

#### Changing Agendas on Sleep, Treatment and Learning in Epilepsy (CASTLE) Sleep-E: A protocol for a randomised controlled trial comparing an online behavioural sleep intervention with standard care in children with Rolandic epilepsy

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# TITLE

Changing Agendas on Sleep, Treatment and Learning in Epilepsy (CASTLE) Sleep-E: A protocol for a randomised controlled trial comparing an online behavioural sleep intervention with standard care in children with Rolandic epilepsy

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## ABSTRACT

**Introduction** Sleep and epilepsy have an established bi-directional relationship yet only one randomised controlled clinical trial has assessed the effectiveness of behavioural sleep interventions for children with epilepsy. The intervention was successful, but was delivered via face-to-face educational sessions with parents, which are costly and non-scalable to population level. The Changing Agendas on Sleep, Treatment and Learning in Epilepsy (CASTLE) Sleep-E trial addresses this problem by comparing clinical- and cost-effectiveness in children with Rolandic epilepsy between standard care and standard care augmented with a novel, tailored parent-led CASTLE Online Sleep Intervention (COSI) that incorporates evidence-based behavioural components.

Methods and analyses CASTLE Sleep-E is a UK-based, multi-centre, open label, active concurrent control, randomised, parallel-group, pragmatic superiority trial. A total of 110 children with Rolandic epilepsy will be recruited in out-patient clinics and allocated 1:1 to standard care (SC) or standard care augmented with COSI (SC + COSI). Primary clinical outcome is parent-reported sleep problem score (Children's Sleep Habits Questionnaire). Primary health economic outcome is the Incremental Cost Effectiveness Ratio (National Health Service and Personal Social Services perspective, Child Health Utility 9D instrument). Parents and children (≥ 7 years) can opt into qualitative interviews and activities to share their experiences and perceptions of trial participation and managing sleep with Rolandic epilepsy.

**Ethics and dissemination** The CASTLE Sleep-E protocol was approved by the Health Research Authority East Midlands (HRA) – Nottingham 1 Research Ethics Committee, reference: 21/EM/0205. Trial results will be disseminated to scientific audiences, families, professional groups, managers, commissioners, and policy makers. Pseudo-anonymised Individual Patient Data will be made available after dissemination on reasonable request.

**Registration details** ISRCTN registry (Trial ID: ISRCTN13202325, prospective registration 09/Sep/2021). See Supplemental Table 1 for the World Health Organisation Trial Registration Data Set (Version 1.3.1).

## Strengths and limitations of this study

- First randomised controlled trial to evaluate the clinical- and cost-effectiveness of a novel, tailored, parent-led CASTLE Online Sleep Intervention (COSI) that incorporates evidence-based behavioural components for children with Rolandic epilepsy
- Extensive Patient and Public Involvement via dedicated CASTLE Advisory Panel
- Embedded health economic evaluation
- **Limitation**: Heavily reliant on parent and child self-report to assess intervention implementation, ameliorated by COSI e-analytics and actigraphy data

#### **INTRODUCTION**

Epilepsy is one of the most common long-term neurological conditions worldwide whose prevalence peaks during childhood (5–9 years) and late in life (over 80 years).[1] Epilepsy in children (5 to <13 years) accounts for the annual loss of 2.6 million disability-adjusted life years, equivalent to 1.8 % of the global burden of disease among children and adolescents.[2] Rolandic Epilepsy (RE) is the most common childhood epilepsy.[3]

In the UK, RE has a stable crude incidence rate of 5 in 100 000 children (<16 years) or 542 new cases annually.[4] Concurrent neuro-developmental disorders are very common (35 %).[5] Seizures are often triggered by sleep fragmentation.[6] Many parents co-sleep or monitor children with nocturnal seizures, and children experience a fear of death during and after a seizure.[7] Problems related to sleep emerge as a top concerns for both children and parents,[8] but are often unaddressed.[9 10]

A recent systematic review and meta-analysis of clinical trials shows that parent-based behavioural sleep interventions are effective for typically-developing children and those with neurological and neuro-developmental disorders.[10] The review concluded that randomised controlled clinical trials assessing functional outcomes (e.g. cognition, emotion, behaviour) and targeting specific populations (e.g. epilepsy) are missing (but see two recent trials).[11 12] Harms capture for cognitive-behavioural and behavioural sleep interventions has been sparse (only 32.3 % of trials address Adverse Events) and predominantly inadequate (92.9 % of trials do not meet adequate reporting criteria).[13] Observed harms of behavioural sleep interventions in adults have been mild (e.g. transient fatigue/exhaustion from sleep restriction in insomnia in 25–33 % of participants).[14] The only published paediatric and adult epilepsy trials did not address harms. [11 12] Based on the existing evidence, the benefits of behavioural sleep interventions in children with epilepsy outweigh potential harms, especially because sleep problems not only affect seizure control, but overall child well-being, learning and memory, and parental quality of life.[9 10] There remains, however, uncertainty whether sleep interventions, which can be resource intensive, are cost-effective in public health systems.

This protocol describes the design for the Changing Agendas on Sleep, Treatment and Learning in Epilepsy (CASTLE) Sleep-E trial, which evaluates the clinical- and costeffectiveness of a novel, tailored, parent-led CASTLE Online Sleep Intervention (COSI) that incorporates evidence-based behavioural components for children with epilepsy. COSI and CASTLE Sleep-E outcome-selection were co-produced by affected children, young people, and their parents, sleep- and epilepsy experts.[8 15-17] The CASTLE Sleep-E protocol follows Standard Protocol Items: Recommendations for Interventional Trials (SPIRIT),[18 19] its extension for Patient Reported Outcomes (SPIRIT-PRO),[20] and the Guidance for Reporting Involvement of Patients and the Public (GRIPP2).[21]

As CASTLE Sleep-E is a pragmatic superiority trial assessing whether UK standard care for children with RE should be augmented with an online behavioural sleep intervention, standard care is the appropriate comparator.[22-24] Current UK clinical guidelines[25-27] recommend that standard care for children with RE consist of a comprehensive care plan with the option of pharmacological treatment with anti-epileptic drugs (AEDs).

The primary objective of CASTLE Sleep-E is to determine if standard care augmented with COSI is superior to standard care alone in reducing sleep problems in children with RE and cost-effective. Implementation details and secondary objectives are reported in Table 1.

Outcome type	Specific measurement variable	Collected for	Participant-level analysis metric	Measurement time-point(s)
Primary				
1. Clinical	Children's Sleep Habits Questionnaire[28]	Child	Total score	Baseline, 3 months
2. Health economic	Cost utility of COSI <sup>a</sup> : National Health Service and Personal Social Services perspective, using outcomes 13–15	Child and Parent	<ul><li>Time integral of utility</li><li>Total costs</li></ul>	Baseline, 3 months, 6 months, (PLICS and HES at 6 months only)
Secondary				
1. Clinical	Children's Sleep Habits Questionnaire[28]	Child	Total score	Baseline, 6 months
2. Clinical	Seizure-free period	Child	Time to first seizure from randomisation (days)	Randomisation, 3 months,
3. Clinical	Seizure remission	Child	Time to 6-months seizure remission from randomisation (days)	6 months
4. Clinical	Knowledge about Sleep in Childhood (unpublished custom-scale)	Parent	Total score	Baseline, 3 months
5. Clinical	Hospital Anxiety and Depression Scale[29]	Parent	Total score	Baseline, 3 months,
6. Clinical	Insomnia Severity Index[30]	Parent	Total score	6 months
7. Clinical	SleepSuite[31] (iPad App)	Child	Reaction time (ms) Executive function (accuracy)	Baseline, 3 months
8. Clinical	<ul> <li>Health-Related Quality Of Life Measure for Children with Epilepsy<sup>[32]</sup></li> <li>World Health Organisation – Five Well- Being Index[33]</li> </ul>	Child Parent	<ul><li>Total score</li><li>Total score</li></ul>	Baseline. 6 months
9. Clinical	Strengths and Difficulties Questionnaire[34]	Child	Total score	Baseline, 3 months, 6 months
10. Clinical	Parenting Self Agency Measure[35]	Parent	Total score	
11. Clinical	Actigraphy[36]	Child and Parent	<ul> <li>Total sleep time (minutes)</li> <li>Sleep latency (minutes)</li> <li>Sleep efficiency (% asleep of sleep period)</li> <li>All 2-week averages</li> </ul>	Baseline, 3 months

Table 1. Outcomes for CASTLE Sleep-E (incl. participant level metrics, time-points, aggregation method). Child measures may be collected by parent proxy.

<sup>&</sup>lt;sup>a</sup> Reported as incremental cost per Quality-Adjusted Life Year (QALY) gained

Outcome type	Specific measurement variable	Collected for	Participant-level analysis metric	Measurement time-point(s)
12. Clinical	Sickness-related school absences	Child	Total number of days	Randomisation,3 months, 6 months
13. Health	Health-utilities derived from:	Child and Parent	Total score	Baseline, 3 months,
economic	• EQ-5D-Y[37]	Child	Utility score	6 months
	<ul> <li>Child Health Utility instrument[38]</li> </ul>	Child	Utility score	
	• EQ-5D-5L[39]	Parent	Utility score	
14. Health	Insomnia Severity Index mapped to	Parent	Total score	Baseline, 3 months,
economic	EQ-5D health state utilities[40]		Utility score	6 months
15. Health economic	<ul> <li>Direct costs: National Health Service and Personal Social Services perspective, measured using</li> <li>Resource Use Questionnaire</li> <li>Case Report Form data</li> <li>Patient Level Information and Costing System (PLICS) data</li> <li>Hospital Episode Statistics (HES) data</li> <li>Serious Adverse Events (assessed at 3 months, 6 months)</li> </ul>	Child	Resource use and total cost	Baseline, 3 months, 6 months, (PLICS and HES at 6 months only)
16. Health economic	<ul> <li>Indirect and direct non-medical costs, measured using:</li> <li>Resource Use Questionnaire</li> <li>Case Report Form data</li> </ul>	Child and Parent	Resource use and total cost	Baseline, 3 months, 6 months
17. Health economic	Cost utility of COSI: Societal perspective, using Quality-Adjusted Life Years and Cost using outcomes 13, 14, and 16	Child and Parent	<ul> <li>Quality-adjusted life years from the time- integral of utility</li> <li>Mean of total costs</li> </ul>	Baseline, 3 months, 6 months
Qualitative	Trial experience	Child and	Qualitative interview transcript	3 months + 3 weeks
		Parent	Activity booklet transcript/photos	6 months + 3 weeks

## METHODS AND ANALYSES

### Trial design

CASTLE Sleep-E is a UK-based, multi-centre, open-label, active concurrent control, randomised (1:1), parallel-group, pragmatic superiority trial (overall trial start date: 14/May/2018, first trial site opened: 12/May/2022, first recruitment: 30/August/2022, planned trial end date: 31/July/2023). Compared are clinical- and cost-effectiveness of standard care (SC) alone and SC augmented with a novel, tailored, parent-led CASTLE Online Sleep Intervention (SC + COSI) in reducing sleep problems in children (5 to <13 years) with RE at 3- and 6 months after randomisation. Parents and children ( $\geq$  7 years) can opt into qualitative interviews and activities to share their experiences and perceptions within 3 weeks of completion of other data collection at 3- and 6 months after randomisation.

### Patient and Public Involvement

The CASTLE programme (which subsumes CASTLE Sleep-E) recruited a dedicated Patient and Public Involvement (PPI) Advisory Panel (AP) through social media and epilepsy charities in 2017. The CASTLE Advisory Panel (CAP) consists of 17 adults with experience of childhood epilepsy and five children with epilepsy (aged 6–15 years). CAP has been involved in CASTLE from the funding application onward (2 CAP members are co-applicants). Full PPI details are provided in GRIPP2 Short Form in Table 2.

### Trial setting and eligibility criteria

Participants will be identified by staff in NHS out-patient general paediatric and paediatric epilepsy clinics in the UK (pre-dominantly urban setting). Eligibility criteria for participants are reported in Supplemental Table 1, field 14 of the World Health Organisation Trial Registration Data Set (Version 1.3.1). In the UK, a clinical RE diagnosis is based on electroclinical criteria defined by the International League Against Epilepsy (<u>https://www.ilae.org/</u>). Semiology and EEG need to be judged as concordant by a consultant neurophysiologist. Neuroimaging does not form part of UK standard care for RE. Eligibility criteria for trial sites include a Capacity and Capability assessment as advised for NHS site set-up by the UK HRA. The expected number of trial sites is 40 (England: 34, Scotland: 4, Wales: 1, Northern Ireland: 1). A list of trial sites can be obtained from the Trial Manager (see Supplemental Table 1).

### Intervention

Participants will be allocated to trial arms (SC or SC + COSI) using minimisation (1:1 ratio). On allocation to SC + COSI, participants will receive an email with access details to COSI. COSI consists of a self-paced, novel, tailored, e-learning package for parents of children with epilepsy that incorporates evidence-based behavioural components. Table 3 provides a brief overview; detailed reports on the development, content, and evaluation of COSI have been published.[15 16] COSI is divided into 13 modules (1 screening for child-specific sleep problems to allow tailoring, 10 content, 1 additional resources, 1 initially hidden evaluation), of which three are compulsory (1 screening, 2 content). The non-compulsory modules are recommended based on screening outcome, but all modules are accessible, repeatable, and printable. The advice in COSI supports parents to implement general prevention techniques (e.g. good sleep hygiene) and specific behavioural change techniques (e.g. bedtime fading) relevant to their child's sleep problems. Three months after first being given access to COSI, parents will be asked by email to complete a COSI evaluation module. At the end of a participant's trial timeline (6 months), access to COSI will be revoked. After the trial, all families (irrespective of trial allocation) have the option to receive the COSI content in electronic format via email.

#### Fidelity, adherence, retention, and acceptability

Fidelity (intervention delivery) will be monitored through e-analytics embedded in the COSI system (modules accessed, and time spent per module). Strategies to improve completion of COSI training in case of non-access include: (1) an automated text-reminder after two days; (2) an email reminder after four days; (3) a phone call from researchers who developed COSI (the Sleep Team) after six days. To improve adherence to the intervention, (1) all participants will receive a phone call from the Sleep Team six weeks after account creation; and (2) children will receive postcards with child-oriented activities (e.g. maze) at three time-points to welcome them to the trial (weeks 1–2), to stay in touch (weeks: 4–5), and to thank them for participating (weeks 4–8 post-trial). To encourage completion of the intervention evaluation, participants will receive: (1) an automated text-reminder after three days of non-completion, (2) and a phone call from the Sleep Team after eight days of non-completion. Fidelity (intervention implementation, acceptability, perceived helpfulness) will be captured jointly by the COSI evaluation module and the qualitative trial component.

#### Discontinuation, withdrawal, concomitant care, or interventions

Participants may discontinue the trial intervention or withdraw from the trial if (1) the parent/child withdraws consent/assent respectively; or (2) a change in the child's condition justifies discontinuation of treatment in their clinician's opinion. Trial site staff will record withdrawal with reason where provided in electronic Case Report Forms (eCRFs). Pseudo-anonymised data up to the time of consent withdrawal will be included in analyses in accordance with General Data Protection Regulation (GDPR)[41] under the UK Data Protection Act 2018[42] — the trial Data Controller relies on the legal bases of 'public interest' and 'research purposes'.

To avoid confounding and to minimise participant burden, co-enrolment into other clinical trials is discouraged. Where recruitment into another trial is considered appropriate, the trial coordinating centre will discuss enrolment with the Chief Investigator (CI). Participation in the Rolandic Epilepsy Genomewide Association International Study (REGAIN: <u>https://childhoodepilepsy.org/research-studies/regain/</u>) is complementary (same CI).

Item
<ul> <li>To contribute to and guide the CASTLE Sleep-E study:</li> <li>To ensure greater relevance and acceptability of the study and study procedures to children with epilepsy and their parents.</li> <li>To ensure the study is communicated to families and the public in an accessible way (e.g. recruitment, dissemination).</li> </ul>
Two adults with experience of childhood epilepsy are co-applicants on the Changing Agendas on Sleep, Treatment and Learning in Childhood Epilepsy (CASTLE) Research Programme National Institute for Health and Care Research (NIHR) Award (https://tinyurl.com/ycyfkc63) and are an integral part of the CASTLE Advisory Panel (CAP). CAP is a dedicated Patient and Public Involvement (PPI) Advisory Panel that was recruited in 2017 through social media and epilepsy charities. CAP consists of 17 adults with experience of childhood epilepsy and five children with epilepsy (aged 6–15 years). CAP members are reimbursed for expenses and offered honorarium payments in acknowledgement of their contributions. Facilitated by a salaried Family Engagement Officer and the PPI lead (LB), CAP members have co-developed working practices (CAP Handbook: Adult version https://tinyurl.com/28u8jex4, child version: https://tinyurl.com/2p8d6bnx) and undertaken research training. CAP members communicate by video conference, telephone, email, social media, and face-to-face. CAP is represented in the Trial Steering Group (TSC, see Supplemental Table 2). CAP feedback and opinion is formally communicated to the CASTLE Sleep-E Trial Management Group (TMG, see Supplemental Table 2) via the CASTLE PPI lead (LB).
<ul> <li>To date (at the recruitment stage of CASTLE Sleep-E), CAP has contributed to the following trial aspects:</li> <li>Initial funding application</li> <li>Two adults with experience of childhood epilepsy are co-applicants on the CASTLE Research Programme NIHR Award (https://tinyurl.com/ycyfkc63)</li> <li>Trial design</li> <li>CAP strongly endorsed the investigation focus (sleep problems) and the focus on non-seizure related issues linked to epilepsy</li> <li>CAP tested and consulted on the trial intervention (CASTLE Online Sleep Intervention [COSI]) in respect to content, format, and acceptability (e.g. knowledge evaluation quiz was changed from compulsory to optional)</li> <li>CAP informed the selection of study questionnaires to ensure relevance to parents and children with epilepsy</li> <li>CAP guided trial design to ensure acceptability of processes (e.g. time, effort, schedule from a family perspective)</li> <li>Trial procedure</li> <li>CAP led the development of a trial flowchart and clinician's guide (top tips for explaining the trial to families to aid recruitment)</li> <li>CAP guided data collection processes (assent/consent procedure, delivery of equipment, instructions, and packaging of Actigraphs and iPads)</li> <li>CAP guided the qualitative interview content and format (e.g. topics, question wording, length, delivery method and format)</li> </ul>

# Table 2. GRIPP2- Short Form (SF)<sup>[21]</sup>: Guidance for Reporting Involvement of Patients and the Public in research

Section and topic	Item
	<ul> <li>CAP guided the development of all participant-facing trial materials including:         <ul> <li>Information Sheets and Consent Forms</li> <li>Child-friendly postcards to update and maintain interest in the trial</li> <li>Wording of trial emails sent to participating families, strap lines for promotional materials (e.g. mugs and pens for trial sites)</li> </ul> </li> <li>Dissemination</li> </ul>
	<ul> <li>CAP informed liaison with stakeholders via social media and direct contact (charities, patient groups)</li> <li>CAP developed lay summaries for completed work as part of the CASTLE programme and helped ensure the CASTLE Sleep-E trial website (https://castlesleepetrial.org.uk/) is accessible to families</li> <li>CAP informed ongoing work to attract new CAP members</li> </ul>
4: Discussion and conclusions Outcomes—Comment on the extent to which PPI influenced the study overall. Describe positive and negative outcomes	<ul> <li>To date (recruitment stage of CASTLE Sleep-E), overall positive outcomes of CAP contributions to CASTLE Sleep-E have resulted in a trial design, procedure, materials, and dissemination that is likely to have greater appeal and relevance to parents of children affected by Rolandic epilepsy and to the children themselves. CAP has made the trial more family- focused, and enabled more direct public involvement (e.g. contact details of the Family Engagement Officer on the CASTLE Sleep-E webpage). This should increase the proportion of eligible patients to assent/consent to trial participation. Materials (including the trial intervention itself) and procedures should be more accessible and more feasible to complete for participants, which should positively affect adherence, compliance, and retention. Throughout their involvement, CAP contributions to the CASTLE programme have exceeded expectations, and taken on a greater, independent purpose (e.g. forming a support group via social media). The Coronavirus (COVID-19) pandemic meant that CAP's work had to move online, and whilst this has facilitated engagement between CAP members across the country, it made it more difficult for the children to join in some of the consultation exercises.</li> </ul>
5: Reflections/critical perspective Comment critically on the study, reflecting on the things that went well and those that did not, so others can learn from this experience	TBC (currently at recruitment stage of CASTLE Sleep-E)

Module	Module Name	Outline content	Compulsory or recommended
A	What is sleep and why is it important	Education about normal sleep physiology and processes	Compulsory
В	Sleep and seizures: a vicious cycle	Information about the relationship between sleep and seizures	Compulsory
С	Personalising this advice for your child	A sleep screening questionnaire to identify key areas of concern or problems around individual child sleep	Compulsory
D	Tips on sleep hygiene for everyone	General advice about key aspects of sleep hygiene	Recommended for all
E	Advanced sleep behaviour training	Introduction to principles of behavioural sleep interventions	Recommended for all
F	Learning difficulties, Attention Deficit Hyperactivity Disorder (ADHD), and Autism Spectrum Disorders	Advice for parents of children with other comorbid conditions	Recommended to parents who highlighted (in module C) their child may have comorbid conditions
G	Solving falling asleep problems	Sleep intervention options for typical falling asleep problems	Recommended to parents who highlighted (in module C) their child may have problems falling asleep
Η	Solving difficult night wakings and early morning waking	Behavioural techniques to address typical night or early waking problems	Recommended to parents who highlight (in module C) their child may have problems with their sleep during night or early morning wakings
I	Solving night-time fears	Behavioural techniques to address typical night-time fears	Recommended to parents who highlight (in module C) their child may have problems with night-time fears
J	Sleep walking, sleep terrors, and nightmares	Information about different sleep behaviours, what causes them and how to identify and manage different conditions	Recommended to parents who highlight (in module C) their child may have problems with sleep walking, sleep terrors, and/or nightmares
К	Troubleshooting and maintaining good sleep	How to deal with common issues, such as the child being ill or parents disagreeing about how to manage sleep and advice about how to maintain any benefits	Recommended to all
L	Resources	Links to additional resources of support, information and advice relating to sleep	Recommended to all
М	Evaluation	Questionnaire in which parents are asked to report on their experiences of using COSI	Recommended to all

Table 3. Content of the CASTLE Online Sleep Intervention (COSI)

### Outcomes and participant timeline

Outcomes are reported in Table 1 and were chosen collaboratively by children and young people with epilepsy and their parents, sleep- and epilepsy experts[8 17] in accordance with Core Outcome Measures in Effectiveness Trials (COMET) guidelines.[43] Psychometric properties and clinical relevance of outcomes are reported in Supplemental Table 3. Each participant will be followed up for 6 months. The participant timeline and estimated time requirement are respectively shown in Table 4 and Supplemental Table 4.

#### Sample size

The target sample size (110 children with RE, 55 per trial arm) was calculated based on achieving 90 % power to detect the minimal clinically important difference (MCID) in the primary clinical outcome (CSHQ) at 3 months after randomisation, accounting for 10 % expected attrition (non-parametric test with two-sided 5% significance level). MCID was defined based on an individual-focused anchor-based method,[44] that is, 'the smallest difference in outcome that patients perceive as beneficial and which mandates a change in patient management'.[45] The MCID value was based on the estimated reduction in total CSHQ score required for children with epilepsy (M = 48.25, SD = 8.91)[7] to fall at or below the diagnostic cut-off score of 41 for sleep disorders in paediatric populations.[28]

### Recruitment, stopping guidelines, interim analyses

An Independent Data and Safety Monitoring Committee (IDSMC) will monitor recruitment and make recommendations to the Trial Steering Committee (TSC) concerning trial continuation, adjustments of recruitment methods, and follow-up optimisation (see Supplemental Table 2). A traffic light approach will determine trial continuation: (1) Green: Continue if at least 30 trial sites have opened and 22 participants have been randomised by end of month 6; (2) Amber: Implement additional recruitment strategies if 15–21 participants have been randomised by end of month 6; (3) Red: If recruitment is <15 participants by end of month 6, then stopping the trial early will be discussed with the TSC. Formal interim analyses of the accumulating data will not be performed.

### Treatment allocation

Participants will be allocated with a 1:1 ratio to either SC or SC + COSI based on a computer generated adaptive restricted randomisation procedure that minimises differences between trial arms in variables likely to affect outcomes. Minimisation algorithm details are not published to avoid subversion of allocation sequence concealment, but include seizure frequency, AED, and sleep medication details. The allocation concealment mechanism is an online, central randomisation service implemented and maintained by the Liverpool Clinical Trial Centre (LCTC). The service will be accessed within four weeks of participant enrolment (once consent and eligibility confirmed, Participant ID issued, baseline dataset completed) by trained, authorised staff at trial sites. Randomisation will trigger allocation emails to the Trial Manager at LCTC and to the relevant trial site as well as enable COSI access for participants allocated to the intervention arm. Trial sites will notify the participant's General Practitioner (GP) of the treatment allocation by letter (electronic or hard copy, depending on preference).

### Blinding

Only quantitative data analysts will be blinded (Participant IDs do not reveal treatment allocation). All other stakeholders (participants, parents, healthcare providers, data

collectors, qualitative researchers) will be aware of the allocated intervention. Emergency unblinding procedures are therefore unnecessary.

	T-4 weeks <sup>a</sup>	<b>TO</b> <sup>b</sup>	T+3 months	T+6 months
	Consent and Baseline	Randomisation	Follow up visit	Follow up visit
Visit No	1	2	3	4
Informed consent/assent	х			
Review of medical history and EEG <sup>c</sup> results	х			
Eligibility confirmation	х	х		
COVID-19 Screener	х		Х	
Review of seizure occurrence		х	Х	Х
Hospital admissions		х	Х	х
Demographics	х			
School absences		х	Х	Х
Check contact details for accuracy		х	Х	х
Children's Sleep Habit Questionnaire[28]	Х		Х	Х
SleepSuite[31] (iPad)	Х		Х	
WHO–Five Well-Being Index[33]	х			х
Health-Related Quality Of Life Measure for Children with Epilepsy[32]	Х			Х
Strengths and Difficulties Questionnaire[34]	Х		Х	Х
CHU-9D <sup>d</sup> /CHU-9D proxy[38]	Х		Х	Х
EQ-5D-Y/EQ-5D-Y proxy[37]	х		Х	х
EQ-5D-5L[39]	Х		Х	х
Parenting Self Agency Measure[35]	Х		Х	х
Insomnia Severity Index[30]	х		Х	х
Hospital Anxiety and Depression Scale[29]	Х		Х	Х
Resource Use Questionnaire	Х		Х	Х
Knowledge about Sleep in Childhood	Х		Х	
Randomisation Standard Care (SC) or (SC + COSI <sup>e</sup> )[16]		Х		
Intervention arm only: COSI[16]				
Actigraphy and sleep diary[36] (14 days)	Х		Х	
Confirm continuing trial participation			Х	Х
Assessment of Serious Adverse Events			Х	Х
Completion of Follow-up Case Report Form			Х	Х
Review of concomitant medications		х	Х	Х
Qualitative Interview <sup>f</sup>			Х	х

Table 4. CASTLE SLEEP-E participant timeline and order of outcome completion.

<sup>&</sup>lt;sup>a</sup> Up to four weeks flexibility between consent and randomisation to allow delivery of actigraph and iPad.

<sup>&</sup>lt;sup>b</sup> Randomisation may be performed once two weeks of actigraphy and the minimum dataset are complete.

<sup>&</sup>lt;sup>c</sup> Electro-encephalogram (EEG)

<sup>&</sup>lt;sup>d</sup> Child Health Utility Index 9D (CHU-9D)

<sup>&</sup>lt;sup>e</sup> CASTLE Online Sleep Intervention (COSI)

<sup>&</sup>lt;sup>f</sup> Optional trial component: Consenting participants are interviewed within 3 weeks of follow up visits 3 and 4

#### Assent and consent

Potentially eligible children will be screened at trial centres by trained site staff. Screening outcome will be documented. Eligible children with interested parents will be invited to participate and provided with a Patient Information Sheet and Consent Form electronically and/or hard-copy (PISC, three versions: Parent, child [5–6 and 7–12 years]). Sufficient time will be allowed for discussion of the trial and the decision to assent/consent to trial entry and the optional qualitative component. Assent (children aged 7-12 years) and consent (parents) may be given face-to-face or remotely and will be electronically captured in a secure Consent Database managed by LCTC. Reasons for declining participation will be asked, but it will be made clear that children and parents do not have to provide a reason.

#### Data collection and management

Data collection will be carried out electronically except for Serious Adverse Events and Participant Transfer Forms (hard copy). At consent/assent, site staff will enter patient medical history (including electro-encephalogram), eligibility confirmation, COVID-19 screening, and demographics (see Table 4) into eCRFs stored in a secure Data Management System managed by LCTC. Trial participation will be added to the patient's medical records alongside their unique Participant ID.

Consent- and Contacts Databases are securely linked. The addition of a new participant will trigger email notifications to the parents containing access links to baseline assessments (see Table 4) and the Sleep Team who will access the Contacts Database to arrange the delivery of an iPad pre-configured by LCTC (optionally fitted with pre-paid SIMs), and two actigraphs with supporting documents. iPads (Generations 7–8, iOS 15.2 or 15.3) will be used to access the SleepSuite App, (V 1.4)[31], which assesses executive functions in child-friendly, interactive games (e.g. popping virtual bubbles with smiling children's faces). Access requires the Participant ID and is only possible at pre-specified trial time-points (see Table 4). Data is only stored on the iPad until the test-session completion, then automatically uploaded to a cloud-based server, and then securely downloaded for analyses by authorised LCTC staff. Families lacking other means of internet access can use iPads fitted with pre-paid SIMS to access other online trial materials (including email).

Actigraphs (Micro Motionlogger<sup>®</sup> Watch and Watchware Software V 1.99.17.4, Ambulatory Monitoring, Inc., NY: USA) will be used to collect 14 days of objective sleep data from child and parent. Concurrent sleep diaries (hard copies) will be completed by the parent with or without child input. At the end of the baseline period, actigraphs will be returned to the Sleep Team via pre-paid courier. The Sleep Team will download and securely store pseudo-anonymised (using Participant IDs) actigraphy data for pre-processing (manual selection of sleep periods cross-checked against sleep diaries) per night at participant-level. Summary variables (sleep latency, total sleep time and sleep efficiency) are then automatically calculated by actigraph software, manually collated, and securely transferred electronically to LCTC for trial-level analyses by the Trial Statistician.

Participants will be randomised to trial arms during a telephone/video call or clinic visit only *after* site staff have confirmed that baseline data (see Table 4) is complete, and eligibility, consent/assent, and contact details are still valid. Data collection will be repeated 3- and 6 months after randomisation, and iPads to LCTC via trial sites (see Table 4).

The Qualitative Research Team will access the Contacts Database to schedule audiorecorded interviews with children and parents who consented/assented to this optional trial component. Interviews (audio- or audio-video) will take place remotely within 3 weeks of completion of other data collection at 3- and 6 months after randomisation. Parents and children will be interviewed together or separately as preferred. Parents and children will have the opportunity to think through their ideas prior to the interview (as proposed by parents and children from the CASTLE Advisory Panel). Children will be invited to complete activity booklets in advance of their interviews (the booklets will be mailed or emailed one week prior to their interview); the content they complete will support the interview. Parents will receive a list of proposed questions/topics. Children will be able to share the booklet with the Qualitative Team (e.g. screen or photograph sharing, verbal description).

The direct costs of health and personal social services, and indirect costs of productivity losses and school absenteeism will be collected using a Resource Use Questionnaire administered at baseline and during follow-up visits. Other data such as concomitant medications, study visits and Adverse Events will be collected using eCRFs. Trial participants' use of secondary care services will be collected from Patient-Level Information and Costing Systems (PLICS) data obtained from the finance departments of each recruiting hospital or from Hospital Episode Statistics (HES) data obtained from NHS Digital at the end of the trial. PLICS and HES data will be pseudo-anonymised and transferred securely to the trial health economists at Bangor University.

#### Data quality, security, and trial oversight

Reliability, validity, and clinical relevance of outcomes are reported in Supplemental Table 3. Processes to promote quality and security of collected data include general local training of site staff and research teams (Good Clinical Practice); and trial-specific training in the use of electronic forms and databases by LCTC. LCTC will request to see evidence of appropriate training and experience of all trial staff. Staff will be signed off as appropriately qualified by the Cl. Electronic data capture provides several in-built validity and security checks (e.g. data type, range, and missingness checks in eCRFs, SleepSuite use/access restrictions). Some electronic and all hard-copy data will be repeat checked (e.g. eligibility, contact details). Data processing requiring more subjective judgement will be performed by minimum of two trained researchers on at least a subset of data (i.e. manually-assisted selection of actigraphy sleep period; thematic and content analysis of qualitative data).

Data will be processed and stored in accordance with GDPR under the UK Data Protection Act 2018. Central data monitoring will be performed by LCTC who will raise and resolve queries with site and research teams within the online system. The University of Liverpool is registered with the Information Commissioners Office. LCTC will receive trial participants' HES identifiers for secure transfer to the Health Economic team, who will access, securely store, and dispose of HES data in accordance with the Bangor University and NHS Digital Data Sharing Framework Contract.

#### Statistical methods

Statistical analyses of all but health economic and qualitative data will be performed by the Trial Statistician (LCTC) using SAS software, Version 9.4 or later. Intention-To-Treat (ITT) will be the main analysis strategy for primary and secondary outcomes (see Table 1 and Table 5). Minimisation variables (including seizure frequency, AED, and sleep medication details) will be adjusted for at baseline. Statistical significance will be set at the conventional two-sided 5 % level; clinical relevance will be based on previous research (see Supplemental Table 3). Point estimates with 95 % two-sided confidence intervals will be reported adjusted and unadjusted for covariates. No multiplicity adjustments will be made (only one primary clinical outcome, uncorrected secondary outcome analyses).

Sensitivity analyses will be carried out if the amount of missing data is greater than 10 %. Multiple imputation will be used to assess the robustness of the analysis to missing primary outcome data. The multiple imputation method will follow published guidelines.[46] PROC MI in SAS will be used to generate 50 complete data sets. The imputation model will include all variables included in the primary outcome analysis model. The overall summary adjusted mean difference will be presented with 95 % confidence intervals, to assess the sensitivity of the primary analysis to missing data. All analyses will be reported in accordance with the Consolidated Standards of Reporting Trials Checklist (CONSORT)[47] and regardless of statistical significance.

#### Health economic evaluation

The economic analysis will be performed in accordance with a Health Economics Analysis Plan, and by the trial health economists at Bangor University. The primary analysis will adopt an NHS and Personal Social Services perspective and, based on Quality-Adjusted Life Years (QALYs) as a measure of health outcome, estimate the incremental cost-effectiveness ratio from an incremental analysis of the mean costs and QALYs for the intervention and control trial arms.[48] Data assumed to be missing at random will be imputed using multiple imputation by chained equations.[49]

Sensitivity analyses will be conducted to test whether, and to what extent, the incremental cost effectiveness ratio is sensitive to key assumptions in the analysis (e.g. unit prices, different utility estimates from CHU-9D[38] vs. EQ-5D-Y[37]). The joint uncertainty in costs and QALYs will be addressed through application of bootstrapping and estimation of cost-effectiveness acceptability curves.[50] Alternative scenarios considering a broader cost perspective (including indirect costs, such as school absences and loss of productivity, valued by reference to published sources), and a range of outcomes (including parental QALYs, measured using the EQ-5D-5L[51] and ISI[30 40]) will be conducted. Inclusion of spillover disutility[52] (impact on parents' utility) will be based on the NICE reference case specification[53] that all QALYs are of equal weight and calculated assuming additive effects. Health-economic findings will be reported according to the Consolidated Health Economic Evaluation Reporting Standards (CHEERS).[54]

#### Qualitative component

Child and parent interviews will be analysed by the Qualitative Research Team using an interpretive, reflexive, and conceptual analytical approach. Audio-recordings of interviews will be transcribed and thematically analysed in discrete sets (e.g. intervention/control, child/parent, engagement/lack of engagement with intervention, types of decision-making, different responses/experiences). Parent and child transcripts will first be analysed separately, and then as dyads. All data will be used for synthesis. Thematic and content analyses will be used for child activity booklets (text and images). Qualitative and selected quantitative data (e.g. anxiety measures, actigraphy data) will be compared, as appropriate.

Outcome type	Specific measurement variable	Hypothesis	Method of analysis
Primary			
Clinical	Children's Sleep Habits Questionnaire[28]	Total score lower in intervention arm at 3 months	<ul> <li>Linear mixed effect regression:</li> <li>Fixed effects: Intervention (binary)</li> <li>Random effects: Trial site (categorical)</li> <li>Co-variates: <ul> <li>Baseline score</li> <li>Use of sleep medication (binary)</li> </ul> </li> </ul>
Health economic	Cost <sup>a</sup> per quality-adjusted life year gained	Not applicable (health economic evaluation)	Cost-effectiveness (utility) analysis
Secondary			
Clinical	Children's Sleep Habits Questionnaire[28]	Total score lower in intervention arm at 6 months	Linear mixed effect regression (as before)
Clinical	Seizure-free period	Time to first seizure (days) differs between trial arms at 3 and 6 months	<ul> <li>Survival analyses</li> <li>Kaplan-Meier curves by trial arm</li> <li>Cox proportional hazards regression (if applicable) <ul> <li>Co-variates:</li> <li>Use of sleep medication (binary)</li> <li>Trial site (categorical)</li> </ul> </li> </ul>
Clinical	Time to 6-months seizure remission from randomisation (days)	Time to 6-months seizure remission (days) differs between trial arms at 6 months	Survival analyses (as before)
Clinical	<ul> <li>Knowledge about Sleep in Childhood</li> <li>Actigraphy[36] (2-week average):         <ul> <li>Total sleep time</li> <li>Sleep latency</li> <li>Sleep efficiency</li> </ul> </li> </ul>	Total score differs between trial arms at 3 months	Linear mixed effect regression (as before)
Clinical	<ul> <li>Hospital Anxiety and Depression Scale[29]</li> <li>Insomnia severity index[30]</li> </ul>	Total score lower in intervention arm at 3 and 6 months	Linear mixed effect regression (as before)
Clinical	Sickness-related school absences	Total days differs between trial arms at 3 and 6 months	Poisson mixed-effects regression

Table 5. Analysis plan for outcome variables in CASTLE Sleep-E. Further analyses details are reported in-text.

<sup>&</sup>lt;sup>a</sup> Perspective: NHS and PSS perspective; Alternative perspective: Societal (Indirect and direct non-medical costs)

Outcome type	Specific measurement variable	Hypothesis	Method of analysis
Clinical	<ul> <li>Health-Related Quality Of Life Measure for Children with Epilepsy[32]</li> <li>World Health Organisation – Five Well-Being Index[33]</li> </ul>	Total score differs between trial arms at 6 months	Linear mixed effect regression (as before)
Clinical	<ul> <li>SleepSuite[31]: Animal task</li> <li>SleepSuite: Bubble task <ul> <li>Shape detection</li> <li>Emotion detection</li> <li>Gender detection</li> </ul> </li> <li>SleepSuite: Maze task</li> </ul>	Executive function, reaction time, and variability differ between trials arm at 3 months	<ul> <li>Poisson/zero-inflated negative binomial regression (depending on presence of overdispersion)</li> <li>2 x 2 multi-variate repeated-measures Analysis of Variance (ANOVA)</li> <li>Factors: Time (PM/AM) x Intervention (Pre/Post)</li> <li>Fitted per detection task (Shape, Emotion, Gender)</li> <li>Linear mixed effect regression (as before)</li> </ul>
Clinical	<ul> <li>Strengths and Difficulties[34] Questionnaire</li> <li>Parenting Self Agency Measure[35]</li> </ul>	Total score differs between trial arms at 3 and 6 months	Linear mixed effect regression (as before)
Qualitative	Trial experience <sup>b</sup>	Not applicable (inductive)	<ul> <li>Thematic analysis (interpretive, reflexive, and conceptual analytical approach)</li> <li>Discrete sets: Intervention/Control, Child/parent, Engagement with intervention/lack thereof, Decision making types, Responses/experiences</li> <li>Separately for child and parent, then jointly (dyad)</li> <li>Comparisons to selective objective data as emerging from analysis (e.g. Anxiety measures, Actigraphy)</li> </ul>

<sup>&</sup>lt;sup>b</sup> Source data for trial experience: Qualitative interviews (parents and children individually and as dyad), activity booklets (children only)

### Harms

A flowchart of Adverse Event (AE) reporting requirements is shown in Supplemental Figure 1. Harms severity and causality will be graded by the investigator responsible for the care of the participant based on categories shown in Supplemental Table 5. If any doubt about causality exists, the local investigator should inform LCTC who will notify the CI. In case of discrepant views, the Research Ethics Committee (REC) will be informed of both views. Seriousness and expectedness of AEs will be defined based on International Council for Harmonisation of Technical Requirements for Pharmaceuticals for Human Use Definitions and Standards for Expedited Reporting (ICH E2A, ref: CPMP/ICH/377/95). Expectedness will be assessed by the CI. The only expected AEs in CASTLE Sleep-E are mild and transient worsening of sleep behaviours targeted by the trial intervention. Safety data will be quality-checked by a statistician not otherwise involved in the trial. Safety analysis will include all patients randomised and starting treatment and be presented descriptively split by treatment arm.

### Auditing

The CI will ensure that the trial team conducts monitoring activities of sufficient quality and quantity (e.g. protocol adherence, consent/assent, data quality). The Sponsor will delegate monitoring duties and activities to LCTC. The CI and LCTC will inform the Sponsor of any concerns. Auditing does not meet the National Institute for Health and Care Research (NIHR) or SPIRIT Statement definitions of independence[19 55] as auditors (LCTC and CI) are part of the trial team.

### Protocol amendments

Substantive protocol amendments will be notified to HRA via the UK's Integrated Research Application System (IRAS). Trial sites will receive an amendment pack of HRA- and RECapproved changes and unless an objection is received within 35 days, the trial will continue at site with a GO LIVE email.

## Ancillary and post-trial care

King's College London (KCL) holds insurance against claims from participants for harm caused by their participation in this clinical study; compensation can be claimed in case of KCL negligence.

## Ethics and dissemination

The CASTLE Sleep-E protocol was approved by the HRA East Midlands – Nottingham 1 REC, reference: 21/EM/0205. Trial results will be disseminated to scientific audiences in peer-reviewed publications and conferences, and — with the help of the CASTLE Advisory Panel (parent and child experts-by-experience), relevant charities (e.g. Epilepsy Action, Epilepsy Society and Cerebra) and professional groups (e.g. Royal College of Paediatrics and Child Health, Epilepsy Specialist Nurses Association) — as plain language summaries to families, other professional groups, managers, commissioners, and policy makers. Pseudo-anonymised Individual Patient Data and associated documentation (e.g. protocol, statistical analysis plan, annotated blank Case Report Form) will be made available after dissemination on reasonable request.

#### Registration details

ISRCTN registry (Trial ID: ISRCTN13202325, prospective registration 09/September/2021). The World Health Organisation Trial Registration Data Set (Version 1.3.1) for CASTLE Sleep-E is shown in Supplemental Table 1.

# **Author Statement**

### Contributorship (alphabetic surname order)

PG, DKP (Chief Investigators); CAP, CTS, HH, LW, BC, CM, DH, and LB (Co-Investigators) conceived the study and are award holders. Topic expertise for the core outcome set development was provided by CAP, LB, BC, AC, HC, PG, DH, CM, DKP, CTS, and PRW. Epilepsy expertise-by-experience is provided by CAP. Topic expertise for epilepsy is provided by DKP. Topic expertise for the health economic evaluation is provided by WASH, DH, and EW. Topic expertise for intervention development was provided by GC, PG, HH, DKP, and LW. Topic expertise for Patient and Public Involvement (Advisory Panel and Family Engagement) is provided by CAP, AR-S, LB, BC, and CM. Responsibility for the selection of Patient-Reported Outcomes lay with CM. Responsibility for Programme management lies with AC. Topic expertise for qualitative research components is provided by CAP, LB, BC, and HS. Topic expertise for sleep is provided by GC, PG, HH, and LW. Topic expertise for Statistical analyses is provided by CTS, VW, and LWh. Responsibility for trial management lies with NA-N, CS, and LS-E. All authors contributed to the design and refinement of the study protocol. The protocol manuscript was written by KCD (including supplemental materials but excluding Figure 1 and Patient Information Sheet and Consent Forms). Authors in the Trial Management Group (TMG) had the opportunity to provide feedback twice (initial and final draft); non-TMG authors had the opportunity to provide feedback once (final draft). Provided feedback was incorporated. The final manuscript was approved for publication by all authors. GRIPP2 content was checked for accuracy by LB. Sponsor name and contact information are provided in Supplemental Table 1. Details of trial committees and other groups and individuals overseeing the trial are listed in Supplemental Table 2. Trial site Principal Investigators will be listed alphabetically in resulting publications as members of the CASTLE Sleep-E Consortium in the Acknowledgements section. There has not been and will not be any use of hired writers.

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**Disclaimer.** To avoid potential bias, neither the funder nor the sponsor of this trial has any role in or authority over the design, execution, analyses, interpretation of data, or result dissemination.

Competing interests. None declared.

Patient consent for publication. Not applicable.

Provenance and peer review. Not commissioned, externally peer reviewed.

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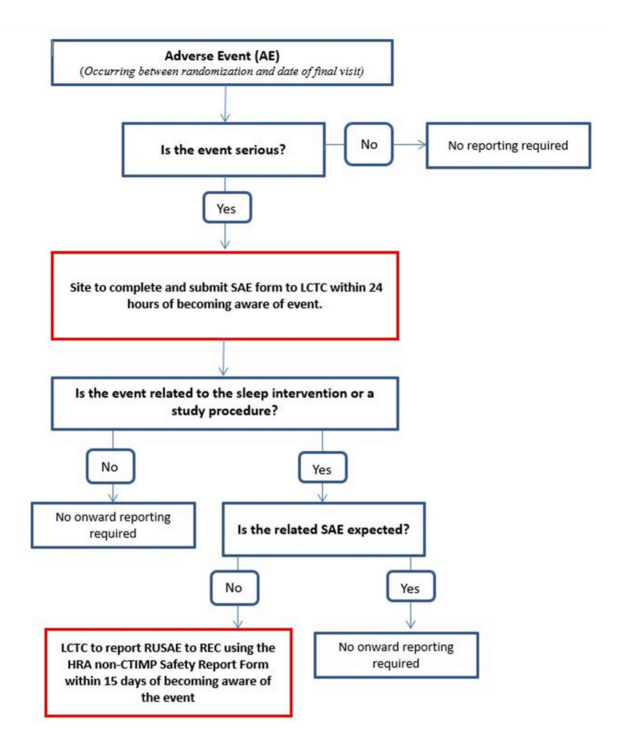
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Data category		Information		
1.	Primary registry and trial identifying number	ISRCTN: ISRCTN13202325		
2.	Date of registration in primary registry	09/September/2021		
3.	Secondary identifying numbers	CPMS 50413 RP-PG-0615-20007 IRAS 289580 21/EM/0205		
4.	Source(s) of monetary or material support	National Institute for Health and Care Research (NIHR)		
5.	Primary sponsor	Ms Jasmine Palmer Research & Innovation Operational Manager King's College Hospital NHS Foundation Trust		
		The Research & Innovation Office First Floor, Coldharbour Works 245a Coldharbour Lane, Brixton London SW9 8RR jasmine.palmer1@nhs.net +44 (0) 7790 950 219		
6.	Secondary sponsor(s)	Professor Reza Razavi Director of Research Management & Director of Administration (Health Schools) Room 5.31 James Clerk Maxwell Building 57 Waterloo Road London SE1 8WA <u>reza.razavi@kcl.ac.uk</u> +44 (0)20 7848 3224		
7.	Contact for public queries	Trial Manager: Lucy Stibbs-Eaton Liverpool Clinical Trials Centre University of Liverpool Liverpool L69 3BX LCTC@liverpool.ac.uk +44 (0)151 795 8751		
8.	Contact for scientific queries	Professor Deb Pal Professor of Paediatric Epilepsy Maurice Wohl Clinical Neuroscience Institute King's College London 5 Cutcombe Road London SE5 9RX <u>deb.pal@kcl.ac.uk</u> +44 (0) 207 848 5762		
9.	Public title	A trial comparing the effectiveness of an online sleep behavioural intervention versus standard care in children with rolandic epilepsy		
10.	Scientific title	Changing Agendas on Sleep, Treatment and Learning in Epilepsy (CASTLE) Sleep-E: A randomised controlled trial comparing an online behavioural sleep intervention with standard care in children with Rolandic epilepsy		

Supplemental Table 6. World Health Organization Trial Registration Data Set (Version 1.3.1) for CASTLE Sleep-E

Data category	Information
11. Countries of recruitment	England Scotland Wales Northern Ireland
<ol> <li>Health condition(s) or problem(s) studied</li> </ol>	Sleep problems in Rolandic epilepsy also known as childhood epilepsy with centro-temporal spikes
13. Intervention(s)	<ul> <li>Intervention arm (SC + COSI): Novel, tailored, parent-led CASTLE Online Sleep Intervention (COSI) that incorporates evidence-based behavioural components. Delivered by parents to enrolled children with Rolandic epilepsy in their own homes after completion of self-paced online training. Standard care (SC) is augmented with the CASTLE Online Sleep Intervention (COSI).</li> <li>Active control arm (SC): UK National Health Service standard care (SC) for children with Rolandic epilepsy, which consists of a comprehensive care plan with the option of pharmacological treatment with anti-epileptic drugs (first- line mono-therapy with lamotrigine, levetiracetam, oxcarbazepine [girls and</li> </ul>
14. Key inclusion and exclusion criteria	<ul> <li>boys], carbamazepine or sodium valproate [both boys only]).</li> <li>Inclusion criteria Main CASTLE Sleep-E study</li> <li>1. Children diagnosed with RE/CECTS (see International League Against Epilepsy Diagnostic Manual at https://www.epilepsydiagnosis.org/syndrome/ects-overview.html)</li> <li>2. EEG showing focal sharp waves with normal background (see International League Against Epilepsy Diagnostic Manual at https://www.epilepsydiagnosis.org/syndrome/ects-eeg.html)</li> <li>3. Aged 5 to &lt;13 years at the time of randomisation</li> <li>4. Parent/Carer reported child sleep problem as defined by mild, moderate or severe score on Hiscock Australian global sleep question (Poor sleeper defined by caregiver responding 'Mild', 'Moderate' or 'Severe' to "Over the last 2 weeks, how much of a problem has your child's sleep been?")</li> <li>5. Documented informed consent received from a person with parental responsibility</li> <li>6. Family have an email address and mobile phone</li> <li>7. Parent and child are to have a good enough understanding of the English language to read and answer study questionnaires</li> <li>Qualitative component</li> <li>1. Consent of care giver to participate and for their child to participate (optional item on main trial consent form)</li> <li>2. Children need to be &gt;=7 years of age</li> </ul>
15. Study type	<ul> <li>Children with moderate/severe learning disability</li> <li>Interventional</li> <li>Allocation: Minimisation using a bespoke LCTC system Allocation concealment: Central web-interface Sequence generation: Randomised, 1:1 ratio Intervention model: Parallel assignment</li> <li>Blinding Child, parent, healthcare providers, data collectors, qualitative researchers: None (open label) Quantitative data analysts: Blinded</li> <li>Primary purpose: Clinical- and cost-effectiveness, process evaluation (qualitative trial component, COSI e-analytics and evaluation module)</li> </ul>

Data category	Information
16. Date of first enrolment	24/June/2022
17. Target sample size	<ul> <li>110 (55 children per arm)</li> <li>Calculation based on:</li> <li>Achieving 90 % statistical power to detect Minimal Clinically Meaningful Difference in primary outcome</li> <li>10 % expected attrition</li> </ul>
18. Recruitment status	Recruiting <ul> <li>First trial site opened: 12/May/2022</li> <li>First recruitment: 30/August/2022</li> </ul>
19. Primary outcome(s)	<ul> <li>Clinical: Children's Sleep Habits Questionnaire at 3 months</li> <li>Health economic: Cost-effectiveness of the intervention over 6 months after randomisation, measured in terms of incremental cost per quality-adjusted life year gained (Child Health Utility instrument or EQ-5D-Y) from the perspective of the National Health Services and Personal Social Services in the UK.</li> </ul>
20. Key secondary outcome(s)	<ul> <li>Clinical Outcome: Sleep problem reduction Metric/method: Children's Sleep Habits Questionnaire Timepoint: 6 months</li> <li>Clinical Outcome: Seizure frequency reduction Metric/method: Time to first seizure (days) Timepoint: 3 months, 6 months</li> </ul>
21. Ethics Review	<ul> <li>Status: Approved</li> <li>Approval reference: 21/EM/0205</li> <li>Health Research Authority East Midlands – Nottingham 1 Research Ethics Committee Chair: Mr Paul Hamilton +44 (0) 207 104 8115 or +44 (0) 207 104 8283 nottingham1.rec@hra.nhs.uk</li> </ul>
22. Completion date	31/July/2023
23. Summary results	ТВС
24. Individual patient data (IPD) sharing statement	<ul> <li>Plan to share IPD: Yes</li> <li>Plan description: At the end of the trial, after the primary results have been published, the pseudo-anonymised Individual Patient Data and associated documentation (e.g. protocol, statistical analysis plan, annotated blank case report form) will be prepared to be shared with external researchers on reasonable request.</li> </ul>
25. Protocol version and date	<ul> <li>Internal protocol: V4.0, 08/December/2021</li> <li>Manuscript for protocol publication: V3.2, 20/December/2022</li> </ul>

Role		Name (Initials)	Affiliation			
Trial management Group (TMG)						
Res	Responsibilities: Day-to-day running and management of the trial.					
Meeting frequency: Bi-weekly to three-monthly, depending on trial stage.						
1.	King's College Hospital Sponsor	Jasmine Palmer	King's College Hospital NHS			
	Representative		Foundation Trust, UK			
2.	Chief Investigator	Deb K. Pal	King's College London, UK			
3.	Co-Chief Investigator	Paul Gringras	Evelina London Children's Hospital, UK			
4.	Co-Investigator Public and Patient Involvement Lead	Lucy Bray	Edge Hill University, UK			
5.	Co-Investigator Qualitative Research Lead Public and Patient Involvement Co-Lead	Bernie Carter	Edge Hill University, UK			
6.	Co-Investigator Health Economics Lead	Dyfrig Hughes	Bangor University, UK			
7.	Co-Investigator Patient Reported Outcome Lead Public and Patient Involvement Co-Lead	Christopher Morris	University of Exeter, UK			
8.	Co-Investigator Lead Statistician	Catrin Tudur Smith	University of Liverpool, UK			
9.	Co-Investigator Intervention Development Lead	Luci Wiggs	Oxford Brookes University, UK			
10.	Supervising Trials Manager	Catherine Spowart	University of Liverpool, UK			
11.	Trial Manager	Lucy Stibbs-Eaton	University of Liverpool, UK			
12.	Trial Statistician	Liam Whittle	University of Liverpool, UK			
13.	CASTLE Programme Manager	Amber Collingwood	King's College London, UK			
14.	Researcher	Georgia Cook	Oxford Brookes University, UK			
15.	Researcher	Kristina C. Dietz	King's College London, UK			
16.	Health economist	Will A. S. Hardy	Bangor University, UK			
17.	Researcher	Holly Saron	Edge Hill University, UK			
Tria	l Steering Committee (TSC)					
Res	ponsibilities: Overall trial supervisi	on and advice, ultimate de	cision for the continuation of the trial.			
	eting frequency: At least annually.					
1.	Chair	Jeremy Parr	Newcastle University, UK			
2.	Medical statistician	Martyn Lewis	Keele University, UK			
3.	Paediatrician	Desaline Joseph	Evelina London Children's Hospital, UK			
4.	Public and Patient Involvement Representative	Jo Conduit-Smith	CASTLE Advisory Panel			
5.	Chief Investigator	Deb K. Pal	King's College London, UK			
6.	Co-Chief Investigator	Paul Gringras	Evelina London Children's Hospital, UK			

Supplemental Table 7. Composition, roles and responsibilities of the Trial Management Group, Programme Steering Committee, and Independent Data and Safety Monitoring Committee for CASTLE Sleep-E.

#### Independent Data and Safety Monitoring Committee (IDSMC) **Responsibilities:** Interim monitoring of safety and effectiveness, trial conduct and external data. Recommendation to TSC about trial continuation. Meeting frequency: At least annually 1. Chair Helen Cross University College London, UK 2. Paediatrician Alberto Verroti University of L'aquila, Italy University College London, UK 3. Medical statistician • Anthony Johnson (to 31/August/2022) • Appointment pending (20/December/2022)

Supplemental Table 8. Psychometrics and clinical relevance/minimal clinically important difference (CR/MCID) for CASTLE Sleep-E outcomes (Table 1). Metrics refer to the single referenced publication. Further validation studies exist, but, due to differences in population, setting, and/or methods, results cannot be merged.

Outcome	Description	Validity	Reliability	CR/MCID
Children's Sleep	Parent-reported, one-	Classification	Test-retest	Cut-off (total score):
Habits	week retrospective sleep	accuracy	2-week delay	41
Questionnaire	screening tool for	Sleep disorder	Pearson's r:	<ul> <li>Sensitivity: 80 %</li> </ul>
(CSHQ)[1]	children (4–10 years)	(yes/no)	0.62-0.79	• Specificity: 72 %
		<b>Receiver Operating</b>		<ul> <li>Accuracy: 80 %</li> </ul>
	35 items (2 duplicated	Characteristic	Internal	
	across subscales)	(ROC) analyses: See	consistency	MCID
	3-point Likert scales	MCID	Cronbach's α	Not assessed
	(rarely, sometimes,		Control	
	usually)	Construct validity	sample: 0.68	
	Total score (33 items):	See MCID	Clinical	
	33–99, lower is better		sample: 0.78	
	8 subscales:	Criterion validity		
	• Bedtime Resistance (6	Not assessed	Inter-rater	
	items)		reliability	
	• Sleep Onset Delay (1		Not assessed	
	item)			
	• Sleep Duration (3			
	items)			
	• Sleep Anxiety (4 items)			
	<ul> <li>Night Wakings (3</li> </ul>			
	items)			
	• Parasomnias (7 items)			
	<ul> <li>Sleep-Disordered</li> </ul>			
	Breathing (3 items)			
	Daytime Sleepiness (8			
	items)			
	,			
	Validation samples			
	Parents of 469 school			
	children (community			
	setting) and 154 children			
	diagnosed with sleep			
	disorder (hospital			
	setting); English			
	language; England, UK.			
	Test-retest: 60 parents			
	from control sample			
EQ-5D-Y[2 3]	Child- or adolescent	Not yet validated in	Not yet	CR/MCID
	reported (4–7 years: EQ-	UK (last updated	validated in	Applicability to utility
	5D-Y proxy; 8–16 years:	07/March/2022)	UK (last	scores debated,
	EQ-5D-Y, $\geq$ 16 years: EQ-	,,,	updated	suggested MCID:
	5D-5L), standardised		07/March/202	difference in index
	measure of current		2)	score between
	('today')		,	baseline health
	<ul> <li>health profile across 5</li> </ul>			profile and single-
	dimensions,			level transitions in
	<ul> <li>self-rated <i>health</i></li> </ul>			single domain (e.g.
	• sen-rated neurin status, and			33333 to 33332).
	• EQ-5D-Y index value,			
	using a country-			
	specific weighting	1		l

Outcome	Description	Validity	Reliability	CR/MCID
	(value set) of a given			
	health profile.			
	Two components:			
	1. Descriptive system			
	5 dimensions with 3			
	response severity			
	options each (tick-box):			
	Mobility			
	Self-care			
	Usual activities			
	Pain/discomfort			
	<ul> <li>Anxiety/depression</li> <li><u>Visual Analogue Scale</u></li> </ul>			
	Self-rated health on a			
	vertical Visual Analogue			
	Scale (VAS) that ranges			
	from 'The best health			
	you can imagine' (100)			
	to 'The worst health you			
	can imagine' (0).			
	Scoring:			
	<ul> <li>Descriptive system: 5-</li> </ul>			
	digit health profile			
	(best health state:			
	11111, indicating no			
	problem in each of the			
	5 dimensions; worst			
	health state: 33333			
	indicating many problems in each of			
	the 5 dimensions; 243			
	possible health states			
	are coded)			
	• VAS: 0–100 subjective			
	health state (worst to			
	best)			
	• EQ-5D-5L index value			
	Single summary			
	number, calculated by			
	subtracting country-			
	specific weighing			
	(value set) of an			
	obtained health profile			
	from 1, where 1			
	represents the best possible health profile			
	of 11111.			
	Value set validation			
	Value set validation sample (UK)			
	Not yet validated in UK			
	(last updated			
	07/March/2022)			

Child-reported (7–11	Due disting a second sec		
	Predictive accuracy	<u>Test-retest</u>	<u>CR/MCID</u>
ears) descriptive system	Standard ordinary	Not assessed	Applicability to utility
or current ('today')	least squares (OLS)		scores debated,
generic health-related	regression: 98.41 %	Internal	suggested MCID:
Juality-of-life		<u>consistency</u>	difference in index
		Utility values	score between
	errors.	are consistent	baseline health
			profile and single-
			level transitions in
	Not assessed	•	single domain (e.g.
			55555555555555 to
	-		555555554).
	Not assessed	-	
-			
-	-		
Activities			
ooring			
-	•	-	
	•		
• • •		-	
•			
-		-	
-	-		
		•	
		,	
_	health for the rest		
each of the 5	of life with a		
dimensions; 1953125	probability <i>p,</i> or		
possible health states	dying with a	Inter-rater	
are coded)	probability 1-p. The	reliability	
CHU-9D index value	utility value of a	Not assessed	
Single summary	given health-state		
number indicating the	is the point of		
utility value of a given			
	-		
_			
• •	-		
tasks.			
/alua aat validati			
	response options.		
esearch team of the			
Centre for Research and			
Evaluation (CRE) at			
Sheffield Hallam			
	uality-of-life dimensions with 5 esponse severity ptions each (circle): Worried Sad Pain Tired Annoyed School-/homework Sleep Daily routine Activities coring: Descriptive system: 9- digit <i>health profile</i> (best health state: 11111111, indicating no problem in each of the 9 dimensions; worst health state: 5555555 indicating many problems in each of the 5 dimensions; 1953125 possible health states are coded) <i>CHU-9D index value</i> Single summary number indicating the utility value of a given health state, established using Standard Gamble (SG) tasks. alue set validation ample (England) 245 households were andomly sampled from database of UK names in daddresses in heffield and uddersfield (England) vere contacted by a esearch team of the enter for Research and valuation (CRE) at	uality-of-lifeNo systematic bias, no auto-correlateddimensions with 5errors.esponse severityerrors.ptions each (circle):Construct validityWorriedNot assessedSadPainTiredNot assessedAnnoyedFace-validitySchool-/homeworkFace-validitySleepPreferenceDaily routineelicitation usingActivitiesStandard Gamble(SG) task, whichgive the choice ofDescriptive system: 9-living in a specificdigit health profilehealth-state until(best health state:that could result in11111111, indicatingcertainty (Choiceno problem in each ofA), or taking agamble (Choice B)gamble (Choice B)worst health state:that could result in55555555 indicatingliving in perfectmany problems inhealth for the resteach of the 5of life with aare coded)given health-stateSingle summarygiven health-stateutility value of a givenindifferencehealth state,between options Aand B.standard Gamble (SG)taks.consistent withhealth profiled anduudersfield (England)vere contacted by aesearch team of theentre for Research andvaluation (CRE) at	uality-of-lifeNo systematic bias, no auto-correlated errors.Consistency Utility values are consistent with health profiles, but required merging of the initial 5 response- levels for all but one of the 9 dimensions as follows:VarriedConstruct validity Not assessedPrefiles, but required merging of the initial 5 response- levels for all but one of the 9 dimensions as follows:Standard Gamble (Sof) task, which porplem in each of the 9 dimensions; the 9 dimensions; that could result in living in perfect health state: are coded)No taking a given health-state is the point of indifference between options A and B.Not assessedVariet talue set validation angple (England) 245 households were and addresses in heffield and luddersfield (England) 245 households were and addresses in heffield and luddersfield (England) zescarch tam of the erter or Research and valuation (CRE) at valuation (CRE) atNo assessed many problem in the entre or Research and valuation (CRE) at valuation (CRE) atNo assessed and addresses in heffield and valuation (CRE) at valuation (CRE) atNo assessed aud asset validation amore contacted by a essearch team of the entre or Research and valuation (CRE) at valuation (CRE) at valuation (CRE) atNo assessed aud asses of UK names and addresses in heffield and valuation (CRE) at valuation (CRE) at valuation (CRE) at valuation (CRE) at valuati

Outcome	Description	Validity	Reliability	CR/MCID
	University. 1195			
	households were			
	approached at the door,			
	of which 661 (55 %)			
	were in, and 300 (25 %)			
	agreed to take part. 282			
	respondents (all adults)			
	were analysed (94 %).			
	Compared to the general			
	UK population, this adult			
	sample was broadly			
	representative, but more affluent and highly			
	restricted			
	geographically.			
	Modelling did not			
	include key demographic			
	characteristics (e.g. age,			
	gender, education,			
	employment, religion			
	and ethnicity). The			
	sample consisted			
	exclusively of adults but			
	was used to derive a			
	paediatric value set.			
EQ-5D-5L[5]	Adolescent or adult-	Classification	Test-retest	CR/MCID
	reported (≥16 years),	<u>accuracy</u>	Not assessed	Applicability to utility
	standardised measure of	Not assessed		scores debated,
	current ('today'):		Internal	suggested MCID:
	<ul> <li>health profile across 5</li> </ul>	Construct validity	<u>consistency</u>	difference in index
	dimensions,	Not assessed	Not assessed	score between
	<ul> <li>subjective health</li> </ul>			baseline health
	<i>status,</i> and	Criterion validity	Inter-rater	profile and single-
	• EQ-5D-5L index value,	Not assessed	reliability	level transitions in
	using a country-		Not assessed	single domain (e.g.
	specific weighting	Face-validity		55555 to 55554).
	(value set) of an	Preference		
	obtained health	elicitation using		
	profile.	time trade-off		
	Two components:	(TTO) and discrete		
	1. <u>Descriptive system</u>	choice experiments		
	5 dimensions with 5	(DCEs).		
	response severity	• TTOs:		
	options each (tick-box):	Confirmation of		
	<ul> <li>Mobility</li> </ul>	negative relationship		
	• Self-care	relationship between level		
	<ul> <li>Usual activities</li> </ul>	sum score and		
	<ul> <li>Pain/discomfort</li> </ul>	average observed		
	<ul> <li>Anxiety/depression</li> </ul>	value.		
	2. Visual Analogue Scale	• DCEs:		
	Self-rated health on a	Confirmation of		
	vertical Visual Analogue			
	vertical Visual Analogue Scale (VAS) that ranges	assumption that health states with		

Outcome	Description	Validity	Reliability	CR/MCID
	you can imagine' (100)	scores are more		
	to 'The worst health you	likely to be		
	can imagine' (0).	chosen.		
	Scoring:			
	• Descriptive system: 5-			
	digit health profile			
	(best health state:			
	11111, indicating no			
	problem in each of the			
	5 dimensions; worst			
	health state: 55555			
	indicating many			
	problems in each of			
	the 5 dimensions;			
	3125 possible health			
	states are coded)			
	• VAS: 0–100 subjective			
	health state (worst to			
	best)			
	• EQ-5D-5L index value			
	Single summary			
	number, calculated by			
	subtracting country-			
	specific weighing			
	(value set) of an			
	obtained health profile			
	from 1, where 1			
	represents the best			
	possible health profile			
	of 11111.			
	Value set validation			
	sample (England)			
	2220 households from			
	66 post-code based			
	primary sampling units			
	in England were			
	contacted by the market			
	research company Ipsos			
	MORI. 2088 participants			
	were invited, of which			
	996 (47.7 %) completed			
	the valuation			
	questionnaire. Only			
	complete responses			
	were analysed (985			
	participants, 98.9 %).			
	Compared to the general			
	population of England,			
	the sample included			
	more people aged over			
	75 years, retired, and			
	with health problems,			
	but fewer younger			

Outcome	Description	Validity	Reliability	CR/MCID
	participants, and fewer males.			
Knowledge About Sleep in Childhood (KASC, custom-scale devised for CASTLE Sleep-E)	13 items Self-reported Likert- scales assessing parental efficacy in managing child sleep and knowledge about child sleep	Not evaluated	Not evaluated	Not evaluated
Hospital Anxiety and Depression Scale (HADS)[6]	Self-reported, one-week retrospective screening tool for anxiety and depression in people aged 16–65. 14 items 5-point Likert scales (0– 3) No total score Subscale score: 0–21, lower is better 2 subscales (7 items each): • Depression • Anxiety <u>Validation samples</u> 2 x 50 patients (16–65 years) with and without psychiatric disorders (hospital setting); English language; England, UK.	ClassificationaccuracyPsychiatricinterview,see CR/MCIDConstruct validitySee CR/MCIDConvergent validitySpearman's ρInterview/self-ratingDepression/Depression: 0.79Anxiety/Anxiety:0.54DiscriminantvaliditySpearman's ρInterview/self-ratingDepression/Depression: 0.79Anxiety/Anxiety:0.54DiscriminantvaliditySpearman's ρInterview/self-ratingDepression/AnxietynsAnxiety/Depressionns	Test-retest Not assessed Internal consistency Spearman's ρ Anxiety: 0.41– 0.76 Depression: 0.30–0.60 Inter-rater reliability Not assessed	Cut-offs (subscales) Depression Absent:≤ 7 Borderline: 8–10 Definite: ≥ 11 • False positives: 1 % • False negatives: 1 % Borderline not counted as error Anxiety Absent:≤ 7 Doubtful: 8–10 Definite: ≥ 11 • False positives: 5 % • False negatives: 1 % Borderline not counted as error <u>MCID</u> Not assessed
		Criterion validity See CR/MCID		

Outcome	Description	Validity	Reliability	CR/MCID
Insomnia Severity	Self-reported, one-	<b>Classification</b>	Test-retest	Control sample (self-
Index (ISI)[7],	month retrospective	<u>accuracy</u>	Not assessed	<u>diagnosis)</u>
patient version	screening tool for	Insomnia (yes/no)		Cut-off (total score):
	insomnia in adults (≥18	ROC analyses, see	Internal	10
	years)	MCID	<u>consistency</u>	<ul> <li>Sensitivity: 86 %</li> </ul>
	7 items		Cronbach's α,	• Specificity: 88 %
	5-point Likert scales (0–	Construct validity	Control	• Accuracy: 87 %
	4, no problem to severe	See CR/MCID	sample: 0.71	
	problem)	Pearson's r	Clinical	Clinical sample
	Total score: 0–28, lower	• Daily sleep diary:	sample: 0.73	Cut-off (total score):
	is better	0.54-0.59		11
	• 0–7: Absence of	<ul> <li>Activity level,</li> </ul>	Inter-rater	• Sensitivity: 97 %
	insomnia	Anxiety (state,	<u>reliability</u>	• Specificity: 100%
	• 8–14: Subthreshold	trait),	Not assessed	Accuracy: 98 %
	insomnia	Depression,		
	• 15–21: Moderate	Fatigue (general,		MCID
	insomnia	physical, mental),		Change required for
	• 22–28: Severe	Motivation: 0.20-		improvement
	insomnia	0.48		Blinded assessor, M,
	Dimensions:			[Cl <sub>95</sub> ]:
	<ul> <li>Severity of sleep onset</li> </ul>	Criterion validity		• Slight: 4.65 [2.61–
	Sleep maintenance	Pearson's r		6.69]
	<ul> <li>Early morning</li> </ul>	Polysomnography		Moderate: 8.36
	awakening problems	<ul> <li>Sleep onset</li> </ul>		[7.20–9.53]
	<ul> <li>Sleep dissatisfaction</li> </ul>	latency: ns		• Marked: 9.89
	<ul> <li>Interference of sleep</li> </ul>	<ul> <li>Wake after sleep</li> </ul>		[8.74–11.04]
	difficulties with	onset: ns		ROC analyses:
	daytime functioning	<ul> <li>Number of</li> </ul>		• Slight: not reported
	<ul> <li>Noticeability of sleep</li> </ul>	awakenings: ns		<ul> <li>Moderate: ≥7</li> </ul>
	problems by others	<ul> <li>Early morning</li> </ul>		<ul> <li>Sensitivity: 60 %</li> </ul>
	<ul> <li>Distress caused by the</li> </ul>	awakening: ns		<ul> <li>Specificity: 70 %</li> </ul>
	sleep difficulties	<ul> <li>Total wake time:</li> </ul>		<ul> <li>Accuracy: not</li> </ul>
		ns		reported
	Validation samples	• Sleep efficiency: -		<ul> <li>Marked: ≥8</li> </ul>
	959 adults with and	0.16		<ul> <li>Sensitivity: 64 %</li> </ul>
	without insomnia			<ul> <li>Specificity: 80 %</li> </ul>
	(community setting), 183			<ul> <li>Accuracy: not</li> </ul>
	adults with insomnia and			reported
	62 controls (clinical			
	setting); English			
	language; Québec,			
	Canada.			

Outcome	Description	Validity	Reliability	CR/MCID
SleepSuite[8]	SleepSuite bubble tasks	<u>Classification</u>	<u>Test-retest</u>	Not assessed
(iPad App):	(iPad games) are	<u>accuracy</u>	Delay	
Bubble task	adapted from a validated	Not assessed	unspecified	
	Balloon Task[9]: The goal		(likely none	
<ul> <li>Executive</li> </ul>	is to burst upward	Construct validity	[immediate	
function	drifting balloons with	Not assessed	retest])	
(accuracy and	children's faces under			
response times	multiple target	Criterion validity	Pearson's r	
[RT])	conditions (e.g. happy	Child Behavior	• Hits: 0.60	
	faces only) and at	Checklist (CBCL) <u>:</u>	• Misses: 0.37	
	increasing presentation	total score, sub-	<ul> <li>Completed</li> </ul>	
	conditions (speed, load:	scales (8), recode to	levels: 0.39	
	number of faces shown	externalising and	• RT: 0.78	
	simultaneously).	internalising		
		behaviours.	Internal	
	Validation sample[9]		<u>consistency</u>	
	134 healthy children (7–	Pearson's <i>r</i> (age	Not assessed	
	12 years, 58 boys, 23	and sex partialled		
	with clinical behavioural	out), across	Inter-rater	
	problems, 40% first-	conditions	<u>reliability</u>	
	born) from middle- and		Not assessed	
	upper-class families of	Completed		
	which 25% included at	levels/RT		
	least one parent who	<ul> <li>Total score: -</li> </ul>		
	immigrated more than	0.24/ns		
	10 years ago. Children	<ul> <li>Delinquency:</li> </ul>		
	lived with their parents	ns/0.18		
	in small households (on	<ul> <li>Aggression: -</li> </ul>		
	average 4.53 members).	0.20/0.23		
	Parents were largely	<ul> <li>Attention</li> </ul>		
	employed full-time	problems: -		
	(fathers: 90.71%,	0.18/ns		
	mothers: 49.31%) and	<ul> <li>Social</li> </ul>		
	well educated (on	withdrawal: -		
	average for 16 years).	0.24/ns		
	Community setting	<ul> <li>Somatic</li> </ul>		
	(school, number	complaints:		
	unspecified); paid	ns/0.18		
	participation (\$15 school	<ul> <li>Thought</li> </ul>		
	supply voucher);	disorders: ns/ns		
	language: Hebrew,	<ul> <li>Anxiety-</li> </ul>		
	Israel.	Depression: -		
		.28/ns		
		<ul> <li>Social problems: -</li> </ul>		
		0.20/ns		
		<ul> <li>Externalising</li> </ul>		
		behaviours: -		
		0.18/0.23		
		<ul> <li>Internalising</li> </ul>		
		behaviours: -		
		0.25/ns		

Outcome	Description	Validity	Reliability	CR/MCID
Health-Related	Quality of life	<u>Classification</u>	Test-retest	Not assessed
Quality Of Life	assessment tool for	<u>accuracy</u>	10– 14 days	
Measure for	children or parents with	Not assessed	delay	
<b>Ch</b> ildren with	epilepsy (no specified		Intraclass	
Epilepsy	time-period); child	Construct validity	correlation	
(CHEQOL)[10]	reported if $\geq$ 8 years,	(child)	coefficient	
	parent proxy-report if	Pearson's r	Child: 0.59–	
	child 5 to <8 years	<ul> <li>Health care</li> </ul>	0.69	
	25 items	utilisation: 0.13–	Parent: 0.60–	
	4-point Likert scales (0–	0.31	0.81	
	4, opposites: true/sort of	<ul> <li>Drug Adverse</li> </ul>		
	true)	Events: 0.18–0.25	<u>Internal</u>	
	Total score: 25–100,	Number of	<u>consistency</u>	
	higher is better	friends: 0.18	Cronbach's α,	
	5 subscales (5 items	<ul> <li>N° of</li> </ul>	subscales	
	each):	extracurricular	Child: 0.63–	
	<ul> <li>Interpersonal/social</li> </ul>	activities: 0.13	0.84	
	consequences	One-way ANOVA (p	Parent: 0.64–	
	<ul> <li>Future worries</li> </ul>	≤ .05)	0.86	
	<ul> <li>Present worries</li> </ul>	<ul> <li>Seizure severity:</li> </ul>		
	<ul> <li>Intrapersonal/emotion</li> </ul>	All 5 subscales	Inter-rater	
	al	Anti-epileptic	<u>reliability</u>	
	• Epilepsy secrecy	drug use: 4	Pearson's r	
		subscales	<ul> <li>Child/mothe</li> </ul>	
	Validation samples	$t$ -tests ( $p \le .05$ )	r: 0.24–0.56	
	381 children (6–15	<ul> <li>Help at school:</li> </ul>	<ul> <li>Child/father</li> </ul>	
	years) with epilepsy and	All 5 subscales	: 0.18–0.54	
	their parents (clinical	Results for parent-	<ul> <li>Mother/fath</li> </ul>	
	setting); English	proxy similar	er: 0.40–	
	language; Ontario,		0.71	
	Canada. Test-retest:	Criterion validity		
	Additional 89, then 31	Not assessed		
	children; additional 48	NOT 85565560		
	parents.			
	Metrics refer to self-			
	report for children 8–15			
	years and parent proxy			
	report for children 5 to			
	<8 years and were			
	assessed for sub-scales,			
	not total score.			
		I	1	1

Outcome	Description	Validity	Reliability	CR/MCID
Outcome World Health Organisation – Five Well-Being Index (WHO- 5)[11]	DescriptionSelf-reported, two-week retrospective tool to assess subjective psychological well-being in people aged 9 years and older.5 items 6-point Likert scales (0– 5, 'at no time' to 'all the time') Raw score: 0–25 Total score multiplied by 4 to give final score: 0– 100, higher is betterValidation samples 446 children analysed (9–12 years, 16 [3.6 %] with depressive 	Validity <u>Classification</u> <u>accuracy</u> Depressive disorder (yes/no) Receiver Operating Characteristic (ROC) analyses: See CR/MCID <u>Construct validity</u> See CR/MCID <u>Criterion validity</u> Diagnostic and Statistical Manual of Mental Disorders (DSM-IV) criteria for depressive disorder (major or minor depression only, dysthymia dropped due to mismatch in time- period of concept definitions), see CR/MCID.	ReliabilityTest-retestNot assessedInternalconsistencyNot assessedInter-raterreliabilityCohen's k =.90	CR/MCID Cut-off (total score): 10 • Sensitivity: 75 % • Specificity: 92 % • Accuracy: 88 % <u>MCID</u> Not assessed

Outcome	Description	Validity	Reliability	CR/MCID
Strengths and	Parent-, teacher-, or	<u>Classification</u>	Test-retest	Cut-off (total score):
Difficulties	child-reported,	<u>accuracy</u>	Not assessed	17
Questionnaire	retrospective screening	Psychiatric disorder		<ul> <li>Sensitivity: 88 %</li> </ul>
(SDQ)[12]	tool of child	(yes/no)	Internal	• Specificity: 59 %
	psychopathology (2–18	Receiver Operating	consistency	• Accuracy: 74 %
	years). Retrospective	Characteristic	Cronbach's α:	
	period: 6 months or	(ROC) analyses: See	0.84	MCID
	current school year	CR/MCID		Not assessed
		Original total score	Inter-rater	
	25 items	cut-offs:	<u>reliability</u>	
	3-point Likert scales (0–	• Normal: 0–13	Not assessed.	
	2,	<ul> <li>Borderline: 14–</li> </ul>		
	not/somewhat/certainly	16		
	true)	<ul> <li>Abnormal: 17–40</li> </ul>		
	Total score: 0–40, lower	transformed to		
	is better	binary:		
	5 subscales (5 items	• No: 0–16		
	each):	• Yes: 17–40		
	<ul> <li>hyperactivity/inattenti</li> </ul>			
	on,	Construct validity		
	<ul> <li>emotional problems</li> </ul>	See CR/MCID		
	<ul> <li>conduct problems</li> </ul>			
	<ul> <li>peer problems</li> </ul>	Criterion validity		
	<ul> <li>prosocial behaviours</li> </ul>	Diagnostic and		
	(omitted from total	Statistical Manual		
	score)	of Mental Disorders		
		(DSM-IV), see		
	Validation samples	CR/MCID.		
	541 children (5–12			
	years) with and without			
	psychiatric disorders			
	(school setting); multiple			
	languages; Italy,			
	Germany, the			
	Netherlands, Lithuania,			
	Bulgaria, Romania, and			
	Turkey. Metrics refer to			
	parent-report, total			
	score, and data			
	aggregated across			
	countries and psychiatric			
	disorders.			

Outcome	Description	Validity	Reliability	CR/MCID
Parenting Self	Self-reported tool	<u>Classification</u>	<u>Test-retest</u>	Not assessed
Agency Measure	assessing overall	<u>accuracy</u>	Not assessed	
(PSAM)[13]	confidence to	Not assessed		
	successfully parent		<u>Internal</u>	
	(including managing the	Construct validity	<u>consistency</u>	
	child's behaviour and	Convergent validity	Cronbach's α:	
	resolving problems with	Pearson's r	0.70	
	the child). The time-	Active coping: 0.31	Comparative	
	period for parental self-	Parenting	Fit Index: 0.94	
	assessment is	acceptance: 0.55		
	unspecified.	Positive re-	Inter-rater	
		interpretation: ns	reliability	
	5 items		Not assessed	
	7-point Likert scales (1–	Discriminant		
	7, rarely to always)	validity		
	Total score: 5–35, higher	Pearson's r		
	is better	Inconsistent		
		parental		
	Validation sample	disciplining: -0.34		
	90 English-speaking	Acceptance coping:		
	mothers (all European-	ns		
	American, median age			
	36–40 years, median	Criterion validity		
	annual income >\$40,000,	Not assessed		
	median education			
	bachelor's degree, 82%			
	•			
	married or co-habiting)			
	of 3–12-year-olds			
	(community setting); 2			
	day-care centres and			
	classes at a large			
	university, 2 churches.			
	English language,			
	southwestern USA.			

Outcome	Description	Validity	Reliability	CR/MCID
Actigraphy: Micro	The Micro-	<u>Classification</u>	Test-retest	Not assessed
Motionlogger <sup>®</sup>	Motionlogger <sup>®</sup> Watch	<u>accuracy</u>	Not assessed	
Watch,	directly measures 3-D	Not assessed		
Watchware	acceleration (in CASTLE		Internal	
Software V	Sleep-E and the	Construct validity	consistency	
1.99.17.4, Action-	referenced validation	Not assessed	Not assessed	
W software, V	study of the non-			
2.7.3285	dominant wrist). Raw	Criterion validity		
(Ambulatory	data (zero-crossing	Agreement of	Inter-rater	
Monitoring, Inc.,	mode) is initially	actigraphy with	reliability	
NY: USA)	recorded as periods of	continuous video-	Not assessed	
combined with	activity and inactivity (1	electroencephalogr	Not assessed	
sleep diaries	min epochs), and then	aphy (24 hours),		
(Child and	recoded into periods of	scored by		
Parent)	wakefulness and sleep	neurologist and		
	using a combination of	neurophysiologist.		
<ul> <li>Total sleep time</li> </ul>	proprietary algorithms			
(minutes)	and manual processing	Bland-Altman plots		
<ul> <li>Sleep latency</li> </ul>	(e.g. sleep periods are	in combination		
(minutes)	visually inspected and	with <i>t</i> -tests for		
<ul> <li>Sleep efficiency</li> </ul>	manually corrected with	significant bias:		
(% asleep of	the aid of participant	<ul> <li>Total sleep time</li> </ul>		
sleep period)	sleep diaries). Sleep- and	(minutes): Bias =		
sicep period)	wake parameters are	8.3 (SD = 31), n.s.		
All 2-week	then calculated	Wake duration:		
	automatically using	Bias = -4.8 (SD =		
averages	validated public	31.1), n.s.		
	algorithms.	,,		
		Pearson's r:		
	Validation sample[9]	<ul> <li>Total sleep time</li> </ul>		
	27 children (3–17 years)	(minutes): 0.96		
	with medically refractory	Wake duration:		
	epilepsy, of which 12	0.93		
	had parent-indicated	0.00		
	sleep problems (44%).			
	Hospital setting (in-			
	patient epilepsy			
	monitoring unit in			
	tertiary paediatric			
	hospital), English			
	language, Toronto,			
	Canada.			
	Culludu.	l		

Table 9. Estimated overall time requirement for CASTLE Sleep-E (participant perspective). Time estimates for questionnaires/instruments are based on published estimates where available, and otherwise on an estimate (indicated by \*) of 30 seconds per item derived from the Children's Sleep Habits Questionnaire (35 items, 10 minutes published completion time), plus an arbitrary estimate of 2 minutes to read instructions and consider responses. The total time requirement for participation in CASTLE Sleep-E varies from minimally 2 hours per month over a 6-month period in the Standard Care arm omitting optional qualitative interviews to maximally 3 hours per month over a 6-month period in the intervention arm including optional qualitative interviews.

Trial component	Time (mins)	Frequency	Overall time (mins)
Study visits (4)			150 minutes
Remote or in-person, combinable with standard care visits			
Consent and baseline data	• 60 minutes	• 1	
Randomisation	• 30 minutes	• 1	
<ul> <li>Follow-up at 3 months</li> </ul>	• 30 minutes	• 1	
Follow-up at 6 months	• 30 minutes	• 1	
Questionnaires/instruments in order of the participant timeline shown in Table 4			246.5 minutes
<ul> <li>Children's Sleep Habits Questionnaire[1], 35 items</li> </ul>	• 10 minutes	• 3	• 30 minutes
<ul> <li>World Health Organisation – Five Well-Being Index[11], 5 items</li> </ul>	• 5 minutes	• 2	• 10 minutes
<ul> <li>Health-Related Quality Of Life Measure for Children with Epilepsy[10], 25 items</li> </ul>	• 12.5 + 2 minutes*	• 2	• 29 minutes
<ul> <li>Strengths and Difficulties Questionnaire[12], 25 items</li> </ul>	• 12.5 + 2 minutes*	• 3	• 43.5 minutes
<ul> <li>Child Health Utility Index 9D (CHU-9D)/CHU-9D proxy[4], 9 items</li> </ul>	• 4.5 + 2 minutes*	• 3	• 19.5 minutes
• EQ-5D-Y/EQ-5D-Y proxy[2], 15 items	• 5 minutes	• 3	• 15 minutes
<ul> <li>EQ-5D-5L[5], 25 items (note: Published time estimate same as for EQ-5D-Y [15 items])</li> </ul>	• 5 minutes	• 3	• 15 minutes
<ul> <li>Parenting Self Agency Measure[13], 5 items</li> </ul>	• 2.5 + 2 minutes*	• 3	• 13.5 minutes
<ul> <li>Insomnia Severity Index[7], patient version, 7 items</li> </ul>	• 3.5 + 2 minutes*	• 3	• 16.5 minutes
<ul> <li>Hospital Anxiety and Depression Scale[6], 14 items</li> </ul>	• 5 minutes	• 3	• 15 minutes
<ul> <li>Resource Use questionnaire (custom instrument), 11 items</li> </ul>	• 5.5 + 2 minutes*	• 3	• 22.5 minutes
<ul> <li>Knowledge About Sleep in Childhood (custom scale), 13 items</li> </ul>	• 6.5 + 2 minutes*	• 2	• 21 minutes
SleepSuite[8] (iPad App)	40 minutes	2	80 minutes
<ul> <li>Morning of single day</li> </ul>	• 20 minutes		
Evening of single day	• 20 minutes		

Trial component	Time (mins)	Frequency	Overall time (mins)
Actigraphy			74 minutes
<ul> <li>Delivery arrangements to participants' home or collection point (incl. SleepSuite iPad)</li> </ul>			
○ Baseline	<ul> <li>15 minutes</li> </ul>	• 1	
<ul> <li>Follow-up at 3 months</li> </ul>	<ul> <li>15 minutes</li> </ul>	• 1	
<ul> <li>Return arrangements to participants' home or collection point (incl. SleepSuite iPad)</li> </ul>			
<ul> <li>Baseline</li> </ul>	<ul> <li>15 minutes</li> </ul>	• 1	
$\circ$ Follow-up at 3 months	<ul> <li>15 minutes</li> </ul>	• 1	
• Use: Removal and re-fitting of device once daily (2 x 0.25 minute) when showering, bathing, or swimming;			
otherwise, the device is worn like a wristwatch without requiring participant interventions.			
<ul> <li>Baseline: 14 days</li> </ul>	• 7 minutes	• 1	
$\circ$ Follow-up at 3 months: 14 days	• 7 minutes	• 1	
Sleep diary			140 minutes
Once daily completion of parent- and child diary (2 x 2.5 minutes)			
• Baseline: 14 days	<ul> <li>70 minutes</li> </ul>	• 1	
<ul> <li>Follow-up at 3 months: 14 days</li> </ul>	<ul> <li>70 minutes</li> </ul>	• 1	
COSI (intervention arm only)			245.5 minutes
<ul> <li>3 mandatory modules (core information about sleep relevant to all families)</li> </ul>	<ul> <li>60 minutes</li> </ul>	• 1	
<ul> <li>3 recommended modules (e.g. sleep hygiene)</li> </ul>	• 60 minutes	• 1	
<ul> <li>5 tailored modules (addressing specific sleep issues indicated by a given parent)</li> </ul>	• 100 minutes	• 1	
<ul> <li>List of additional resources, optional, 10 webpages, not included in time estimate</li> </ul>	• 0 minutes	• 1	
<ul> <li>Evaluation questionnaire, 3 sections, 47 items overall</li> </ul>	• 23.5 + 2 minutes*	• 1	
A parent assigned to COSI (i.e. the intervention arm) would be expected to look at minimally 7 and			
maximally 11 modules. All modules are self-paced (i.e. do not have a fixed duration). To read and engage			
with a single module could take anywhere between 5–20 minutes depending on how quickly one reads,			
whether one watches the videos, does the quizzes, etc. Consequently, the estimated time requirement for			
initial material completion not including breaks or re-visits is 35–220 minutes for modules alone.			
To be conservative, maximal estimates are used in calculations.			

Trial component	Time (mins)	Frequency	Overall time (mins)
Qualitative interviews (optional)			140 minutes
Two time-points (Follow-up at 3 months + 3 weeks, at 6 months + 3 weeks)			
<ul> <li>Interview date and time arrangement</li> </ul>	• 10 minutes	• 2	• 20 minutes
<ul> <li>Interview preparation using supplied interview guide</li> </ul>	• 10 minutes	• 2	• 20 minutes
Actual interview	• 40 minutes	• 2	• 80 minutes
De-brief	• 10 minutes	• 2	• 20 minutes
For the qualitative interviews with parents, we typically expect that the total time burden for each of the			
two interviews would range from 30–70 minutes. However, we will tailor the core interview to fit with the			
time the parent has available, so some interviews may be a little longer or shorter.			
To be conservative, maximal estimates are used in calculations.			
Total time for participation over a 6-months period			
<ul> <li>Standard Care arm (SC), not participating in optional qualitative interviews</li> </ul>			• 690.5 minutes
<ul> <li>Standard Care arm (SC), participating in optional qualitative interviews</li> </ul>			• 830.50 minutes
<ul> <li>Intervention arm (SC + COSI), not participating in optional qualitative interviews</li> </ul>			• 936 minutes
<ul> <li>Intervention arm (SC + COSI), participating in optional qualitative interviews</li> </ul>			• 1076 minutes

Supplemental Table 10. Categories used to define the causality and severity of Adverse Events in CASTLE Sleep-E

Category	Definition			
Causality				
Almost Certainly	There is clear evidence to suggest a causal relationship, and other possible contributing factors can be ruled out.			
Probably	There is evidence to suggest a causal relationship, and the influence of other factors is unlikely.			
Possibly	There is some evidence to suggest a causal relationship (e.g. the event occurred within a reasonable time after administration of the study procedure). However, the influence of other factors may have contributed to the event (e.g. the participant's clinical condition, other concomitant events).			
Unlikely	There is little evidence to suggest there is a causal relationship (e.g. the event did not occur within a reasonable time after administration of the study procedure). There is another reasonable explanation for the event (e.g. the participant's clinical condition).			
Not related	There is no evidence of any causal relationship.			
Severity				
Mild	The Adverse Event does not interfere with the participant's daily routine and does not require further procedure; it causes slight discomfort.			
Moderate	The Adverse Event interferes with some aspects of the participant's routine, or requires further procedure, but is not damaging to health; it causes moderate discomfort.			
Severe	The Adverse Event results in alteration, discomfort or disability which is clearly damaging to health.			

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