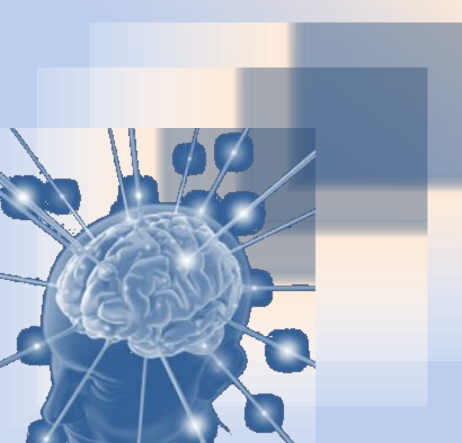
DRUGS USED IN HEADACHE & MIGRAINE



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- Differentiate between types of headache regarding their symptoms, signs & pathophysiology.
- Recognize drugs used to prevent migraine
- Identify drugs used to rescue & abort migraine
- Elaborate on the pharmacokinetics, dynamic & toxic profile of some of these drugs.



It is caused by disturbance of the **Pain – Sensitive Structures** around the brain

Within the cranium

(blood vessels, meninges, cranial nerves)

Outside the cranium

(muscles, nerves, arteries, veins, subcutaneous tissues, eyes, ears & other tissues).

MIGRAINE

Recurrent attacks of throbbing headache Unilateral / or on both sides Lasting from > 2 up to 72 hrs. + Preceded (or accompanied) by AURA

Perceptual disturbance of motor < sensory nature visual [Photophobia (↑sensitivity to light)] auditory [Phonophobia (↑ sensitivity to sound)] olfactory unpleasant smell Sensory; abnormal sensation at face, extremities. Develops over 5-20 min & last fewer than 60 min.

Aura: flashes of light, blind spots or tingling in your arm.

Migraine pain is usually on one side of head with facial & neck pain, nausea & vomiting.

Phases of Migraine

1. Prodrom Phase; a change in mood or behavior (irritability, neck stiffness) that starts hours or days before headache. It is experienced by 60% of migraineurs.

2. Aura Phase; Sensory > motor symptoms starts 5-20 min before the migraine attack. It is experienced by 20% of migraineurs.

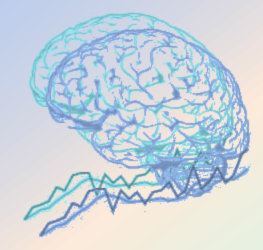
3. Headache Phase; moderate to severe pain, ↑ with activity + anorexia, vomiting, Intolerance to light, sounds, odors Blurry vision /Blocked nose /Pale face Sensations of heat or coldness /Sweating /Tenderness of the scalp

4. Postdrom Phase: still not normal, either;
 More likely fatigued → irritability /impaired concentration /scalp tenderness /mood changes / GIT symptoms,

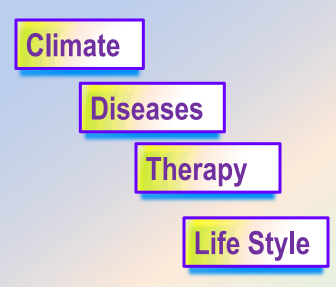
Migraine Triggers



Aged cheese, Alcohol, Chocolate, Caffeine, Hot dogs, Avocado, Fermented or pickled foods, Yeast or protein extracts.



Hormonal changes: Menstrual migraine



Antibiotics, Antihypertensives, H₂ blockers, Vasodilators, Oral contraceptives.

Migraine Causal Theories

Vascular

Criggers Spreading Depression

Neurovascular theory ? on \rightarrow migraine aura

focal ischemia $\rightarrow \uparrow$ mediators \rightarrow rebound vasodilatation $\rightarrow \uparrow$ permeability & leak in j inflammatory reaction \rightarrow activates perivascular nociceptive nerves \rightarrow migraine headache

Dopaminergic Hypersensitivitysensitive area with each heart beat

Migraine Causal Theories

Vascular

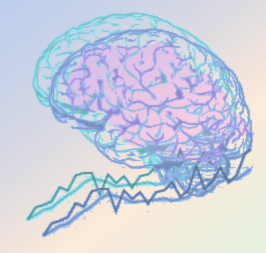
Triggers

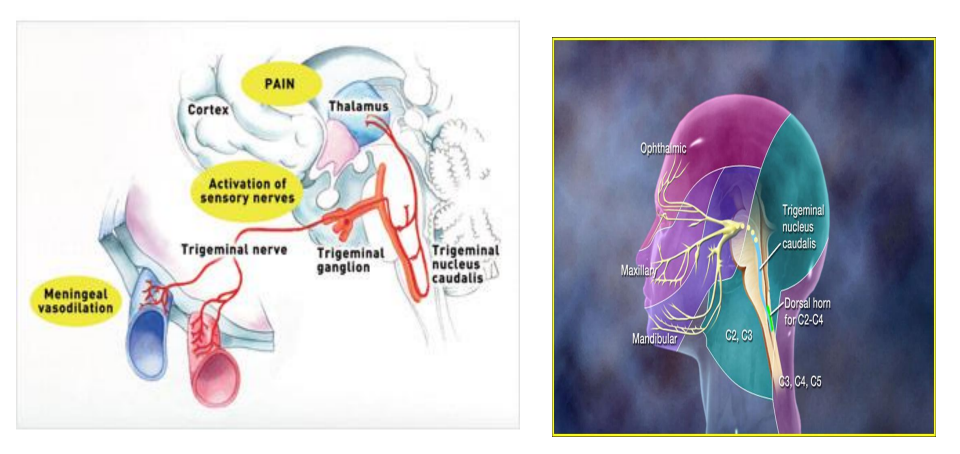
Řelease K / glutamates

Neurovascular theory ?

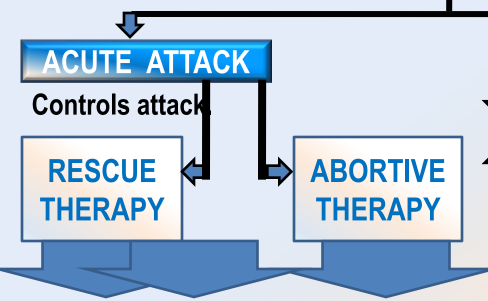
Creates a slowly well-defined depolarizing wave \rightarrow depolarize adjacent tissues oto propagating at a rate of 2-6 mm/min \rightarrow vasoconstriction \rightarrow migraine aura

 \rightarrow activate trigeminovascular complex \rightarrow vasodilation \rightarrow migraine headache. Dopaminergic Hypersensitivity





Stimulation of the trigeminal nerve causes the release of vasoactive peptides; this is responsible for the head pain, as well as the facial & neck pain, experienced during migraine.



TREATMENT STRATEGY

PREVENT RECURRENCE

 recurrence frequency, severity, duration & / or disability
 responsiveness to abortive therapy
 N.B. Full effect of therapy needs several weeks to manifest
 & should continue for 6 m.
 & can be repeated

Non-specifically target individual symptoms i.e. alleviating pain, emesis & associated symptoms

Mild-Moderate

They specifically target pathways of migraine by Ψ meningeal dilatation & Ψ neural activation via 5HT₁ agonism \rightarrow i.e. stopping headache as it is evolving.

Abortive medications > effective if taken early, just before the pain starts, losing effectiveness once the attack has begun Severe/ Disabling idly acting 10

TREATMENT of Acute Attack RESCUE THERAPY

➔ Analgesics

- NSAIDs / Aspirin< Acetaminophen
 (ibuprofen, naproxen for mild to moderate attack with no nausea & vomiting)
- >Opioid-like drugs: μ agonist; e.g. Tramadol.

➔ Antiemetics

Oppamine Antagonists + Gastro-prokinetic
Dependence Antagonists + Gastro-prokinetic

Domperidone

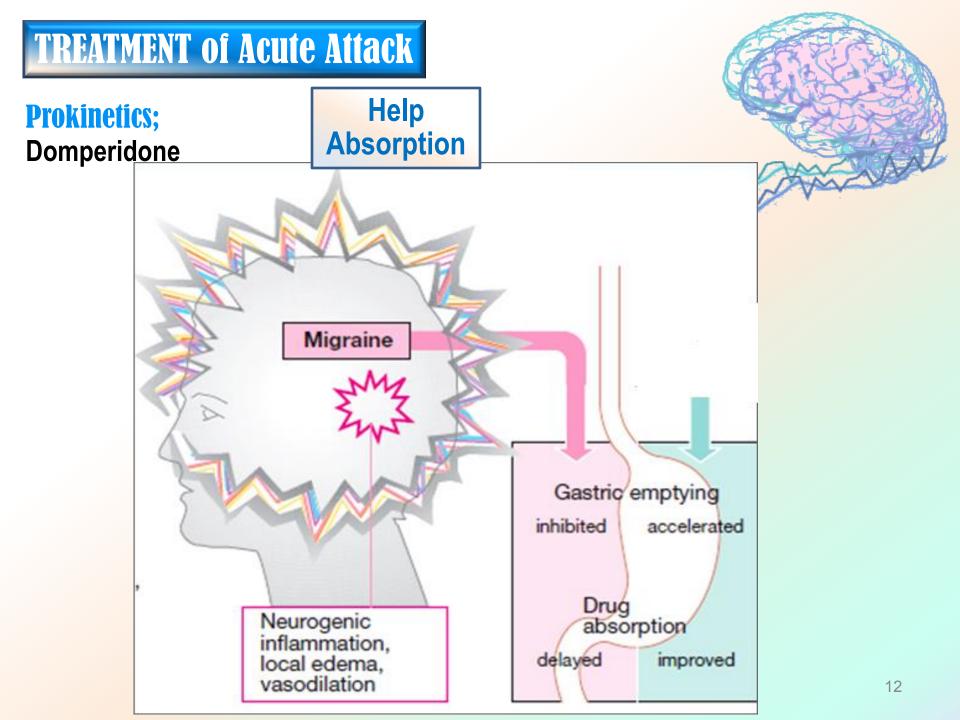
↑ Absorption & bioavailability of <u>abortive therapy</u>

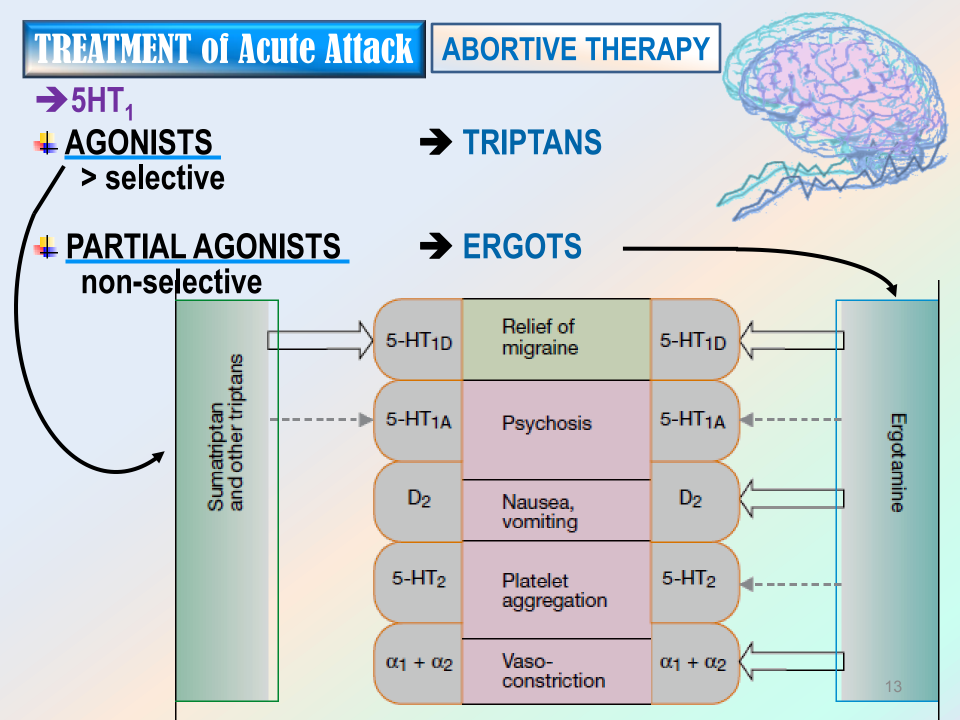
Phenothiazines Promethazine

Dopamine antagonists + Sedation

- ♦ 5HT₃ antagonists (for severe nausea & vomiting
 - Ondanseteron Granisetron
 - H₁ antagonist Meclizine, diphenhydramine

Antihistamine +sedation Anticholinergic





 TREATMENT Of Acute Attack
 ABORTIVE THERAPY
 ERGOTS

 Product of Claviceps purpurea; a fungus growing on rye/ grains
 Image: Claviceps purpurea in the second secon

Non-Selective

Partial agonism at 5HT₁ receptors (5HT-1D/1B found in cerebereal & menigeal vessels)

↓ release of vasodilating peptides

↓excessive firing of nerve endings

At blood vessels $\rightarrow \downarrow$ vasodilation & stretching of the pain endings

Partial agonist effect on α -adrenoceptors \rightarrow vasoconstriction

Ergotamine tartarate (resticted use) Oral, sublingual, rectal suppository, inhaler Caffeine - Cafergot

Dihydroergotamine (DHE)

Nasal spray, inhaler & injectable forms (good to use if patient is vomiting)

Ergotamine tartarate (rare clinical use due to sever
adverse effectsERGOTSOral absorptionIncomplete (erratic) + slow \rightarrow low bioavailability

Despite $t_{1/2}$ nearly 2 hours, ergotamine produces vasoconstriction \rightarrow 24 hours or longer due to high & long tissue binding ability. <u>Ergotamine tartrate</u> has significant side effects, & may worsen the nausea & vomiting associated with migraine.

DHE (preferred in clinical setting)

infrequent but severe attacks.

Given parenterally, DHE is eliminated more rapidly than ergotamine, presumably due to its rapid hepatic clearance & has less adverse effects.

Indications

They are only used to abort the attacks [Exception DHE can be given for severe, recurrent attacks not responding to other drugs] Their use is restricted to patients with frequent, moderate attack or





- GIT upset
- Feeling of cold & numbness of limbs, tingling
- Anginal pain due to coronary spasm, & disturbed cardiac rhythm (tachycardia or bradycardia)
- ➡ Prolong use → rebound headache due to vasodilatation followed by vasoconstriction.
- ♣ Prolong use & high dose → paraesthesia (tingling or burning sensation).

Contraindications

- Pregnancy; fetal distress & miscarriage (ergot is uterine stimulant & vasoconstrictor)
- Peripheral & coronary vascular diseases
- **Hypertension**
- Liver & kidney diseases
- prophylaxis of migraine
- In concurrent use with triptans (at least 6 hrs from last dose of triptans or 24 hrs from stopping ergotamine & β-blockers.

TREATMENT of Acute Attack ABORTIVE THERAPY

TRIPTANS

Selective

- Agonism at 5HT₁ receptors
- Same as discussed for ergotamine except that triptans are more selective as serotonergic agonist.
- <u>No α_1 , α_2 , β –adrenergic, dopamine or muscarinic receptors.</u>
- Triptans inhibit the release of vasoactive peptides, promote vasoconstriction, & block pain pathways in the brainstem. Triptans inhibit transmission in the trigeminal nucleus caudalis. **SUMATRIPTAN** Present in \rightarrow oral, nasal spray, and injectable forms Oral bioavailability low / Subcutaneous (SC) bioavailability is 97%, peaks after 2 min & t_{1/2} nearly 2 hours (fast action with Sc, good for patient with vomiting) **ZOLMITRIPTAN** Present in \rightarrow nasal spray, and injectable forms Oral bioavailability 40%, peaks after 2 hrs & t_{1/2} nearly 3 hours **NARATRIPTAN** *Present in addition* \rightarrow + *Oral preparations* Oral bioavailability 70%, peaks after 2 hrs & t_{1/2} nearly 6 hours (slower
- onset, less side effects).

Indications

To abort attacks in patients with frequent, moderate or infrequent but severe attacks. 🔸 In cluster headache

ADRS

- **4** most of ADRs are the same as with ergot but triptans are better tolerated.
- **4**Mild pain & burning sensation at the site of injection.
- **4** Vasospasm, Ischemic heart; Angina & Arrhythmias

Contraindications

- Peripheral vasospastic diseases
- Uncontrolled hypertension
- **History of ischemia**
- Cerebrovascular disorders
- ↓ In concurrent use with ergots or others inducing vasospasm ↓ In concurrent use with MAOIs, lithium, SSRIs,→(5HT increased)

to toxic level)

Renal or hepatic impairment.

Zolmitriptan Chest & neck tightness Coronary vasospasm **4** Somnolence.

DECIDING WHETHER BETTER WITH A TRIYPTAN OR WITH DHE.

- For patients with headache episodes lasting 2 or 3 days at a time, DHE is often the optimal choice because it has longer t_{1/2}
- For patients with migraines a day or less & need rapid relief of pain,
 Triptans are often a better choice
- For pregnant women: paracetamol or intranasal sumitriptan & or diphenhydramin, meclizine are safe to be used.

The form of drug preparation could influence the choice Injectable sumatriptan reaches T_{max} the fastest followed by DHE nasal spray & rizatriptan. Differences in the time to peak blood concentration T_{max}, equates with faster relief of pain.

4Differences in $t_{1/2} \rightarrow a$ clinical effect in terms of recurrence of headache

Pharmacokinetics

Medication T_{1/2}(h) For extremely fast relief within 15 min. injectable sumatriptan is the only choice.

- If expected re-dosing is needed & / or recurrence of headache Naratriptan, frovatriptan, have slower onset, fewer side effects, & a lower recurrence rate
- Menstrual migraine: Frovatriptan (longer half life (26 hrs) 2.5 mg twice per day beginning 2 days before the anticipated onset of menstrual migraine & continuing for 6 days.

TREATMENT STRATEGY

ACUTE ATTACK

PREVENT RECURRENCE

Antiepileptics;

Block Na channel & augment GABA at GABA-A receptors

e.g. Topiramate;

Valproic;

Antidepressants

TCA; amitryptylin & nortryptyline

Antihypertensives

- β-blockers
- e.g. propranolol
- .Propranolol is commonly
- used in pophylaxis of migraine attack.