Copper deficiency myeloneuropathy and pancytopenia secondary to overuse of zinc supplementation

J Rowin, S L Lewis

The haematological complications of acquired copper deficiency have been well documented, but the neurological complications have only recently been reported. An illustrative case of copper deficiency myeloneuropathy with pancytopenia is presented and the potential aetiologies and neurological manifestations of this deficiency state discussed.

CASE REPORT
A 53 year old woman presented with progressive gait imbalance. Four months before presentation she began to develop tingling and numbness in the fingers and feet. One month later she visited an outside emergency room for recurrent sinusitis. She was found to have an anaemia with a haemoglobin concentration of 8.3 g/dl and a white blood cell count of 6 × 10³ with a mean corpuscular volume of 112.5 μm³ (normal 80 to 100). The electrolytes, liver enzymes, glucose, thyroid stimulating hormone, fatty acid profile, vitamin B-12, folic acid, homocysteine, methylmalonic acid, vitamin E, serum protein immunoelectrophoresis, serum protein immunofixation, angiotensin converting enzyme, free fatty acids, and arylsulphatase A were all normal. Antinuclear antibody, SS-A and SS-B antibodies, antigliadin antibody, and antiphospholipid antibodies were negative. Syphilis, Lyme, and HIV serology were negative.

The serum copper level was markedly low at 7 μg/dl (normal 70 to 155), as was the serum caeruloplasmin at 2.1 mg/dl (normal 22.9 to 43.1). Serum zinc was raised at 2.28 μg/ml (normal 0.66 to 1.10). The patient’s zinc supplement was inspected. Each capsule contained 50 mg of elemental zinc in the form of zinc gluconate. The recommended dosage, according to the label, was one capsule daily (333% of the recommended daily dose of 15 mg). The patient described taking four to eight capsules daily; she would increase the dose according to the severity of her symptoms, ingesting up to 400 mg/day.

A diagnosis of zinc induced copper deficiency myeloneuropathy associated with pancytopenia was made. The zinc was discontinued. She received one dose of intravenous copper chloride (2.0 mg elemental copper) and was subsequently started on oral copper sulphate supplementation (2.0 mg/day elemental copper). A bone marrow biopsy taken 12 days after the start of copper supplementation was normocellular with slightly left shifted maturation in all cell lines. No tumour cells were present. Her copper level, white blood cell count, and mean corpuscular volume normalised after three months of treatment. There was significant but incomplete improvement in her gait and spasticity at six months. There was also some improvement in her nerve conduction studies, with the appearance of a normal sural sensory response (latency 4.0 ms (normal < 4.2), amplitude 6.5 μV (normal > 5.0)). There was no improvement in the motor conduction times.

DISCUSSION
We believe that our patient developed copper deficiency myeloneuropathy and pancytopenia secondary to overuse of zinc supplements, as in a case reported by Kumar et al. Our patient was ingesting approximately 20 times the recommended daily allowance of zinc. Both copper and zinc are absorbed in the stomach and proximal duodenum. Excess...
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zinc levels cause an upregulation of metallothionein production in the enterocytes. Copper has a higher affinity for metallothionein than zinc, so it displaces zinc from metallothionein. Copper then remains in the enterocytes and is sloughed off into the intestinal tract and eliminated.

Other aetiologies of copper deficiency myeloneuropathy have been reported, including five cases secondary to gastric resection. Other cases have an unclear aetiology with no known external sources of zinc, but with high plasma zinc concentrations. Copper deficiency with haematological manifestations has been reported in malnutrition, prematurity, and parenteral or enteral feeding that does not include copper.

There are some features in common between the neurological manifestations of copper deficiency and multiple sclerosis. Interestingly, the copper chelating agent cuprizone is used as a neurotoxicant in a mouse model of CNS demyelination. Although our patient did not show central demyelinating lesions on MRI, Prodan and Holland reported CNS white matter lesions in the brains of their patients with copper deficiency. Also, hyperintensity on T2 weighted MR images can be seen in the dorsal spinal cord of some patients with copper deficiency myelopathy.

Gastrectomy surgery for obesity is associated with neuropathy as well as other neurological complications. It appears that approximately 40% of the cases of post-gastric reduction surgery neuropathy are associated with thiamin or vitamin B-12 deficiency, but in the remaining 60% no vitamin deficiency is found. As there are reports of copper deficiency myeloneuropathy after gastric bypass surgery for various indications in the absence of parenteral or enteral nutrition, copper deficiency should be considered in cases of neuropathy after weight reduction surgery.

As with vitamin B-12 deficiency, the neurological manifestations of copper deficiency may be seen with or without the haematological manifestations, and with or without abnormalities on MRI imaging of the brain and spinal cord. The associated anaemia of copper deficiency may be macrocytic, as in our case, microcytic, or normocytic. Therefore a high index of suspicion is necessary in patients at risk. Copper therapy (2 mg/d) generally leads to an early recovery of the haematological abnormalities, followed by variable recovery of the neurological symptoms.

Copper deficiency should be considered in the differential diagnosis of multiple sclerosis, subacute combined degeneration of the cord, optic myeloneuropathy, post-gastric reduction surgery neuropathy, or in other cases of myelopathy, optic neuropathy, or polynuropathy where nutritional deficiency or overuse of zinc supplementation is suspected. Prompt recognition and treatment may improve the prognosis for neurological recovery.

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