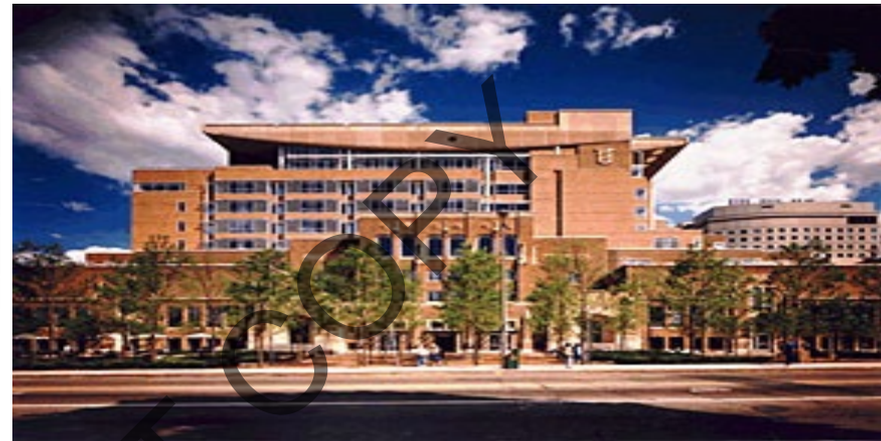




Beth Israel Deaconess
Medical Center



HARVARD MEDICAL SCHOOL
TEACHING HOSPITAL



Introduction to Seizures

Trudy D. Pang, MD, MMSc
Assistant Professor of Neurology
Harvard Medical School
Comprehensive Epilepsy Center
Beth Israel Deaconess Medical Center

Outline

- Definitions and epidemiology
- Differential diagnosis and seizure types
- Diagnostic workup
- Seizure first aid
- Seizure management

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Seizures and TMS

- They are the most dramatic and medically dangerous acute complication of TMS
- IRB/ethics boards expect them to be addressed as a risk of TMS research
- The world of TMS research has expanded:
 - To researchers who are not physicians or who are not familiar with clinical neurological disorders
 - To labs that are not located proximate to medical facilities
 - To subject populations with known epilepsy or with neurological disorders that have an increased risk of seizures

What is a Seizure?

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Seizure

- A clinical episode of neurologic dysfunction caused by the abnormal hypersynchronous activity of a group of neurons

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What is Epilepsy?

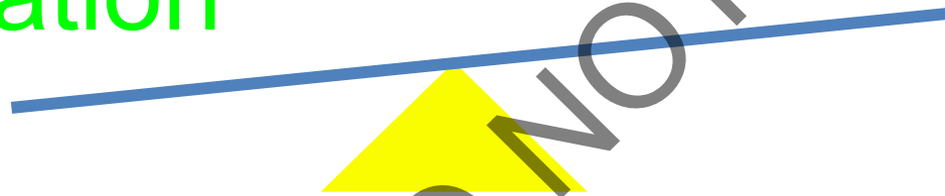
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- Any disorder characterized by a tendency toward recurrent, unprovoked seizures
- A disease of the brain defined by any of the following 3 conditions:
 1. 2 or more seizures occurring >24h apart
 2. 1 unprovoked seizure and a probability of further seizures of at least 60% occurring over the next 10 years
 3. Diagnosis of an epilepsy syndrome
- In practice, diagnosed after two unprovoked seizures

Seizures occur when an imbalance of excitation and inhibition exists in the nervous system

Excitation

Inhibition



Examples

hypoxic-ischemic brain injury

developmental brain malformation

traumatic brain injury

neurosurgery

brain tumors

alcohol-related

strokes

CNS infections

neurodegenerative diseases

CNS demyelination/inflammation

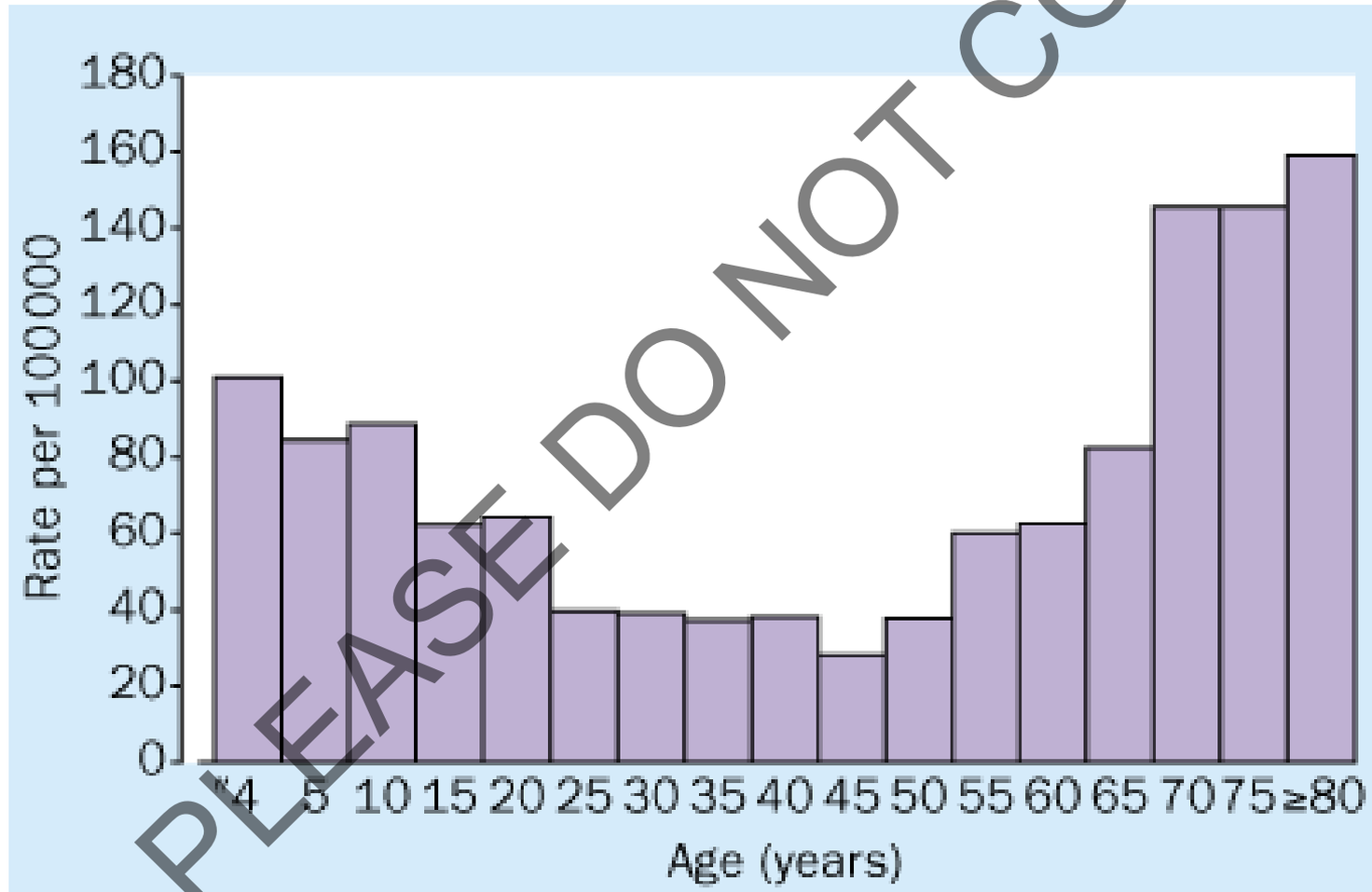
inborn errors of metabolism

systemic illness

Seizures and epilepsy are quite common

- Prevalence of epilepsy in the general population is about 0.5% to 1%
- Cumulative lifetime incidence of one or more seizures is 5-10%, including febrile seizures

The incidence of epilepsy is highest in the young and in the old



Seizures are classified by their origin in the brain and associated clinical features

- Focal-onset
 - ▣ Focal aware (simple partial)
 - ▣ Focal impaired awareness (complex partial)
- Generalized-onset
 - ▣ Generalized tonic-clonic
 - ▣ Absence
 - ▣ Myoclonic
- Focal onset seizures have the potential to progress to secondarily generalized seizures

ILAE 2017 Classification of Seizure Types Basic Version ¹

Focal Onset

Aware

**Impaired
Awareness**

**Motor Onset
Nonmotor Onset**

focal to bilateral tonic-clonic

Generalized Onset

Motor

Tonic-clonic
Other motor

Nonmotor (Absence)

Unknown Onset

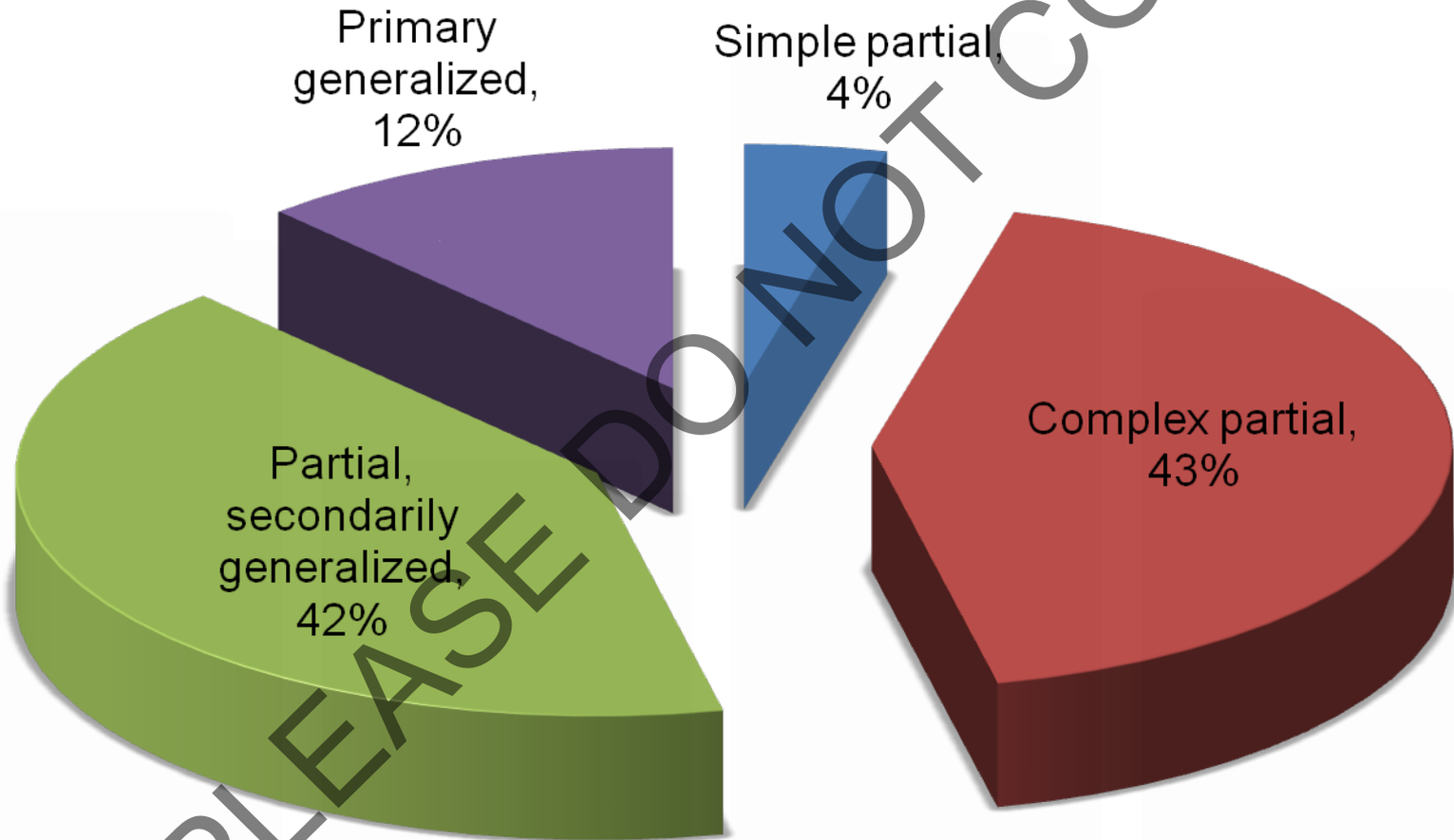
Motor

Tonic-clonic
Other motor

Nonmotor

Unclassified ²

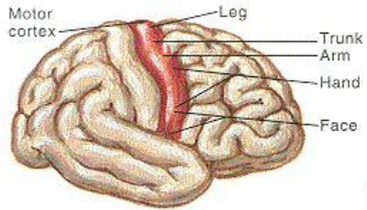
Most seizures in adults are partial-onset



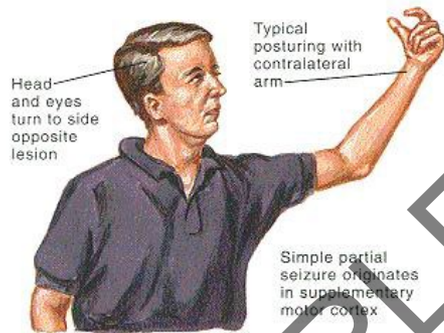
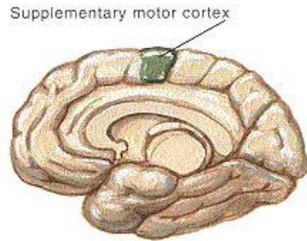
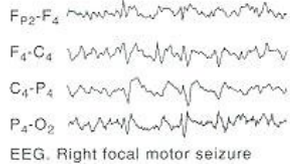
Seizure types in the elderly population

Holt-Seitz et al., 1999

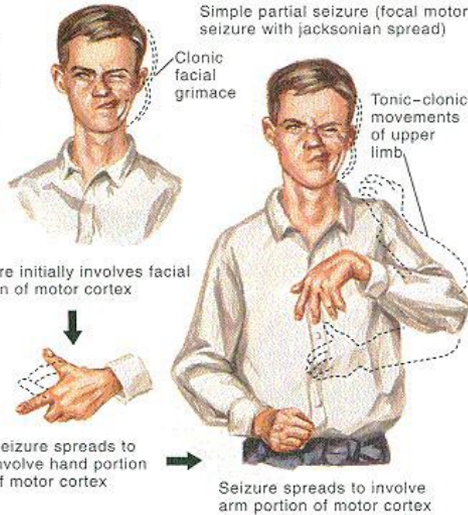
Partial Motor and Somatosensory Seizures



Motor cortex arranged in specific zones. Body areas involved in seizure may help localize seizure focus



Involvement of supplementary motor cortex results in versive movements

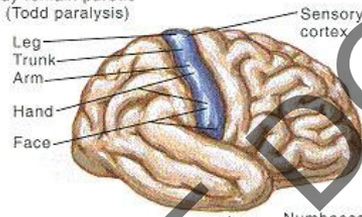


Seizure initially involves facial portion of motor cortex

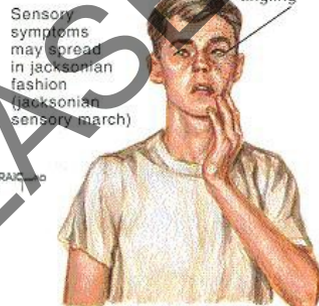
Seizure spreads to involve hand portion of motor cortex

Seizure spreads to involve arm portion of motor cortex

Affected areas may remain paretic for several hours (Todd paralysis)

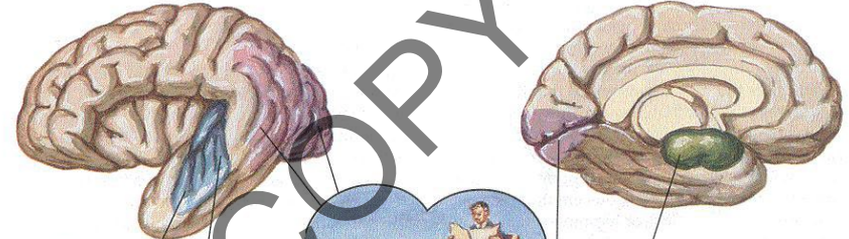


Numbness and tingling



Somatosensory cortex also arranged in anatomic zones

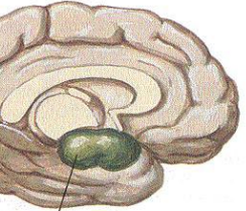
Partial Sensory and Autonomic Seizures



Formed visual hallucinations (posterior temporal, parietal, occipital, visual association cortex, temporal limbic cortex)



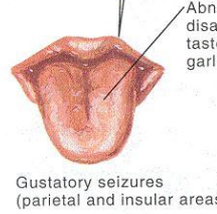
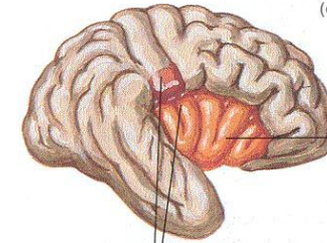
Unformed visual hallucination (occipital lobe)



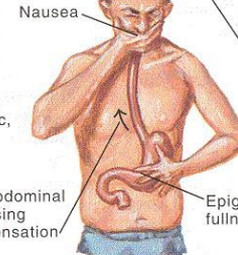
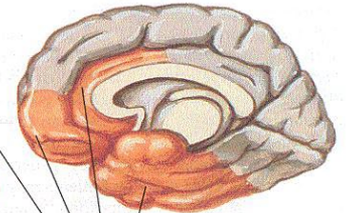
Usually disagreeable odor, such as burnt rubber



Olfactory seizures (uncinate gyrus)



Gustatory seizures (parietal and insular areas)

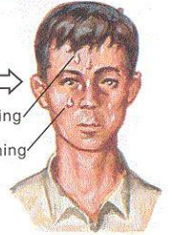


Nausea

Abdominal rising sensation

Epigastric fullness

Visceral



Sweating

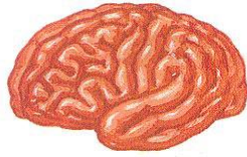
Flushing

Autonomic

Visceral and autonomic seizures (insular, cingulate gyrus, and frontal temporal areas)

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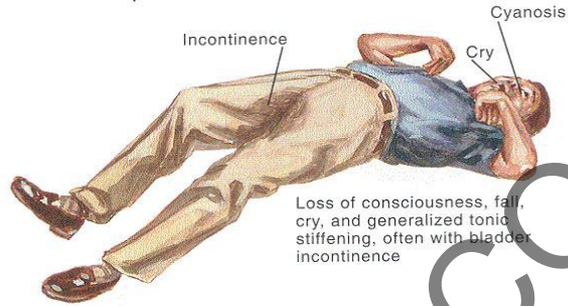
Generalized Tonic-Clonic Seizures



Simultaneous bilateral cortical seizure activity

JOHN A. CRAIG, M.D.
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Tonic phase



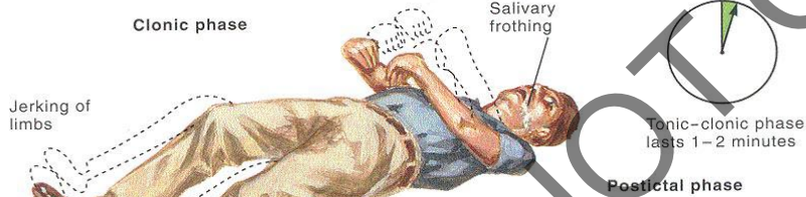
Incontinence

Cyanosis

Cry

Loss of consciousness, fall, cry, and generalized tonic stiffening, often with bladder incontinence

Clonic phase

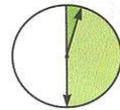


Jerking of limbs

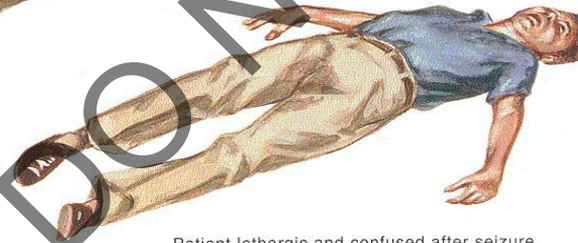
Salivary frothing

Tonic-clonic phase lasts 1-2 minutes

Postictal phase

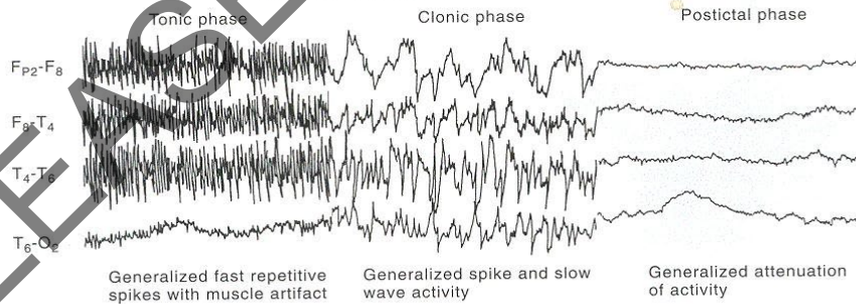


Postictal period may last minutes to hours

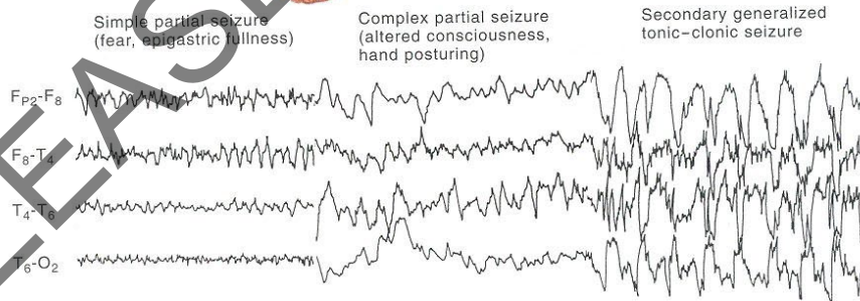
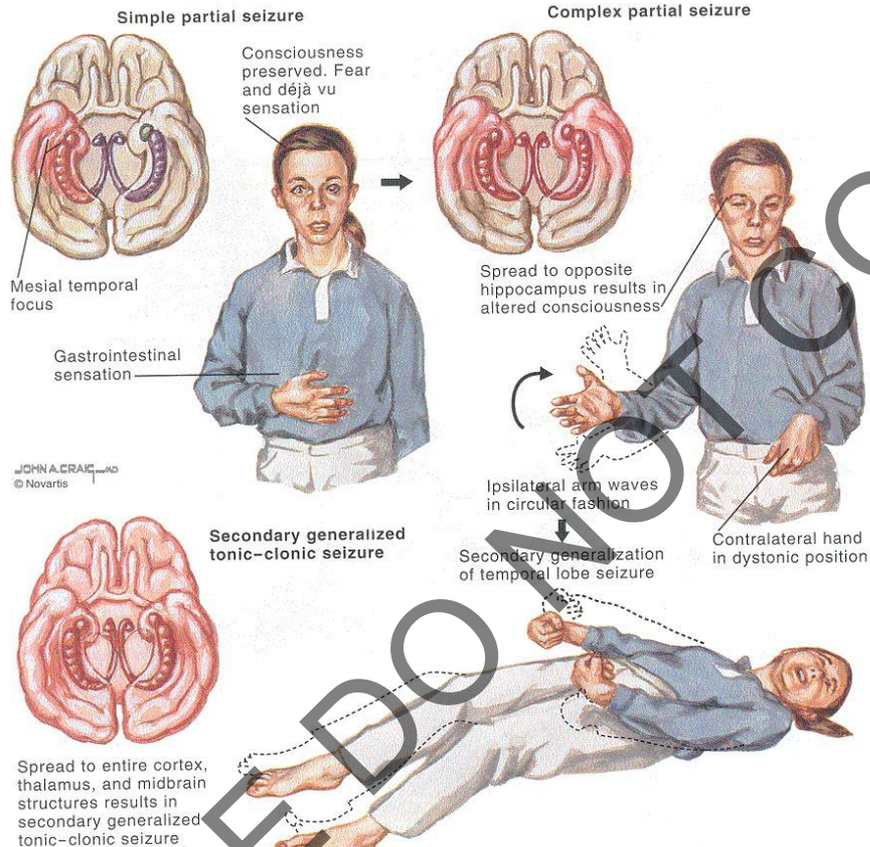


Patient lethargic and confused after seizure. Often sleeps

Stages of generalized tonic-clonic seizure



Temporal Lobe Epilepsy



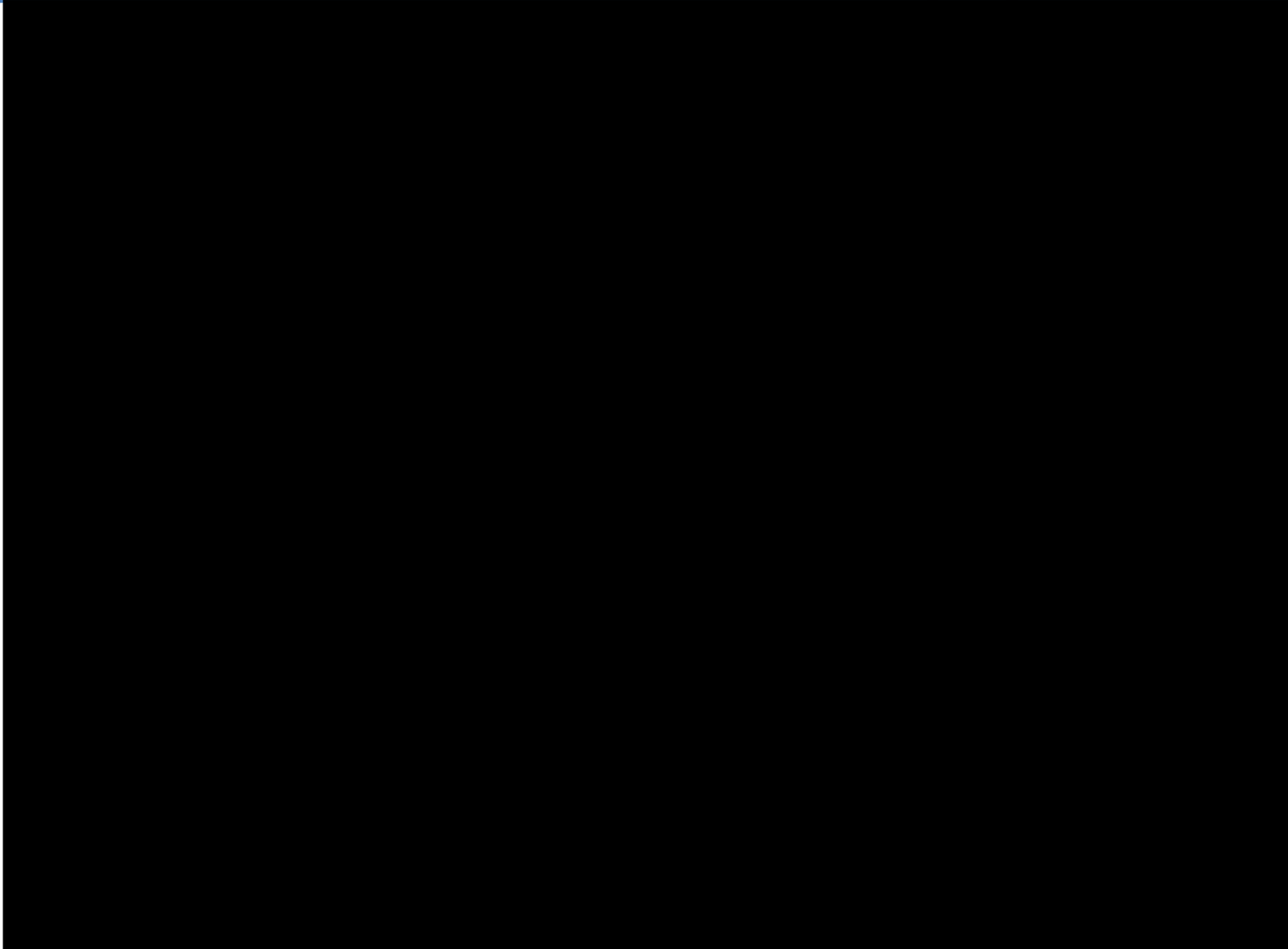
EEG. Progression of seizures in temporal lobe epilepsy

Focal aware (Simple partial seizures)
do not impair awareness or consciousness

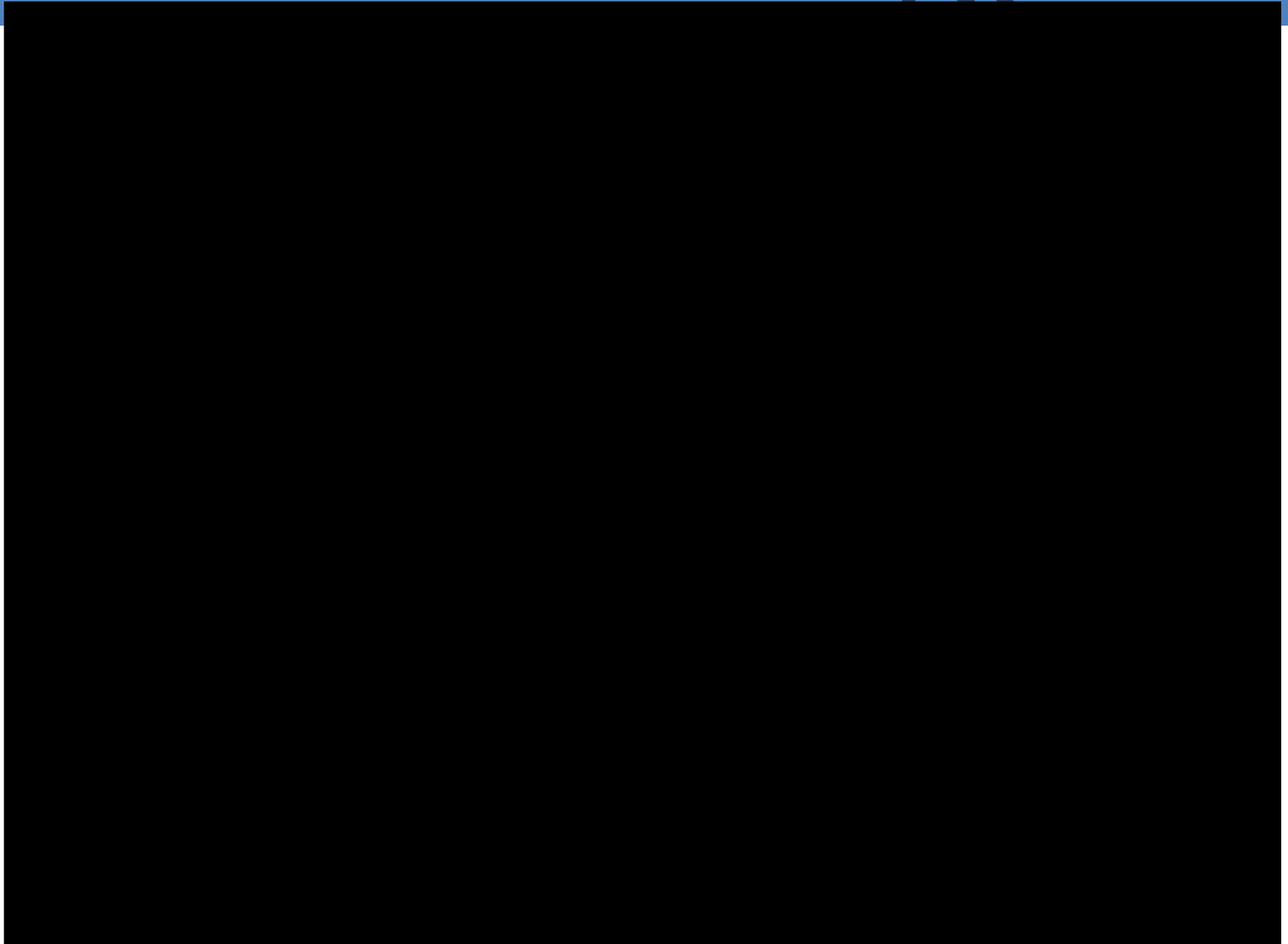


Focal impaired awareness (complex partial seizures) of temporal lobe origin have distinct characteristics

Copy



Some focal impaired awareness have minimal associated motor activity



Focal-onset seizures can progress to generalized seizures



The generalized tonic-clonic phase
has a very typical appearance

DRY



Acute response to seizures

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There is little to do acutely during a seizure for most types of seizures

- Absence, myoclonic, focal aware (simple partial) seizures
 - ▣ Usually no intervention necessary except reassurance when event ends
- Focal impaired awareness (complex partial) seizures
 - ▣ Allow event to run its course while preventing patient from encountering harm
 - ▣ Patients may become hostile or violent if actively restrained
- Generalized tonic-clonic seizures
 - ▣ Lay patient on side
 - ▣ Remove nearby objects that may cause harm
 - ▣ Do not place anything inside the mouth

Seizure First Aid

How to help someone having a seizure

1

STAY with the person until they are awake and alert after the seizure.

- ✓ Time the seizure
- ✓ Remain **calm**
- ✓ Check for **medical ID**



2

Keep the person **SAFE**.

- ✓ Move or guide away from **harm**



3

Turn the person onto their **SIDE** if they are not awake and aware.

- ✓ Keep **airway clear**
- ✓ **Loosen tight clothes** around neck
- ✓ Put **something small and soft** under the head



Call
911
if...

- ▶ Seizure lasts longer than 5 minutes
- ▶ Person does not return to their usual state
- ▶ Person is injured, pregnant, or sick
- ▶ Repeated seizures
- ▶ First time seizure
- ▶ Difficulty breathing
- ▶ Seizure occurs in water

Do
NOT

- ✗ Do **NOT** restrain.
- ✗ Do **NOT** put any objects in their mouth.
- ✓ **Rescue medicines can be given** if prescribed by a health care professional

Learn more: epilepsy.com/firstaid



epilepsy.com

24/7 Helpline: 1-800-332-1000

Life-threatening complications of isolated seizures are rare

- Vast majority of generalized tonic-clonic seizures last 1-2 minutes
- Emesis, aspiration, face-down positioning
- Cardiac arrest or prolonged respiratory arrest, anoxia

Generalized seizures > 2 minutes

- ABC's:
 1. Airway
 2. Breathing
 3. Circulation
- Lorazepam 2mg IV/IM/IN
- Call neurology

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What are the initial elements of evaluating a possible seizure?

- History
 - ▣ Details of the event
 - ▣ Past history of seizure-like symptoms or similar events
 - ▣ History of head trauma, febrile seizures, CNS infection
 - ▣ Family history of seizures
- Exam
 - ▣ General exam: evidence of head injury, meningismus, tongue bite
 - ▣ Neurologic exam: evidence suggesting a focal brain lesion
- Labs
 - ▣ Evidence of infection or metabolic disturbance: CBC, electrolytes, toxicologic screen, drug levels

Many paroxysmal events can appear similar to seizures clinically

- ❑ Syncope / orthostatic hypotension
- ❑ TIA
- ❑ Confusion/delirium
- ❑ Medication side effects
- ❑ Cardiac arrhythmia
- ❑ Migraine (without headache)
- ❑ Hallucinations
- ❑ Myoclonus
- ❑ Transient global amnesia
- ❑ Vertigo
- ❑ Movement disorder
- ❑ Nonepileptic seizure

The initial clinical diagnosis is based on some distinguishing features

Seizure

- ❑ Sudden onset
- ❑ Possible warning / “aura”
- ❑ Possible postictal state
- ❑ Automatisms
- ❑ “Positive” neurological symptoms
- ❑ Possible tongue bite/incontinence/limb jerking

Syncope

- ❑ Gradual onset
- ❑ Presyncopal warning
- ❑ Change in color / appearance
- ❑ Brief loss of consciousness, with rapid recovery
- ❑ Loss of tone

TIA

- ❑ Sudden onset
- ❑ Rapid recovery of focal neurological deficit
- ❑ Preserved consciousness
- ❑ “Negative” neurological symptoms

Acute confusion

- ❑ Waxing and waning
- ❑ Inattention
- ❑ No focal neurological deficit
- ❑ Drowsiness / decreased alertness / delirium
- ❑ Asterixis / myoclonus

Further neurodiagnostic testing could be indicated in certain cases

❑ Neuroimaging (MRI/CT)

- All new partial-onset seizure patients should have a nonurgent MRI
- If acute neurologic lesion is suspected, or injury sustained during a seizure, obtain an urgent CT or MRI

❑ EEG

- New onset seizure patients
- Can help to clarify partial- vs. generalized-onset and prognosticate risk of recurrence

What about the neurology triad?



□ MRI / CT

- If acute neurologic lesion is suspected, obtain an urgent CT or MRI
- All new partial-onset seizure patients should have a non-urgent MRI

■ EEG

- All new seizure patients should have an EEG
- Can help to clarify partial- vs. generalized-onset and prognosticate risk of recurrence

■ LP

- Should be performed if CNS infection is suspected
- Does not need to be automatically performed after any unexplained seizure

A single seizure does not generally warrant antiepileptic drug treatment

- The risk of recurrence after a single unprovoked seizure in next two years is 25-40%
 - ▣ Depends on seizure type, EEG findings
- The risk of recurrence after two unprovoked seizures is 80% or more
 - ▣ Most neurologists do treat after two episodes



There are many antiepileptic drugs, some of which have multiple indications

Classical

- 1857 – Bromides
- 1912 – Phenobarbital (PB)
- 1937 – Phenytoin (PHT)
- 1954 – Primidone
- 1958 – ACTH
- 1960 – Ethosuximide (ESM)
- 1963 – Diazepam
- 1974 – Carbamazepine (CBZ)
- 1975 – Clonazepam (CZP)
- 1978 – Valproate (VPA)

Newer

- 1993 – Felbamate (FBM), Gabapentin (GBP)
- 1995 – Lamotrigine (LTG)
- 1997 – Topiramate (TPM), Tiagabine (TGB)
- 1999 – Levetiracetam (LEV)
- 2000 – Oxcarbazepine (OXC), Zonisamide (ZNS)
- 2005 - Pregabalin (PGB)
- 2008 – Lacosamide (LCM), Rufinamide (RUF)
- 2009 – Vigabatrin (VGB)
- 2011 – Ezogabine
Clobazam
- 2012 – Perampanel
- 2013 – Eslicarbazepine
- 2016 – Brivaracetam
- 2018 - Epidiolex (6/25/2018)

Status epilepticus is a medical emergency

- Either a state of continuous seizure activity or a state in which seizures are recurring so frequently that there is no recovery in between
- The operational definition (when to begin acting) is 5 minutes

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There are many precipitating risk factors for status epilepticus

- Preexisting epilepsy
 - ▣ Medication noncompliance
 - ▣ Sleep deprivation or alcohol
 - ▣ Worsening underlying disease

- Metabolic / toxic disturbances
 - ▣ Hyperglycemia, hyponatremia, etc.
 - ▣ Drug intoxication

- Structural neurological causes
 - ▣ Acute stroke, hemorrhage
 - ▣ Head trauma

Summary

- Seizures are quite common in the population, but rare as a direct complication of TMS
- Most seizures in adults are focal-onset, but can become secondarily generalized
- Seizures have some distinguishing characteristics, but can still be confused with other types of events
- There is little to do other than ensure safety in the setting of an acute seizure
- The vast majority of seizures stop by themselves, but any lasting 5 minutes or more should be treated as a medical emergency

Questions

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ILAE 2017 Classification of Seizure Types Expanded Version ¹

