

Efficacy and Tolerability of RAP-219, a Potential First-in-Class Negative Allosteric Modulator of γ 8 Transmembrane AMPA Receptor Regulatory Protein (TARPy8): Impact on RNS Long Episodes and Focal Seizures

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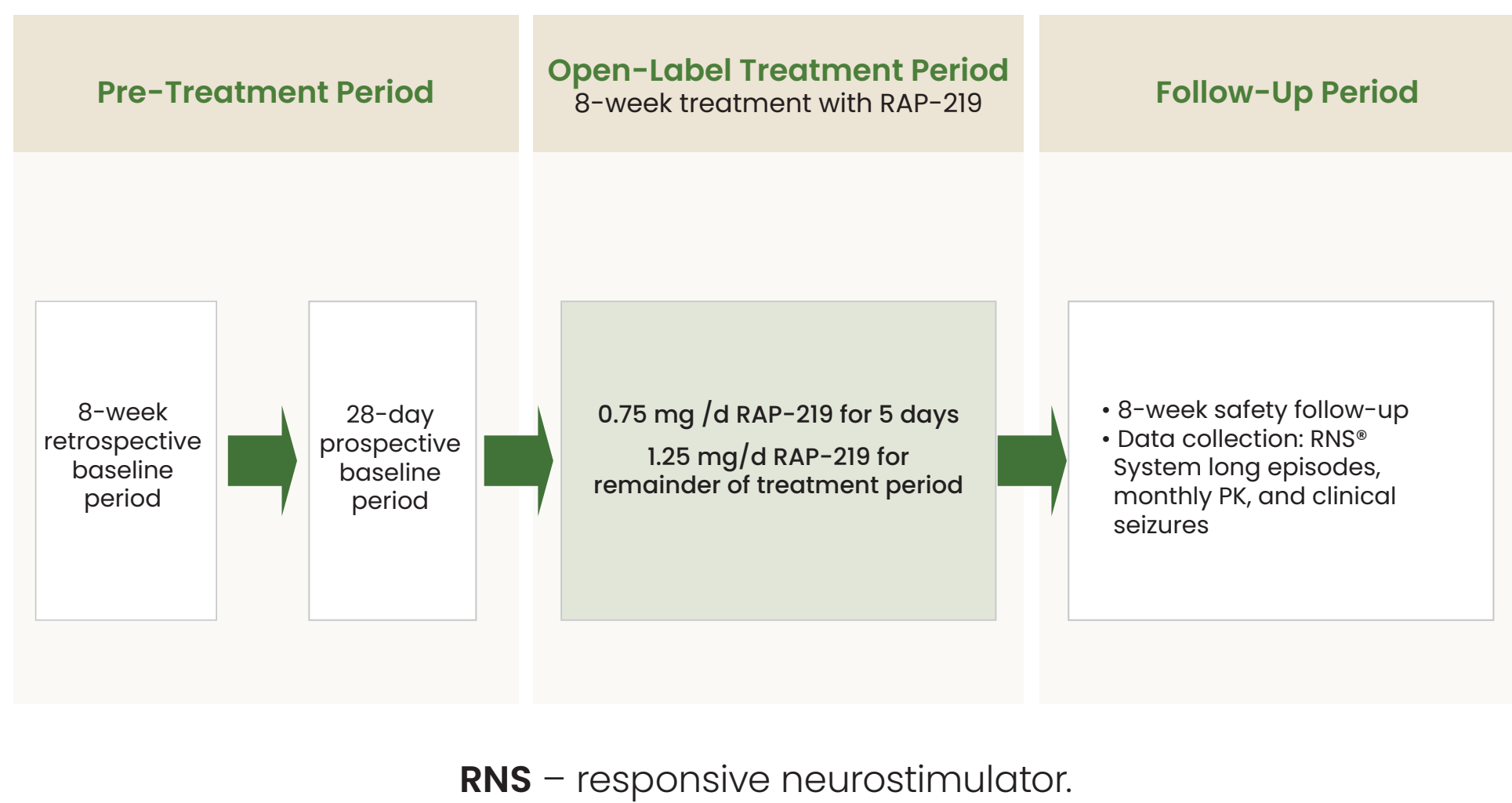
Background

- New antiseizure medications (ASMs) for drug-resistant focal epilepsy with novel mechanisms of action and improved tolerability are needed
- γ 8 transmembrane AMPA receptor regulatory protein (TARPy8) is highly expressed in the neocortex and mesial temporal lobe (MTL), regions of the brain where nearly all focal onset seizures (FOS) originate¹⁻³
- RAP-219, a selective and potent negative allosteric modulator of TARPy8, offers a precision approach for the treatment of focal epilepsy via targeted AMPA receptor inhibition
- The responsive neurostimulator (RNS® System, NeuroPace) continually monitors electrocorticographic intracranial EEG activity and responds to abnormal activity⁴
 - Long episodes (LEs), measured by the RNS System, are organized epileptiform activity exceeding a clinician-specified duration
 - LEs often represent electrographic seizures and may potentially supplement seizure diaries⁵
 - Recent analyses suggest that a $\geq 30\%$ reduction in LEs is associated with a $\geq 50\%$ reduction in clinical seizures (CSs)⁶
- In this novel Phase 2 proof-of-concept (POC) study, the effect of RAP-219 in patients with drug-resistant FOS and an implanted RNS System was assessed
 - Key outcomes were change in LEs and CSs

Methods

- Adults (18–65 y) with drug-resistant focal epilepsy and an implanted RNS device with ≥ 1 lead implanted in the MTL were enrolled in this Phase 2 study
- LEs were recorded by an RNS System; CSs were recorded in seizure diaries

Figure 1. Study Design



RNS – responsive neurostimulator.

Table 1. Inclusion and Exclusion Criteria

Inclusion Criteria ^a	Exclusion Criteria ^b
- Adults 18–65 years old with drug-resistant focal epilepsy - RNS System device requirements: ≥ 1 electrode implanted within the MTL ≥ 15 months before screening - Stable device settings for ≥ 8 weeks before screening - Had ≥ 16 LEs and ≥ 1 CS during an 8-week retrospective analysis period $> 50\%$ electrographic seizure and LE concordance^c ≤ 4 concomitant ASMs with stable dose for ≥ 8 weeks before screening	- Epilepsy surgery ≤ 12 months before screening - Use of > 4 concomitant ASMs - Use of perampanel ≤ 12 weeks before screening - Ketogenic diet with any regimen changes within 12 weeks before screening - Stable ketogenic diet allowed

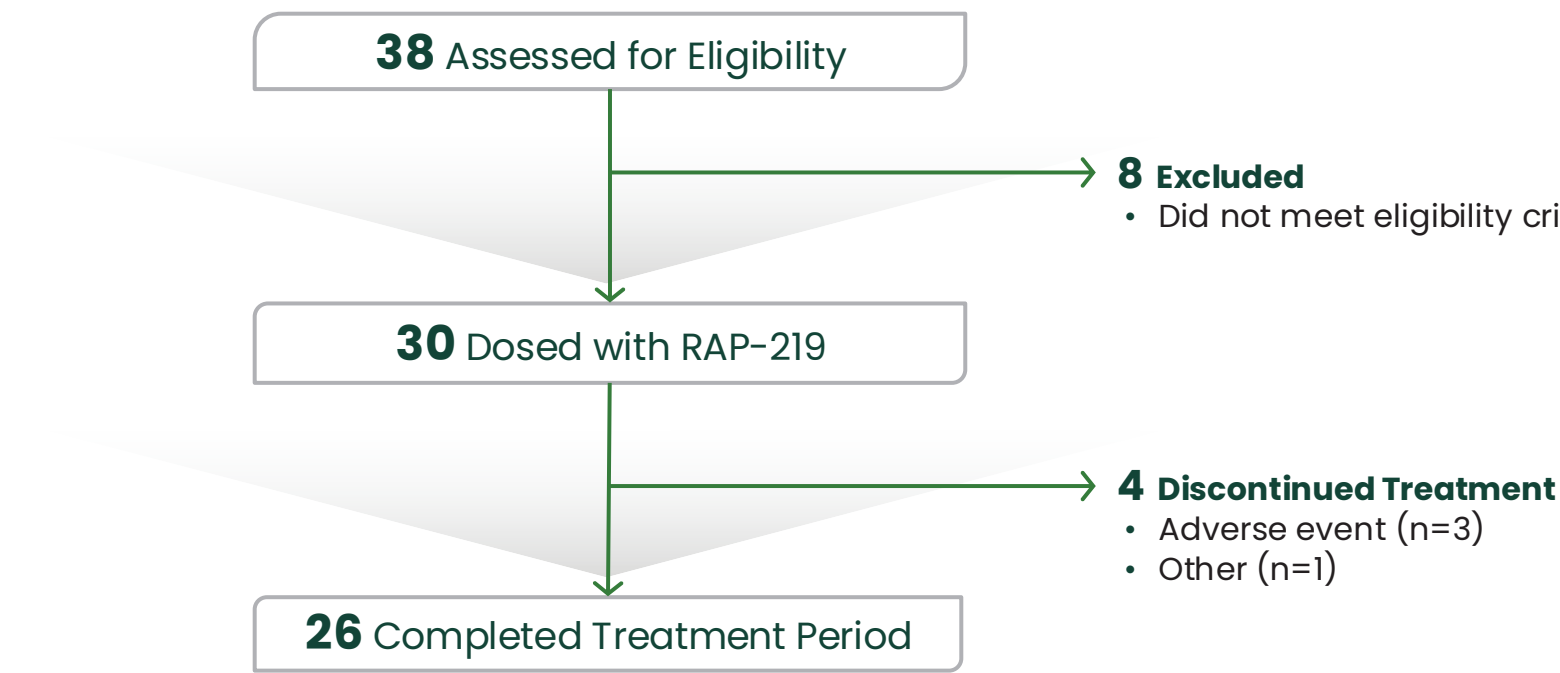
^aDoes not include all inclusion criteria. ^bDoes not include all exclusion criteria. ^cEvaluated by the Central Epileptologist Review Team, consisting of a panel of epileptologists from the ESC. ASM – antiseizure medication; CS – clinical seizure; ESC – Epilepsy Study Consortium; LE – long episode; MTL – mesial temporal lobe; RNS – responsive neurostimulator.

Outcomes

- Median percentage change in LE and CS frequency from baseline to Weeks 1–8
 - The combined 8-week retrospective and 4-week prospective periods served as the baseline for LEs
 - The 4-week prospective period served as the baseline for CSs
- Responder rates for LE and CS reductions, with LE thresholds of $\geq 30\%$, $\geq 50\%$, $\geq 75\%$, and 100% and CS thresholds of $\geq 50\%$, $\geq 75\%$, and 100%, were analyzed
- Incidence of AEs

Results

Figure 2. Patient Disposition



Analysis Population

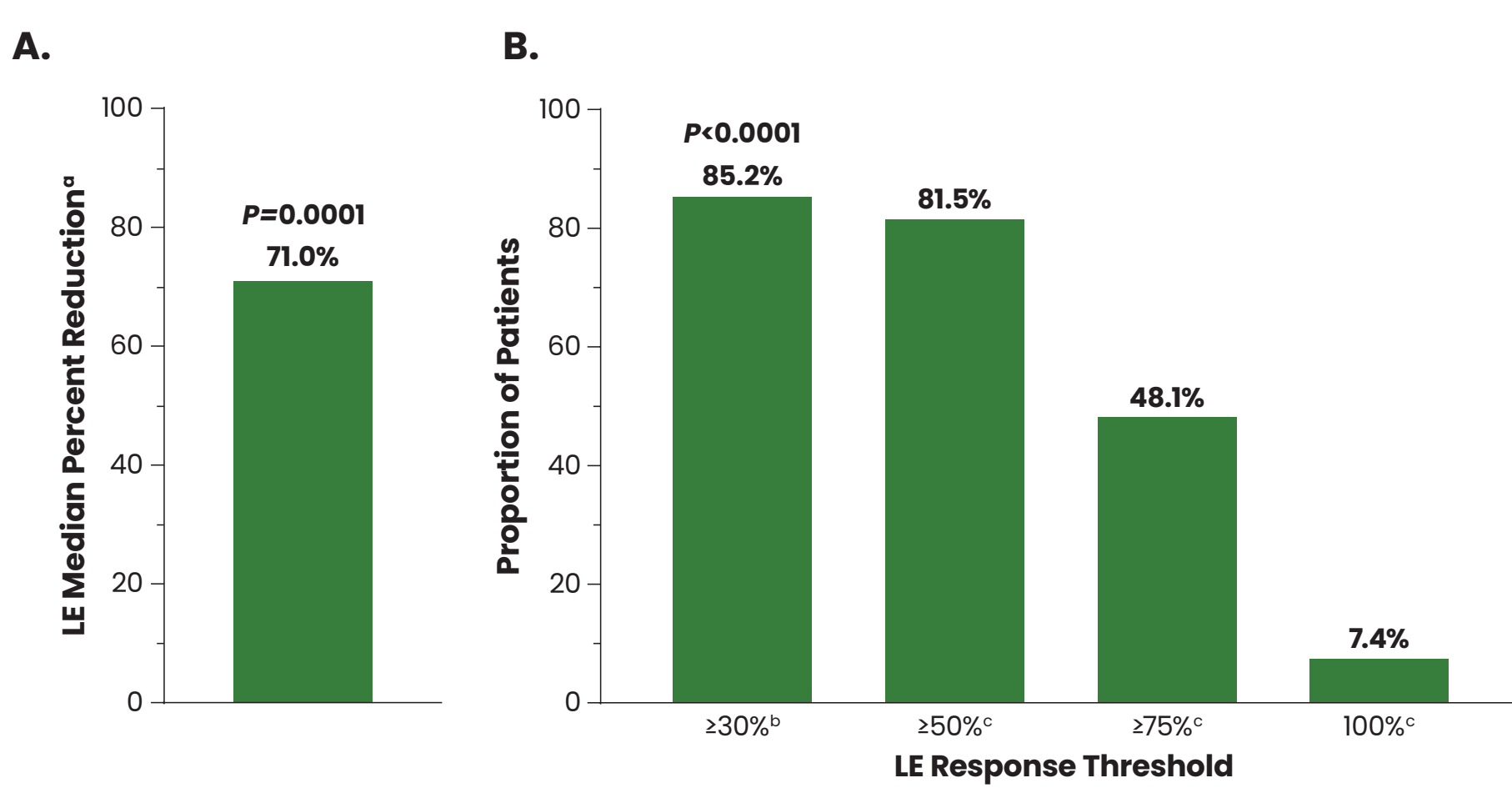
- Safety population: N=30
- Modified intent-to-treat (mITT) population (LE efficacy): N=27
 - Not included from the safety population: 2 patients with < 3 weeks of treatment; 1 patient with an RNS setting change
- mITT-CS population (CS efficacy): N=25
 - Not included from the mITT population: 2 patients with no CSs during the prospective baseline

Table 2. Demographics and Baseline Characteristics

	Safety Population N=30
Male, n (%)	18 (60)
Age at study entry, y, mean $\pm$ SD	40.1 $\pm$ 10.4
Age at first seizure, y, mean $\pm$ SD	15.8 $\pm$ 9.3
LE frequency per 28 days ^a , median (range)	48 (8–751.7) ^a
CS frequency per 28 days ^b , median (range)	10 (0.8–314.6) ^b
Number of concomitant ASMs, median (range)	3 (1–4)
1 concomitant ASM, n (%)	2 (7)
2 concomitant ASMs, n (%)	7 (23)
3 concomitant ASMs, n (%)	18 (60)
4 concomitant ASMs, n (%)	3 (10)
Most commonly used ASMs, n (%)	
Lamotrigine	15 (50)
Levetiracetam	12 (40)
Cenobamate	11 (37)
Zonisamide	9 (30)
Clobazam	7 (23)
Lacosamide	7 (23)
Years since RNS System implantation, median (range)	4.6 (2–11)
Concordance between LEs and electrographic seizures, median (range)	90 (30–100) ^c

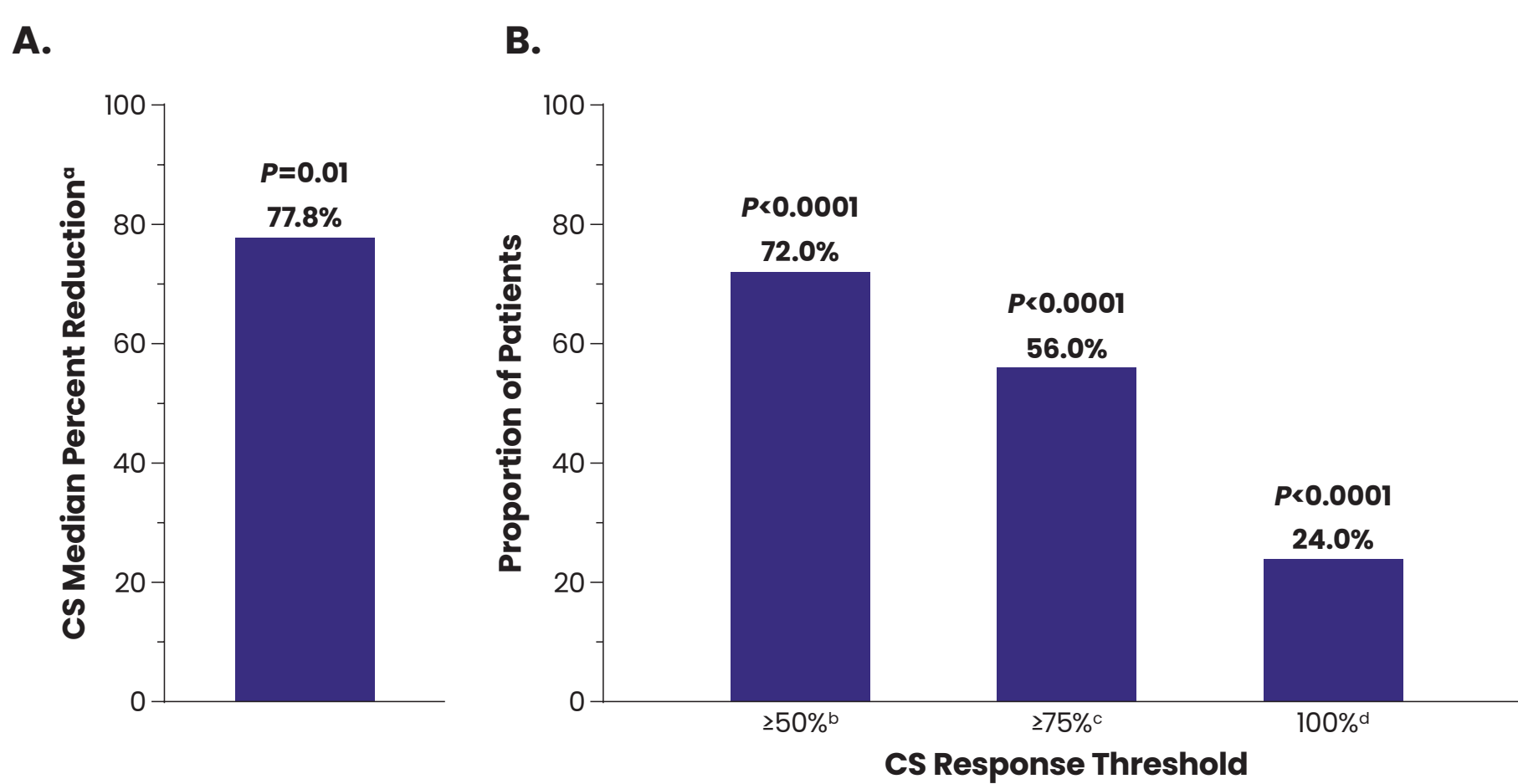
^amITT population (N=27). ^bmITT-CS population (N=25). ASM – antiseizure medication; CS – clinical seizure; LE – long episode; RNS – responsive neurostimulator; SD – standard deviation.

Figure 3. Weeks 1–8 A. Median LE Percent Reduction From Baseline and B. $\geq 30\%$, $\geq 50\%$, $\geq 75\%$, and 100% Responder Rates Following Treatment With RAP-219



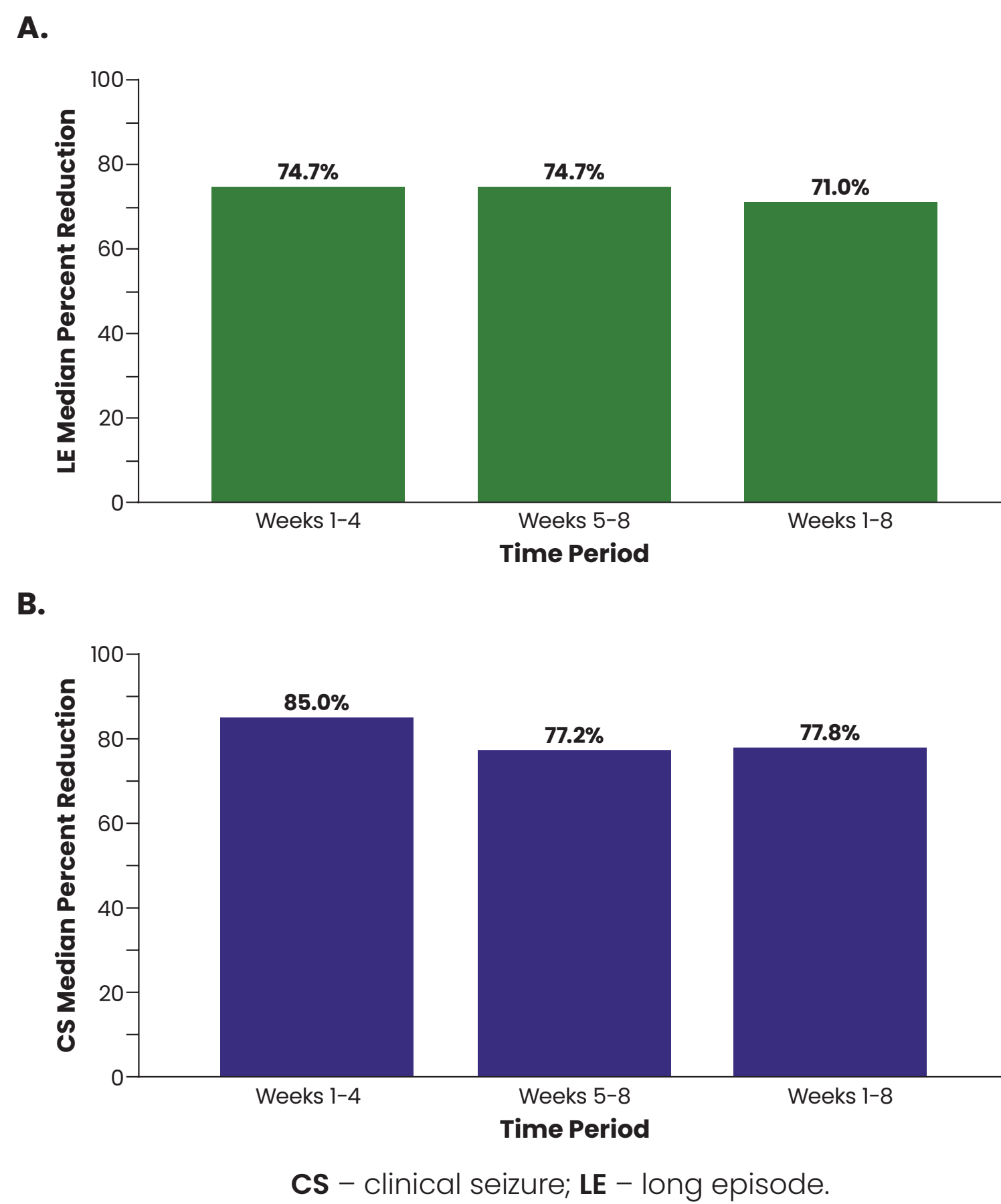
^aMedian percent change statistical comparison used the Wilcoxon signed rank test to determine if the percent change in LE was greater than 0%. ^bResponder analysis statistical comparison was based on a one-sample exact test to determine whether the proportion of responders was $> 10\%$. ^cStatistical comparisons were not made for other cut points. 95% confidence intervals for responder analysis were based on Clopper-Pearson exact binomial: $\geq 50\%$, (61.9, 93.7); $\geq 75\%$, (28.7, 68.1); 100%, (0.9, 24.3). LE – long episode.

Figure 4. Weeks 1–8 A. Median CS Percent Reduction From Baseline and B. $\geq 50\%$, $\geq 75\%$, and 100% Responder Rates Following Treatment With RAP-219



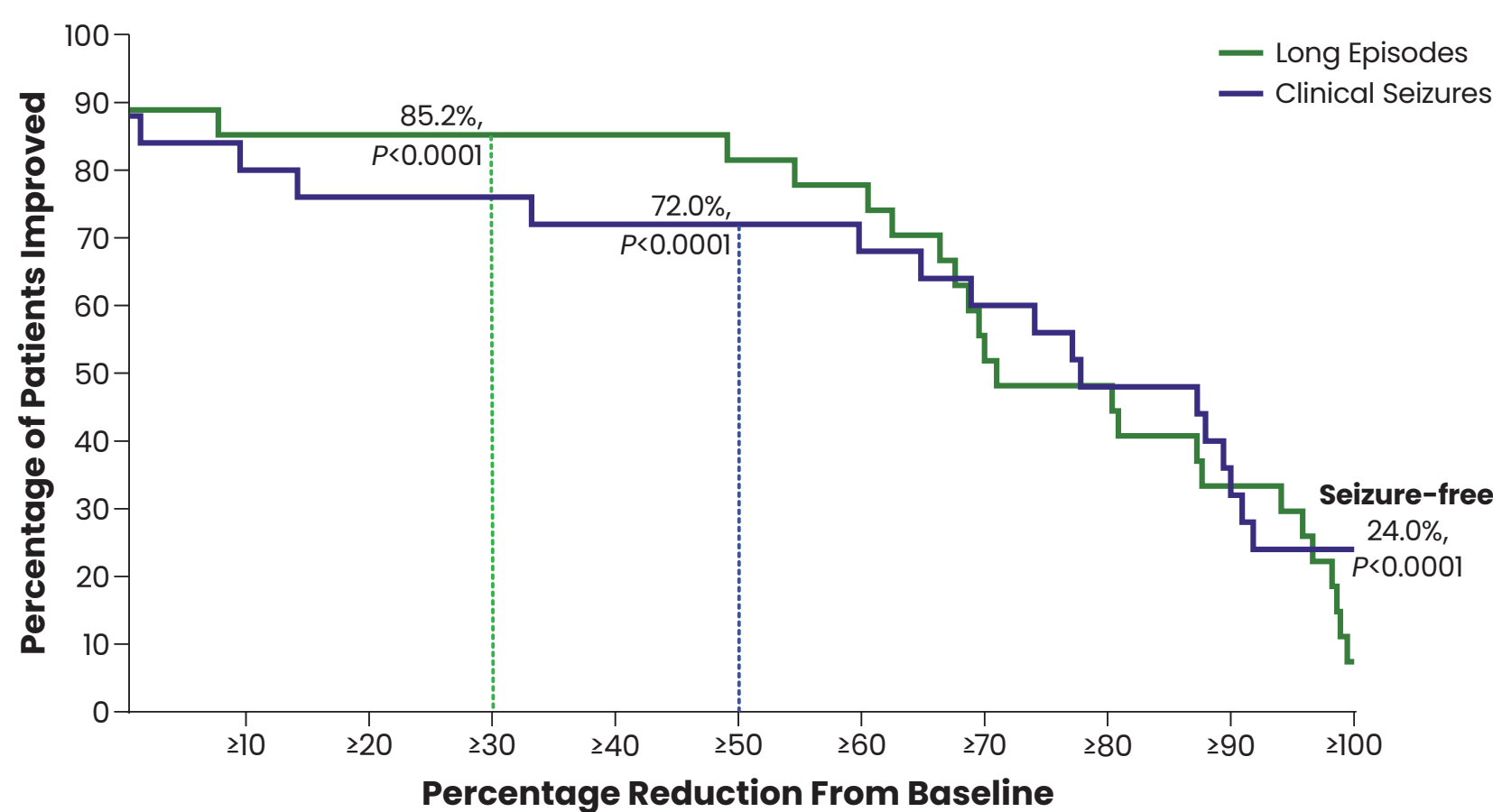
^aStatistical comparison for median percent change used the Wilcoxon signed-rank test to determine if the median percent reduction from baseline in CS was greater than 20%. ^{b,c,d}Responder analysis statistical comparison was based on a one sample exact test to determine whether the proportion of responders was: (b) $> 20\%$; (c) $> 7\%$; (d) $> 1.5\%$ for the $\geq 50\%$ responder, $\geq 75\%$ responder, and 100% (seizure freedom) groups, respectively. CS – clinical seizure.

Figure 5. Effect of RAP-219 Observed During Weeks 1–4 and 5–8 on A. LEs and B. CSs



CS – clinical seizure; LE – long episode.

Figure 6. Cumulative Response for Long Episodes and Clinical Seizures



Safety

Table 3. Treatment-Emergent Adverse Events (Weeks 1–8)

	Safety Population (N=30)
Any TEAE, n (%)	25 (83)
TEAE leading to study drug discontinuation ^a	3 (10)
Grade 1 TEAE (mild)	15 (50)
Grade 2 TEAE (moderate)	10 (33)
Grade ≥ 3 TEAE (severe)	0
TEAEs reported in $\geq 10\%$ of patients, n (%)	
Dizziness	8 (27)
Headache	6 (20)
Fatigue	4 (13)
Fall	3 (10)
Nausea	3 (10)
Somnolence	3 (10)

^aTEAEs leading to study discontinuation: worsening of preexisting memory impairment (Grade 1); panic attack (Grade 1); worsening of preexisting anxiety (Grade 2). TEAE – treatment-emergent adverse event.

Conclusions

- Treatment with RAP-219 provided statistically significant and clinically meaningful improvement in LE frequency (objective biomarker) and CS frequency
 - Improvements were consistent across the entire treatment period
- Clinically meaningful improvements in LE and CS frequencies were observed in 85% and 72% of patients, respectively (Figure 6)
 - Complete seizure freedom over the entire treatment period (Titration + Maintenance) was achieved by 24% of patients
- RAP-219 treatment was generally well tolerated, with a 10% discontinuation rate secondary to TEAEs and no severe or serious AEs or clinically significant laboratory, vital signs, or ECG abnormalities noted during the treatment period
- These results support advancing RAP-219 into Phase 3 studies

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