The use of zinc in medicinal skin cream was mentioned in Egyptian papyri from 2000 BC (for example, the Smith Papyrus), and zinc has apparently been used fairly steadily throughout Roman and modern times (for example, as the American lotion named for its zinc ore, ‘Calamine’). It is, therefore, somewhat ironic that zinc is a relatively late addition to the pantheon of signal ions in biology and medicine. However, the number of biological functions, health implications and pharmacological targets that are emerging for zinc indicate that it might turn out to be ‘the calcium of the twenty-first century’. Here neurobiological roles of endogenous zinc is summarized. J Clin Neurol 1(2):121-133, 2005

Key Words : Ischemia, Stroke, Alzheimer, Amyotrophic lateral sclerosis

INTRODUCTION AND OVERVIEW

In medicine and biology, zinc has several connotations. It is an essential micronutrient, a component of enzymes and other proteins, and a toxic pollutant as well.

To neuroscientists, zinc is also an ionic signal, \( \text{Zn}^{2+} \) enters cells through gated channels, and moves among various organelles and storage depots within cells, modulating protein function by binding to and detaching from zinc-dependent proteins. Like calcium, excess free zinc in tissue is toxic.

\( \text{Zn}^{2+} \) is selectively stored in, and released from, the presynaptic vesicles of a specific type of neurons in the mammalian brain. These zinc-releasing neurons also release glutamate, so the term “gluzinergic” has been proposed to describe them. By and large, the gluzinergic neurons all have their cell bodies in either the cerebral cortex or in the limbic structures of the forebrain. Thus the gluzinergic neuronal system comprises a vast cortical-limbic associational network that unites limbic and cerebrocortical functions. The gluzinergic message is the exclusive voice of the cerebrocortical and limbic systems.

In the fifty years since the first identification of chelatable zinc in the brain, a broad outline of the function of gluzinergic neurons has slowly come into focus. First, zinc appears to modulate the overall excitability of the brain via effects on glutamate, and probably GABA receptors. Clinical links to epileptic disorders have been a major theme in the literature of zinc neurobiology. Secondly, perhaps because it is preferentially located in cerebrocortical associative pathways, zinc may be important in synaptic plasticity.

However, excess free zinc ion is toxic. Indeed, a major portion of current interest in the neurobiology of zinc is driven by the idea that the zinc ion is a causal contributor in both the acute brain injury of stroke, head trauma, seizures, or cardiac arrest, and the slow and
relentless brain injury of the neurodegenerative disorders such as Alzheimer’s disease (AD) and possibly amytrophic lateral sclerosis (ALS). In the present paper, current evidence that implicates endogenous zinc in pathophysiology of both acute brain damage and degenerative brain diseases is reviewed.

**BASIC NEUROPHYSIOLOGY OF ZINC**

Maske first identified the zinc-containing mossy-fiber terminals. Subsequently, it was found that many of the intrinsic, glutamatergic pathways of the cerebral cortex are comprised of gluzinergic neurons. The “intrinsic” is here emphasized because corticofugal and corticopedal long-axon pathways, though also glutamatergic, are generally devoid of zinc.

Surprisingly little is known about the biological life cycle of zinc in the glutamatergic vesicles of the forebrain. There is a protein, zinc transporter-3 (ZnT-3), that co-localizes with zinc vesicles, and mice lacking the gene for that protein (znt-3 knockouts) show no staining for zinc in their presynaptic terminals. These data indicate that the ZnT-3 protein plays a role in sequestering vesicular zinc, but exactly what that role is remains uncertain.

Like other neurotransmitters packed in vesicles, zinc is released with neuronal activity. A number of groups have found very robust and reliable release of zinc from boutons upon electrical stimulation or simple elapsed time. Most recently, the release of zinc has been elegantly demonstrated on a pulse-by-pulse basis, with each action potential releasing zinc.

The soma and dendrites of mammalian neurons are studded with a variety of zinc-permeable ion channels. These include the NMDA channel, voltage-gated calcium channels, and the calcium-permeable AMPA/Kainate (Ca-A/K) channel. Zinc influx through these channels has been demonstrated.

Because presynaptic terminals release zinc and the postsynaptic soma and dendrites have zinc-permeable channels, it follows that zinc ions will travel from inside a presynaptic neuron to inside a post synaptic neuron (translocate) under favorable conditions. Because both glutamate and depolarization open the zinc-permeable channels, one expects the maximum zinc translocation during intense neuronal activity with depolarization. Much evidence discussed below indicates that such translocation contributes to zinc-induced cell injury in excitotoxicity. There is also evidence that a smaller-volume translocation may occur during normal physiological synaptic signaling, with the translocated zinc perhaps triggering further signal cascades in the postsynaptic neuron.

In addition to the zinc that can be released from presynaptic terminals into the extracellular fluid, it is clear that there is also a pool of zinc stored in perikarya that can be “released” into the cytoplasm. One source of this zinc is the metallothionein family of proteins (MTs), from which zinc can be released at an especially high rate by nitrosylation of the thiol ligands by NO. Of three isoforms of MTs, MT-3 has only been found in brain and testes, whereas the others are more widespread. In brain injury, the absence of MT-3 significantly reduces cell injury in hippocampal field CA1 and the thalamus, implying that zinc released off MT3 can contribute to cell injury. In contrast, in hippocampal field CA3, the absence of MT-3 increases cell death in excitotoxic injury, presumably because the presynaptic release of zinc is so pronounced in CA3, that the postsynaptic MT-3 serves more as a zinc sink than a zinc source.

One of the very first neuronal receptors found sensitive to Zn\(^{2+}\) was the NMDA-type glutamate ionophore, which was shown to be inhibited by Zn\(^{2+}\) in 1987. The sensitivity of the NMDA-type receptor-ionophore is now understood to be mediated by two separate mechanisms, a voltage-independent site on the NR2A subunit that has an IC\(_{50}\) in the single-digit nanomolar range and a less sensitive, voltage-dependent site on the NR2B subunit where ionic current is decreased by low-micromolar concentrations of Zn\(^{2+}\). Another potentially critical aspect of the zinc-NMDA relationship is that prior exposure to zinc apparently causes a delayed increase in the sensitivity of the receptor to agonists. This delayed effect (over hours) is mediated by increased phosphorylation of the NR2A and NR2B subunits, thereby decreasing their sensitivity to the tonic
The second receptor that has been studied intensively for zinc sensitivity is the GABA receptor, which was first shown to be inhibited by Zn\(^{2+}\) in 1987.\(^{41,47}\) Although most of the data concerning the GABA sensitivity to zinc have come from experiments in which exogenous zinc was added to tissue baths, several exemplary experiments have used the blockade (chelation) paradigm to reveal effects of endogenous zinc signals.\(^{48,49}\) Changes in the zinc modulation of GABA receptor have been implicated in the etiology of epilepsy. Mody, Coulter and others\(^{50-52}\) have suggested that the seizure-induced sprouting of zinc-releasing axons into ectopic locations could result in ectopic release of zinc, thus reducing GABA-A receptor-mediated inhibition, and enhancing seizure susceptibility.\(^{51}\) In addition to the sprouting of zinc-releasing axons, there are additional changes in zinc modulation of the GABA receptor that could contribute to the progressive epileptogenesis.\(^{53,54}\)

**ZINC IN ACUTE BRAIN INJURY**

1. Zinc accumulation as a cause of neuronal death

Although zinc lacks redox activity, and was traditionally regarded as relatively non-toxic,\(^{55}\) an increasing body of evidence demonstrates that zinc is in fact a potent killer of neurons and glial cells. As mentioned earlier, the toxicity of free zinc (even 1 µM) in streams and oceans is well known to environmental scientists.\(^{3,56}\)

In 1986, we have demonstrated that brief (15 min) exposure to 300-600 µM zinc results in extensive neuronal death in cortical cell culture.\(^{57}\) Combined with the realization that neurons store up to 300 µM of free zinc in their terminals\(^{58}\) and release that zinc when they are depolarized,\(^{59-61}\) the fact that zinc was cytotoxic suggested the possibility that zinc might play an active role in neuronal injury.

The facts that (i) minutes of exposure to µM zinc kills brain cells in culture, and (ii) zinc is massively released in acute brain injury, suggest that zinc toxicity could contribute to neuronal injury in vivo. Staining of brain sections of ischemia- or epilepsy-subjected animals with a zinc fluorescent dye and acid fuchsin, revealed a striking correlation between zinc accumulation in cell bodies and their death. It was demonstrated that both neuronal death and zinc accumulation in transient cerebral ischemia, were reduced or prevented by an extracellular zinc chelator, CaEDTA.\(^{62}\) Subsequently, the principle of endogenous zinc toxicity as a contributing mechanism has been examined and determined valid in other injury models such as blunt head trauma,\(^{63}\) focal ischemia,\(^{64}\) oxygen-glucose deprivation in vitro\(^{65}\) and glucose deprivation in vivo.\(^{66}\)

Because histochemically-reactive zinc in synaptic vesicles was initially considered the only releasable pool of zinc,\(^{67}\) it was postulated that the zinc that appeared in postsynaptic neuronal somata was likely presynaptic zinc that had been released and “translocated” into the post-synaptic neurons. However, while useful for a time, the “zinc translocation” hypothesis is now recognized as incomplete. First, zinc accumulation in degenerating neurons has always been observed to some extent in areas only lightly innervated by glutamergic fibers. For instance, thalamic neurons are surrounded by terminals that lack vesicle zinc.\(^{11,68}\) Still, these neurons exhibit zinc accumulation following ischemia and seizures.\(^{62,68}\) Second, even in znt3-null mice that lack synaptic zinc, extensive zinc accumulation in degenerating CA1 and thalamic neurons was observed.\(^{69}\) Finally, the fairly recent discovery that extracellular CaEDTA can remove zinc from inside of cells and even presynaptic vesicles\(^{70}\) (presumably by creating extremely steep transmembrane gradients) brought the interpretation of CaEDTA data into direct question. Specifically, blockade by CaEDTA could no longer be accepted as evidence that the zinc had traveled through the extracellular fluids.

Zinc accumulation in degenerating neurons of znt-3-null mice indicates that there are other zinc sources besides synaptic vesicle zinc. One such source is the zinc that can be mobilized off MT-3 (and possibly from mitochondria) discussed above. As mentioned previously, such intracellular zinc release could lead to a somatic release of zinc into the extracellular fluid with subsequent zinc translocation into neighboring cells. The direct role of Nitric oxide in releasing this MT3 pool of zinc during excitotoxicity was recently demonstrated by Wei.\(^{71}\)
2. Zinc-initiated cell death pathways

Regardless of specific sources or routes involved, increased levels of reactive or “free” zinc inside cells is toxic. This toxic effect of zinc was initially puzzling to some because zinc had been considered relatively innocuous metal, and zinc was known to inhibit apoptosis in diverse cell systems.72

Although zinc is not an oxidizer, several lines of evidence have shown that zinc toxicity is mediated largely by oxidative stress. First zinc-induced cell death is accompanied by increased levels of superoxides and lipoperoxides, markers for oxidative injury.73-75 Second, zinc-induced cell death is attenuated by various antioxidative measures.76,77 Third, free radical-generating enzymes such as NADPH oxidase are induced and activated after zinc exposure, and their inhibitors attenuate zinc toxicity.78

With brief exposure to high concentrations of zinc, neurons exhibit signs of necrosis, such as cell body swelling and destruction of intracellular organelles.73 However, in less fulminant zinc toxicity, signs of apoptosis such as DNA fragmentation and caspase activation, are also observed.76,79 The fact that zinc exposure induces apoptosis was puzzling, since depletion of zinc also induces caspase activation and apoptosis.80,81 However, elevated zinc does indeed produce apoptosis, and mechanisms for zinc-triggered apoptosis are now being identified. For example, in zinc-exposed neurons, both p75NTR and p75NTR-associated death executor (NADE), are induced,82 a combination that can induce caspase activation and apoptosis.83 In addition to this pathway, zinc can trigger the release of pro-apoptotic proteins such as cytochrome C and apoptosis inducing factor (AIF) from mitochondria.84 Whether and how much apoptosis contributes to zinc-related acute brain injury is unknown. However, in rat models of ischemia and seizures where the role of zinc as a neurotoxin is likely, p75NTR and NADE are co-induced in neurons that undergo cell death,82,85 strengthening the possible involvement of this apoptogenic cascade in vivo.

Another pivotal factor in zinc toxicity is nitric oxide (NO). NO releases 7 zinc ions from each single MT molecule,86-88 and the brain-specific MT-3 isoform has a considerably lower threshold for zinc release by NO than the other isoforms.89,90 Because inhibition of NO synthase (NOS) dramatically reduces the release of zinc from brain slices,71 and reduces the appearance of zinc staining after hypoglycemic brain injury,66 it is clear that NO release of zinc from MT plays a crucial role in excitotoxic zinc toxicity. NO also rapidly releases zinc from presynaptic terminals,91 thus contributing to cell death via the zinc translocation mechanism. Whereas NO releases zinc, elevated Intracellular zinc also induces and activates neuronal NOS in cultured cortical neurons.92 Thus regardless whether zinc or NO is the initial trigger, a destructive cycle is easily induced. The final pathway to zinc-induced cell necrosis seems to be poly-ADP-ribose polymerase (PARP) activation,92 as in other cases of predominantly necrotic cell death.93 DNA damage induced by oxidative and nitrosative stresses activates PARP, an enzyme that transfers the ADP-ribose moiety from NAD$^+$ to various target proteins. Since up to several hundred moieties are transferred to a molecule of protein, continued activation of PARP results in a drastic depletion of NAD$^+$ and ATP.94 Consistent with the idea that PARP activation is limited only to necrosis type cell death,93 chronic exposure to low concentrations of zinc, which preferentially induces apoptosis,79 is not attenuated by deletion of PARP-1.95

ZINC IN NEURODEGENERATIVE DISEASES

1. Alzheimer’s Disease

Alzheimer’s disease (AD) is characterized by loss of cortical neurons and progressive deterioration of cognitive function, memory, and self-care. The pathological hallmark of AD is marked accumulation of amyloid-β (Aβ) protein, neurofibrillary tangles (NFTs) and neuropil threads in the neocortex.96 Aβ (39-43 amino acid residues, ~4 kDa), is the main constituent of both senile plaques and cerebrovascular amyloid deposits.97,98 The Aβ peptide is produced from the proteolytic cleavage of a much larger transmembrane precursor, the Amyloid Protein Precursor (APP).99 Mutations of APP (on
chromosome 21) within or adjacent to the Aβ domain cause aggressive familial AD, indicating that abnormal Aβ and APP metabolism can give rise to the disease. Since the discovery that Zn\(^{2+}\) precipitates Aβ,\(^{95,100}\) considerable evidence has emerged that free Zn\(^{2+}\) in the extracellular fluid induces amyloid deposition. Aβ1-40 specifically and saturably binds zinc with a 1:1 (zinc:Aβ) stoichiometry. Because zinc concentrations of the extracellular brain milieu are apparently in 1 to 10 nM range, one would expect Aβ1-40 to bind very little zinc under normal conditions. However, events leading to a sustained increase in extracellular zinc levels, such as a transient hypoperfusion, head trauma, or even local paroxysmal neuronal firing\(^{101}\) could easily lead to zinc binding to Aβ.

The zinc binding site was mapped to a stretch of contiguous residues between positions 6-28 of the Aβ sequence, and the histidine at residue 13 plays a critical role in Zn\(^{2+}\) mediated aggregation.\(^{102}\) Occupation of the zinc binding site, which straddles the lysine 16 position of α-secretase cleavage,\(^{103}\) by zinc inhibits β-secretase type cleavage and so may influence the generation of Aβ from APP, and may increase the biological half-life of Aβ by protecting the peptide from proteolytic attack.\(^{90}\) Zinc concentrations above 300 nM rapidly precipitate synthetic human Aβ1-40.\(^{100}\) Importantly, Zn\(^{2+}\)-induced precipitation is completely reversed with chelation treatment.\(^{104}\)

Zinc-induced Aβ precipitation at pH 7.4 is highly specific for zinc; however, Cu\(^{2+}\) and Fe\(^{3+}\) can induce partial aggregation at pH 7.4 which increases substantially under mildly acidic conditions (pH 6.6).\(^{105}\) Raman spectroscopy has recently shown that Zn\(^{2+}\) binds to the N(tau) atom of the histidine imidazole ring and that the peptide aggregates through intermolecular His(N(tau))-Zn\(^{2+}\)-His(N(tau)) bridges.\(^{106}\)

Aβ binds Cu\(^{2+}\) and Zn\(^{2+}\) through selective binding sites. When synthetic Aβ is co-incubated with excess but equal amounts of Cu\(^{2+}\) and Zn\(^{2+}\), ≈1.5 equivalents of each metal ion binds to each mole of peptide. Because the affinity of the Cu\(^{2+}\) binding sites on Aβ is much higher than that of Zn\(^{2+}\) binding sites, the finding that Cu\(^{2+}\) does not compete for all of the available metal binding sites when co-incubated with Zn\(^{2+}\) implies that Aβ possesses separate and selective Cu\(^{2+}\) and Zn\(^{2+}\) binding sites.\(^{107}\) Zn\(^{2+}\), Cu\(^{2+}\) and Fe\(^{2+}\) are markedly enriched in amyloid plaques,\(^{108}\) but only Cu and Zn co-purify with Aβ extracted from post-mortem human brain\(^{109}\) and have been determined by Raman spectroscopy to coordinate with Aβ in plaques.\(^{110}\)

In mouse brain Cu\(^{2+}\) and Fe\(^{2+}\) levels rise with age.\(^{111}\) One idea is that Aβ, which can bind up to 3.5 moles of Cu and Zn per monomer\(^{108}\) becomes hypermetallated (overloaded) with age, and abnormally oxidized while handling Cu\(^{2+}\) physiologically.\(^{18}\) Such a hypothetical abnormal binding of Cu\(^{2+}\) to Aβ would yield two adverse outcomes: (i) toxicity mediated by redox activity, and (ii) oxidative modification of Aβ.

Aβ:Cu\(^{2+}\) complexes are strongly reducing, and generate H\(_2\)O\(_2\) catalytically from biological reducing agents including cholesterol.\(^{109,112,113}\) The redox activity is stronger for human Aβ1-42, than human 1-40 or the rat Aβ peptide, correlating with the toxicity of the peptide in cell culture.\(^{114}\)

Cu\(^{2+}\)-mediated oxidation of Aβ causes damage to histidine and tyrosine side-chains,\(^{115}\) dityrosine cross-linking\(^{116}\) and sulfoxidation of the sole methionine at residue 35.\(^{117}\) This latter methionine is essential for keeping metallated Aβ in its normal (redox-silent) location within lipid membranes.\(^{118,119}\) Therefore, oxidation of the Aβ by Cu\(^{2+}\) may be the first step in liberating Aβ species that can later be precipitated by Zn\(^{2+}\). This may explain why virtually all of the Aβ that deposits in the brain in AD is oxidized.\(^{120}\)

Generation of H\(_2\)O\(_2\) by soluble, but oxidized forms of Aβ,\(^{121}\) may explain the association of brain Aβ accumulation with the severe peroxidative damage that is characteristic of the AD-affected brain.\(^{122}\)

Events leading to a sustained increase in extracellular zinc in combination with oxidative stress, such as stroke, head trauma, cardiac arrest, or epilepsy, would increase the likelihood of soluble Aβ precipitation into plaque and are indeed risk factors for AD.

Zn/Cu chelators reverse Zn/Cu-induced aggregation of synthetic Aβ in vitro\(^{104}\) inhibit Aβ-mediated H\(_2\)O\(_2\) formation,\(^{109,113,123,124}\) and solubilize Aβ from amyloid deposits in post-mortem AD-affected brain tissue.\(^{222}\) Studies of the impact of the genetic ablation of ZnT3
in the Tg2576 mouse model of AD have provided evidence that synaptically-released zinc underlies amyloid pathology in this model. We found that the complete absence of any staining for synaptic vesicle zinc in the knockout mouse was accompanied by a reduction of the cerebral plaque load by approximately 80%.\textsuperscript{125} Interestingly, synaptic zinc levels (as measured by histo-fluorescence for zinc) as well as plaque loads increased to a greater degree with age in female mice than in male, suggesting the influence of sex hormones on synaptic zinc levels.\textsuperscript{125} Preliminary evidence suggests that estrogen may reduce the level of synaptic vesicle zinc, perhaps by modulating the expression level of the adaptor protein 3 (AP3) complex, which is required for the correct insertion of ZnT3 into vesicular membrane.\textsuperscript{126}

Cerebral amyloid angiopathy (CAA) is also decreased in ZnT-3 knock out/Tg2576 compared to Tg2576 controls.\textsuperscript{2}

2. Amyotrophic Lateral Sclerosis (ALS)

Two abnormalities of zinc-metalloproteins have implicated zinc in the pathophysiology of ALS (Lou Gehrig’s disease). First, it is the well-established fact that familial form of ALS in man is accompanied by mutations in the metalloenzyme Cu- Zn-superoxide dismutase (SOD).\textsuperscript{127,128} Mutations in SOD are also associated with ALS-resembling spinal motor defects in mice, with different mutants having different amounts of wild-type enzymatic activity, ranging from 0% (e.g. H46R and G85R) to 100% (e.g. G37R). SOD1 knockout mice do not develop the ALS phenotype,\textsuperscript{129} and the age of onset and duration of disease in ALS transgenic mice is unaffected by levels of wild-type SOD1 activity.\textsuperscript{129} Thus, the toxicity of mutant SOD1 (mSOD1) is a gain-of-function.

Several gain-of-function redox reactions have been proposed for mSOD1, and at least two currently appear plausible. Increased peroxidase activity has been reported \textit{in vitro}\textsuperscript{130,131} in the H48Q, A4V, and G93A variants, although not consistently.\textsuperscript{132} Increased peroxidase activity in vivo has been reported in the A4V and G93A\textsuperscript{132} species. Cu replete, Zn deficient SOD1 has been reported to confer toxicity by producing peroxynitrite according to these reactions, and loss of Zn from mSOD1 has been proposed as a primary pathogenic event.\textsuperscript{133} The second zinc metalloprotein that is aberrant in ALS patients is metallothionein, immunoreactivity to which is elevated in the brain and liver.\textsuperscript{127,128} The same pattern of elevated metallothionein immunoreactivity occurs in a transgenic model of ALS: SOD1-G93A transgenic mice demonstrate increased MT-1, MT-2, and MT-3 expression in astrocytes and increased MT-3 in neurons.\textsuperscript{134} Metallothionein elevation is likely compensatory and protective. In the G93A mutant SOD1 transgenic model of ALS, deficiency of MT-1, MT-2 or MT-3 exacerbates the ALS phenotype.\textsuperscript{135,136}

1. Buffering Free Zinc

There are three general directions for effective zinc-based drug development. (i) Zinc buffers with equilibrium constants at the optimal value, preventing excess zinc damage while avoiding zinc deficiency of the brain; (ii) for acute brain injuries (stroke, trauma, ischemia, hypo-perfusion), a very short-lived chelator with tighter-binding compounds that allow some control of zinc toxicity with minimal untoward effects of lowered zinc; (iii) “pro-buffers” or “tethered buffers” which could be designed to act upon zinc only whenever or wherever such zinc buffering is therapeutically required.

The first strategy, that of using a relatively weak chelator, has already produced promising results. The quinoline compound clioquinol, which binds zinc in the mid nanomolar range, has been shown to reduce the amount of amyloid plaque in transgenic mice dramatically\textsuperscript{137,138} and to slow the rate of cognitive decline in human patients\textsuperscript{18,139} with AD. Unfortunately, the phase III trial with clioquinol was discontinued due to a problem in the manufacturing process. Other candidate chelators are considered as alternatives.

Another promising use of the low-affinity approach has been reported for acute zinc-toxicity. In those studies it was shown that, pyrithione (Kd~1 µM) can rescue cultured cells from zinc toxicity if administered at the right time.\textsuperscript{140,141} Pyrithione moves freely through
membranes and presumably transports free zinc down its concentration gradient, thus rescuing cells from zinc toxicity when intracellular zinc levels are higher than extracellular zinc levels.

The idea of a “pro-drug chelator” is also under active investigation as a treatment for Alzheimer’s disease. In this case, a classical strong chelator (BAPTA) has been rendered lipophilic and inactive by the addition of alkyl chains. Once through the blood brain barrier and embedded in a cell wall (lipid membrane) the non-chelating drug (DP-109) can be transformed into the active BAPTA by membrane lipases. Hence it is expected that DP-109 will chelate metals predominantly in the vicinity of cell membranes. In Tg2576 mice, DP-109 significantly reduced Aβ plaque load by about 60-80% without noticeable side effects. A related compound (DPb99) has also been tested in small samples of human patients as a neuroprotective against the zinc-mediated injury in stroke, and during the coronary bypass surgery.

2. Downstream Control of Zinc-Triggered Toxic Signals

Therapies targeting later events are also promising. As discussed above, diverse serial and parallel events contribute to zinc-induced cell death. First, as zinc toxicity is largely mediated by oxidative and nitrosative stress73,75,92,142 antioxidants and NOS inhibitors may be useful.

Another approach would target inhibition of PARP, which appears to be a key downstream event in zinc toxicity,92,95 may be effective in reducing zinc toxicity. Third, anti-apoptosis measures such as caspase inhibition may be a possibility. Although these mechanisms have been demonstrated to contribute to zinc toxicity in cell culture, they are considered more or less general mechanisms of cell death in acute brain injury. At the moment, it is not known whether any particular neuroprotectant is better against zinc toxicity than other injury mechanisms. Hence, more studies may be needed to zoom in on drug targets that are more specific to zinc toxicity.

Pyruvate protects against zinc-induced cell death in cortical culture144 Pyruvate protection is somewhat specific to zinc toxicity, because pyruvate does not attenuate calcium-overload excitotoxicity in the same cortical cell culture.145 Consistently, in a rat model of transient global ischemia where the role of zinc is established,62 pyruvate almost completely blocks zinc accumulation as well as neuronal death throughout the brain.145 Pyruvate also reduces retinal cell death following zinc exposure in culture or following pressure-induced ischemia in rats.146 Protection by pyruvate against zinc-triggered cell death is applicable not only to neurons and glial cells, but also to pancreatic beta cells. Streptozotocin-induced beta cell death to which paracrine toxic effect of endogenous zinc contributes, is markedly attenuated by pyruvate administration.147 Direct antioxidative effect and/or normalization of NAD+ levels may contribute to cytoprotection by pyruvate.143,148

Another possible neuroprotectant with specificity against zinc-mediated injury is tPA, which is currently used for thrombolysis in human patients.149 Although most of tPA’s biological effect, including its excitotoxicity-potentiating effect, is mediated by its protease action,150 blockade of zinc toxicity by tPA takes place even in the presence of excess protease inhibitors.151 Although the protective mechanism is still unclear, tPA had no effect on zinc influx into cells, excluding the possibility that the protection occurs by the chelation of zinc in the media. Rather, a subsequent study showed that tPA increases zinc influx into cells.152 A preliminary result suggests that certain membrane receptors with tyrosine kinase activity may mediate this effect, since EGF receptor tyrosine kinase inhibitor C56 can reverse the protection (Koh unpublished). If the effective moiety and its cognate membrane receptors can be identified, development of tPA-derived peptides that prevent zinc toxicity, may be a possibility.

CONCLUSION

Like calcium, zinc is proving to be an essential and ubiquitous ionic signal in a myriad of cells and tissues. Because fluorescent calcium probes frequently respond to zinc as well, separating calcium signals from zinc
signals will be mandatory in future research. Therapies based on manipulating zinc signals by preventing release, blocking channels, altering transport and buffering zinc of target tissues are all likely to have increasingly important roles in twenty-first century medicine.

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