Electroencephalography (EEG) remains the single most important physiologic test of the cerebral cortex. In temporal lobe epilepsy (TLE), focal interictal and ictal epileptiform discharges help in identifying the involved cortical areas. Several nonepileptiform EEG abnormalities can also be seen in TLE. Scalp and sometimes invasive EEG recordings are needed for accurate localization of the seizure onset. The development of video-EEG monitoring has allowed careful correlation of clinical semiology with simultaneous EEG recordings. This paper will review the EEG characteristics of TLE.

**EEG Recording**

**I. Scalp EEG recording**

The 10-20 electrode system is an internationally accepted standard method of measurement and application of EEG scalp electrodes. The standard electrodes can detect only up to 58% of all spikes. Therefore, additional measured and more closely spaced scalp electrodes placed midway between the standard electrodes of the 10-20 system can provide further localization of epileptiform discharges in TLE. Several “non-standard” electrodes can be used to further evaluate the EEG abnormalities including sphenoidal, nasopharyngeal, anterior temporal...
(T1/T2), and surface electrodes applied over the mandibular notch (MN) or zygoma\textsuperscript{7,9}. Sphenoidal electrodes are inserted bilaterally through the skin below the zygomatic arch, 2-3 cm anterior to the tragus and directed posterosuperiorly in the direction of the foramen ovale\textsuperscript{10}. This procedure should be performed with local anesthesia by a qualified physician. Flexible wire electrodes, insulated and bared at the tip, are placed about 3 cm deep by attaching them to the external shaft of a 22 gage special (lumbar puncture like) needle. The needle is then removed leaving the recording electrode in place; the external wire is coiled to relieve tension and taped to the skin.

Nasopharyngeal electrodes are uncomfortable and have not consistently demonstrated superior sensitivity for spike detection when compared to the other non-standard electrodes\textsuperscript{11,12}. Therefore, nasopharyngeal electrodes are no longer in widespread use. Sphenoidal, T1/T2, and MN electrodes can record spikes not detected by standard electrodes (Figure 1). Sphenoidal spikes are almost always detected by T1/T2 or MN electrodes, although with slightly lower amplitudes\textsuperscript{13}. Recording with multiple non-standard electrodes simultaneously to obtain a more comprehensive evaluation of the electrical field has been recommended\textsuperscript{7}.

Few studies have evaluated the ictal (as opposed to interictal) merits of these electrodes or have addressed the issue of montage design for interictal and ictal TLE abnormalities\textsuperscript{14}. The ability to perform off-line montage reformating with digital EEG technology diminishes the importance of initial montage selection. Ives et al studied ictal EEGs in patients with sphenoidal electrodes by using a coronal sphenoidal montage as compared to an anterior-posterior temporal montage\textsuperscript{14}. The sphenoidal montage was superior in the detection of ictal EEG changes.

II. Invasive EEG recording

Invasive intracranial techniques using subdural or intracerebral depth electrodes can be used to improve diagnostic accuracy if surface EEG recording yields inconclusive or ambiguous results\textsuperscript{15}. Recent shifts in surgical candidacy toward younger age groups resulted in increased utility of invasive monitoring especially in patients with atypical clinical semiology or normal imaging studies\textsuperscript{16}. Subdural electrodes are inserted surgically to record over the cerebral cortex. Electrode grids are square or rectangular in shape with small platinum or stainless steel disks embedded into soft silastic with several contact points. Different sizes and shapes can be selected dependant on the anatomic area to be covered and require a craniotomy for insertion. Electrode strips, of various sizes, consist of a row of contacts and are usually inserted through a burr hole.

Intracerebral or depth electrodes are used to record from within the temporal lobe using rigid or flexible probes. Multiple-lead intracerebral electrodes can be implanted stereotactically (stereoelectroencephalography) to allow sampling from multiple medial (amygdala, hippocampus, entorhinal cortex, insular cortex) and lateral (temporal pole, gyr, parietotemporal and

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure1.png}
\caption{Anterior temporal spike with phase reversal at F7 and shown in three montages, (A) antero-posterior, (B) common average reference (CAR), and (C) coronal with mandibular notch (MN) electrodes. Notice that the spike is better illustrated when MN electrode was used. (A = low frequency filter [LFF] 0.3 sec, high frequency filter [HFF] 70 Hz, Sensitivity 7.5 µV/mm / B&C = LFF 0.3 sec, HFF 70 Hz, Sensitivity 10 µV/mm).}
\end{figure}
occipitotemporal junctions) temporal lobe structures. It is important to note that scalp electrodes detect activity generated from a large cortical surface whereas subdural or depth electrodes will detect potential changes which occur over only a few millimeters of cortex. Interictal spikes not otherwise seen on scalp EEG can be easily identified with invasive electrodes (Figure 2). Invasive electrodes are more sensitive in detecting spikes and seizures from a localized area but may miss epileptiform activity which occurs in adjacent regions only a short distance away.

Indications for invasive recording include lack of seizure lateralization and/or localization, seizure onset discrepant with other data (clinical, EEG, and/or imaging), and relation of seizure localization to eloquent cortex. Advantages of invasive monitoring in TLE include improved spatial resolution and increased sensitivity (early signal), no attenuation from scalp and skull, reduced ictal muscle and movement artifact, and utility for direct electrical stimulation of cortex. Disadvantages include limited cortical sampling or risk for sampling error (tunnel vision), and risk of significant complications (2-3%). These complications include aseptic meningitis (subdural more than depth electrodes), cerebrospinal fluid leak, cerebral edema, brain contusion, scalp infection, and bleeding (depth more than subdural electrodes).

Foramen ovale electrodes are occasionally used to record from mesial temporal structures without requiring penetration of the skull. A multi-contact flexible electrode is placed in the ambient cistern with the aid of a needle inserted through the foramen ovale. These electrodes are not as close to hippocampal structures as intracerebral electrodes and do not allow as large a recording field as grids and strips but detect mesial temporal EEG discharges better than sphenoidal and scalp electrodes.

**Figure 2:** Right anterior temporal spikes on subdural electrodes (top channels) using referential montage. Simultaneous scalp recording (bottom 2 channels) did not reveal the interictal spikes. (low frequency filter [LFF] 0.1 sec, high frequency filter [HFF] 70 Hz, Sensitivity 100 µV/mm for subdural channels / LFF 0.3 sec, HFF 70 Hz, Sensitivity 20 µV/mm for scalp channels).

**Figure 3:** Benign epileptiform transients of sleep (BETS) maximally expressed on the right temporal region on antero-posterior (AP) bipolar montage (A). Their low voltage and wide field is best seen on common average reference (CAR) (B). Notice the central spindles of sleep seen in both figures. (A&B = LFF 0.3 sec, HFF 70 Hz, Sensitivity 3 µV/mm).
When extracranial recordings are equivocal, foramen ovale electrodes offer a less invasive alternative to a more complete intracranial evaluation or can be used in association with grids and strips.

**ARTEFACTS**

The differentiation of artefacts from EEG abnormalities can be problematic on video-EEG recordings. Unusual artefacts not seen in standard laboratory EEG recording include mechanical artefacts from altered electrode/scalp contact, intermittent lead wire disconnection, body movement, or movement of the electrode cable. Artefacts produced by rubbing or scratching of the scalp and other rhythmic movements of head or extremity can result in patterns that must be differentiated from ictal discharges. Some artefacts are characteristic of the oro-alimentary automatisms of TLE (e.g. rhythmic chewing and swallowing). Rhythmic slow waves can also be due to chipped electrode coating, instability of the electrode scalp interface, and electrode wire movement. A conservative interpretation of unusual or equivocal EEG events is mandatory, particularly when the patient’s activity cannot be verified by video review. Interictal sharp waves noted only during active wakefulness should be interpreted with caution. Ictal rhythms should have a recognizable field with typical ictal progression (evolution of rate and voltage), spread to other regions, or postictal slowing. Intracranial recordings tend to have fewer physiologic artefacts, such as chewing and eye blinks.

**NORMAL EEG VARIANTS**

The differentiation of some normal or benign EEG transients from EEG abnormalities poses a problem in EEG interpretation and its clinical correlation. Normal EEG variants refer to waveforms that mimic epileptiform abnormalities. In TLE, three important normal variants needs to be recognized to avoid confusion; small sharp spikes of sleep, wicket spikes, and psychomotor variant.

Small sharp spikes of sleep, also known as benign epileptiform transients of sleep, are sharp, small waves occurring unilaterally or bilaterally (often asynchronously), especially in the temporal and frontal regions (Figure 3). Benign epileptiform transients of sleep have large, generally sloping fields in contrast to the restricted field of temporal lobe spikes. They are most often noted in adults during light sleep and have no association with TLE.

Wicket spikes have no pathological significance. They occur almost exclusively in adults and are thought to be due to harmonics of two or more waveforms that combine to form pseudospikes. They have a typical arciform, sharply contoured appearance (Figure 4). When temporal wicket spikes have large amplitude, they may appear similar to pathological spikes.

![Figure 4: Wicket spikes involving T4 on AP bipolar montage. Notice that there is no after coming slow wave or disruption in the background rhythm. (LFF 0.3 sec, HFF 70 Hz, Sensitivity 5 µV/mm).](https://www.cambridge.org/core/core)

When extracranial recordings are equivocal, foramen ovale electrodes offer a less invasive alternative to a more complete intracranial evaluation or can be used in association with grids and strips.

![Figure 5: Rhythmic mid-temporal discharge shown on AP bipolar montage. Notice the asymmetry with onset on the right mid temporal region (A), and offset on the other side (B). (A&B = LFF 0.3 sec, HFF 70 Hz, Sensitivity 7.5 µV/mm).](https://www.cambridge.org/core/core)
However, they can be easily distinguished by their morphology, lack of after coming slow wave, and superimposition on normal background rhythms. Psychomotor variant (rhythmic mid-temporal discharge) can be confused with an ictal rhythm. This benign rhythm occurs as asymmetrical runs of theta activity primarily in the temporal regions, lasting for a few seconds or as long as 45 seconds (Figure 5). This waveform is a harmonic of two or more rhythms. The waves often have a bifid appearance, start suddenly on one side and last for several seconds before terminating abruptly. The rhythm does not evolve in frequency or amplitude and has no associated clinical symptoms, all of which help in differentiating it from an ictal rhythm19.

INTERICTAL EEG ABNORMALITIES

I. Background abnormalities

Focal and regional background slowing (theta and delta) is a nonspecific finding that can be seen in a variety of central disorders, including epilepsy. Persistent focal or regional slowing usually suggests an underlying "structural" abnormality while intermittent slowing is a "functional" abnormality that may indicate an underlying epileptic zone. Lateralized arrhythmic (irregular) delta activity may be found in up to 66% of patients with TLE and is highly concordant with temporal spiking20. These authors concluded that their lateralizing value is similar to that of temporal spiking. Temporal intermittent rhythmic delta activity (TIRDA) is a more specific and accurate interictal indicator of temporal lobe seizures21. Trains of TIRDA were found in up to 90% of patients with MRI demonstrated hippocampal atrophy and TLE22. Others confirmed a significant association with mesial rather than lateral TLE23. The delta lateralized to the side of atrophy with accuracy equal to spikes24. These authors concluded that focal TIRDA is a reliable indicator of the side of the epileptogenic focus and is considered an EEG marker of an epileptogenesis that involves the mesial temporal structures. Therefore, focal or regional polymorphic delta in patients with established TLE should not be ignored when assessing the side of ictal onset.

II. Epileptiform discharges

The classic interictal EEG abnormality of temporal lobe seizures is a spike (or sharp wave) that has an electronegative peak over the anterior temporal region (F7/F8) as shown in Figure 125. Anterior temporal spikes are seen in most (94%) patients with medial TLE26. Computer field mapping show that the peak electronegative field of these spikes is anterior and inferior to the standard 10-20 electrode positions (F7/F8); this field accounts for the improved sensitivity of the additional non-standard electrodes described in the EEG recording section27. Most patients with sphenoidal maximum spikes have hippocampal originating seizures28. Sphenoidal spikes are also seen in temporal neocortical and orbital frontal onset seizures28. Thus, while sphenoidal maximum spikes are sensitive to mesial TLE, they are not highly specific. Mid temporal (T3/T4) and posterior temporal (T5/T6) spikes are more likely to originate from the temporal neocortex.

Sleep increases the spike frequency in TLE where spike foci unapparent during wakefulness may be found29. In young patients, a natural sleep recording is always superior to a drug induced sleep as spike activation may occur mainly in the lighter stages of sleep30. Sleep deprivation is therefore recommended to achieve this goal, however, this procedure should not be requested routinely because of the possible risk of seizure provocation.

It seems intuitively obvious that patients with a unifocal or very strongly unilaterally predominant interictal temporal spikes have a strong likelihood of ictal onset from the same temporal region31. Several studies have demonstrated a significant correlation between the side of ictal onset and the side with a preponderance of interictal spikes32,33. This concordance is best if unilateral hippocampal atrophy is associated33. Although many publications support the value of the site of maximum interictal spike frequency for predicting the site of ictal onset, a predominant lateralized surface interictal finding may still incorrectly predict the temporal lobe of ictal origin. Ictal recording is therefore recommended before surgical management is considered34. Spikes associated with TLE are not always confined to one temporal lobe. Up to 30% of patients

Figure 6: Ictal rhythm in the theta range with right anterior temporal onset (A) on AP bipolar montage. In the postictal phase (B) lateralized polymorphic delta rhythm is shown 45 seconds from seizure offset. (A = LFF 0.3 sec, HFF 70 Hz, Sensitivity 5 µV/mm / B = LFF 0.3 sec, HFF 70 Hz, Sensitivity 7.5 µV/mm).
with TLE demonstrate independent bitemporal discharges, particularly when prolonged telemetry recording is used. These patients could still have unilateral pathology and subsequent successful surgical outcome, particularly when the neuroimaging abnormality and ictal events were concordant. Bisynchronous spike waves are occasionally encountered in patients with focal temporal EEG abnormalities. The mechanism of production of these diffuse waves is unclear; however, their presence should not deter the search for a surgically resectable focus.

**Ictal EEG Abnormalities**

### Ictal Scalp Recorded Rhythms

Ictal EEG recording is a major component of presurgical evaluation. Ictal rhythms include background attenuation, start-stop-start phenomenon, irregular 2-5 Hz lateralized activity, 5-10 Hz sinusoidal waves or repetitive epileptiform potentials (Figures 6,7). As well, no or minimal EEG change may be noted (see later section on invasive EEG recording). The multiple scalp EEG expressions of temporal lobe originating seizures are likely a consequence of several integrated factors including the location of ictal onset (mesial vs. lateral), the underlying pathology (hippocampal sclerosis or atrophy vs. neoplasm), routes and speed of seizure propagation. Factors that can alter seizure propagation and therefore the EEG expression include the state of the patient (wake vs. sleep), antiepileptic drug use, and whether the patient had unifocal vs. multifocal epilepsy.

Focal or regional background attenuation as the initial EEG change of scalp-recorded temporal lobe seizure is seen in up to 25% of seizures. In a smaller percentage, the seizure may appear to start, stop, and restart “start-stop-start phenomenon.” The initial “start” typically has a more restricted field than the restart. If the initial start is overlooked, the restart may be misinterpreted as the actual seizure onset and therefore considered non-localizing, while in fact the initial start is typically focal and localizing.

Other initial ictal EEG features of TLE include sinusoidal waves and repetitive epileptiform potentials (47% of cases). As seizures evolve, there is a change in frequency (usually an increase) with spread to other EEG channels as a result of seizure propagation. Only in the final stages of the seizures is there a decrease in wave frequency. A lateralized ictal change of irregular polymorphic 2-5 Hz temporal rhythm is most often associated with TLE originating from the neocortex. A lateralized ictal change of rhythmic 5-10 Hz sharp activity, maximally at F7/F8 or sphenoidal electrode position is most commonly seen in hippocampal onset TLE (Figures 6,7). This rhythm usually occurs within 30 seconds of seizure onset and has high specificity for hippocampal onset. However, this rhythm is not directly caused by seizures in the hippocampus (see later section on invasive EEG recording).

Another abnormality of possible significance is the increased heart rate just before the onset of ictal EEG discharges. This phenomenon is associated with EEG pattern of TLE originating from the mesial structures, suggesting activation of the neuronal circuits involved in sympathetic regulation.

There are no prospective studies to answer the question of how many ictal recordings should be obtained in a given patient for definite lateralization and localization of TLE. Patients with unilateral interictal spikes were much more likely to have all seizures “concordant” than patients with bilateral interictal spikes. Among the group of patients with “conflicting” seizures, all patients demonstrated the conflicting seizures by the fourth recorded seizure. These results were similar to those of Blum who determined, using a statistical modeling technique that five recorded seizures were required to have a 95% chance of avoiding a conflicting seizure and only four seizures were needed if all interictal spikes were unilateral.

Potential difficulties with scalp recordings include an inability to interpret the EEG because of excessive muscle and movement artifacts, a paucity of unequivocal changes, inter-observer disagreement, and false lateralization. The overall accuracy of scalp lateralization ranges between 60 to 83%. Other causes of such errors include temporal ‘plus’ epilepsy and extratemporal originating seizure. Temporal ‘plus’ epilepsy is characterized by temporal lobe seizures that also involve the neighboring structures such as the orbitofrontal cortex, insula, frontal and parietal opercula, and tempo-parieto-occipital junctions. These are currently most accurately identified by invasive EEG monitoring but scalp EEG can be useful in differentiating TLE from temporal ‘plus’ epilepsy. In temporal ‘plus’ epilepsy, the interictal EEG more frequently exhibited...
Figure 8: 1) Ictal scalp EEG showing no changes at seizure onset (A) on coronal montage. After 60 seconds, an ictal rhythm is seen on the left side (B), falsely lateralizing the seizure because of "burnt out" right hippocampus (see also fig 8.2). (A&B = LFF 0.3 sec, HFF 70 Hz, Sensitivity 10 µV/mm). 2) Ictal subdural EEG of the same patient in Figure 8.1 confirming that the ictal onset was indeed from the right side (A&B). As the seizure terminates, propagation is seen to the left side (C). This is when the falsely lateralizing scalp rhythm (Fig 8.1) was seen. (A = LFF 0.1 sec, HFF 70 Hz, Sensitivity 75 µV/mm / B&C = LFF 0.1 sec, HFF 70 Hz, Sensitivity 100 µV/mm).
bilateral or precentral abnormalities, while the ictal EEG more frequently suggested early involvement of the anterior frontal, temporoparietal, and central regions.

Occasionally, a seizure starts on one side but terminates ipsilaterally as the seizure propagates to the other temporal lobe (Figure 7). False lateralization of ictal scalp EEG can also be rarely secondary to the so called "burnt out hippocampus"48. Severe advanced hippocampal atrophy and sclerosis results in failure of ipsilateral neocortical propagation and recognition on scalp EEG ipsilaterally49. Several seconds after seizure onset, the ictal discharge appears on the contralateral side resulting in false lateralization (Figure 8). Therefore, the ictal rhythm cannot be used in isolation for definite ictal localization.

II. Intracranial recorded rhythms

Scalp EEG incompletely records many aspects of intracranial activity50. The sensitivity, specificity, and interobserver reliability of scalp ictal EEG for localization of the epileptogenic zone have been disputed. If presurgical surface EEG recording, with additional non-standard EEG electrodes (e.g. sphenoidal or mandibular notch electrodes) cannot provide an adequate answer, invasive intracranial techniques (primarily subdural or intracerebral electrode recordings) can be used to improve localization of the epileptogenic zone51. Intracranial EEG directly records paroxysmal activity with high sensitivity, relatively artefact free recording, and excellent temporal resolution in the order of milliseconds52,53. In the pre-MRI era the need for such intracranial recording in the determination of an epileptogenic focus was emphasized at some centers where invasive procedures were required in most patients prior to epilepsy surgery. It is now recognized that in most cases of medial temporal lobe epilepsy, standard EEG combined with sphenoidal electrodes and evidence of mesial sclerosis on MRI, will suffice to define the surgical focus.

In addition to the ictal rhythms described in the scalp EEG section, a low voltage fast (>20 Hz) rhythm can be seen at the onset of intracranial recorded TLE54. It has been demonstrated using a combination of surface and depth electrodes in patients with TLE that a significant proportion of spatially restricted seizures seen with depth recordings produce no visible changes in surface EEG recordings (Figure 8)55. Only 19% of auras and 10% of subclinical seizures are accompanied by scalp EEG changes and occasionally, seizures produce bilateral scalp abnormalities which are not indicative of the side from which depth recorded seizure activity was initiated56. A comprehensive systematic study of scalp-intracranial ictal EEG findings in TLE was reported by Pacia and Ebersole55. Simultaneous scalp electrodes, mesial temporal depth probes, subdural strip electrodes, and subdural grids were used. Patients with seizures beginning at the hippocampal contacts of the depth electrode had ictal rhythms recorded from the mesial subdural strip electrodes. Simultaneous scalp recordings showed either no change or a diffuse disruption of background. Only when the seizure spread to the basal and inferolateral subdural contacts did a 6-7 Hz rhythm appear in the inferior and standard scalp electrodes. The authors concluded that a scalp recorded 5-9 Hz rhythm is highly associated with hippocampal onset seizures but is not a direct manifestation of hippocampal ictal activity. This characteristic rhythm appears to reflect recruitment of adjacent temporal cortex and cortico-hippocampal interactions as seizures initiated in the neocortex did not demonstrate this scalp pattern until the hippocampus became involved. Seizures confined to the mesial-basal cortex produce a different scalp rhythm that is manifest on the scalp as a 5 Hz rhythm maximally expressed, perhaps unexpectedly, at the vertex. This rhythm is of opposite polarity from that seen with the intracranial basal electrodes and although seemingly unilaterализed, is in fact quite localizing because of the particular cortical orientation required for its generation57. The intracranial correlate of this seizure type was a focal or regional fast rhythm (beta frequency or higher). Such high frequency changes were never seen with scalp recordings in this study. The corresponding scalp recordings consisted of a high voltage, irregular, delta frequency pattern. A further study showed that the location of seizure onset in the temporal lobe was related to the degree of hippocampal pathology56. Marked hippocampal atrophy and high grade hippocampal sclerosis were both associated with initial ictal discharge restricted to the hippocampus on depth recording56.

Other authors studied the ictal onset of TLE secondary to other pathologies, such as temporal dysembryoplastic neuroepithelial tumor, using invasive subdural EEG recording57. These patients frequently had more than one ictal onset zone, which were detected more frequently in the tissue adjacent to the tumor rather than within the tumor or in the mesial temporal structures. Such accurate localization of the ictal onset is critical in dysembryoplastic neuroepithelial tumor patients as the surgical outcome was better when all ictal onset zones were completely resected57.

POSTICTAL EEG ABNORMALITIES

Excess muscle and movement artefacts may obscure the EEG features of scalp recordings resulting in difficulties in identifying the seizure onset. Several postictal features are described in TLE including polymorphic delta activity, regional attenuation, or activation of focal spikes. Of these, regional delta was most frequently observed by Kaibara et al in 57% of patients with scalp EEG recorded seizures58. Other authors identified lateralized delta in up to 69% of postictal EEGs59. Postictal change, whatever its nature appeared principally or exclusively ipsilateral to the side of seizure onset in all cases (Figure 6B). In the Williamson et al study of patients rendered seizure-free after a temporal lobectomy, 67% of patients had lateralized postictal slowing that was concordant to the side of seizure origin in 100% of instances60. Similarly, Walczak et al found that when postictal changes were present and could be lateralized (50% of patients), these findings were concordant with the side of seizure onset in 96-100% of patients who were at least two years seizure-free after surgery61. The postictal changes may be situated principally over the region most intensely involved in the seizure, which may not always coincide with the region of seizure onset. We evaluated the relationship of postictal delta to the side of seizure onset in patients with TLE by reviewing postictal EEGs with the EEGers blind to clinical and EEG data62. Lateralized postictal delta was present in 64% of all ictal EEGs and in at least one record from 22 patients (76%). The two EEGers agreed on the postictal EEG findings in 73 of the 80 seizures (kappa=0.88). Lateralized postictal delta, when present, was concordant with the side of surgery in 96% of the EEGs. The results confirmed
that lateralized postictal delta strongly predicts the side of seizure onset in TLE. In another analysis, the postictal EEG features of simple partial (SP) seizures were compared to those of complex partial (CP) and secondary generalized (SG) seizures in patients with documented TLE. Postictal delta was identified in 55%, 85%, and 100% of postictal EEG segments of SP, CP, and SG seizures respectively. Unilateral postictal delta was identified in 36%, 76%, and 44% of postictal EEG segments of SP, CP, and SG seizures respectively. The EEGs of the partial seizures (SP and CP) were seven times more likely to reveal unilateral PDA when compared to the EEGs of SG seizures, however, unilateral postictal delta was more likely seen in the CP group. Unilateral postictal delta was ipsilateral to the side of temporal lobeectomy in 94% of EEGs. These findings confirm the potentially important role of postictal EEG in seizure lateralization, particularly if the ictal rhythm was missed or obscured. If polymorphic delta activity emerges in the EEG, it should raise the suspicion of a preceding ictal event.

**SUMMARY AND CONCLUSIONS**

Scalp EEG is needed for accurate lateralization and localization of the seizure onset in TLE. Important EEG background abnormalities include TIRDA, which is a reliable indicator of epileptogenesis that involves the mesial temporal structures. Standard EEG can detect up to 58% of all interictal spikes, therefore, additional “non-standard” electrodes can be used to further evaluate the EEG abnormalities. Ictal rhythms in TLE includes background attenuation, start-stop-start phenomenon, irregular 2-5 Hz lateralized activity, and 5-10 Hz sinusoidal waves or repetitive epileptiform potentials. Postictal delta can be lateralized in 60% of patients with TLE and is concordant with the side of seizure onset in most patients. Invasive EEG is needed in patients with atypical clinical semiology or normal imaging studies. Invasive electrodes can be implanted on the surface of the brain or stereotactically to allow sampling from multiple medial and lateral temporal lobe structures. Electroencephalography abnormalities should be correlated with the clinical and imaging data as accurate identification of the seizure origin is more likely to be achieved if focal EEG onset and MRI findings were concordant with the clinical semiology.

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