Contamination of water, air, and food by chemicals and heavy metals is an unfortunate consequence of our industrialized, high-tech society. The resultant accumulation of heavy metals in the human body poses a significant health risk, leading to a wide array of symptomatology and disease states. Although environmental lead levels have decreased in the United States since lead was eliminated from gasoline and lead-based paint, lead continues to be a significant problem, particularly in urban areas and areas of lead mining and smelting. In addition, mercury, cadmium, and arsenic toxicity from occupational and environmental exposure also continue to pose significant threats to public health. For these reasons, diagnostic testing for heavy metals and the subsequent decrease in the body’s burden of these substances has become a necessity. Meso-2, 3-dimercaptosuccinic acid (DMSA) is a sulfhydryl-containing, water-soluble, non-toxic, orally administered, metal chelator that has been in use as an antidote to heavy metal toxicity since the 1950s. Recent clinical use and research substantiates this compound’s efficacy and safety, and establishes it as the premier oral metal chelation compound when compared to other available chelating substances.

Biochemistry and Pharmacokinetics

The ability of sulfhydryl-containing compounds to chelate metals is well-established. DMSA is a dithiol (containing two sulfhydryl, or S-H, groups) and an analogue of dimercaprol (BAL, British Anti-Lewisite), a lipid-soluble compound also used for metal chelation. DMSA’s water solubility, oral dosing, large therapeutic window, and low toxicity make it superior to other chelating agents available. Approximately 20 percent of an oral dose of DMSA is absorbed from the gastrointestinal tract of healthy individuals. Ninety-five percent of the DMSA that makes it to the bloodstream is bound to albumin. It is suggested that one of the sulfhydryls in DMSA binds to a cysteine residue on albumin, leaving the other S-H available to chelate metals.

Clinical Indications

Lead Toxicity

Lead exposure continues to be a public health problem in the United States. It is still found in millions of pre-1940s homes. Lead toxicity causes numerous malfunctions in calcium uptake and utilization, and also interferes with calcium-facilitated cellular metabolism. Lead is particularly toxic to the central nervous system (CNS), as evidenced by its particularly deleterious effects on mental development and intelligence in children with lead toxicity. In addition, neurobehavioral deficits resembling attention deficit disorder have been attributed to lead exposure. In a child, blood lead concentrations of
20-25 mg/100 ml can cause irreversible CNS damage. In adults, acute lead exposure leads to renal proximal tubular damage, while chronic exposure causes renal dysfunction characterized by hypertension, hyperuricemia, gout, and chronic renal failure.

**Mercury Toxicity**

Human exposure to mercury is primarily in two forms: mercury vapor and methylmercury compounds. Mercury vapor in the atmosphere makes its way into fresh and salt water by falling in precipitation. Methylmercury compounds are created by bacterial conversion of inorganic mercury in water and soil, and are subsequently concentrated in seafood. Dietary fish intake has been found to have a direct correlation with methylmercury levels in blood and hair. "Silver" amalgam dental fillings are the major source of inorganic mercury exposure in humans. The most commonly used dental amalgam material contains approximately 50-percent liquid metallic mercury; thus, amalgam preparation and placement in the patient’s mouth results in mercury vapor exposure for the patient, dentist, and technician. Mercury vapor continues to be released as the patient chews, brushes, or drinks hot beverages, after which it is inhaled into the lungs and enters the bloodstream. Studies have shown a direct correlation between the number of amalgam fillings and mercury concentration in the blood and urine. Amalgam removal results in decreased blood-mercury levels below the pre-removal baseline.

Research has demonstrated that DMSA, when compared to treatment with other chelating agents, resulted in the greatest urinary excretion of mercury, as well as being the most effective at removing mercury from the blood, liver, brain, spleen, lungs, large intestine, skeletal muscle, and bone. Another study indicated that mercury excretion was greatest in the first 8-24 hours after oral DMSA administration. Animal studies show that following intravenous administration of methylmercury, DMSA was the “most efficient chelator for brain mercury,” removing two-thirds of the brain mercury deposits.

**Cadmium and Arsenic Toxicity**

Environmental cadmium exposure comes from pollutants discharged by industries utilizing it, including herbicide and battery manufacturers. It is also found in cigarette smoke. Cadmium, like lead and mercury, can interact metabolically with nutritionally essential metals. Cadmium interacts with calcium in the skeletal system to produce osteodystrophies, and competes with zinc for binding sites on metallothionein, which is important in the storage and transport of zinc during development. Cadmium poisoning can lead to rhinitis, nephropathy, and osteomalacia, and has a possible link to cardiomyopathy, hypertension, and hepatic and prostate disorders.

Arsenic toxicity is also usually a result of exposure to industrial pollutants or cigarette smoke. Symptoms include eczema, dermatitis, malaise, muscle weakness, and “garlic breath.” The use of DMSA in arsenic chelation is more effective in cases of acute poisoning than in those of long-term exposure. This may be due, in part, to the possibility that DMSA is a more effective chelator of arsenic in the bloodstream than it is of the tissue-bound arsenic seen in long-term exposure. In addition to the metals mentioned above, DMSA and other dithiol agents have a binding affinity for bismuth, tin, nickel, and thallium.

**Side-Effects and Toxicity**

DMSA is very safe and generally well tolerated, with few side-effects being noted. Some patients may experience slight gastrointestinal disturbances or urticaria, but it is not usually necessary to discontinue treatment. Any detoxification regimen requires the bowels to be fully functioning.
patient is constipated, normal bowel function should be restored prior to DMSA chelation. Chelation of heavy metals can also result in chelation of copper, manganese, molybdenum, and zinc, resulting in deficiencies. Although DMSA does not directly bind magnesium, cysteine, and glutathione, heavy metal detoxification can result in depletion of these nutrients as well. Therefore, deficiencies of these essential elements should be screened for and corrected. Sulfhydryl compounds in DMSA will make the urine smell sulfurous, necessitating adequate communication with the patient regarding this issue.

**Dosage/Protocols**

Dosing protocols for DMSA vary depending on physician preference and individual patient need, but currently two protocols are most often used. In one protocol, 10-30 mg/kg per day is given in three divided doses, using a three-days-on, 11-days-off cycle with a minimum of eight cycles. A second protocol involves giving 500 mg per day (in two or three divided doses) every other day for a minimum of 5 weeks. DMSA is best taken between meals.

**References**


