Eslicarbazepine acetate in the treatment of adults with partial-onset epilepsy: an evidence-based review of efficacy, safety and place in therapy

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Introduction: Up to 30% of the patients diagnosed with epilepsy will continue suffering from seizures despite treatment with antiepileptic drugs, either in monotherapy or polytherapy. Hence, there remains the need to develop new effective and well-tolerated therapies.

Aim: The objective of this article was to review the evidence for the efficacy and safety of eslicarbazepine acetate (ESL) as adjunctive treatment in adult patients with focal onset seizures.

Evidence review: ESL is the newest, third-generation, single enantiomer member of the dibenzazepine family. Following oral administration, ESL is rapidly and extensively metabolized by hepatic first-pass hydrolysis to the active metabolite eslicarbazepine, which has linear, dose-proportional pharmacokinetics and low potential for drug-drug interactions. Eslicarbazepine works as a competitive blocker of the voltage gated sodium channels; unlike carbamazepine (CBZ) and oxcarbazepine (OXC), it has a lower affinity for the resting state of the channels, and reduces their availability by selectively enhancing slow inactivation. Efficacy and safety of ESL have been assessed in four randomized, Phase III clinical trials: the median relative reduction in standardized seizure frequency was 33.4% and 37.8% in the ESL 800 and 1,200 mg daily dose groups, and the responder rates were 33.8% and 43.1%, respectively. The incidence of treatment-emergent adverse events (TEAEs) increased with raising the dosage (ESL 400 mg: 63.8%, ESL 800 mg: 67.0%, ESL 1,200 mg: 73.1%). The TEAEs were generally mild to moderate in intensity, and the most common were dizziness, somnolence, headache and nausea.

Open-label studies confirmed the findings from the pivotal trials and demonstrated sustained therapeutic effect of ESL over time and improvement of tolerability profile in patients switching from OXC/CBZ. No unexpected safety signals emerged over >5 years of follow-up.

Conclusion: Once-daily adjunctive ESL at the doses of 800 and 1,200 mg was effective to reduce the seizure frequency and was fairly well tolerated in adults with focal onset epilepsy. Starting treatment at 400 mg/day, followed by 400 mg increments every 7–14 days, could provide the optimal balance of efficacy and tolerability.

Keywords: eslicarbazepine acetate, epilepsy, focal seizures, review

Clinical impact summary for eslicarbazepine acetate as adjunctive treatment in patients with focal onset seizures

<table>
<thead>
<tr>
<th>Evidence</th>
<th>Implications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Randomized, placebo-controlled, double-blind, clinical trials and observational, open-label studies</td>
<td>Once-daily eslicarbazepine acetate at the dosage of 800 and 1,200 mg consistently demonstrated to be effective in the adjunctive treatment of adult patients presenting with focal onset seizures.</td>
</tr>
</tbody>
</table>
Outcome measure | Evidence | Implications
---|---|---
A dose-dependent response occurred with an increase in effect by increasing the dose. In the pooled analysis of the Phase III pivotal trials, the median relative reduction in standardized seizure frequency was 33.4% and 37.8% in the eslicarbazepine acetate 800 mg and 1,200 mg daily dose groups and the responder rates were 33.8% and 43.1%, respectively. The drug efficacy was demonstrated regardless of whichever concomitant baseline, antiepileptic drugs were used. Open-label studies showed similar results and demonstrated sustained therapeutic effect of eslicarbazepine acetate.

Patient-oriented evidence
Tolerability and safety | Randomized, placebo-controlled, double-blind, clinical trials and observational, open-label studies
---|---
Once-daily eslicarbazepine acetate was well tolerated overall as adjunctive treatment in adults with focal onset epilepsy. In the pooled analysis of the Phase III trials, the incidence of treatment-emergent adverse events increased with increase of the dose (eslicarbazepine acetate 400 mg: 63.8%, eslicarbazepine acetate 800 mg: 67.0%, eslicarbazepine acetate 1,200 mg: 73.1%); most of the treatment-emergent adverse events were mild to moderate in intensity. The most common were dizziness, somnolence, headache and nausea. The incidence of adverse events was higher in patients co-treated with carbamazepine than with other antiepileptic drugs. Changes in mean laboratory parameters were not associated with clinically relevant findings, and there were no variations in vital signs, body weight or electrocardiography of clinical concern. Hyponatremia leading to treatment discontinuation occurred in less than 1% of the patients taking eslicarbazepine acetate. Similar results were observed in the open-label studies. Tolerability profile improved in patients switching from oxcarbazepine/carbamazepine to eslicarbazepine acetate due to side effects. No unexpected safety signals emerged over >5 years of follow-up.

Net benefit | Randomized, placebo-controlled, double-blind, clinical trials and observational, open-label studies
---|---
Starting treatment at the 400 mg/day dose, followed by 400 mg increments every 7–14 days until the optimal dose is reached, can maximize the balance of efficacy and tolerability.

Economic evidence | Not available
---|---

Introduction
Epilepsy is one of the most common neurological disorders affecting approximately 70 million people worldwide. It has an annual incidence of about 80 cases per 100,000 people and a point prevalence between 4 and 10 cases per 1,000 people.¹ The treatment of epilepsy is mainly symptomatic and nearly 70% of all patients have a good prognosis and achieve a long-term remission. However, up to 30% of adults diagnosed with
epilepsy will continue suffering from seizures despite treatment with antiepileptic drugs (AEDs), either in monotherapy or polytherapy.² Notably, the inadequate seizure control, which may be related to either pharmacoresistance or intolerable adverse effects, is associated with physical risks and impairments in social opportunities and has a negative influence on overall quality of life.³ Hence, there remains the need to develop new effective and well-tolerated treatment options.

Eslicarbazepine acetate (ESL) is a novel AED, which has been approved by the European Medicines Agency and the US Food & Drug Administration as monotherapy and adjunctive therapy for focal onset seizures (FOS), with or without secondary generalization, in adults, and was recently authorized also in children and adolescents as adjunctive treatment.

This paper provides a review of the available evidence on the efficacy and safety of ESL as adjunctive treatment in adult patients with FOS, and highlights major issues about its actual place in therapy.

Brief pharmacology: chemical structure, pharmacodynamics and pharmacokinetics

ESL is the newest, third-generation, single enantiomer member of the dibenzazepine family, which also includes carbamazepine (CBZ) (first generation) and oxcarbazepine (OXC) (second generation), and works as a competitive blocker of the voltage gated sodium channels (VGSC).⁴ The structural variation at the 10,11 position of the dibenzazepine nucleus conveys ESL unique pharmacodynamics and pharmacokinetics.⁵ Compared to CBZ, ESL has similar affinity for the inactivated state and a threefold lower affinity for the resting state of the VGSC. Unlikely traditional sodium channel blockers, which interfere with the fast inactivation pathway, ESL reduces the VGSC availability by selectively enhancing slow inactivation, similarly to lacosamide.⁶,⁷ These properties result in stabilization of hyper-excitabile neuronal membranes, inhibition of sustained repetitive production of action potentials characteristic of neurons involved in epilepsy and reduction of long-term channel availability, with a low propensity to disturb physiological function.⁸

ESL is an oral pro-drug that is well absorbed following ingestion, with a bioavailability of approximately 94%, and is rapidly and extensively bio-transformed to its major active metabolite eslicarbazepine (S-licarbazepine) by hepatic first-pass hydrolysis. Accordingly, plasma levels of ESL usually remain below the limit of quantification.⁹,¹⁰ The binding of eslicarbazepine to plasma proteins is relatively low (<40%) and independent of concentration. Plasma peak dose is attained at 2–3 hours post-dose, and the steady state levels are achieved after 4–5 days of once-daily dosing, consistent with an effective half-life of 20–24 hours.⁹,¹⁰ Minor metabolites in plasma include the right isomer of eslicarbazepine (R-licarbazepine) and OXC, which were shown to be active, and the glucuronic acid conjugates of ESL, S-licarbazepine, R-licarbazepine and OXC.⁹ The pharmacokinetics of ESL avoids the toxic product of CBZ degradation (i.e., carbamazepine-10,11-epoxide),⁴ and the early peak in plasma and cerebrospinal fluid (CSF) concentrations following immediate-release OXC administration.¹¹ Further, ESL does not affect its own metabolism or clearance; eslicarbazepine is a weak inducer of CYP3A4 and UDP-glucuronyl transferases, and has inhibiting properties with respect to CYP2C19. ESL metabolites are eliminated from the systemic circulation primarily by renal excretion, in the unchanged and glucuronide conjugate forms; S-licarbazepine and its glucuronide correspond to more than 90% of total metabolites excreted in urine.⁹ Overall, the pharmacokinetics of ESL are linear, dose-proportional in the 400–1,200 mg range and unaffected by age, sex and food intake.⁵,¹⁰,¹²

Drug formulation and dosing

ESL is an oral AED available in a liquid suspension or tablet form, which may be taken with or without food. The recommended starting dose of ESL is 400 mg once daily, which should be increased to 800 mg once daily after 1 or 2 weeks; according to individual response, the dose as adjunctive treatment may be increased to 1,200 mg once daily.⁹ The plasma levels show a linear correlation with the posology, but a therapeutic range has not been defined.

Efficacy, tolerability and safety of adjunctive ESL in randomized, placebo controlled, double-blind trials

The adjunctive therapy program to evaluate the efficacy and tolerability of ESL in adult patients with FOS included one Phase II study (study BIA-2093-201)¹³ and four Phase III, multicenter, randomized, double-blind and placebo controlled clinical trials (studies BIA-2093-301 [NCT00957684], BIA-2093-302 [NCT00957047], BIA-2093-303 [NCT00957372] and BIA-2093-304 [NCT00988429]).¹⁴–¹⁷ All the studies recruited patients presenting with at least four FOS per month, with or without secondary generalization, despite treatment with one to three AEDs at a stable dose; notably, OXC was not allowed as a concomitant AED as it has the same pharmacologically active main metabolite with ESL.

In the Phase II study,¹³ 143 patients were randomized to treatment with ESL once daily, ESL twice daily, or placebo.
The ESL daily dose was gradually titrated up to 1,200 mg/day in three steps at 4-week intervals. A statistically significant difference from placebo in the response rate at the end of the 12-week treatment period was found only for the once-daily group, whereas the incidence of adverse events was similar across the arms. Accordingly, ESL has been administered once a day in all the subsequent trials.

All the Phase III trials followed a similar design with ESL administered once daily for 12 weeks (maintenance period); the major differences in trial designs included the number of doses tested and the titration and tapering-off regimens. Studies BIA-2093-301 and BIA-2093-302 had three ESL daily dose groups (400, 800 and 1,200 mg), whereas studies BIA-2093-303 and BIA-2093-304 had only two ESL doses (800 and 1,200 mg). In each of the individual studies, the primary efficacy variable was the 4-week standardized seizure frequency (SSF) over the maintenance period, and the predefined key efficacy outcomes included the median reduction and the 50% or greater reduction in SSF (response rate) over the maintenance period compared to baseline. Safety assessment included treatment-emergent adverse events (TEAEs), clinical laboratory parameters, vital signs and electrocardiography. The main characteristics and results of the Phase III trials are summarized in the Table 1.

The population in the pivotal four trials was predominantly Caucasian with a mean age of 38 years and a mean disease duration of 22 years. The baseline seizure frequency was approximately 15 seizures/4 weeks, and all patients but five were taking at least one concomitant AED at the end of the baseline period. The most common concomitant drugs were CBZ, lamotrigine and valproic acid. Overall, results of these studies clearly demonstrated that once-daily ESL at the dosage of 800 or 1,200 mg is effective and well tolerated as adjunctive treatment in patients with focal onset epilepsy; furthermore, starting treatment at the 400 mg/day dose, followed by 400 mg increments, provides optimal balance of efficacy and tolerability. In a pooled analysis of the integrated datasets, the median relative reduction in SSF was 17.6% in the placebo group compared to 23.4%, 33.4% and 37.8% in the ESL 400, 800 and 1,200 mg groups, respectively; the responder rates were 22.9%, 33.8% and 43.1% for the ESL 400, 800 and 1,200 mg daily doses, respectively, and 22.2% for the placebo group. The differences to placebo were statistically significant only for the ESL 800 and 1,200 mg doses; the drug efficacy was demonstrated regardless of whichever concomitant baseline AEDs were used. The overall incidence of TEAEs increased with the increase of ESL doses; the drug efficacy was demonstrated regardless of whichever concomitant baseline AEDs were used. The overall incidence of TEAEs increased with the increase of ESL doses; the drug efficacy was demonstrated regardless of whichever concomitant baseline AEDs were used. The overall incidence of TEAEs increased with the increase of ESL doses; the drug efficacy was demonstrated regardless of whichever concomitant baseline AEDs were used.

### Table 1 Main characteristics and results of the Phase III, randomized, double-blind, placebo-controlled trials of adjunctive eslicarbazepine acetate in patients with refractory focal onset seizures

<table>
<thead>
<tr>
<th>Study (reference)</th>
<th>AEDs allowed</th>
<th>Titration</th>
<th>Tapering-off</th>
<th>Treatment arms</th>
<th>Median relative reduction in SSF</th>
<th>Reduction in SSF ≥50%</th>
<th>Any TEAEs</th>
</tr>
</thead>
<tbody>
<tr>
<td>BIA-2093-30114 1–2</td>
<td>At 400 mg weekly steps</td>
<td>At 400 mg weekly steps until 0 mg (PBO)</td>
<td>N=402</td>
<td>16.0%</td>
<td>20.0%</td>
<td>31.4%</td>
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<td></td>
<td></td>
<td></td>
<td>PBO=102</td>
<td></td>
<td>26.0%</td>
<td>23.0%</td>
<td>44.0%</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>ESL 400 mg=100</td>
<td></td>
<td>36.0% (p&lt;0.05)</td>
<td>34.0% (p&lt;0.05)</td>
<td>50.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>ESL 800 mg=98</td>
<td></td>
<td>45.0% (p=0.001)</td>
<td>43.0% (p&lt;0.001)</td>
<td>60.8%</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>ESL 1,200 mg=102</td>
<td></td>
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</tr>
<tr>
<td>BIA-2093-30215 1–3</td>
<td>No titration in the</td>
<td>No tapering-off</td>
<td>N=395</td>
<td>0.8%</td>
<td>13.0%</td>
<td>68.0%</td>
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<tr>
<td></td>
<td>400 and 800 mg/</td>
<td></td>
<td>PBO=100</td>
<td></td>
<td>18.7%</td>
<td>16.7%</td>
<td>78.1%</td>
</tr>
<tr>
<td></td>
<td>starting with 800</td>
<td></td>
<td>ESL 800 mg=96</td>
<td></td>
<td>2.6% (p&lt;0.001)</td>
<td>4.0% (p&lt;0.001)</td>
<td>83.2%</td>
</tr>
<tr>
<td></td>
<td>mg in the 1,200 mg arm</td>
<td></td>
<td>ESL 800 mg=101</td>
<td></td>
<td>32.8% (p&lt;0.001)</td>
<td>37.1% (p&lt;0.01)</td>
<td>79.6%</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>ESL 1,200 mg=98</td>
<td></td>
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<tr>
<td>BIA-2093-30316 1–2</td>
<td>Starting with half</td>
<td>At 2 weeks: half of the maintenance dose; at 2 weeks: 0 mg (PBO)</td>
<td>N=252</td>
<td>17.0%</td>
<td>22.6</td>
<td>39.1%</td>
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<tr>
<td></td>
<td>of the assigned maintenance dose</td>
<td></td>
<td>PBO=87</td>
<td></td>
<td>37.9% (p&lt;0.05)</td>
<td>34.5</td>
<td>52.9%</td>
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<td></td>
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<td>ESL 800 mg=85</td>
<td></td>
<td>41.9% (p&lt;0.05)</td>
<td>37.7 (p&lt;0.05)</td>
<td>61.3%</td>
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<td></td>
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<td>ESL 1,200 mg=80</td>
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<tr>
<td>BIA-2093-30417 1–2</td>
<td>At 400 mg weekly steps</td>
<td>At 400 mg weekly steps until 400 mg</td>
<td>N=650</td>
<td>21.8%</td>
<td>23.1%</td>
<td>55.8%</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>PBO=224</td>
<td></td>
<td>29.7%</td>
<td>30.5%</td>
<td>67.1%</td>
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<tr>
<td></td>
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<td></td>
<td>ESL 800 mg=216</td>
<td></td>
<td>35.6% (p&lt;0.05)</td>
<td>42.6% (p&lt;0.001)</td>
<td>77.6%</td>
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<td></td>
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<td>ESL 1,200 mg=210</td>
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</table>

**Abbreviations:** AED, antiepileptic drug; PBO, placebo; ESL, eslicarbazepine acetate; SSF, standardized seizure frequency; TEAEs, treatment-emergent adverse events.
normal at baseline to low (i.e., <135 mEq/L) at the end of the maintenance treatment phase was found in 1.8% of patients treated with placebo, and in 6.1%, 4.8% and 6.6% of patients treated with ESL 400, 800 and 1,200 mg daily, respectively; hyponatremia leading to treatment discontinuation occurred in less than 1% of the patients taking ESL.18

The incidence of TEAEs was lower in patients who initiated ESL at 400 mg/day, followed by 400 mg increments, than in those starting treatment at higher dosages.18 Changes in mean laboratory parameters were not associated with clinically relevant findings, and there were no variations in vital signs, body weight or electrocardiography of clinical concern.

Open-label extension and real-world studies

Two 1-year, open-label extension studies19,20 including patients who completed the randomized pivotal Phase III trials14,15 were performed. The starting ESL daily dose was 800 mg, which could be individualized thereafter within the 400–1,200 mg range, while keeping stable the dosages of concomitant AEDs. Overall, these studies demonstrated sustained therapeutic effect and favorable long-term safety and tolerability of once-daily adjunctive ESL. A total of 462 patients concluded the 1 year of treatment; compared to the baseline period of the corresponding double-blind trial, the median seizure frequency decreased by 32–39% in weeks 1–4, and between 37% and 56% thereafter. The responder rate was 37–41% during the first 4 weeks, and then ranged from 38% to 53% per 12-week interval. The proportion of seizure-free patients per 12-week interval ranged between 5.0% and 12.5%. Adverse events were reported by 51–83% of the patients; the most common adverse events were dizziness and headache and they were usually of mild to moderate intensity. Significant improvements in quality of life domains and depressive symptoms were also observed.

Different prospective and retrospective real-world studies have been performed after ESL’s authorization as adjunctive treatment in FOS;21–30 they have provided reassuring evidence of its efficacy and safety in routine clinical care and across different baseline patient conditions, and have complemented data from clinical trials. The characteristics and findings of the main real-life studies are summarized in Table 2.

<table>
<thead>
<tr>
<th>Study</th>
<th>Study design</th>
<th>Patient characteristics</th>
<th>Main outcome measures</th>
<th>Authors’ conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Euro-Esli21</td>
<td>Pooled analysis of 14 European, prospective and retrospective clinical audit studies6</td>
<td>N=2,058</td>
<td>Retention rate: 73.4% (12 months)</td>
<td>Euro-Esli is the largest ESL clinical practice study available to date.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Male: 52.1%</td>
<td>Responder rate: 75.6% (12 months)</td>
<td>It provides strong and reassuring evidence of ESL’s safety profile. No unexpected safety signals emerged over a median duration of follow-up of &gt;5 years.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Age: 44.0 (15) years</td>
<td>Seizure freedom: 41.3% (12 months)</td>
<td>ESL showed good retention rates, elicited a significant reduction in seizure frequency and was well tolerated when used in clinical practice.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Epilepsy duration: 20.9 (16.4) years</td>
<td>AEs: 34.0%</td>
<td>The administration of ESL after the first monotherapy failure was associated with an optimal seizure control and tolerability profile. Over half of the patients were converted to ESL monotherapy during follow-up.</td>
</tr>
<tr>
<td>ESLADOBA22</td>
<td>Prospective, multicenter (Portugal); ESL as add-on therapy to one AED</td>
<td>N=52</td>
<td>Retention rate: 73.0% (6–9 months)</td>
<td>Seizure freedom: 39.5% (months)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Male: 48.1%</td>
<td>Responder rate: 71.1% (6–9 months)</td>
<td>AEs: 23.1%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Age: 41.5 (13.3) years</td>
<td>Seizure freedom due to AEs: 9.6%</td>
<td>Discontinuation due to AEs: 6.7%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Epilepsy duration: 18.5 (14.8) years</td>
<td>Reduction in Clinical Global Impression of Severity: 42.1%</td>
<td>Most frequent AEs: dizziness (5.4%), somnolence (5.1%)</td>
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<tr>
<td></td>
<td></td>
<td>Baseline monthly seizure frequency: 13.6 (49.9)</td>
<td></td>
<td>ESL more effective in patients aged &lt;65 years, not receiving treatment with other sodium channel blockers, receiving &lt;2 concomitant AEDs</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Number of concomitant AEDs: 1.7 (1.0)</td>
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<tr>
<td>EARLY-ESLI23</td>
<td>Retrospective, multicenter (Spain), 1-year observational; ESL as add-on after first monotherapy failure</td>
<td>N=253</td>
<td>Retention rate: 92.9% (12 months)</td>
<td>The administration of ESL after the first monotherapy failure was associated with an optimal seizure control and tolerability profile. Over half of the patients were converted to ESL monotherapy during follow-up.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Male: 57.3%</td>
<td>Responder rate: 70.3% (3 months), 79.8% (6 months), 82.5% (12 months)</td>
<td>Seizure freedom: 37.3% (12 months)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Age: 48.7 (18–87) years</td>
<td>AEs: 31.6%</td>
<td>Discontinuation due to AEs: 3.6%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Epilepsy duration: 3 (1–10) years</td>
<td>Most common AEs: somnolence (8.7%), dizziness (5.1%)</td>
<td>After starting ESL, 54.2% patients withdrew the prior monotherapy and converted to ESL alone</td>
</tr>
<tr>
<td></td>
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<td>Baseline monthly seizure frequency: 2.9 (16.1)</td>
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<td></td>
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<td>Final median ESL dose: 800 mg/day</td>
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</table>

(Continued)
**Table 2 (Continued)**

<table>
<thead>
<tr>
<th>Study</th>
<th>Study design</th>
<th>Patient characteristics</th>
<th>Main outcome measures</th>
<th>Authors’ conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>EPOS [26,27]</td>
<td>Prospective, multicenter (Europe); ESL as add-on therapy to one AED</td>
<td>N=219 Male: 57.5%, Age: 43 (18–83) years Epilepsy duration: 12.3 (0–57.3) years Seizure frequency over 3 baseline months: 29.3 (95.8), Non-responder to CBZ: N=45, Age: &gt;60 yrs N=41 ESL 800 mg/day: 74.3%, Target dose achieved with one titration step: 79.3%</td>
<td>Retention rate: 89.0% (3 months), 82.2% (6 months) Responder rate: 69.9% (3 months), 81.8% (6 months) Seizure freedom: 25.9% (3 months), 39.2% (6 months) AEs: 26.0% Discontinuation due to AEs: 11.4% Most frequent AEs: dizziness (4.6%), headache (3.2%), convulsion (3.2%), fatigue (2.7%) Decrease (i.e., improvement) of QOLiE-10 from 2.9 (baseline) to 2.4 (3 months) and 2.1 (6 months)</td>
<td>Adjunctive ESL therapy to one AED showed favorable retention, seizure control and tolerability, and improvement in quality of life. ESL in patients less refractory than those included in clinical trials led to higher responder and seizure freedom rates. ESL was effective and generally well tolerated in elderly patients and in those who had previously not responded to CBZ therapy.</td>
</tr>
<tr>
<td>Correia et al [28]</td>
<td>Retrospective, single-center (Portugal), 2-year observational</td>
<td>N=152 Male: 50.6%, Age: 38.5 (14.2) years Epilepsy duration: 26.8 (13.1) years Baseline monthly seizure frequency: 19.7 (32.9), Two or more AEDs at baseline: 57.9%</td>
<td>Retention rate: 82.9% (6 months), 71.3% (12 months), 65.1% (18 months), 62.8% (24 months) Responder rate: 25.7% (6 months), 19.0% (18 months), 17.1% (24 months) AEs: 42.1% Discontinuation due to AEs: 21.1% Most frequent AEs: dizziness, somnolence AEs more common in regimens with CBZ</td>
<td>ESL appears to be a clinically useful add-on AED, with good safety profile and high retention rates throughout 2 years, even in a very refractory group of patients.</td>
</tr>
<tr>
<td>ESLIBASE [28]</td>
<td>Retrospective, multicenter (Spain), non-interventional</td>
<td>N=327 Male: 48.0%, Age: 41.9 (14–87) years Two or more AEDs at baseline: 78% Median ESL daily dose: 800 mg (3 months), 1,200 mg (6 and 12 months)</td>
<td>Retention rate: 89.3% (3 months), 80.1% (6 months), 72.5% (12 months) Responder rate: 46.3% (3 months), 57.9% (6 months), 52.3% (12 months) Seizure freedom: 21.0% (3 months), 28.0% (6 months), 25.3% (12 months) AEs: 40.7% (12 months) Discontinuation due to AEs: 16.2% (12 months) Most common AEs: dizziness/nausea (11.3%), somnolence (6.1%), ataxia (5.1%)</td>
<td>ESL was well tolerated and effective over 1 year. The responder rate increased when ESL was combined with a non-sodium channel-targeting drug. Tolerability profile improved in &gt;50% of the patients switching from OXC/CBZ to ESL due to AEs.</td>
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<td>Massot et al [29]</td>
<td>Observational, single-center (Spain), descriptive, cross-sectional</td>
<td>N=61 Male: 31.5%, Age: 43.6 (14.4) years Baseline monthly seizure frequency: 5.5 (1.13–30), ESL 800 mg/day: 42.6%, Mean follow-up duration: 4.7 (3.2) months</td>
<td>Retention rate: 75.4% (3 months) Reduction of monthly median seizure frequency by 63.6% AEs: 57.4% Most common AE: dizziness (34.4%) Twelve patients (19.7%) switched overnight to ESL from OXC (ratio 1:1), and 13 patients (21.3%) switched overnight from CBZ to ESL (ratio 1:1.3). At 3 months after the transition, median frequency decreased by 20%.</td>
<td>ESL is effective in the treatment of focal epilepsies and its early retention rate is &gt;70%. AEs mostly occurred during the titration phase.</td>
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<td>Serrano-Castro et al [30]</td>
<td>Observational single-center (Spain)</td>
<td>N=105 Male: 51.4%</td>
<td>Responder rate: 58.4% (6 months) Seizure freedom: 20.7% (6 months) AEs: 18.1% (6 months) Most common AE: cognitive disorders. Treatment discontinuation: 11.5%</td>
<td>ESL is well tolerated and effective as an add-on therapy with most of the AEDs. Add-on to lacosamide was less effective; association with other sodium channel inhibitors had similar efficacy than others combinations with other AEDs.</td>
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**Notes:** *Including the ESLADOBA, Early-Esli, EPOS, Correia et al, ESLIBASE and Massot et al studies. Data are presented as mean (standard deviation) or median (interquartile range).*

**Abbreviations:** AE, adverse event; AED, antiepileptic drug; CBZ, carbamazepine; ESL, eslicarbazepine acetate; OXC, oxcarbazepine; QOLiE-10, Quality of Life in Epilepsy Inventory-10.
Clinical use
Renal and hepatic impairment
ESL metabolites are eliminated from the systemic circulation primarily by renal excretion, and clearance is dependent on kidney function in patients with mild to severe renal impairment. Dose adjustment is not required if creatinine clearance (CrCl) is >60 mL/min; in patients with CrCl between 30 and 60 mL/min, the initial ESL dose should be 200 mg once daily or 400 mg every other day for 2 weeks, followed by a once-daily dose of 400 mg, which may be increased up to 600 mg/day on the basis of individual response. ESL is not recommended in patients with severe renal impairment (CrCl<30 mL/min) due to insufficient data. Hemodialysis partially removes ESL and its metabolites from plasma. The liver function has less critical effects on the pharmacokinetics of ESL, and patients with mild to moderate hepatic impairment do not need dosage adjustment. Conversely, the drug has not been evaluated in the presence of severe liver dysfunction, and its use is not recommended in this setting.9

Drug interactions
An increase in the dose of the medicinal products that are mainly metabolized through CYP3A4 (e.g., simvastatin) or eliminated by conjugation through the UDP-glucuronyl transferases may be required when co-administered with ESL due to its inducer effect, albeit weaker than CBZ. Eslicarbazepine has inhibiting properties with respect to CYP2C19, and interactions can arise when ESL is taken with drugs that are substrates of this enzyme (e.g., diazepam).

Oral anticoagulants and digoxin
ESL at the 1,200 mg daily dose decreases plasma exposure to S-warfarin by 23%, without significant effects on the R-warfarin pharmacokinetics or coagulation: due to interindividual variability in the interaction, it is, however, advisable to carefully monitor the international normalized ratio during the first weeks after initiation or ending concomitant treatment of warfarin and ESL. No studies have specifically assessed whether pharmacological interactions exist between ESL and the direct oral anticoagulants. No influence of ESL on digoxin pharmacokinetics has been observed, suggesting the lack of meaningful interactions with the transporter P-glycoprotein.9

Antiepileptic drugs
The co-administration of ESL and CBZ results in an average decrease of 32% in exposure to eslicarbazepine, most likely caused by an induction of glucuronidation; this may require an increase of ESL dose. Remarkably, the concomitant treatment with ESL and CBZ can increase the risk to develop diplopia, abnormal coordination and dizziness.

The concomitant use of ESL and phenytoin determines a reduction of one-third in exposure to the active metabolite eslicarbazepine, due to an induction of glucuronidation, and an increase of 31–35% in exposure to phenytoin, caused by CYP2C19 inhibition. Accordingly, based upon individual response, the dose of ESL may need to be increased and the dose of phenytoin to be decreased.

Data from healthy subjects revealed minor pharmacokinetic interactions between ESL and lamotrigine (15% decrease in exposure to lamotrigine) or topiramate (18% decrease in exposure to topiramate). ESL does not appear to interact significantly with levetiracetam and valproate.9 Notably, concomitant use of ESL with OXC is not recommended since it may cause overexposure to the active metabolites.

Oral contraceptives
Eslicarbazepine may decrease the effectiveness of hormonal contraception: the administration of ESL (1,200 mg/day) to female subjects using combined oral contraceptives showed an average decrease in systemic exposure to levonorgestrel of 37% and to ethinylestradiol of 42% owing to the CYP3A4 induction. Therefore, to avoid inadvertent pregnancy, women of childbearing potential may need an increase in oral contraceptive dose or use alternative methods of contraception during ESL treatment and up to the end of the current menstrual cycle after treatment has been stopped.9

Fertility, pregnancy and lactation
Epilepsy is a common neurologic disorder and the majority of affected people are expected to participate fully in life experiences, including childbearing. The prevalence of epilepsy among pregnant women is up to 0.7%, and from three to five births per thousand will be to women with epilepsy.11,12 Worldwide registries and observational studies have provided consistent findings on the risks related to both epilepsy itself and many of the older antiepileptic medicinal products, while there remains a large gap in the knowledge of most of the newer ones, including ESL.33 In studies in mice, impairment of fertility has been shown after treatment with ESL, like a decrease in implantations and live embryos. ESL has no teratogenic effects in the rat or rabbit, but induced skeletal abnormalities in the mouse. Ossification delays, reduced fetal weights, increases in minor skeletal and visceral anomalies
were observed at maternal toxic doses in embryo-toxicity studies in mice, rats and rabbits. Additionally, animal studies have shown excretion of eslicarbazepine in breast milk. To date, there are insufficient data to allow conclusions about the safety of ESL in pregnant women, and it is unknown whether it is excreted in human milk. Accordingly, specialist advice should be given to women who become or plan to become pregnant while receiving ESL.

**Elderly population (over 65 years of age)**

A post-approval commitment, multicenter, open-label, non-controlled, single-arm study (BIA-2093-401 [NCT01422720]) with flexible dose of ESL between 400 and 1,200 mg daily was performed to address the safety and efficacy of ESL as adjunctive treatment in the elderly. Among 72 patients treated with one or two concomitant AEDs, the ESL dose did not exceed 800 mg/day in the majority of the cases. The treatment was associated with a 54% relative reduction in the standardized baseline seizure frequency and did not raise major safety concerns. The overall frequency of TEAEs (65.3%) was similar to the rate observed in the pooled Phase III studies, and the most common were dizziness (12.5%) and somnolence (9.7%); notably, the incidence of hyponatremia (8.3%) was slightly higher than the proportion observed in the randomized clinical trials. Overall, 25% of the subjects discontinued prematurely due to side effects.

A retrospective survey that included elderly patients with focal seizures who started ESL in 12 Spanish hospitals, according to real-life experience between 2010 and 2012, was reported as part of the ESLIBASE study. In all, 29 patients with a mean age of 71.2 years were included, of whom 18 were pharmacoresistant. At 1 year, the mean ESL daily dose was 850 mg and the retention rate was 69%. Sixty-two percent of the patients were seizure responders, and 24.1% were free from seizures. Adverse effects were experienced by 55.2% of the study cohort and led to treatment discontinuation in seven (24.1%) cases. The most common side effects were dizziness, nausea and ataxia; the tolerability profile improved in most of the patients who switched from CBZ or OXC to ESL. In a single-center, retrospective study of patients treated with ESL, there was no significant difference in the prevalence of mild adverse effects between young and elderly (≥60 years old) adults, whereas adverse effects leading to treatment discontinuation were more frequent in the elderly group. Notably, the pharmacokinetic profile of ESL is unaffected in the elderly and no dose adjustment is needed, provided that renal function is preserved (CrCl >60 mL/min).

**Transitioning from OXC or CBZ to ESL**

The switch from OXC to ESL is easy to perform: a dose ratio of 1:1 should be employed to estimate the ESL target dose, transition can take place overnight and no changes to co-medication are required. Conversely, the change from CBZ to ESL is less straightforward and requires careful considerations on a patient-by-patient basis: a dose ratio of 1:1.3 should be considered, the conversion period should last a minimum of 1–2 weeks, although longer intervals may be advised, and concomitant medications metabolized by cytochrome enzymes may require dose adjustment.

Transition to ESL might be appropriate in patients who experience or are at risk to develop CBZ- or OXC-related adverse events, including metabolic disorders resulting from enzyme induction like hypercholesterolemia, osteoporosis or sexual dysfunction, in patients who are poorly compliant with two- or three-daily dosing regimen, frequently forget to take their medication or work rotating shifts, and in patients who are poly-medicated or affected by cognitive dysfunction or liver disease. Notably, epilepsy is common among patients with cognitive impairment and dementia, and the good profile of ESL with respect to neurocognitive functioning can make it a preferable option over AEDs, like CBZ, characterized by deeper effects on psychomotor performance. In this regard, a recent network meta-analysis has compared the tolerability of the new-generation sodium blocker AEDs in drug-resistant epilepsies from double-blind, placebo-controlled trials. At the highest recommended doses, treatment withdrawal was more frequently observed in patients treated with OXC than with ESL and lacosamide, and vestibulocerebellar side effects including abnormal coordination, ataxia and diplopia were more common among patients taking OXC rather than ESL.

Overall, ESL has been shown to be less frequently associated with cutaneous reactions in comparison to CBZ, and the lack of a toxic epoxide derivative could reduce the risk of developing rash. In patients known to be positive for HLA-B*1502 or HLA-A*3101, medications other than CBZ should be used and, due to possible cross-reactivity, CBZ-related compounds such as OXC should also be avoided. In this respect, the official summary of product characteristics contraindicates ESL in patients with hypersensitivity reactions to CBZ. A recent report described the case of a patient with the HLA-A*31:01 haplotype who, after the development of a severe cutaneous reaction following CBZ administration, was successfully prescribed ESL without experiencing any adverse event, and it suggested that ESL may be considered if the benefits are thought to exceed the risks. The safety of
ESL in subjects with HLA-B*15:02 or HLA-A*31:01 haplotypes and a history of CBZ- or OXC-induced hypersensitivity reactions remain, however, to be fully addressed in larger studies before drawing definitive conclusions.

The overnight OXC-ESL transition resulted in improvements in side effects, quality of life and alertness, while maintaining seizure control. In addition, compared with the first- and second-generation dibenzazepine as well as the traditional inducer AEDs, ESL may be preferred in patients at higher vascular risk by exhibiting a safer profile according to lipid metabolism and atherosclerosis development.

Apart from the selective interaction with the inactive state of the VGSC and the reduction of their availability by interfering with the slow inactivation pathway, distinctive properties of ESL include the blockade of high- and low-affinity hCaV3.2 inward currents with a 10–60-fold higher potency than CBZ, the lack of effect upon KV7.2 outward currents and the block of T-type CaV3.2 channels. Moreover, while CBZ and OXC show pro-epileptic actions at clinically relevant concentrations through the enhancement of the excitatory post-synaptic transmission, ESL has no effects on post-synaptic currents, explaining its lack of seizure exacerbation. ESL may not only suppress seizure activity, but also inhibit the generation of a hyperexcitable network, and it can overcome cellular mechanisms of resistance to sodium-channel acting AEDs. In this respect, a real-world clinical experience has suggested that ESL may represent an effective therapeutic option even in those patients who have previously tried, but discontinued the older carboxamides, regardless of the reasons for discontinuation.

**Conclusion**

Once-daily ESL at the doses of 800 and 1,200 mg showed consistent results across efficacy and safety endpoints both in randomized-controlled and open-label studies. Notably, starting treatment at 400 mg/day, followed by 400 mg increments every 7–14 days up to the optimal dose, could maximize the balance between seizure control and tolerability.

Despite the lack of direct comparisons, the distinctive chemical structure of ESL compared to the older members of the dibenzazepine family, CBZ and OXC, can convey it unique advantages, including the once-daily regimen, the lower potential for drug-drug interactions and the more favorable tolerability profile, while enjoying similar efficacy. Additionally, ESL has shown the potential to be effective in syndromes characterized by high refractoriness to drug treatment, like the mesial temporal lobe epilepsy associated with hippocampal sclerosis.

In conclusion, ESL represents an interesting improvement on a classic drug family for the treatment of FOS, and it has the potential to enhance the quality of life for drug-refractory patients.

**Disclosure**

FB acted as consultant for Eisai. GZ has received speaker or consultancy fees from Eisai, Sanofi-Aventis and UCB Pharma. The authors report no other conflicts of interest in this work.

**References**


