Epidemics and outbreaks of peripheral nervous system disorders: II. Toxic and nutritional causes

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Abstract
Peripheral neuropathies have various causes, both infectious and non-infectious. When we think of “epidemics”, we often refer to an infectious or even post-infectious origin. Nevertheless, the history of mankind is marked by episodes of epidemics of peripheral neuropathies of non-infectious nature, either of nutritional or toxic origin: we present here the main causes of such epidemics.

Keywords Alcohol · Beriberi · Tropical ataxic neuropathy · Lead · Arsenic · Hexacarbons

Abbreviations
ARW American revolutionary war
EMS Eosinophilia–myalgia syndrome
FDA Food and drug administration
JG Jamaica ginger
L-TRP l-Tryptophan
PN Peripheral neuropathy
PNS Peripheral nervous system
PP Pellagra preventis
TAN Tropical ataxic neuropathy
TOCP Tri-ortho-cresyl-phosphate
WHO World Health Organization

Introduction
Although many epidemics of peripheral neuropathy (PN) were due to infectious or post-infectious diseases, non-infectious disorders may also be incriminated. For example, two of the main current causes of PN worldwide are diabetes mellitus and alcohol consumption, both being considered as real epidemics (or even endemics/pandemics) from an epidemiological point of view. Other frequent forms of PN are drug-induced PN (chemotherapy, etc.). However, this paper will not deal with drug-induced and diabetic PN, but only focus on alcohol-induced PN and other lesser known causes of toxic and nutritional PN giving rise to epidemics in the past.

Alcohol consumption and peripheral neuropathy

The history of alcoholism
Since ancient times, alcoholic beverages play an important social role across the globe. The first archaeologic evidence of production and consumption of fermented beverage may go back 13,000 years (Neolithic), cereal-based beer having been developed by the Natufians in the Near East (Israel) [1]. Fermented beverages such as beer and wine were an essential part of the gracious way of life in many ancient civilizations (Egypt, Greece, India, China, Roma, etc.) [2], and even today. Primitive forms of distillation were developed in ancient Mesopotamia (first millenium BC), and some forms of distilled alcohol were probably known to the ancient Indians and Chinese, as well as ancient Greeks [3]. There was also evidence of true alcohol distillation (and the discovery of the flammable properties of alcohol vapor) in the Arabian alchemical treatises of the Middle Ages [4]. It
The main steps in the history of alcohol-induced peripheral neuropathy

It had long been observed that drinking alcoholic beverages may result in mental or physical health problems, including neurological disorders such as sensory deficits, tremors, seizure, delirium tremens or dementia [5]. In 1822, the American physician James Jackson (1777–1867) described, under the term “arthrodyinia a potta”, what could be considered as the first reports of alcohol-induced PN after consumption of ardent spirits, the patients usually presenting symptoms as follows: “It commences with pains in the lower limbs, but especially in the feet, and afterwards extend to the hands and arms”; “the pain is more severe in the feet and hands, than in the upper parts of the limbs”; “at length the hands and feet become nearly useless, the flexor muscles manifesting, as in other disease, greater power than the extensors”. He added that the effect of “abstinence from spirituous liquors of every kind, either in the form of medicine, or in any other form” is “gradual” and “great”, and used also “opium in sufficient quantities to relieve the pain at night and procure sleep” [8]. Finally, it was also probably one of the first detailed report of PN, although Jackson said that “the paralysis is to be referred to some affection of the muscles, and not to the nerves, as in common paralytic cases” [8]. A few years later, having not observed any abnormality by macroscopically studying the peripheral nerves of five patients with chronic alcoholism and neuropathic pain, Huss suggested that the origin of what he described as “hyperesthesic alcoholism” could be in the spinal cord [6]. Although Huss mentioned the symmetric pattern of these painful symptoms (described as “tinglings” and “burning”), he did not note motor weakness that was mild in his cases [6].

Etienne Lancereaux (1829–1910) was the first to thoroughly study “alcoholic paralysis”, giving more details on the pathological lesions of the nervous system and being the first to suggest a link between it and his observed alterations of peripheral nerves [5, 9]. Emile Théodore Leudet (1825–1887) also studied many cases of “alcoholic paralysis” associated with “motor incoordination” (probably in part from cerebellar origin), he thought to be of spinal origin (“spinal paralysis”); however, he added that “in at least some cases of chronic alcoholism, the peripheral nervous accidents were due to an anatomically demonstrable lesions of these nerve branches”, as well as in muscles [5, 10]. The same observation of “drunkard’s” or “alcoholic paraplegia” was made in England since 1867 by Samuel Wilks (1824–1911). In 1868, Reginald Edward Thompson (1834–1912) described five patients: autopsy was performed in one case showing “limited swelling of the posterior columns” of the spinal cord and observing that the peroneal nerve was “diminished in size” [11]. In 1883, Thomas Robinson Glynn (1841–1931) noted the abolition of the patellar reflexes [12]. In 1884, Carl Franz Moeli (1849–1919) observed pathological lesions of the femoral nerves [13], Walter Baugh Hadden (1856–1893) reported lesions in sciatic nerves (with absent deep tendon reflexes) [14] and Julius Dreschfeld (1845–1907) observed such lesions in both femoral and sciatic nerves [15]. The same year, Jean-Martin Charcot (1825–1893) highlighted that such patients also frequently presented vasomotor disturbance of the lower limbs and also sometimes cognitive disorders such as loss of memory [16]. For the latter, it is probable that Charcot, like Wilks in 18 [17, 68] described some cases of what will be later called Wernicke-Korsakoff’s syndrome, a pathology...
that was fully detailed as “diffuse encephalitis” (ophthalmoplegia and drowsiness but no sign of PN) by Charles Gayet (1833–1904) [18], “polioencephalitis haemorrhagica superioris” by Carl Wernicke (1848–1905) [19], then “psychosis polyneuritica seu cerebropathia psychica toxoemica” by Sergei Korsakoff (1864–1900) [20–22]. Nowadays, we know that this complex brainstem syndrome (frequently associated with alcohol-induced PN) is caused by thiamine deficiency (Gayet-Wernicke’s encephalopathy) or the combined effects of thiamine deficiency and excessive alcohol consumption (Korsakoff syndrome) [23]. In 1885, William Marc Oettinger (1856–?) clearly considered “alcoholic paralysis” as “multiple neuritis” (polyneuropathy) with pathological lesions strictly affecting the peripheral nerves (sparing the spinal cord) [24]. Alfred Gombault (1844–1904) was one of the first to fully study the pathological stages in the peripheral nerves of patients with “alcoholic paralysis”, showing Wallerian degeneration and rapid destruction of both myelin and axons (Fig. 1) [25].

Alcohol-related PN usually presents as chronic polyneuropathies involving sensory, autonomic, and motor nerves [23]. Usually classified as a progressive, predominantly sensory axonal length-dependent polyneuropathy (with motor weakness in the more severe forms), acute/subacute PN or small fiber neuropathy were also described [23]. Another special neurological complication of alcohol consumption (even if not the only cause) was described in 1957, sometimes referred as “acrodistrophie of Bureau and Barrière” [27]. The most important risk factor for alcohol-related PN is the total lifetime dose of ethanol (although other risk factors have been identified: genetic, male gender, and type of alcohol consumed); nutritional deficiency (thiamin, niacin, cobalamin) can also exacerbate alcoholic PN [23]. The overall prevalence of polyneuropathy (whatever the cause) in the general population is around 1%, rising to 7% in the elderly: [29] the prevalence of alcohol-induced PN is high in chronic alcohol abusers (44.2–46.3%) and represents 10% amongst all patients with PN [28].

**Epidemics of nutritional peripheral neuropathy**

**Vitamin B1 deficiency**

During the first third of the seventeenth century, the Dutch physician Jacobus Bontius (1592–1631) reported on mysterious epidemics of paralysis that natives of India called “beriberi” [30], although John MacGowan (1835–1922) thought it was mentioned under the term “Kioh-Ki” in a fundamental treatise on Chinese medicine (“Huángdì Nèijīng”) dating from 2600 BC: a more detailed description of beriberi as “Kak-ke” (from the Chinese word “Kiaku” for “leg” and “ki” for “disease”), appeared in another book (“Sen-Kin-ho”) written by the Chinese physician Son-Shi-Baku (640 AD) [31, 32]. The most commonly accepted hypothesis about the word “beriberi” is that it would mean “weak, weak” in Sinhalese (from “bhayree” meaning “weak”), although alternative explanations were also suggested [33]. This disease was called “bharbari” in India and “buhr bari” (meaning “marine asthma”) in Arabia and Madras (India), later translated into “barbers” in English then “barbiers” in French [33], this later word being also used at least during two centuries in the European medical literature [31]. These various terms had unfortunate consequences that contributed to maintaining (if not increasing) the initial misunderstanding.
about beriberi. Actually, Bontius used “beriberi” only for patients with paralysis, but never used it for those with generalized edema (due to cardiac involvement), the other main symptom of the disease that Bontius, however, mentioned as “frequent in India” [30]. In fact, beriberi has long time been confused with other diseases, so various other terms were proposed to describe it, such the “hydrome asthamatico” of the Scottish Surgeon Colin Rogers (referring to the generalized edema then dyspnea observed in beriberi): [34] it was not until the beginning of the nineteenth century that “beri-beri” was linked to generalized edema [35]. At that time, many authors called the severe paralytic forms “barbiers” (in Europe) or “kak-ke” (in Japan), whereas the less severe motor forms with edema were considered as “beriberi” [35]. Nowadays, beriberi is divided into three forms: ‘dry beriberi’ (characterized by a sensorimotor, distal, axonal PN often associated with calf cramps, muscle tenderness, and burning feet), ‘wet beriberi’ (association of PN and high output congestive heart failure), and ‘infantile beriberi’ (observed between 2 and 6 months of age, it may present with the cardiac, aphonic, or pseudomeningitic forms) [36].

Despite the fact that beriberi was endemic in many areas over many centuries (especially in Asia, but also in Africa then America; Fig. 2), its origin was unknown before the turn of the twentieth century. In the seventh century, Son-Shi-Baku thought it was produced by a gaseous poison coming from the ground (under the influence of cold, heat, wind and humidity), penetrating through the

Fig. 2 Major beriberi outbreaks in the world in 1906. This map is extracted from “Jeanselme E. Le béribéri. Paris: Masson & Cie–Gauthier-Villars, 1906; pages 16-17” (freely available via the “Medical Heritage Library”)
feet, up to the legs and then to the rest of the body [35]. Many cases were also reported as “sailing polineuritis” (“|nautical beriberi|”) in the European sailboats coming from endemic areas and making long non-stop crossings without replenishing supplies of fresh food [35]. In 1872, Baron Kanehiro Takaki (1849–1920) entered the Japanese Navy where he observed many fatal cases of beriberi, a subject he became more interested in after 1880, when he came back from a 5-year stay in England where he was a student of William Willis (1837–1894). Although Albrecht Wernich (1843–1896) was probably the first to clearly suggest a causal connection between beriberi and an exclusive diet of rice [37], Takaki discovered that “the nitrogenous substances contained in the food were not sufficient to maintain nitrogen metabolism, but that the food contained too much carbohydrate” and that “finally, the greater the difference in these proportions (between nitrogen and carbon) the more beriberi occurred, and the lesser the difference the less beriberi occurred” [38]. From 1884, Takaki eliminated white rice from the Navy’s diet, tested a better balanced nitrogen–carbon diet and observed a dramatic reduction in the number of cases of beriberi (which he mistakenly attributed, 1 year later, to the nitrogen-rich diet) [39]. When barley (mixed with rice) was introduced into their rations, the toll that beriberi took on Japanese soldiers declined dramatically [40]. However, in 1883, while beriberi was still considered by many physicians as an infectious disease, João Batista de Lacerda (1846–1915) thought he found the “|Bacillus beribericus|” [41], as did Masanori Ogata who claimed he discovered “Micrococcus beribericus”; this was quickly refuted by Robert Koch (1843–1910) and Shibasaburo Kitasato (1853–1931) who failed to repeat the experiment and concluded that such organisms did not cause beriberi [40]. The Dutch physician Christiaan Eijkman (1856–1930), was initially convinced that beriberi might be of bacterial origin, and decided to study it in chickens. He accidentally discovered that chickens that were fed polished rice (purchased for the army) developed “polineuritis gallinarium”, a disease whose symptoms resembled beriberi [42]. Eijkman and Adolphe Vordermann (1844–1902) thought there was an “anti-beriberi factor”, but failed to isolate it [43]. In 1911, Casimir Funk (1884–1967) finally isolated the substance (from rice husk) he called “|amine|” (corresponding to thiamine) and thought there were many other similar substances (essential to life) he called “|vitae aminae|” (later shortened in “vitamine”) [44]. But only Eijkman was honoured with the Nobel Prize (1929) [45] for his work on beriberi [43]. Nowadays, we know that thiamine deficiency may be observed in other conditions such as chronic alcoholism, persistent vomiting, anorexia, or after bariatric surgery [36–46].

**Pellagra**

At the beginning of the eighteenth century, the first cases of a new disease called “mal de la rosa” (“rose sickness”) were reported in Asturias (Spain), first by François Thiéry (1719–?) in 17 [47, 55] then by Gaspar Casal in 1762 [48]. This disease became one of the major health and demographic problems of Italy at the end of the eighteenth century, where it was named “pellagra” (“pel” meaning “skin” and “agra” meaning “rough” in the Italian dialect from Bergamo) by Francesco Frapolli (?-1773) [49]. This affection is characterized by the “|four D’s|”: dermatitis (reddish-brown hyperkeratotic rash), diarrhea, dementia and death. Also called “plague of corn”, pellagra has been quickly linked to high maize and low protein diets, and was rampant in some parts of Europe, especially in Southern Europe, where maize became more common as a staple food during the eighteenth century, as in Northern Italy where the poor diet of the Italian peasantry was almost exclusively based on maize flour such as “polenta”. In the early 1900s, pellagra was also rampant in the southern USA [50].

Pellagra, as a disease of poverty and social inequality, was initially associated with scrobut and leprosy because of the skin lesions, and such patients (as with leprosy) were victims of social exclusion. Many theories were put forward to explain it (maize diet with little or no milk, meat or fresh vegetable supplements; spoiled or rotten grain; disease spread by insects; bad heredity; overall consequence of poverty) before the true etiology of pellagra was found in 1914–1915 by the American physician Joseph Goldberger (1874–1929), although his conclusions were only accepted nearly 20 years later, in the 1940’s (when pellagra was completely eliminated from the US): pellagra is caused by long-term insufficient niacin (also called vitamin-PP, for “|Pellagra Preventis|”) and its amino acid precursor (tryptophan intake), high-maize being usually deficient in such substances. In 1952, pellagra was also recognized as a complication of isoniazid therapy [51]. Moreover, because excessive alcohol consumption is also a known risk factor of pellagra, it may be difficult to diagnose pellagra in alcohol-dependent patients where it frequently coexists with PN (32%), Wernicke’s encephalopathy (29%) and seizure (16%) [52].

Peripheral nerve changes have been associated with pellagra, although PN seems to be more characteristic of beriberi than pellagra. Sensory disturbances (usually mild) consist of paraesthesiae and numbness. Occasionally with muscular wasting and loss of deep tendon reflexes [53]. In 1881, Jules Dejerine (1849–1917) was the first to observe extensive degeneration of the myelin sheath in the cutaneous nerves of two patients with pellagra [54]. In 1940, Samuel Alexander Kinnier Wilson (1878–1937) studied 13 patients with pellagra, reporting practically constant peripheral nerve
changes: swelling or thinning of myelin sheaths, fragmentation of axons, oedema of nerve fibers and bundles, and thickening of the epi- and peri-neurium [55]. Despite these observations, PN in pellagra may stem from other concomitant B-group vitamin deficiencies, such as thiamine [36].

**Tropical ataxic neuropathy**

Between 1883 and 1897, Henry Strachan (Senior Medical Officer in Jamaica) observed 510 cases of “a form of multiple neuritis prevalent in the West Indies” [56, 57], characterized by “numbness and burning heat in the palms of the hands and the soles of the feet”, often “accompanied by cramps”; hyperpigmentation of the skin was also sometimes observed [56]. “Impaired vision and hearing”, “extreme wasting of the muscular system” (involvement of lower then upper extremities, claw hand deformity, depressed deep tendon reflexes) and sensory deficits were observed in the more advanced stage of the disease (sometimes with respiratory impairment) [56]. The enigmatic “Strachan’s syndrome” (or “Strachan-Scott syndrome”) was characterized pathologically by variable combinations of damage to the anterior horns and posterior columns of the spinal cord, peripheral nerves (predominantly the sensory nerves; Fig. 3), optic nerves and cochlear nerves [58]. Called “tropical ataxic neuropathy” (TAN) in 19 [59] it is characterized by predominantly sensory ataxic polyneuropathy, the cases with predominant spastic paraplegia nowadays being ascribed to “Konzo” [60]. A longstanding condition in the Caribbean, but other cases have been reported in Africa, Asia and Latin America (including European prisoners of war) during the twentieth century [59]. Its nutritional origin was suggested since 1911, and thiamine/riboflavin deficiency was proposed in 1959 [59]. But a role of the consumption of roots of cassava (Manihot esculenta), one of the largest source of carbohydrates in the tropics, was also suspected since 1930: so the chronic, monotonous consumption of cassava meals associated with minimal protein supplementation was proposed to explain TAN [60].

**Cuban neuropathy**

From 1991 to 1993, 50,000 patients were after by the so-called “Cuban neuropathy” (spreading from West to East of Cuba), characterized by the subacute occurrence of symptoms including retrobulbar optic neuropathy, predominantly sensory PN and dorsolateral myeloneuropathy [61], sometimes mimicking TAN. On sural nerve biopsy, moderate or severe axonal damage was observed in 73% of the cases, with a predominant loss of myelinated fibers in 92% of the cases [61]. Evidence of thiamine deficiency was found in up to 70% of the patients, and the majority of them responded biochemically to a daily oral multivitamin supplement containing thiamine [62]. Two types of neuropathy were observed: the optic form (prevalent in men; subacute onset) and a peripheral form, (prevalent in women; predominantly sensory PN, with posterior spinal cord involvement in some cases, with or without concurrent optic neuropathy). Cuban and international scientist proposed a toxic and nutritional hypothesis to explain it (injuries to the mitochondrial oxidative phosphorylation pathway, nutritional deficiencies, excitotoxicity, and dysfunction of the blood–brain barrier) [63].

**Myanmarese neuropathy**

Between 2008 and 2014, an increasing number of Myanmarese refugees admitted in Malaysia with acute/subacute predominantly sensorimotor axonal PN was observed: most of them were malnourished, with a preceding history of starvation of 2–4 weeks before onset of symptoms [64]. This clinical picture was closed to the neurological complications of thiamine deficiency, as observed in other refugees [65] or prisoners [66], so a nutritional cause was suggested. Moreover, folate and vitamin B12 deficiencies were also detected in 31.5% of the patients, and most of them improved after supportive treatment with appropriate vitamins supplementation [64]. However, the same patients also usually had at least one additional presenting symptom such as fever, lower limb swelling, vomiting, abdominal pain, or difficulty in breathing. All these findings suggested the presence of PN related to nutrition against a backdrop of other possible
environmental factors such as infections, metabolic disorders, or exposure to unknown toxin [64].

**Epidemics of toxic peripheral neuropathy**

**Lead**

Naturally present in the Earth’s crust, traces of lead exposure have been found in Neanderthal children [67]. However, lead becomes highly toxic once mined, transformed into man-made products and dispersed throughout the environment. In the Middle East and Egypt, lead was used 6000 years ago, and lead toxicity was recognized since at least 2000 BC with endemic “chronic plumbism” (the main reported manifestations being colic, anemia and gout, although neurological manifestations are usual). This was also observed by the ancient Greeks and Romans, the latter being the first mass distributors of lead (using it for cooking utensils and pots, wine urns, plumbing and aqueducts, etc.) [68]. Later, lead-based additives were used to sweeten wine, giving epidemics of “colic” during the Middle Ages, and lead poisoning was considered as a “plague” in Europe and America between the fifteenth and eighteenth centuries [68]. Since the nineteenth century and the industrial revolution of Western countries, many workers also absorbed lead from inhalation of fine dust or through the skin [68]. Occupational exposure to lead nowadays remains a problem in developing countries, but is also still of concern in some areas of Western countries, especially in children [69].

Nicander of Colophonius (second century BC) was probably the first to report lead paralysis, and Paracelsus (1493–1541) wrote about “morbis metallicis”, but the first detailed clinical descriptions of lead paralysis were made by Louis Tanquerel des Planches (1810–1862) [70], Guillaume Duchenne de Boulogne (1806–1875) [71] then Augusta Dejerine-Klumpke (1859–1927) in the second half of the nineteenth century [72]. Lead paralysis may be focal or generalized: for the focal forms, the most common one affects the upper limbs, usually first beginning by involving common extensors of the fingers, giving the classical aspect of a patient “making the horns”, as described by Duchenne de Boulogne [71, 72].

**Arsenic**

Arsenic is known to be a traditional poison, called “poudre de succession” (“powder of inheritance”) in France under the reign of Louis XIV [73], and was also used in traditional Chinese medicine since at least 4000 years ago [74]. In Europe, it was part of the treatment of syphilis or other disorders in the nineteenth century [74]. Today, it is still widely used in industry (especially in the production of pesticides, herbicides, wood preservatives, and semiconductors) and represents a public health issue in some areas [74].

Although acute arsenic poisoning produces arterial hypertension, encephalopathy and organ failure (leading to death at high doses), chronic exposure is characterized by skin involvement (keratosis and pigmentation) and other disorders including PN [74]. One of the most famous epidemics of arsenic-induced polyneuropathy was described in England in 1900–1901: first considered as alcohol-induced PN (because “peripheral neuritis is unfortunately a very common affection in Lancashire among alcoholics, and for at least 20 years has been a prolific cause of paralysis. The subjects affected are almost invariably beer drinkers”), this mysterious epidemic (enough to fill hospitals with beer drinkers) was finally due to the consuming of unsuspected and undetected quantities of arsenic in the beer because of contaminated barley malt used unwittingly by brewers [75].

**Hexacarbon solvents**

There are many examples of outbreaks of PN due to hexacarbon solvents (such as n-hexane and methyl-n-butyl-kenone), first observed in Japan in the 1960’s then in Europe and the US. Hexacarbon exposure leads to subacute-chronic dying-back PN, sometimes with giant axons and demyelinating signs on nerve biopsy (Fig. 4) [76]. These solvents were especially used in shoe-making and the printing industry. Another common cause of hexacarbon PN is the deliberate inhalation of the vapors of lacquers or glue (“glue-sniffing”) [77].
**Tri-ortho-cresyl-phosphate**

Tri-ortho-cresyl-phosphate (TOCP) was a famous cause of an outbreak of polyneuropathy (PN) in the 1930’s. “Jamaica ginger” was an alcoholic extract of ginger available as “medicine” in US drug stores since 1863. During prohibition, some Jamaica ginger (JG) extract was legally marketable as a carminative [78]. But illicit brands (made with legal JG) were created to circumvent prohibition (especially in the Southern and Midwestern US): it was highly adulterated JG beverages (“Jake”) containing various substances, sometimes including TOCP used as a liquid plasticizer (TOCP was present in up to 2% of the beverages).

The tragic consequence of this massive poisoning was severe PN with gait disturbance, called “Jake leg”: it also inspired some American popular music as the “Jake Walk Blues” of the Allen Brothers (1930) [78]. Similar cases were observed in Sri Lanka in 1981, due to the consumption of gingili oil contaminated with TOCP [79].

**Coyotillo**

Due to the consumption of the fruits of coyotillo (Karwinskia humboldtiana), a poisonous shrub of the buckthorn family, outbreaks of “buckthorn polyneuropathy” (giving generalized flaccid paralysis clinically mimicking GBS, sometimes leading to death by respiratory failure) were observed in 106 Mexican soldiers in 1918 and 16 Mexican children in 1951 [80]. It may result from a disruption of the metabolic activity of Schwann cells [80].

**Refined aniline**

The “toxic oil syndrome” was another outbreak observed in nearly 20,000 people in Spain (1981), due to the consumption of oils containing refined aniline denatured rape seed oil: the main symptoms were eosinophilia, myalgia and pulmonary edema in the acute period, then a multisystemic disease including PN in some cases [81].

**L-Tryptophan**

During the summer of 1989, an epidemic of what was called “eosinophilia-myalgia syndrome” (EMS) was observed in New Mexico (affecting around 1500 individuals, with 36 deaths): the patients presented eosinophilia and first complained of intense myalgia, followed by chronic cutaneous lesions, progressive PN and myopathy [82]. EMS was associated with ingestion of L-tryptophan (L-TRP) contained in dietary supplements manufactured in a Japanese plant using genetically engineered bacteria: removal of L-TRP from the market was followed by swift resolution of the EMS outbreak. Since the Food and Drug Administration (FDA) lifted the L-TRP import alert in 2005, only one sporadic case of EMS was reported in 2011 [83].

**Agent orange**

Agent Orange (AO) was an herbicide used during the Vietnam War by the US military to clear plants and trees: it was named “orange” after the orange-striped barrels in which it was shipped [84]. AO was a 1:1 mixture of 2,4-dichlorophenoxyacetic (2,4-D) acid and 2,4,5-trichlorophenoxyacetic (2,4,5-T) acid, but its toxicity was mainly due to the high concentrations of toxic contaminant, 2,3,7,8 tetrachlorodibenzo[p]dioxin (dioxin). In 1961, the program code named “RANCH HAND” was approved by President John Fitzgerald Kennedy (1917–1963). The objective was to aerially disseminate herbicides in the Republic of Vietnam, with two missions: defoliation and crop destruction [84]. Six herbicides were used (Orange, Purple, Pink, Green, White and Blue), but AO was the primary defoliant: [84] between 1965 and 1970, > 40 million liters of AO were sprayed by the US Air Force in South Vietnam [85].

Health concerns about the use of AO emerged in the 1950’s, mainly in workers (in 2,4-D and 2,4,5-T plants) who developed various manifestations: chloracne and mucous membrane irritation, hepatotoxicity, neuromuscular problems, psychologic alterations and porphyria cutanea tarda [86]. In October 1978, the US Air force conducted an epidemiological study of AO exposure on their “Ranch Hand” personnel in Vietnam, finally published in December 1982 [87]. However, despite it was observed signs/symptoms consistent with PN in US veterans exposed to AO [87], it concluded that “There is insufficient evidence to support a cause and effect relationship between herbicide exposure and adverse health in the Ranch Hand group at this time” [84]. In a later study (follow-up of the “Ranch Hand veterans” between 1992 and 1997), a statistically significant increased odds of probable PN was observed but the authors remained cautious in their interpretation “about the possible association between exposure to dioxin in Operation Ranch Hand and subsequent development of peripheral neuropathy” [88]. Two years later, a Korean study observed high frequency of PN (odds ratio: 2.39) in Korean Vietnam veteran compared to non-Vietnam veterans [89]. In another Korean study, exposure to AO was “associated with a significantly higher prevalence of cancers (colon cancer, leukemia, and multiple myeloma), circulatory diseases (hypertension, cerebral infarction, and peripheral vasculopathy), neuromuscular diseases (peripheral neuropathy, multiple nerve palsy, and multiple sclerosis), skin diseases, and lipedemias” [90].
Conclusion

The causes of PN are numerous, both sporadic and epidemic. Through the writings of our forefathers, including those from ancient times, we know that many epidemics of PN have affected mankind, from various origins (infectious or not) and to a greater or lesser extent. In view of the current COVID-19 pandemic affecting the entire world, it is, therefore, highly likely that other epidemics (of whatever the cause) may re-occur, and it is to be hoped that the lessons of the past will enable us to better understand the future.

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Compliance with ethical standards

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