons after 24h, as indicated by TUNEL and silver staining (Ikonomidou et al., 1999). This accelerated pattern of programmed cell death found to be identical to that observed during the naturally occurring apoptosis that occurs during brain maturation. In Ikonomidou et al. 1999, it has also been shown that the effect of MK-801 is dose-dependent above a threshold dose of 0.25 mg/kg, with higher doses inducing larger apoptotic responses. Besides this key experimental work, a significant number of in vivo studies that were carried out afterwards further support the MK-801 induced apoptosis in the immature rat and mouse brain that have been reviewed in Lim et al. 2012. MK-801 has also been shown to increase apoptosis in other cellular models like in motor neurons of a chick embryo preparation (Hsu et al., 2000) or in rat primary cortical cultures (Terro et al., 2000). In the latter cellular model, MK-801 induced apoptosis is more evident at early stage of culture (9 DIV) and gradually decreases as the culture matures (13-15 DIV), suggesting that less mature neurons are more sensitive to NMDA receptor blockage induced cell death. Systemic administration of MK-801 to 7 PND rats at doses that causes apoptosis has been found to lead to sustained depletion of BDNF mRNA levels (Hansen et al., 2004). Furthermore, in the same study, cultured immature neocortical neurons exhibit a significant transcriptional loss of BDNF following MK-801 (5 μM) exposure up to 60% compared to controls. The downregulation of BDNF mRNA levels could not be further decreased with increasing MK-801 concentrations up to 50 μM of MK-801. However, the higher concentration of MK-801 (50 μM) has caused the most severe increased (2.5 fold) in apoptosis of neurons (Hansen et al., 2004).

**Ketamine:** Data from developing rats and other rodent cell models suggest that ketamine can cause dose-dependent apoptosis (Ikonomidou et al., 1999; Scallet et al., 2004; Wang et al., 2005). In rhesus monkey frontal cortical cultures (PND 3) treatment with 1, 10, or 20 μM ketamine for 24 h causes a marked increase in DNA fragmentation and release of lactate dehydrogenase (Wang et al., 2006). In 5-day-old rhesus monkeys, administration of ketamine at doses that produce light surgical plane of anesthesia has been given for either 9 or 24 h resulting in neuroapoptosis (Slikker et al., 2007). Similarly, neuroapoptosis has been recorded even in the brain of the fetus after exposure of pregnant rhesus monkeys to ketamine for 24 h on day 122 of gestation (equivalent to the third trimester of human pregnancy), but no neuroapoptosis has been found following administration of ketamine on PND 35 (Slikker et al., 2007). Ketamine when is administered by i.v. infusion for 5 h to PND 6 rhesus neonates or to pregnant rhesus females at 120 gestation day causes apoptosis, that is 2.2 times greater in fetuses than in neonates brains (Brambrink et
al., 2012). Other NMDA receptor antagonists: In organotypic neocortical slices derived from newborn (P0) C56/balbC mice treatment with 20 μM(±) 3-(2-carboxypiperazin-4-yl)propyl-1-phosphonic acid (CPP), an NMDA receptor antagonist, induces significant increase in the number of apoptotic cells after all the time points tested (6, 12, and 24 h) (Heck et al., 2008). In the same study, the blockage of the NR2A subunit cells with 400 nM NVP-AMM077 causes increase in the apoptosis rate after 6, 12, and 24 h compared to the untreated controls. It is worth mentioning that the number of apoptotic cells is highest after 6 h treatment with NVP-AMM077 and significantly decreases after longer exposure times probably because NR2A-independent process is upregulated after 12 and 24 h. In contrast, the NR2B subunit specific antagonist ifenprodil has no significant effect on the apoptosis rate after 6 h application time, but the number of apoptotic neurons increase significantly after 12 and 24 h exposure to this antagonist (Heck et al., 2008).

**Various anaesthetics:** Treatment of immature mouse neurons to propofol can cause neuroapoptosis through disruption of proBDNF/p75NTR pathway. Knockout of p75NTR or administration of TAT-Pep5 (a specific p75NTR inhibitor) prevent propofol-induced neuroapoptosis (Pearn et al., 2012). In vivo studies with newborn rats (P7) exposed to highly pro-apoptotic anesthesia protocol (midazolam, isoflurane, nitrous oxide) for a period of 2, 4 or 6 h demonstrate that this type of anaesthesia modulates key steps in BDNF-activated apoptotic cascade in time-dependent fashion by activating both Trk-dependent in thalamus and Trk-independent p75NTR in cerebral cortex neurotrophic pathways (Lu et al., 2006). In the same rodent models, the involvement of the NMDA receptor in mediating the programmed cell death in developing brain has been confirmed by the use of some other non-competitive and competitive NMDA receptor antagonists, including ketamine, phencyclidine and carboxypiperazin-4-yl-propyl-1-phosphonic acid that have demonstrated similar results (Wang et al., 2001; Wang et al., 2003; Fredriksson et al., 2004).

**Ethanol:** The majority of studies measuring BDNF levels, TrkB, or BDNF signaling suggest that ethanol inhibits the neurotrophic activity of BDNF in the cerebellum and that this effect may contribute to cerebellar Purkinje cell loss (reviewed in Davis, 2008). In a well-designed study, ethanol induced disruptions in BDNF-TrkB neurotrophic signaling by altering the mRNA levels of BDNF and its receptors that happened as early as 1 h after treatment, whereas the loss of apoptotic suppression and thus death of Purkinje cells occur after 6 h (Ge et al., 2004), supporting temporal concordance between the two KEs. Ethanol, which toxicity is mainly induced by inhibition of NMDA receptor, can trigger widespread apoptosis in the brain of rodents after exposure during synaptogenesis (Ikonomidou et al., 1999, 2000; Dikranian et al., 2001). This neuropathological finding has also been found to be dose- and developmental age-dependent, depending on whether ethanol exposure happens in early, mid or late synaptogenesis (Ikonomidou et al., 1999, 2000; Dikranian et al., 2001). Widespread apoptotic cell death has been observed throughout many regions of the brain following a single exposure to ethanol in third trimester fetal monkeys (Farber et al., 2010).

**Uncertainties or Inconsistencies**

**Pb2+:** A number of studies demonstrate that deletion of BDNF does not lead to significant apoptotic cell death of neurons in the developing CNS (reviewed in Dekkers et al., 2013). In an in vivo Pb2+ exposure study, where female rats received 1,500 ppm prior, during breeding and lactation shows no changes at mRNA levels of BDNF in different hippocampus section derived from their pups (Guillarte et al., 2003). Regarding Pb2+, the pre- and neonatal exposure of rats to Pb2+ (Pb2+ blood levels below 10 μg/dL) show a decreased number of hippocampus neurons but no morphological or molecular features of severe apoptosis or necrosis have been detected in tested brains (Baranowska-Bosiacka et al., 2013). In contrast to the lack of apoptotic signs, reduced levels of BDNF have been recorded in forebrain cortex (39%) and hippocampus (29%) (Baranowska-Bosiacka et al., 2013). Pregnant rats have been exposed to lead acetate (0.2% in the drinking water) after giving birth until PND 20. At PND 20, blood Pb2+ levels in pups reached at 80 μg/dl. In these animals, the gene expression in different brain regions has been assessed and demonstrated that hippocampus is most sensitive with alterations beginning at PND 12 when caspase 3 mRNA increases after Pb2+ exposure (Chao et al., 2007). However, bcl-x and BDNF mRNA in the hippocampus have been significantly increased after caspase 3 increase, suggesting that the apoptotic signal activates a compensatory response by increasing survival factors like BDNF and that the temporality suggested in this AOP may not be accurate (Chao et al., 2007).

**MK-801:** Furthermore, in a study aiming to elucidate the mechanism underlying MK-801 induced apoptosis...
in the developing brain, reduced mRNA expression has been recorded not only for BDNF but also for some more neurotrophic factors (Hansen et al., 2004). Interestingly, the downregulation in cortical gene transcription of BDNF persists for up to 30 h after MK-801 treatment. The same has been noted for glial derived neurotrophic factor (GDNF), whereas mRNA levels of neurotrophin-3 have been found to be similar to control levels (Hansen et al., 2004), meaning that in the mechanism of apoptosis in developing brain induced by NMDA receptor antagonists BDNF might not be the only neurotrophic factor involved (Heck et al., 2018).

**Ethanol:** Interestingly, in a study assessing the effects of ethanol in developing brain, increase of BDNF levels has been recorded in rats and at the same time approximately equal numbers of pro-apoptotic and pro-survival changes have been produced (Heaton et al., 2003). Furthermore, most of the pro-apoptotic alterations occur rapidly following termination of ethanol treatment, which has been considered a critical period for initiation of apoptosis. At this specific time point, increased BDNF levels have been measured (Heaton et al., 2003). Several studies addressing apoptosis mainly in the developing cerebral cortex have shown that more mechanism besides neurotrophic factors may be involved. Cytokines, as well as neurotransmitters can potentially activate a number of intracellular proteins that execute cell death (Henderson, 1996; Kroemer et al., 2009), meaning that further branches to this AOP might be added in the future.

### Quantitative Understanding of the Linkage

*Is it known how much change in the first event is needed to impact the second? Are there known modulators of the response-response relationships? Are there models or extrapolation approaches that help describe those relationships?*

### Evidence Supporting Taxonomic Applicability

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<td>The survival and antiapoptotic role of BDNF has been investigated not only in rodents but also in developing chicken neurons (Hallbook et al., 1995; Frade et al., 1997; Reinprecht et al., 1998). In invertebrates, only recently a protein with possible neurotrophic role has been identified but its influence and function in neuronal cell death of developing neurons has not been investigated yet (Zhe et al., 2008).</td>
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### References


Presynaptic release of glutamate, Reduced Indirectly Leads to Synaptogenesis, Decreased

How Does This Key Event Relationship Work

The presynaptic release of glutamate causes activation of NMDA receptors and initiates synaptogenesis through activation of downstream signalling pathways required for synapse formation (reviewed in Ghiani et al., 2007). Lack or reduced release of glutamate affects the transcription and translation of molecules required in synaptogenesis (reviewed in Ghiani et al., 2007).

Weight of Evidence

Biological Plausibility
The NMDA receptor activation by glutamate during development increases calcium influx, which acts as a secondary signal. Eventually, immediate early genes (IEG) activation is triggered by transcription factors and the proteins required for synapse formation are produced (reviewed in Ghiani et al., 2007).

Glutamate released from entorhinal cortex neurons has been shown to promote synaptogenesis in developing targeted hippocampal neurons (Mattson et al., 1988). Similarly, glutamate has been found to regulate synaptogenesis in the developing visual system of frogs (Cline and Constantine-Paton, 1990).

The ratio of synaptic NR2B over NR2A NMDAR subunits controls spine motility and synaptogenesis, and it has been suggested a structural role for the intracellular C terminus of NR2 in recruiting the signaling and scaffolding molecules necessary for proper synaptogenesis (Gambrill and Barria, 2011).

**Empirical Support for Linkage**

*Include consideration of temporal concordance here*

**Ethanol:** After ethanol (4 g/kg bw) oral administration to guinea pigs for one hour, hippocampal slices showed persistent decrease of stimulated glutamate release in the fetus but had no effect in adults (Reynolds and Brien, 1994). Further to acute in vivo ethanol treatment, in vitro ethanol (48 mM) exposure also decreased stimulated glutamate release only in the hippocampus of the immature fetus (Reynolds and Brien, 1994).

**Uncertainties or Inconsistencies**

**Ethanol:** In vivo exposure to elevated ethanol doses (322-395.6 mg/dL) during the third trimester-equivalent period impaired synaptic plasticity in the developing CA1 hippocampus of rat pups, without affecting glutamate release after being measured by paired-pulse plasticity of AMPAR-mediated EPSPs frequency (Puglia and Valenzuela, 2010).

The objective of another study was to test the hypothesis that acute in vivo ethanol exposure alters basal glutamate release in the fetal cerebral cortex. The experimental approach involved measuring fetal cortical extracellular glutamate concentration using the technique of in vivo microdialysis. Near-term fetal sheep were chronically instrumented with a microdialysis probe and it was observed that ethanol causes increase of extracellular glutamate levels in the fetal cerebral cortex in a dose-dependent manner (Reynolds et al., 1995).

**Quantitative Understanding of the Linkage**

*Is it known how much change in the first event is needed to impact the second? Are there known modulators of the response-response relationships? Are there models or extrapolation approaches that help describe those relationships?*

**Evidence Supporting Taxonomic Applicability**

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As mentioned above (Empirical Support for linkage) the effects of ethanol on glutamate release has been studied in hippocampus slices of guinea pigs (Reynolds and Brien, 1994) and in sheep cortex (Reynolds et al., 1995), however, contradictory results have been described. Administration of ethanol in guinea pig caused decreased stimulated glutamate release, whereas in sheep caused dose-dependent increase in extracellular glutamate levels.


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**Overall Assessment of the AOP**

The aim of the present AOP is to construct a linear pathway that captures the KEs and KERs that occur after binding of antagonist to NMDA receptor in neurons during development in hippocampus and cortex. All KEs of the AOP are characterised by STRONG essentiality for the AO (learning and memory impairment). Similarly, the biological plausibility in the majority of KERs is rated STRONG as there is extensive mechanistic understanding. However, the empirical support for the present KERs cannot be rated high as in most occasions the KEup and KEdown of a KER have not been investigated simultaneously under the same experimental protocol.
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<td>Directly Leads to</td>
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<td>Indirectly Leads to</td>
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</tbody>
</table>

The table provides a summary of the biological plausibility and empirical support for each KER described in this AOP based on "Annex 1: Guidance for assessing relative level of confidence in the overall AOP based on rank ordered elements" found in User's Handbook.

More information about the evidence that support these KERs and the relevant literature can be found in each KER description.

The main reason for the overall scoring is that for the majority of KERs, the KEup and KEdown have not been investigated simultaneously in the same study.
influx, Decreased Calcium influx, Decreased Indirectly Leads to Release of BDNF, Reduced Release of BDNF, Reduced Indirectly Leads to Dendritic morphology, Aberrant Release of BDNF, Reduced Indirectly Leads to Cell death, N/A Release of BDNF, Reduced Indirectly Leads to Presynaptic release of glutamate, Reduced Cell death, N/A Indirectly Leads to Synaptogenesis, Decreased Dendritic morphology, Aberrant Indirectly Leads to Synaptogenesis, Decreased Presynaptic release of glutamate, Reduced Indirectly Leads to Synaptogenesis, Decreased Synaptogenesis, Decreased Directly Leads to Neuronal network function, Decreased Neuronal network function, Decreased Indirectly Leads to Learning and Scientific understanding is not completely investigated Yes Not investigated Limited conflicting data
Essentiality of the Key Events

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<th>Key Event</th>
<th>Support for Essentiality</th>
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<tr>
<td>NMDARs, Inhibition</td>
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<td>Release of BDNF, Reduced</td>
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<tr>
<td>Presynaptic release of glutamate, Reduced</td>
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<tr>
<td>Cell death, N/A</td>
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</tr>
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</table>

1) Essentiality of the MIE: binding of antagonist to NMDAR in neurons during synaptogenesis in hippocampus and cortex

The MIE is defined and described above as the binding of antagonist to NMDA receptor in neurons during development in hippocampus and cortex (the critical brain structures for learning and memory formation). Activation of NMDA receptors results in long-term potentiation (LTP), which is related to increased synaptic strength and memory formation in the hippocampus (Johnston et al., 2009). LTP induced by activation of NMDA receptors has been found to be elevated in the developing rodent brain compared to the mature brain, partially due to "developmental switch" of the NMDAR 2A and 2B subunits (Johnston et al., 2009).

Essentiality of MIE (binding of antagonist to NMDAR in neurons during synaptogenesis in hippocampus and cortex) for AO (Impairment of learning and memory) is STRONG: It is well documented that learning and memory processes rely on physiological functioning of NMDA receptors. The essentiality of the MIE has been demonstrated in both animal and human studies investigating NMDA itself, NMDA receptors antagonists and mutant mice lacking NMDA receptor subunits (reviewed in Rezvani, 2006 and Granger et al., 2011). NMDA systemically administered in rats, has been shown to potentiate cognitive functions (Rezvani, 2006). There are various studies dealing with specific NMDA receptor subunit gene knock-out leading to a variety of phenotypes. Depending on the endogenous levels of NMDAR subunits, the pattern of their expression and their importance in developmental processes, the loss of a subunit may lead from early embryonic lethality, to mild neurobehavioral impairment up to neuronal disorders that manifest learning and memory deficits (reviewed in Rezvani, 2006 and Granger et al., 2011). Mutant mice lacking NR1 gene have shown perinatal lethality, whereas transgenic mice lacking NR1 subunit in the CA1 region of the hippocampus show both defective LTP and severe deficits in both spatial and nonspatial learning (Shimizu et al., 2000; Tsien et al., 1996). A similar impairment of LTP, long-term depression (LTD), and spatial memory has been seen with CA1-specific NR2B deletion (Brigman et al. 2010). However, LTP has been normal in postnatal forebrain knock-out of NR2A in mice, even though spatial memory has been impaired, probably because of the severe reduction observed in overall excitatory transmission (Shimshek et al., 2006), while the inactivation of the same gene has led to reduced hippocampal LTP and spatial learning (Sakimura et al., 1995). Furthermore, a NR2B transgenic (Tg) line of mice has been developed, in which the NMDA-receptor function has been increased, showing both larger LTP in the hippocampus and superior learning and memory (Tang et al., 1999). Finally, depletion of both NR2A and NR2B in single neurons has shown alteration in synaptic development (Gray et al., 2011). Interestingly, during development, especially
during postnatal days (PND) 7-14 in rodents, the central nervous system (CNS) exhibits increased susceptibility to toxic insults that affect NMDA receptors (Haberny et al., 2002). This increased susceptibility has been suggested to be related to the elevated expression of specific NMDA receptor subunits (Miyamoto et al., 2001). Because of the critical role of the NMDA receptor system in brain development, the exposure to antagonists of NMDA receptors can have long-lasting and severe effects (Behar et al., 1999). NMDA-receptor antagonists such as MK-801, ketamine, phencyclidine (PCP) and 2-amino-5-phosphonopentanoate (AP5 or APV) have been extensively used to study the role of NMDA in learning and memory in developing organisms. Both acute and subchronic administration of NMDA-receptor antagonists in several laboratory animals has been shown to impair performance on tasks that seem to depend upon hippocampal functions (reviewed in Rezvani, 2006; Haberny et al., 2002). The developmental neurotoxicity of several agents, including methylmercury, lead, and ethanol is also thought to result from interaction of these substances with the NMDA receptor system (Guilarte, 1997; Guilarte and McGlothran, 1998; Ikonomidou et al., 2000; Kumari and Ticku, 1998; Miyamoto et al., 2001).

Essentiality of MIE (binding of antagonist to NMDAR in neurons during synaptogenesis in hippocampus and cortex for KE (aberrant dendritic morphology) is MODERATE: The cortex-restricted knockout of NR1 causes refinement in dendritic arborisation in cortex and loss of patterning (Iwasato et al., 2000; Lee et al., 2005). Similar alteration in dendritic arbor has also been identified after depletion of both NR2A and NR2B subunits in isolated neurons (Espinosa et al., 2009). Blockade of NMDA receptor with APV has shown decrease of dendritic growth rate in some in vivo experimental approaches (Rajan et al., 1998; 1999). However, other studies have reported increase in dendritic spine number and dendritic branching after chronic APV-treatment both in vivo and in vitro (Rocha and Sur, 1995; McAllister et al., 1996). This discrepancy is possibly attributed to the different developmental expression of NMDA receptor subunits that triggers distinct intracellular signaling pathways linking NMDAR function to different morphological findings.

Essentiality of MIE (binding of antagonist to NMDAR in neurons during synaptogenesis in hippocampus and cortex) for KE (cell death) is STRONG: The essential role of NMDA receptors in survival during early cortical development has been pointed out both in in vitro (Hwang et al., 1999; Yoon et al., 2003) and in vivo rodent studies (Ikonomidou et al., 1999; Tome et al., 2006). NMDA receptor deficient mice have revealed the importance of this receptor for neuronal survival during development as an approximately 2-fold increase in developmental cell death has been observed in these transgenic mice, which was caspase-3 and Bax dependent (Adams et al., 2004; Rivero Vaccari et al., 2006).

Essentiality of MIE for KE (decreased neuronal network function) is STRONG: The NMDA receptor is associated with circuit formation and function at the developmental stage of an organism as a number of antagonists of this receptor importantly disrupt the neuronal circuit (Simon et al., 1992). Hence, the nature of evidence for the essentiality of the MIE is High (Strong).

2) Essentiality of the KE (Inhibition of NMDA receptors)

Essentiality of KE (Inhibition of NMDA receptors) for AO (Impairment of learning and memory) is STRONG: The noncompetitive antagonist MK-801 has been shown to induce dose-dependent impairment of learning and memory (Wong et al., 1986) and data on rodent models has been recently reviewed in van der Staay et al. 2011. Learning impairments induced by NMDA receptor blockade using MK-801 have also been reported in nonhuman primates (Ogura and Aigner, 1993). Moreover there are human studies demonstrating that NMDA-receptor inhibition impairs learning and memory processes (reviewed in Rezvani, 2006).

3) Essentiality of the KE (Decreased Calcium influx)

Essentiality of KE2 (Decreased Calcium influx) for AO (Impairment of learning and memory) is STRONG: In the nervous system, many intracellular responses to modified Ca2+ levels are mediated by calcium/calmodulin-regulated protein kinases (CaMKs), a family of protein kinases that are initially modulated by binding of Ca2+ to CaM and subsequently by protein phosphorylation (Wayman et al., 2008). Multifunctional CaMKs, such as CaMKII and members of CaMK cascade (CaMKK, CaMKI and CaMKIV) are highly abundant in CNS and regulate different protein substrates (Soderling, 1999). Mice with a mutation in
the alpha- CaMKII that is abundantly found in the hippocampus have shown spatial learning impairments, whereas some types of non-spatial learning processes have not been affected (Silva et al., 1992).

4) Essentiality of KE (Decreased levels of BDNF)

**Essentiality of KE (Decreased levels of BDNF) for AO (Impairment of learning and memory)** is STRONG: BDNF serves essential functions in the brain development and more specific in synaptic plasticity (Poo, 2001) and is crucial for learning and memory processes (Lu et al., 2008). The action of BDNF signaling on synapses happens within seconds of its release (Kovalchuk et al., 2004) and strengthens LTP processes, a cellular model for learning and memory, via sustained TrkB activation as a result of elevated transcription of BDNF (Kang and Schuman, 1996; Nagappan and Lu, 2005). This positive transcriptional feedback happens through TrkB-mediated CREB activation and increases gene transcription of BDNF (Lu et al., 2008). Furthermore, there are experimental evidence showing that loss of BDNF through transgenic models or pharmacological manipulation leads to impaired LTP (Patterson et al., 1996; Monteggia et al., 2004) and decreased learning and memory (Lu et al., 2008). The important role for BDNF in LTP and learning and memory is suggested from numerous studies in rodents. Hippocampal LTP is impaired in mice lacking BDNF in their neurons, and BDNF enhances LTP in the hippocampus and visual cortex (reviewed in Mattson, 2008). BDNF can also be released from neurons during LTP and possibly recycled and used for LTP maintenance. In learning and memory enhancement studies, it has been found that dietary energy restriction (which enhances synaptic plasticity) increases the production of BDNF and glial cells derived neurotrophic factor (reviewed in Mattson, 2008). In humans, a common single-nucleotide polymorphism in the Bdnf gene results in poor performance on memory tasks and may contribute to the pathogenesis of depression and anxiety disorders (reviewed in Cohen and Greenberg, 2008). Similarly, the transgenic mice with such mutation display defects in learning and memory tasks as well as anxiety-related behaviours (reviewed in Cohen and Greenberg, 2008). BDNF has also been shown to play pivotal role in a variety of learning paradigms in a variety of animal models such as mice, monkeys, zebra finches and chicks (reviewed in Tyler et al., 2002).

5) Essentiality of KE (Cell death)

**Essentiality of KE (Cell death) for AO (Impairment of learning and memory)** is STRONG: Several experimental studies dealing with postnatal administration of NMDA receptor antagonists such as MK-801, ketamine or ethanol have shown a devastating cell apoptotic degeneration in several brain regions of animals models, resulting in learning deficits (reviewed in Fredriksson and Archer, 2004; Creeley and Olney, 2013). The apoptosis induced in developing brain after exposure to NMDA receptor antagonists is not reversible although the developing brain has plasticity properties that may allow to a certain degree to compensate for neuronal losses. This severe bilaterally symmetrical neuronal losses in both hemispheres that occurs by treatment with NMDA receptor antagonists leads to neurobehavioral disorders including learning and memory deficits (Creeley and Olney, 2013).

6) Essentiality of the KE (Decreased presynaptic release of glutamate)

**Essentiality of KE (Decreased presynaptic release of glutamate) for AO (Impairment of learning and memory)** is STRONG: Riedel et al. 2003 have reviewed data available that is related to the understanding of the role of glutamate and its different receptor subtypes in learning and memory, focusing mainly in psychopharmacological in vivo studies conducted in rodents and primates. Furthermore, this review has included literature on long-term potentiation (LTP) and long-term depression (LTD), the most commonly used models for studying the cellular mechanisms underlying memory formation in relation to glutamate rather than exploring relevant mechanistic data. Classical conditioning of a tone-shock association (commonly used to study learning and memory) causes a lasting increase in glutamate release in dentate gyrus synaptosomes, whereas blockade of NMDA receptors during learning prevents conditioning and the change in glutamate release (Redini-Del Negro and Laroche, 1993). It is worth mentioning that there are two types of LTP, the NMDA receptor-dependent and the NMDA receptor-independent. The later type of LTP is induced presynaptically and strongly activates presynaptic Ca2+ channels, which results in an increase in cAMP and activation of protein kinase A that is believed to be involved in the long-lasting enhancement of glutamate release from the presynaptic terminal. This type of LTP has been observed at mossy fiber-CA3 synapses in the hippocampus or at parallel fiber-Purkinje cell synapses in the cerebellum.
7) Essentiality of the KE (Aberrant dendritic morphology)

**Essentiality of KE (Aberrant dendritic morphology) for AO (Impairment of learning and memory)** is *STRONG*: Spine morphology is considered to be an important morphological unit for establishing learning and memory (Sekino et al., 2007). As dendrites are the postsynaptic site of most synaptic contacts, dendritic development determines the number and pattern of synapses received by each neuron (McAllistair, 2000). Defects induced in dendritic growth are often leading to severe neurodevelopmental disorders such as mental retardation (Purpura, 1975). Thus, the proper growth and arborization of dendrites are crucial for proper functioning of the nervous system. Changes in spine formation have been found to be involved in impairment of learning and memory in live animals (Yang et al. 2009; Roberts et al. 2010). Electrical activity-dependent changes in the number as well as in the size and shape of dendritic spines have been strongly related to some forms of learning (reviewed in Holtmaat and Svoboda, 2009). In mouse, motor cortex learning leads to dendritic spine remodeling associated with the degree of behavioral improvement suggesting a crucial role for structural plasticity during memory formation (Yang et al., 2009 and Fu et al., 2012). Furthermore, accumulating evidence indicates that experience-dependent plasticity of specific circuits in the somatosensory and visual cortex involves structural changes at dendritic spines (Holtmaat and Svoboda, 2009).

8) Essentiality of the KE (Decreased synaptogenesis)

**Essentiality of KE (Decreased synaptogenesis) for AO (Impairment of learning and memory)** is *STRONG*: Learning and memory result from plastic events that modify the way neurons communicate with each other (Bear, 1996). Plastic events are considered changes in the structure, distribution and number of synapses and it has been suggested that morphological events like these underlie memory formation (Rusakov et al., 1997; Woolf, 1998; Klintsova and Greenough, 1999). In mutant mice lacking PSD-95, it has been recorded increase of NMDA-dependent LTP, at different frequencies of synaptic stimulation that cause severe impaired spatial learning, without thought affecting the synaptic NMDA receptor currents, subunit expression, localization and synaptic morphology (Migaud et al., 1998). Furthermore, recent genetic screening in human subjects and neurobehavioural studies in transgenic mice have suggested that loss of synaptophysin plays important role in mental retardation and/or learning deficits (Schmitt et al., 2009; Tarpey et al., 2009).

### Quantitative Considerations

<table>
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<th>Description</th>
<th>Triggers</th>
<th>Quantitative Understanding</th>
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<tr>
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Synaptogenesis, Decreased to Decreased Neuronal network function, Decreased Indirectly Leads to Learning and memory, Impairment Release of BDNF, Reduced Indirectly Leads to Cell death, N/A Presynaptic release of glutamate, Reduced Indirectly Leads to Synaptogenesis, Decreased

A quantitative structure activity relationship (QSAR) model has been developed based on various molecular parameters that have been calculated for a series of competitive NMDA antagonists with known activity values and these parameters have been applied to make a regression analysis which provides a model that relates the computationally calculated parameters to experimentally determined activity values (Korkut and Varnali, 2003).

Recently, they have established a QSAR model for non-competitive antagonists of NMDA receptor, by studying a series of 48 substituted MK-801 derivatives (Chtitaa et al., 2015). In this paper, a quantitative model has been proposed, and then there has been an attempt to interpret the activity of the compounds relying on the multivariate statistical analyses. By this approach, they have been able to predict the inhibitory activity of a set of new designed compounds (Chtitaa et al., 2015).

2D- and 3D-QSAR models have also been developed to establish the structural requirements for pyrazine and related derivatives for being NR2B selective NMDA receptor antagonists (Zambre et al., 2015).

**Applicability of the AOP**

<table>
<thead>
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<tr>
<td>Female</td>
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**Life Stage Applicability:** This AOP is applicable only for specific period of brain development that is the time of synaptogenesis. This vulnerable period of synaptogenesis appears to happen in different developmental stages across species. For example, in rodents primarily synaptogenesis occurs during the first two weeks after birth. For rhesus monkeys, this period ranges from approximately 115-day gestation up to PND 60. In humans, it starts from the third trimester of pregnancy and continues 2-3 years following birth (Bai et al., 2013). Furthermore, synaptogenesis does not happen in a uniform way in all brain regions and there are important differences between the times of appearance of the main two types of synapses (reviewed in Erecinska et al., 2004). For example, in rat hippocampus excitatory synapses are well established or fully mature within the two first postnatal weeks, whereas inhibitory synapses cannot be found prior to PND 18, after which it increases steadily to reach adult levels at PND 28. In addition, in rat neostriatal neurons the excitatory responses to both cortical and thalamic stimuli can be observed by PND 6, but the long-lasting hyperpolarization and late depolarization is never seen before PND 12.
**Taxonomic Applicability:** The data used to support the KERs in this AOP derives from experimental studies conducted in primates, rats and mice or cell cultures of similar origin as well as from human epidemiological studies. The majority of the KEs in this AOP seem to be highly conserved across species. It remains to be proved if these KERs of the present AOP are also applicable for other species rather than human, primates, rats and mice.

**Sex Applicability:** The majority of the studies addressing the KEs and KERs of this AOP were carried out mainly in male laboratory animals. Few studies are available in females and some of them compare the effects between females and males. It appears that this AOP is applicable for both females and males.

**Considerations for Potential Applications of the AOP (optional)**

Exposure to xenobiotics can potentially affect the nervous system resulting in neurobehavioral alterations and/or neurological clinical symptoms. To assess the neurotoxic properties of compounds, current testing largely relies on neurobehavioural tests in laboratory animals, histopathological analysis, neurochemical and occasionally electrophysiological observations.

Throughout the years, a significant number of methods have been developed to assess neurobehaviour in laboratory animals and a comprehensive summary of them can be found in OECD Series on testing and assessment, number 20, Guidance Document for Neurotoxicity Testing (2004). Learning and memory is an important endpoint and a wide variety of tests to assess chemical effects on cognitive functions is available and used for the study of neurotoxicity in adult and young laboratory animals. Some of these tests that allow the appreciation of cognitive function in laboratory animals are: habituation, ethologically based anxiety tests (elevated plus maze test, black and white box test, social interaction test), conditioned taste aversion (CTA), active avoidance, passive avoidance, spatial mazes (Morris water maze, Biel water maze, T-maze), conditional discrimination (simple discrimination, matching to sample), delayed discrimination (delayed matching-to-sample, delayed alternation) and eye-blink conditioning.

During the investigation of developmental neurotoxicity induced by chemicals, learning and memory, through the above mentioned tests, is evaluated in OECD Test Guideline for Developmental Neurotoxicity (426), OECD Test Guideline for Combined Repeated Dose Toxicity Study with Reproduction/Developmental Toxicity Screening Test (422) and OECD Test Guideline for Extended One-Generation Reproductive Toxicity Study (443).

**References**


cholinergic drugs. J Pharmacol Exp Ther. 266: 60-64.


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