American Clinical Neurophysiology Society

Guideline 7: Guidelines for EEG Reporting
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The purpose of this guideline is to provide a standardized format for reporting the results of adult routine scalp electroencephalography (rsEEG). The only-moderate interobserver reliability of EEG interpretation may be partly explained by the different reporting styles utilized (Beniczky S et al., 2013), and there is significant variability in the observation of guidelines for EEG reporting (Tatum WO, 2013). Computer-based remote access technology has become more sophisticated, and video is now “routine” during rsEEG, prompting the need to revise and update the earlier ACNS guideline on “Writing an EEG Report” (ACNS Guideline 7, J Clin Neurophys 2006). To assist in producing useful information for clinical and research purposes, standardized terminology and following an orderly approach to EEG reporting is recommended (Benbadis SR, Kaplan PW, 2013).

This guideline is designed to outline the conditions and parameters of EEG recording, including a description of the record obtained and the final impression that summarizes the EEG’s visual analysis. Its framework is intended to be useful to a clinician providing general neurologic care (including in the primary care setting) who may not be an expert on the technical aspects of EEG or on the terminology in the rsEEG report (Chatrian et al., 1974). Proper interpretation of the results reported depends on minimum technical standards for the performance of an EEG (see “Minimum Technical Requirements for Performing Clinical EEG” available at http://www.acns.org/practice/guidelines). Also, it is clear that consistently higher inter-observer agreement occurs when there is a forced choice paradigm using a limited set of EEG terms (Gerber et al., 2008). This guideline intends to provide a framework for the EEG report to address the features as normal or abnormal, with subsequent specification of their clinical importance. The significance should be evident to the clinician, and the findings readily interpretable within the patient-specific context of the rsEEG recording.

When reporting specialized types of EEG (e.g., electrocerebral inactivity, or neonatal EEGs) or EEG recordings in special settings, or for prolonged durations (e.g., continuous EEG in critical illness or during video-EEG monitoring), there may be modification with special formats more applicable to those specific settings. In these situations, the description of technical details should be enhanced and more complete than is required for standard rsEEG reporting. Also, the guidelines described below are not intended to be the sole means of reporting for institutions where research indexing may apply. Guidelines have been developed by the American Clinical Neurophysiology Society (ACNS) to provide the appropriate means of recording in some special situations. Standardized reporting of clinical neurophysiologic procedures has been implemented successfully (Hirsch LJ et al., 2012). The current guideline on “EEG Reporting” complements the earlier “Writing an EEG Report” and supplements those on non-routine recording situations and locations, available at http://www.acns.org/practice/guidelines.
Format for Reporting

A standard format for rsEEG reporting should include five sections:  History, Technical Description, EEG Description, Impression, and Clinical Correlation.

**History.**  The history section is an aid to interpretation of the rsEEG and should include the reason for obtaining the recording and any relevant clinical information, as well as identification of the patient and EEG recording.

Templates for reporting EEGs should supply information and demographics about the patient.  Personal information should include patient identification, including the medical record number (or social security number) and clinical/hospital EEG record number, in addition to the last name, first name, gender, and date of birth and age at the time of the recording.  This essential information should be entered by the person who prepares the recording for final interpretation.  A worksheet should provide additional information prepared by the technologist performing the EEG recording and should include relevant medical history, pertinent medications including antiseizure drugs, neuroimaging results, note of any cranial operations, the reason for obtaining the recording, and whether prior EEGs have been performed.

**Technical Description.**  The technical description should detail the conditions and parameters of the recording, including the duration of the record, as well as the date and location of acquisition and interpretation -- to ensure that both are identified in case the rsEEG is interpreted days after the recording.  Technical parameters should include the number of electrodes used and that the placement was in accordance with the 10-20 or 10-10 International System of Electrode Placement.  Minimum standards for performing an EEG (e.g. head measurement) are required (see also Guideline 1: Minimum Technical Requirements for Performing Clinical EEG.)  Additional electrodes (e.g. T1/T2, sphenoidal or subtemporal electrodes), special electrodes (e.g. eye movement monitors) and modifications of the 10-20 and 10-10 System (e.g. ‘prime’ electrodes used when there are skull defects or alteration) should be included.  Special parameters used during post-hoc review of the rsEEG should be included in this section.  An example would be “This record was reviewed under the following settings:  bandpass filters of 1-70 Hz, sensitivity of 7 uV/mm, a display speed of 30 mm/sec, with a 60 Hz notched filter applied as appropriate.  Modifications in these parameters were made as necessary for further waveform resolution.  Supplemental ECG leads for cardiac rhythm monitoring were utilized during the recording.”

The conditions of the recording should be elucidated.  A statement regarding the use of pre-medication should include the drug and dose (e.g. “lorazepam 1 mg was administered prior to the recording”).  Other conditions that can influence the EEG should also be documented, including sleep deprivation, potential dietary influences (e.g. fasting or NPO status) and modality used (rsEEG, ambulatory EEG etc.).  Reporting the total recording time is also advisable, particularly when it is shorter or longer than recommended in the ACNS Guideline 1, Sec. 3.7, for compliance purposes.

The patient’s state of consciousness at the onset of the record should be clear at the beginning of the recording, including if the patient is awake, drowsy, asleep, or in a
compromised level of consciousness such as coma or coma-like states. This information on the patient's state, i.e. the level of consciousness, helps guide the electroclinical correlation.

**EEG Description.** This section should include a description of the background electrocerebral activity, including all the essential characteristics of waveforms in the record, detailed as objectively as possible.

EEG signals are complex, and extraction of clinically relevant features by visual analysis alone is subject to individual variability. While automated software application enhances our ability to detect and quantify the power of specific bandwidths of EEG, human extraction of the clinically relevant features requires identification, integration of various bandwidths, and interpretation of the significant features in the context of the overall recording.

Description of the record should provide an objective means of analysis for review at a different time or by another interpreter. It should utilize technical terminology and metrics to detail the waveforms present for the duration of the recording. The aim is to provide a complete, objective, and orderly description of the state of the patient, the background activity, and the most salient features of the EEG to allow a conclusion of 'normal' or 'abnormal.' It should also identify and describe normal variants in addition to abnormal findings. When an abnormality is identified, the degree of abnormality should be stated.

The rsEEG description should begin with a complete description of the background activity, including the posterior dominant rhythm additional features of the background, and special features. The description of the best posterior dominant rhythm in units of frequency (Hertz (Hz), or cycles per second) and amplitude (microvolts/mm) should be reported with the patient in the most alert state. Records obscured by artifact, recordings in infants, and some normal records may not have a clearly defined posterior dominant frequency, and the report should reflect this. Response or reactivity to external stimuli should also be noted. Subsequent non-dominant background activity should be identified by principal frequencies, amount of each present, degrees of symmetry, location/distribution, morphology, amplitude, and rhythmicity, using the same units as for the posterior dominant frequency. Description of non-dominant background activity should include beta, theta and delta activity. Terms such as "low," "medium" and "high" voltage may be used but should be quantified with numerical measures.

State changes in the patient should be documented. The level of alertness, the organization of the EEG background frequency and amplitude over time, and the spatial features of the recording should be noted. Sleep patterns and architecture should be reported to reflect all sleep stages attained during the EEG recording. Abnormal patterns such as rapid sleep cycling, sleep-onset rapid eye movements, and asymmetry or attenuation in the normal sleep elements (e.g. spindles) should be noted.

Hyperventilation and intermittent photic stimulation are routine activating procedures, used to trigger abnormalities during rsEEG. They should be performed and their effects noted. When omitted, the reason for their omission should be stated. If augmentation of slowing or any epileptiform abnormalities are encountered in the rsEEG during or after activating procedures, these responses should be detailed. Documentation
of poor effort with hyperventilation is relevant to rsEEG interpretation. If additional methods are used to enhance EEG abnormalities, they should also be documented.

Any special characteristics present in the background, such as voltage attenuation or augmentation, suppression-burst activity, or electrocerebral inactivity should be detailed using the same terminology used to describe the background. Descriptions should note morphology (monomorphic, polymorphic, or irregular), rhythmicity, voltage, continuous vs intermittent features, laterality (e.g. left or right; bilateral or diffuse), region of involvement (e.g. frontal, temporal, central, parietal, occipital) and frequency (e.g. theta, delta slowing). For epileptiform and non-epileptiform features with bilateral localization, amplitude symmetry (e.g. > 50%) and synchrony (e.g. secondary bilateral synchrony vs bilateral synchronous) should be included in the description, as well as the temporal pattern of their occurrence (e.g. bursts, prolonged runs, or sporadic).

Salient abnormal features should be noted following identification of the state and background activity. When interictal epileptiform discharges are present, one should document the location, morphology (e.g. spike, sharp, polyspike +/- slow wave), pattern (e.g. single, run, random, rhythmic, periodic), and incidence (e.g. rare, intermittent, occasional, frequent, continuous). Further description of the frequency in Hz should be included as well as the pattern of occurrence (e.g. single, couplet, bursts, a train), as well as their duration. In addition, some abnormal findings may be influenced by external stimulation (e.g. stimulus-induced, rhythmic periodic ictal-like discharges). Quantifying paroxysmal abnormalities is often expressed in a subjective manner and is relative to individual reporting designs (Hirsch LJ et al., 2013; Beniczky et al., 2013).

Snapshots of a reported abnormality are encouraged to be included in the EEG report. This will help facilitate an understanding of what is being identified as an abnormality (Figure). By providing a patient sample of the EEG abnormality, more universal validation will be enabled (beyond the text document).

The presence of an electroclinical seizure or electrographic (i.e. without clinical manifestations) seizure should include description of the electrographic onset, field of propagation, and postictal period, defining the temporal and spatial characteristics and using terminology to define the location, distribution, morphology, amplitude, and rhythmicity, in addition to the duration and frequency of recurrence. When present, any clinical changes and the qualitative nature of the change should be documented, as described below under video recording.

Artifacts are present in virtually every rsEEG. They should be reported when they mimic cerebral activity, when they are unusual or excessive (e.g. eye movements or muscle activity), when they interfere with interpretation of the record (> 50% of the tracing involving >50% of the electrodes), or when they provide valuable diagnostic information (e.g., myokymia, nystagmus, etc.).

A single channel electrocardiogram (ECG) should be included in every EEG. Reporting ECG findings in the EEG description will vary and depend upon the interpreter’s level of expertise. Other channels such as eye movement monitors (used by some laboratories routinely), channels monitoring respiration, movement, EMG and non-cephalic monitors should be reported and described when applicable and when their significance is questioned. [Guideline 1: Minimum Technical Requirements for Performing Clinical EEG].

Video recording is a routine part of most proprietary systems for performing rsEEG. Including video descriptions in the report is important for providing additional information
involving electroclinical episodes and seizures, and in assessing artifacts [as well as having an impact on CPT coding for billing purposes (https://www.aan.com/go/practice/coding)]. The main features should include a description of the clinical event and, when possible, terminology used by the International League Against Epilepsy seizure classification system (Berg A et al., 2010) reporting the duration, level of responsiveness/consciousness, and any intervention provided.

**Impression.** The impression (or interpretation) is a synthesis of the significance of the EEG findings. It is written primarily for the referring clinician and should, therefore, be as succinct as possible, and readily understandable to a clinician of any level of expertise or specialty. It should include an initial, clear summary statement as to whether it is a normal or an abnormal record. When the EEG is abnormal, the reasons why should be listed in a clear and concise line item format, in part to simplify the results for comparison among successive records. It is desirable to list the abnormalities by degrees of importance [examples: (1) a left focal temporal electrographic seizure; (2) left anterior temporal spike-and-waves; (3) left temporal delta slowing; (4) mild slowing of the background activity.]

The summary of the findings should be succinctly stated in lay terminology. When reporting several types of abnormalities, the list should be limited to the most salient findings and to the minimum number necessary to convey the significance of those findings (preferably no more than 3 or 4). Often, the impression will be the only part of the report of major interest to the clinician, so the importance of this section is stressed. The impression should avoid confusing terminology and technical jargon, but “epileptiform discharges,” including “spikes and sharp waves” are universally accepted terms.

**Clinical Correlation.** The clinical correlation is the ultimate translation of the EEG. It should integrate the reason for referral for the EEG and the findings, to be used jointly to assist with patient management. The clinical correlation should clearly express the relevance of the findings to the clinician. Avoiding technical terminology is helpful to convey the message to the least experienced clinician on the team caring for the patient. A good litmus test is that this section should be understandable to a general practitioner or nurse [Kaplan & Benbadis 2013].

Phrases such as “no focal or lateralizing abnormality”, “no epileptiform abnormality” and “No electrographic seizures or evidence of status epilepticus were present” are helpful in the clinical correlation when the clinical request is explicit. Some common scenarios will be consistent from patient to patient. While individual reporting styles vary and wording differs slightly, the following are examples of clinical correlations that may be used to express such concepts:

1. “A normal interictal EEG does not exclude nor support the diagnosis of epilepsy.”
2. “Focal slowing suggests an underlying lesion involving the white matter of the ipsilateral hemisphere.”
3. “Diffuse slowing of the background activity reflects a [include degree: mild, moderate, severe] diffuse cortical dysfunction, which can be seen with toxic-metabolic or systemic causes or neurodegenerative disorders, and also with cortical atrophy.”

4. “The generalized spike-and-waves seen in this tracing imply a generalized mechanism in a patient with a clinical diagnosis of epilepsy but may also exist as an inherited trait independent of clinical seizures.”


6. “The suppression-burst pattern following normothermic cardiac arrest (in the absence of anesthetic drugs) suggests a poor prognosis for neurologic outcome.”

While the clinical correlations in these cases may help standardize reporting, specific therapeutic suggestions such as “this pattern warrants antiseizure drugs” or “clinical correlation is strongly advised” should be avoided, recognizing the diagnostic limits of a rsEEG. Suggestions for further testing may be made within this section, e.g. suggesting a repeat EEG with sleep-deprivation, ambulatory EEG, video-EEG monitoring, referral to a sleep laboratory when sleep apnea is suspected or further cardiologic evaluation when the ECG is abnormal. When previous EEGs are available, comparison of the current record to previous tracings should be included.

This standardized reporting format is intended to maximize clear communication between different reviewers of the same patient’s rsEEG. It is hoped that this will facilitate inter-observer reliability of EEG reporting for clinical care of patients and increase consistency for research studies.

References

1. Tatum WO. How not to read an EEG: Introductory statements. Neurology 2013(suppl 1);80(1):S1-S3.


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Figure: A single right temporal spike-and-wave with a regional temporal field during drowsiness

This statement is provided as an educational service of the American Clinical Neurophysiology Society (ACNS). It is based on an assessment of current scientific and clinical information. It is not intended to include all possible proper methods of care for a particular problem or all legitimate criteria for choosing to use a specific procedure. Neither is it intended to exclude any reasonable alternative methodologies. ACNS recognizes that specific patient care decisions are the prerogative of the patient and the physician caring for the patient, based on all of the circumstances involved. The clinical context section is made available in order to place the evidence-based guidelines into perspective with current practice habits and challenges. Formal practice recommendations are not intended to replace clinical judgment.