

Lead Poisoning

Among the natural substances that man concentrates in his immediate environment, lead is one of the most ubiquitous. A principal cause for concern is the effect on children who live in decaying buildings

by J. Julian Chiodo, Jr.

Lead has been mined and worked by man for millennia. Its chemical, high resistance to corrosion and other properties make it one of the most useful of metals. The inappropriate use of lead has, however, resulted in outbreaks of lead poisoning in humans from time to time since antiquity. The disease, which is sometimes called "plumbism" (from the Latin word for lead) or "saturnism" (from the alchemical term), was first described by the Greek poet-physician Nemesius more than 2,000 years ago. Today our concern about human health and the dissemination of lead into the environment are twofold: (1) there is a need to know whether or not the current level of lead absorption in the general population presents some subtle risk to health; (2) there is an even more urgent need to control this hazard in the several subgroups within the general population that run the risk of clinical plumbism and its known consequences. In the young children of urban slums, lead poisoning is a major source of brain damage, mental deficiency and serious behavior problems. Yet it remains an insidious disease: it is difficult to diagnose, it is often unrecognized and until recently it was largely ignored by physicians and public health officials. Now public attention is finally being turned on childhood lead poisoning, although the difficult task of eradicating it has just begun.

Symptomatic lead poisoning is the re-

sult of very high levels of lead in the tissues. Is it possible that a constant of lead in the body that is insufficient to cause obvious symptoms can nevertheless give rise to slowly evolving and long-lasting adverse effects? The question is at present unanswered but is more pertinent. There is much evidence that lead wastes have been accumulating during the past century, particularly in congested urban areas. Increased exposure to lead has been shown in populations exposed to lead at an old paint mill. Post-mortem examinations show a higher lead content in the organs of individuals in highly industrialized societies than in the organs of most individuals in primitive populations. Although no population group is apparently yet being subjected to levels of exposure associated with the symptoms of lead poisoning, it is clear that a continued rise in the pollution of the human environment with lead could eventually produce levels of exposure that could have adverse effects on human health. Efforts to control the dissemination of lead into the environment are therefore indicated.

The more immediate and urgent problem is to control the exposure to lead of well-defined groups that are known to be directly at risk: young children who live in dilapidated housing where they can nibble chips of leaded paint, which they drink who consume quantities of lead-contaminated foodstuffs, people who eat or drink from improperly lead-glazed

earthenware, workers in certain small-scale industries where exposure to lead is not controlled. Of these the most disturbing group is the large group of children between about one and three to five years of age who live in deteriorating buildings and have the habit of eating nonfood substances including peeling paint, plaster and putty containing lead. (This behavior is termed pica, after the Latin word for magpie.) The epidemiological data are still scanty; large-scale screening programs now in progress in Chicago and New York City indicate that between 3 and 10 percent of the children tested show evidence of asymptomatic increased lead absorption and that between 1 and 2 percent have unsymptomatic plumbism. Small-scale surveys in the worst housing areas of a few other cities reveal even higher percentages.

There is little doubt that childhood lead poisoning is a real problem in many of the urban slum areas of the U.S. and perhaps in rural communities as well. Current knowledge about lead poisoning and its long-term effects in children is adequate to form the basis of a rational attack on this particular problem. The ubiquity of lead-pigment paints in older substandard housing and the prevalence of pica in young children indicate, however, that any effective program will require the concerted and sustained effort of each community. Furthermore, the continued use of lead-pigment paints on housing surfaces that are accessible to

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young children and will at some future date fall into the category of only perpetuate the problem.

Traces of lead are almost ubiquitous in nature and minute amounts are found in normal diets. According to the extensive studies of Robert A. Kehoe and

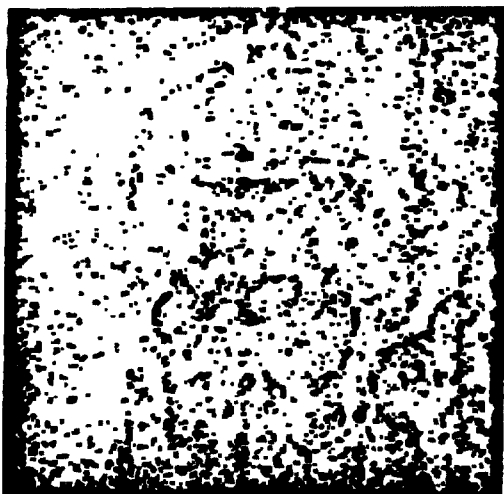
his associates during the past 35 years at the Kettering Laboratories of the University of Cincinnati, the usual daily dietary intake of lead in adults averages about .3 milligram. Of this, about 90 percent passes through the intestinal tract and is not absorbed. Kehoe's data

indicate that the small amount absorbed is also excreted, so that under "normal" conditions there is no net retention of lead in the body. In addition the usual respiratory intake is estimated at between .25 and .50 micrograms of lead per day. These findings must be re-ex-

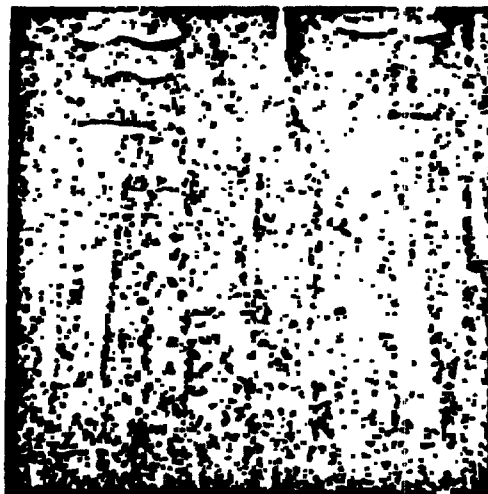


EXCESS LEAD combined with protein forms inclusion bodies in the nuclei of certain cells in lead-poisoned animals and man. In an electron micrograph made by Robert A. Coyer and his colleagues at the University of North Carolina School of Medicine the surface

of a cell from a proximal renal tubule of a lead-poisoned rat is enlarged 25,000 diameters. The large structure with a dense core and a fibrous outer zone is an inclusion body; below it to the left is a smaller one. The dark area below the large body is the nucleus.



X-RAY PLATES may show evidence of lead ingestion as of an over-the-body burden of the metal. The abdominal X ray (left) shows a number of bright opaque particles in the large intestine;



tion of lead-containing paint that had been eaten by the 16-month-old subject. The X ray of the same child's legs (right) shows bright "lead lines" across lead stored at the end of the long bones.

cified with postmortem analyses, which indicate that the concentration of lead in bone increases with age, although its concentration in the soft tissues is relatively stable throughout life. The physiological significance of increasing storage in bone is not entirely clear, but it has caused considerable concern. It is quite clear that at the level of intake of lead increases, the rate of absorption may exceed the rate at which lead can be excreted or stored in bone. And when the rates of excretion and storage are exceeded, the levels of lead in the soft tissues rise. Studies in adults indicate that as the estimated daily intake of lead rises above one milligram of lead per day, higher levels of lead in the blood result, and metabolic, functional and clinical responses follow (see illustration on pages 22 and 23). The reversible effects abate when the rate and amount of lead absorbed are reduced again to the usual dietary range.

As far as is known, lead is the trace element essential to no child, but this particular question has not been adequately examined. Some of the adverse effects of lead on metal ions have been studied in considerable detail. These effects are related to the concentration of lead in the soft tissues. At the level of cellular metabolism, the best-known adverse effect of lead is its inhibition of the activity of enzymes that are dependent on the presence of free sulfhydryl (SH) groups for their activity. Lead interacts with sulfhydryl groups in such a way that they are not available to certain enzymes that require them. In the living organism, under most conditions, this inhibition is apparently partial. Inhibitory effects of lead on other aspects of cellular metabolism have been demonstrated in the test tube. Such studies are preliminary. Most of the effects reported are produced with concentrations of lead considerably higher than are likely to be encountered in the tissues of man, so that speculation about such effects is unwarranted at this point.

The clearest manifestation of the inhibitory effect of lead on the activity of sulfhydryl-dependent enzymes is the disturbance it causes in the biosynthesis of heme. Heme is the iron-containing constituent that combines with protein to form hemoglobin, the oxygen-carrying pigment of the red blood cells. Heme is also an essential constituent of the other respiratory pigments, the cytochromes, which play key roles in energy metabolism. The normal pathway of heme synthesis begins with an isolated succinate (produced by the Krebs cycle, a major stage in the conversion of food energy to

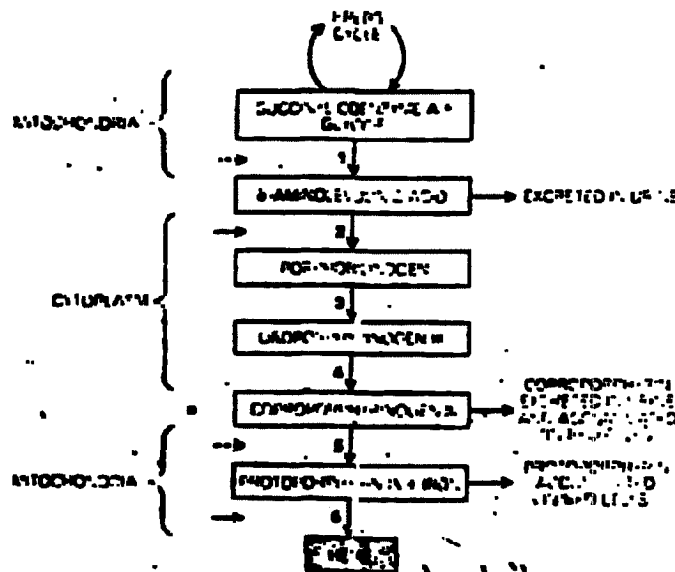
PATIENTS	LEAD OUTPUT (MILLIGRAMS PER 24 HOURS)		
	MEAN	MEDIAN	RANGE
UNEXPOSED CONTROLS	132	157	012-175
HOUSEHOLD CONTROLS	832	651	017-153
INCREASED LEAD ABSORPTION, NO SYMPTOMS	216	111	316-974
LEAD NEUTRALIZATION WITH AND WITHOUT BRAIN DAMAGE DURING EXPOSURE AFTER TREATMENT:	410 302	270 240	508-111 102-622

SECRETION OF LEAD in feces is an index of exposure to lead. These results of a study by the author and Harold E. Harrison illustrate the marked exposure seen in lead poisoning. Unexposed controls were children with no known exposure to lead. The other groups were children with increased lead absorption (high blood lead), children with lead poisoning and members of their households with neither high blood values nor overt symptoms.

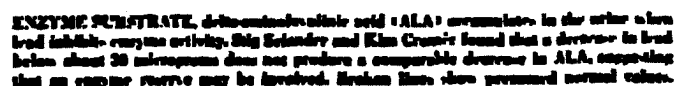
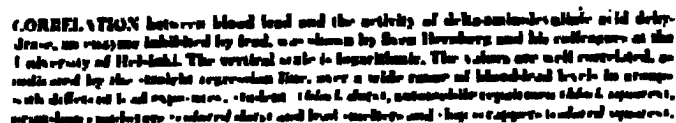
biological energy) and proceeds through a series of steps (see illustration below). Two of these steps are inhibited by the presence of lead; two others may also be inhibited, but at higher lead concentrations.

Lead is implicated specifically in the metabolism of delta-aminolevulinic acid (ALA) and in the final formation of heme from iron and protoporphyrin. Both of these steps are mediated by enzymes that are dependent on free sulfhydryl

groups for their activity and are therefore sensitive to lead. The two steps at which lead may possibly be implicated are the formation of ALA and the conversion of coproporphyrinogen to protoporphyrin. Although the exact mechanism is not known, coproporphyrin (an oxidized product of coproporphyrinogen) accumulates in the urine and the red cells in lead poisoning. Whatever the mechanism, the increased excretion of ALA and coproporphyrin is almost al-



BIOSYNTHESIS OF HEME, a constituent of hemoglobin, is inhibited by lead, resulting in accumulation of intermediates in the synthetic pathway. Of six steps in the pathway, the first and the last two take place in mitochondria, the others elsewhere in the cell cytoplasm. Lead inhibits two steps (solid colored arrows) and may inhibit two others (broken arrows).



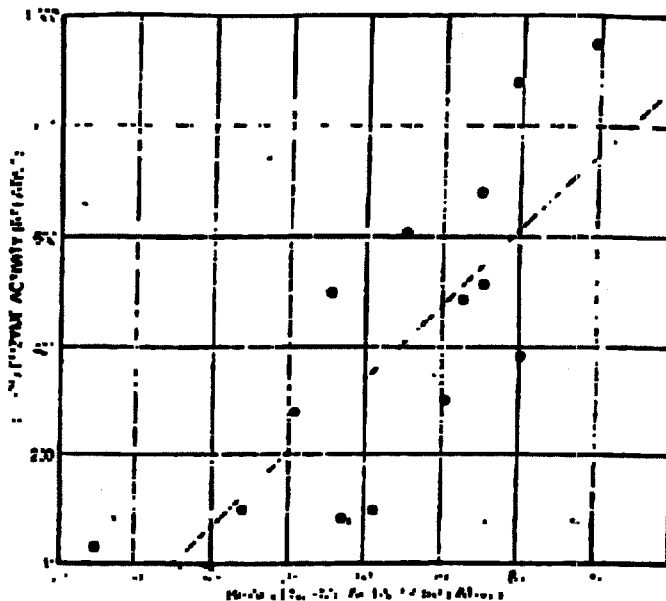
This does not seem to be the case. Sug Ratanjy and Kim Cramer in Sweden, comparing blood lead and urine-ALA values, found that the first measurable increase in urine ALA is observed only after blood lead rises above approximately 30 micrograms of lead per 100 milliliters of whole blood [see bottom illustration at left]. The apparent discrepancy between the effect of lead on the activity of an enzyme in the test tube and the accumulation of the enzyme's substrate in the body might be explained by the presence of an enzyme reserve. This hypothesis is consistent with the functional reserve exhibited in many human enzymes.

Almost all the information we have on the effect of lead on the synthesis of heme comes from observations of red blood cells. Yet all cells synthesize their own heme-containing enzymes, notably the cytochromes, and ALA dehydratase is also widely distributed in tissues. The observations on red blood cells may therefore serve as a model of lead's possible effects on heme synthesis in other organ systems. Even so, the degree of inhibition in a given tissue may vary and will depend on the concentration of lead within the cell, on its access to the heme synthetic pathway and on other factors. For example, J. A. Miller and his colleagues in Goldberg's group found that ALA-dehydratase activity is inhibited in the brain tissue of heavily lead-poisoned laboratory rats at about the same rate as it is in the blood (see blood section on opposite page). When these workers used amounts of lead that produced an aver-

age blood-lead level of 30 micrograms per 100 milliliters of blood, the level of ALA-dehydrase activity in the brain did not differ significantly from the levels found in control rats that had not been given any added lead at all. It is now established experimentally that lead does interfere with heme synthesis in those preparations from the kidney, the brain and the liver as well as in red cells but the concentrations of lead that may begin to cause significant inhibition in these organs are not yet known.

Only in the blood is it as yet possible to see a direct cause-and-effect relation between the metabolic disturbance and the functional disturbance in animals or people. In the blood the functional effect is obvious. The decrease in heme synthesis leads at first to a decrease in the life-span of red cells and later to a decrease in the number of red cells and in the amount of hemoglobin per cell. In compensation for the shortage, the blood-forming tissue steps up its production of red cells: immature red cells, reticulocytes and haemoglobin-stuffed cells (known as their stippled appearance after absorbing a basic dye) appear in the circulation. The presence of stippled cells is the most characteristic finding in the blood of a patient with lead poisoning. The stippling represents remnants of the cytoplasmic constituents of red cell precursors, including mitochondria. Normal mature red cells do not contain mitochondria. The anemia of lead poisoning is a reversible condition: the metabolism of heme returns to normal, and the anemia improves with removal of the patient from exposure to excessive amounts of lead.

The toxic effect of lead on the kidneys is under intensive investigation but here the story is less clear. In acute lead poisoning there are visible changes in the kidney and kidney function is impaired. Again the mitochondria are implicated: their structure is visibly changed. Much of the acute lead is concentrated in the form of dense inclusion in the nuclei of certain cells, including those lining the proximal renal tubules. Robert A. Goyer of the University of North Carolina School of Medicine isolated and analyzed these inclusions and found that they consist of a complex of protein and lead (see upper illustration on page 18). He has suggested that the inclusions are a protective device: they tend to keep the lead in the nucleus, away from the vulnerable mitochondria. Involvement of the mitochondria is also suggested by the fact that lead-poisoned kidney cells consume more oxygen than normal cells in laboratory cultures, which indicates



CORRELATION between the activity of ALA dehydrase in the blood and in the brain of normal rats (solid dots) and lead-poisoned rats (open circles) suggests that the enzyme may be implicated in brain damage, according to J. A. Miller and his colleagues. These data are for severely poisoned rats; in others with blood-lead values of about 30 micrograms per 100 milliliters of blood, brain enzyme activity was not significantly less than in controls.

that their energy metabolism is affected.

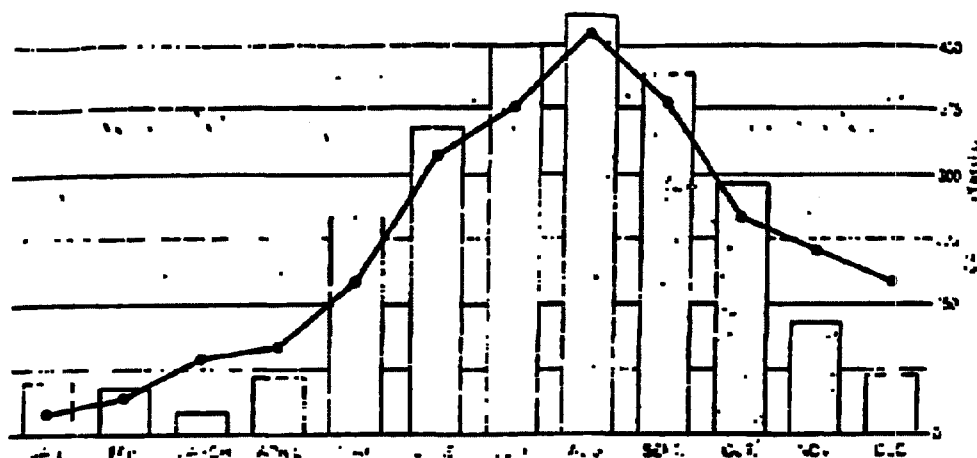
Kidney dysfunction, apparently due to this impairment to energy metabolism, is expressed in what is called the Fournier syndrome: there is an increased loss of amino acids, glucose and phosphate in the urine because the damaged tubular cells fail to reabsorb these substances as completely as normal tubular cells do. The excessive excretion of phosphates is the important factor because it leads to hypophosphatemia, a low level of phosphate in the blood. There is considerable evidence that, when phosphate is mobilized from bone for the purpose of maintaining an adequate level in body fluids, lead that is stored with relative safety in the bones may be mobilized along with the phosphate and enter the soft tissues where it can do harm. The effect of acute lead poisoning on the kidney can be serious but, like the effect on blood cells, it is reversible with removal of abnormal exposure. Furthermore, the Fournier syndrome is seen only at very high levels of lead in blood (greater than 150 micrograms of lead per 100 milliliters of blood) and only in patients with severe acute poisoning.

In the central nervous system the toxic effect of lead is least understood. Little

is known at the metabolic level; most of the information comes from clinical observation of patients and from post-mortem studies. Two different mechanisms appear to be involved in lead neuropathy, or brain damage: edema and direct injury to nerve cells. The walls of the blood vessels are somehow affected so that the capillaries become too permeable; they leak, causing edema (swelling of the brain tissue). Since the brain is enclosed in a rigid container, the skull, severe swelling destroys brain tissue. Moreover, it appears that certain brain cells may be directly injured, or their function inhibited, by lead.

The effects I have been discussing are all those of acute lead poisoning, the result of a large accumulation of lead in a relatively short time. There are chronic effects too, either the aftereffects of acute poisoning or the result of a more buildup of a burden of lead over a period of years. The first-known effect is chronic nephritis, a disease characterized by a slowing and shrinking of kidney tissue. This complication of lead poisoning came to light in Australia in 1929, when L. J. J. Mee became aware of a pattern of chronic nephritis and early death in

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SEASONAL PATTERN of lead-poisoning cases is striking. The bars show the average number of cases reported monthly in Illinois

more from 1951 through 1953 (numbers at left). Curve shows cases reported monthly in New York City last year (numbers at right).

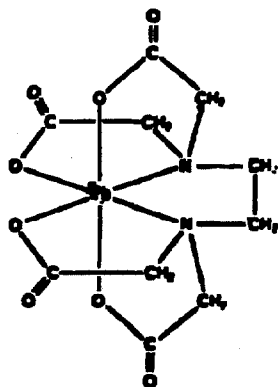
the state of Queensland. Investigation revealed that Queensland children drank quantities of rainwater that was collected by runoff from loose roof's shingles with sloughs covered with lead-jungles. In 1971 D. A. Henderson found that of 332 adults in Queensland who had had childhood lead poisoning 15 to 40 years earlier, 165 had died, 94 of chronic nephritis. Glaucoma had neuropathy, which is sometimes accompanied by great, is also seen in peripheral, heavy metal-line disorders and in some people who have had severe industrial exposure. In all these cases, however,

the abnormal intake of lead prevails for more than a decade or so before the onset of neuropathy. Most of the patients have a history of reported episodes of acute poisoning, which suggests that they have levels of lead in the tissues far above those found in the general population. Furthermore, there is the suspicion that factors in addition to lead may be involved.

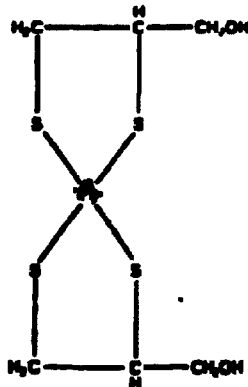
The other known result of chronic overexposure to lead is peripheral nerve disease, affecting primarily the motor nerves of the extremities. Here the tissue damage appears to be to the myelin

sheath of the nerve fiber. Specifically, according to animal studies, the motor-nerve sheath of the Schwann cells, which synthesize the sheath, seem to be affected. Various investigators, including Pamela Fothergill of Middlesbrough Hospital in London, have found that conduction of the nerve impulses may be impaired in the peripheral nerves of industrial workers who have had a long exposure to lead but who have no symptoms of acute lead poisoning.

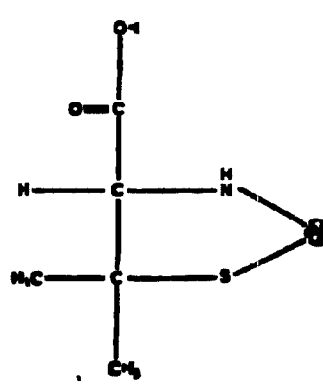
These findings and others make the case a question. It is clear that a single attack of acute neuropathy can cause pro-



CHELATING AGENTS used in treating lead poisoning bind lead atoms (Pb) firmly in one or more six-membered chelate rings. This



grams show lead chelates formed by EDTA (left), BAL (middle) and d-penicillamine (right). The last structure is still hypothetical.



found mental retardation and other forms of neurological injury that is permanent. Similarly, in young children repeated bouts of symptomatic plumbism can result in permanent brain damage ranging from subtle learning deficits to judgment, cerebral invagination and epilepsy. Can a level of absorption that is insufficient to cause obvious acute symptoms nevertheless cause "latent" brain damage? This question remains unanswered, in part because of the difficulty in recognizing mild symptoms of lead poisoning in children and in part because the experimental studies that might provide some answers have not yet been undertaken.

Central plumbism—the acute disorder—now opens today primarily in children with the pica habit. Before discussing these cases in some detail I shall briefly take up two other current environmental sources of lead: earthenware improperly glazed with lead and lead-contaminated alcoholic beverages.

Michael Klein and his colleagues at McGill University recently reported two cases of childhood lead poisoning, one of which was fatal, that they traced to an earthenware jug in which the children's mother kept a continuously replenished supply of apple juice. The slightly acidic juice was leaching lead out of the glaze, the thin layer of glossy material fused to the ceramic surface of the jug. The investigators then analyzed 117 commercial earthenware food and beverage containers and 147 samples made with 49 different commonly used glazes in the McGill ceramics laboratory. Excessive amounts of lead—more than the U.S. maximum permissible amount for glazes of seven parts per million—was leached out of half the vessels. (The maximum permissible amount should probably be reevaluated, since just outlines of testing have not taken account of such variables as the quantity of the lead or beverage consumed, its acidity, the length of time it is stored and whether or not it is cooked in the pot.) As the McGill report points out, the danger of poisoning from lead-glazed pottery has been well-known practically since antiquity. The Greeks knew about the danger but the Romans did not; they made the mistake of storing wine in earthenware. James Lind, who in 1753 recommended brass or flow pipe as a preservative for squary, also warned that the pipes should not be stored in earthenware jugs. Now the index of suspicion has fallen too low: one physician poisoned himself recently by drinking a cold beverage (and 3.2 milligrams of

POPULATION	EXPOSURE (MICROGRAMS PER CUBIC METER OF AIR)	MEAN BLOOD LEAD (MICROGRAMS PER 100 GRAMS)
RURAL U.S.	0.5	0.5
INDUSTRIAL U.S.	1.5	1.5
CONCRETE PC. CEMENT	1.5	1.5
CONCRETE TRAFFIC CONCRETE	1.5	1.5
INDUSTRIAL U.S. CONCRETE	1.5	1.5
INDUSTRIAL U.S. CONCRETE	1.5	1.5

RESPIRATORY EXPOSURE to lead is reflected in the mean blood-lead values of various groups, according to John H. Goldsmith and Alfred C. Hester of the California Department of Public Health. Groups apparently exposed to more lead in the air have generally higher blood-lead values; whether these indicate higher body burdens of lead is not known.

lead) every evening for two years (even a long life span had made for him. Do these cases represent isolated instances? How many other people are unduly exposed? Clearly the first step is the testing of earthenware and a reevaluation of its fabrication and use for food and drink.

In the manufacture of medicinal whiskey, lead solder is used in the tubing of distillation units. Moreover, discarded automobile radiators that contain lead often serve as condensers. Lead is therefore found in many samples of commercial medicinal whiskey. Lead neurotoxicity, especially with food and other lead-related substances, have been reported in medicinal consumers, largely in the southeastern part of the U.S. The problem of diagnosis is complicated by the fact that the symptoms of acute alcoholism and acute lead poisoning are similar in many ways. (Again there is a historical record. The McGill report noted that the Massachusetts Bay Colony failed to use distillation in leaded stills in 1723 as an effort to prevent "dry gripes," an intestinal condition. In 1767 Sir George Baker blamed "the endemic colic of Devonshire" on the use of lead-lined troughs in the making of apple cider.)

Childhood lead poisoning in the U.S. is now almost exclusively in children of prewar age who live in deteriorated housing built before 1910 (when the nation double began to replace lead in the paint of many urban areas). The cumulative factors are, however, a thick atmosphere of lead, a toddler with pica and parents with inadequate resources (emotional, intellectual, informa-

tional and/or economic) to cope with the family's needs. The three factors interact to increase the likelihood that the child will eat chips of leaded paint. A chip of paint about the size of an adult's thumbnail can contain between 50 and 100 milligrams of lead, and as a child eating a few small chips a day easily ingests 100 or more times the tolerable adult intake of the metal. In one study conducted some years ago at the Baltimore City Hospital and the Johns Hopkins Hospital, Harold E. Harrison and I found that the average daily lead excretion of lead by children with severe plumbism was 44 milligrams. In a group of normal unexposed children we found a daily lead excretion of less than 3 milligrams of lead. In other words, pica for leaded paint results in grossly massive exposure. And when the abnormal intake ceases, it may be several months or years before blood-lead levels return to normal.

The reported ingestion of leaded paint chips for about three months no longer can lead to clinical symptoms and eventually to the absorption of a potentially lethal body burden of lead. During the first four to six weeks of abnormal ingestion there are no symptoms. After a few weeks minor symptoms such as decreased appetite, irritability, constipation, unwillingness to play, fatigue, headache, abdominal pain and vomiting begin to appear. Then, of course, are all acute neurologic symptoms, easily treated as infectious diseases in children as well as childhood diseases. In a few weeks the lethargy may progress to incontinence and stupor; the vomiting may become persistent and forceful.

Thus patients of fulminating erythematism in conjunction with diffuse hives between 15 and 30 months of age; older children tend to suffer recurrent but less severe acute episodes and are usually brought to the hospital with a history of spasmodic convulsions, behavior problems, hyperactivity or mental retardation. The symptoms tend to wax and wane, usually becoming more severe in summer. (Some 65 percent of all head-painful cases are reported from May through October. This remarkably clear seasonal pattern is still not understood. It may be due at least in part to the fact that the ultraviolet component of sunlight increases the absorption of food from the intestine.)

The symptoms of even acute encephalopathy are nonspecific, resembling those of brain abscesses and tumors and of viral and bacterial infections of the brain. Diagnosis depends, first of all, on a high level of suspicion. To make a positive diagnosis it is necessary to show high lead absorption as well as the adverse effects of lead. This requires the measurement of lead in blood and other

found in the presence of values of between 80 and 800 micrograms of lead per 100 milliliters of blood. As the blood-lead level rises above 80 micrograms the risk of acute symptoms increases sharply. Even in the absence of symptoms, in children blood-lead levels exceeding 80 micrograms call for immediate treatment and separation of the child from the source of lead.

Treatment is with gastric compounds known as chelating agents (from the Greek *chele*, meaning claw): molecules that tend to bind a metal atom firmly, overpowering it and then rendering it highly mobile (see "Chelation," by Harold F. Walton; SCIENTIFIC AMERICAN, June, 1955). Chelating agents remove lead atoms from tissues for excretion through the kidney and through the liver. With chelating agents very high blood levels of lead can be rapidly reduced to levels approaching normal, and the adverse metabolic effects can be promptly suppressed. Initially two agents are administered by injection: EDTA and BAL. (EDTA, or ethylenediaminetetraacetic acid; BAL is "British Anti-Lewisite," developed during World War II as an antidote for arsenic, an arsenic-containing poison gas.) After the blood level has been re-

may be administered orally as a follow-up therapy.

Before these new agents were available, about two-third of all children with lead encephalopathy died. Now the mortality rate is less than 5 percent. Unfortunately the improvement in therapy has not substantially reduced the incidence of brain damage in the survivors. Meyer A. Perleman and R. Attala of the Northwestern University Medical School found that of 20 children with developmental encephalopathy, 62 percent were left with permanent injury: mental retardation, convulsive disorders, cerebral palsy or blindness. This high incidence of permanent damage suggests that some of these children could have had occasional episodes of phlebotomy; we have found that if a child who has been treated for acute encephalopathy is returned to the same hazardous environment, the risk of permanent brain damage rises to virtually 100 percent. In Baltimore, with the help of the Health Department and through the efforts of dedicated medical social workers, we are able to make it an absolute rule that no victim of lead poisoning is ever returned to a dangerous environment. The child goes from the hospital to a convalescent home and does not return to his family until all hazardous lead

	I NO DEMONSTRABLE EFFECTS	II MINIMAL SUBCLINICAL EFFECTS DETECTABLE	III COMPENSATION	IV FUNCTIONAL INJURY (SHORT, INTENSE EXPOSURE)
HEARING LOSS	NO HEARING LOSS	HEARING LOSS DETECTABLE BY AURAL TONOMETRY	HEARING LOSS DETECTABLE BY AURAL TONOMETRY	HEARING LOSS DETECTABLE BY AURAL TONOMETRY
REDUCED RED CELL LIFE-SPAN INCREASED PRODUCTION	NO EFFECT	NO EFFECT	REDUCED RED CELL LIFE-SPAN INCREASED PRODUCTION	REDUCED RED CELL LIFE-SPAN INCREASED PRODUCTION
REDUCED FUNCTION	NO EFFECT	NO EFFECT	SOMETIMES FUNCTIONAL DYSFUNCTION	FUNCTIONAL DYSFUNCTION (REVERSIBLE)
PERMANENT DAMAGE TO BRAIN	NO EFFECT	NO EFFECT	NO EFFECT	PERMANENT DAMAGE TO BRAIN
PERMANENT EFFECTS	NO EFFECT	NO EFFECT	NO EFFECT	PERMANENT EFFECTS
NO EFFECT	NO EFFECT	NO EFFECT	SOMETIMES MILD EFFECTS DETECTABLE BY AURAL TONOMETRY	ANALYSIS OF CASES OF PERMANENT DAMAGE TO BRAIN
PERMANENT EFFECTS	NO EFFECT	NO EFFECT	NO EFFECT	RANGE FROM MINIMAL LEARNING DISABILITY TO PROFOUND MENTAL AND BEHAVIORAL DEFICIENCY, CONVULSIVE DISORDERS, EPILEPSY

• **EFFECTS OF LEAD** are associated in a general way with the levels of exposure and rates of absorption of the metal. Level I is associated with blood-lead concentrations of less than 50 micrograms of lead per 100 milliliters and Level II with the 50-80 microgram range. Level III, at which compensatory mechanisms apparently

minibus or prevent driver functional injury, may be correlated with concentrations of between 30 and 100 micrograms. Level IV is usually associated with concentrations greater than 60 micrograms but impairment may be evident at lower levels, particularly if compensatory resources are impaired with a range other disease state.

Baltimore has taken a "case-finding" approach to these tests. Free diagnostic surveys were established by the city Health Department in the 1930's. Physicians took advantage of the service, and increasing numbers of cures were discovered. Since 1951 the removal of leaded paint has been required in any dwelling where a child is found with a blood-lead value of more than 60 micrograms. The number of cures reported each year rose for some time as diagnostic methods and cures improved, but recently it has leveled off. In order to reach children before they are poisoned, however, more is required than

Since World War II the incidence of food poisoning (usually in the form of

With regard to child labor and prisoning, however, we know enough to act. It is imperative for a human society to fail to do what is necessary to eliminate a wholly preventable disease.

What one can say is that the risk of functional injury increases as the concentration of lead in the blood exceeds 50 micrograms per 100 milliliters. The residual effects persist after blood lead levels return to normal.