PEDIATRIC AND ADOLESCENT HEADACHES

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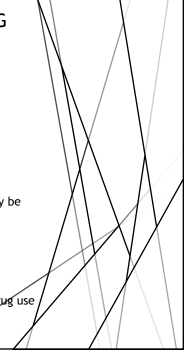
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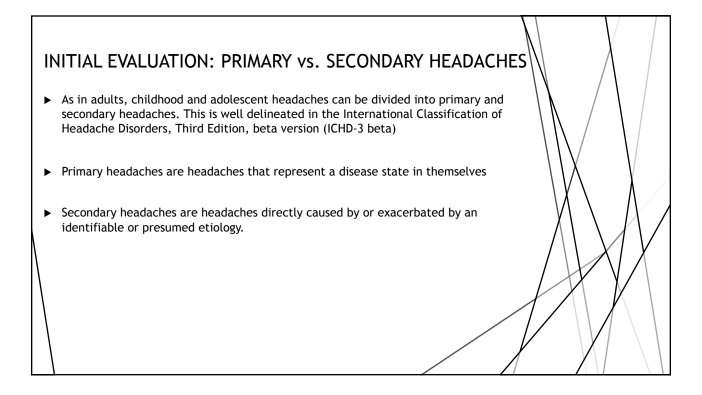
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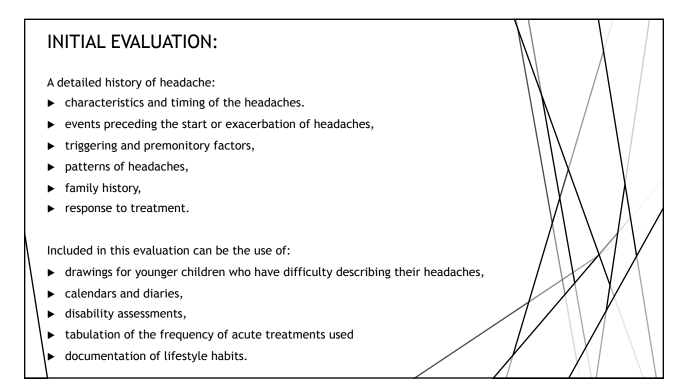
- ▶ APPROACH TO PEDIATRIC HEADACHE HISTORY TAKING
- ► INITIAL EVALUATION
- ► RED FLAGS FOR SECONDARY HEADACHE PATHOLOGY IN CHILDREN AND ADOLESCENTS
- ▶ INTRACRANIAL NEOPLASMS
- ▶ NEUROIMAGING
- ▶ PEDIATRIC HEADACHE FEATURES NOT RED FLAGS FOR SECONDARY PATHOLOGY
- ▶ PRIMARY HEADACHE DISORDERS AFFECTING CHILDREN AND ADOLESCENTS
- ► TREATMENT OF PEDIATRIC AND ADOLESCENT MIGRAINE
- **▶** CONCLUSION

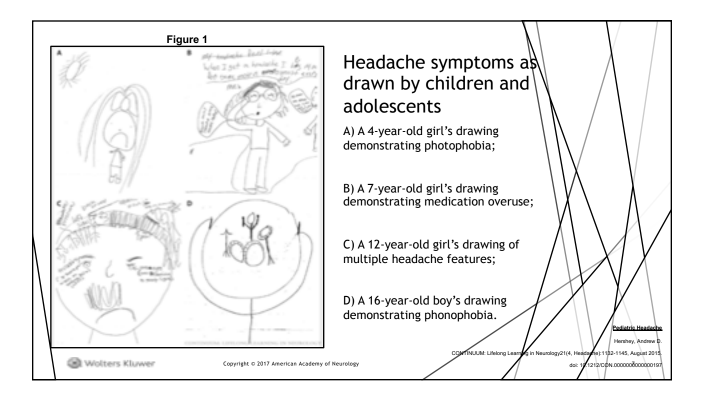
APPROACH TO PEDIATRIC HEADACHE HISTORY TAKING

- ▶ Step 1: Assign seats
- ▶ Step 2: Set expectations at the outset
- ▶ Step 3: "Good headache histories are taken, not given."
 - ▶ Very Young Children less than 6 years old
 - ▶ information from parent/guardian
 - ▶ Have children draw a picture of themselves and how they feel
 - ▶ Preadolescent School aged children
 - ▶ Time is a challenging concept details on duration and frequency may be obtained from parents, calendars, diaries
 - ► Medication names/dosages
 - ▶ Adolescents
 - ▶ Can usually give complete history with minimal parental assistance
 - ▶ May ask parent to step out for a confidential discussion on alcohol, drug use



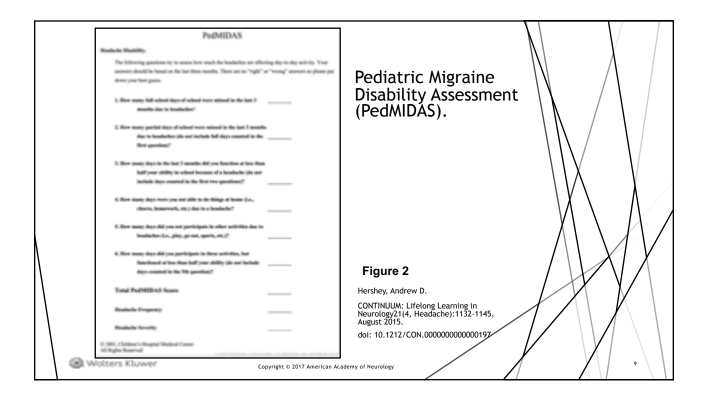


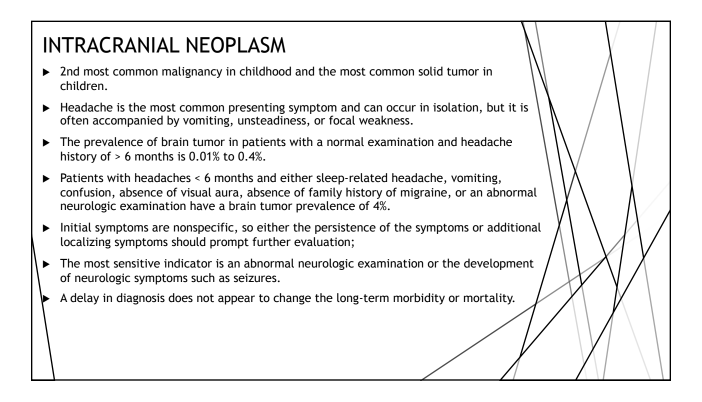




DISABILITY ASSESSMENT:

- ► The assessment of impact and disability caused by the headaches is a subjective but important tool.
- ▶ This can include both disease-specific and nonspecific features.
- For children and adolescents, disease nonspecific disability has been demonstrated using the Pediatric Quality of Life Inventory (PedsQL).
- ► For migraine and other primary headaches, disease-specific features may be the associated symptoms (eg, vomiting) or characteristics of the aura. Additionally, disease-specific components of migraine may impact functioning socially, at school, and at home.
- ► The Pediatric Migraine Disability Assessment (PedMIDAS) was developed as a modification of the Migraine Disability Assessment (MIDAS) to assess migraine-related disability in children and adolescents.
- ▶ It is not intended to be an exact measure, but has been demonstrated to correlate well with the overall impact of migraine and is a useful tool to follow treatment response.





NEUROIMAGING:

One aspect of secondary headache evaluation is the role of neuroimaging. Several key components of guidelines (from AAN, CNS, AHS) that indicate the need for neuroimaging include:

- ▶ neurologic symptoms,
- any abnormality on the neurologic examination,
- an exclusively occipital location to the headaches.
- ▶ lack of family history
- frequent early morning headaches that awaken the child from sleep or result in vomiting, suggestive of a pressure effect;
- not meeting ICHD-3 beta criteria;
- recent onset of headaches without a history of headaches;
- ▶ a recent change in the headaches, suggestive of a new type of headache;
- ▶ children aged 6 or younger who may have difficulty describing their headaches.
- ▶ brain MRI is the preferred modality choice in children and adolescents.
- neurophysiologic testing (EEG) and blood chemistry testing not useful

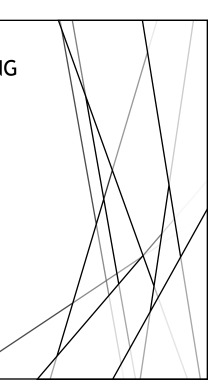
RED FLAGS FOR SECONDARY HEADACHE PATHOLOGY IN CHILDREN AND ADOLESCENTS Persistent headaches of less than 6-months' duration that do not respond to medical treatment Headache associated with abnormal neurologic findings, especially if accompanied by papilledema, nystagmus, or gait or motor abnormalities Persistent headaches associated with a negative family history of migraine Persistent headaches associated with substantial episodes of confusion, disorientation, or emesis Headaches that repeatedly awaken a child from sleep or occur immediately on awakening Family and medical history of disorders that predispose to CNS lesions and clinical laboratory findings that suggest CNS involvement Reprinted with permission from Medina J, Pinter J, Zurakowski D, et al. Children with headache: clinical CONTINUUM: Lifelong Neurology14(4, Neurol 2008 predictors of surgical space-occupying lesions and the role of neuro/maging. Radiology 1997;202:819. CONTINUUM: LIFELONG LEARNING IN NEUROLOGY Wolters Kluwer Copyright © 2017 American Academy of Neurology

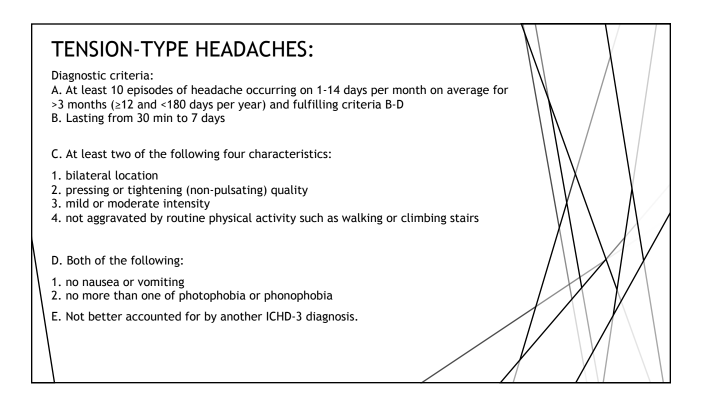
PEDIATRIC HEADACHE FEATURES NOT RED FLAGS FOR SECONDARY PATHOLOGY

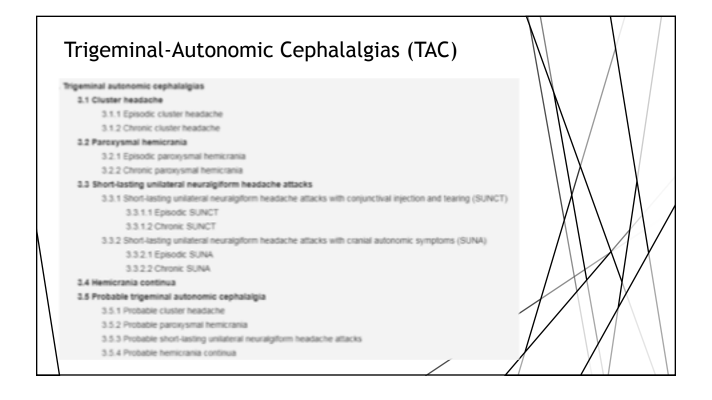
- ▶ Occipital Headache Location
- ▶ Headaches Accompanied by Nasal Congestion, Itchy Eyes, or Ear Pressure
- ► The list of cranial autonomic symptoms recognized in the International Classification of Headache Disorders, Third Edition (ICHD-3) is as follows:
 - ► Conjunctival injection and/or lacrimation
 - ▶ Nasal congestion and/or rhinorrhea
 - ▶ Eyelid edema
 - ▶ Forehead and facial sweating
 - ▶ Forehead and facial flushing
 - ▶ Sensation of fullness in the ear
 - ▶ Miosis and/or ptosis

PRIMARY HEADACHE DISORDERS AFFECTING CHILDREN AND ADOLESCENTS

- ► Tension-Type headaches
- ► Trigeminal-Autonomic Cephalalgias (TAC)
- ▶ Post-Traumatic headaches
- Primary stabbing headaches
- ▶ Migraine







Posttraumatic Headache

- ▶ According to the ICHD-3, posttraumatic headache must begin within 7 days of head trauma to be attributed to that injury
- ► The topic of concussion and its management is a burgeoning and important area of research and is beyond the scope of this presentation.
- ▶ In brief, the phenotype of posttraumatic headache in children and adolescents can be featureful (ie, migrainous) <u>or</u> featureless (ie, similar to tension-type headache).
- ▶ In the absence of randomized trials guiding posttraumatic headache treatment in this age group, treating the headache based on the underlying phenotype seems reasonable.

Primary Stabbing Headache

- ▶ Brief attacks of sharp pain.
- ▶ The pain is typically described as a stab or series of stabs. It can be quite severe. Some children will be brought to their knees by the pain.
- Duration is typically just a few seconds, although some children may experience attacks lasting several minutes.
- Location can change from attack to attack or be fixed.
- ► The complete absence of cranial autonomic symptoms is important in distinguishing primally stabbing headache from trigeminal autonomic cephalalgias.
- Migratory location is also a helpful feature in distinguishing primary stabbing headache from trigeminal autonomic cephalalgia.
- Usually, attacks are rare and short enough that they do not require any specific treatment.
 Reassurance about the benign nature of the headaches is typically all that families need.
- ► However, preventive treatment may be considered in cases where the attacks are frequent and distressing.
 - ▶ Indomethacin is useful for some adult patients.
 - ▶ Some patients will respond to nightly melatonin.

How Common is Migraine?

36 million Americans are affected by migraine

18% women and 6% of men in the US have migraine

1 in every 4 American households includes an individual affected by migraine









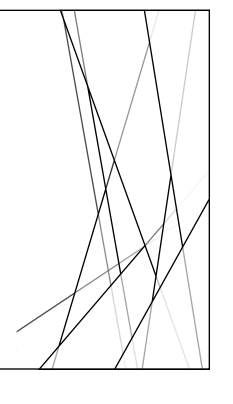
American Migraine Prevalence and Prevention (AMPP) Study

Identifying Migraine

The 3 Question ID Migraine Screen (a validated screen) for migraine

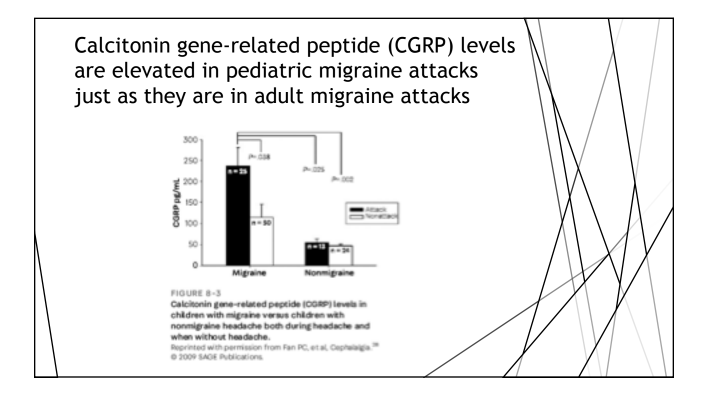
- Has a headache limited your activities for a day or more in the last three months?
- 2 Are you nauseated or sick to your stomach when you have a headache?
- 3 Does light bother you when you have a headache?

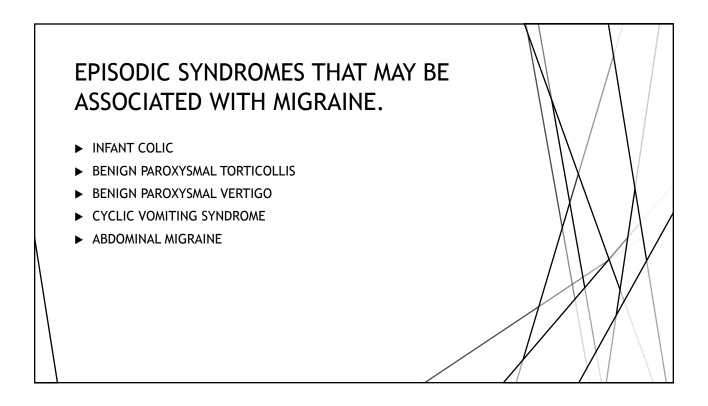
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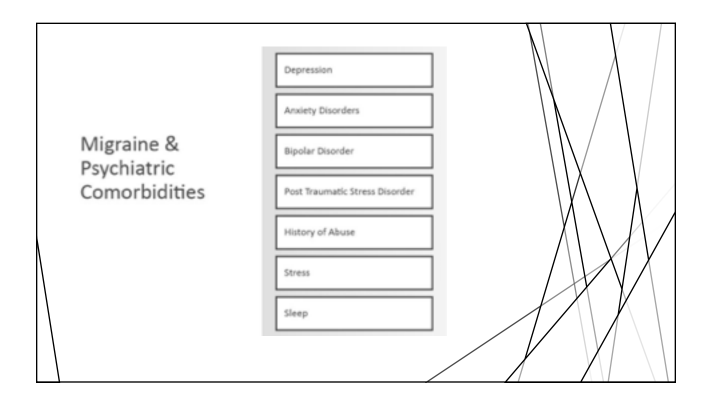


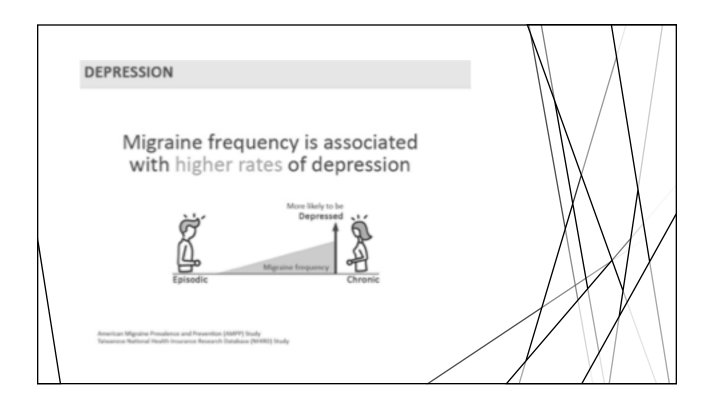
PHENOTYPIC FEATURES OF MIGRAINE THAT DIFFER IN CHILDREN AND ADOLESCENTS VERSUS ADULTS

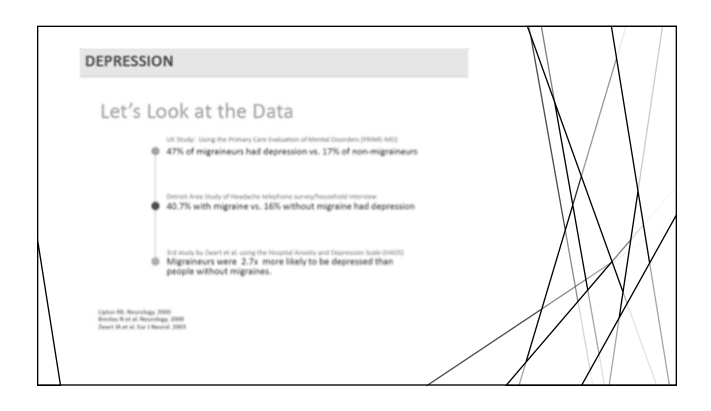
- ▶ PREMONITORY PHASE OF PEDIATRIC AND ADOLESCENT MIGRAINE
 - ▶ Fatigue, irritability/mood change, neck stiffness, and facial changes
- ► ICTAL PHASE OF PEDIATRIC AND ADOLESCENT MIGRAINE
 - ► Migraine duration in children can be shorter, particularly in children younger than 7 years of age.
 - ▶ In the ICHD-3, the lower margin of duration for untreated or unsuccessfully treated attacks in children is 2 hours versus 4 hours in adults.
 - ▶ The majority (more than 80%) of children and adolescents report bilateral migraine headache. This is the phenotype through late adolescence.
- ▶ POSTDROME PHASE OF PEDIATRIC AND ADOLESCENT MIGRAINE
 - ▶ thirst, somnolence, visual disturbances, and food cravings
 - ▶ In the vast majority of patients, postdrome symptoms resolved within 12 hours.

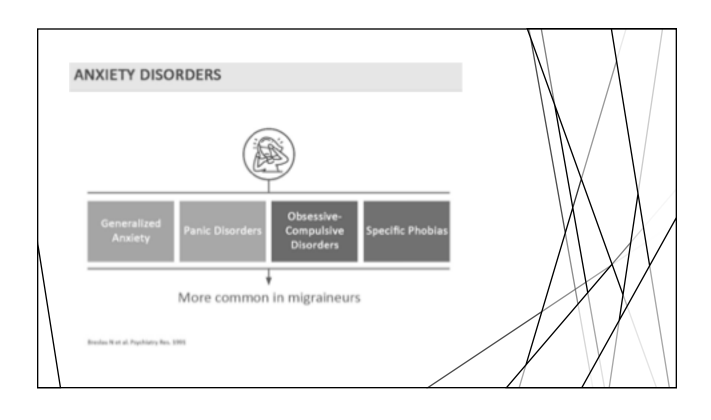


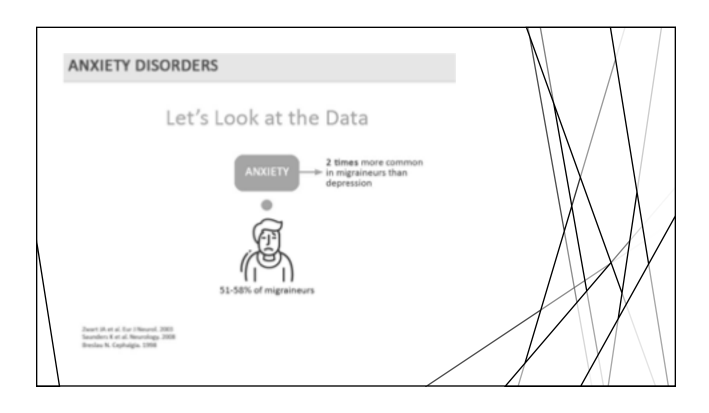


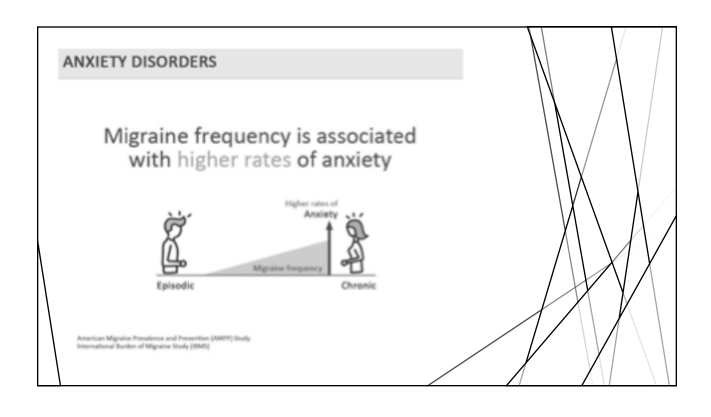


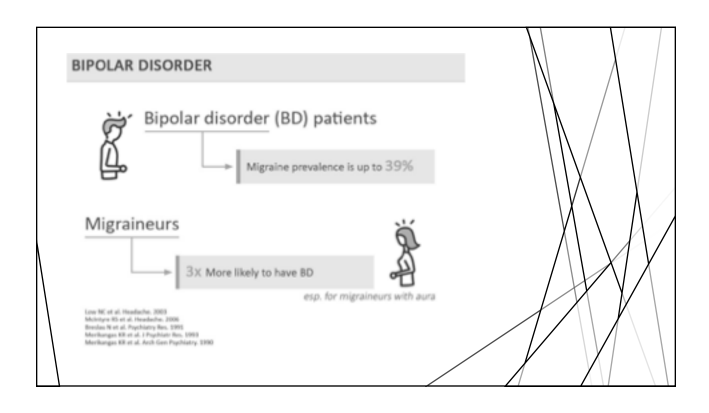


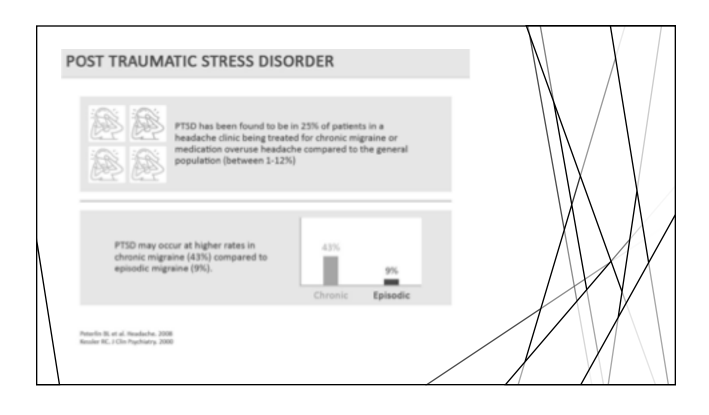


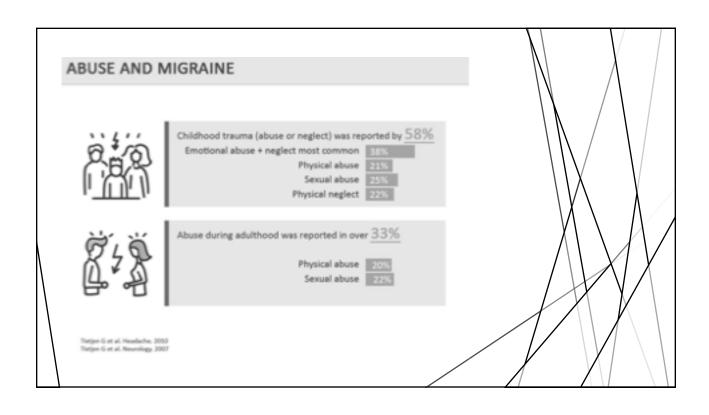




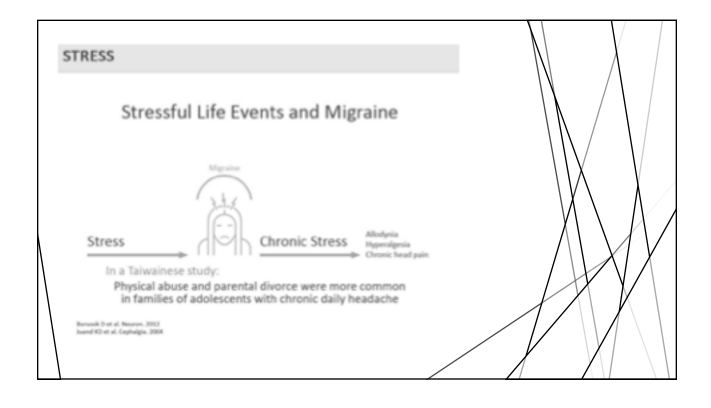


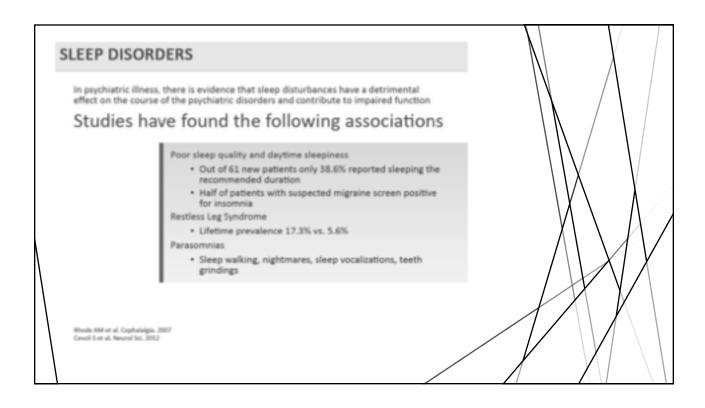


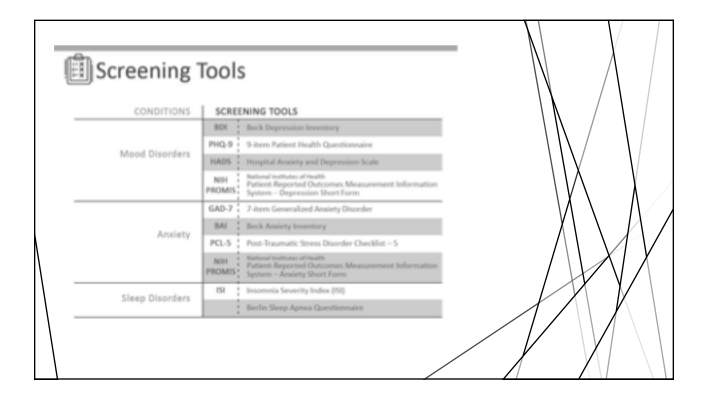


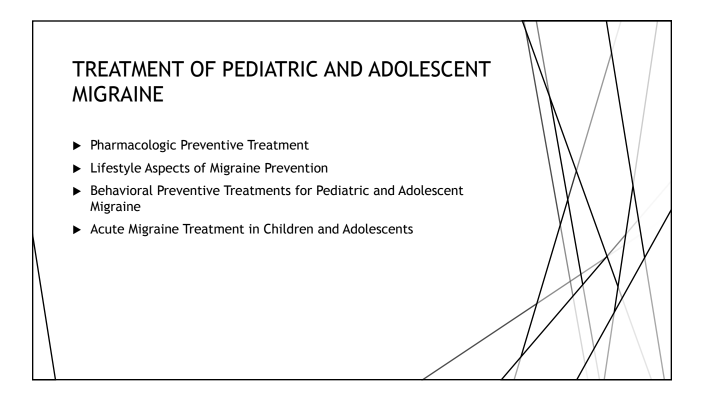


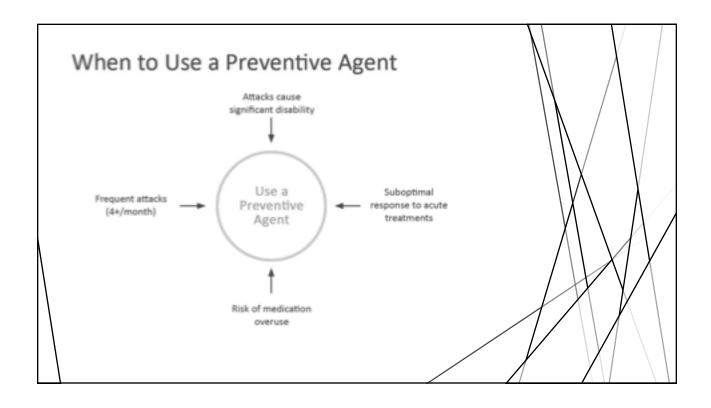
* Hormonal changes * Nocturnal peak in prolactin * Delayed nocturnal peak in melatonin * Increased amount of cortisol * Menstrual Cycle * Medication overuse can also affect allostasis









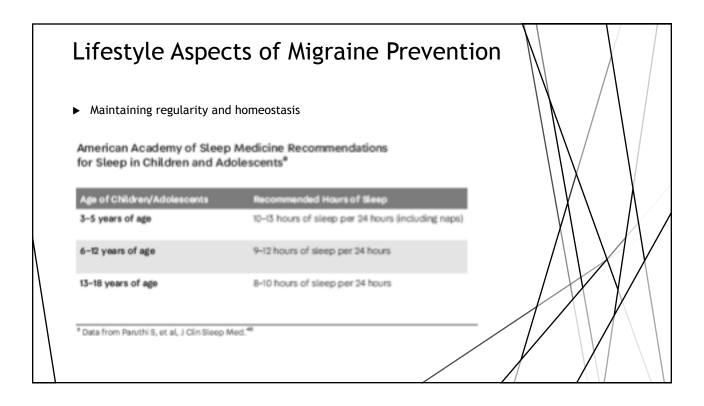


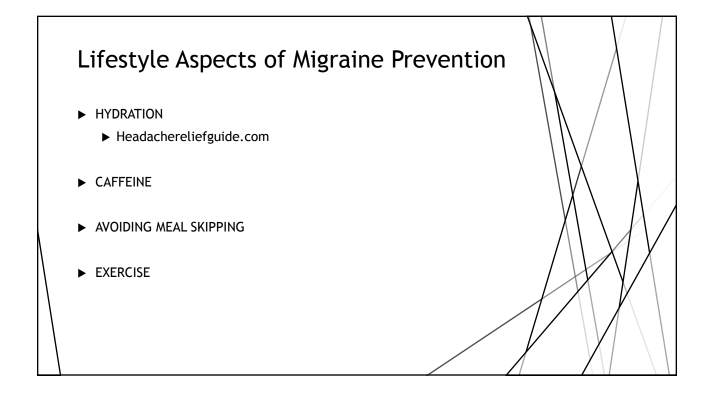
Pharmacologic Preventive Treatment

- ► The CHAMP trial published in 2017 has helped to reframe the approach to migraine prevention in children and adolescents
- ▶ NIH-funded multisite trial designed to identify a first-line preventive for pediatric migraine prevention in children and adolescents ages 8 to 17 years, participants could have episodic or chronic migraine
- ► The three treatment arms of the CHAMP trial were amitriptyline (goal dose of 1 mg/kg), topiramate (2 mg/kg), and placebo.
- ▶ In all three treatment arms, approximately 60% of the participants met the primary end point of a 50% or more reduction in headache days 24 weeks after starting preventive therapy
- Topiramate currently remains FDA-labeled for migraine prevention in adolescents 12 to 17 years of age, and no preventive therapies labeled for children younger than age 12 exist

High placebo response rate seen in the CHAMP trial:

- ► Lifestyle migraine management advice on sleep, exercise, hydration/eating, and caffeine. This advice was reinforced at monthly study visits.
- All participants received evidence-based optimal acute therapy, specifically nonsteroidal anti-inflammatory drugs (NSAIDs) and triptans, dosed appropriately and with use frequency guidance so as to avoid medication overuse.





Behavioral Preventive Treatments for Pediatric and Adolescent Migraine

- ► Cognitive Behavioral Therapy
 - ▶ incorporates identification and modification of thinking patterns and behaviors to improve perception of pain
- Biofeedback
 - ► Subconsciously controlled body functions (skin temperature, heart rate) are monitored with a technical device and patients learn to control them
- ▶ Relaxation Techniques
 - ► Mindfullness Stress Reduction Therapy
 - ▶ Aerobic exercise
 - ► Acupuncture

NEUTRACEUTICALS

- The evidence for the use of nutraceuticals is low or conflicting.
- For migraine prevention, Level B evidence, at best, exists for the use of feverfew, magnesium, and riboflavin (vitamin B2) in adults.
- Level C evidence exists for coenzyme Q10 (CoQ10)
- · Level U evidence for melatonin.
- The evidence level for IV magnesium for acute migraine treatment is B.
- Recent AAN guidelines, concluded that relaxation training, thermal biofeedback combined with relaxation training, EMG biofeedback, and cognitive-behavioral therapy all have Grade A evidence for episodic migraine prevention.

Acute Migraine Treatment in Children and **Adolescents**

- **ACETAMINOPHEN**
- NONSTEROIDAL ANTI-INFLAMMATORY DRUGS
 - ▶ Ibuprofen
 - ▶ Naproxen
 - ► Ketorolac
- TRIPTANS
 - ▶ almotriptan (oral),
 - ▶ zolmitriptan (nasal spray),
 - ▶ rizatriptan (melt),
 - ► sumatriptan/naproxen (oral)

Counseling to ensure the patient does not develop medication-overuse beadache

Acute Migraine Treatment in the Emergency and the Inpatient setting:

- Sumatriptan 4 mg to 6 mg subcutaneously
- Antiemetics plus dihydroergotamine (0.5 mg to 1 mg IV, repeat in 1 hour)
- ▶ Neuroleptics

Chlorpromazine (0.1 mg/kg) 12.5 mg to 37.5 mg IV/IM Prochlorperazine 5 mg to 10 mg IM, 25 mg per rectum

Haloperidol 5 mg IV in 500 mg normal saline over 20 to 30 minutes

- ▶ Ketorolac 30 mg to 60 mg IM; may treat cutaneous allodynia if not complicated by opioid use
- ▶ Magnesium 1 g to 2 g IV; limited evidence; may treat photophobia/phonophobia
- Valproate 300 mg to 500 mg IV; open-label trials
- ► Corticosteroids (eg, dexamethasone 10 mg to 24 mg IV)
- Metoclopramide 20 mg IV; may repeat

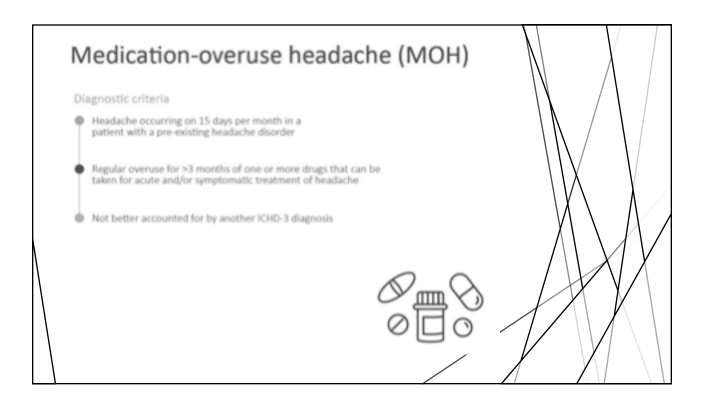
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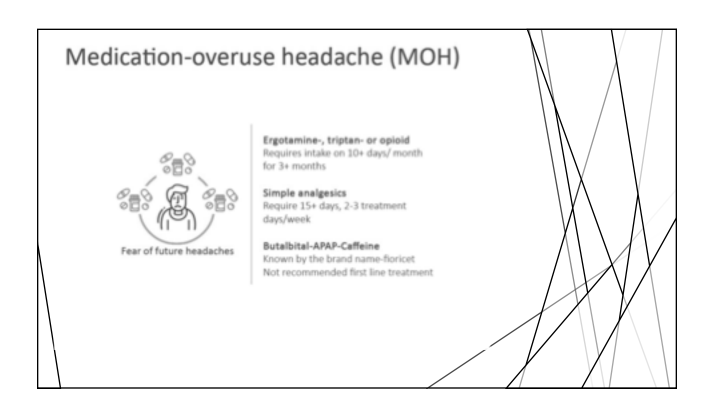
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CONCLUSION

- ▶ Headache is common in children and adolescents.
- Most children who come to see the neurologist for headaches will have a primary headache disorder, with migraine being most common in this setting.
- ► For acute migraine treatment, acetaminophen and NSAIDs have been studied in children age 4 and older and have been found to be effective.
- ► Triptans are also effective in children and adolescents. Four triptans are now FDA-labeled for acute migraine treatment in adolescents, and rizatriptan is labeled for use in children age 6 and older.
- For preventive migraine treatment, the recent CHAMP trial indicates that approximately 60% of children and adolescents with migraine will improve with a three-pronged treatment approach that includes:
 - ▶ lifestyle management counseling
 - evidence-based optimally dosed acute therapy, specifically NSAIDs and triptans
 - ▶ a daily preventive treatment that has some evidence for efficacy and a side effect profile that is similar to that of placebo

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- Rizzoli, Paul B. CONTINUUM: Lifelong Learning in Neurology18(4, Headache):764-782, August 2012. doi: 10.1212/01.CON.0000418641.45522.3b
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