Expanding the Treatment Landscape for Lennox-Gastaut Syndrome: Current and Future Strategies

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Abstract

Lennox-Gastaut syndrome (LGS), a childhood-onset severe developmental and epileptic encephalopathy (DEE), is an entity that encompasses a heterogenous group of aetiologies, with no single genetic cause. It is characterised by multiple seizure types, an abnormal EEG with generalised slow spike and wave discharges and cognitive impairment, associated with high morbidity and profound effects on the quality of life of patients and their families. Drug-refractory seizures are a hallmark and treatment is further complicated by its multiple morbidities, which evolve over the patient’s lifetime. This review provides a comprehensive overview of the current and future options for the treatment of seizures associated with LGS. Six treatments are specifically indicated as adjunct therapies for the treatment of seizures associated with LGS in the US: lamotrigine, clobazam, rufinamide, topiramate, felbamate and most recently cannabidiol. These therapies have demonstrated reductions in drop seizures in 15%–68% of patients across trials, with responder rates (≥ 50% reduction in drop seizures) of 37%–78%. Valproate is still the preferred first-line treatment, generally in combination with lamotrigine or clobazam. Other treatments frequently used off-label include the broad spectrum anti-epileptic drugs (AED) levetiracetam, zonisamide and perampanel, while recent evidence from observational studies has indicated that a newer AED, the levetiracetam analogue brivaracetam, may be effective and well tolerated in LGS patients. Other treatments in clinical development include fenfluramine in late phase III, perampanel, soticlestat–OV953/TAK-953, carisbamate and ganaxolone. Non-pharmacologic interventions include the ketogenic diet, vagus nerve stimulation and surgical interventions; these are also expanding, with the potential for less invasive techniques for corpus callosotomy that have promise for reducing complications. However, despite these advancements, patients continue to experience a significant burden. Because LGS is not a single entity, tailoring of treatment is needed as opposed to a ‘one size fits all’ approach. Further research is needed into the underlying aetiologies and pathophysiology of LGS, together with advancements in treatments that encompass the spectrum of seizures associated with this complex syndrome.

1 Introduction

Lennox-Gastaut syndrome (LGS) is a childhood-onset severe developmental and epileptic encephalopathy (DEE), associated with high morbidity and profound effects on the quality of life (QoL) of patients and their families [1, 2]. LGS is a rare disease; it has an estimated incidence rate of 0.1 to 0.28 per 100,000 overall, with an incidence rate of 2 per 100,000 in children, accounting for approximately 2–5% of all childhood epilepsies, with a slight male preponderance [3].

The onset of LGS typically occurs by 12 years of age, with a peak between 3 and 5 years [1]; late-onset cases have been reported, albeit rarely [4]. A universally agreed-upon definition of LGS does not yet exist, and it has a wide variety of clinical presentations that continue to change and evolve over time [2, 5, 6]. However, it is traditionally defined as having a ‘triad’ of features encompassing (i) epilepsy with multiple pharmaco-resistant seizure semiologies, (ii) a generalised spike wave discharge pattern on EEG (Fig. 1) and (iii) cognitive and behavioural impairments (Fig. 2) [1, 2, 5, 6]. Tonic seizures are a hallmark of LGS, while other common seizure types include atypical absences and tonic or
Lennox-Gastaut syndrome (LGS) is a complex syndrome that is challenging to treat.

LGS is an entity with a range of underlying causes that is characterised by multiple seizure types that are drug-refractory (with tonic seizures being a hallmark feature), an abnormal EEG with generalised slow spike and wave discharges and cognitive impairment.

There are a range of pharmacological and non-pharmacological options for the treatment of seizures associated with LGS, which differ regarding mechanism of action, safety and tolerability.

Valproate in combination with the licensed pharmacotherapies lamotrigine or clobazam are a mainstay of treatment; other licensed adjunct therapies include rufinamide, topiramate, felbamate and most recently cannabidiol.

A personalised approach, tailored to the individual symptoms and responses of the patient during all stages of care, with regular assessment of treatment options, is particularly important for LGS.

Atonic drop attacks; other less common types include clonic, myoclonic and generalised tonic–clonic seizures. The characteristic EEG abnormalities include interictal slow spike waves at < 3 Hz, occurring while awake, typically in series and often evolving into subclinical seizure patterns, and paroxysmal fast rhythms (10–20 Hz) during sleep (Fig. 1) [2, 6]. Most patients with LGS experience progressive cognitive impairment, marked by developmental delay, intellectual disability and other behavioural problems such as hyperactivity, aggression and autistic behaviours [2, 6].

Early and accurate diagnosis is crucial for effective management [5, 6], which is especially important given the compelling data in other DEEs that early treatment is associated with improved outcomes [7]. However, early diagnosis is significantly hampered by a number of challenges including a lack of agreed-upon clinical criteria and assessment measures, absence of specific biological markers and the diverse and evolving manifestations of the syndrome, which may have overlapping clinical features with other DEEs (Fig. 2) [2, 5, 6]. With regard to the latter points, it can take time for the characteristic symptoms to appear—often a year or more. Furthermore, because tonic seizures can be absent at the initial stages, differential diagnosis from other DEEs may be difficult, especially with epilepsies with myoclonic–tonic–atonic/astatic seizures. An additional impediment to early diagnosis is that there is no single underlying cause of LGS, and indeed in 25–33% of cases the cause is unknown. The known aetiologies are varied and include the result of a brain abnormality (an ischaemic or haemorrhagic stroke, developmental malformation [e.g. cortical dysplasia], congenital infections, central nervous system [CNS] infections [e.g. meningitis], or tumours) and development from other severe infantile encephalopathies such as West syndrome (infantile spasms) (Fig. 2) [2, 8]. In addition, more rarely, LGS may be associated with the rare genetic disorder tuberous sclerosis complex (TSC), hereditary metabolic disorders, as well as other genetic disorders, with possible, albeit poorly defined, relationships with Down’s syndrome and Miller–Dieker syndrome (Fig. 2) [2, 8, 9]. Indeed, LGS may represent the end-stage of various epilepsies.

LGS imposes a substantial burden on people living with the condition and their families [10–12]. Atonic seizures are common in LGS; they are also known as ‘drop attacks’ or ‘drop seizures’ because they are characterised by a sudden
The triad of symptoms characteristic of Lennox-Gastaut syndrome (LGS), diagnostic challenges and aetiology. The characteristic LGS EEG pattern with slow SSW (Fig. 1) is a key diagnostic criteria. In contrast, cognitive impairment (intellectual disability and associated behavioural problems) is not always present at the outset of LGS and therefore this part of the “triad” is not always included in the diagnostic criteria. Brain damage can be the result of hypoxia at birth or head injuries, among others. DEEs developmental and epileptic encephalopathies, EEG electroencephalogram, NCSE non-convulsive status epilepticus, SSW slow spike wave, TSC tuberous sclerosis complex.
Though it is not specifically licensed for LGS, one uncontrolled, observational study involving 336 total patients with epilepsy, 38 of whom had LGS, showed that 21% of the LGS patients experienced total seizure control on valproate [29]. Another 55% showed a ≥ 50% improvement in seizure frequency, while only mild adverse events (AEs) were reported. Valproate proved particularly effective against drop attacks, atypical absences and myoclonic seizures, similar to what was reported in a separate study in patients with epilepsy [30]. Safety considerations to be aware of and notable drug–drug interactions are shown in Table 1; of note, valproate can result in hepatotoxicity and pancreatitis, resulting in a black box warning in the US label, and due to its teratogenicity, contraception must be ensured in females of childbearing potential [23, 24, 31]. However, overall, valproate is the mainstay treatment in the first-line setting (Fig. 3), underpinned by an abundance of physician experience, cost effectiveness and its positive mood-stabilising effects [17].

2.1.2 Clobazam

Clobazam, a 1,5-benzodiazepine compound that has a better safety and tolerability profile compared with other benzodiazepines [32], acts by binding to postsynaptic GABA receptors to increase the action potential threshold and reduce the frequency of action potentials, therefore decreasing the likelihood of seizures [24]. Although clobazam has been available as an AED for decades, it has only recently been approved in the US specifically for LGS, while it is approved as adjunctive therapy in epilepsy in the European Union (EU) (Table 1). After a phase II clinical trial showed that clobazam was well tolerated and reduced the weekly rates of drop and non-drop seizures in LGS patients [33], a phase III, double-blind, placebo-controlled, multicentre clinical trial was conducted to further evaluate the safety and efficacy of clobazam to treat LGS [34]. During this trial, patients were given a low (0.25 mg/kg/day), medium (0.5 mg/kg/day), or high (1 mg/kg/day) dose of clobazam for 12 weeks, with the option to continue treatment after this point. The average weekly rate of total seizures and drop seizures alone decreased in a dose-dependent manner, while responder rates increased with the dosage (Table 2; Fig. 4). Along with these promising results, all clobazam dosages resulted in an improvement of symptoms according to physicians’ and caregivers’ global evaluations (Table 2). There were no new safety signals during this trial, though nine developed pneumonia as a serious AE (SAE). Following these promising results, 267 of the patients involved in the phase II and phase III
trials continued adjunctive treatment with clobazam in a multicentre, open-label extension (OLE) study [35]. Over the course of 2–6 years, 16% of patients showed seizure control with clobazam monotherapy, while the decrease in the weekly rate of total seizures and drop seizures alone was maintained to year 5 in 85% and 85–91% of patients, respectively. The percentage of responders to clobazam was consistent with the original trials, and there was no indication of tolerance as the mean modal clobazam dose did not increase with time. After year 3 of the study, 80% of patients’ caregivers considered symptoms “very much improved” or “much improved”, suggesting a considerable improvement in QoL. Although 60% of patients experienced an AE, most were mild or moderate, with only 5% of patients reporting treatment-related SAEs. One patient had a convulsion with a fatal outcome during this study that was considered by the investigator as “possibly related” to clobazam treatment. Overall, clobazam showed an ability to elicit seizure freedom or improvement in patients treated over several years. However, some studies have shown that approximately one-third of patients may develop tolerance, while both cognitive and behavioural AEs have been associated with the drug [2, 24, 36]. In addition, physical and/or psychic dependence may develop, especially during prolonged use (Table 1). Despite these issues, clobazam is a useful combination therapy, with few reported harmful drug–drug interactions (Table 2), and it is considered a first-line treatment option (Fig. 3).

Table 1 Summary of pharmacotherapies widely used, licensed or upcoming for Lennox-Gastaut syndrome (LGS)

| Indication for LGS | Indication for seizuresa | Safety considerations | Key drug-drug interactions with other AEDs |
|--------------------|--------------------------|----------------------|----------------------------------------|
| US     | EU     | US     | EU     | US     | EU     |
| VPA    | x      | x      | ✓      | ✓      | ✓      | ✓      |
| LTG    | ✓      | ✓      | ✓      | ✓      | ✓      | ✓      |
| CLB    | ✓      | x      | ✓      | ✓      | ✓      | ✓      |
| RUF    | ✓      | ✓      | ✓      | ✓      | ✓      | ✓      |
| CBDa   | ✓      | ✓      | ✓      | ✓      | ✓      | ✓      |
| TPM    | ✓      | ✓      | ✓      | ✓      | ✓      | ✓      |
| FFAb   | x      | x      | ✓      | ✓      | ✓      | ✓      |
| FLB    | ✓      | x      | ✓      | ✓      | ✓      | ✓      |

The colours correspond to the colours used for the equivalent pharmacotherapies in Fig. 4

a Associated with epilepsy and/or other DEEs
b Approved for the treatment of seizures associated with DS, LGS and TSC in patients aged ≥1 year
c Approved for the treatment of seizures associated with DS in patients aged ≥2 years in the US and in the EU
d In phase III development for LGS

AE adverse event, AED antiepileptic drug, CBD cannabidiol, CLB clobazam, DEEs developmental and epileptic encephalopathies, DS Dravet syndrome, FFA fenfluramine, FLB felbamate, LTG lamotrigine, PK pharmacokinetic, RUF rufinamide, STP stiripentol, TPM topiramate, TSC tuberous sclerosis complex, VPA valproate.
### Table 2  Efficacy of pharmacotherapies from RCTs in Lennox-Gastaut syndrome (LGS)

| Study design | Seizures: drop attacks | Seizures: total | QoL/behaviour |
|--------------|------------------------|-----------------|---------------|
|              | Median percent reduction vs PBO | 50% responder rate | Median percent reduction vs PBO | 50% responder rate |  |
| **Licensed pharmacotherapies** | | | | | |
| LTG (Motte et al. 1997) | Phase III PBO-controlled RCT (n = 169) | − 34 vs − 9; *p* = 0.01 | 37 vs 22; *p* = 0.04 | − 32 vs − 9; *p* = 0.002 | 33 vs 16; *p* = 0.01 |
| CLB CONTAIN trial (Ng et al. 2011) [34] | Phase III PBO-controlled RCT (n = 238) | Low*: − 41.2 vs − 12.1; *p* = 0.0120 | Medium: − 49.4 vs − 12.1; *p* = 0.0015 | High: − 68.3 vs − 12.1; *p* < 0.0001 | Low*: − 34.8 vs − 9.3; *p* = 0.0414 | Medium: − 45.3 vs − 9.3; *p* = 0.0044 | High: − 65.3 vs − 9.3; *p* < 0.0001 | Percentages of patients at least minimally improved: 71.2% to 80.7% (physicians’ assessments) and 79.2% to 81.6% (caregivers’ assessments) for CLB vs 47.3% (physicians’ assessments) and 45.5% (caregivers’ assessments) for placebo |
| RUF: Study 022 (Glauser et al., 2008) [67] | Phase III PBO-controlled RCT (n = 138 aged 4–37 years) | − 42.5 vs + 1.4; *p* < 0.0001 | 42.5 vs 16.7; *p* = 0.002 | − 32.7 vs − 11.7; *p* = 0.0015 | 31.1 vs 10.9; *p* = 0.0045 | Mean CBCL total problems score: difference in LS mean total score between treatment groups across time (across weeks 24, 56, 88 and 106): RUF vs any other AED: − 1.197 (95% CI − 7.6 to 5.3); *p* = 0.7083 |
| RUF: Study 303 (Arzimanoglou et al., 2016 and 2019) [68, 69] | Phase III, PBO-controlled RCT (n = 37 paediatric patients aged ≥1 to <4 years) | Mean total score between treatment groups across time (across weeks 24, 56, 88 and 106): RUF vs any other AED: − 1.197 (95% CI − 7.6 to 5.3); *p* = 0.7083 |  |  |  |  |  |  |
Expanding the Treatment Landscape for Lennox-Gastaut Syndrome: Current and Future Strategies

AED antiepileptic drug, CBCL Child Behaviour Checklist, CBD cannabidiol, CGI Clinical Global Impressions scale, CGI-I Clinical Global Impression of Improvement. CI confidence interval, CLB clobazam, FFA fenfluramine, FLB felbamate, LS Likert scale, LTG lamotrigine, PBO placebo, QoL quality of life, RCT randomised controlled trial, RUF rufinamide, TPM topiramate

Table 2 (continued)

| Seizures: drop attacks | Seizures: total | QoL/behaviour |
|------------------------|----------------|---------------|
| Median percent reduction vs PBO | Median percent reduction vs PBO | CGI (patient/caregiver) proportion of patients improved: |

CBD GWPCARE3 (Devinsky et al., 2018) [86]  
Study design: Phase III, PBO-controlled RCT  
CBD10: 37.2 vs 17.2; p = 0.002  
CBD20: 41.9 vs 17.2; p = 0.005  
Subgroup with CLB:  
CBD10: 45.6 vs 22.7; p = 0.0355  
CBD20: 64.3 vs 22.7; p < 0.0001

CBD GWPCARE4 (Thiele et al., 2018) [87]  
Study design: Phase III, PBO-controlled RCT  
CBD20: 43.9 vs 21.8; p = 0.0135  
Subgroup with CLB:  
CBD10: 40.5 vs 21.6; p = 0.0584  
CBD20: 55.6 vs 21.6; p = 0.0021

TPM (Sachdeo et al., 1999) [45]  
Study design: PBO-controlled RCT (n = 98)  
− 14.8 vs + 5.1; p = 0.041  
28 vs 14; p = 0.071  
− 25.8 vs + 5.2; p = 0.015  
33 vs 8; p = 0.002

FLB (Felbamate Study Group in LGS, 1993) [53]  
Study design: PBO-controlled RCT (n = 73)  
− 19 vs + 4; p = 0.002  
− 34 vs − 9%; p = 0.01

Late clinical development

FFA Study 1601 (Zogenix 2020) [105]  
Study design: Phase III, PBO-controlled RCT  
FFA0.7: 26.5 vs 7.8; p = 0.0012  
FFA0.7: 25.3 vs 10.3; p = 0.0165

CGI-I proportion of patients improved: 44.8% vs 33.8%; p = 0.0567

CGI-I proportion of patients much improved or very much improved: 26.3% vs 6.3%; p = 0.0007

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AED antiepileptic drug, CBCL Child Behaviour Checklist, CBD cannabidiol, CGI Clinical Global Impressions scale, CGI-I Clinical Global Impression of Improvement, CI confidence interval, CLB clobazam, FFA fenfluramine, FLB felbamate, LS Likert scale, LTG lamotrigine, PBO placebo, QoL quality of life, RCT randomised controlled trial, RUF rufinamide, TPM topiramate

aLow dose = CLB 0.25 mg/kg/day; medium dose = CLB 0.5 mg/kg/day; high dose = CLB 1 mg/kg/day

bValues are mean

cPhysicians’ and caregivers’ global evaluations of the patients’ overall changes in symptoms over time (using a 7-point Likert scale, with 1 = very much improved and 7 = very much worse)

dGlobal evaluations were made on the caregivers’ impressions of QoL with respect to alertness, verbal responsiveness, general wellbeing and seizure control

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2.2 Second-Generation Antiseizure Treatment Licensed for LGS

2.2.1 Lamotrigine

Lamotrigine’s mechanism of action is not fully understood; however, it is thought that the drug selectively stabilises voltage-sensitive sodium channels, inhibiting the release of excitatory neurotransmitters [37, 38]. It is possible that lamotrigine acts to reduce the number of spike-and-wave events that are typical of LGS [39, 40], and the drug has been shown to be effective at treating refractory epilepsy when used as an adjunctive therapy [41]. These early findings prompted a double-blind, placebo-controlled, randomised controlled trial (RCT) to determine the efficacy of lamotrigine against LGS, involving 169 patients, which led to the approval of lamotrigine specifically for LGS in the US and EU [37]. The lamotrigine-treated group were titrated up to 100–200 mg/day if on concomitant valproate or 300–400 mg/day if not, resulting in a 32% median reduction in all major seizures, compared with just 9% in the placebo group (Table 2; Fig. 4). Similar results were seen with tonic-clonic seizure or drop attack frequency alone (Table 2; Fig. 4). Of note, valproate increases lamotrigine concentrations more than two-fold, and there have also been suggestions that these two therapies may work synergistically to provide enhanced seizure control over each drug independently [42].

The most common AEs of lamotrigine seen in the study were pharyngitis and fever, common childhood ailments, and only three patients prematurely discontinued treatment due to AEs. However, lamotrigine is associated with the development of rash, common particularly in younger patients [31], which can progress, albeit rarely, into the serious and occasionally fatal Stevens-Johnson syndrome (Table 1) [43]. This risk can be reduced by the mandatory slow titration of lamotrigine (combination therapy starting dosage 0.3 mg/kg/day), which is even slower with valproate (starting dosage 0.15 mg/kg/day) due to the enzyme inhibition of valproate and consequent increase in lamotrigine levels (Table 1). With these precautions, lamotrigine in combination with valproate remains a mainstay of treatment (Fig. 3).

2.2.2 Topiramate

Topiramate, a fructose derivative, is a broad-spectrum AED able to prevent the onset of multiple seizure types [44]. Topiramate is specifically approved for use in LGS patients (Table 1). Though it is not known which of its mechanisms of action is most important for its anticonvulsant effects,
topiramate is known to modulate voltage-gated sodium channels, reduce neuronal excitation by mediating N-methyl-D-aspartate receptors, enhance inhibition of GABA-mediated neurotransmission, inhibit carbonic anhydrase, and modulate voltage-gated calcium channels [27, 44].

The efficacy of topiramate as an adjunctive therapy in LGS patients has been demonstrated in a double-blind, placebo-controlled RCT involving 98 patients (Table 2; Fig. 4) [45]. Patients were given up to 6 mg/kg/day (titrated up from 1 mg/kg/day) of topiramate for 11 weeks, while seizure frequency and severity were monitored. Topiramate significantly reduced the average monthly rate of both drop attacks (defined as tonic and atonic seizures in this study) and major motor seizures by 14.8% and 25.8%, respectively, compared with a 5.1% and 5.2% increase in drop attacks and major motor seizures in the placebo-treated group (Table 2; Fig. 4). Importantly, there was a ≥ 50% reduction in major motor seizure frequency from baseline frequency in one-third of topiramate-treated patients, compared with just 8% of the placebo-treated patients (Table 2; Fig. 4). Along with the reduction in frequency, topiramate also reduced seizure severity, with topiramate-treated patients approximately twice as likely to show an improvement, while one patient became seizure free. The AEs reported during this trial were mostly mild or moderate; however, 23 patients on topiramate experienced severe AEs that were most often CNS-related. Overall, this trial demonstrated topiramate to be a well-tolerated add-on treatment for LGS and the results of this study have since been mirrored in open-label studies, showing long-term reduction in seizure frequency in patients with LGS [46, 47]. The American Academy of Neurology (AAN) and the American Epilepsy Society (AES) support the use of topiramate in conjunction with lamotrigine, as this combination seems to be particularly effective at reducing tonic/ataonic seizures in LGS [48]. However, it should be noted that topiramate is associated with cognitive side effects including mental slowing and dysphasia [49], which may lead to poor retention rates [50].

### 2.2.3 Felbamate

Although felbamate is licensed in the US (Table 1), it is not commonly used as it carries a black box warning owing to the risk of aplastic anaemia and hepatic failure, two life-threatening AEs [51]. Due to this risk, it has not been approved by the European Medicines Agency (EMA) for any indication, although it may be available in some countries on a patient-by-patient basis [2]. The mechanism of action of the carbamate-type anticonvulsant is also not well understood. Early seizure models indicated that felbamate increases seizure threshold and prevents the spread of seizures in the brain [52], which may make the drug particularly effective against generalised tonic-clonic or partial seizures [53]. More recent studies suggest the drug primarily operates by reducing glutamatergic transmission, though it may also inhibit GABA-receptor binding along with voltage-gated sodium and calcium channels [54].

The approval of felbamate in the US is based on a double-blind, placebo-controlled RCT involving 73 LGS patients (Table 2) [53]. Felbamate, which was titrated up over 14 days from 15 mg/kg/day to a maximum of 45 mg/kg/day or 3600 mg/day, was associated with a 34% reduction in the frequency of atonic seizures, with five patients experiencing no atonic seizures while receiving their maximum dosage, compared with 9% in the placebo group (Table 2; Fig. 4). This decrease of atonic seizure counts showed a linear correlation with an increase in felbamate plasma concentrations as dosage increased. The drug was also effective against other seizure types with the frequency of total seizures reduced by 19% and four patients becoming seizure free in the felbamate-treated group, compared with a 4% increase in the placebo group (Table 2; Fig. 4). Importantly, evaluations revealed a significant increase in QoL for the patients taking felbamate compared with the placebo group, based on caregivers’ impressions (Table 2). AEs were primarily mild or moderate, although the study had a small sample size and a relatively short 10-week period of treatment.

Seventy of the patients who completed this trial continued in an open-label follow-up study, where those who were originally treated with placebo showed similar levels of improvement as the felbamate-treated patients [55]. During the original study, only five of 36 patients in the placebo-treated group showed a ≥ 50% reduction in total seizure frequency, which increased to 21 patients after a month on felbamate. Only two of 22 patients in the placebo-treated group showed a > 50% reduction in atonic seizure frequency during the original trial [53], which then increased to 12 when these patients were treated with felbamate during the open-label study. In the follow-up study, control of atonic seizure frequency was maintained at 12 months, and 51% of patients maintained a ≥ 50% reduction in total seizure frequency at this timepoint, suggesting relative long-term efficacy and a lack of tolerance. Due to the previously mentioned risk for felbamate-associated fatal AEs, it is generally recommended as a last resort for patients with highly refractory epilepsy (Fig. 3) [56]. However, despite the black-box warning, several retrospective real-world clinical studies have found felbamate to be safe in the context of close monitoring of hepatic and haematologic functions, confirming that felbamate does have a place in the LGS treatment pathway [57–59].
Steroids, including prednisone, prednisolone, methylprednisolone, adrenocorticotropic hormone (ACTH) and hydrocortisone also have an important role during periods of heightened seizure activity. While the efficacy of steroids for infantile spasms has been established from RCTs [60], evidence for the treatment of other types of seizures is more limited, generally from observational studies [61]. In studies evaluating steroids in patients with refractory epilepsies other than infantile spasms that have included at least a few patients with LGS, responder rates of 30%–79% have been reported [62, 63]. However, the well documented side effects of prolonged steroid use, including hypertriglyceridaemia, osteoporosis and suppression of childhood growth means that their use should be curtailed to short-term ‘crisis’ periods [2].

### 2.3 Third-Generation Antiseizure Treatment Licensed for LGS: Rufinamide

Rufinamide is a triazole derivative structurally unrelated to other AEDs. The best characterised mechanism of action of rufinamide is the modulation of the activity of sodium channels, prolonging their inactive state, although it may have additional mechanisms through which it exerts its therapeutic effects [64–66]. Rufinamide failed to show efficacy in clinical studies in a general epilepsy population, while efficacy was observed in LGS patients. As such, rufinamide was approved in the EU in 2007 and in the US in 2008 as adjunctive therapy for the treatment of seizures associated with LGS in children and adults aged 4 years and older, which was expanded to include patients 1 year of age and older in 2015 in the US and in 2018 in the EU [65, 66].

#### 2.3.1 Pivotal Clinical Trials

The FDA and EMA approvals in LGS were based on data from a phase III, double-blind, placebo-controlled RCT in patients with LGS aged 4–37 years (Study 022) [67], with the expanded indication from a phase III, open-label RCT in paediatric patients (≥ 1 to < 4 years old) (Study 303; ClinicalTrials.gov identifier NCT01405053) [68, 69].

Study 022 and OLE (Study 022E), conducted in 138 patients (74 in the rufinamide group and 64 in the placebo group), consisted of a 4-week baseline period, a 12-week, double-blind treatment period, a 2-week, double-blind conversion phase, followed by an open-label single-arm extension period of up to 3 years [67, 70]. Compared with placebo at 12 weeks, the median percentage reduction in both total seizure frequency and drop seizures was significantly greater in the rufinamide group compared with the placebo group (Table 2; Fig. 4). The rufinamide group had a larger improvement in seizure severity and a significantly higher 50% responder rate compared with placebo (Table 2; Fig. 4) [67]. The results are also supported by a phase III trial in Japan in patients aged 4–30 years, which showed that the frequency of epileptic seizures was significantly decreased in the rufinamide group compared with the placebo group; the median percent change in frequency of tonic-atonic seizures was −24.2% and −3.3%, respectively, \( p = 0.003 \) and that of total seizures was −32.9% and −3.1%, respectively \( p < 0.001 \) [71].

In Study 303, 37 paediatric patients randomised to rufinamide \( n = 25 \) or any other AED \( n = 12 \) were evaluated across a 106-week treatment phase for safety/tolerability and behavioural effects assessed via the Child Behaviour Checklist (CBCL) [68, 69]. Overall, behavioural outcomes were comparable between the rufinamide and ‘any other AED’ groups, although the small sample size and difficulties assessing behaviour are important caveats that should be acknowledged (Table 2) [68]. In addition, in a post-hoc analysis, the mean number of seizure-free days was 42.2% greater post-baseline compared with baseline in patients treated with rufinamide \( p < 0.0001 \), with only one rufinamide patient experiencing a decrease in the number of seizure-free days post-baseline [72]. The median time to reach the baseline number of seizures increased by 10.5 days for rufinamide and 0.5 days for the ‘any other AED’ group during the treatment phase, to 46.0 and 54.0 days, respectively.

#### 2.3.2 Longer-Term Data

In Study 022E, 124 of the original 139 patients from Study 022 entered the extension study; patients were treated with rufinamide for a median (range) of 432 (10–1149) days [70]. Efficacy was sustained for both seizure frequency reduction and responder rates, with reductions in seizure frequency observed throughout the study; during the last 12 months of treatment, 41.0% and 47.9% of patients had ≥ 50% reduction in total and tonic–atonic seizure frequency, respectively. In addition, in another OLE, this time in Japanese patients with LGS who participated in the RCT, reduction of seizure frequency was maintained to 52 weeks [73].

### 2.3.3 Safety

Rufinamide is a generally well tolerated treatment. In Study 022, the most common AEs were somnolence (24.3% with rufinamide vs 12.5% with placebo) and vomiting (21.6% vs 6.3%), and in Study 022E they were vomiting (30.6%) and pyrexia (25.8%) [67, 70]. In Study 022/022E, three patients in the rufinamide group experienced status epilepticus compared with none in the placebo group. In the paediatric population of patients aged ≥ 1 to < 4 years (Study 303), the frequency, type and severity of AEs were similar to that in...
children 4 years of age and older, adolescents and adults; the most frequently reported AEs in the rufinamide treatment group (occurring in ≥10% of subjects) were upper respiratory tract infection and vomiting (28% each), pneumonia and somnolence (20% each), diarrhoea, pyrexia, cough, sinusitis and otitis media (16% each), and rash, irritability, decreased appetite, constipation, bronchitis and nasal congestion (12% each) [68].

2.3.4 Additional Prospective and Retrospective Clinical Studies

Further to the pivotal phase III studies, the effectiveness and safety/tolerability of rufinamide in treating seizures associated with LGS has been supported by other clinical trials and several ‘real-world’ clinical practice studies; responder rates (≥ 50% reduction in seizures) ranging from 33.3 to 60.5% have been reported across the studies in LGS patients, and no new safety concerns have been identified [74–79]. The additional studies have been recently comprehensively reviewed by Balagura et al. [80] (2020) and by Striano et al. (2018) [81].

2.3.5 Other Considerations: Drug–Drug Interactions

The presence of possible interactions with other AEDs and other medications are also important aspects to be taken into consideration. Significant increases in rufinamide plasma concentrations may occur with concomitant valproate, and consequently a dose reduction of rufinamide may be required (Table 1) [65, 66]. In contrast, no significant changes in rufinamide concentration have been observed following co-administration with lamotrigine, topiramate or benzodiazepines. In addition, rufinamide appears not to have a clinically relevant effect on carbamazepine, lamotrigine, phenobarbital, topiramate, phenytoin or valproate steady-state concentrations [65, 66].

2.4 Recently Approved Pharmacologic Treatment for LGS: Cannabidiol

The full mechanisms of action by which cannabidiol exerts its anti-seizure effects are still being explored, although it is known that it possesses affinity for diverse targets that decrease the neuronal excitability related to the pathogenesis of the disease; cannabidiol targets include the G protein-coupled receptor 55 (GPR55) and transient receptor potential vanilloid 1 (TRPV-1) channels that modulate intracellular calcium, and the equilibrative nucleoside transporter 1 (ENT-1) involved in adenosine-mediated signalling [82, 83]. The anticonvulsant effect does not work via cannabinoid receptors, and this lack of appreciable affinity or activity at these receptors means that it is not associated with the psychoactivity of the archetypal cannabinoid, tetrahydrocannabinol [84].

Cannabidiol has been approved as adjunctive therapy for seizures associated with LGS or Dravet syndrome (DS) for patients 2 years of age and older since 2018 in the US and 2019 in the EU [83, 85]; in the US the FDA has recently expanded the indication to include patients 1 year of age and older, as well as TSC patients (Table 1). In the EU it is approved in conjunction with clobazam because in the key pivotal trial this combination resulted in greater efficacy than in the subgroup of patients not taking clobazam (Table 2; Fig. 4) [83]. The recommended maintenance dose of the oral solution is 10 mg/kg/day, which can be increased to 20 mg/kg/day.

2.4.1 Pivotal Clinical Trials

The approvals from the FDA and EMA for LGS were informed by data from two phase III, double-blind, placebo-controlled, parallel-group RCTs. GWPCARE3 [86] and GWPCARE4 [87], which both consisted of a 14-week treatment period (2-week titration period and 12-week maintenance period), were conducted in LGS patients (aged 2–55 years) who were inadequately managed on at least two AEDs. In GWPCARE3, patients (n = 255) were randomised to receive cannabidiol at 20 mg/kg of body weight (CBD20 group; n = 76) or 10 mg/kg (CBD10 group; n = 73) or matching placebo (n = 76). In GWPCARE4, 171 patients were randomised to receive cannabidiol at 20 mg/kg (n = 86) or matching placebo (n = 85). Across both trials, commonly used conventional AEDs that were used in >25% of patients were valproate, clobazam, lamotrigine, levetiracetam and rufinamide, with approximately 50% of the patients taking concomitant clobazam.

Across the trials, cannabidiol was associated with statistically significant and clinically meaningful improvements in seizure frequency compared with placebo (Table 2; Fig. 4). Compared with placebo, cannabidiol was associated with a significantly higher percentage reduction in the monthly frequency of drop seizures during the 14-week treatment period in both trials (Table 2; Fig. 4). Furthermore, significantly higher percentages of patients in the cannabidiol groups achieved ≥50% reductions in the monthly frequency of drop seizures than in the placebo group (Table 2; Fig. 4). A few patients in the cannabidiol groups were drop-seizure free during the entire 12-week maintenance period, although not during the whole 14-week period that included the titration period, suggesting that cannabidiol might have a delayed effect on drop attacks. In addition, cannabidiol may have efficacy over a range of seizures types, with significant reductions in the median frequency for total seizures and non-drop seizures. In addition, cannabidiol treatment led to an increase in the number of drop seizure-free days of
concomitant use of clobazam, with this AE generally being resolved by reductions in the dose of cannabidiol and/or concomitant valproate [83, 85]. In addition, the incidence of somnolence and sedation has also been shown to be dose-related, and is more common in patients on concomitant clobazam [83, 85].

2.4.4 Other Considerations: Drug–Drug Interactions

As discussed above, in the EU, cannabidiol is only indicated in conjunction with clobazam due to the increased efficacy observed in this subgroup of patients (Table 2; Fig. 4). A bidirectional drug–drug interaction occurs with the combination of cannabidiol with clobazam leading to increased levels of active metabolites of both compounds. While there has been speculation that the efficacy of cannabidiol may be due solely to an increase in the plasma concentration of clobazam, recent meta-analyses of trials in patients with DS and LGS have demonstrated that cannabidiol has benefits beyond the interaction with clobazam; although improvement in seizure control was indeed greatest in patients with cannabidiol and concomitant clobazam, efficacy was also improved compared with placebo in patients with other concomitant medications [90–92].

Increases in serum levels of topiramate and rufinamide (in adults and children), and zonisamide and eslicarbazepine (in adults) have been reported with increasing doses of cannabidiol, but were within the accepted therapeutic range [85]. In addition, a case series of five patients reported a marked increase in brivaracetam levels of 95% to 280%, although only two patients reported AEs, which were mild, but resulted in a dose reduction of brivaracetam in one patient [93]. As discussed in the safety section, a clinically and statistically significant interaction with clobazam and its active metabolite N-desmethyleclobazam has been reported, resulting in increased sedation [83, 85, 94, 95] (Table 1).

In addition, an interaction has been noted in patients taking concomitant valproate and cannabidiol with regard to elevated liver function test results (Table 1) [83, 85, 94]. Also of potential interest is a case report of a possible interaction with the oral anticoagulant warfarin, requiring a reduction in the dosage of warfarin to prevent potential bleeding complications [96]. Other considerations related to drug–drug interactions include reducing the cannabidiol dose when coadministered with a moderate or strong inhibitor of CYP3A4 or CYP2C19, and increasing the dose when coadministered with a strong CYP3A4 or CYP2C19 inducer [83, 85]. In addition, dose reductions of substrates of UGT1A9, UGT2B7, CYP2C8, CYP2C9, and CYP2C19 (e.g., clobazam) should be considered, and substrates of CYP1A2 and CYP2B6 may also require dose adjustment [83, 85].

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2.5 Upcoming Pharmacologic Treatments: Agents in Phase III Clinical Development

2.5.1 Fenfluramine

Fenfluramine is a derivative of amphetamine that was initially developed as an appetite suppressant [97]. Fenfluramine acts via multiple receptors to exert its therapeutic effects for the treatment of seizures, although the exact mechanisms are still being elucidated. Fenfluramine increases extracellular levels of serotonin (5-HT) through interaction with serotonin transporter proteins, and exhibits agonist activity at serotonin 5HT-2 receptors (5HT1D and 5HT2C) [97–100].

In addition, at the in vitro and in vivo level, fenfluramine activity at 5-HT receptors is complemented synergistically by functional activation of sigma-1 receptors, resulting in improved cognitive functions of spatial and contextual learning via activity at sigma-1 receptors in mouse models [101]. Fenfluramine was previously prescribed as an anorexigen, but it was withdrawn in 1997 based on reports of increased risks of cardiac valvulopathy and pulmonary hypertension in adult patients treated for obesity; however, the fenfluramine doses were much higher compared with a licensed maximum dose for DS, and it was frequently given off-label with phentermine, which itself has an impact on valvular disease [102–104]. In contrast, the benefit/risk profile of fenfluramine given at low dose for the treatment of seizures appears to be positive, and fenfluramine has been approved by the FDA in the US for the treatment of seizures associated with DS in patients age 2 years and older [98]. In December 2020 fenfluramine was approved by the EMA in the EU for the treatment of DS.

2.5.1.1 Pivotal Clinical Trial

Fenfluramine is currently in late-stage phase III clinical development as a therapy for the treatment of uncontrolled seizures in children and adults with LGS. The phase III, multicentre, global, double-blind, placebo-controlled study (Study 1601 [NCT03355209]) enrolled 263 patients with LGS aged 2–35 years whose seizures were currently uncontrolled despite treatment with one or more AED(s) [105]. Patients were randomised into three treatment groups: fenfluramine 0.7 mg/kg/day (26 mg maximum daily dose; n = 87; FFA0.7), fenfluramine 0.2 mg/kg/day (n = 89; FFA0.2) and placebo (n = 87). With a median age of 13 years, 29% of patients were 18 years or older. Patients had had an average of seven prior AEDs and were taking between one and four AEDs at baseline. The median baseline drop seizure frequency across the study groups was 77 seizures per month. During Part 1 of the trial, baseline seizure frequency was established during the first 4 weeks, after which there was a 14-week treatment period consisting of a 2-week titration period followed by a 12-week fixed-dose maintenance period. Patients who completed Part 1 could enrol in an ongoing 12-month OLE study (Part 2), which was conducted to establish the long-term safety, tolerability and effectiveness.

Compared with placebo, patients in the FFA0.7 group had a significant median percent reduction in the frequency of monthly drop seizures from baseline, thereby meeting the study’s primary endpoint (Table 2; Fig. 4). Using a parametric analysis, FFA0.7 was associated with a 26.5% greater reduction in mean monthly drop seizure frequency compared with placebo (p = 0.0034). On the other hand, the median percent reduction in monthly drop seizures for FFA0.2 did not reach statistical significance compared with placebo (13.2%; p = 0.0915). Clinically meaningful (≥ 50%) reductions in mean convulsive seizure frequency in monthly drop seizures were experienced in significantly more patients treated with FFA0.7 compared with placebo (Table 2; Fig. 4). In addition, using the Clinical Global Impression of Improvement (CGI-I) that assesses improvement or worsening relative to baseline as judged by the investigator, significantly more patients in the FFA0.7 group than the placebo group were considered to be much improved or very much improved (Table 2).

2.5.1.2 Longer-Term Data

The phase III study supports the results from an earlier, phase II, open-label, dose-finding study of add-on fenfluramine in patients with LGS (NCT02655198), which now has data for 15 months of treatment [99]. During the 20-week core study, there was a 53% median reduction (n = 13) in convulsive seizures, with eight patients (62%) having a ≥ 50% reduction in convulsive seizures, while at 15 months (long-term extension; n = 9), the median reduction in convulsive seizures was 58%, with six (67%) patients having a ≥ 50% reduction.

2.5.1.3 Safety

Fenfluramine was generally well tolerated, with no new safety signals to those observed in the two DS studies (reviewed by Strzelczyk and Schubert-Bast 2020 [106]). Overall, 89.7% experienced at least one AE in the FFA0.7 group, 76.4% in the FFA0.2 group and 79.3% in the placebo group. Decreased appetite, somnolence, fatigue, vomiting, diarrhoea and pyrexia were the most common AEs (≥10%) reported in the fenfluramine groups. SAEs occurred in 11.5% of patients (n = 10) in the FFA0.7 group, 4.5% (n = 4) in the FFA0.2 group and 4.6% (n = 4) in the placebo group. AEs leading to study discontinuation occurred rarely, in six patients in the FFA 0.7 group, four in the FFA0.2 group and one in the placebo group, although the majority were considered as treatment-related. One patient died during the trial (FFA0.7 group) due to SUDEP (sudden unexpected death in epilepsy) that was unrelated to the study drug as assessed by the investigator. A total of 247 (93.9%)
patients entered the OLE phase. Similarly, in the phase II study, the most common AEs included decreased appetite (31% \( n = 4 \)) and decreased alertness (15% \( n = 2 \)). No cases of valvular heart disease or pulmonary hypertension were observed in either the phase III trial (both Part 1 and Part 2) or in the phase II study.

2.5.1.4 Other Considerations: Drug–Drug Interactions No significant impact on the pharmacokinetics of clobazam, valproate and cannabidiol has been observed with fenfluramine [107, 108]. A combination of stiripentol with valproate + clobazam led to an increase in the dose plasma levels and systematic exposure of fenfluramine, requiring a reduction in the dose of fenfluramine (to a maximum dose of 0.4 mg/kg/day) when administered in combination with the stiripentol regimen, although this is more relevant to the treatment of DS [107, 109].

2.5.2 Perampanel

Perampanel is a selective, non-competitive antagonist of the ionotropic α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) glutamate receptor on post-synaptic neurons, although the precise mechanisms by which it exerts its antiepileptic effects are not yet fully understood [110, 111]. Perampanel is currently approved by the EMA and the FDA in patients with epilepsy aged 12 years and older as an adjunctive therapy for the treatment of partial-onset seizures with or without secondarily generalised seizures and for primary generalised tonic-clonic seizures [110, 111].

Perampanel is currently being investigated in a phase III, multicentre, double-blind, placebo-controlled RCT as adjunctive therapy in patients with inadequately controlled seizures associated with LGS (NCT02834793) [112]. With an estimated enrolment of 142 patients, and a primary endpoint of median percent change in drop seizure frequency per 28 days during double-blind treatment, the primary completion date is estimated to be in January 2022 [112].

While no results have been reported from the clinical trial to date, the efficacy and safety of perampanel in LGS patients has been demonstrated in retrospective, open-label, observational studies [113–118]. Response rates (≥ 50% seizure reduction) of 64.8% (46/71 adult patients) [113] and 69.2% (9/13 children/adolescents [117]) have been reported in LGS patients. In addition, in a large real-world study of 2396 individuals with treatment-resistant epilepsy who were treated with add-on perampanel, 9% were seizure free for ≥ 6 months, and the 1-year retention rate was 48% [118]. In 388 individuals with available data, responder rates were 42% at 3 months, 46% at 6 months and 39% at 12 months, and no new safety signals were observed. Further studies in mixed epileptic populations have also reported response rates of 46% (34/74 children/adolescents with various refractory epilepsies [5 with LGS] [116]) and 31% (18/58 difficult-to-treat epilepsy patients [4 with LGS] [114]). Perampanel was generally well tolerated, with typical AEs being dizziness, somnolence and fatigue [113–117]. In addition, improvements in cognitive function and/or behaviour were reported in a proportion of patients: 7/13 (53.8%) children/adolescents with LGS [117], and 4/71 (5.6%) adult patients with LGS [113], although conversely, perampanel can also be associated with psychiatric AEs including irritability and aggression that may be more marked in those with cognitive impairment [119]. Perampanel is already used in LGS patients off-label (Fig. 3), while positive data from the phase III study may lead to a specific indication for LGS.

2.6 Off-Label AEDs

Some other AEDs, including levetiracetam and zonisamide, may also be useful in treating multiple seizure types because of their broad spectrum in their modes of action. Zonisamide, which is approved by the FDA and EMA as an adjunctive therapy for the treatment of partial seizures in adults and children aged 6 years and above, is thought to act through various mechanisms, including blocking sodium and T-type calcium channels, resulting in increased dopamine and serotonin neurotransmission, as well as blocking glutamate release [120]. In a study of 62 children with LGS treated with zonisamide, three patients were seizure free and 51.6% of patients experienced a ≥ 50% reduction in seizure frequency [120]. In addition, among 55 children with LGS, levetiracetam was associated with a reduction in seizure frequency of ≥ 50% in 58.2%, with 15 patients (27.3%) being seizure free [121]. Levetiracetam in particular is a common choice owing to its generally good safety and tolerability profile as well as having few interactions with other medications, although it can be associated with psychobehavioural AEs, especially in children and in patients with cognitive impairment [122]. The therapeutic indications for levetiracetam in the US and EU include the treatment of partial-onset seizures, and it may exert its effects through various mechanisms, including affecting intraneuronal \( \text{Ca}^{2+} \) levels and/or by its interaction with synaptic vesicle protein 2A [122]. However, it should be noted that overall there is a paucity of evidence regarding zonisamide and levetiracetam in LGS, generally garnered from a few observational studies (reviewed in Cross et al. 2017 [2]), and robust evidence from RCTs is lacking.

There has also been recent evidence of a newer AED, brivaracetam, showing efficacy in patients with epileptic encephalopathies, including LGS [123–125]. Brivaracetam is a levetiracetam analogue that binds to the synaptic vesicle protein 2A, and inhibits excitatory neurotransmitters’ release [126, 127]. It was approved in the US and the EU in 2016 as adjunctive treatment for focal-onset seizures in patients over 16 years of age.
and in 2018 for children over 4 years of age. In a cohort study in Germany (n = 44, 20 [45.5] with LGS), a 50% responder rate was reported in 19 patients (43%); nine patients (20%) were seizure free for > 12 months, and two were seizure free for > 6 months [125]. AEs, observed in only 16% of patients, were predominantly of psychobehavioural nature; the rate was lower than had been reported in patients on levetiracetam at switch (32%), suggesting that brivaracetam may be a suitable alternative for those who experience psychobehavioural AEs associated with levetiracetam. Similarly, in a large population of patients with predominantly drug-resistant epilepsy (n = 575), brivaracetam was shown to be effective and well tolerated, with a ≥50% responder rate of 39.7%, and 17.5% being seizure free; tolerability was not highly affected in patients with learning disability or psychiatric comorbidity [124]. Finally, in another observational study that included 31 patients, of whom 11 had epileptic syndromes (5 with LGS), the responder rate was 45.2%, and eight patients had better response to seizures compared with levetiracetam. In addition, AEs were rare (mild somnolence [6.4%], psychosis [3.2%] and nausea [3.2%]) [123].

Other AEDs including phenobarbital, ethosuximide and bromides are also sometimes used, despite a lack of gold-standard evidence from RCTs [31]. Carbamazepine, oxcarbazepine, eslicarbazepine, tiagabine, phenytoin, vigabatrin, gabapentin and gabapentin are infrequently used, and there may be potential for aggravating certain seizure types with these AEDs [2, 17].

### 2.7 Pharmacologic Agents in Earlier Clinical Development

#### 2.7.1 Soticlestat–OV953/TAK-953

Soticlestat (TAK-935/OV935) is a novel, highly selective first-in-class inhibitor of the brain-specific enzyme cholesterol 24-hydroxylase (CH24H) that dose-dependently reduces plasma levels of 24S hydroxy cholesterol (24HC) [128]. 24HC is an endogenous positive allosteric modulator of NMDA receptors [128], which are a subtype of glutamate receptors that play important roles in excitatory neurotransmission and synaptic plasticity implicated in several neurological diseases, including epilepsy [129]. Soticlestat has shown anticonvulsant activity in animal seizure models, and a correlation between reduction in 24HC levels and reduced seizure frequency was observed in a phase Ib/IIa study [130].

Soticlestat is being evaluated in two phase II trials for DEEs including LGS—ELEKTRA and ENDYMION. The ELEKTRA trial (NCT03650452) is a phase II, multicentre, double-blind, placebo-controlled RCT designed to evaluate the efficacy, safety and tolerability of soticlestat in paediatric patients, aged 2–17 years old, with DS (n = 51) or LGS (n = 88) [131]. The top-line results have recently (late August 2020) been announced, reporting that the study met its primary endpoint, with a 27.8% median reduction from baseline in convulsive seizure (DS) and drop seizure (LGS) frequency compared with a 3.1% median increase in patients taking placebo during the 12-week maintenance period (median placebo-adjusted reduction: 30.5%; p = 0.0007, n = 120 patients with seizure data in the maintenance period) [132]. In the LGS cohort, soticlestat was associated with a 20.6% median reduction in drop seizure frequency versus 6.0% with placebo (median placebo-adjusted reduction in seizure frequency: 14.8%; p = 0.1279). In addition, 27.9% of LGS patients showed an improvement according to the investigator-assessed Global Impression of Change. Soticlestat was well tolerated, with the most frequent AEs with a ≥5% difference from placebo being lethargy and constipation.

The long-term safety and tolerability of soticlestat is also being evaluated in ENDYMION (NCT03635073), a phase II, multicentre, open-label, long-term extension study of soticlestat in adult patients (18–65 years) with DEEs who participated in the phase Ib/IIa study [133]. Following on from the phase Ib/IIa study which showed a 36% median reduction in seizure frequency from baseline to day 85 (n = 16) [134], initial results from ENDYMION have reported a median seizure frequency reduction of 46.4% following 13–24 weeks (n = 7), 82% following 25–36 weeks (n = 7), and 90% following 37–48 weeks (n = 4) of treatment. In addition, the longest seizure-free consecutive duration by two different participants was 264 days (out of 350 treatment days) and 150 days (out of 336 treatment days), respectively. Soticlestat was generally well tolerated, with the majority of reported AEs being mild in severity, and no SAEs were observed.

#### 2.7.2 Carisbamate

Carisbamate is being evaluated in a phase I, open-label, multicentre study in adult and paediatric patients with LGS (NCT03731715), with an estimated study completion date of September 2020 [135]. Carisbamate previously failed to show efficacy in a phase III study in patients with partial-onset seizures; however, given its wide spectrum of anticonvulsant activity in preclinical models, it is now being investigated in LGS patients [22, 130, 136]. Carisbamate belongs to the carbamate drug family, characterised by a key structural motif [137], which also includes felbamate (described above) and cenobamate; the latter was FDA approved in late 2019 for the treatment of partial seizures in adult patients, and is currently being assessed by the EMA [138].

#### 2.7.3 Ganaxolone

Ganaxolone is a GABA-A receptor modulator, which activates synaptic and extrasynaptic GABA-A receptors at a site
distinct from benzodiazepines and barbiturates [139]. In an open-label trial, ganaxolone was assessed in eight children with LGS (aged 2–15 years) with severe, treatment-resistant generalised tonic-clonic and drop seizures [139]. The median percentage change at 26 weeks for major seizures was −32% (range −80 to +40%; p = 0.195), and the median percentage change for seizure-free days was +33% (range 0 to +170%; p = 0.031). No SAEs were reported. Marinus Pharmaceuticals is currently conducting studies evaluating ganaxolone in CDKL5 deficiency disorder, PCDH19-related epilepsy and refractory status epilepticus [140]; however, it is not clear if any further studies in LGS patients are planned.

3 Non-Pharmacologic Agents

3.1 Ketogenic Diet

The ketogenic diet (KD) is a diet that is high in fat and low in carbohydrate with adequate protein, with strictly controlled amounts and ratios of fat to a combination of protein and carbohydrates [141]. The KD mimics the fasting state, characterised by systemic ketosis, with elevations in the concentrations of the ketone bodies, beta-hydroxybutyrate, acetoadecate and acetone [142]. The KD and related diets have been shown to be useful in various pharmacoresistant childhood epilepsies, including LGS [141]. While the mechanism of action for the KD’s anticonvulsant effects are unknown, theories include modulation by the ketone bodies on glutamatergic and GABAergic neurotransmitter metabolism, glycolytic restriction/diversion, improved cellular bioenergetics, mitochondrial function and reduced oxidative stress, direct inhibitory effects of fatty acids, and enhancement of the tricarboxylic acid cycle [141–143].

Recent systematic literature reviews and meta-analyses have provided a useful synthesis of the evidence regarding the efficacy and safety of the KD in patients with refractory epilepsy, albeit not specific to LGS. A Cochrane Database Systematic Review conducted in 2018 identified 11 RCTs evaluating the KD for drug-resistant epilepsy: the proportion of patients achieving ≥50% reduction in seizure frequency ranged from 35% to 56% in the KD groups compared with 0% to 18% in the control groups, and the proportion achieving seizure freedom ranged from 0 to 15% in the KD groups versus 0% to 9% in the control groups [144]. Further to this, a meta-analysis of 16 observational studies in adult patients with intractable epilepsy reported that the combined responder rate (≥50% seizure reduction) was 53% and the rate of seizure freedom was 13% [145]. The adverse reactions of the KD were mild. In addition, Lyons et al. recently (2020) conducted a systematic review of the KD specifically in infants aged <2 years with drug-resistant epilepsy, that identified 33 studies (2 RCTs and 31 uncontrolled cohort studies), with a total of 534 infants with efficacy data [146]. Meta-analyses of the uncontrolled studies estimated that 59% of infants achieved ≥50% seizure reduction and 33% of infants achieved seizure freedom. The KD was well tolerated in this population, with the most common AEs being dyslipidaemia (12%), vomiting (6%), constipation (4%), gastroesophageal reflux (4%), and diarrhoea (4%). However, this latter study may have limited generalisability to LGS, since a firm diagnosis of LGS is only rarely made at this young age, when associated with specific aetiologies such as anoxic-ischaemic encephalopathy that may develop into LGS.

With respect to studies conducted solely in patients with LGS, there has been a number of open-label studies that have reported ≥50% responder rates of 40–51%, and seizure freedom in 1–15% of patients, and have generally found the KD to be well tolerated [147–150]. The KD may also provide benefits beyond seizure control, with a systematic review reporting that the KD was associated with subjective improvements in alertness, attention, and global cognition in patients with refractory epilepsy [151]. However, diet adherence can be an obstacle to successful KD implementation, particularly in adults [152, 153].

3.2 Vagus Nerve Stimulation

Vagus nerve stimulation (VNS), which consists of intermittent electrical stimulation of the left cervical vagus nerve by an implanted helical electrode that is connected to a pulse generator, is a well-established procedure in drug-resistant epilepsy, with some evidence of efficacy in LGS. An evidence-based guideline update regarding VNS for the treatment of epilepsy from the AAN concluded that VNS could possibly achieve >50% seizure reduction in 55% of patients with LGS, based on four studies [154]. However, it should be noted that corpus callosotomy (discussed in the following section) may have a better outcome than VNS for reduction of atonic seizures [155–157].

3.3 Corpus Callosotomy

Corpus callosotomy is a palliative surgical procedure for patients with refractory epilepsy, which involves cutting the corpus callosum, the bundle of nerve fibres that connects the two cerebral hemispheres, thereby preventing epileptic discharges and seizure patterns from propagation to both hemispheres. It is especially targeted for those patients with drop attacks [2]. Corpus callosotomy is performed either as an anterior two-thirds disconnection of corpus callosum or as a complete disconnection [158], generally performed either through an open approach via a standard craniotomy with the aid of an operating microscope, or alternatively via a mini-craniotomy with endoscope assistance, with the latter having the benefit of a smaller incision, minimised...
brain retraction, and lower postoperative pain [158]. More recently, a minimally invasive method—magnetic resonance imaging (MRI)-guided stereotactic laser interstitial thermal therapy (LITT)—has shown promising results in case reports [159–167]. In the largest study to date, investigating MRI-guided stereotactic laser anterior corpus callosotomy (SLACC) in 10 patients with LGS (median age 33 years, range 11–52 years), eight (80%) patients had > 80% reduction in drop attacks, of whom five (50%) became free of drop attacks, and six (60%) achieved > 80% seizure reduction, with two (20%) becoming seizure free [167]. In addition, the complications of SLACC were minimal (one asymptomatic tract haemorrhage and no cases of disconnection syndrome), and patients had short hospital stays and minimal postoperative discomfort. However, it should be noted that the decision to perform corpus callosotomy should only be taken by medical teams with expertise in rare epilepsies and performed by neurosurgeons with the appropriate experience in epilepsy surgery.

### 3.4 Resective Epilepsy Surgery

Resective surgery in LGS patients is generally only recommended in a very few select patients with structural aetiology who have lesions predominantly in one hemisphere or TSC [1, 2, 168]. A recent study from Korea reported on the long-term outcomes of resective epilepsy surgery in 90 patients with LGS with focal epileptic pathology (mean age ± standard deviation: 9.3 ± 4.4 years, range 3–23.5 years) [169]. With an average post-operative follow-up of 6.1 ± 2.2 years (range 2.1–11.4 years), 45 patients (50.0%) had no seizures, and 15 (16.7%) reported infrequent seizures. Of note, seizure-free patients achieved better adaptive behaviour and social competence than did patients with persistent seizures at the second follow-up (2–3 years after surgery) and third follow-up (4–6 years after surgery), and a shorter latent period from seizure onset to surgery led to a better level of adaptive behavioural functioning. The authors concluded that early and intensive investigations to determine eligible patients with focal aetiology are important for achieving improvements in behavioural functioning. However, overall, the evidence regarding resective epilepsy surgery in LGS patients is limited, and no controlled trials have been reported.

### 4 Discussion

Despite valproate never being specifically licensed for use in LGS, it is generally recommended as a good first-line option due to its broad spectrum and a low risk of seizure aggravation [1, 2, 23] (Fig. 2). However, in the vast majority of cases, valproate therapy is insufficient for adequate seizure control. Evidence of efficacy from placebo-controlled clinical trials, leading to approvals specifically for LGS in the US, has been available for lamotrigine, topiramate and felbamate since the 1990s, and more recently from rufinamide (2008) and clobazam (2011). Cannabidiol is the most recent, and welcome, addition to the treatment landscape. Fenfluramine, which was recently approved for the treatment of seizures associated with DS, is also in late-stage development for LGS, and perampanel, which is approved for seizures of focal and generalised onset already used off-label in LGS patients, is also being evaluated in LGS patients in a phase III trial. Evidence from observational studies suggests that brivaracetam may also have benefits for patients with drug-resistant epilepsy, including LGS patients. Non-pharmacologic interventions are also expanding, with the potential for less invasive techniques for corpus callosotomy that have promise for reducing complications, while recent systematic reviews have provided further consolidated evidence of the benefits of the KD.

There has been a lack of direct head-to-head studies to determine the comparative efficacy of the various treatments [170]. Indeed, this may not be easy or appropriate for LGS; because of the heterogeneous nature of the syndrome, both between patients and over time in the same patient, there cannot be a ‘one size fits all’ approach to treatment [2, 21–24]. A personalised approach, tailored to the individual symptoms and responses of the patient during all stages of care, with regular assessment of treatment options, is particularly important for a syndrome such as LGS. In this respect, having more choices that will fit patient’s needs is clearly beneficial, although it also provides challenges to clinicians in selecting the best treatment or combination of treatments for their patients, whereby they must balance trade-offs between efficacy and side effects, taking into account the evidence from clinical trials (Table 2) and real-world studies, personal experience, as well as country availability. In this review, we have provided a comprehensive overview of the recent literature surrounding the efficacy and safety of treatments for LGS, with the aim of providing a useful resource for scientists and clinicians.

There are some limitations regarding the trials for LGS across the board that warrant some discussion. The trials were designed to evaluate the control of countable motor seizures, as opposed to specifically evaluating tonic or myoclonic seizures, which would need continuous video EEG monitoring. Therefore, the results regarding the total number of seizures (Table 2, Fig. 4) should be interpreted with the caveat that they do not encompass all the seizure types associated with LGS, and are therefore likely to represent an estimate of the actual total seizures.

It should also be noted that the therapeutic goals are shifting from symptom control (i.e. seizure freedom) towards QoL outcomes [2, 6]. Measuring QoL can be challenging in complex childhood syndromes, especially in instances where...
children are young and/or cognitively impaired [171, 172]; however, there has been more emphasis on outcomes that have an impact on QoL in more recent trials. Cannabidiol and fenfluramine are both associated with improvements in the patient’s overall condition as per the Global Impression of Change, and cannabidiol was also associated with an increase in the number of drop seizure-free days. Further analysis of these newer treatments, as well as the ‘older’ treatments, are needed to assess their long-term impact on the QoL of patients, and also their caregivers.

The currently available treatments for LGS are primarily focussed on the control of seizures, with little evidence of an impact on the intellectual disability and behavioural aspects. This may require advances in two areas in particular; firstly, improvements in early diagnosis so that patients can be appropriately treated before there is irreparable damage to brain structures, which occurs in infants and children in the early stages of the syndrome [6]; and secondly, a more comprehensive understanding of the neurobiology of the syndrome is required with the aim of developing targeted therapies to the electro-clinical complex of LGS itself as well as the underlying aetiologies that can improve cognition and behaviour in addition to seizures. Of note, fenfluramine has shown improvements in emotion and behaviour regulation in DS, which may provide some hope in LGS patients, although there is no evidence to date of this benefit in LGS [101, 173–175]. A recent observational prospective study found that rufinamide was not associated with any statistically significant changes in cognitive, adaptive function and emotional aspects in LGS patients (N = 16, mean age 22 years) with a follow-up of 1 year, demonstrating neither a positive nor a negative effect on these outcomes [176].

5 Conclusion

Despite recent breakthroughs, with newly approved and up-coming treatments, patients continue to experience a significant burden. Because LGS is not a single entity, tailoring of treatment is needed as opposed to a ‘one size fits all’ approach. Further research is needed into the underlying aetiologies and pathophysiology of LGS, together with advancements in treatments that encompass the spectrum of seizures associated with this complex syndrome.

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