Toxic Neuropathies Associated with Pharmaceutic and Industrial Agents

Zachary London, MD, James W. Albers, MD, PhD*

Department of Neurology, University of Michigan, 1324 Taubman Center,
Ann Arbor, MI 48109-0322, USA

The diagnosis of peripheral neuropathy can be established readily on the basis of clinical and electrodiagnostic criteria. Clinicians generally think of neuropathy, as a manifestation of an underlying systemic disorder, such as diabetes mellitus, or as a hereditary disorder of myelination. Investigations aimed at identifying the source of a neuropathy are fueled appropriately by the desire to find a reversible cause or at least to provide information about prognosis. The most common causes of peripheral neuropathy are genetic, inflammatory, or systemic, and many of these conditions can be diagnosed with simple blood tests.

In up to 20% of cases of neuropathy, however, the standard battery of laboratory tests is unrevealing. In these instances, it is natural for patients and physicians to wonder whether or not an unsuspected toxicant could be responsible. This suspicion can be heightened when patients have

* Corresponding author.

E-mail address: jwalbers@med.umich.edu (J.W. Albers).
unexplained systemic manifestations or when other close contacts also have developed symptoms of neuropathy. In other cases, exposure to a specific agent is suspected as the cause of peripheral neuropathy. At times like this, it can be tempting to equate exposure (or opportunity for exposure) with causation. Patients may be anxious to know if they need to take steps to eliminate an ongoing industrial or environmental exposure, and in some cases litigation is involved. Perhaps an even more common situation is patients who have idiopathic neuropathy and who undergo exhaustive work-up and are found to have abnormal levels of an unsuspected toxicant. For example, routine screening for urine arsenic is a common practice, and elevated levels periodically are discovered in patients who do not have a known exposure. The physician then is left with the task of determining if these values represent true toxicity.

In all of these situations, it helpful to have an understanding of the clinical, laboratory, and electrodiagnostic features of specific toxic neuropathies. Whenever a toxic neuropathy is suspected, it also is essential to understand the scientific methodology necessary to verify or refute this suspicion. The purpose of this article is to discuss the clinical investigation of a suspected toxic neuropathy, to review some of the more common or representative toxicants, and to identify the methods for establishing causation.

**The clinical evaluation of suspected toxic neuropathy**

Perhaps the most compelling reason to be familiar with the various toxic neuropathies is that these conditions are, by nature, reversible with removal of the offending agent. Thus, even though these conditions are rare, there is an urgency to arrive at the correct diagnosis so measures can be taken to eliminate ongoing toxic exposure. Early diagnosis also can identify other individuals who may be at risk. Unfortunately, identifying a toxicant as the causative agent for a patient’s neuropathy can be a difficult task, as there are no neurologic or electrodiagnostic features that distinguish toxic neuropathy from other causes of peripheral neuropathy reliably. Thus, toxic neuropathies always must be considered once a clinical diagnosis of neuropathy is established.

The first step in working up a suspected toxic neuropathy is to establish that a patient does, in fact, have clinical and electrodiagnostic evidence of peripheral neuropathy. The history should focus on sensory, motor, and autonomic complaints, including temporal profile, magnitude, and description of the symptoms. Most toxic neuropathies involve the longest and largest axons, causing numbness, paresthesias, or weakness in a stocking or stocking-glove distribution [1].

If a toxic neuropathy is suspected based on a patient’s symptoms and signs and an apparent lack of a systemic or hereditary cause, the diagnostic yield can be increased by asking directed questions about potential
pharmaceutic, industrial, recreational, or environmental exposures. It is useful particularly to become familiar with common pharmaceutic agents that can be associated with neuropathy. Examples include amitriptyline, cimetidine, colchicine, dapsone, disulfiram, ethambutol, gold, hydralazine, isoniazid, lithium, paclitaxel, phenytoin, nitrofurantoin, metronidazole, thalidomide, and vincristine. It also is important to ask about over-the-counter preparations and vitamins, because some agents, such as the essential vitamin, pyridoxine, can be potent neurotoxicants at high doses. It is also worthwhile to ask about potential occupational exposures, including specific chemicals, whenever possible. Industrial exposures have become increasingly uncommon compared with their reported frequency in the early twentieth century. This almost certainly is the result of increased awareness of the neurotoxic properties of the chemicals used in manufacturing and the institution of preventive measures to reduce exposure. Exposures to known neurotoxicants still occur, however, often in epidemic form, and new industrial chemicals that may cause neuropathy periodically are introduced into society.

It is important, also, to solicit information about recreational exposures, such as use of chemicals in hobbies. Recreational drugs, such as alcohol, nitrous oxide, and n-hexane (from sniffing glue) are well-established causes of neuropathy. These causes of neuropathy easily can be missed, because patients may hesitate to volunteer information about recreational drug use. Even a negative response to direct questioning does not exclude such agents from further consideration.

Finally, it is important to take a thorough review of systems and perform a thorough general examination, because many neurotoxicants also cause systemic toxicity (Table 1). For example, arsenic poisoning can cause abnormal skin pigmentation and nail abnormalities (Mees’ lines), and thallium intoxication can cause alopecia. Often, patients complain of nonspecific systems suggestive of gastrointestinal, cardiovascular, hepatic, or renal toxicity. These features should raise clinical suspicion for a toxic neuropathy, even though they may not suggest a specific toxicant.

Physical examination should focus foremost on findings that are suggestive of peripheral neuropathy. Objective findings, such as motor weakness and loss of reflexes, are more significant than subjective findings, such as sensory loss, without supportive evidence of impaired sensation (eg, a positive Romberg’s sign). Examination of the skin, hair, and nails may help identify features of systemic toxicity. Most patients found to have toxic neuropathy do not exhibit any of these cardinal features. Furthermore, features that are believed the most suggestive of a toxic exposure, such as Mees’ lines, often appear long after patients develop symptoms of neuropathy, limiting their initial diagnostic usefulness.

Electromyography (EMG) and nerve conduction studies are an essential part of the investigation for neuropathy, because they can confirm the diagnosis of peripheral neuropathy and give valuable information about
pathophysiology and severity [2]. Nerve conduction studies, in particular, provide a degree of information that cannot be determined on a clinical basis alone. First, nerve conduction studies can determine whether or not the neuropathy involves sensory axons, motor axons, or both. Second, they can determine whether or not there is conduction slowing. Any neuropathy that causes loss of large-caliber myelinated axons can cause mildly reduced conduction velocities, but conduction slowing out of proportion to axonal loss can help focus the differential diagnosis significantly. Third, nerve conduction studies can identify the pattern of neuropathy, such as a typical stocking-glove polyneuropathy or a mononeuritis multiplex. Needle EMG is of

<table>
<thead>
<tr>
<th>Neurotoxicant</th>
<th>Systemic feature</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acrylamide</td>
<td>Irritant dermatitis, palmar erythema, desquamation, hyperhydrosis, axonal swellings</td>
</tr>
<tr>
<td>Arsenic</td>
<td>Gastrointestinal symptoms, hyperpigmentation, hyperkeratosis, Mees’ lines, cardiomyopathy, hepatomegaly, renal failure, anemia, basophilic stippling of red blood cells</td>
</tr>
<tr>
<td>Colchicine</td>
<td>Myopathy (neuromyopathy)</td>
</tr>
<tr>
<td>Dapsone</td>
<td>After decades of use, possibly slow acetylators</td>
</tr>
<tr>
<td>Ethyl alcohol</td>
<td>Nutritional factors, Wernicke’s syndrome (dementia, ophthalmoplegia, and ataxia), midline cerebellar degeneration, abnormal liver function, cirrhosis</td>
</tr>
<tr>
<td>n-hexane</td>
<td>Irritant dermatitis, axonal swellings</td>
</tr>
<tr>
<td>Lead</td>
<td>Gastrointestinal symptoms, musculoskeletal complaints, weight loss, gum lead line, bone lead line, Mees’ lines, renal failure, anemia, basophilic stippling of red blood cells</td>
</tr>
<tr>
<td>Lithium</td>
<td>Postural tremor</td>
</tr>
<tr>
<td>Mercury, elemental</td>
<td>Anorexia, gingivitis, hypersalivation, papular rash, hyperkeratosis, lens opacities, postural tremor, nephrotic syndrome, respiratory tract irritation, metal fume fever</td>
</tr>
<tr>
<td>Nitrofurantoin</td>
<td>Elderly with impaired renal function</td>
</tr>
<tr>
<td>Nitrous oxide</td>
<td>Myelopathy</td>
</tr>
<tr>
<td>Organophosphate pesticides</td>
<td>Irritant dermatitis, acute cholinergic effects, corticospinal tract residua, noncardiogenic pulmonary edema</td>
</tr>
<tr>
<td>Phenytoin</td>
<td>Gingival hyperplasia, cerebellar ataxia</td>
</tr>
<tr>
<td>Thallium</td>
<td>Gastrointestinal symptoms, irritant dermatitis, alopecia, noncardiogenic pulmonary edema</td>
</tr>
<tr>
<td>Trichloroethylene</td>
<td>Vasodilation with ethanol ingestion, irritant dermatitis, elevated liver function tests, cirrhosis</td>
</tr>
<tr>
<td>Toluene</td>
<td>Respiratory tract irritation, irritant dermatitis</td>
</tr>
<tr>
<td>L-tryptophan</td>
<td>Peau d’orange, eosinophilia</td>
</tr>
</tbody>
</table>

secondary importance in the evaluation of neuropathy. It is useful predomi-
antly for identifying the degree of axonal involvement and ruling out pol-
yradiculopathy, which can mimic peripheral neuropathy.

Many neurotoxicants cause similar clinical manifestations, but electro-
diagnostic testing can be used to divide the neuropathies produced by neuro-
toxicants into broad categories. These categories are not exclusive, and some
toxicants can cause more than one type of neuropathy, often depending on
whether or not patients are subjected to a massive acute exposure or a low-
dose chronic exposure. Nevertheless, this classification scheme is useful be-
cause it incorporates the suggested pathophysiology of the abnormality and
can reduce an exhaustive list of potential neurotoxicants to a more manage-
able differential. These major categories include motor or motor greater
than sensory neuropathy with or without conduction slowing, sensory neu-
ropathy or neuronopathy, sensory greater than motor neuropathy with or
without conduction slowing, and mononeuritis multiplex.

Discussion of every known toxic neuropathy is beyond the scope of this
article. Rather, it focuses on selected examples that are the most common or
most characteristic of each of the categories described.

**Motor and motor greater than sensory neuropathy, with conduction slowing**

Some neurotoxicants can mimic acute inflammatory demyelinating poly-
radiculoneuropathy (AIDP), with a motor or motor greater than sensory
neuropathy with conduction slowing. Examples include arsenic (shortly af-
ter exposure), n-hexane, amiodarone, carbon disulfide, cytosine arabinoside,
methyl n-butyl ketone, perhexiline, saxitoxin, and suramin.

**Arsenic**

Arsenic is a metalloid best known for its use as a poison in homicide and
suicide. Industrial exposure may occur in lead and copper smelting, mining,
and pesticide manufacturing [3]. Sources of environmental exposure include
tainted well-water, wood preservatives, and arsenic-contaminated fossil
fuels [4,5].

The neurotoxic effects of arsenic differ, depending on whether or not pa-
tients are subjected to an acute massive exposure or a chronic, low-level ex-
posure. High-dose exposure can lead to a syndrome that can mimic AIDP
[6]. Neuropathy begins 5 to 10 days after exposure and progresses over
weeks. As with many toxic neuropathies, there may be a “coasting” effect,
with continued progression of disease for a period of weeks after removal
from exposure. Patients can develop flaccid areflexic quadriparesis, bifacial
weakness, and even diaphragm paralysis, requiring ventilatory support [7,8].

Clinical manifestations can help distinguish arsenic toxicity from inflam-
matory demyelinating neuropathy, including a variety of systemic symp-
toms that may develop before the onset of neuropathy. Gastrointestinal
disturbance with abdominal pain and vomiting, tachycardia, and hypoten-
sion are common, although these symptoms also may be seen in AIDP [9]. Nonspecific systemic manifestations may follow, including hepatomeg-
aly, renal failure, anemia, and cardiomyopathy, which are atypical for
AIDP. Other systemic features are more specific to arsenic toxicity, such
as brownish desquamation of the hands and feet (arsenical dermatitis)
and Mees’ lines on fingernails and toenails. Unfortunately, the skin and
nail changes may not appear until a month or more after isolated ingestion
of arsenic, by which time the diagnosis rarely is still in question.

If an initial EMG is performed within days, decreased motor unit recruit-
ment may be the only finding. Over the first few weeks, EMG may show
motor greater than sensory neuropathy with reduced amplitudes, borderline–low conduction velocities, prolonged F waves, and even partial conduc-
tion block in several motor nerves. The nerve conduction studies even may
fulfill criteria for the diagnosis of an acquired demyelinating neuropathy
[10]. Cases even are reported in which the sural sensory response is normal,
but the median sensory response is absent, a finding often found in AIDP
[11]. Follow-up studies are more consistent with a typical dying-back axon-
opathy, with absent sensory and motor responses and denervation/reinner-
vation changes on needle EMG.

Chronic exposure to low levels of arsenic, such as can be seen in industrial
settings, often results in dermatologic manifestations first, including hyperpig-
mentation, hyperkeratosis, and mucosal irritation [12]. Ongoing exposure
may lead to a painful length-dependent neuropathy with proportionally fewer
motor symptoms [3].

Laboratory testing can aid in the diagnosis of arsenic poisoning in acute
and chronic settings. Urine arsenic levels greater than 25 μg in a 24-hour
specimen generally are considered abnormal, although levels may be ele-
vated falsely by the ingestion of seafood, in particular bottom-feeding fin-
fish. Small amounts of arsenic bind to keratin in growing tissues, allowing
diagnosis to be made by measuring levels in hair and nails. This is useful
particularly in the setting of chronic or low-level exposure or for detecting
a remote exposure that has since ceased. Blood arsenic levels are not helpful,
because serum arsenic is cleared within 2 to 4 hours [13].

Chelation with penicillamine or dimercaprol should be started as soon as
possible after exposure [14,15]. The usefulness of chelation in preventing
progression of acute arsenical neuropathy is unknown, however.

Hexacarbons

N-hexane and methyl-n-butyl ketone are hexacarbons used in industrial
solvents and household glues. Most of the industrial exposures producing
neuropathy occurred in the cabinet- and shoe-making industries, where hex-
acarbon solvents were used extensively until the 1970s. More recently, cases
of n-hexane neuropathy were reported in automotive technicians using
degreasing solvents and cleansers. The most common source of n-hexane exposure today is the intentional inhalation of household glues for the purpose of intoxication [16,17].

Both n-hexane and methyl-n-butyl ketone are metabolized to 2,5-hexanedione in the liver, which probably is the neurotoxic agent [18]. Substantial n-hexane toxicity causes central nervous system depression and narcosis, but with repeated exposures, peripheral neurotoxicity can develop. Specifically, n-hexane causes a neuropathy consisting of length-dependent distal sensory loss, weakness, atrophy, reduced or absent distal reflexes, and autonomic dysfunction. Among “huffers,” who inhale massive quantities of n-hexane volitionally, motor and cranial nerve symptoms may predominate. Furthermore, nerve conduction studies often demonstrate reduced amplitudes with conduction velocities in the range of a primary demyelinating disorder [19]. As with arsenic, n-hexane can create a clinical picture that mimics AIDP. Chronic, low-level exposures are associated with a more typical dying-back sensorimotor neuropathy with sensory loss, distal weakness, and absent ankle reflexes [20,21]. Chronic occupational and recreational exposures also are associated with degeneration of distal corticospinal and dorsal column pathway and impairment of color vision [22].

The classic neuropathologic finding in n-hexane neuropathy is giant axonal swellings, which consist of neurofilamentous aggregates that accumulate secondary to abnormalities of axonal transport. The slow conduction velocities found on nerve conduction studies are believed to represent secondary myelin sheath damage from these focal axonal swellings [17,23–27].

Cessation of exposure is the primary means of treatment, and in one large case series, all 102 patients who had identified cases of n-hexane intoxication recovered completely without other intervention [28].

Amiodarone

Amiodarone is a di-iodinated benzofuran derivative used for refractory or life-threatening ventricular arrhythmias. Amiodarone is associated with several neurotoxic effects, including peripheral neuropathy, action tremor, myopathy, optic neuropathy, basal ganglia dysfunction, encephalopathy, and pseudotumor cerebri.

The neuropathy associated with chronic amiodarone use varies in various reports. The clinical presentation may be a symmetric, subacute to chronic sensorimotor polyneuropathy with distal predominance [29–37]. Other investigators have reported a rapidly evolving motor-predominant neuropathy that can be difficult to distinguish from AIDP [38]. Reports of nerve conduction studies vary from a sensory-predominant axonopathy with reduced amplitudes to a demyelinating picture with prominent conduction slowing [39]. The findings in sural nerve biopsies also vary, with some specimens demonstrating axonal loss and others showing almost pure demyelination [40]. Amiodarone is highly lipophilic and forms intralysosomal
lipid complexes, leading to inclusions in multiple tissues. These inclusions are seen in neural structures, suggesting a possible mechanism of toxicity.

There is no specific treatment for amiodarone-induced neuropathy other than lowering or discontinuing the drug. Recovery can be slow after removal of the exposure, because amiodarone has a long half-life (25–100 days).

**Motor and motor greater than sensory neuropathy, without conduction slowing**

Some neurotoxicants are known to produce a motor-predominant neuropathy without conduction slowing, including certain organophosphates, vincristine and other vinea alkaloids, nitrofurantoin, cimetidine, dapsone, and possibly lead.

**Organophosphates**

Unlike most chemicals that cause neuropathy, organophosphates have been used widely because they are poisonous, rather than in spite of it. The neurotoxic properties of organophosphate compounds have led to their use as insecticides and “nerve” gases. Less toxic forms also have been used in hydraulic fluids, lubricants, fuel additives, plastic modifiers, and flame retardants. Today, most cases of acute neurotoxicity are in the setting of intentional ingestion of insecticides as a suicide attempt.

One of the most remarkable illustrations of large-scale organophosphate poisoning was the jake leg epidemic of 1930. Jamaica ginger, or jake, was an alcohol-based patent medicine that was popular among poor city-dwellers trying to circumvent Prohibition laws. To fool Prohibition chemists into thinking the medicine had a higher percentage of solids, one supplier contaminated Jamaica ginger with triorthocresyl phosphate (TOCP), an organophosphate that was believed harmless. By the time the source of the contamination was identified and removed, tens of thousands of Americans had developed disabling neuropathy [41–43].

Organophosphates exert their primary neurotoxic effect by inactivating acetylcholinesterase. This leads to an accumulation of acetylcholine in muscarinic and nicotinic receptors. Symptoms of cholinergic excess include bradycardia, salivation, nausea, bronchospasm, miosis, diarrhea, sweating, central nervous system dysfunction, muscle weakness, and fasciculations. After 1 to 4 days, an intermediate syndrome develops, characterized by weakness of proximal limb, neck, extraocular, bulbar, and respiratory muscles. This syndrome resembles myasthenia gravis and most likely represents a depolarizing blockade of neuromuscular transmission. A subacute motor greater than sensory neuropathy develops as the symptoms of the acute and intermediate syndromes resolve. Organophosphate-induced delayed neurotoxicity (OPIDN), as it has been called, occurs 7 to 21 days after exposure.
and occurs only with certain exposure to certain organophosphates. Weakness follows a length-dependent pattern, with early foot drop and weakness of hand intrinsic muscles followed by more proximal weakness. Some patients go on to develop spasticity and upper motor neuron signs, indicative of superimposed distal corticospinal tract dysfunction [44]. TOCP, the agent implicated in jake leg, is a unique organophosphate that causes OPIDN without causing overt cholinergic symptoms first [45,46].

OPIDN is a distal axonopathy that affects the peripheral nerves and the long tracts in the spinal cord. Nerve conduction studies show evidence of sensorimotor neuropathy without conduction slowing [47,48]. The pathophysiology of OPIDN is believed the result of organophosphate-induced modification of a neuronal membrane protein called neuropathy target esterase, although the exact mechanism is unknown [49,50].

**Vincristine**

Vincristine is a vinca alkaloid chemotherapeutic agent used for treatment of solid tumors, lymphoma, and leukemia. Peripheral neuropathy is the dose-limiting side effect of all vinca alkaloids, with vincristine the most neurotoxic [51]. The mechanism of vincristine neuropathy is believed related to impairment of the function of microtubules involved in axonal transport [52].

Pain and small-fiber sensory loss predominate early, usually occurring at 4 to 5 weeks [53]. Autonomic dysfunction also may occur early, with constipation, orthostatic hypotension, and impotence. Distal symmetric weakness invariably occurs with continued exposure, and in some patients, these symptoms may develop fulminantly [54]. There also are cases of vincristine therapy leading to a severe acute neuropathy by unmasking a subclinical inherited neuropathy, such as Charcot-Marie-Tooth disease type 1 or hereditary neuropathy with liability to pressure palsies [55–57]. Patients who have these neuropathies may be susceptible to developing severe weakness with low or even single doses of vincristine.

In vincristine neuropathy, the EMG demonstrates axonal neuropathy, with decreased sensory and motor amplitudes on nerve conduction studies. The presence of motor involvement on electrodiagnostic testing correlates with the degree of clinical weakness.

Muscle weakness usually recovers rapidly after the drug is discontinued. Up to two thirds of patients continue to have residual sensory symptoms and absent deep tendon reflexes, however. Electrodiagnostic measures of neuropathy may persist indefinitely, with low or absent sensory nerve action potentials.

**Sensory neuropathy or neuronopathy, without conduction slowing**

Cisplatin, pyridoxine, thallium, metronidazole, ethyl alcohol, nitrofurantoin, and thalidomide are shown to induce a pure sensory neuropathy or neuronopathy without conduction slowing.
**Cisplatin**

Cisplatin is a chemotherapeutic agent used for ovarian and small cell lung cancers that causes a cumulative dose-limiting axonal sensory neuropathy. The primary site of damage is the dorsal root ganglion (sensory neuronopathy), but the large myelinated sensory axons also may be affected [58,59]. Manifestations include numbness, paresthesias, and occasionally pain in the distal extremities, with loss of deep tendon reflexes and position sense. The most important condition to consider in the differential diagnosis is paraneoplastic sensory neuronopathy, which can present identically. Paraneoplastic neuropathies may be associated with autoantibodies in the serum and may continue to progress despite discontinuation of the cisplatin. In cisplatin neuropathy, nerve conduction studies show decreased sensory nerve action potential amplitudes and prolonged sensory latencies.

The toxicity of cisplatin can persist long after the medication is discontinued. In one study of patients who had been treated with cisplatin 13 or more years prior, 38% were found to have nonsymptomatic neuropathy, 28% symptomatic neuropathy, and 6% disabling polyneuropathy [60].

The mechanism of cisplatin neurotoxicity is unknown, but it is believed to relate to disruption of fast axonal transport and induction of apoptosis in dorsal root ganglion cells [61,62].

**Pyridoxine**

Pyridoxine, or vitamin B₆, is an essential vitamin that has neuroprotective effects when used to treat isoniazid overdose, Gyromitra mushroom or false morel (monomethylhydrazine) poisoning, and hydrazine exposure [63]. Pyridoxine also is a potent neurotoxicant, however, with low-dose, chronic exposure and with acute massive exposure. Pyridoxine toxicity produces a pure sensory neuropathy, with numbness and loss of position sense but no dysfunction of motor nerves or the central nervous system [64]. In most cases, the neuropathy is slowly reversible with discontinuation of the pyridoxine, but large acute doses may be associated with permanent profound sensory loss and pseudoathetosis [65].

**Thallium**

Thallium is a neurotoxic metal that has been used in the manufacture of optical lenses, semiconductors, scintillation counters, some fireworks, insecticides, and rodenticides [66]. The most common causes of thallium poisoning are homicidal, suicidal, or accidental ingestion of rat poison, although the number of cases has declined substantially since thallium was banned in the United States as a rodent poison in 1972 [67–70].

Thallium causes a predominantly small-fiber neuropathy, with painful dysesthesias in the distal lower extremities. Dysautonomia often is present and may precede neuropathy [71]. Reflexes often are preserved, a feature
that helps localize the lesion to the small-fiber nerve and differentiate thallium poisoning from AIDP and other toxic neuropathies. Although most cases of thallium poisoning cause sensory symptoms only, there also are reports of motor manifestations [72].

Early signs of systemic toxicity include gastrointestinal symptoms, cardiac and respiratory failure, encephalopathy, and renal insufficiency [73]. The most pathognomonic manifestation of thallium poisoning is alopecia, which appears 15 to 39 days after intoxication and, therefore, is not useful in the acute setting [74]. Mees’ lines (white stria of the nails) also may develop as a late sign.

Early in the course of the disease, there may be involvement only of the small nerve fibers. At this point, nerve conduction studies may be normal or show only mild abnormalities, such as absent plantar sensory responses. In more severe cases, evidence of axonal loss is found on electrodiagnostic testing and sural nerve biopsy [75,76].

The exact mechanism of thallium toxicity is unknown. Thallium enters cells through potassium channels and may compete with potassium in intracellular reactions and interfere with energy metabolism in the Kreb’s cycle, oxidative phosphorylation, and glycolysis [77].

**Sensory greater than motor (sensorimotor) neuropathy, with conduction slowing**

Saxitoxin and tetrodotoxin, although not pharmaceutic agents, are biologic neurotoxicants that produce a sensory greater than motor neuropathy with conduction slowing.

**Saxitoxin**

Saxitoxin, otherwise known as red tide, is the neurotoxicant implicated in paralytic shellfish poisoning. The most common source of saxitoxin poisoning is the consumption of bivalve mollusks, in particular those harvested in the months of May, June, and July [78,79]. Clinical manifestations include gastrointestinal symptoms, cerebellar ataxia, and a sensorimotor neuropathy, which may be severe enough to cause respiratory depression.

Saxitoxin exerts its effects by blocking sodium channels, reducing the local currents associated with propagation of the action potential [80]. Nerve conduction studies show prolonged distal sensory and motor latencies with slowed conduction velocities and moderately diminished response amplitudes [81].

**Tetrodotoxin**

Tetrodotoxin is a poison derived from puffer fish, where it is found in various concentrations in the liver, ovaries, intestines, and skin [82]. Fugu, or puffer fish fillet, is a delicacy in Japan, where it is the most common cause
of fatal food poisoning [83]. Clinical manifestations depend on the dose of the exposure, but neurologic symptoms and signs may range from perioral numbness to flaccid paralysis, dilated pupils, and respiratory failure without loss of consciousness. Symptoms may begin within 1 hour after ingestion of the contaminated fish and generally abate after 5 days [84].

Like saxitoxin, tetrodotoxin causes blockade of voltage-sensitive sodium channels, leading to conduction failure. Nerve conduction studies show prolongation of sensory and motor latencies, prolongation of F waves, slowing of conduction velocities, and reduced sensory and motor amplitudes [85].

**Sensory greater than motor (sensorimotor) neuropathy, without conduction slowing**

The majority of peripheral neurotoxincants produce a length-dependent sensorimotor neuropathy without conduction slowing. Some examples of neurotoxicants known to fit this pattern include acrylamide, amitriptyline, arsenic (chronic), carbon monoxide, ethambutol, ethyl alcohol, ethylene oxide, gold, hydralazine, isoniazid, lithium, elemental mercury, metronidazole, nitrofurantoin, nitrous oxide, paclitaxel, perhexiline, phenytoin, thallium, and vincristine.

**Acrylamide**

Acrylamide is a vinyl polymer used to synthesize polyacrylamide, which has many applications as a soil conditioner, flocculator, and waterproofing agent and in the cosmetic, paper, and textile industries. Although polyacrylamide is nontoxic, it can be contaminated with the toxic acrylamide, especially when it is used as a flocculator [86,87].

Acrylamide causes a typical axonal neuropathy with weakness, sensory loss, and areflexia involving primarily large axons. Systemic symptoms include irritant dermatitis, palmar erythema, and encephalopathy. Nerve conduction abnormalities generally are mild and show low amplitude sensory responses, and, to a lesser extent, low amplitude motor responses without significant conduction slowing [88]. Subclinical neuropathy also is detected in patients who have low-level industrial exposures [89,90].

The pathologic hallmark of acrylamide neurotoxicity, like that of n-hexane, is giant axonal swellings. Classically, acrylamide is believed to exert its toxicity through disruption of anterograde and retrograde axonal transport [91]. Some studies suggest, however, that the nerve terminal may be the primary site of neurotoxicity rather than the axon itself [92,93].

**Nitrofurantoin**

Nitrofurantoin is an antibacterial agent specific for urinary tract infections. Nitrofurantoin is implicated in the development of axonal sensorimotor...
neuropathy, with sensory-predominant (common) and motor-predominant (less common and typically developing after onset of the sensory neuropathy) neuropathies reported. Patients who have pre-existing renal dysfunction or diabetes mellitus are at greater risk for developing nitrofurantoin-induced neuropathy, but neuropathy also is described in otherwise healthy patients taking standard doses of this medication. [94,95]. The pathologic changes in nitrofurantoin-induced neuropathy are those of acute, severe axonal degeneration [95]. It is hypothesized that nitrofurantoin exerts its toxic effects through dose-dependent depletion of glutathione [96].

**Phenytoin**

Peripheral neuropathy long has been recognized as a side effect of chronic phenytoin use, especially at higher doses [97,98]. Although it is not well studied in prospective trials, most reports of phenytoin toxicity suggest that it probably does not live up to its reputation as a serious peripheral neurotoxicant. With standard doses and close monitoring of levels, phenytoin-induced neuropathy is rare. When present, patients usually are asymptomatic and the neuropathy can be detected only by physical examination or electrophysiologic studies [99–101].

**Mononeuritis multiplex**

Rarely, neurotoxicants can present with a clinical picture suggestive of multiple mononeuropathies, rather than a symmetric, length-dependent peripheral neuropathy. Examples include trichloroethylene, dapsone, lead, and L-tryptophan.

**Trichloroethylene**

Trichloroethylene is a chlorinated hydrocarbon that has been used as a cleaner, solvent, degreasing agent, and surgical anesthetic. It is unusual among neurotoxicants in that it is associated with a cranial mononeuritis multiplex with little evidence of sensory or sensorimotor neuropathy [102]. Patients develop ptosis, extraocular muscle dysfunction, facial and bulbar weakness, and signs of trigeminal dysfunction [103]. Like other neurotoxicants, trichloroethylene also is a systemic poison. It is shown to cause irritant dermatitis, cirrhosis, and cardiac failure [104].

In association with a trichloroethylene cranial mononeuritis multiplex, facial motor nerve distal latencies and blink reflex R1 latencies are prolonged. Blink reflexes also are used to suggest the presence of a subclinical trigeminal neuropathy in patients who have chronic, low-dose exposure to trichloroethylene through contaminated well water [105,106].

It is controversial whether or not trichloroethylene is directly toxic to nerves or if neurologic symptoms actually are the result of toxicity from...
dichloroacetylene, a metabolite of trichloroethylene that is formed only in certain conditions, such as high heat or extreme alkalinity [107]. It also is hypothesized that trichloroethylene does not actually cause a toxic neuropathy but rather triggers the reactivation of a latent herpes virus [108]. Necropsy examination of one patient showed degeneration of the brainstem nuclei and tracts, the trigeminal nerve, and cranial sensory roots [103].

**Dapsone**

Dapsone is an antiparasitic and antimycobacterial agent used to treat leprosy, toxoplasmosis, malaria, and the skin condition, dermatitis herpetiformis. Most descriptions of dapsone neuropathy are those of a motor-dominant neuropathy, often with an asymmetric presentation, suggesting a mononeuritis multiplex [109–112]. Mixed sensorimotor neuropathy and mild, sensory neuropathies also are described [113,114].

Neuropathy generally is seen only in patients who have been taking dapsone for several years, with most cases developing within 5 years of initiation. Ironically, dapsone has been used widely to treat leprosy, an infectious cause of peripheral neuropathy. In spite of this, most cases in the literature occur when patients are taking dapsone for dermatitis herpetiformis. Slow acetylators of dapsone likely are at additional risk for developing neuropathy [110,115].

Electrophysiologic and pathologic studies suggest a distal motor axonopathy without features of demyelination [116]. Most patients recover completely over the course of many months after the drug is withdrawn [115].

**Methodology used to establish causation**

The most important tenet of establishing a neurotoxic cause of any neurologic problem is that the opportunity for exposure does not prove that the symptoms were caused by the exposure. Given the widespread prevalence of peripheral neuropathy, a patient who works with industrial chemicals or takes a medication known to cause neuropathy still could develop neuropathy from a different cause. In fact, in many situations, the alternative explanation is more likely. Knowledge of the biologic effects of the toxicant and the circumstances of the exposure can help differentiate causation from mere association.

Most clinicians apply general scientific principals in the formulation of differential diagnosis without giving thought to the process, but formal criteria exist for evaluating the role of a suspected toxin in a specific case [117]. The purported effect of the toxicant needs to be biologically plausible. The relative risk of neuropathy varies significantly between neurotoxicants, and large epidemiologic studies, in particular cohort or case control studies, that demonstrate a strong association between the toxicant and neuropathy confer more support for the hypothesis than isolated case reports or
cross-sectional studies. The existence of a well-studied animal model also can be helpful in identifying potential mechanisms of neurotoxicity.

The circumstances of the exposure can provide additional information. For instance, the temporal nature of the exposure is important in establishing causation. Obviously, the neuropathy cannot precede the exposure. Conversely, a neuropathy that develops months or years after a single acute exposure is not consistent with causation.

In almost all toxic neuropathies, there should be a dose-response relationship between the level of toxicant exposure and the severity of the neuropathy. For many toxicants, biologic markers can be measured in the blood, urine, or hair. Established reference levels can be used to suggest whether or not the degree of exposure is substantial enough to account for a patient’s symptoms. Resolution of the neuropathy and normalization of the biologic markers after removal of the exposure provide some of the strongest evidence of causation. Some toxic neuropathies, however, continue to progress for a few weeks after removal of exposure before stabilizing and eventually improving, a phenomenon known as “coasting.”

The most difficult task in establishing causation is eliminating other causes from the differential diagnosis. Although a patient’s symptoms, signs, and electrodiagnostic findings may be “consistent with” a toxic neuropathy, it is likely that the same findings are “consistent with” several other causes also. Eliminating all competing causes from the differential diagnosis requires a working knowledge of systemic, genetic, inflammatory, infectious, and nutritional causes of neuropathy and other neurotoxicants that could produce a similar form of neuropathy.

Industrial, environmental, and pharmacologic causes of neuropathy are uncommon, and may account for only a small fraction of neuropathies in which no underlying cause is identified with routine tests. Most likely, physicians who attribute a neuropathy to a toxic cause as a diagnosis of exclusion simply are not generating an accurate or complete differential diagnosis. Nevertheless, it is important to become familiar with the most common and representative neurotoxicants, because toxic neuropathies are among the most treatable forms of peripheral nervous system dysfunction.

References

275

[99] Shorvon SD, Reynolds EH. Anticonvulsant peripheral neuropathy: a clinical and electrophysiological study of patients on single drug treatment with phenytoin, carbamazepine or barbiturates. J Neurol Neurosurg Psychiatry 1982;45:620–6.


