Community-based parent-delivered early detection and intervention programme for infants at high risk of cerebral palsy in a low-resource country (Learning through Everyday Activities with Parents (LEAP-CP): protocol for a randomised controlled trial)

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ABSTRACT

Introduction Cerebral palsy (CP) is the most common childhood physical disability, with 80% estimated to be in low-middle-income countries. This study aims to (1) determine the accuracy of General Movements (GMs)/Hammersmith Infant Neurological Examination (HINE) for detecting CP at 18 months corrected age (CA); (2) determine the effectiveness of a community-based parent-delivered early intervention for infants at high risk of CP in West Bengal, India (Learning through Everyday Activities with Parents for infants with CP; LEAP-CP).

Methods This study comprises two substudies: (1) a study of the predictive validity of the GMs and HINE for detecting CP; (2) randomised, double-blinded controlled trial of a novel intervention delivered through peer trainers (Community Disability Workers, CDW) compared with health advice (15 fortnightly visits). 142 infants at high risk of CP (‘absent fidgety’ GMs; ‘high risk score’ on HINE) aged 12–40 weeks CA will be recruited to the intervention substudy, with infants randomised based on a computer-generated sequence. Researchers will be masked to group allocation, and caregivers and CDWs naïve to intervention status. Visits will include therapeutic modules (goal-directed active motor/cognitive strategies and LEAP-CP games) and parent education. Health advice is based on the Integrated Management of Childhood Illness, WHO. Infants will be evaluated at baseline, post intervention and 18 months CA. The primary hypothesis is that infants receiving LEAP-CP will have greater scaled scores on the Pediatric Evaluation of Disability Inventory—Computer Adaptive Test (mobility domain) at 18 months compared with health advice. Secondary outcomes include infant functional motor, cognitive, visual and communication development; infant growth; maternal mental health.

Ethics and dissemination This study is approved through appropriate Australian and Indian ethics committees (see in text) with families providing written informed consent. Findings from this trial will be disseminated through peer-reviewed journal publications and conference presentations.

Strengths and limitations of this study

► This study is an adequately powered randomised double-blinded controlled trial of a novel parent-delivered early intervention for infants at risk of cerebral palsy in a low-middle-income country.
► Outcomes are evaluated with standardised measures and evaluate a range of infant and family domains, including functional motor, cognitive, visual and communication developmental outcomes; infant growth and maternal mental health.
► This is a pragmatic Randomised Controlled Trial; as such the potential contamination of other ‘care as usual’ interventions and other ‘real world’ factors may influence the data and its interpretation.
rates and larger populations, it is proposed that 80% of CP cases globally are in LMICs, where individuals and their families are frequently trapped in the negative downward cycle of disability and poverty. Individuals with a disability and their families in LMIC have increased rates of premature mortality and associated morbidities and are economically disadvantaged by productivity loss, costs of interventions and equipment as well as consequences of social stigma. Significant gains have been made in the past decade in reducing infant mortality in LMIC. With a renewed global international development strategy pledged by the United Nations Sustainable Development Goals (2015–2030), it is pertinent to broaden attention from infant survival and to improved quality of life and developmental outcomes for children with a disability.

In many LMICs, there is a shortage of specialised health workers to attend to the often larger populations and greater disease/disability rates. A recent multisite early intervention study (spanning Africa and Asia) for infants with birth asphyxia advocated for the need to train non-professional individuals to address this workforce gap. Lay health workers have been used as effective change agents across Asia, sub-Saharan Africa and Latin America, to improve outcomes for both communicable and non-communicable diseases. The current project aims to adapt this model for an early disability intervention for infants at risk of CP. The Indian National Rural Health Mission has been instrumental in establishing a community health worker programme to meet primary health needs in both urban and rural areas across the country (known as USHA/ASHA, Urban/Accredited Social Health Activists). West Bengal is an easterly state in India and with a population of 90 million people is considered one of the most densely populated geographies in the world. These factors collectively make West Bengal an important location for piloting and implementing innovative interventions and service delivery models that are highly scalable and transposable for disability intervention in other LMICs.

Recently published International Clinical Practice Guidelines advocate for CP detection from as young as 3 months using the General Movements (GMs) Assessment, and there is growing evidence to support the effectiveness of early cognitive and early active motor interventions in the first year of life for infants at high risk of CP. Despite this state of science, many children in LMICs only receive their diagnosis at the age of school entry, missing a significant window of opportunity for improved outcomes. Children with CP in these settings also face economic, geographical and social barriers to accessing medical and rehabilitation interventions. In addition to the impact of disability on the child, there is strong evidence of higher prevalence of poor mental health in mothers of children with CP, for which there are demonstrated improvements consequent to early intervention for their child. In order to impact on the social inclusion and workforce productivity of individuals with a disability in these contexts, it is essential to first establish innovative, accessible and feasible means to detect infants at risk of CP and subsequently develop early intervention programmes that are accessible for families of high-risk infants in these settings, are cost-effective and can be widely delivered. The GMs Assessment and Hammersmith Infant Neurological Examination (HINE) are two gold standard early detection tools based on systematic review, with demonstrated predictive validity for identifying later CP (GMs sensitivity: 95%–100%, specificity: 96%–98%; HINE sensitivity: 90%–96%, specificity: 85%–91%). Both have simple administration (the GMs a 3 min video taken on a smart phone, the HINE a truncated 26-item neurological examination) which we believe can be feasibly administered in the community context in LMIC and scored by expert certified raters.

A community-based intervention of active goal-directed strategies and environmental enrichment delivered peer to peer in the home also presents a viable solution for accessible and scalable intervention in this context. The aims of this study are twofold; to determine the predictive validity of a community-based early detection programme for infants at risk of CP in LMIC and to determine the effectiveness of a home-based peer delivered early intervention.

**METHODS AND ANALYSIS**

**Study design**

This study consists of two substudies: (1) a study of the validity of an early community-based detection programme for identifying infants at high risk of CP and (2) a randomised, double-blinded controlled trial of a novel intervention (LEAP-CP: Learning through Everyday Activities with Parents for infants at high risk of Cerebral Palsy) compared with health advice (standard care). Field work will be conducted from March 2017 to March 2019.

**Aims and hypotheses**

**Early intervention for infants at high risk of cerebral palsy in low-middle-income countries**

1. To determine the effectiveness of a community-based parent-delivered intervention on infant’s developmental outcomes for those at high risk of CP.

**Hypothesis 1**: Infants with CP who receive the LEAP-CP intervention will have higher scores on the mobility domain of the Pediatric Evaluation of Disability Inventory Computer Adaptive Test (PEDI-CAT) compared with infants of caregivers receiving health advice.

2. To determine the effectiveness of a community-based parent-delivered intervention on caregiver’s mental health outcomes.

**Hypothesis 2**: Caregivers who receive the LEAP-CP intervention will have reduced depression and anxiety scores compared with caregivers receiving health advice.

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Early detection of infants at high risk of CP in LMIC

3. To determine the predictive validity of GMs assessment administered at 12–17 weeks for detecting CP at 18 months in high-risk infants in West Bengal.

4. To determine the predictive validity of the HINE when administered from 18 to 40 weeks for detecting CP at 18 months in high-risk infants in West Bengal.

Hypothesis 3: The GMs and HINE assessments will have predictive validity equivalent to that in high-income countries to detect CP at 18 months.

Recruitment

Participants

A total of 142 infants at high risk of CP aged 12–40 weeks corrected age (CA) will be recruited to the intervention trial. Infants will be screened until the target recruitment for the intervention substudy is achieved (recruitment pathways are shown in figure 1). The caregiver and their infant will be referred to the detection substudy by health professionals or Community Health Workers, from regional and tertiary hospitals, community health centres, routine immunisation clinics or in the community. Three implementing partner organisations were selected based on their existing expertise in the field of childhood disability, community-based maternal child health or neonatology. Corresponding geographical areas were selected based on the geographical catchment of the three partner organisations.

i. *Asha Bhavan Centre* is a community-based non-government (non-profit) organisation providing multidisciplinary rehabilitation and support to underprivileged children with a disability in rural West Bengal. The catchment for this site is Uluberia I, Bagnan II, Shyampur II (Howrah District), reaching a population of 575,961 and 240 villages (Census of India 2011).21
At 12–17 weeks CA infants with absent/abnormal fidgety movements on GMs assessment are considered high risk. The diagnostic accuracy of the GMs has been well documented, with sensitivity ranging from 95% to 100% and specificity 96% to 98%, with the measure also shown to have robust reliability (inter-rater kappa=0.91–0.92, average agreement 90%). GMs will be videoed by trained Community Coordinators/Community Disability Workers (CDWs) and videos rated independently by two GMs Trust accredited scorers, of whom there are >20 advanced and >100 basic raters in India alone. In the case of disagreement, a third accredited rater will provide consensus.

At 18–40 weeks, CA infants scoring below the established HINE cut-points will be considered high risk of CP (<56 points at 3 months (SE 96%; sp 85%), <59 points at 6 months (SE 90%; sp 89%), <62 points at 9 months (SE 90%; sp 91%). The HINE has been shown to have excellent intrarater and inter-rater reliability for the global scores (Intra-Class Correlation Coefficient=0.97 for both) when administered to high-risk 12-month-old infants in India. The scoring form provides simple instructions for administration and scoring and is accessible for all clinicians, regardless of experience. All HINE examinations will be videoed and conducted by trained Community Coordinators (Indian trained therapists). Videos will be rated independently by two HINE trained scorers. In the case of disagreement (ie, of ‘high risk’ status), a third rater will provide consensus.

### Inclusion/exclusion criteria

To participate, infants must live in one of the study geographical areas and be 12–40 weeks CA. Infants with known or suspected congenital or chromosomal abnormalities which are likely to affect their neurodevelopmental outcome; those diagnosed with neurodegenerative conditions and those that are considered medically fragile will be excluded. Infants must also have one or more risk factors:

- Maternal infection (antenatal).
- Low birth weight (<2.5 kg).
- Preterm delivery (<37 weeks).
- Hypoxic ischaemic encephalopathy.
- Perinatal asphyxia.
- Neonatal jaundice requiring treatment.
- Prolonged hypoglycaemia.
- Seizures after birth.
- Admission to neonatal intensive care unit or special newborn care unit.
- Postneonatal complications in infant (infection, head injury, near drowning).
- Altered tone or delayed motor milestones for the infant.

To be eligible for the intervention substudy, infants must be assessed in the detection substudy to be at ‘high risk’ of CP based on the GMs or HINE, as follows:

- At 12–17 weeks CA infants with absent/abnormal fidgety movements on GMs assessment are considered high risk. The diagnostic accuracy of the GMs has been well documented, with sensitivity ranging from 95% to 100% and specificity 96% to 98%, with the measure also shown to have robust reliability (inter-rater kappa=0.91–0.92, average agreement 90%). GMs will be videoed by trained Community Coordinators/Community Disability Workers (CDWs) and videos rated independently by two GMs Trust accredited scorers, of whom there are >20 advanced and >100 basic raters in India alone. In the case of disagreement, a third accredited rater will provide consensus.

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### Sample size calculation

#### Detection substudy

The primary hypothesis of the detection substudy is that the GMs and HINE will have equivalent predictive validity to that in Western literature. With estimates of the GM sensitivity=0.95 and specificity=0.96, a total of 19 infants identified as high risk of CP and 15 identified as low risk of CP (ie, rated present fidgety on the GMs) will be needed to demonstrate equivalent diagnostic accuracy to 10% precision either side (and 95% confidence). Similarly, with estimates of the HINE sensitivity=0.90 and specificity=0.89, a total of 35 infants identified as high risk of CP and 38 infants identified as low risk of CP (ie, ‘normal’ scores using the HINE) will be needed to demonstrate equivalent diagnostic accuracy to 10% precision either side (and 95% confidence).

#### Intervention substudy

The primary hypothesis of the intervention substudy is that infants receiving the LEAP-CP intervention will have significantly greater scaled scores on the mobility domain of the PEDI-CAT at 18 months compared with those receiving health advice. The PEDI-CAT mobility domain has been shown to detect a 1SD change in scaled scores following 3 or more months of physiotherapy in young children (aged 1 month–3 years). With 64 children per group, a difference of at least 0.5SD between groups will be detectable ($\alpha$=0.05, $\beta$=0.8). Accounting for 10% attrition, this equates to 142 infants (71 in each group).

#### Detection substudy

Infants who have been screened as ‘low risk’ on the GMs and HINE will not be eligible to participate in the intervention substudy. A random sample of 53 infants with negative case status on the screening tests at 12–40 weeks CA (15 GMs screened, 38 HINE screened) will be contacted when the child is 18 months CA for CP diagnosis (as described in the outcome measures). Infants
screened as ‘high risk’ on the GMs or HINE will be invited to participate in the intervention substudy and CP diagnosis provided at 18 months CA. Data from all 142 case infants will be used to calculate sensitivity.

**Intervention substudy**

The LEAP-CP intervention is a multidomain family-centred best practice intervention consisting of infant goal-directed therapeutic strategies and learning games and caregiver educational modules. It is based on effectiveness shown in systematic reviews and early intervention trials. In the development phase of LEAP-CP, in-depth consultation was held with the developers of GAME (Goals-Activity-Motor Enrichment). LEAP-CP includes many of the key ingredients of GAME but adapted for low-income settings and have been used with permission. The components shown necessary for effective interventions for infants with CP include (1) goal-directed tasks; (2) home-based delivery and include (3) active motor learning and (4) strategies to enrich the home environment. LEAP-CP is based on principles of parent coaching which promote caregiver problem solving and self-determination. Specifically, LEAP-CP includes:

- Activity-based motor and cognitive skills training, based on goals identified by parents.
- Practice is structured using motor-learning principles of repetition and variation. Functional motor skills, such as reach/grasp and attaining independent mobility, will be coached and parents given visual supports (photo/video) for ongoing practice through the week.
- Enrichment, which facilitates enhanced cognitive, motor and multisensory learning (eg, visual and auditory), will be encouraged within the home environment using resources based on the Abecedarian Learning Games curriculum modified for CP and adapted for the context. The Abecedarian approach has strong empirical evidence from >16 Randomised Controlled Trials (RCTs) in at-risk children. This includes early play-based learning and literacy activities and promotes use of materials readily and cheaply available in the home and community.

The parent educational modules (table 1) are evidence-based discussion topics which cover three broad areas: (1) ‘learn’—enabling active play and learning for babies with CP; (2) ‘grow’—feeding, nutrition (breastfeeding, complementary feeding, balanced diet) and health; (3) ‘love’—caregiver mental health based on acceptance and commitment therapy and responsive parenting.

**Dose**

The LEAP intervention will commence at 3–9 months CA at a dose of 20 min per day for 5 days per week (1.6 hours) up to 6 months CA (total dose 19.2 hours); then graduate to 30 min per day for 5 days per week (2.5 hours per week) from 6 to 9 months CA (total 30 hours); then 40 min per day for 5 days per week (3.3 hours per week) from 9 to 12 months CA (total 40 hours). In addition, there will be approximately 15 hours of direct intervention administered during home visits by either the parent or CDW. The overall dose will be 104.2 hours for the entire intervention up to 18 months CA.

**Peer to peer service delivery**

The service delivery model for each geographical catchment is represented in figure 2. The programme adopts an iterative coaching model as represented visually in figure 3. The Community Coordinator (health professional) oversees each site and coaches the CDW on new goals and activity targets for the infant at a fortnightly training session (forward loop) as well as providing supported problem solving of previous goals and activities via video recorded sessions (back loop). CDWs are peer trainers from their local communities, with priority for employment given to mothers of a child with a disability. CDWs will coach the caregiver (or other significant people in the infant’s life) on the goal or activity target at fortnightly home visits (forward and back loops). Caregivers are the primary change agent for their infant.

### Table 1. Parent educational modules in the LEAP-CP programme

| Learn:33 73–75 | Grow:58 53 76 77 | Love:73 76 79 |
|----------------|-------------------|---------------|
| Importance of play | Breastfeeding* | Infant and family strengths |
| Providing new experiences (tummy time, being upright, weight-bearing) | Observation of feed | Understanding CP |
| Motivation and success | Complementary feeding | Parent-infant bonding |
| Learning from everyday life | Observation of solids | Responsive parenting |
| Scaffolding play | Balanced diet | Dealing with grief |
| Perseverence vs stress | Preparing safe weaning foods | Support from family |
| Positioning toys | Introducing textured foods | Values and finding joy |
| Problem solving approach | Growth monitoring | Dealing with negative thoughts |
| Learning through repetition | Making the home safe | Creating a parent support group |
| Talking through the day | Health check | Questions from my community |
| Sharing books | Going to the doctor | Self-care |
| Waiting for my baby to respond | | |

*Infants who are not breastfeeding at the time of this module will receive education discussing hydration.

LEAP-CP: Learning through Everyday Activities with Parents-Cerebral Palsy.
The CDWs will receive a 3-day training package at the onset of the programme (with the Chief Investigator, an Australian Government funded Post-Doctoral Fellow and speech pathologist). This will include topics such as:

- Building rapport and a positive therapeutic relationship with caregivers.
- Exploring customs, beliefs and family culture.
- Using everyday opportunities and routines to encourage infant development.
- Observation skills and coaching.
- Motor training and therapeutic principles.
- Understanding typical development and development in children with CP.
- Ethics and research practices.

**Health advice (standard care)**

The health advice is based on the WHO’s Integrated Management of Childhood Illness Key Family Practices. This includes counselling on breastfeeding and introduction of complementary nutrition, hygiene practices, vaccination counselling and management of the sick child. It also includes clinical signs indicating the need for referral to existing health services. It was considered necessary to provide nutrition and health advice to all families in the study, as undernutrition in conjunction with CP has been shown to increase mortality within this context. The same service delivery model and visiting schedule will be used as for the intervention arm (a fortnightly home visit for 15 visits), with a different CDW visiting standard care group families to avoid contamination. There will not be a direct intervention dose delivered to infants in this study arm.

**Concurrent therapies (care as usual)**

Infants from both study arms are able to continue to access medical and therapy support as per their family’s preference. Frequency and duration of access to local medical/therapy services will be recorded fortnightly on the Health Resource Use Form and included in the analysis.

**Randomisation and blinding**

Infants will be randomised to the LEAP-CP intervention arm or health advice (standard care) arm using simple randomisation based on computer-generated sequences, generated and stored centrally. If twins are both eligible, they will be randomised as a single family unit. Allocation concealment will be ensured by using the REDCap V.8.0.2 database (Vanderbilt University, USA 2009) for group assignment at the time of randomisation. The clinician completing the eligibility assessment will be unaware of group allocation. Caregivers receiving the intervention and CDWs administering the intervention will be naïve to intervention status. Researchers assessing the outcomes and analysing the data will be masked to group allocation.

**Figure 2** Service delivery model for LEAP-CP Study. Infographic represents one geographical area, with centralised community coordinator who trains approximately five Community Disability Workers, each working with approximately six infant-caregiver dyads. LEAP-CP, Learning through Everyday Activities with Parents-Cerebral Palsy.

**Figure 3** Coaching model for supporting goals in the LEAP-CP Study. CC, Community Coordinator; CDW Community Disability Worker; LEAP-CP, Learning through Everyday Activities with Parents-Cerebral Palsy.
Fidelity

Intervention fidelity will be monitored at each level of programme delivery, including its delivery to the community coordinator, their delivery to the Community Disability Worker, the CDW delivery to the caregiver and finally implementation of the intervention with the infant (table 2). These levels of fidelity will be monitored to ensure consistency within and across intervention sites for (1) study design (active ingredients); (2) training providers; (2) delivery of treatment; (4) receipt of treatment; (5) patient enactment.41

Adverse events

Any serious adverse events such as injury, prolonged hospitalisation or mortality occurring during programme delivery will be monitored by the Data Safety Monitoring Representative, a non-treating senior medical professional from the Indian context. They will review study retention, compliance/quality of treatment and monitor any adverse or unintended effects on a 12 monthly basis and advise the Chief Investigators regarding whether the adverse events are likely related to the intervention provided in the trial.

Outcome measurement

Baseline, postintervention and final outcome assessments (at 18 months CA) will be conducted by an assessor masked to intervention status, as shown in figure 1. If the postintervention outcome is within 1 month of the final outcome, this assessment point will be excluded. All questionnaires will be translated to Bengali to ensure a single consistent translation is used. All written translations have been back-translated (English to Bengali and back) to ensure translation accuracy.

The literature lacks consensus for the use of Western developmental norms in an Indian/Asian population. Some suggest that cognitive and physical milestones are appropriate, but that self-care skills such as toilet training, washing and dressing are considered to have greater cultural determination.42 Others propose that using western norms will significantly overestimate delayed development in socialisation and motor domains, but underestimate delay for communication and daily living skills.43 Raw scores will be compared in secondary analysis to account for these possible cultural and ethnic differences from standard scores.

Primary infant outcome measures

The infant’s functional outcomes will be assessed using the Pediatric Evaluation of Disability Inventory-Computer Adaptive Test (PEDI-CAT). This is a parent-reported measure of their child’s independence in self-care, mobility and social function (aged birth–20 years).44 The PEDI-CAT has been Rasch-analysed in both children with disabilities and those with typical development. The computerised adaptive version, based on Item-Response Theory, has been shown to increase accuracy and efficiency of administration.45 The raw scores will be converted to standardised scores using normative data (0–100) to measure change in function. The PEDI has been used in non-Western cultural contexts46 and has undergone cultural adaptation in conjunction with the present study based on the PEDI-CAT author’s guidelines (with n=13 Bengali clin-

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**Table 2** Fidelity evaluation plan for the LEAP-CP trial

| Data source                                      | Components of fidelity                          | Coordinator to CDW | CDW to caregiver | Caregiver to infant |
|-------------------------------------------------|-------------------------------------------------|--------------------|------------------|--------------------|
| Study design                                    | Study design                                    | Coordinator to CDW | CDW to caregiver | Caregiver to infant |
| Training providers                              | Training providers                              | Coordinator to CDW | CDW to caregiver | Caregiver to infant |
| Delivery of treatment                           | Delivery of treatment                           | Coordinator to CDW | CDW to caregiver | Caregiver to infant |
| Receipt of treatment                            | Receipt of treatment                            | Coordinator to CDW | CDW to caregiver | Caregiver to infant |
| Patient enactment                               | Patient enactment                               | Coordinator to CDW | CDW to caregiver | Caregiver to infant |

*Checklist based on fidelity tool developed by Sakzewski, Boyd and Ziviani for the REACH trial.80

LEAP-CP, Learning through Everyday Activities with Parents-Cerebral Palsy.
The infant’s motor outcomes will be assessed using the Peabody Developmental Motor Scales—Second edition (PDMS-2), a commonly used measure of motor skills in infants and children aged birth to 6 years. It has demonstrated validity (discriminative and concurrent with the Bayley47 and Gross Motor Function Measure48) and responsiveness in infants with CP.49 50 The PDMS-2 requires a semistandardised kit and standard forms for its administration and should be administered by an allied health professional or special education professional.

The infant’s cognitive and communication outcomes will be assessed using the Bayley Scales of Infant Development III (BSID-III), the gold standard norm-referenced assessment of infant development (0–3 years).51 The BSID-III requires an extensive kit of manipulatives and standard forms for its administration and should be administered by an independent rater at 18 months CA. The COPM is administered using either a paper-based or online form, with training videos for its administration available online.

The infant’s motor outcomes will be assessed using the Canadian Occupational Performance Measure (COPM) will be used to assist caregivers in setting and prioritising goals and measuring parent-perceived change of their infant’s performance of the goal and their own satisfaction with progress. This assessment will be administered by the Community Disability Worker (trained in its administration) in a semistructured interview as part of Educational Module 2 (goal setting). Postintervention assessment will be administered by an independent rater at 18 months CA. The COPM is administered using either a paper-based or online form, with training videos for its administration available online.

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Nutritional status will be determined using length/height and weight which will be converted to z scores using the WHO age and gender referenced data.53 Head circumference and mid-upper arm circumference will also be recorded.

The Health Resource Use Form will provide health usage outcome data as well as being included as a covariate in analyses. It was developed for a large population-based study in Australia54 and has been previously modified and used by our team in Bangladesh.55

Hammersmith Infant Neurological Examination (HINE) will be used to assess infant neurological status and CP severity. The HINE at 3–6 months has been shown to have strong and significant correlations with the GMFCS at 2 years.56 For infants who had the HINE administered for their eligibility assessment, only a postintervention HINE will be administered.

Home Observation for Measurement of the Environment (HOME) Inventory: Infant and Toddler Version is a measure of the quality and quantity of parent and home stimulation, covering six domains of parent responsivity, acceptance and involvement and the home physical environment including availability of learning materials and variety of stimulation.57 The HOME has been used in contexts of disability and low-income countries, including Bangladesh.58 59–61 Parenting Sense of Competence Scale (PSCS) is a commonly used measure of perceived parenting competence and self-esteem (Gilbad-Wallston and Wanderson, 1978, cited in62). It consists of 17 items rated by parents on a 6-point response scale from ‘strongly agree’ to ‘strongly disagree’.

Differential diagnosis of cerebral palsy at 18months CA will be provided by an Australian qualified paediatrician according to published guidelines,53 based on clinical history (on the Physician Checklist) and videoed HINE and Gross Motor Function Classification System (GMFCS) semistructured play session. Children will be classified as ‘definite CP’, ‘suspected CP’ or ‘no CP’. This method has been used in our previous research.56 Confirmed or suspected diagnoses other than CP will be identified by the physician, based on the clinical examination and medical history.

Primary caregiver outcome measure
Caregiver outcomes will be assessed using the Depression, Anxiety, Stress Scale—Short Form (DASS), a self-reported norm-referenced measure of depression, anxiety and stress.64 An official Bengali translation is available on the measure website.65

Covariates and descriptive measures
Physician checklist (PC) was developed for a large population-based study in Australia54 and used by the CIA in Bangladesh.55 It gathers birth and developmental history from the caregiver. Questions include preterm status, birth complications, presence of seizures and medications.

GMFCS is a five-level classification of children’s functional gross motor function. The <2year old age band will be used to classify gross motor function at the completion of the study (18 months CA).66 This will be classified by an Australian physiotherapist from video, as per our previous work in Bangladesh.55 Motor type (spasticity, dyskinesia, hypotonia) and distribution (number of limbs) will be classified by an Australian physiotherapist from video according...
to the Surveillance of CP in Europe. This methodology has been used in this research team’s previous research in Bangladesh. 

- Poverty Measurement Tool will provide a measure of poverty/economic status. This scale was developed in rural Bangladesh to provide a measure of poverty, defined as ‘inadequate fulfilment of basic needs, such as food, clothing, shelter, health, education and social involvement’. Scores range from 24 to 72 with increasing values indicating increasing poverty. The scoring cut-points were validated against wealth rankings of households using participatory rural appraisal methods. It has excellent test-retest reliability and strong internal consistency.

- Multidimensional Scale of Perceived Social Support (MSPSS) measures caregiver’s cognitive social capital, defined as a subjective measure of what people feel, such as those of trust and reciprocity. The MSPSS is a 12-item scale with four items for each source of support, with items rated on a seven-point scale. The measure has good cross-cultural stability, strong internal consistency when tested in a range of samples in a developing country and was significantly associated with two measures of depression and anxiety (the Beck Depression Inventory and the State-Trait Anxiety Inventory).

**Statistical analysis**

Study data will be collected and managed using REDCap (Research Electronic Data Capture) electronic data capture tools hosted The University of Queensland. REDCap is a secure, web-based application designed to support data capture for research studies. All analyses will be undertaken using Stata V.13.1 (Stata, College Station, Texas, USA) with significance set at p<0.05. Professor Robert Ware, lead biostatistician, will provide expert input for the analysis.

**Detection substudy**

The predictive validity of the GMs and HINE will be tested using receiver operating characteristic curves, and sensitivity and specificity calculated. This will be based on the positive/negative ‘high risk of CP’ status on each screening tool (at 12–40 weeks CA) compared with definite or suspected CP/no CP clinical diagnosis at 18 months CA.

**Intervention substudy**

Analyses will adhere with the standard principles for an RCT, primarily focusing on differences in outcome between intervention and standard care arms. Intention-to-treat analysis will be undertaken. A generalised linear model will be run to compare the primary outcome (PEDI-CAT mobility domain score) between intervention groups. The model’s main effects will be intervention group and time (baseline/end of intervention/end of study) and a group-by-time interaction will be included. A generalised linear model will also be run to compare the primary caregiver outcome (DASS) between intervention groups. Again, intervention group and time will be included as main effects as well as a group-by-time interaction. First, the full data set will be analysed and then analyses rerun after stratifying using the covariable of neurological severity (binary HINE score <40). The distributional family will be Gaussian and the identity link will be used. Secondary analyses will use similar methods to compare outcomes at 18 months, considering differences in goal attainment (COPM), motor development (PDMS-2), cognition/communication (BSID-III Scales), growth (anthropometry), caregiver outcomes (PSCS) and health economics. Skewed continuous data will be transformed, and when indicated, non-parametric tests used.

Validity of results will be determined using baseline and descriptive data, including systematic differences between those completing the intervention and drop-outs. Sensitivity analyses will be conducted using multiply imputed data sets to investigate the effect of possibly differential drop-out, as appropriate.

**Patient and public involvement**

This study was conceptualised directly from the author’s (KAB) engagement with children with CP and their families living in Bangladesh. Families were predominately rural-based from across all districts in Bangladesh and attending the Paediatric Unit at the Centre for the Rehabilitation of the Paralysed (Savar, Bangladesh). The service delivery model drew on both formal discussions regarding barriers restricting access to therapy for these families as well as informal observations of families attending direct centre-based therapy programmes. It also builds on findings from the distance learning package trial (NZK).

The CDW employed as project staff are all mothers residing in the same communities as the participants. Four of these CDWs are mothers of a child with a disability, and one worker herself, a person with a disability. These women are all actively involved in the everyday recruitment and conduct of the study and represent the user’s voice. An info-graphic summary of the key findings of the study will be developed at the conclusion of the study and distributed to participating families by their CDW.

**Outcomes and significance**

The proposed study is expected to result in a number of important outcomes spanning several tiers of society, including child, caregiver/family and health systems. Studies in LMIC have shown positive effects of early interventions for the general population and other high risk groups (eg, low birth weight) with significant improvements in children’s cognition, language and social-emotional development. It is anticipated that the proposed intervention package, designed specifically for infants at risk of CP, will result in significant improvements in their language, cognition, mobility and activities of daily living. Consequently, these changes to children’s developmental trajectories are likely to improve their participation in education, employment and community. By reducing the average age of CP detection to the first year of life, high-risk infants are expected to have greater improvements to brain development and
motor/cognitive outcomes. The detection programme is also anticipated to better direct limited health resources to those that are at an increased risk of later diagnosis. Early interventions for infants with a disability have also been associated with reductions in maternal anxiety and depression. In a country where women are often more socially isolated, which is further compounded by the stigma of having a child with a disability, such improvements in maternal mental health are expected to be significant for families and communities. By empowering mothers as disability resource champions in their local communities, the intervention is also expected to have a lasting and far-reaching benefit, beyond the duration of the study. South Asia has been at the heart of the Community Health Worker model development, which has been subsequently up-scaled around the globe. Building on this model, of up-skilling community members with varied levels of formal/technical education to deliver community-based healthcare, the proposed project is expected to result in a cost-effective and feasible model of care for infants with cerebral palsy that is highly scalable and transposable to other LMICs.

ETHICS AND DISSEMINATION
This study is registered with the Australia and New Zealand Clinical Trials Register (12616000653460p). A two-stage consent process will be adopted for this study; caregivers will first provide informed consent for the eligibility assessment (detection substudy), and subsequently, those who are eligible for the intervention (‘high risk of CP’) will then provide informed consent for the clinical trial (intervention substudy). Participant data will be stored and managed according to universal privacy and confidentiality standards. There are no known risks associated with the interventions, and both interventions (LEAP-CP and health advice) are anticipated to provide benefit to the infant and their family. Findings from this trial will be disseminated through peer-reviewed publications and at national and international conference presentations.

STRENGTHS AND LIMITATIONS
This study is an adequately powered randomised double-blinded controlled trial of a novel parent-delivered early intervention for infants at risk of CP in LMICs. Eligibility has been determined with gold standard tools (the GMs assessment (detection substudy), and subsequently, those who are eligible for the intervention (‘high risk of CP’) will then provide informed consent for the clinical trial (intervention substudy). Participant data will be stored and managed according to universal privacy and confidentiality standards. There are no known risks associated with the interventions, and both interventions (LEAP-CP and health advice) are anticipated to provide benefit to the infant and their family. Findings from this trial will be disseminated through peer-reviewed publications and at national and international conference presentations.

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Contributors KB conceptualised the study, secured funding for the study, drafted the manuscript and approved the final manuscript as submitted. She is Chief Investigator overseeing all aspects of the project delivery in West Bengal. RNB conceptualised the study, secured funding for the study, provided critical review.
of the manuscript and approved the final manuscript as submitted. IN, CM and KW conceptualised the study, provided critical review of the manuscript and approved the final manuscript as submitted. RSW advised on statistical design of the study, provided critical review of the manuscript and approved the final manuscript as submitted. AS provided expert advice on the vision components of the programme, including intervention design and measurement. She provided critical review of the manuscript and has approved the final manuscript as submitted. SBL provided expert advice on the nutritional components of the programme, including intervention design and measurement. She provided critical review of the manuscript and approved the final manuscript as submitted. NZK provided expert advice on the cultural aspects of the programme for delivery in South Asia. She provided critical review of the manuscript and approved the final manuscript as submitted. AKG is overseeing the project delivery in West Bengal and provided expert advice on the cultural aspects of the programme for delivery in West Bengal. He provided critical review of the manuscript and approved the final manuscript as submitted. AB hosted the Endeavour scholarship, is the study lead at Apollo Gnieagles Hospital and provided expert advice on the cultural aspects of the programme for delivery in West Bengal. He provided critical review of the manuscript and approved the final manuscript as submitted. GM is the study lead at Child In Need Institute which is an implementing partner. He provided expert advice on the cultural aspects of the programme for delivery in West Bengal. He provided critical review of the manuscript and approved the final manuscript as submitted. DB is the study lead at Asha Bhavan Centre which is an implementing partner. He provided expert advice on the cultural aspects of the programme for delivery in West Bengal. He provided critical review of the manuscript and approved the final manuscript as submitted. ST is the Data Safety Monitoring Representative and provided critical input into the study design and cultural aspects of the programme as submitted. AKG is overseeing the project delivery in West Bengal and provided expert advice on the cultural aspects of the programme for delivery in West Bengal. He provided critical review of the manuscript and approved the final manuscript as submitted. IN, CM and KW conceived of the study, provided critical review of the manuscript and approved the final manuscript as submitted. ST is the Data Safety Monitoring Representative and provided critical input into the study design and cultural aspects of the programme as submitted. ST is the Data Safety Monitoring Representative and also a member of the Apollo Gnieagles Hospital Ethics Committee.

Patient consent Not required.

Ethics approval Children’s Health Queensland Hospital and Health Service Human Research Ethics Committee (HREC/16/QCH/214). The University of Queensland Medical Research Ethics Committee (2016001073), the Apollo Gnieagles Hospital Kolkata Institutional Ethics Committee (IEC/2016/12/35), Dr BC Roy Postgraduate Institute Institutional Ethics Committee (BCH/ME/PR/PS/54).

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