

Comprehensive Guide to EEG Interpretation, Seizure Semiology, and Epilepsy Management

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The study of clinical neurophysiology is a cornerstone of modern neurology, providing the objective data necessary to diagnose, classify, and treat various seizure disorders and encephalopathies. At the heart of this field lies **Electroencephalography (EEG)**, a non-invasive method for recording electrical activity of the brain. When combined with **Seizure Semiology**—the detailed study of the physical signs and symptoms during a seizure—clinicians can achieve a level of diagnostic precision that informs everything from pharmacological intervention to neurosurgical planning. This guide provides an in-depth technical analysis of these components, drawing from foundational neurophysiology and modern clinical management strategies.

The Neurophysiological Foundations of EEG

To understand EEG, one must first grasp the cellular mechanics that generate the signals recorded at the scalp. The EEG does not record action potentials; instead, it measures the **summation of excitatory and inhibitory postsynaptic potentials (EPSPs and IPSPs)** occurring primarily in the dendrites of pyramidal cells in the cerebral cortex. These cells are oriented perpendicularly to the cortical surface, creating a dipole.

Neuronal Membrane Physiology

The electrical potential of a neuron is maintained by the selective permeability of the cell membrane and the action of ion pumps. The **Resting Membrane Potential (RMP)**, typically around -70mV, is a product of the concentration gradients of Potassium (K⁺), Sodium (Na⁺), and Chloride (Cl⁻). The **Goldman-Hodgkin-Katz equation** provides the mathematical framework for this potential, accounting for the relative permeability of each ion:

$$V_m = (RT/F) \ln [(P_K[K^+]_{out} + P_{Na}[Na^+]_{out} + P_{Cl}[Cl^-]_{in}) / (P_K[K^+]_{in} + P_{Na}[Na^+]_{in} + P_{Cl}[Cl^-]_{out})]$$

When synaptic transmission occurs, neurotransmitters bind to receptors, causing local shifts in membrane potential. For a signal to be detectable on the scalp, thousands of these pyramidal neurons must fire synchronously. This **synchrony** is what allows the relatively weak electrical field to penetrate the meninges, cerebrospinal fluid, and skull via **volume conduction**.

Technical Specifications and EEG Performance

Executing a high-quality EEG requires strict adherence to technical standards. The **International 10-20 System** is the gold standard for electrode placement, ensuring that electrode locations are proportional to skull size and shape. The "10" and "20" refer to the fact that the actual distances between adjacent electrodes are either 10% or 20% of the total front-back or right-left distance of the skull.

Hardware and Signal Processing

Modern digital EEG systems utilize sophisticated amplification and filtering to isolate brain activity from environmental noise. Key technical parameters include:

- **Input Impedance:** Must be high (ideally >100 MOhms) to minimize signal loss.
- **Common Mode Rejection Ratio (CMRR):** A high CMRR (e.g., >100 dB) is vital to cancel out external electrical noise that is common to both the active and reference electrodes.

- **Sampling Rate:** According to the Nyquist-Shannon sampling theorem, the sampling rate must be at least twice the highest frequency of interest. For clinical EEG, a sampling rate of 256 Hz or 512 Hz is standard.
- **Filters:** Standard settings include a Low-Pass Filter (usually 70 Hz) and a High-Pass Filter (usually 0.5-1.0 Hz), along with a Notch Filter (50 or 60 Hz) to eliminate power line interference.

Montage Selection

The display of EEG data is organized into "montages." A **Bipolar Montage** compares the potential difference between two adjacent electrodes, which is excellent for identifying **phase reversals**—the hallmark of focal epileptiform discharges. A **Referential Montage** compares each electrode to a common reference (like the earlobe or a calculated average), which is superior for determining the absolute amplitude of a signal.

Seizure Semiology: The Philosophy of Signs

While EEG provides the "electro" data, **semiology** provides the "clinical" data. Derived from the Greek word *semeion* (sign), semiology in medicine is the systematic study of the signs observed during a paroxysmal event. It is a critical tool for **seizure localization** (identifying where the seizure starts) and **lateralization** (identifying which hemisphere is involved).

The Temporal Evolution of Semiology

A seizure is not a static event; it evolves through phases:

1. **Aura:** The subjective sensory or psychic phenomenon that precedes the observable seizure. This represents the initial focal discharge.
2. **Ictus:** The main seizure event, characterized by motor activity, loss of consciousness, or autonomic changes.
3. **Post-ictal State:** The recovery period, which may include confusion, Todd's paralysis (focal weakness), or aphasia.

Localization through Semiology

Specific signs act as "red flags" for certain cortical regions. For example, **ictal pouting** (dacrystic features) often points to frontal lobe involvement, while **gastric auras** (a rising sensation in the stomach) are strongly associated with the mesial temporal lobe. **Automated movements** (automatisms) like lip-smacking or hand-fumbling are common in temporal lobe epilepsy but are often more complex and violent in frontal lobe epilepsy.

Differential Diagnosis: Epileptic vs. Non-Epileptic Events

One of the most challenging tasks in neurophysiology is distinguishing true epileptic seizures from mimics.

Physiologic mimics include syncope (fainting), migraines, and sleep disorders like narcolepsy. **Psychogenic Non-Epileptic Seizures (PNES)** are events that resemble seizures but lack the corresponding electrographic discharges of epilepsy.

Comparison Matrix: Seizure vs. Syncope vs. PNES

Feature	Epileptic Seizure	Syncope	PNES
Onset	Sudden or preceded by aura	Gradual (blacking out)	Variable; often gradual
Duration	1-3 minutes	Seconds	Often >5 minutes
Eyes	Open (usually)	Closed or rolling up	Tightly closed
EEG Correlate	Spikes, sharp waves, slowing	Generalized flattening	Normal (no ictal change)
Post-ictal State	Prolonged confusion/fatigue	Rapid recovery	Variable; often rapid
Trigger	Usually none (spontaneous)	Orthostatic, vasovagal	Emotional stress

Advanced EEG Pattern Interpretation

Effective management requires the ability to distinguish between normal variants and pathological abnormalities. Normal variants, such as **Wicket Spikes** or **Rhythmic Mid-temporal Theta of Drowsiness (RMTD)**, can easily be mistaken for epilepsy by the untrained eye.

Epileptiform Abnormalities

Pathological markers are divided into interictal (between seizures) and ictal (during seizures) discharges. **Interictal Spikes** are transients lasting less than 70ms, while **Sharp Waves** last between 70ms and 200ms. The presence of a following slow wave is highly suggestive of an epileptogenic zone. In **Generalized Epilepsy**, patterns like the 3 Hz Spike-and-Wave are classic for absence seizures.

Quantitative EEG (qEEG) and Trend Analysis

In Intensive Care Units (ICU), continuous EEG (cEEG) monitoring generates massive amounts of data. **qEEG** uses mathematical algorithms like the **Fast Fourier Transform (FFT)** to convert raw EEG into color-coded trends. This allows clinicians to quickly identify subtle changes in brain state, such as **non-convulsive status epilepticus (NCSE)** or focal ischemia, which might be missed on intermittent visual inspection.

The Integrated Management Approach

Managing epilepsy is an iterative process that begins with accurate classification. The International League Against Epilepsy (ILAE) classification system relies on both the seizure type and the epilepsy syndrome.

Step-by-Step Diagnostic Workflow

1. Clinical History: Detailed interview to identify semiological clues.
2. Routine EEG: Initial 20-30 minute recording with activation maneuvers (photic stimulation and hyperventilation).
3. Video-EEG Monitoring (VEM): The gold standard. Simultaneous recording allows for the correlation of physical signs with electrical activity.
4. Neuroimaging: MRI (3 Tesla or higher) using specialized epilepsy protocols to find structural lesions like mesial temporal sclerosis or cortical dysplasia.
5. Pharmacological Selection: Choosing Anti-Seizure Medications (ASMs) based on the seizure type (e.g.,

Ethosuximide for absence seizures, Levetiracetam for focal seizures).

Troubleshooting EEG Artifacts

Artifacts can obscure real brain activity or, worse, mimic epileptiform discharges. Clinicians must be adept at identifying and neutralizing them:

- Biological Artifacts: EKG (heartbeat), EMG (muscle tension), Glossokinetic (tongue movement), and Eye Blinks.
- Environmental Artifacts: 60 Hz interference, electrode "pop" (sudden impedance change), and IV drip interference.

The Future of EEG and Seizure Analysis

The field is moving toward **automated seizure detection** using machine learning (ML) and artificial intelligence (AI). Deep learning models, specifically **Convolutional Neural Networks (CNNs)**, are being trained on thousands of hours of EEG data to identify subtle pre-ictal patterns that precede a seizure by minutes. This could lead to wearable devices that warn patients of an impending seizure, significantly improving safety and quality of life.

Furthermore, the integration of **Magnetoencephalography (MEG)** and **high-density EEG** (up to 256 channels) is refining our ability to perform non-invasive source imaging. By mapping the electrical dipoles onto the patient's individual MRI, surgeons can pinpoint the seizure focus with sub-millimeter accuracy, increasing the success rates of epilepsy surgery.

Ultimately, the marriage of neurophysiological science and clinical semiology remains the most potent tool in the epileptologist's arsenal. By meticulously analyzing the electrical "language" of the brain and the physical "signs" of the body, medical professionals can navigate the complexities of epilepsy with authority and precision, leading to better patient outcomes and a deeper understanding of the human brain's electrical landscape. The evolution from the first "Atlas of EEG" to today's digital, AI-enhanced landscape reflects a commitment to technical excellence and clinical rigor that continues to define the frontier of neurology.

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