Seizures and movement disorders: phenomenology, diagnostic challenges and therapeutic approaches

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ABSTRACT
Seizures and movement disorders (MDs) are distinct neurological conditions presenting with abnormal movements. Despite sharing an overlap in phenomenology, these movements have different origins. In order to explore the overlaps and the narrow boundaries between these two conditions, we performed a review of the literature to explore the risk of seizures in MDs. We discussed the mimics and chameleons including MDs that look like seizure (eg, paroxysmal dyskinesia, status dystonicus) and seizures that look like MDs (eg, epilepsy partialis continua, nocturnal frontal lobe epilepsy). Additionally, we examined the therapeutic challenges as well as the anatomical and chemical pathways relevant in the interplay between epilepsy and MDs. Finally, we proposed an algorithm to guide clinicians towards the final diagnosis of conditions characterised by the co-occurrence of MDs and seizures.

INTRODUCTION
Movement disorders (MDs) are a group of neurological conditions characterised by abnormal movements that commonly arise from an altered function in the nuclei of the basal ganglia or their connections.1 Seizures are defined as ‘a transient occurrence of signs and/or symptoms due to abnormal excessive or synchronous neuronal activity in the brain’, whereas epilepsy is a brain disease characterised by one or more seizures with a relatively high recurrence risk.2

In spite of being caused by several different conditions, both MDs and seizures display abnormal movements with overlapping phenomenology. Given these phenomenological similarities as well as possible aetiological commonalities, clinicians are often challenged by conditions at the boundaries between epilepsy and MDs. In fact, several syndromes may exhibit coexisting abnormal movements from epileptic or non-epileptic origin regardless of the causes, which can overlap or be distinct.

This review aims to gather all the most updated literature with the ultimate goal of navigating clinicians through the complex interplay between MDs and seizures. We will discuss the seizure semiology and the risk of epileptic activity in MDs primarily presenting hypokinetic or hyperkinetic movements (or a combination of both). Most importantly, we will address the mimics that often pose challenges, including MDs that look like epileptic disorders and seizures that look like MDs.

Lastly, we will discuss the anatomical and chemical pathways involved in the overlap between seizures and MDs, as well as the treatment approaches. Moreover, we propose an algorithm to guide clinicians towards the final diagnosis of conditions characterised by the co-occurrence of MDs and seizures.

The most recent nomenclature of genetic MDs and the new revised operational classification by the International League Against Epilepsy will be used in this review.3 4 In each section, the disorders will be listed in order of prevalence, starting with the most frequent ones.

SEARCH METHODS
This review was conducted through a literature review in Medline and Embase databases of articles published between 1 January 2013 and 1 April 2018. Only articles in English and on humans were included. The search strategy included the combination of the following terms: ((seizures) OR (epilepsy)) AND ((movement disorders) OR (chorea) OR (tremor) OR (stereotypies) OR (parkinsonism) OR (ataxia) OR (dystonia) OR (dyskinesia) OR (tics) OR (mimics) OR (paroxysmal movement disorders)). The initial search identified 604 articles. Among those, 57 articles were selected according to the presence of pertinent terms in the abstract and title. Additional references, identified from the reference list of included articles but that were not part of the original search results, were included at authors’ discretion. We focused on the scientific literature of the past 5 years, but also included older publications of high merit or originality.

RISK OF SEIZURES IN MDs
Among the hypokinesias, Parkinson’s disease (PD) is one of the most prevalent neurodegenerative disorders worldwide. The relation between PD and seizures has been controversial, with small studies showing a positive correlation, but generally with a low prevalence of seizures in patients with PD.5 However, a more recent study demonstrated that patients with PD who did not have any seizure-provoking comorbidity had an adjusted OR of epileptic seizures of 2.24 compared with PD-free individuals without any seizure-provoking comorbidity.6 Although the association between PD and seizures remains unclear, superimposed brain disorders such as cerebrovascular disease, infections, surgery and trauma tend to be more strongly associated with seizures in patients with PD.6

DYT-ATP7B or Wilson’s disease is a hepatolenticular degeneration due to mutations in the ATP7B gene leading to an impaired copper metabolism. Majority of patients exhibit neurological and hepatic symptoms. DYT-ATP7B may display hypokinetic or hyperkinetic movements. Seizures can occur in approximately 10% of patients, including generalised tonic-clonic (GTC) and focal onset seizures. Patients with seizures more often have white matter changes than those without (figure 1A). 

Idiopathic basal ganglia calcification, formerly known as Fahr’s disease, is characterised by abnormal deposition of calcium in the basal ganglia and other parts of the brain (figure 1B), which leads to hypokinetic or hyperkinetic movements combined with dementia, neuropsychiatric symptoms and GTC seizures. It is a rare inherited or sporadic neurological condition due to genetic causes or secondary to an endocrine disorder (eg, hypoparathyroidism or pseudohyoparathyroidism). Mutations in SLC20A2, PFBC, PDGFB and PDGFRB have been associated with Fahr’s disease, with SLC20A2 being the most common causative gene. 

Presence of parkinsonism is more often seen in SLC20A2 mutations, headache in PDGFB and GTC seizures in patients with pseudohyoparathyroidism. 

Cerebrotendinous xanthomatosis (CTX) is a rare autosomal recessive inborn disorder of bile acid metabolism due to mutations in the CYP27A1 gene. GTC seizures and parkinsonism are the initial neurological features of CTX. Xanthomas are one of the hallmarks of this condition and usually begin to form in early adulthood and most commonly seen in tendons. 

Neurodegeneration with brain iron accumulation (NBIA) is a group of disorders characterised by excess iron accumulation in the globus pallidus and other areas, depending on the underlying genetic abnormality (figure 1C–D). NBIA clinically presents as progressive hypokinetic and/or hyperkinetic MDs with a variable degree of cognitive and psychiatric involvement. The most common NBIA mutations causing seizures are PLA2G6-associated neurodegeneration, beta-propeller protein-associated neurodegeneration, mitochondrial membrane protein-associated neurodegeneration and fatty acid hydroxylase-associated neurodegeneration. Although rare, seizures have also been reported in NBIA/DYT/PANK2 and aceruloplasminaemia. 

Huntington’s disease (CHOR-HTT) is a hereditary neurodegenerative condition caused by a CAG repeat expansion in the HTT gene leading to chorea, cognitive deterioration and psychiatric problems. Seizures are more common in juvenile Huntington’s disease (onset before age 21), whereas chorea is less common. Seizures have also been reported in Huntington’s disease-like syndromes, such as CHOR-PRN or HDL1, and SCA-TBP or HDL4/SCA17, but not in CHOR-JPH3 or HDL2. 

Chorea-acanthocytosis (CHOR-VPS13A) and McLeod syndrome (CHOR-XK) are characterised by neurodegeneration of the basal ganglia and red cell acanthocytosis. CHOR-VPS13A is a rare autosomal recessive neurodegenerative disorder due to mutations in the vacuolar protein sorting 13 homolog A (VPS13A). Seizures are rare, but can be the first symptom in unusual presentations. CHOR-XK is inherited in an X linked manner and is caused by mutations in the XK gene encoding an antigen of the Kell blood group system. Cardiomyopathy and arrhythmia are distinguishing features and patients can also display seizures. 

Spinocerebellar ataxias (SCAs) are a group of clinically and genetically heterogeneous neurodegenerative disorders characterised by progressive ataxia and a broad spectrum of other neurological findings. Seizures are rare, being reported in 5% of patients at disease onset. Dentatorubral-pallidoluysian atrophy (DRPLA) is the condition with the highest prevalence of seizures among the dominant cerebellar ataxias; however, seizures have also been reported in SCA 10, 17, 19 and 22. DRPLA is caused by a CAG trinucleotide repeat expansion and is characterised by various combinations of cerebellar ataxia, choreoathetosis, myoclonus, epilepsy, dementia and psychiatric symptoms. Epileptic seizures are a common feature in all patients with onset before the age of 20, whereas the occurrence of seizures in patients with onset after the age of 40 is rare. Tonic, clonic or GTC seizures are observed in DRPLA.

Autosomal dominant cortical tremor, myoclonus and epilepsy are a non-progressive disorder with an age of onset of 12–50 years characterised by the variable combination of ‘cortical tremor’, myoclonus and seizures (mostly GTC). Cortical tremor is an action and postural fine shivering movement consisting of continuous, arrhythmic, mainly distal, fine twitches in the hands resembling essential tremor. A neurophysiological study can help to distinguish cortical tremor versus essential tremor. Usually, patients have significant clinical improvement with antiepileptic drugs including valproate and clonazepam. 

Tourette syndrome (TS) is characterised by multiple motor tics with at least one vocal tic and a high rate of psychiatric comorbidities. In a recent retrospective cohort study, the TS group had an 18.4-fold increased risk of seizures than the control group. Even after adjusting for comorbidities, the risk of seizures in the TS group remained high. Although this single study suggested a higher risk of seizures, the aetiological link between seizures and TS remains to be investigated and further studies are warranted. 

Alternating hemiplegia of childhood (AHC) is associated with a variety of neurological features including episodes of hemiplegia and choreoathetosis. AHC is associated with mutations in ATPase Na+/K+ transporting subunit alpha 3 (ATP1A3) gene. Epileptic seizures may occur around the time of onset of hemiplegic attacks, including GTC, focal seizures and status epilepticus (SE). Approximately 50% of patients with AHC experience a seizure at least once in their lifetime. More recently, two cases of AHC phenotype were described with mutations in ADCYS gene (see below). 

Stiff-person spectrum disorder comprises the stiff-person syndrome (SPS), a segmental form called stiff-limb syndrome and the more severe disease called progressive encephalomyelitis with rigidity and myoclonus (PERM). SPS presents muscle rigidity and painful spasms that occur spontaneously or are triggered by diverse stimuli. Overall, seizures are a rare symptom: a few cases have been described in SPS, but it is most often
associated with PERM, in which seizures have been found in approximately 13% of patients.21 N-methyl-D-aspartate receptor (NMDAR) antibody-associated encephalitis is a disorder associated with the presence of autoantibodies directed against the extracellular domain of the NR1 subunit of the NMDAR.22 NMDAR encephalitis can present a broad spectrum of hyperkinetic movements including oro-lingual-facial dyskinesias (most common MDs), limb and trunk choreoathetosis, elaborate motions of arms and legs, oculogyric crisis, dystonia, rigidity, and opisthotonic postures.23 Seizures can take place at any time during the disease, but tend to occur earlier in males.24 The most common type of seizures seen are GTC, focal impaired awareness seizures (FIAS) and focal to bilateral tonic-clonic seizures. Refractory SE and epilepsy partialis continua (EPC) can occur at any time during the illness. Seizure frequency and intensity tend to decrease as the disease evolves. Residual epilepsy is very infrequent after recovery.

Tonic-dystonic seizures (TDS), previously called facioabracial dystonic crises, are considered the prototype of the borderline between MD and seizures. TDS are associated with antibodies against the neuronal target LGI1 protein and are characterised by very brief and frequent episodes of abnormal posturing affecting mostly the arm and ipsilateral face and leg.25 TDS often overlap with other types of seizures, such as autonomic, temporal lobe and GTC.25 These type of seizures often precede the development of limbic encephalitis and other symptoms of LGI1 antibody-mediated encephalitis; therefore, prompt recognition and treatment with immunotherapy of TDS may prevent the development of full-blown encephalitis.25 Contactin-associated protein-like 2 (CASPR2) is another specific target protein in cases of autoimmune encephalitis. Some clinical features may differentiate patients with antibodies against these two components (LGI1 and CASPR2) of the voltage gated potassium channel complex (VGKC).22 For example, the presence of TDS and hyponatraemia strongly suggests LGI1 reactivity, whereas the presence of neuromyotonia or other features of Morvan’s syndrome (eg, thymoma) suggests CASPR2-directed autoantibodies (sometimes accompanied by LGI1 antibodies).

Most antibody-mediated encephalitis can manifest with seizures and less frequently with MDs.22 The ones more frequently associated with seizures are those related to antibodies against inhibitory receptors such as neurotransmitter gamma-aminobutyric acid receptors type A and B (GABAa and GABAb receptors).22 In young children, anti-GABAA receptors encephalitis can mimic anti-NMDAR encephalitis.26 In young patients with encephalitis and prominent symptoms of basal ganglia dysfunction (chorea, parkinsonism, dystonia), the encephalitis associated with antibodies against dopamine 2 receptor should be considered, although this is extremely rare.22

Lastly, among the context of autoimmune MDs, Hashimoto’s encephalopathy is defined by the combination of neuropsychiatric symptoms, laboratory evidence of antithyroid antibodies and prominent clinical improvement with steroids, along with lack of evidence of other disorders.25 Clinical features include confusion, hallucinations, seizures, SE, tremor, myoclonus, gait ataxia, palatal myoclonus or chorea.27 High titres of antithyroid peroxidase antibodies are usually observed; however, most patients are euthyroid or have subclinical hypothyroidism; overt hypothyroidism occurs in 20% of cases. Back-averaged electroencephalogram (EEG) does not show a cortical correlate with myoclonus, suggesting a subcortical origin.

Subacute sclerosing panencephalitis (SSPE) typically results from persistent measles virus infection at a young age. Myoclonus is the most common and frequently the first symptom.28 Rare and less typical presentations of SSPE include seizures, EPC, acute encephalopathy, hemiparesis, acute ataxia, unilateral dystonia or psychiatric disturbances.

Creutzfeldt-Jakob disease (CJD) is a rapidly progressive transmissible spongiform encephalopathy that can be familial, sporadic or acquired. The typical clinical triad consists of rapidly progressive dementia, startle myoclonus, and characteristic periodic sharp complexes on the EEG and peculiar findings on MRI (figure 1E).29 Seizures are an uncommon finding in CJD, occurring in less than 15% of patients, and atypical presentations have been previously reported including non-convulsive SE and EPC.30

**MIMICS AND CHAMELEONS: MDs that look like seizures**

These are MDs that look like seizures due to the occurrence of episodic involuntary movement. Subcortical mechanisms are involved, thus explaining why EEG does not show epileptic activity in most cases.

Hemifacial spasm (HFS) is an MD of the seventh cranial nerve which is characterised by either brief or persistent, intermittent twitching of the muscles innervated by the facial nerve.31 Primary HFS likely results from compression of the seventh nerve at the root exit zone in the posterior cranial fossa by an aberrant or ecstatic vessel, whereas secondary causes are seen after VII nerve palsy or brainstem lesions.32 HFS may be mistaken for EPC involving the face (see below).32

Periodic limb movements (PLMS) are sleep-related phenomena characterised by periodic episodes of repetitive and highly stereotyped limb movements, which most often occur in the lower extremities. PLMS can be associated with brief arousals in sleep and autonomic reaction, and is defined as the presence of >15 movements per hour and a complaint of insomnia and/or excessive daytime sleepiness with no other explanation for these symptoms.33 PLMS are frequently seen in restless legs syndrome but can also occur in sleep disordered breathing, narcolepsy and rapid eye movement (REM) sleep behaviour disorder, as well as in several medical conditions, like renal failure, essential hypertension and PD, or associated with medication intake (antidepressants and neuroleptic agents).33 PLMS may mimic focal seizures, and in some cases further investigation (eg, EEG) is required.34

Status dystonicus (SD) is currently defined as ‘a movement disorder emergency characterized by severe episodes of generalized or focal hyperkinetic movements that have necessitated urgent hospital admission because of the direct life-threatening complications of these movements’.35 36 SD can be triggered by precipitating factors such as infections, medication adjustments, surgical procedures, metabolic disorders and failure of deep brain stimulation. SD usually occurs as a result of a continuum of worsening dystonia with a preserved level of consciousness and normal EEG. Cerebral palsy, NBIA/DYT-PANK2, DYT-TOR1A, DYT-ATP7B and mitochondrial disorders are the main aetiologies of SD.35

Rare tremor syndromes may be episodic and mimic seizures; the differential diagnosis may include EPC and myoclonus.36 These syndromes are limb-shaking transient ischaemic attacks syndrome, hereditary chin tremor, paroxysmal head tremor, bobble-head doll syndrome, spasmus nutans and shuddering attacks (table 1). Paroxysmal dyskinesias are an important paradigm of the overlap between MDs and seizures.37 It is a heterogeneous
group of disorders characterised by episodes of abnormal involuntary movements (dystonic, choreiform, ballistic) without loss of consciousness and sometimes preceded by a sensory aura. These movements can be mistaken for focal seizures including focal aware seizure (FAS) or FIAS. Although in paroxysmal dyskinesias scalp EEG does not show ictal discharges, many frontal lobe seizures that look like dyskinesias can also present normal EEG. Paroxysmal dyskinesia can be subdivided into three clinical syndromes: paroxysmal exercise-induced dyskinesia (PED), paroxysmal kinesigenic dyskinesia (PKD) and paroxysmal non-kinesigenic dyskinesia (PNKD), each of which is associated with the known causative genes SLC2A1, PRRT2 and PNKD, respectively, although a certain genetic heterogeneity is recognised (table 2). Glucose transporter type-1 deficiency syndrome (GLUT1DS) is also caused by mutations in SLC2A1 presenting early-onset refractory seizures, PED and MDs. Approximately 90% of patients have clinical seizures, mainly GTC, followed by absence, myoclonic and focial onset seizures. PRRT2 mutations can also produce epileptic disorders. Benign familial infantile seizure (BFIS) is an autosomal dominant syndrome characterised by brief seizures featuring motor arrest, cyanosis, hypertonia and limb jerks with onset between 3 and 12 months of age that generally remit before the age of 2 years. Up to 80% of cases with BFIS have PRRT2 mutations. Analogously, almost 90% of cases with ICCA syndrome (infantile convulsions and choreoathetosis) carry PRRT2 mutations. This syndrome is characterised by infantile convulsions that are followed by the development of PKD, usually by the age of 5 years. More recently, a number of different genetic syndromes have also been reported to cause paroxysmal dyskinesia associated with epilepsy. This is the case of mutations in the KCNMA1 gene that produce a syndrome of PNKD and epilepsy, either in the form of absence or GTC. Mutations in the SCN8A cause a variety of seizure types along with episodes of paroxysmal dystonia, sometimes resembling PKD. Furthermore, ECHS1 mutations and pyruvate dehydrogenase complex deficiency can also produce both epilepsy and paroxysmal dyskinesia, usually embedded in a more complex syndrome of lactic acidosis and encephalopathy.

Startle syndrome is another example of paroxysmal disorder and should be included in the differential diagnosis of epileptic seizures. It includes hyperekplexia, stimulus-induced disorders (which can be epileptic or not), and neuropsychiatric syndromes such as startle-induced tics, culture-specific disorders (eg, Latah) and functional startle syndromes. The most relevant disorder is hyperekplexia, which is characterised by generalised stiffness at birth, excessive startle reflexes and generalised stiffness following startle; it is usually associated with mutations of the glycine receptor. The preserved consciousness and normal EEG distinguish it from epileptic seizures.

Patients with ADCY5 gene mutations (CHOR/DYT-ADCY5) have been associated with heterogeneous syndromes with a broader variety of MDs, such as dyskinesias (chorea, myoclonus, ballism or choreoathetosis); axial hypotonia that leads to a broader variety of MDs, such as dyskinesias (chorea, myoclonus, ballism or choreoathetosis); axial hypotonia that leads to an ‘gonadal-like’ ambulation; dystonic spasms; spasticity; or eye movement abnormalities. A notable characteristic of these patients is that dyskinesias have nocturnal exacerbations, which often lead to the suspicion of a seizure disorder, although video EEG telemetry is invariably negative in all the patients reported. A relation between CHOR/DYT-ADCY5 and seizures is not completely established, but only 1 of the 20 patients described in the largest cohort published to date had associated seizures.

Epilepsy that looks like MDs

These are seizures (with EEG-confirmed cortical origin) causing involuntary movements resembling MDs from a phenomenological standpoint.

Focal seizures are classified according to the level of awareness (FAS or FIAS) and can display motor onset or non-motor onset, reflecting the first prominent sign or symptom in the seizure. 1

### Table 1: Episodic tremor syndromes

<table>
<thead>
<tr>
<th>Syndrome</th>
<th>Age at onset</th>
<th>Cause</th>
<th>Phenotype</th>
<th>Duration</th>
<th>Recurrence</th>
<th>Triggers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Limb shaking</td>
<td>Adulthood.</td>
<td>Vascular: severe carotid stenosis (&gt;80%).</td>
<td>Typically unilateral, upper limb jerking tremor.</td>
<td>Transient—may last several minutes.</td>
<td>Episodes occur monthly or up to several times per day.</td>
<td>Orthostatic positional change, hypotension, hyperventilation, neck extension or walking.</td>
</tr>
<tr>
<td>Hereditary chin tremor</td>
<td>Adolescence.</td>
<td>AD linked to chromosome 9q13-q21 in one family.</td>
<td>Oscillatory rhythmic movements of the chin muscles, mainly the mentalis muscle. Frequency varies between 2 and 11 Hz.</td>
<td>Seconds to hours.</td>
<td>Daily.</td>
<td>Emotion or anxiety.</td>
</tr>
<tr>
<td>Paroxysmal head tremor</td>
<td>Adulthood.</td>
<td>Heterogeneity: missense mutation in CACNA1A gene, encoding the calcium voltage-gated channel.</td>
<td>Progressive disorder with ‘no-no’ pattern and frequency of 3–5 Hz involving the splenius and sternocleidomastoid muscles.</td>
<td>5 and 60 min.</td>
<td>Up to several times a week.</td>
<td>No specific triggers.</td>
</tr>
<tr>
<td>Bobble-head doll syndrome</td>
<td>Childhood.</td>
<td>Expansion in the region of the III ventricle resulting from a suprasellar arachnoid or a cyst.</td>
<td>Episodic 2–3 Hz ‘yes-yes’ tremor of the head with occasional ‘no-no’ movements. Other areas: neck, shoulders, trunk or upper limbs.</td>
<td>Variable.</td>
<td>Variable.</td>
<td>Absent during sleep. Amplitude increases with walking or excitement and decreases or disappears under volitional control or concentration.</td>
</tr>
<tr>
<td>Shuddering attacks</td>
<td>Childhood.</td>
<td>Idiopathic and benign.</td>
<td>Brief bursts of rapid, shivering-like movements of the head and both arms.</td>
<td>Seconds.</td>
<td>Up to 100 episodes per day.</td>
<td>No triggers.</td>
</tr>
</tbody>
</table>

**AD**, autosomal dominant.
Focal seizures with motor-onset symptoms can be mistaken for different MDs. For example, automatisms in FIAS may mimic orofacial dyskinesia, focal tonic seizures may be interpreted as dystonia, and focal clonic seizures may be interpreted as transient tremor. Ictal EEG may be negative in FAS.

EPC is a rare type of focal SE that may last from days to years, and its characteristic semiological features are spontaneous regular or irregular muscular twitching that may wax and wane and affects a limited part of the body. EPC is sometimes aggravated by action or sensory stimuli, occurring for a minimum of 1 hour and recurring at intervals of no more than 10 s. Common causes of EPC are cerebrovascular disease and tumours in adults and Rasmussen encephalitis in children, followed by metabolic, toxic and other immune-mediated causes. Of note, hyperosmolar hyperglycaemic (a treatable metabolic emergency that requires prompt and accurate diagnosis) can present a broad spectrum of neurological manifestations including EPC, focal seizures and may evolve into Lennox-Gastaut syndrome. Infantile spasms are often difficult to differentiate on the clinical ground alone unless EEG polysomnography is performed. Adding to the diagnostic difficulty, EEG is typically normal or may be masked by excessive movement artefact during the events. The high incidence of uninformative EEG emphasises the importance of other characteristics in the differential diagnosis (eg, the highly stereotyped nature of the episodes). Mutations in CHRNA4, KCNT1 and DEPDC5 have been associated with NFLE; however, most of the cases are non-familial and often the genetic cause is not found.

Progressive myoclonus epilepsies (PME) comprise a disabling group of rare disorders characterised by the coexistence of myoclonus, ataxia and epilepsy (table 3). Myoclonus is present at rest, but it can be triggered by movement, noise, light or touch. Recently, new PME genes have been reported, such as Neuronal, AFG3L2, GOSR2, SCARB2 and PRICKLE1.

Head drops as a seizure manifestation are abrupt and often repetitive episodes of neck atonia occurring from one to several times per day in a regular or irregular fashion, and are often part of epileptic syndromes like myoclonic-astatic epilepsy, myoclonic epilepsy of infancy or severe epileptic encephalopathies. Infantile spasms or West syndrome is a severe condition often associated with intractable epilepsy and developmental delay, and may evolve into Lennox-Gastaut syndrome. Infantile spasms

<table>
<thead>
<tr>
<th>Principal causative gene</th>
<th>SLC2A1</th>
<th>PRRT2</th>
<th>PNKD</th>
<th>KNCMA1</th>
<th>SCN8A</th>
<th>ECHS1 and PDC deficiency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gene characteristics</td>
<td>Solute carrier family 2 member 1. This gene encodes a major glucose transporter in the blood–brain barrier.</td>
<td>Proline-rich transmembrane protein 2. This gene encodes a transmembrane protein in the brain and spinal cord.</td>
<td>Paroxysmal non-kinesigenic dyskinesia. This gene is thought to play a role in the regulation of myofibrillogenesis.</td>
<td>Potassium calcium-activated channel subfamily M alpha 1 involved in the control of smooth muscle tone and neuronal excitability.</td>
<td>Sodium voltage-gated channel alpha subunit B involved in the rapid membrane depolarisation that occurs in excitable neurons.</td>
<td>Enoli-CoA hydratase, short chain 1 caused by mutations in one of the genes encoding the 3 PDC subunits (PDHA1, DLAT, PDHX).</td>
</tr>
<tr>
<td>Main clinical syndrome</td>
<td>PED.</td>
<td>PKD.</td>
<td>PNKD.</td>
<td>PNKD + epilepsy.</td>
<td>PKD + epilepsy.</td>
<td>PED + epilepsy.</td>
</tr>
<tr>
<td>Phenomenology and clinical features</td>
<td>Duration of episodes: few minutes up to 2 hours. Frequency: once per day or few times per month. Lower limb dystonia.</td>
<td>Duration of episodes: seconds to minutes. Frequency: up to 100 times per day. Dystonia, chorea, or athetosis—usually unilateral.</td>
<td>Duration of episodes: minutes to hours, sometimes up to a day. Frequency: few episodes per week or just a few episodes in a lifetime. Dystonia, chorea or athetosis involving one limb and gradually spreading to other limbs and face.</td>
<td>Duration of episodes: minutes to hours, either in the form of absence or GTSeizures.</td>
<td>FNKD and epilepsy, either in the form of absence or GTSeizures.</td>
<td>Focal, tonic, clonic, myoclonic and absence of seizures, along with episodes sometimes resembling PNKD.</td>
</tr>
<tr>
<td>Triggers</td>
<td>Prolonged exercise, and rarely by muscle vibration, cold or passive movements.</td>
<td>Sudden movement, such as sudden acceleration or change in direction of movement, or startle.</td>
<td>Caffeine, alcohol, fatigue or emotional stress.</td>
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</tr>
<tr>
<td>Additional disorders related to causative gene</td>
<td>GLUT1DS: early-onset refractory seizures and MDs (dystonia, chorea, myoclonus and PED).</td>
<td>BFIS: AD syndrome characterised by brief seizures featuring motor arrest, cyanosis, hypertonia and limb jerks. ICCA: infantile convulsions followed by PNKD.</td>
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</table>

AD, autosomal dominant; BFIS, benign familial infantile seizures; ECHS1, gene encoding PDC subunits; GLUT1DS, glucose transporter type-1 deficiency syndrome; GTC, generalised tonic-clonic seizures; ICCA, infantile convulsions and choreoathetosis syndrome; KNCMA1, gene encoding for a subunit of a calcium-activated potassium channel; MDs, movement disorders; PDC, pyruvate dehydrogenase complex; PED, paroxysmal exercise-induced dyskinesia; PKD, paroxysmal kinesigenic dyskinesia; PNKD, paroxysmal non-kinesigenic dyskinesia; SCN8A, encodes the alpha subunit of the sodium channel Nav1.
are characterised by brief and symmetrical contraction of neck, trunk and arms several times per day. The head appears to drop while the arms often elevate. Epileptic head drops and infantile myoclonus can be treated with levetiracetam, valproic acid and clonazepam as first-line agents, whereas subcortical and brainstem myoclonus can be treated with levetiracetam, valproic acid and clonazepam.

**TREATMENT APPROACHES**

The treatment of focal seizures if associated with an MD should begin with the medications demonstrated in randomised clinical trials to be effective and safe, in particular seizure type or epilepsy syndrome. Some antiepileptic drugs may exacerbate myoclonus when used in patients with generalised onset epilepsy; this effect is generally seen in patients with an epilepsy syndrome that includes myoclonus, for example, juvenile myoclonic epilepsy. Drugs reported to exacerbate myoclonus include carbamazepine, phenytoin, vigabatrin, gabapentin, pregabalin and lamotrigine.

Certain antiepileptic drugs may have adverse effects, making them less desirable in a patient with a known MD. Several antiepileptic drugs can produce tremor, most notably valproate, with a clear dose-related effect. Valproate has also been associated with drug-induced parkinsonism. The following antiepileptic drugs have tremor listed in prescribing information: lacosamide, lamotrigine, topiramate, oxcarbazepine and tiagabine. The following antiepileptic drugs have tremor listed in prescribing information: lacosamide, lamotrigine, topiramate, oxcarbazepine and tiagabine.

Additionaly, there is a significant overlap between the treatment of epilepsy and MDs as antiepileptic drugs can be beneficial to both. Essential tremor is often treated with primidone, topiramate (both first-line treatment) or gabapentin. Cortical myoclonus can be treated with levetiracetam, valproic acid and clonazepam as first-line agents, whereas subcortical and brainstem movements are characterised by brief and symmetrical contraction of neck, trunk and arms several times per day. The head appears to drop while the arms often elevate. Epileptic head drops and infantile myoclonus can be treated with levetiracetam, valproic acid and clonazepam. Subcortical and brainstem myoclonus can be treated with levetiracetam, valproic acid and clonazepam. Subcortical and brainstem movements are characterised by brief and symmetrical contraction of neck, trunk and arms several times per day. The head appears to drop while the arms often elevate. Epileptic head drops and infantile myoclonus can be treated with levetiracetam, valproic acid and clonazepam.
myoclonus can be treated with clonazepam as a first-line agent, but levetiracetam and valproic acid can be tried as well.53 PKD usually responds well to anticonvulsants like carbamazepine and phenytoin, often in low doses. Other anticonvulsants can also be tried, such as topiramate, levetiracetam, lamotrigine, valproate, oxcarbazepine and phenobarbital.37 Conversely, in PNKD, antiepileptic drugs are not always effective.37

PATHOPHYSIOLOGY

Traditionally, epilepsy has been considered a disorder originating from the cerebral cortex, while MDs have mostly been construed to reflect a dysfunction in subcortical areas. However, a net distinction between the pathophysiology of purely subcortical and cortical events risks to be an oversimplification owing to strong reciprocal connections between the basal ganglia and the cerebral cortex, particularly within the frontostriatal pathway. For instance, in one case with epilepsy and paroxysmal dyskinesia, the epileptogenic source was identified within the supplementary sensory motor cortex, but the ictal discharge rapidly spread to the basal ganglia.58 In patients with SLC2A1 mutations manifesting with epilepsy and PED, ictal single-photon emission computed tomography (SPECT) demonstrated the involvement of a network including both basal ganglia and cortical motor areas.59 Similarly, electrodecremental events have been described as the ictal EEG correlates of TDS associated with LGI1 antibodies.25 Some authors argue that simultaneous pathogenic involvement of the basal ganglia and the cortex may explain the overlap of cortical (epileptic) and subcortical (MDs) features of these spells, as further suggested by a positron-emission tomography (PET) study showing a concomitant involvement of the motor cortex, the hippocampus and of the striatum.60 It should be noted, however, that metabolic connectivity does not necessarily reflect anatomical connectivity but rather the coupled energy demand of different brain areas.61 Therefore, it is not surprising that disorders in which energy failure represents the main pathomechanism (eg, GLUT1DS, ECHS1 mutations or PDC deficiency) might display both epilepsy and MDs, as a function of energy failure of both cortical and subcortical areas, regardless of their interconnectivity.

These notions can be applied to many other epileptic conditions. For example, different distribution of subcortical dopaminergic dysfunction can be seen in epileptic syndromes using PET studies with dopamine transporter ligand. Patients with juvenile myoclonic epilepsy have shown lower tracer binding in the midbrain, whereas patients with GTC had reduced binding in the putamen.62 Similarly, the regional distribution of the products of mutated genes at both cortical and subcortical levels has been suggested.
to represent another mechanism whereby epilepsy and MDs might coexist in conditions genetically determined. This might be the case, for instance, of PRRT2 that is highly expressed at both sites.37

In contrast to PRRT2 mutations, PNKD mutations encompass paroxysmal MDs but not seizure, although they are also expressed at the cortical level.38 Other mechanisms are therefore possible. In fact, striatal neurons are more susceptible than cortical ones to PNKD dysfunction, suggesting that neuronal vulnerability to specific pathomechanisms might also account for the relative risk of seizure across different MDs.39

Furthermore, mutation type might further modulate epilepsy risk. As an example, SCN8A mutations associated with gain-of-function and high channel activity typically result in epileptic encephalopathy, whereas loss-of-function SCN8A mutations, which impair trafficking of the channel protein, are associated with isolated intellectual disability or MDs without seizures.40

CONCLUSIONS AND FUTURE DIRECTIONS

Despite being distinct neurological conditions, seizures and MDs present abnormal and involuntary movements with a well-defined phenomenology overlap, but with different pathophysiology. The clinical diagnosis can be challenging, and while a positive EEG is a useful tool to provide supporting evidence towards a diagnosis of seizure, a negative EEG has limited utility. In a few cases, the boundaries are well established, for instance in EPC and NFLE (epileptic origin) or in paroxysmal dyskinesia and SD (well-known MDs). However, these boundaries are blurred in disorders manifesting seizures and MDs concomitantly. These conditions represent the majority of cases discussed in our review, and although the clinical diagnosis can be more challenging, the correct definition of the MDs associated with seizure can adequately inform the diagnostic process. Our extensive review of the literature allowed us to propose an algorithm to facilitate the clinical approach, improve the diagnostic accuracy and guide clinicians who face these challenges in their daily practice (figure 2).

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