



Adverse drug reactions affecting PNS

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OBJECTIVES

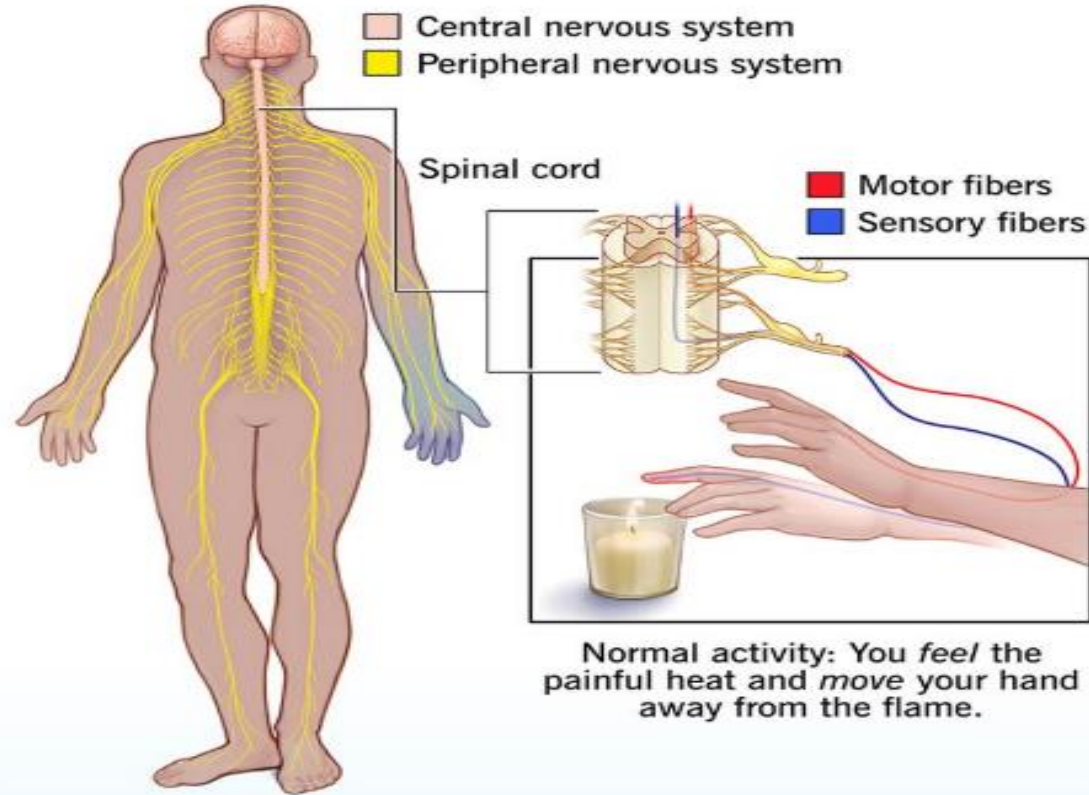
- Classification of adverse drug reactions affecting PNS
- Drug-induced peripheral neuropathy (DIPN): definition, causes, management
- Ototoxicity
- Optic neuropathy
- Neuroleptic malignant syndrome

- **Adverse drug reaction affecting PNS include:**
- 1- Drug-induced peripheral neuropathy DIPN
- 2- Ototoxicity
- 3- Optic neuropathy
- 4- Neuroleptic malignant syndrome

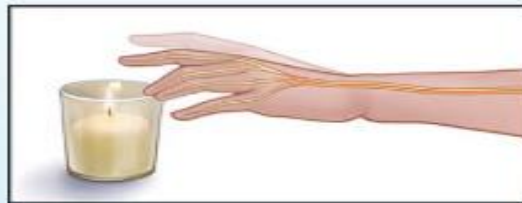
Drug-induced peripheral neuropathy (DIPN)

- **Toxic neuropathy** occurs when a chemical substance (drug) causes damage to the peripheral nervous system
 - The most common cause of **toxic neuropathy** is drug toxicity, particularly associated with chemotherapy treatments
 - **Manifestations of Toxic neuropathies are primarily characterized as:**
 - length-dependent, symmetric, sensory polyneuropathies with possible motor or autonomic involvement.
 - affect longest fibers → more vulnerable
 - distal adverse effects
 - same in both sides
- ⊗ Diabetes, cardiopathy

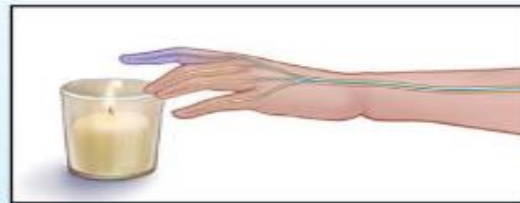
Peripheral Neuropathy



Possible symptoms of peripheral neuropathy



Muscle weakness: Your finger can barely move away from the painful flame.



Numbness: Your finger does not feel the heat of the flame.

Causes of DIPN (toxic neuropathy)

- There are more than 200 chemicals known to be neurotoxic to humans.
- There are a number of prescribed medications, including chemotherapeutic agents, that cause **neurotoxicity**
- **Peripheral neurotoxicity** can be a limiting factor in the use of many chemotherapy agents
- According to the WHO, 5.1 % of DIPN is due to alcohol.
- Some ^{الأعشاب} **herbal medicine** products commercially available have been shown to contain heavy metals such as **lead**, **mercury**, and **arsenic**.
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- Using these herbal products may lead to **heavy metal toxicity** and secondary peripheral neuropathy.

Classification of DIPN

- **Drugs associated with peripheral neuropathies**

- 1- Chemotherapeutic agents – cisplatin, paclitaxel, vincristine, bortezomib

- 2- TNF-alpha inhibitors (infliximab, etanercept)

- 3- Antiretroviral agents (stavudine)

- 4- Cardiac drugs (amiodarone, statins)

- 5- Thalidomide
 ↳ Arrhythmia ↳ hyperlipidemia

- 6- Antimicrobials (metronidazole, fluoroquinolones, isoniazid)

- 7- Disulfiram → alcohol withdrawal

- 8- Pyridoxine **high doses**

↳ Vit. B6

- 9- Colchicine → Gout

- 10- Phenytoin, Lithium

↳ Anti-epilepsy

- 11- Chloroquine, hydroxychloroquine

↳ Anti-malarial drugs



Chemotherapy-induced peripheral neuropathy

⊗ axonal degeneration → irreversible **neuropathy** → dose-dependent

Agent	Symptoms and signs	Reversible after discontinuation
<p>خفيفة →</p> <p>Bortezomib</p>	<u>Sensory, painful</u>	<u>Gradually</u>
<p>ثقيلة ←</p> <p><u>Paclitaxel (Taxol)</u></p>	<u>Paresthesias, dysesthesias (weakness), ataxia</u>	<u>Partially</u> → no complete recovery
<u>Vincristine</u>	<u>Sensorimotor and autonomic</u>	<u>Yes</u>
<u>Cisplatin</u>	<u>Sensory axonopathy, ataxia (neuronopathy), L'hermitte sign</u>	<u>May worsen after discontinuation</u>

→ abnormal unpleasant feel in touch

→ barber sign → dorsal column disorder

The relationship between the occurrence of neuropathy and the use of cytostatic drugs is usually dose-related.

If axonal or neuronal degeneration has occurred, permanent symptoms and signs remain.

Drug	Mechanism of neuropathy	Drug	Mechanism of neuropathy	Drug	Mechanism of neuropathy
Metronidazole	<u>Axonal</u> degeneration, shown to bind to neuronal <u>RNA</u>	Amiodarone	<u>Demyelination</u> , loss of large <u>axons</u>	Interferons	immune mediated <u>myelin</u> degradation, <u>vessel occlusion</u> leading to nerve ischemia
Linezolid	<u>Mitochondrial</u> toxicity	Thalidomide	<u>Mitochondrial</u> dysfunction in <u>axons</u>	Biologics	T cell and humoral immune attack on peripheral <u>myelin</u> and <u>vasculitis</u>
Vincristine ↓ same as colchicine	<u>Microtubule-mediated</u> dysfunction	Statins	<u>alterations of membrane function</u> , <u>disruption of energy utilization in nerves</u>		

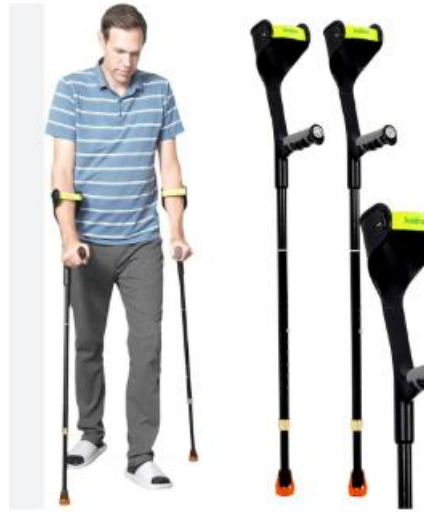
Management of toxic neuropathy

- Includes:
 - 1- Prevention
 - 2- Rehabilitation of functional impairments
 - 3- Dosage reduction or change in the drug
- **Neuropathic pain management:**
- Anticonvulsants/nerve membrane stabilizers (**gabapentin**, **pregabalin**):
- Block voltage-gated calcium channels, to inhibit the release of excitatory neurotransmitters (**glutamate and substance P**) in the presynaptic area: block pain by affecting the pain messages travelling through the brain and down the spine.
- Tricyclic antidepressants (**amitriptyline**): inhibits re-uptake of norepinephrine and serotonin, thereby increasing their concentration at the synaptic clefts of the brain.
- Serotonin-noradrenalin reuptake inhibitor (SNRI) drugs (**duloxetine**)
- Topical Capsaicin; Topical lidocaine
- Opiate analgesics and mixed opioids (**tramadol**).

Management

أدوات تقويم العظام

- **Orthoses and assistive devices**: can aide weakness and impaired balance and sensation.
- **Ankle Foot Orthosis** ^{دعامة} can serve as a brace to prevent foot drop and also aide in postural instability.
- Assistive devices such as ^{عصا} **canes**, ^{عكازات} **walkers**, **crutches**, etc. can also help with weakness and/or balance impairment.
- **Protective footwear** is of importance for anyone with sensory impairments for peripheral neuropathy to ensure no skin breakdown or wounds occur.
- **Splinting** and **casting** may also be of benefit to prevent or treat joint contractures occurring from weakness and immobilization in peripheral neuropathy.



Ototoxicity

→ can't regenerate

- **1- All aminoglycosides** are ototoxic due to destruction of the hair cell in organ of Corti.
- **Manifestations:** Affection of cochlear part causes tinnitus and hearing loss (high-pitched sound is affected first), affection of vestibular part causes nausea, vertigo and disturbance in gait.
pitch: - tone of sound → تقول صباح الخير وأنت منعدا وفرحان
- The effect may does not appear until several days after stoppage of drug administration and may progress to complete and permanent hearing loss (as no regeneration in the hair cell).
- **Streptomycin and gentamicin are more toxic to the vestibular division.**
- **Neomycin, kanamycin and amikacin are more toxic to the cochlear division.**
- **Tobramycin is toxic to both divisions.**
- **They can cross placenta causing damage to eighth cranial nerve of fetus.**
- **Factors enhance the ototoxicity of aminoglycosides: (also enhance nephrotoxicity):**
 - a) Use of high doses for long duration as ototoxicity is a dose-dependent.
 - b) Renal failure as there is failure of drug excretion.
 - c) Elder patients as more susceptible to toxicity.
 - d) Use of loop diuretics especially ethacrynic acid (less with furosemide) as they potentiate the ototoxicity of aminoglycosides

- **2- Loop diuretics:** can cause dose-related hearing loss that is usually reversible.
- It is most common in patients who have reduced renal function or who are also receiving other ototoxic agents such as aminoglycoside antibiotics.
- Ototoxicity occurs most frequently with rapid intravenous administration.
- Ethacrynic acid appears to induce ototoxicity more than do other loop diuretics

Optic neuropathy

- Phosphodiesterase type 5 (PDE-5) inhibitors, amiodarone, linezolid, isoniazid, ethambutol.
- PDE-5 Inhibitors: treatment of erectile dysfunction (ED) in males.
- Currently there are three agents available—sildenafil (Viagra).
- Mechanism: Inhibitory effects on PDE 6, an isoenzyme expressed in the rods and cones of the eye.
- Amiodarone: This drug, a class III antiarrhythmic agent: exact mechanism of optic neuropathy is unknown.
- The condition is typically insidious in onset, takes months to resolve, and presents bilaterally
↳ very very slow

Optic neuropathy

- Optic neuropathy has been associated with the long-term use of linezolid, with reported duration of treatment ranging from 5 to 11 months in doses of 600 to 1,200 mg/day.
- **Ethambutol**: This drug, a first-line agent in the treatment and prevention of tuberculosis
 - . It has been well documented to cause optic neuropathy in up to 5% of patients taking the drug.
 - The mechanism is related to related to chelation of copper in retinal cells.
- **Isoniazid-induced optic neuropathy** is thought to be less frequent and reversible.

Side Effects of Antipsychotic Medications (neuroleptic drugs)

Neuroleptic Malignant Syndrome (NMS): life-threatening

Due to autonomic disturbances

• Hyperthermia, muscular rigidity, rhabdomyolysis, tachycardia, hyper or hypotension, confusion

• **Complications**: Coma and death

• **Treatment**:

• Stop drug

• Supportive management and

• Sever cases: ICU

⊗ Haloperidol

⊗ chlorpromazine

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- **Thank you**