Operational Classification of Seizure Types by the International League Against Epilepsy

Robert S. Fisher¹, J. Helen Cross², Jacqueline A. French³, Norimichi Higurashi⁴, Edouard Hirsch⁵, Floor E. Jansen⁶, Lieven Lagae⁷, Solomon L. Moshe⁸, Jukka Peltola⁹, Eliane Roulet Perez¹⁰, Ingrid E. Scheffer¹¹, Sameer M. Zuberi¹²

robert.fisher@stanford.edu; h.cross@ucl.ac.uk; Jacqueline.french@nyumc.org; nori@jikei.ac.jp; edouard.hirsch@chu-strasbourg.fr; F.E.Jansen@umcutrecht.nl; lieven.lagae@uzleuven.be; solomon.moshe@einstein.yu.edu; jukka.peltola@pshp.fi; eliane.roulet-perez@chuv.hospvd.ch; scheffer@unimelb.edu.au; sameer.zuberi@nhs.net;

¹ Stanford Department of Neurology & Neurological Sciences, Stanford, CA, USA
² UCL-Institute of Child Health, & Great Ormond Street Hospital for Children, London, UK
³ Department of Neurology, NYU Langone School of Medicine, NY USA
⁴ Department of Pediatrics, Jikei University School of Medicine, Tokyo, Japan
⁵ Unite Francis Rohmer, Strasbourg France
⁶ Department of Pediatric Neurology, Brain Center Rudolf Magnus, University Medical Center, Utrecht, The Netherlands
⁷ Pediatric Neurology, University Hospitals KU Leuven, Leuven, Belgium
⁸ Saul R. Korey Department of Neurology, Department of Pediatrics and Dominick P. Purpura Department Neuroscience, Montefiore Medical Center, Bronx, NY USA
⁹ Department of Neurology, Tampere University Hospital, Tampere, Finland
¹⁰ Unité de Neurologie et Neuroréhabilitation Pédiatrique CHUV-1011 Lausanne Switzerland
¹¹ Florey Institute and University of Melbourne, Austin Health and Royal Children’s Hospital, Melbourne, Australia
¹² The Paediatric Neurosciences Research Group, Royal Hospital for Children, Glasgow, UK & College of Medicine, Veterinary & Life Sciences, University of Glasgow, United Kingdom

Corresponding author:
Robert S. Fisher, MD, PhD
Neurology, SNHC, Room 4865
213 Quarry Road
Palo Alto, CA 94304
robert.fisher@stanford.edu
Phone: (650) 721-5289
Fax: (650) 721-4865
Summary

Objectives: The International League Against Epilepsy (ILAE) presents a revised operational classification of seizure types. The purpose of such a revision is to recognize that some seizure types can have either a focal or generalized onset, to allow classification when the onset is unobserved, to include some missing seizure types and to adopt more transparent names.

Methods: Because current knowledge is insufficient to form a scientifically-based classification, the 2017 classification is operational (practical) and based upon the 1981 Classification, extended in 2010.

Results: Changes include:
1. “partial” becomes “focal”; 2. Awareness is used as a classifier of focal seizures; 3. The terms dyscognitive, simple partial, complex partial, psychic, secondarily generalized are eliminated; 4. New focal seizure types include automatisms, behavior arrest, hyperkinetic, autonomic, cognitive and emotional. 5. Atonic, clonic, epileptic spasms, myoclonic, tonic seizures can be either of focal or generalized onset. 6. Focal to bilateral tonic-clonic seizure replaces secondarily generalized seizure. 7. New generalized seizure types are: absence with eyelid myoclonia, myoclonic absence, myoclonic-atonic, myoclonic-tonic-clonic. 8. Seizures of unknown onset may have features that can still be classified.

Significance: The new classification does not represent a fundamental change, but allows greater flexibility and transparency in naming seizure types.
Introduction

The International League Against Epilepsy (ILAE), through the Commission for Classification and Terminology, has developed a working classification of seizures and epilepsy. Following the proposed reorganization in 2010\textsuperscript{1,2}, further clarification has been discussed and feedback sought from the community. One area that required further elucidation was the organization of seizure types. A Seizure Type Classification Task was established in 2015 to prepare recommendations for classification of seizure types, which are summarized in this document. A companion document guides the intended use of the classification.

Descriptions of seizure types date back at least to the time of Hippocrates. Gastaut\textsuperscript{3,4} proposed a modern classification in 1964. Various basic frameworks for seizure classification can be considered. Manifestations of certain seizures are age-specific and depend on maturation of the brain. Previous classifications have been based on anatomy, with temporal, frontal, parietal, occipital, diencephalic or brainstem seizures. Modern research changed our view of the pathophysiological mechanisms involved and has shown epilepsy to be a network disease and not only a symptom of local brain abnormalities.\textsuperscript{5} From a network perspective, seizures could arise in neocortical, thalamo-cortical, limbic, and brainstem networks. Although our understanding of seizure networks is evolving rapidly\textsuperscript{6} it is not yet sufficient to serve as a basis for seizure classification. In 1981, an ILAE Commission led by Dreifuss and Penry\textsuperscript{7} evaluated hundreds of video-EEG recordings of seizures to develop recommendations that divided seizures into those of partial and generalized onset, simple and complex partial seizures and various specific generalized seizure types. This classification remains in widespread use today, with revisions in terminology and classification of seizures and epilepsy by the ILAE\textsuperscript{2,8-13,14} and with suggested insights, modifications and criticisms by others.\textsuperscript{15-24} We chose not to develop a classification based solely on observed behavior - instead, reflecting clinical practice, the 2017 classification is interpretive, allowing the use of additional data to classify seizure types.

The intention of the 2001\textsuperscript{12} and 2006\textsuperscript{13} reports on reclassification was to identify unique diagnostic entities with etiologic, therapeutic, and prognostic implications, so that when a syndromic diagnosis could not be made, then the therapy and prognosis would be based on seizure type. Such a classification would permit grouping of reasonably pure cohorts of patients for discovery of etiologies, including genetic factors, research into fundamental mechanisms, involved networks and clinical trials. The ILAE Seizure Type Classification Task Force (hereafter called “the Task Force”) chose to use the phrase “operational classification,” because it is impossible at this time to base a classification fully on the science of epilepsy. In the absence of a full scientific classification, the Task Force chose to use the basic organization initiated in 1981 and subsequently modified\textsuperscript{1,2} as a starting point for the revised operational classification.
Methods

What is a seizure type?

A seizure is defined as “a transient occurrence of signs and/or symptoms due to abnormal excessive or synchronous neuronal activity in the brain” \(^{25}\) It is the clinician’s first task to determine that an event has the characteristics of a seizure and not one of the many imitators of seizures. \(^{26}\) The next step is classification into a seizure type.

The Task Force operationally defines a seizure type as a useful grouping of seizure characteristics for purposes of communication in clinical care, teaching and research. Mention of a seizure type should bring to mind a specific entity, albeit sometimes with subcategories and variations on a theme. Choices must be made by interested stakeholders to highlight groupings of seizure characteristics that are useful for specific purposes. Such stakeholders include patients, families, medical professionals, researchers, epidemiologists, medical educators, clinical trialists, insurance payers, regulatory agencies, advocacy groups, and medical reporters. Operational (practical) groupings can be derived by those with specific interests. A pharmacologist, for example, might choose to group seizures by efficacy of medications. A researcher doing a clinical trial might consider seizures as disabling or non-disabling. A surgeon might group by anatomy in order to predict the eligibility for and likely success of surgical therapy. A physician based in an ICU with predominantly unconscious patients might group seizures in part by EEG pattern \(^{27}\). The principal aim of this classification is to provide a communication framework for clinical use. Seizure types are relevant to clinical practice in humans; whereas, it is acknowledged that seizure types in other species, experimental and natural, may not be reflected in the proposed classification. One goal was to make the classification understandable by patients and families and broadly applicable to all ages, including neonates. The Commission on Classification & Terminology recognizes that seizures and epilepsies in the neonate can have motor manifestations or alternatively little or no behavioral manifestations. A separate Neonatal Seizure Task Force is working to develop a classification of neonatal seizures. The 2017 seizure classification is not a classification of electroencephalographic ictal or subclinical patterns. The guiding principle of the Seizure Type Task Force was a quotation from Albert Einstein: “Make things as simple as possible, but no simpler.”

Motivation for change

Adapting to a change of terminology can be effortful and needs to be motivated by a rationale for change. Seizure type classification is important for several reasons. First, the classification becomes a worldwide shorthand communication among clinicians caring for people with epilepsy. Second, the classification allows grouping of patients for therapies. Some regulatory agencies approve drugs or devices indicated for specific seizure types. A new classification should gracefully map to existing indications for drug or device usage. Third, the
seizure type grouping might provide a useful link to specific syndromes or etiologies, for example, by noting an association between gelastic seizures and hypothalamic hamartoma or epileptic spasms with tuberous sclerosis. Fourth, the classification allows researchers to better focus their studies on mechanisms of different seizure types. Fifth, a classification provides words to patients to describe their disease. Motivations for revising the 1981 Seizure Classification are listed below.

1. Some seizure types, for example, tonic seizures or epileptic spasms, can have either a focal or generalized onset.

2. Lack of knowledge about the onset makes a seizure unclassifiable and difficult to discuss with the 1981 system.

3. Retrospective seizure descriptions often do not specify a level of consciousness, and altered consciousness, while central to many seizures, is a complicated concept.

4. Some terms in current use do not have high levels of community acceptance or public understanding, such as “psychic,” “partial,” “simple partial,” “complex partial” and “dyscognitive.”

6. Some important seizure types are not included.

Results

Classification of Seizure Types

Figure 1 depicts the basic and Figure 2 the expanded 2017 seizure classification. The two represent the same classification, with collapse of the subcategories to form the basic version. Use of one versus the other depends upon the desired degree of detail. Variations on the individual seizure theme can be added for focal seizure types according to level of awareness.
Figure 1: The Basic ILAE 2017 Operational Classification of Seizure Types.

ILAE 2017 Classification of Seizure Types Basic Version

- **Focal Onset**
  - Aware
  - Impaired Awareness
  - Motor Onset
  - Non-Motor Onset
  - focal to bilateral tonic-clonic

- **Generalized Onset**
  - Motor
    - Tonic-clonic
    - Other motor
    - Non-Motor (Absence)

- **Unknown Onset**
  - Motor
    - Tonic clonic
    - Other motor
    - Non-Motor

**Unclassified**

---

1. Definitions, other seizure types and descriptors are listed in the accompanying paper & glossary of terms
2. Due to inadequate information or inability to place in other categories
Figure 2: The Expanded ILAE 2017 Operational Classification of Seizure Types. The following clarifications should guide the choice of seizure type. For focal seizures, specification of level of awareness is optional. Retained awareness means the person is aware of self and environment during the seizure, even if immobile. A focal aware seizure corresponds to the prior term simple partial seizure. A focal impaired awareness seizure corresponds to the prior term complex partial seizure, and impaired awareness during any part of the seizure renders it a focal impaired awareness seizure. Focal aware or impaired awareness seizures optionally may further be characterized by one of the motor-onset or non-motor onset symptoms below, reflecting the first prominent sign or symptom in the seizure. Seizures should be classified by the earliest prominent feature, except that a focal behavior arrest seizure is one for which cessation of activity is the dominant feature throughout the seizure. A focal seizure name also can omit mention of awareness when awareness is not applicable or unknown and thereby classify the seizure directly by motor onset or non-motor onset characteristics. Atonic seizures and epileptic spasms would usually not have specified awareness. Cognitive seizures imply impaired language or other cognitive domains or positive features such as déjà vu, hallucinations, illusions or perceptual distortions. Emotional seizures involve anxiety, fear, joy, other emotions or appearance of affect without subjective emotions. An absence is atypical because of slow onset or termination or significant changes in tone supported by atypical, slow, generalized spike and wave on the EEG. A seizure may be unclassified due to inadequate information or inability to place the type in other categories.
Structure of the classification

The classification chart is columnar, but not hierarchical, so arrows are intentionally omitted. Seizure classification begins with the determination of whether the initial manifestations of the seizure are focal or generalized. The onset may be missed or obscured, in which case the seizure is of unknown onset. The words “focal” and “generalized” at the start of a seizure name are assumed to mean of focal or generalized onset.

For focal seizures, the level of awareness optionally may be included in the seizure type. Retained awareness means that the person is aware of self and environment during the seizure, even if immobile. A focal aware seizure (with or without any subsequent classifiers) corresponds to the prior term “simple partial seizure.” A focal impaired awareness seizure (with or without any subsequent classifiers) corresponds to the prior term “complex partial seizure.” Impaired awareness during any part of the seizure renders it a focal impaired awareness seizure.

Focal seizures also are sub-grouped as those with motor and non-motor signs and symptoms. If both motor and non-motor signs are present at the seizure start, the motor signs will usually dominate, unless non-motor (e.g., sensory) symptoms and signs are very prominent.

Focal aware or impaired awareness seizures optionally may further be characterized by one of the listed motor onset or non-motor onset symptoms, reflecting the first prominent sign or symptom in the seizure, for example, focal impaired awareness automatism seizure. Seizures should be classified by the earliest prominent motor onset or non-motor onset feature, except that a focal behavior arrest seizure is one for which cessation of activity is the dominant feature throughout the seizure, and any significant impairment of awareness during the course of the seizure causes a focal seizure to be classified as having impaired awareness. A focal seizure name can omit mention of awareness when awareness is not applicable or unknown; thereby classifying the seizure directly by motor onset or non-motor onset characteristics. The terms motor onset and non-motor onset may be omitted when a subsequent term generates an unambiguous seizure name.

The classification of an individual seizure can stop at any level: a “focal onset” or “generalized onset” seizure, with no other elaboration, or a “focal sensory seizure,” “focal motor seizure,” “focal tonic seizure” or “focal automatism seizure,” etc. Additional classifiers are encouraged, and their use may depend upon the experience and purposes of the person classifying the seizure. The terms focal onset and generalized onset are for purposes of grouping. No inference is made that each seizure type exists in both groups; including absence seizures in the generalized onset category does not imply existence of “focal absence” seizures.

When the primacy of one versus another key symptom or sign is unclear, the seizure can be classified at a level above the questionably applicable term with additional descriptors of seizure semiology relevant to the individual seizure. Any signs or symptoms of seizures,
suggested descriptor terms in the companion paper or free text descriptions can optionally be appended to the seizure type as descriptions, but they do not alter the seizure type.

The seizure type “focal to bilateral tonic-clonic” is a special seizure type, corresponding to the 1981 phrase “partial onset with secondary generalization.” Focal onset bilateral tonic-clonic reflects a propagation pattern of a seizure, rather than a unitary seizure type, but it is such a common and important presentation that the separate categorization was continued. The term “to bilateral” rather than “secondary generalized” was used to further distinguish this focal onset seizure from a generalized onset seizure. The term “bilateral” is used for propagation patterns and “generalized” for seizures that engage bilateral networks from onset.

Seizure activity propagates through brain networks, sometimes leading to uncertainty about whether an event is a unitary seizure or a series of multiple seizures starting from different networks (“multifocal”). A single unifocal seizure can present with multiple clinical manifestations as a result of propagation. The clinician will have to determine (by observation of a continuous evolution or stereotypy form seizure-to-seizure) whether an event is a single seizure or a series of different seizures. When a single focal seizure presents with a sequence of signs and symptoms, then the seizure is named for the initial prominent sign or symptom, reflecting the usual clinical practice of identifying the seizure onset focus or network. For example, a seizure beginning with sudden inability to understand language followed by impaired awareness and clonic left arm jerks would be classified as a “focal impaired awareness (non-motor onset) cognitive seizure” (progressing to clonic left arm jerks). The terms in parentheses are optional. The formal seizure type in this example is determined by the cognitive non-motor onset and presence of altered awareness during any point of the seizure.

Generalized seizures are divided into motor and non-motor (absence) seizures. Further subdivisions are similar to those of the 1981 classification, with addition of myoclonic-atonic seizures, common in epilepsy with myoclonic-atonic seizures (Doose syndrome), myoclonic-tonic-clonic seizures common in juvenile myoclonic epilepsy, myoclonic absence and absence seizures with eyelid myoclonia seen in the syndrome described by Jeavons and elsewhere. Generalized manifestations of seizures can be asymmetrical, rendering difficult the distinction from focal onset seizures. The word “absence” has a common meaning, but an “absent stare” is not synonymous with an absence seizure, since arrest of activity also occurs in other seizure types.

The 2017 classification allows appending a limited number of qualifiers to seizures of unknown onset, in order to better characterize the seizure. Seizures of unknown onset may be referred to by the single word “unclassified” or with additional features, including motor, non-motor, tonic-clonic, epileptic spasms and behavior arrest. A seizure type of unknown onset may later become classified as either of focal or generalized onset, but any associated behaviors (e.g., tonic-clonic) of the previously unclassified seizure will still apply. In this regard, the term “unknown onset” is a placeholder - not a characteristic of the seizure, but of ignorance.
Reasons for decisions

The terminology for seizure types is designed to be useful for communicating the key characteristics of seizures and to serve as one of the key components of a larger classification for the epilepsies, which is being developed by a separate ILAE Classification Task Force. The basic framework of seizure classification used since 1981 was maintained.

Focal vs. partial

In 1981, the Commission declined to designate as “focal” a seizure that might involve an entire hemisphere, so the term “partial” was preferred. The 1981 terminology was in a way prescient of the modern emphasis on networks, but “partial” conveys a sense of part of a seizure, rather than a location or anatomical system. The term “focal” is more understandable in terms of seizure onset location.

Focal vs. generalized

In 2010 the ILAE defined focal as “originating within networks limited to one hemisphere. They may be discretely localized or more widely distributed. Focal seizures may originate in subcortical structures.” Generalized from onset seizures were defined as “originating at some point within, and rapidly engaging, bilaterally distributed networks.” Classifying a seizure as having apparently generalized onset does not rule out a focal onset obscured by limitations of our current clinical methods, but this is more an issue of correct diagnosis than of classification. Furthermore, focal seizures may rapidly engage bilateral networks, while classification is based on unilateral onset. For some seizure types, for example epileptic spasms, the distinction of a focal versus generalized onset may require careful study of a video-EEG recording or the type of onset may be unknown. A distinction between focal and generalized onset is a practical one, and may change with advances in ability to characterize the onset of seizures.

Focality of seizure onset can be inferred by pattern matching to known focal onset seizures, even when the focality is not clear strictly in terms of observable behavior. A seizure is focal, for example, when it starts with déjà vu and then progresses to loss of awareness and responsiveness, lip smacking and hand rubbing for a minute. There is nothing intrinsically “focal” in the description, but video-EEGs of countless similar seizures have previously shown focal onsets. If the epilepsy type is known, then the onset can be presumed even if it is unwitnessed, for example, an absence seizure in a person with known juvenile absence epilepsy.

Clinicians have long been aware that so-called generalized seizures, for example, absence seizures with EEG generalized spike-waves, do not manifest equally in all parts of the brain. The Task Force emphasized the concept of bilateral, rather than generalized involvement of some
seizures, since seizures can be bilateral without involving every brain network. The bilateral manifestations need not be symmetric. The term “focal to bilateral tonic-clonic” was substituted for “secondarily generalized.” The term “generalized” was maintained for seizures generalized from onset.

**Unknown Onset**

Clinicians commonly hear about tonic-clonic seizures for which the onset was unobserved. Perhaps, the patient was asleep, alone or observers were too distracted by the manifestations of the seizure to notice the presence of focal features. There should be an opportunity to provisionally classify this seizure, even in the absence of knowledge about its origin. The Task Force therefore allowed further description of seizures of unknown onset when key characteristics, such as tonic-clonic activity or behavior arrest are observed during the course of the seizure. The Task Force recommends classifying a seizure as having focal or generalized onset only when there is a high degree of confidence (e.g., ≥ 80%, arbitrarily chosen to parallel the usual allowable beta error) in the accuracy of the determination; otherwise, the seizure should remain unclassified until more information is available.

It may be impossible to classify a seizure at all, either because of incomplete information or because of the unusual nature of the seizure, in which case it is called an unclassified seizure. Categorization as unclassified should only be used for the exceptional situation in which the clinician is confident that the event is a seizure but cannot further classify the event.

**Consciousness and awareness**

The 1981 classification and the revision in 2010 \(^1\)\(^{10}\);\(^{32}\) suggested a fundamental distinction between seizures with loss or impairment of consciousness and those with no impairment of consciousness. Basing a classification upon consciousness (or one of its allied functions) reflects a practical choice that seizures with impaired consciousness should often be approached differently from those with unimpaired consciousness, for example, with respect to allowing driving in adults or interfering with learning in children. The ILAE chose to retain impairment of consciousness as a key concept in the grouping of focal seizures. However, consciousness is a complex phenomenon, with both subjective and objective components.\(^{33}\) Multiple different types of consciousness have been described for seizures.\(^{34}\) Surrogate markers for consciousness usually comprise measurements of awareness, responsiveness, and memory. The 1981 classification specifically mentioned awareness and responsiveness, but not memory for the event.

Retrospective determination of state of consciousness can be difficult. An untrained classifier might assume that a person must be on the ground, immobile, unaware and unresponsive (e.g., “passed out”) for a seizure to show impaired consciousness. The Task Force
adopted state of awareness as a relatively simple surrogate marker for consciousness. “Retained awareness” is considered to be an abbreviation for “seizures with no impairment of consciousness during the event.” We employ an operational definition of awareness as knowledge of self and environment. In this context, awareness refers to perception or knowledge of events occurring during a seizure, not to knowledge of whether a seizure occurred. In several languages, “unaware” translates as “unconscious,” in which case changing the seizure designation from “complex partial” to “impaired awareness” will emphasize the importance of consciousness by putting its surrogate directly in the seizure title. In English, “focal aware seizure” is shorter than is “focal seizure without impairment of consciousness” and possibly better understood by patients. As a practical issue, retained awareness usually includes the presumption that the person having the seizure later can recall and validate having retained awareness; otherwise, impaired awareness may be assumed. Exceptional seizures present with isolated transient epileptic amnesia in clear awareness but classification of an amnestic seizure as focal aware would require clear documentation by meticulous observers. Awareness may be left unspecified when the extent of awareness cannot be ascertained.

Responsiveness may or may not be compromised during a focal seizure. Responsiveness does not equate to awareness or consciousness, since some people are immobilized and consequently unresponsive during a seizure, but still able to observe and recall their environment. Additionally, responsiveness often is not tested during seizures. For these reasons, responsiveness was not chosen as a primary feature for seizure classification, although responsiveness can be very helpful in classifying the seizure when it can be tested. The term “dyscognitive” was not carried into the current classification as a synonym for “complex partial” because of lack of clarity and negative public and professional feedback.

Awareness is not a classifier for generalized onset seizures, because the large majority of generalized seizures present with impaired awareness or full loss of consciousness. However, it is recognized that awareness and responsiveness can be at least partially retained during some generalized seizures, for example, with brief absence seizures, including absence seizures with eyelid myoclonias or myoclonic seizures.

**Etiologies**

A classification of seizure types can be applied to seizures of different etiologies. A post-traumatic seizure or a reflex seizure may be focal with or without impairment of awareness. Knowledge of the etiology, for instance, presence of a focal cortical dysplasia, can aid in classification of the seizure type. Any seizure can become prolonged, leading to status epilepticus of that seizure type.

**Supportive information**
As part of the diagnostic process, a clinician will commonly use supportive evidence to help to classify a seizure, even though that evidence is not part of the classification. Such evidence may include videos brought in by family, EEG patterns, lesions detected by neuroimaging, laboratory results such as detection of anti-neuronal antibodies, gene mutations, or an epilepsy syndrome diagnosis known to be associated with either focal or generalized seizures or both, such as Dravet syndrome. The seizures usually can be classified on the basis of symptoms and behavior, provided that good subjective and objective descriptions are available. Use of any available supportive information to classify the seizure is encouraged. Availability of supportive information may not exist in the resource-poor parts of the world, which may lead to a less specific, but still correct classification.

ICD 9, 10, 11, 12

The World Health Organization International Classification of Diseases (ICD) is used for inpatient and outpatient diagnoses, billing and many other purposes. Concordance between ICD epilepsy diagnoses and ILAE seizure types is desirable for clarity and consistency. This is possible only to a limited extent with existing ICD terms, since ICD 9, 10, 11 are already formulated. The ILAE proposals will always lead ICD standards. ICD 9 and 10 make use of old seizure terminology, including terms such as petit mal and grand mal. ICD11 does not name seizure types at all, but focuses on epilepsy etiologies and syndromes, as do ILAE epilepsy classifications. For this reason, there is no conflict between our proposed seizure type classification and ICD11. Efforts can be made to incorporate new classifications of seizure types and syndromes into the development of ICD12.

Discussion

Discontinued terms

Simple/complex partial: After approximately 35 years of use, the terms “simple partial seizure” and “complex partial seizure” may be missed by some clinicians. The reasons for changing are three. First, a decision was previously made to globally change partial to focal. Second, “complex partial” has no intrinsic meaning to the public. The phrase “focal impaired awareness” can convey meaning to a lay person with no knowledge of seizure classification. Third, the words “complex” and “simple” can be misleading in some contexts. Complex seems to imply that this seizure type is more complicated or difficult to understand than other seizure types. Calling a seizure “simple” may trivialize its impact to a patient who does not find the manifestations and consequences of the seizures to be at all simple.

Convulsion: The term “convulsion” is a popular, ambiguous and unofficial term used to mean substantial motor activity during a seizure. Such activity might be tonic, clonic, myoclonic
or tonic-clonic. In some languages, convulsions and seizures are considered synonyms and the motor component is not clear. The word “convulsion” is not part of the 2017 seizure classification, but will undoubtedly persist in popular usage.

**Added terms**

*Aware/impaired awareness:* As discussed above, these terms designate knowledge of self and environment during a seizure.

*Hyperkinetic:* Hyperkinetic seizures have been added to the focal seizure category. Hyperkinetic activity comprises agitated thrashing or leg pedaling movements. Hypermotor is an earlier term introduced as part of a different proposed classification by Lüders and colleagues in 1993. The term hypermotor, which contains both Greek and Roman roots, was supplanted in the 2001 ILAE glossary and 2006 report by “hyperkinetic,” and to be both etymologically and historically consistent, “hyperkinetic” was chosen for the 2017 classification.

*Cognitive:* This term replaces “psychic” and refers to specific cognitive impairments during the seizure, for example, aphasia, apraxia or neglect. The word “impairment” is implied since seizures never enhance cognition. A cognitive seizure can also comprise positive cognitive phenomena, such as déjà vu, jamais vu, illusions or hallucinations.

*Emotional:* A focal non-motor seizure can have emotional manifestations, such as fear or joy. The term also encompasses affective manifestations with the appearance of emotions occurring without subjective emotionality, such as may occur with some gelastic or dacrystic seizures.

**New focal seizure types:** Some seizure types that previously were described only as generalized seizures now appear under seizures of focal, generalized and unknown onset. These include epileptic spasms, tonic, clonic, atonic, myoclonic seizures. The list of motor behaviors constituting seizure types comprises the most common focal motor seizures, but other less common types, for example, focal tonic-clonic, may be encountered. Focal automatisms, autonomic, behavior arrest, cognitive, emotional and hyperkinetic are new seizure types. Focal to bilateral tonic-clonic seizure is a new type as the renamed secondarily generalized seizure.

**New generalized seizure types:** Relative to the 1981 classification, new generalized seizure types include: absence with eyelid myoclonia, myoclonic-atonic, and myoclonic-tonic-clonic (although clonic onset of tonic-clonic seizures was mentioned in the 1981 publication). Seizures with eyelid myoclonia and myoclonic absence seizures potentially have features of both absence and motor seizures, and could have been placed in either group. Seizures with eyelid myoclonia may even rarely display focal features. Epileptic spasms are seizures represented in focal, generalized and unknown onset categories, and the distinction may require video-EEG
recording. The term “epileptic” is implied for every seizure type, but explicitly stated for epileptic spasms, because of the ambiguity of the single word “spasms” in neurological use.

**What is different from the 1981 classification?**

Table 1 summarizes the changes in the ILAE 2017 seizure type classification from the 1981 classification. Note that several of these changes were already incorporated into the 2010 revision of terminology and subsequent revisions.\(^1\);\(^3\)\(^2\).

Table 4: Changes in Seizure Type Classification from 1981 to 2017

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Change of “partial” to “focal.”</td>
</tr>
<tr>
<td>2.</td>
<td>Certain seizure types can be either of focal, generalized or unknown onset.</td>
</tr>
<tr>
<td>3.</td>
<td>Seizures of unknown onset may have features that can still be classified.</td>
</tr>
<tr>
<td>4.</td>
<td>Awareness is used as a classifier of focal seizures.</td>
</tr>
<tr>
<td>5.</td>
<td>The terms dyscognitive, simple partial, complex partial, psychic, secondarily generalized were eliminated.</td>
</tr>
<tr>
<td>6.</td>
<td>New focal seizure types include automatisms, autonomic, behavior arrest, cognitive, emotional, hyperkinetic, sensory and focal to bilateral tonic-clonic seizures. Atonic, clonic, epileptic spasms, myoclonic and tonic seizures can be either focal or generalized.</td>
</tr>
<tr>
<td>7.</td>
<td>New generalized seizure types include absence with eyelid myoclonia, myoclonic absence, myoclonic-tonic-clonic, myoclonic-atonic, epileptic spasms.</td>
</tr>
</tbody>
</table>

Compared to the 1981 classification, certain seizure types now appear in multiple categories. Epileptic spasms can be of focal, generalized or unknown onset. Represented both in focal and generalized columns are atonic, clonic, myoclonic and tonic seizures, although the pathophysiology of these seizure types may differ for the focal onset versus generalized onset seizure type of that name.

The net effect of updating the Classification of Seizures should be the following: 1. Render the choice of a seizure type easier for seizures that did not fit into any prior categories; 2. Clarify what is meant when a seizure is said to be of a particular type; 3. Provide more
transparency of terminology to the nonmedical and medical community. A companion paper provides guidance on how to apply the 2017 classification. Employment of the 2017 classification in the field for a few years likely will motivate minor revisions and clarifications.

Key Points

- The ILAE has constructed a revised classification of seizure types. The classification is operational and not based on fundamental mechanisms.
- Reasons for revision include clarity of nomenclature, ability to classify some seizure types as either focal or generalized, and classification when onset is unknown.
- Seizures are divided into seizures of focal, generalized, unknown onset, with subcategories of motor, non-motor, with retained or impaired awareness for focal seizures.

Acknowledgements

Funding for this study was provided by the International League Against Epilepsy. The lead author (RSF) was supported by the Maslah Saul MD Chair, the James & Carrie Anderson Fund for Epilepsy, the Susan Horngren Fund and the Steve Chen Research Fund.

Special thanks are given to the Revision Task Force appointed to revise the classification after receipt of public comments. Members of this Revision Task Force do not necessarily concur with all details of the classification or the publication, since opinions were not always concordant. These Task Force members were: Carol D’Souza, Sheryl Haut, Ernest Somerville, Michael Sperling, Andreas Schulze-Bonhage, Elza Marcia Yacubian. Additional key comments were received from Soheyl Noachtar, Kimford Meador and Kevin Graber.

Disclosures

Disclosures relevant to classification: Dr. Fisher has stock options from Avails Pharmaceuticals, Cerebral Therapeutics, Applied Neurometrics, Zeto, Smart Monitor and research grants from Medtronic and the NSF.

J. A. French discloses support via The Epilepsy Study Consortium, which pays Dr French’s university employer for her consultant time related to Acorda, Alexza, Anavex, BioPharm Solutions, Concert, Eisai, Georgia Regents University, GW Pharma, Marathon, Marinus,
Neurelis, Novartis, Pfizer, Pfizer-Neusentis, Pronutria, Roivant, Sage, SciFluor, SK Life Sciences, Takeda, Turing, UCB Inc., Ultragenyx, Upsher Smith, Xenon Pharmaceuticals, Zynerba grants and research from Acorda, Alexza, LCGH, Eisai Medical Research, Lundbeck, Pfizer, SK life sciences, UCB, Upsher-Smith, Vertex, grants from NINDS, Epilepsy Therapy Project, Epilepsy Research Foundation, Epilepsy Study Consortium; She is on the editorial board of Lancet Neurology, Neurology Today and Epileptic disorders, and was an Associate Editor of Epilepsia, for which she received a fee.

Sheryl Haut is a consultant for Acorda and Neurelis.

Edouard Hirsch has received honoraria for lectures and/or advice from Novartis, EISAI, and UCB.

Dr. Moshé is the Charles Frost Chair in Neurosurgery and Neurology and funded by grants from NIH NS43209, CURE, US Department of Defense, the Heffer Family and the Segal Family Foundations and the Abbe Goldstein/Joshua Lurie and Laurie Marsh/ Dan Levitz families and receives from Elsevier an annual compensation for his work as Associate Editor in Neurobiology of Disease and royalties from 2 books he co-edited. He received a consultant’s fee from Lundbeck, Eisai and UCB.

Jukka Peltola has participated in clinical trials for Eisai, UCB, and Bial; received research grants from Eisai, Medtronic, UCB, and Cyberonics; received speaker honoraria from Cyberonics, Eisai, Medtronic, Orion Pharma, and UCB; received support for travel to congresses from Cyberonics, Eisai, Medtronic, and UCB; and participated in advisory boards for Cyberonics, Eisai, Medtronic, UCB, and Pfizer.

Dr Scheffer serves on the editorial boards of Neurology and Epileptic Disorders; may accrue future revenue on pending patent WO61/010176 (filed: 2008): Therapeutic Compound; has received speaker honoraria/consultant fees from GlaxoSmithKline, Athena Diagnostics, UCB, Eisai and Transgenomics.

Dr Zuberi is Editor-in-Chief of the European Journal of Paediatric Neurology for which he receives an annual honorarium from Elsevier Ltd. He has received research funding from Dravet Syndrome UK, Epilepsy Research UK, UCB Pharma and Glasgow Childrens’ Hospital Charity. The remaining authors listed no disclosures relevant to the classification of seizure types.

References


41. Bergen DC, Beghi E, Medina MT. Revising the ICD-10 codes for epilepsy and seizures. *Epilepsia* 2012;53 Suppl 2:3-5.


