Excitatory amino acid neurotoxicity—a broader horizon for cerebral protection?

Substantial experimental evidence accumulated over the past decade indicates that an endogenous mechanism of toxicity, which results in a selective neuronal lesion more severe in dendrites than in axons (with sparing of glia), may be significant in the process of neuronal degeneration seen after brief neurological insults. Much of the evidence has been pharmacological with protection from the development of morphological neuronal injury by the use of antagonists acting specifically at the glutamate family of neuronal receptors. 1 2 The powerful protective effects of these antagonists are potentially of great clinical significance, and could have a role in the treatment of cerebral ischaemia, profound hypoglycaemia, and status epilepticus. In the immature brain, however, the potential usefulness of these agents may be precluded by their adverse effects on behaviour and brain development, because of the importance of excitatory neurotransmission in neural outgrowth, plasticity, and cell interaction. 3 Despite such a potential limitation for treatment, an understanding of the mechanism of excitatory amino acid neurotoxicity takes us significantly closer to conceptualising the process of neuronal death induced by acute insults, which must be central to any cerebral protective measures.

The 'excitotoxic' hypothesis

In its simplest form the so called 'excitotoxic' hypothesis of neuronal injury proposes that glutamate or other related endogenous excitatory neurotransmitters become toxic in their interaction with glutamate receptors, resulting in a cascade of intracellular events that culminate in neuronal death. 4-8 What makes an endogenous transmitter (which is normally released into and cleared from the synaptic cleft) become a neurotoxin is still a matter of much speculation, but these events may be attributable to:

(1) an excessive build up of extracellular glutamate because of abnormal neurotransmitter release, or abnormal uptake by neurons and glia;
(2) abnormal glutamate receptor activation because of abnormal postsynaptic sensitivity;
(3) abnormal induction and amplification of intracellular cytotoxic events.

Glutamate receptors

There is evidence for multiple subtypes of excitatory neurotransmitter receptors, which are commonly called glutamate receptors, as glutamate is the most widely accepted endogenous excitatory transmitter. These receptors can be divided into three groups. There are two classes which possess an intrinsic ion channel, called ionotropic receptors. These are the N-methyl-D-aspartate (NMDA) receptor and the non-NMDA, kainate/quisqualate receptor, which are presumed to be large, multisubunit integral membrane protein complexes, which possess receptor binding sites with a central ion channel. The third class of receptor is the metabotropic glutamate receptor, which instead of possessing an intrinsic ion channel exerts its action through second messenger systems by activation of a G-protein.

An understanding of the extent to which these glutamate receptor subtypes contribute to the neuronal injury induced by acute cerebral insults has been facilitated by experimental
studies that have evaluated the relative neurotoxicity produced by intracerebral injection of specific receptor agonists, NMDA, quisqualate, and kainate. The results indicate major ontogenic alterations in the excitatory amino acid pathways, with different brain regions and neuronal types exhibiting their own developmental profile of susceptibility to action of the glutamate-receptor agonists. Comparing NMDA with non-NMDA agonists, on an equimolar basis, the greater degree of injury in the immature brain is produced by NMDA, and in the developed brain by non-NMDA agonists. Furthermore, in the immature brain, there appears to be a transient overproduction of glutamate-receptor synaptic terminals, which may not only account for periods of heightened synaptic plasticity and consolidation of synaptic connections, but also provide a molecular substrate for increased susceptibility to particular insults.

Gluatamate-receptor agonists

Of all the major classes of glutamate receptor, the best antagonists were first found for the NMDA receptor subtype. Consequently, most is known about the potential role of this receptor in mediating the neuronal injury of acute brain insults. It should, however, be remembered that many of the antagonists are not ‘pure’ in their differentiation between the receptor subtypes, particularly at higher concentrations, and so any beneficial effect may not be solely mediated by antagonism of a specific receptor subtype. In addition to the obvious interest in the pharmacologically ‘designed’ antagonists, which have for the time being only proved to be useful experimental tools, there has been some recent experimental and clinical interest in those agents already in clinical usage that have a known antagonistic effect on the NMDA receptor channel complex. These drugs are considerably less potent than the experimental agents, and include ketamine and dextromethorphan (phencyclidine site ligands) and magnesium.

Excitotoxicity and acute cerebral insults

Cerebral ischaemia, profound hypoglycaemia, and status epilepticus, result in a complex interaction between changes in cerebral energy metabolism, cerebral blood flow, blood-brain barrier function, neuronal depolarisation, and glial-neuronal interaction. As already mentioned, the main evidence that an excitotoxic mechanism provides a crucial link in the sequence of events leading to neuronal loss after these insults is pharmacological. Several studies have demonstrated that glutamate-receptor blockade can prevent neuronal injury induced by a variety of acute insults in both the mature and immature brain. Although the most well characterised insult is ischaemia, interaction at a single subtype of glutamate receptors cannot explain the findings in all the experimental models of hypoxic-ischaemic insult. As an example, it has been observed that there is a relative lack of neuroprotection with NMDA antagonists in somatosensory-evoked potential models of cerebral ischaemia. These findings would be consistent with a model of glutamate-receptor mediated toxicity that takes into account the ontogenic profiles of neuronal injury due to exogenous glutamate-receptor agonists; ischaemia induced injury would be expected to be predominantly mediated by NMDA receptors in the immature brain and by non-NMDA receptors in the adult brain.

An excitotoxic process has also been implicated in profound hypoglycaemia and status epilepticus. In hypoglycaemia it seems likely that neurotoxicity is mediated by the NMDA receptor with aspartate as the endogenous agonist. That glutamate-receptor antagonists are also potent anticonvulsants suggests involvement of these receptors in the mechanism of toxicity induced by status epilepticus. This speculative hypothesis, however, would be impossible to test with receptor antagonists, as blockade of glutamatergic neurotransmission for neuroprotection would also stop the epileptic activity. There has therefore been much effort in trying to demonstrate an abnormal accumulation of excitatory amino acids in the extracellular space during the period of insult.

The wider horizon for cerebral protection

This annotation has so far presented the barest model of excitatory amino acid neurotoxicity induced by acute cerebral insults, with the NMDA receptor providing a central role in the process of neuronal injury. Having illustrated some of the pharmacological evidence supporting the recent interest in the NMDA receptor, it is also worth noting that this receptor has unusual physiological properties. Because the receptor ion channel is blocked by normal concentrations of extracellular magnesium, the NMDA receptor is difficult to activate. Only as the membrane becomes depolarised and the magnesium blockade is lifted, can the receptor be activated. The ion channel then allows calcium to enter the cell. The resultant rise in cytosolic calcium is considered to be the initiating step in the cascade of intracellular events that culminate in neuronal death.

The excitotoxic mechanism of neuronal injury has gained a broader interest beyond acute cerebral insults because of recent observations that indicate that neurons can be more susceptible to the excitotoxic effects of glutamate and other related excitatory amino acids by manipulations of cellular energy metabolism and membrane potential. It has been suggested that impairment of cellular energy metabolism leads to failure of the Na+/K+ ATPase that maintains the normal membrane potential, resulting in membrane depolarisation, reduced magnesium blockade, and easier activation of NMDA receptors. The process of neuronal death could be initiated by impairment of cellular metabolism or membrane potential with an excitotoxic process being the final common pathway of neuronal death: an attractive hypothesis for chronic neurodegenerative disorders such as Huntington’s disease and Parkinson’s disease.

As our understanding of neuronal injury produced by excitatory amino acids has grown, experimenters have used glutamate receptor antagonists as pharmacological tools to explore the role of endogenously triggered excitotoxicity in the complex pathophysiology of brain injury following acute cerebral insults. Although the knowledge gained from such experimental models and tissue culture may ultimately be translated into practical clinical strategies for the treatment of acute brain injury, there is potentially a far wider application in chronic neurodegenerative disease, and some inherited metabolic disorders.

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