Clinical aspects of trace elements: Zinc in human nutrition – A biochemical and physiological perspective

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T HIS REVIEW DISCUSSES THE TISSUE distribution, concentration and intracellular binding of zinc in healthy humans. It is the first of a five-part series that examines zinc in terms of its biochemistry and physiology, metabolism, dietary requirements, nutritional assessment, and states of excess and deficiency.

The term ‘trace element’ was coined half a century ago when scientists began describing a number of mineral elements that occur in minute amounts (‘traces’) in biological materials (1). These low concentrations made measurement of the elements virtually impossible given the analytical methods available at the time. However, the tremendous technological advances in recent years have provided the analytical means for trace elements to be measured with speed, sensitivity and precision.

Trace elements, as noted above, can be distinguished from major mineral elements in that they exist and function in living organisms in very small concentrations. ‘Trace’ should not be taken to mean ‘unimportant’ because a number of trace elements act as vital catalysts for essential enzymes, stabilize membranes, provide tissue structure and are involved in hormonal function. At present 22 mineral elements are considered essential for life (2).
Trace metals are considered essential when an inadequate intake produces an impairment of function, and when supplementation with the element reverses the impaired function or prevents impairment (3,4). It is likely that some of the 20 to 30 elements considered to be toxic substances or contaminants will be found to be essential in small amounts. Trace element deficiencies may arise from both inadequate dietary intake and decreased bioavailability, or they may be associated with disease states in which impaired absorption, excessive excretion, and/or excessive utilization occurs.

**BIOCHEMISTRY OF ZINC**

Zinc has been recognized as an essential nutrient for humans since the early 1960s (5,6). The human body normally contains 2 to 3 g of zinc (2,7), and zinc is found in all tissues in varying concentrations; the highest concentrations occur in the retina, prostate, prostatic secretions and spermatozoa. Three-quarters of the total bodily amount can be found in the skeleton, from which it is slowly removed over time (2). Significant amounts also appear in the liver, kidney and skeletal muscle (8,9) (Table 1). In the blood, zinc is concentrated within the erythrocytes. A small amount appears in blood serum (2) which contains a zinc concentration of approximately 18.4 mol/L (10). Zinc in the form of an intracellular ion accounts for over 95% of total body zinc (11). Intracellular zinc is predominantly found in the cytosol; about 10 to 20% is seen in the nucleus, and smaller amounts are present in the microsomal and mitochondrial fractions (12). A smaller amount of zinc is found in the cell membrane, and it has been suggested that the fraction of intracellular zinc specifically bound to membranes may play an important role in the onset of zinc deficiency symptoms (13).

The chemical state of zinc within cells is not clearly understood. Biologically, zinc is found attached to organic ligands and proteins rather than free in solutions as a metallic ion. However, it appears that zinc may play a controlling role in some enzymes rather than being firmly bound as an integral part of the molecule (11). This finding may imply that a readily available zinc ‘pool’ exists within cells or it may suggest that binding substances assist in enzyme activation by zinc (11). The characteristics of these pools or substances remain unclear. A number of researchers have reported that cytosolic zinc of a number of different tissues is distributed in three or four pools within the cell, and that these pools consist of mainly larger weight molecular proteins, including many of the zinc metalloenzymes (14). It appears that there are age-related differences in zinc distribution. There are considerable differences in the relative sizes of tissues, as well as in their zinc content, between neonates and the adult human. In particular, the zinc content of liver and bones is much higher in newborns (about 25 to 40% of the total body zinc) and the relative size of the liver is much larger (11,15). Much of this zinc may be used for soft tissue growth since neonatal bones undergo comprehensive remodelling after birth (11).

There is little information on zinc nutriture in the elderly. Some studies have shown a decline in plasma zinc with age (16), while others have failed to observe such a change (17,18). Whether age affects zinc requirements is not known. A few studies have investigated the relationship between zinc nutriture and the immune function of elderly persons. This subject is important because the deterioration of T cell immune function is associated with both zinc deficiency and ageing. There is, thus, the possibility that poor zinc status contributes to T cell immune dysfunction (16).

**TABLE 1**

Approximate zinc content of major tissues in the normal human

<table>
<thead>
<tr>
<th>Tissue</th>
<th>Zinc content (mg/kg dry weight)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liver</td>
<td>141-245</td>
</tr>
<tr>
<td>Kidney</td>
<td>184-230</td>
</tr>
<tr>
<td>Lung</td>
<td>67-86</td>
</tr>
<tr>
<td>Skeletal muscle</td>
<td>197-226</td>
</tr>
<tr>
<td>Pancreas</td>
<td>115-135</td>
</tr>
<tr>
<td>Bone</td>
<td>218</td>
</tr>
<tr>
<td>Prostate</td>
<td>520</td>
</tr>
<tr>
<td>Eye (retina)</td>
<td>571</td>
</tr>
</tbody>
</table>

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tivities of the DNA-synthesizing enzymes DNA polymerase and thymidine kinase. Numerous investigators have showed that decreased DNA synthesis is associated with the decreased specific activity of both enzymes (24-26), and Duncan and Hurley (27) have demonstrated that zinc deficiency results in reduced DNA polymerase activity in rats. Many other investigators have shown that zinc deficiency in animals impairs the incorporation of labeled thymidine into DNA (9). This decreased enzyme activity may ultimately be responsible for the growth retardation linked to zinc deficiency.

Pardee et al (28) proposed that zinc plays a role in cell division. These researchers have suggested that zinc may have a role in stabilizing a multiprotein ‘replicase’ complex consisting of such enzymes as DNA polymerase, thymidine kinase and thymidine synthase before the incorporation of deoxynucleotides into DNA. Without zinc, the spatial relationships among the various components may be modified in such a way as to prevent proper DNA synthesis. In this same article, Pardee et al provide evidence that shows that a final regulatory protein must be synthesized and united with the replicase complex before the replicase can function. It is possible that insufficient quantities of zinc may impede the transcription of mRNA that is required for synthesis of adequate amounts of this protein.

Zinc’s regulation of the cell cycle may occur after DNA synthesis at the point of chromatin decondensation before mitosis and during assembly of the mitotic spindle via zinc’s effects on microtubule assembly. Sen and Crothers (29) showed that zinc has the ability to modulate the condensation/decondensation of chromatin. Zinc can stimulate the polymerization of purified tubulin in vitro (30). It is thus likely that zinc plays a critical role in regulation of the cell cycle, and that a deficiency can disrupt the normal cell synthetic processes.

Recent research has focused on zinc’s interaction with various chromatin components. It has been demonstrated that zinc can stabilize the DNA double helix with respect to thermal denaturation (31). In addition, zinc can promote recombination of single-stranded DNA into its native double-stranded form through its ability to bind to both the phosphate backbone and nucleoside bases of DNA (31). Zinc also promotes conformational transformations of DNA (32). Because zinc deficiency can alter the conformation of chromatin, RNA polymerase access to various genes may be impeded by a lack of zinc (30). Castro and colleagues (33) demonstrated that an alteration in the amounts of histone H1 occurs in zinc-deficient rat liver. Because histones are necessary to the proper structure and functioning of chromatin, these alterations may explain some of the abnormal metabolic processes occurring in zinc-deficient organisms. However, the mechanism by which zinc affects histone metabolism is not known.

Nonhistone proteins responsible for gene expression, such as RNA polymerase, can also be affected by zinc deficiency. In one study, Euglena gracilis cells grown in a zinc-sufficient medium produced three distinct RNA polymerases, while cells grown in zinc-deficient media synthesized only one RNA polymerase, which was different from the other three (34). It is not known whether zinc affects the transcription of RNA polymerase genes directly or indirectly. Some researchers have been investigating a direct role for zinc in gene transcription. A Xenopus transcription factor has been identified as a zinc metalloenzyme (30). This suggests that some genes may be partially or fully regulated by zinc enzymes. Given the past and current evidence, it appears realistic to presume that some of the defects caused by zinc deficiency occur as a result of altered chromatin metabolism. This possibility may be of particular importance in the case of the fetus, where synchronized timing and expression among genes are critical for normal development.

RNA polymerases contain more than 1 mol of zinc/mol protein (30). It is reasonable to suggest that zinc has both structural and catalytic functions in RNA polymerases, and that a deficiency of dietary zinc can lead to a reduction in the activity or function of RNA polymerases (30). Terhune and Sanstead (35) have demonstrated that RNA polymerase activity is significantly lower than normal in the liver of postnatal zinc-deficient rats. Zinc is now known to be a normal constituent of RNA; it is also important to the stabilization of ribosome conformation (30). Both of these macromolecules are present in pre-initiation and initiation complexes; it is, therefore, likely that zinc plays an essential role in stabilizing these structures. In a number of studies, zinc-deficient organisms were found to have decreased amounts of, and reduced synthetic rates for, RNA (30). It has been suggested that these lowered RNA rates and amounts are mainly the result of increased ribonuclease activity, which in turn is regulated by zinc (or, in this case, a lack thereof). Taken as a whole, the information on abnormal gene expression and RNA metabolism in zinc deficiency states strengthens the hypothesis that zinc deficiencies alter protein metabolism. Defects in protein synthesis are detrimental to the human fetus, as well as to children and adults.

Wound healing: Zinc’s involvement in the cell cycle, as discussed above, is integral to its role in wound healing. Studies have confirmed that zinc accumulates around the wound (where cell division takes place most vigorously), particularly during the initial week of healing (36). Supplementary zinc, however, only appears effective in promoting healing when small amounts are ingested and low plasma zinc levels are present. There is no evidence that high levels of zinc ingestion provide any additional advantages (2,36).

Growth and reproduction: Because zinc has its most profound influence on rapidly growing tissues, its effect on growth and reproduction is important. Zinc is necessary for adequate gonadal development and growth response from infancy to adulthood (2). Zinc deficiency causes growth retardation in infants, children and adolescents. A mirror relationship exists between plasma zinc levels and the growth index, an indicator of growth velocity (9). Animal studies have shown a graded response in the impairment of growth ve-
progressively decreased from the minimum level required to maintain maximal growth velocity (37). Studies conducted by both Hambidge (37) and Walravens et al (38) have shown consistent and significant positive effects of zinc supplementation (in physiological doses of 5 mg/day) on linear growth increments for mildly zinc-deficient children. A significant difference in linear growth rates between zinc-supplemented and placebo-treated children was invariably observed for males but not females (12). The reason for this sex-linked difference is unclear. Additionally, the direct local effects of zinc deficiency on bone epiphyses are thought to reduce longitudinal growth (39). Depletion of zinc body stores may occur during growth spurts. Thus, children and adolescents would benefit from an increased zinc intake during such periods (9).

Zinc deficiency may depress growth rates through several different mechanisms. At a molecular level, several of the enzymes required for nucleic acid synthesis either contain zinc or are zinc-dependent, and relatively large amounts of zinc are required for new tissue synthesis. Additionally, decreased food intake is an early and notable feature of experimental zinc deficiency (1). Some of zinc’s effects on growth may be mediated through changes in the production or secretion of somatomedin. Somatomedin is thought to mediate the skeletal growth-promoting effect of growth hormone. It has been proposed that zinc deficiency may both decrease the effectiveness of growth hormone in stimulating somatomedin production and directly decrease the biological effectiveness of somatomedin in stimulating cartilage growth (39).

Although there have been a substantial number of investigations of male sex hormones and spermatogenesis, little is known about the effects of zinc on the sex hormones of the non-pregnant female. Zinc is important to male testicular function, affecting both spermatogenesis and the production of testosterone by the Leydig’s cells. Zinc supplementation has been shown to reverse abnormalities of sexual maturation and hypogonadism in human males (9). Although hypogonadism in zinc-deficient males appears to be more common than ovarian dysfunction in zinc-deficient females (9), zinc does play an important role in pregnancy (40) due to its involvement in cellular growth and maturation, its action as an antibacterial and antiviral agent in amniotic fluid, and the positive correlations that exist between maternal serum zinc concentration and birth weight (41).

**Insulin function:** Zinc is involved in the action of insulin, thereby affecting glucose tolerance (2,9). A delayed absorption of glucose has been demonstrated in zinc-deficient patients (42). It has been suggested that the rate of insulin secretion in response to glucose stimulation is reduced during zinc deficiency (9). Additionally, it is thought that zinc participates in the storage of insulin in the beta cells and that the amount of insulin stored in zinc-deficient subjects is decreased (9). Alternatively, an increase in the degradation of insulin may account for the decrease in glucose tolerance (9). Depressed serum zinc and hyperzincuria are associated with type I and type II diabetes (43,44). Increased urinary excretion of zinc is also positively related to glucose excretion.

**Immunological competence:** Numerous clinical reports have suggested an association between zinc deficiency and impaired immune competence. In humoral immunity, the interleukin-1 signal, which acts to stimulate helper T cells and pre-beta cells, causes a rapid redistribution of zinc from the plasma pool to the liver, bone marrow and thymus (45). The reduction in plasma zinc may be critical for phagocytic cell activation, while the stimulation of zinc uptake by the thymic and bone marrow tissues may suggest a critical need for this element during T and beta cell generation (45). The production and/or action of some cytokines is thought to be zinc-dependent (45). For example, the production and/or membrane-binding of interleukin-1, interleukin-2 and interferon may be zinc-dependent (46-48). In the case of cell-mediated immunity, zinc deficiency has not been shown to affect the complement system negatively (45). However, zinc at higher levels of the normal physiological range inhibits the complement cascade (49). Thus, the reduction in plasma zinc levels seen following an infection may be a positive consequence for this branch of the immune system.

The production of T lymphocytes is impaired in zinc deficiency states (4,45). Whether zinc deficiency has a differential effect on specific T cell subsets is currently an area of controversy. Zinc deficiency often presents with a loss of lymphoid tissue mass exceeding that of other body tissues (50). Chronic zinc deficiency will result in atrophy of the entire organ (51). The effects of zinc deficiency on T cell maturation have been attributed to a possible reduction in the activity of deoxyribonucleotide transferase, a zinc-containing DNA polymerase found in high concentrations in mature thymocytes (52). It is also possible that zinc deficiencies may alter thymic epithelial function and impair thymic hormone production, the activities of which, in turn, would inhibit T cell maturation in the thymus and periphery (53). In addition, overall responsiveness to T cell-independent antigens, as well as to T cell-dependent mitogens, may be noticeably inhibited in cases of zinc deficiency (45,51).

A zinc deficiency may directly influence lymphocyte proliferation. The in vitro stimulation of blast transformation to mitogens such as phytohemagglutinin can be inhibited by zinc-chelating agents (45). Zinc acts as a mitogen to both T and beta cells (5,9,45). Flynn (47) has suggested that zinc deficiency may affect T cell prolif-
eration by reducing the production of cytokines responsible for proliferation, by interfering with the processing of antigens by accessory cells or by causing a loss of cell function or ‘activation state’ (45).

Delayed-type hypersensitivity, which depends on the interaction between effector T cells and macrophages, is reduced in zinc-deficient animals (51, 54). Macrophage function can be inhibited in zinc-deficient animals (55). Defects in macrophage function include a decreased ability to take up the target and kill it; this decrease is probably due to the limitation of zinc activity in a critical process such as maintaining membrane integrity or tubulin-mediated phagocytosis (55). Decreased natural killer cell activity and depressed cytotoxic responses have been reported in zinc-deficient mice, although increased natural killer activity with zinc deficiency has also been reported. This defective chemotaxis in mammals indicates that neutrophil function is also affected by zinc deficiency (56,57).

The immunological changes observed in zinc-deficient humans and animals may be largely responsible for increased susceptibility to infection. Data obtained by numerous researchers demonstrate an increased susceptibility of zinc-deficient animals to viral, parasitic and bacterial challenges (45). Virtually all immunological abnormalities discussed above are rapidly corrected following zinc repletion (51). There is, however, a notable exception: immunological memory that is lost due to postnatal zinc deficiency may not be fully recovered upon zinc repletion (55).

The importance of zinc as a micronutrient can be seen in the vast variety of essential physiological functions it performs (Table 2). This importance has served to further the interest in the biological roles of zinc and its nutritional significance. Preceding reports have given an overview of current knowledge regarding the role of zinc in basic physiological and biochemical processes. Further investigation is necessary if the precise mechanisms underlying the role of zinc in the normal function of the human organism, as well as its role in disease processes, are to be clarified.

REFERENCES
