Common terms for rare epilepsies: Synonyms, associated terms, and links to structured vocabularies

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SUMMARY

Identifying individuals with rare epilepsy syndromes in electronic data sources is difficult, in part because of missing codes in the International Classification of Diseases (ICD) system. Our objectives were the following: (1) to describe the representation of rare epilepsies in other medical vocabularies, to identify gaps; and (2) to compile synonyms and associated terms for rare epilepsies, to facilitate text and natural language processing tools for cohort identification and population-based surveillance. We describe the representation of 33 epilepsies in 3 vocabularies: Orphanet, SNOMED-CT, and UMLS-Metathesaurus. We compiled terms via 2 surveys, correspondence with parent advocates, and review of web resources and standard vocabularies. UMLS-Metathesaurus had entries for all 33 epilepsies, Orphanet 32, and SNOMED-CT 25. The vocabularies had redundancies and missing phenotypes. Emerging epilepsies (SCN2A-, SCN8A-, KCNQ2-, SLC13A5-, and SYNGAP-related epilepsies) were underrepresented. Survey and correspondence respondents included 160 providers, 375 caregivers, and 11 advocacy group leaders. Each epilepsy syndrome had a median of 15 (range 6–28) synonyms. Nineteen had associated terms, with a median of 4 (range 1–41). We conclude that medical vocabularies should fill gaps in representation of rare epilepsies to improve their value for epilepsy research. We encourage epilepsy researchers to use this resource to develop tools to identify individuals with rare epilepsies in electronic data sources.

KEY WORDS: Rare epilepsy, Terminology, Classification, Natural language processing, Synonyms.

One obstacle to epilepsy research is limited representation in diagnostic coding systems like the International Classification of Diseases (ICD) system. Only a few entities appear in versions 9 and 10, such as tuberous sclerosis (ICD-9-CM 759.5, ICD-10 Q85.1) and infantile spasms (ICD-9-CM 345.6, ICD-10 G40.82). Although version 10 adds juvenile myoclonic epilepsy (G40.B), absence epilepsy (G40.A), and Lennox-Gastaut syndrome (G40.81), many rare epilepsies appear in neither system. For example, Dravet syndrome is coded with the non-specific ICD-9 345.1 “Generalized Convulsive Epilepsy” or ICD-10 G40.4 “Other generalized epilepsy and epileptic syndromes.” This limits the utility of large administrative and clinical datasets for epilepsy research, despite the clear value of such data for research in epidemiology, comparative effectiveness, health services, and quality improvement.1

Given the increasing availability of electronic health records for research, computerized analysis of clinical
notes may be useful for finding patients with specific epilepsy syndromes. For example, text processing can identify children with complex febrile seizures, and natural language processing can identify candidates for epilepsy surgery.

One challenge for using these techniques is the diversity of terms for epilepsies. Dravet syndrome, for example, is “severe myoclonic epilepsy of infancy” for some clinicians and “intractable childhood epilepsy with generalized tonic clonic seizures” for others. Although existing standardized medical vocabularies document some synonyms, many vocabularies have gaps. For example, OMIM (Online Mendelian Inheritance in Man) lacks well-defined entries for infantile spasms, Lennox-Gastaut syndrome, and migrating partial seizures of infancy.

To facilitate the development of text and natural language processing tools for epilepsy, we compiled synonyms and associated terms for 33 rare epilepsies, with links to 3 standard medical vocabularies.

Vocabularies

Two investigators (ZG and BH) manually reviewed 3 standardized vocabularies: OrphaNet (www.orpha.net), Unified Medical Language System (UMLS) Metathesaurus (utslogin.nlm.nih.gov/cas/login), and Systematized Nomenclature of Medicine Clinical Terms (SNOMED-CT; browser.ihtsdotools.org). In SNOMED-CT and UMLS, we included terms and subterms (ie, “infantile spasms” and “refractory infantile spasms”) as well as pathognomonic physical examination findings (“Ash leaf spot, tuberous sclerosis”). We limited the search to terms in English.

Advocacy groups

We individually contacted advocacy group leaders via email. Each email contained a working list of synonyms and associated terms for the specified epilepsy, and asked the group leader to review, amend, and/or expand the list.

Clinical experts

Five clinical pediatric epilepsy specialists (EY, TM, PM, AN, and SW) reviewed the drafted compendium and made additional recommendations and edits.

Methods

Study design

We compiled synonyms and associated terms for rare epilepsies via 6 sources: a survey of pediatric neurology clinicians (“provider survey”), a survey of caregivers of individuals with rare epilepsies (“caregiver survey”), manual review of online resources, a manual review of 3 structured medical vocabularies, correspondence with leaders of rare epilepsy advocacy groups, and independent clinician review. We selected 33 rare epilepsies based on the parent-led advocacy groups that constitute the Rare Epilepsy Network (REN; ren.rti.org), an umbrella organization that fosters research collaboration.

Surveys

For each survey, we designed, piloted, and iteratively refined questions using online software (SurveyMonkey, Inc., San Mateo, CA, U.S.A.). The “provider survey” was distributed to members of the Child Neurology Society (CNS) via email. Five versions were distributed, each asking respondents to list synonyms for 5–6 of the 33 epilepsies. The “caregiver survey” was distributed to REN participants. It asked respondents to name the epilepsy affecting the individual, and then list synonyms used by clinicians and family members.

Websites

One investigator (ZG) manually reviewed several websites for more synonyms and associated terms. These included websites focused on epilepsy (epilepsy.com, dup15q.org), rare diseases (rarediseases.org, rarechromo.org, and ghr.nlm.nih.gov), and general knowledge (wikipedia.com).

Results

There were 160 respondents to the provider survey of the 1982 emailed CNS members (response rate 8%). These included 107 (67%) pediatric neurologists, 40 (25%) pediatric epilepsy specialists, and 13 (8%) other clinicians (ie, nurse, nurse practitioners, adult epilepsy specialist, physician assistant, EEG technician, or medical student). Twenty-six (16%) were physician trainees (ie, residents or fellows). Most worked in the United States (153; 96%), including responses from 33 states and the District of Columbia. The 7 international responses were from Canada (5), Lebanon (1), and the United Kingdom (1).

There were 375 respondents to the caregiver survey, of 1162 members of the REN (response rate 32%). Most (356; 95%) were parents of an individual with epilepsy. The remainder included 2 individuals with rare epilepsy (both progressive myoclonic epilepsy), 4 unspecified caregivers, and 13 unknown (left blank). The majority lived in the...
United States (319; 85%), including 48 states (none from Alaska or North Dakota). There were 55 responses from 20 countries, and 1 unknown.

Leaders of 11 advocacy groups added more terms for the following disorders: Aicardi syndrome, Doose syndrome, Dravet syndrome, hypothalamic hamartoma, Lennox-Gastaut syndrome, PCDH19, Phelan-McDermid syndrome, Ring chromosome 20, SCN8A, SYNGAP, and tuberous sclerosis.

The final list included all 33 rare epilepsies (Table 1), including 16 defined by phenotype and 17 by genotype. Across all 33 rare epilepsies, there was a median of 14 synonyms (range 6–28). Fifteen had no associated terms. The remaining 19 epilepsies had a median of 4 associated terms (range 1–41). For example, Aicardi syndrome had 6 synonyms (AS, Aicardi’s syndrome, Aicardi disease, Aicardi’s disease, Aicardi, and Aicardi’s) and 12 associated terms (microcephaly, retinal lacunae, agenesis of corpus callosum, absent corpus callosum, infantile spasms, polymicrogyria, porencephalic cysts, coloboma, optic disc, ACC, retinal lesions, and lacunae) (Table S1).

The UMLS-Metathesaurus had entries for all 33 epilepsies. Half (15) had 1 entry; the remaining had multiple entries, including 11 for Lennox-Gastaut syndrome, 18 for infantile spasms, and 25 for neuronal ceroid lipofuscinosis. Several entries were redundant—for example, there were 7 entries with overlapping descriptions of “refractory infantile spasms,” and 2 nearly identical entries for nonrefractory Lennox-Gastaut syndrome (Table 2). Five epilepsies (KCNQ2, SCN2A, SCN8A, SYNGAP, and SLC13A5) were not linked to disease entities, but rather to entries entitled “Caused by a mutation in [gene].”

OrphaNet had entries for 32 epilepsies. The missing epilepsy (SLC13A5) was linked with “ORPHA442835 Undetermined early-onset epileptic encephalopathy” but did not have its own entry. Three contained information about the relevant gene (SCN2A, SCN8A, and SYNGAP) only, without phenotype information (Table 2). Thirty had 1 entry, 2 had 7 entries: Angelman syndrome and neuronal ceroid lipofuscinosis.

SNOMED-CT had entries for 25 rare epilepsies (missing: CDKL5, Dup15q Syndrome, Glut1 deficiency, KCNQ2, SCN2A, SCN8A, SLC13A5, and SYNGAP; Table 2). Of these 25 rare epilepsies, 2 had multiple entries (6 for infantile spasms and 4 for tuberous sclerosis).

**Discussion**

**Summary**

We compiled synonyms and associated terms for 33 rare epilepsies, including links to 3 standardized vocabularies. The compilation is available freely online. There are gaps in these vocabularies, including poor representation of emerging epilepsies (eg, SCN2A, SCN8A, SYNGAP, KCNQ2, and SLC13A5), and redundancies of clinical concepts (treatment resistance).

**Significance**

This work builds an important resource to help clinicians and researchers find epilepsy subpopulations in electronic health record systems. Currently, to collect large cohorts of children with epilepsy, clinical researchers are building registries via recruitment of patients one by one in clinical settings. In contrast, text processing and/or natural language processing tools can quickly identify cohorts for observational research and clinical trials. Such tools may also support care coordination and quality improvement initiatives.

Three examples follow. First, case finding may be possible with simple text searches of clinical notes. This approach was used successfully (in combination with chart review) to identify children with febrile seizures. However, simple text search may fail when terms are ambiguous (“SCN1A pending”), negated (“SCN1A testing unrevealing”), or included in boilerplate text (“Our epilepsy gene panel includes SCN1A, SCN1B, SCN2A...”). Second, customized natural language processing algorithms may be developed for specific populations. For example, in epilepsy, recent work identified surgical candidates by applying support vector machines to the text of physician notes. Similar work in cardiology can identify individuals with heart failure. However, such algorithms are often tailored to a single center, and may be difficult to disseminate. Third, published epilepsy classification systems and ontologies could be integrated into existing natural language processing systems. This approach would require evaluation of baseline performance, identification of gaps (as we have done here), and ongoing review and maintenance.

The specific deficiencies in currently available vocabularies merit additional commentary. The phenotype of SCN2A encephalopathy is only beginning to emerge, largely based on international registry data. These phenotypes should be incorporated into the existing medical informatics infrastructure to support ongoing research into these diseases. In contrast, the epilepsies associated with SLC13A5 and SYNGAP are described only through small case series. Their full phenotype is not well understood.

Finally, redundancies in the epilepsy vocabularies can reduce their utility. Redundancies make it harder to use and maintain a structured vocabulary, and erode the premise that a one-to-one relationship ties each clinical concept to a term. For example, the redundant entries for “treatment resistance” is particularly important, given that (1) research often focuses on this population, and (2) treatment-resistant patients may benefit from referral to a comprehensive epilepsy center.
| Category                                      | Rare epilepsy                   | Synonyms | Associated terms | SNOMED-CT entries | OrphaNet entries | UMLS meta-theaurus |
|-----------------------------------------------|---------------------------------|----------|------------------|-------------------|-----------------|-------------------|
| Epilepsies primarily defined by phenotype     |                                 |          |                  |                   |                 |                   |
| Aicardi syndrome                              | 6                               | 12       | 1                | 1                 | 1               | 1                 |
| Dravet syndrome                               | 23                              | 0        | 1                | 1                 | 3               |                   |
| Hypothalamic hamartoma                        | 10                              | 5        | 1                | 1                 | 2               | 1                 |
| Infantile spasms                              | 9                               | 8        | 6                | 1                 | 18              |                   |
| Landau-Kleffner                               | 18                              | 0        | 1                | 1                 | 3               |                   |
| Lennox-Gastaut syndrome                       | 8                               | 2        | 1                | 1                 | 11              |                   |
| Migrating partial seizures of infancy         | 15                              | 0        | 1                | 1                 | 3               |                   |
| Myoclonic epilepsy with ragged red fibers     | 20                              | 2        | 1                | 1                 | 3               |                   |
| Neuronal ceroid lipofuscinosis                | 25                              | 1        | 1                | 1                 | 7               | 25                |
| Ohtahara syndrome                             | 8                               | 0        | 1                | 1                 | 3               |                   |
| Rasmussen syndrome                            | 9                               | 1        | 1                | 1                 | 3               |                   |
| Rett syndrome                                 | 17                              | 2        | 1                | 1                 | 5               |                   |
| Sturge-Weber                                  | 16                              | 6        | 1                | 1                 | 2               |                   |
| Tuberous sclerosis                            | 10                              | 41       | 4                | 1                 | 7               |                   |
| Epilepsies primarily defined by genotype      |                                 |          |                  |                   |                 |                   |
| Alpers disease (POLG mutations)               | 25                              | 1        | 1                | 1                 | 1               |                   |
| Angelman syndrome                             | 13                              | 0        | 1                | 1                 | 7               | 1                 |
| Epilepsy due to CDKL5 mutations               | 19                              | 0        | 0                | 1                 | 3               |                   |
| Dup15q                                        | 13                              | 0        | 0                | 1                 | 2               |                   |
| Fragile X syndrome                            | 17                              | 8        | 1                | 1                 | 1               |                   |
| Glut 1 deficiency                            | 17                              | 0        | 0                | 1                 | 1               |                   |
| KCNQ2 Encephalopathy                         | 28                              | 0        | 0                | 1                 | 2               |                   |
| PCDH19                                        | 21                              | 0        | 1                | 1                 | 2               |                   |
| Phelan-McDermid                               | 13                              | 25       | 1                | 1                 | 1               |                   |
| Prader-Willi                                  | 15                              | 3        | 1                | 1                 | 1               |                   |
| Ring chromosome 14                            | 6                               | 0        | 1                | 1                 | 1               |                   |
| Ring chromosome 20                            | 15                              | 0        | 1                | 1                 | 1               |                   |
| SCN2A encephalopathy                          | 18                              | 4        | 0                | 1                 | 2               |                   |
| SCN8A encephalopathy                          | 13                              | 0        | 0                | 1                 | 2               |                   |
| Epilepsy due to SLC13AS mutations             | 8                               | 0        | 0                | 1                 | 1               |                   |
| Epilepsy due to SYNGAP mutations              | 9                               | 17       | 0                | 1                 | 1               |                   |
| Unverricht-Lundborg Disease (CSTB mutations)  | 14                              | 0        | 1                | 1                 | 1               |                   |
| Deficiency       | Clinical concept                  | Vocabulary          | Terms                                                                 |
|------------------|-----------------------------------|---------------------|----------------------------------------------------------------------|
| Redundancy       | Lennox-Gastaut syndrome, not intractable | UMLS Metathesaurus | C3648103 “Lennox-Gastaut syndrome, not intractable”                   |
|                  |                                    |                     | C3494904 “Lennox-Gastaut syndrome, non-refractory”                    |
|                  | Infantile spasms, Intractable     | UMLS Metathesaurus | C0154716 “Infantile spasms, with intractable epilepsy”               |
|                  |                                    |                     | C1 827396 “Refractory infantile spasms”                              |
|                  |                                    |                     | C2712779 “Infantile spasms, poorly controlled”                        |
|                  |                                    |                     | C2712780 “Infantile spasms, refractory (medically)”                   |
|                  |                                    |                     | C2712781 “Infantile spasms, pharmacologically resistant”              |
|                  |                                    |                     | C2712782 “Infantile spasms, treatment resistant”                      |
|                  |                                    |                     | C3648801 “Infantile spasms with intractable epilepsy with status epilepticus” |
|                  |                                    |                     | C3837134 “Infantile spasms, with intractable epilepsy, without status epilepticus” |
|                  | SCN2A encephalopathy              | OrphaNet            | ORPHA118500 “SCN2A—sodium voltage-gated channel alpha subunit 2”     |
|                  |                                    | UMLS Metathesaurus | C2727014 “Caused by mutation in the alpha-1-subunit of the voltage-gated type II sodium channel gene (SCN2A, 182390.0002)” |
|                  |                                    | UMLS Metathesaurus | C27279128 “Caused by mutation in the voltage-gated, type II sodium channel, alpha subunit (SCN2A, 182390.0008)” |
|                  | SCN8A encephalopathy              | UMLS Metathesaurus | C2800426 “Caused by mutation in the voltage-gated sodium channel, type VIII, alpha subunit gene (SCN8A, 600702.0001)” |
|                  |                                    | UMLS Metathesaurus | C3553209 “Caused by mutation in the voltage-gated sodium channel, type VIII, alpha subunit gene (SCN8A, 600702.0002)” |
|                  | Epilepsy due to SYNGAP            | UMLS Metathesaurus | C3808213 “Caused by mutation in the synaptic Ras GT Pase activating protein 1 gene (SYNGAP1, 603384.0001)” |
|                  | Epilepsy due to SLC13A5            | UMLS Metathesaurus | C4014623 “Caused by mutation in the solute carrier family 13 (sodium-dependent citrate transporter), member 5 gene (SLC13A5, 608305.0001)” |
|                  | KCNQ2 encephalopathy              | UMLS Metathesaurus | C1 852593 “Caused by mutation in the potassium voltage-gated channel, KQT-like subfamily, member 2 gene (KCNQ2, 602235.0001)” |
|                  |                                    | UMLS Metathesaurus | C27279124 “Caused by mutation in the voltage-gated potassium channel, KQT-like subfamily, member 2 gene (KCNQ2, 602235.0007)” |
| No entry         | CDKL5 mutations                   | SNOMED-CT           | (missing)                                                            |
|                  | Dup 15q syndrome                  | SNOMED-CT           | (missing)                                                            |
|                  | Glu1 deficiency                   | SNOMED-CT           | (missing)                                                            |
|                  | KCNQ2 encephalopathy              | SNOMED-CT           | (missing)                                                            |
|                  | SCN2A encephalopathy              | SNOMED-CT           | (missing)                                                            |
|                  | SCN8A encephalopathy              | SNOMED-CT           | (missing)                                                            |
|                  | Epilepsy due to SYNGAP            | SNOMED-CT           | (missing)                                                            |
|                  | Epilepsy due to SLC13A5            | SNOMED-CT           | (missing)                                                            |
|                  | Epilepsy due to SLC13A5            | OrphaNet            | (missing)                                                            |
Limitations
First, our selection of rare epilepsies was guided by patient advocacy groups, each of which is unified by a specific feature of the affected individuals; for some a gene, and others a clinical phenotype. This may cause some individuals to be classified in multiple groups. PCDH19 mutations, for example, may cause the clinical phenotype of Dravet syndrome. Second, the survey respondents were largely US based, and the terms were limited to the English language. Further work is required to add non–English-language terms. Third, language evolves over time. Thus the compendium will need updating as terms arise or fade. Fourth, we selected only a subset of several hundred rare epilepsies. Fifth, the survey response rates were low. However, we generated a robust list of terms by supplementing the 2 surveys with other sources, thereby meeting the study objectives. Finally, our work is silent on how terms are used in clinical practice. Further work is needed to describe how terms appear in clinical notes to understand their value for case finding.

Conclusions
Epilepsy terms in structured medical vocabularies have gaps and redundancies, which should be addressed. These collected terms may help researchers and clinicians find individuals with rare epilepsies in electronic data sources. Further work is needed to evaluate their utility in identifying affected individuals.

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We confirm that we have read the Journal’s position on issues involved in ethical publication and affirm that this report is consistent with those guidelines.

Disclaimer
The findings and conclusions of this report are those of the authors and do not necessarily represent the official position of the Centers for Disease Control and Prevention.

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**Supporting Information**

Additional Supporting Information may be found in the online version of this article:

*Table S1. The Compendium of Terms.*