Clobazam and its use in epilepsy

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Abstract

Clobazam (CLB) is an older anti-epileptic drug, with a slightly different chemical structure from that of the classic benzodiazepines currently used in the treatment of epilepsy, which confers less sedative properties in terms of negative adverse effects. It is also thought to be better tolerated than other anti-epileptic drugs, whilst maintaining a very similar level of efficacy. It has been tested extensively in over 50 studies on more than 3000 patients with epilepsy and is now approved as an adjunctive treatment of epilepsy in >100 countries. The aim of this review is to evaluate several existing studies on the effectiveness of CLB as an adjunctive therapy in the treatment of epilepsy and whether this therapy is more useful in particular types of epilepsy or seizure prevention. This is not a systematic review but a general overview of some of the most recent studies on the effectiveness of CLB as an adjunctive therapy. Additionally, the benefits of having an oral suspension of CLB will be evaluated with regards to patient groups benefiting from this formulation. The last issue addressed is that of the importance of prescribing CLB by brand, along with the benefits and risks of not doing so.

The use of clobazam as an adjunct therapy in the treatment of epilepsy

Clobazam (CLB) is a 1,5-benzodiazepine that has been introduced in 1975 as an anxiolytic drug and shortly after, it was discovered that it has strong anti-epileptic properties as well. It is distinguished from other classic 1,4-benzodiazepines in that its nitrogen atoms are placed in 14th and 5th positions in the B ring. Its mechanism of action is similar to other benzodiazepines however CLB is believed to be a partial agonist rather than non-selective full receptor agonists, which is what 1,4-benzodiazepines are. Moreover, CLB has lower affinity for the GABA_A subunits and greater selectivity for a2 subunits over a1 subunits, than the 1,4-benzodiazepines and it is thought that these properties confer to CLB less sedative effects than other benzodiazepines. The standard treatment for epilepsy involves using a single anti-epileptic drug at the minimally effective dose, up to the maximum tolerated dose. However, the numerous seizure types that a patient may experience render treatments with one agent ineffective so combination therapy is often required. Breakthrough seizures are often experienced by patients; hence continuous adjustments need to be made to their medications regimes over the course of their lifetime, both in terms of dosage and number of agents used.

Due to its less sedative effects and its very similar effectiveness in comparison with other agents, CLB is very frequently selected as an add-on agent when polytherapy is needed, particularly in the case of intractable epilepsy. Several studies have shown that CLB is an effective adjunctive anti-epileptic drug (AED) for a few specific types of epilepsy, most importantly Lennox-Gastaut syndrome (LGS). These include both retrospective studies and more importantly randomized, double-blind studies.

A randomized, double-blind, dose-ranging study evaluated safety and efficacy of CLB as adjunctive therapy for drop seizures in patients with LGS. LGS is an epileptic encephalopathy characterized by multiple types of seizures and developmental delay. The presence of a characteristic triad described typical LGS: i) tonic axial, atonic, and/or atypical absence seizures; ii) electroencephalography (EEG) abnormalities with bursts of diffuse slow spike-wave pattern of 1.5-2.5 Hz; and iii) impaired intellectual growth. Atonic or drop seizures are frequent in patients with LGS and are responsible for most injuries associated with falls. Seizures in LGS are refractory to most AEDs hence the need for combinational therapy.

The study conducted was a phase II, randomized, double-blind, dose-ranging multicenter study which comprised a 4-week baseline period, a 3-week titration period and a 4-week maintenance period. A daily seizure diary record was used to record the number of seizures, specifically any drop seizures. They were recorded as single drop seizures (defined as a drop seizure occurring 15 min or more before and after the next seizure or cluster) or as clusters (defined as two or more drop seizures, with less than 15 min between any two consecutive seizures); non-drop seizures have also been recorded but on a smaller number of patients. Six weight groups have been defined and patients with two or more drop seizures per week during the baseline period were placed in one of six weight groups and randomly assigned to either low-dose CLB (target dose of 0.25 mg/kg/day; maximum 10 mg/day if weight was lower than 37.6 kg) or high-dose CLB (target dose of 1.0 mg/kg/day; maximum 40 mg/day if weight was higher than 37.6 kg). A significant reduction in drop seizure rates was observed both in the low-dose group (mean reduction=12%; P=0.0162) and in the high-dose group (mean reduction=85%; P<0.0001). Importantly, high-dose CLB was significantly more effective in reducing drop seizure rates compared with low-dose CLB (P<0.0001). Eighty-nine percent of responders in the high-dose group and 56% in the low-dose group experienced a ≥50% reduction in drop seizures (P=0.0025); 83 and 38% experienced a ≥50% reduction (P=0.0001); 67 and 25% experienced a ≥75% reduction (P=0.0006); and 6 and 22% experienced a 100% reduction (P=0.0629). The study showed that CLB reduces the non-drop seizure rates as well, particularly in the high-dose group. The percent change in the low-dose group was not significant, however in the high-dose group the percent change from baseline (59±55%, n=22) was significant. The dose-dependent manner of reducing drop seizure rates was also recorded for non-drop seizure rates, the reduction being significantly greater in the high-dose group compared with the low-dose group (P=0.0222). Parent/caregiver and investigator global evaluations have both demonstrated that the high-dose CLB group showed significantly greater improvements in overall symptoms compared to low-dose CLB group. At one investigation site, the quality of life of four children receiving CLB was greatly improved, as they discontinued wearing helmets and therefore they were able to move freely with...
out the constant adult supervision previously needed to prevent injury from drops. A more advanced phase III, double-blind, placebo-controlled study on the safety and efficacy of CLB in patients with LGS aged 2-54 was conducted at 51 sites in the United States, India, Europe and Australia between August 2007 and December 2009 and further assessed CLB’s role as adjunctive therapy.7 This study evaluated the efficacy of 3 CLB dosages in decreasing weekly frequencies of drop and total seizures and also assessed its safety when administered ≥18 weeks at these 3 dosages. Patients aged 2-60 years were eligible to participate if they had onset of LGS before 11 years age and currently weighed ≥12.5 kg. The study included 4-week baseline, 3-week titration and 12-week maintenance periods, followed by either continuation in an open-label study or a 2- or 3-week taper period. Patients were randomly assigned to one of 4 groups, depending on weight (12.5 kg to ≤30 kg, >30 kg): i) placebo; ii) low-dosage CLB: target of 0.25 mg/kg/day (maximum, 10 mg/day); iii) medium-dosage CLB: target of 0.5 mg/kg/day (maximum, 20 mg/day); or iv) high-dosage CLB: target of 1.0 mg/kg/day (maximum, 40 mg/day). The mean patient age was 12.4 years. Importantly, approximately 50% of all patients were receiving concomitant valproic acid, valproate semisodium, or valproate sodium.

The mean percentage decrease in average weekly rate of drop seizures from baseline to maintenance period was 12.1% for placebo vs 41.2% (P=0.0120), 49.4% (P=0.0015) and 68.3% (P<0.0001) for the 0.25, 0.50 and 1 mg/kg/day dosage groups respectively. Mean difference from the placebo group increasing with increasing CLB dosage (mean differences of 29.1, 37.3 and 56.1% for the low, medium and high respectively). A linear trend has been noted in that increasing dosage of CLB lead to increased efficacy in reducing the drop seizure rates (P<0.0001). The mean percentage decrease in average weekly rate of total (drop and non-drop) seizures was 9.3% for placebo vs 34.8% (P=0.0014), 45.3% (P=0.0044), and 65.3% (P<0.0001) for the CLB 0.25-, 0.5-, and 1.0-mg/kg/day groups.

A 40.0% decrease in the average weekly rate of non-drop seizures was observed for the high-dosage group, however this was not statistically significant by analysis of covariance model. An increase of 76.3, 53.3 and 3.3% has been noted in the average weekly rate of non-drop seizures for the placebo, low-dosage and medium-dosage CLB group.

Increasing CLB dosage lead to increase response rates in patients with LGS. The percentage of patients with ≥50% decrease from baseline to maintenance period in average weekly rate of drop seizures was 31.6% for placebo, 43.4, 58.6 and 77.6% for the low-, medium-, and high-dosage CLB groups, respectively. In comparison with the placebo group, the likelihood of achieving ≥50% response was greater for the medium-dosage and high-dosage CLB groups. Seizure-free patients have also been reported: 2 patients in the placebo group (3.5%) were seizure-free, compared with 4 (7.5%), 7 (12.1%) and 12 (24.5%) patients for the low-, medium-, and high-dosage CLB groups.

Global evaluations of patients’ overall changes in symptoms from physicians and caregivers during the observed period have also shown the CLB as adjunctive therapy led to improvements: percentages of patients who were at least minimally improved range from 71.2 to 80.7% (physicians’ assessments) and 79.2 to 81.6% (caregivers’ assessments) for CLB vs 47.3 and 45.5% respectively for placebo.

Although retrospective studies are statistically less significant than randomized controlled trials, they still provide valuable information about the effectiveness and safety of CLB in patients with epilepsy. A retrospective study was conducted between January 2013 and January 2015 in patients suffering from status epilepticus (SE). SE is defined as seizures lasting >5 min or multiple seizures without recovery of consciousness in between. Refractory status epilepticus (RSE) is defined as SE that persists despite adequate treatment with benzodiazepines and at least one AED, or SE requiring general anesthesia. About 12-43% of the cases with SE become refractory, and 50% of those requiring anesthesia will become super-refractory. Patients from all age groups in whom CLB was administered for the management of SE were included in the study. Over a period of 24 months, 17 patients received CLB for the treatment of RSE. In all these patients, CLB was used as add-on therapy after failure of two or more AEDs in adequate dosing and it was the last AED added in 94% of the patients. Thirteen patients reported a successful response to CLB (76.5%).

Another retrospective study conducted at the Hospital de Clínicas da Unicamp on 97 patients ranged 15 to 70 years who were evaluated for surgery and had been followed-up for ≥1 year has evaluated the effectiveness of CLB as add-on therapy. Of these 97 patients, 74.2% had temporal lobe epilepsy, 8.2% had extra-temporal epilepsy and in 17.6% patients the epileptic syndrome could not be identified. CLB was introduced after previous failure of at least two mono-therapies, with carbamazepine, phenytoin or valproate used up to maximum tolerated dose. The dosage of CLB ranged from 10 to 60 mg twice a day and the period of usage ranged from 1 month to 7 years and 9 months. The study brought proof to the effectiveness of CLB as adjunctive therapy: 7.2% patients were seizure-free, 49.4% had ≥50% improvement in seizure control and 40.2% patients had <50% improvement in seizure control. In 3.1% no data were available. A review study conducted in 2011 evaluated several studies on the effectiveness of CLB, both prospective and retrospective studies. In pediatric patients with refractory epilepsy, six open-label prospective studies have shown that at least 54%-85% of patients experienced at least a 50% drop in seizure rates (Table 1).7,11-20

Additionally, two retrospective studies have also reported significant decrease in seizure rates for pediatric patients using CLB as add-on therapy (Table 1).11,17

Clinical studies of LGS were identified in a 2009 Cochrane review and by electronic database search and indirect comparison of the relative efficacies of CLB, felbamate, lamotrigine, topiramate and rufinamide as adjunctive treatments for LGS was performed. These indirect comparisons were performed by transforming the primary efficacy endpoint from each trial into Cohen’s d effect size. The results have also shown that high-dosage CLB (1.0 mg/kg/day) was the most effective as placebo, whereas medium-dosage CLB (0.5 mg/kg/day) and rufinamide had moderate effects. Felbamate, lamotrigine and topiramate had low effect sizes. Numbers of total seizures and tonic-atonic seizures (drop attacks) were indirectly compared and both comparisons proved that medium- and high-dosage CLB are superior to the other adjunctive LGS therapies.21

A study that investigated potential drug interactions between CLB and other AEDs, including phenytoin, phenobarbital, carbamazepine, valproate, lamotrigine, felbamate, and oxcarbazepine, found no clinically meaningful drug pharmacokinetic interactions, which makes this drug suitable for the management of LGS as an adjunctive therapy due to its pharmacokinetic properties and less aggressive side effects.22

Why should clobazam be prescribed by brand?

The prescription of anti-epileptic medication can become an issue in the treatment of epilepsy. Practitioners are often encouraged to prescribe the cheapest drugs available and this is often inappropriate for the management of epilepsy.23 A research study conducted in 2003 suggests that even small differences between two versions of the same drug can become very problematic for the patient who is switching them.24 Crawford et al. suggest that these problems include additional side-effects or seizures frequency.24 However, the necessity of prescribing CLB by brand is a hypothesis based on the available evidence on other AEDs.

The claimed advantage of prescribing gener-
Table 1. Clobazam as add on therapy in refractory pediatric epilepsy.

<table>
<thead>
<tr>
<th>Study</th>
<th>Trial design</th>
<th>Participants and included diagnoses</th>
<th>Dosage</th>
<th>Results</th>
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</thead>
<tbody>
<tr>
<td>Conry et al.6</td>
<td>Phase II, multi-center, randomized, double-blind, dose-ranging</td>
<td>68 patients; 2-26 years; LGS</td>
<td>0.25 mg/kg/day, or 1.0 mg/kg/day</td>
<td>0.25 mg/kg/day: 38% of patients had a ≥50% decrease in drop seizure rates; 1.0 mg/kg/day: 83% of patients had a ≥50% decrease in drop seizure rates</td>
</tr>
<tr>
<td>Conry et al.7</td>
<td>Phase III, multi-center, randomized, double-blind, dose-ranging, placebo-controlled</td>
<td>238 patients; 2-54 years; LGS</td>
<td>0.25 mg/kg/day, 0.5 mg/kg/day, or 1.0 mg/kg/day</td>
<td>0.5 mg/kg/day: 58% of patients had a ≥50% decrease in drop seizure rates; 1.0 mg/kg/day: 77% of patients had a ≥50% decrease in drop seizure rates</td>
</tr>
<tr>
<td>Da Silveira et al.11</td>
<td>Retrospective</td>
<td>100 patients; 1-18 years; refractory local epilepsy</td>
<td>5-60 mg/day</td>
<td>33% of patients had a ≥50% decrease in seizure rates</td>
</tr>
<tr>
<td>Farrell12</td>
<td>Open-label, prospective</td>
<td>50 patients, 33 with LGS2; 16 years; refractory epilepsy</td>
<td>5-40 mg/day</td>
<td>50% of patients had a ≥50% decrease in seizure rates</td>
</tr>
<tr>
<td>Jan and Shaabat13</td>
<td>Open-label, prospective</td>
<td>31 patients, 14 with LGS; 2 months to 15 years intractable childhood epilepsy</td>
<td>5-40 mg/day</td>
<td>80% of patients had a ≥50% decrease in seizure rates</td>
</tr>
<tr>
<td>Kalra et al.14</td>
<td>Open-label, prospective</td>
<td>88 patients; 7 months to 12 years refractory epilepsy</td>
<td>0.3-2.0 mg/kg/day</td>
<td>89% of patients had a ≥50% decrease in seizure rates</td>
</tr>
<tr>
<td>Keene et al.15</td>
<td>Double-blind, placebo-controlled, crossover</td>
<td>21 patients; 2-19 years; refractory epilepsy</td>
<td>0.25-1.0 mg/kg/day</td>
<td>54% of patients had a ≥50% decrease in seizure rates</td>
</tr>
<tr>
<td>Munn and Farrell16</td>
<td>Open-label, prospective</td>
<td>115 patients, 25 with LGS; 15 months to 17 years refractory epilepsy</td>
<td>0.36-3.8 mg/kg/day</td>
<td>62% of all patients had a ≥50% decrease in seizure rates; 64% of LGS patients had a ≥50% decrease in seizure rates</td>
</tr>
<tr>
<td>Silva et al.17</td>
<td>Retrospective</td>
<td>97 patients, 26 with LGS; 2 with LGS and West syndrome; 1-17 years; epileptic encephalopathy</td>
<td>5-60 mg/day</td>
<td>37% of patients had a ≥50% decrease in seizure rates</td>
</tr>
<tr>
<td>Sheth et al.18</td>
<td>Open-label, prospective</td>
<td>63 patients, 14 with LGS; 3-20 years; intractable epilepsy</td>
<td>Average 0.8 mg/kg/day</td>
<td>69% of patients had ≥50% decrease in seizure rates</td>
</tr>
<tr>
<td>Sugai19</td>
<td>Open-label, prospective</td>
<td>Short-term: 55 patients; 8 with LGS; long-term: 31 patients, 4 with LGS, refractory epilepsy</td>
<td>0.28-1.25 mg/kg/day</td>
<td>Short-term: 71% of all patients and 62% of LGS patients had a ≥50% decrease in seizure rates; Long-term: 81% of all patients and 50% of LGS patients had a ≥50% decrease in seizure rates</td>
</tr>
<tr>
<td>Vadja et al.20</td>
<td>Open-label, prospective or double-blind, placebo-controlled, crossover</td>
<td>14 patients,*7 with LGS; 6-38 years; refractory epilepsy</td>
<td>15-60 mg/day</td>
<td>40% of patients had a ≥50% decrease in seizure rates</td>
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LGS, Lennox-Gastaut syndrome; *results were not reported for 4 patients.
ically is that large amounts of money can be saved. However, the hidden consequences of generic prescribing is that costs may actually increase due to increased doctor visit (as a result of patient anxiety), increased sick leave, worse health for the patient and even in some cases potential loss of employment (Table 1).23

A survey of 1851 patients with epilepsy conducted by Epilepsy Action revealed that in the previous year, 33% of responders were given a different version of brand of their regular AED. Of these, almost 25% experienced an increase in seizure frequency as a result and 33% experienced more or distinct side effects from the ones previously experienced.23

The survey also showed that a significant number of people (24%) reported that they received a variety of versions of their medication in one single prescription.25

The recommended guideline from the National Institute for Health and Care Excellence (NICE) with regards to this issue states as follows: Changing brand of AED is not recommended due to variances in bioavailability, difference in pharmacokinetic profiles, which leads to increased potential for reduced effect or excessive side-effects (NICE, 2004).25

Patient’s anxiety is a factor of major importance in epilepsy, as it can easily trigger seizures. Patients with changed medication may be anxious to take them, which in turn may lead to the loss of seizure control. Suffering a seizure after a long seizure-free period could have dramatic consequences on the well being of the patient, both in terms of the impact it has on his life but also in terms of the damage a seizure can cause itself. The impact that a slightly different version of an AED can have on a patient’s life is to be taken into consideration and must not be underestimated.25,26

What patient groups may benefit most from a prescription of clobazam?

The NICE recommends CLB as an adjunctive treatment for epilepsy where first-line antiepileptic drug treatment has failed.27

NICE recommends CLB as an adjunctive treatment option for seizures: i) focal seizures; ii) generalized tonic-clonic seizures; and epilepsies: i) benign epilepsy with centrotemporal spikes; ii) Panayiotopoulos syndrome; iii) late-onset childhood occipital epilepsy (Gastaut type); iv) Dravet syndrome; v) epilepsy with generalized tonic-clonic seizures only.

NICE recommends CLB as an option on referral to tertiary care for seizures: i) generalized myoclonic seizures; ii) generalized absence seizures; and epilepsies: i) childhood absence epilepsy or other absence epilepsy syndromes; ii) juvenile absence epilepsy or other absence epilepsy syndromes; iii) juvenile myoclonic epilepsy; iv) idiopathic generalized epilepsy.

What patient groups may benefit most from a licensed liquid formulation of clobazam?

Accurate Dosing

Children

The ability to prescribe and administer safe and accurate doses of anti-epileptic drugs is fundamentally important in the treatment of epilepsy. The potential, however, for dosing accuracy in young children using CLB tablets is limited, as the smallest dose that can be accurately administered is 5 mg; it is for this reason that CLB tablets are not licensed for children under 6 years of age. Children between the age of 1 month and 6 years require small, weight-based doses, beginning at 125 mcg/kg twice a day. CLB oral suspension allows for the simple measurement of accurate doses, which will support compliance and offer the opportunity for optimal seizure control.

Other patient groups

CLB suspension may be beneficial for patients who require small starting doses of CLB (doses <5 mg), for example elderly patients or in those known to be poor CYP 2C19 metabolizers.

Ease of Administration

Swallowing difficulties (or the inability to tolerate solid oral dosage forms) are over-represented in both adults and children with severe seizure disorders. Known risk factors for refractory epilepsy include diffuse brain injury, genetic and metabolic disorders and underlying brain abnormalities, all of which are likely to be associated with additional neurological and/or behavioral deficit that may preclude, or at least complicate, dosing with solid oral dosage forms. Oral CLB suspension offers these patients a more acceptable formulation, which removes the need to crush tablets, saving patient/caregiver time, while potentially supporting compliance. The scientific evidence backing the administration of CLB orally is lacking, however this is a pragmatic recommendation that can benefit a lot of patients who have swallowing difficulties.

Anxious patients

While epilepsy represents the majority of CLB prescribing, it is also used in the short-term treatment of anxiety in adults, where it is reserved for the management of anxiety that is severe, disabling or subjecting the individual to unacceptable distress. Although drug administration in this patient group might not pose the same complexity as is associated with the epileptic group this is, none-the-less, a severely unwell cohort for whom compliance may be compromised and for whom availability of CLB oral suspension is advantageous. This group is likely to include some elderly patients, for whom the possibility of using smaller doses would be desirable.

References

12. Farrell K. Benzodiazepines in the treat-