# STUDY PROTOCOL



# The Brain Imaging for Global Health (BRIGHT) Project:

# Longitudinal cohort study protocol [version 1; peer review: 1

# approved, 1 approved with reservations]

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# Abstract

There is a scarcity of prospective longitudinal research targeted at early postnatal life which maps developmental pathways of earlystage processing and brain specialisation in the context of early adversity. Follow up from infancy into the one-five year age range is key, as it constitutes a critical gap between infant and early childhood studies. Availability of portable neuroimaging (functional near infrared spectroscopy (fNIRS) and electroencephalography (EEG)) has enabled access to rural settings increasing the diversity of our sampling and broadening developmental research to include previously underrepresented ethnic-racial and geographical groups in low- and middle- income countries (LMICs). The primary objective of the Brain Imaging for Global Health (BRIGHT) project was to establish brain function - using longitudinal data from mother - for-age reference curves infant dyads living in the UK and rural Gambia and investigate the association between context-associated moderators and developmental trajectories across the first two years of life in The Gambia. In total, 265 participating families were seen during

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pregnancy, at 7-14 days, 1-, 5-, 8-, 12-, 18- and 24-months postpartum. An additional visit is now underway at 3-5 years to assess pre-school outcomes. The majority of our Gambian cohort live in poverty, but while resource-poor in many factors they commonly experience a rich and beneficial family and caregiving context with multigenerational care and a close-knit supportive community. Understanding the impact of different factors at play in such an environment (i.e., detrimental undernutrition versus beneficial multigenerational family support) will (i) improve the representativeness of models of general cognitive developmental pathways from birth, (ii) identify causal pathways of altered trajectories associated with early adversity at both individual and group level, and (iii) identify the context-associated moderators (i.e. social context) that protect development despite the presence of poverty-associated challenges. This will in turn contribute to the development of targeted interventions.

# **Keywords**

Gambia, UK, infancy, development, undernutrition, longitudinal, neuroimaging, global health, fNIRS, EEG

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### Introduction

# Background

The first 1000 days of life, which describes the developmental period between conception and two years of age, is characterised by prodigious physiological, psychological and physical change. As such, this period represents a critical window for brain development, during which plasticity to environmental factors is greatest. According to UNICEF, 19.5% of the world's children live in poverty, the majority of whom reside within sub-Saharan Africa (51.7%) and South Asia (35.7%). Infants and children growing up in poverty may be exposed to a range of biological and/or psychosocial risk factors both preand postnatally. Such risk factors include lower parental income and educational level, parental mental health issues, reduced access to recreational and educational activities (particularly in rural communities), undernutrition, food insecurity, environmental hazards and poor sanitation (Giovanelli et al., 2016; Jensen et al., 2017; Smith et al., 2015; Worku et al., 2018). While resource poor in many factors, there can also be many beneficial context-associated moderators, for example some communities who grow up in poverty also experience a rich and beneficial family and caregiving context with multigenerational social support and a close-knit supportive community. The impact of environment on neurocognitive development is therefore dynamic and multi-faceted, affecting biological, social, and behavioural developmental processes. For example, undernourished infants may seek, and consequently receive, less stimulation from caregivers. This lack of social stimulation is, in turn, linked to changes in brain function which are likely to precede changes in behaviour (East et al., 2017). Despite these multifaceted links between different factors, evidence examining the impact of poverty-related risk and developmental outcomes oftentimes focusses on only narrow subsets of relevant factors, which leaves open questions regarding the interplay across domains. As noted by Nobel Laureate Esther Duflo in 2019, 'Our goal is to make sure that the fight against poverty is based on scientific evidence. It starts from the idea that often the poor are reduced to caricatures and often even people who try to help them do not actually understand what are the deep roots of the problems' (Cho, 2019). Here, we describe the rationale behind and cohort characteristics of the Brain Imaging for Global Health (BRIGHT) project, which works with family cohorts in the UK and The Gambia. We first present relevant literature that informed the design of the BRIGHT project, before providing details on the study protocol and characteristics of the two cohorts.

Childhood poverty has been associated with lower performance on language, memory and cognitive control tasks (Farah *et al.*, 2006). At a global level, this is reflected by one third of pre-school-aged children in low-and-middle-income countries (LMICs) failing to reach age-appropriate milestones in cognitive and/or socio-emotional development (McCoy *et al.*, 2016). Within the first years of life, one major povertyassociated risk factor that poses a considerable risk to early child development is stunting (low length/height for age against an international reference), which occurs as the result of chronic growth failure and affects one in five children under five years of age globally (Development Initiatives, 2018). Furthermore, the interplay between the impact of undernutrition and compensatory factors (i.e., positive parenting practices), thought to scaffold early child development, is complex. A recent study in rural Cambodia looking at the joint role of parenting and nutritional status - in relation to inequities in family wealth - found that, while more stimulating and supportive parenting practices were associated with improved developmental outcomes in three to five year olds, this was strongest for non-stunted children (Berkes et al., 2019). We therefore urgently need to further our understanding of brain and cognitive development during early childhood in the context of poverty-associated risk factors. This is especially relevant as compromised development of a core set of age-appropriate skills in childhood has a significant impact on subsequent academic achievement, mental health and economic status - and consequently the potential to lead full and productive lives and support future generations (Alderman et al., 2014; Hackman & Farah, 2009; Martorell et al., 2010; Victora et al., 2008). The United Nations Sustainable Development Goals have consequently identified the reduction of poor cognitive development during childhood in LMICs as a key priority for global health research and interventions (UN, 2015). All LMICs fall within what are also known as Majority World countries, where 85% of the world's population live (Alam, 2008). However, over the past 15-30 years, only 3 - 17% of published child development journal articles (Moriguchi, 2022; Nielsen et al., 2017) are from Majority World countries and only 5% of child development interventions (Draper et al., 2023), meaning that the world's child population are under-represented in our theoretically driven understanding of development (Draper et al., 2022). In the following section we review some of the existing literature on links between poverty and brain development as well as describe how a small number of studies are beginning to address the under-representation of child development research in LMIC (Majority World) contexts.

Over the last decade several large-scale studies from the United States have shown links between poverty and brain development in childhood (Barch et al., 2016; Hair et al., 2015; Luby et al., 2013; Noble et al., 2015); for example, children of parents with high school education had roughly 3% less cortical volume than those with university level education, and those with parental incomes below \$25,000 had 6% less than those making over \$150,000 (Noble et al., 2015). These findings have led to models relating specific components of socioeconomic status (SES), stress and brain structural and functional development (Farah, 2017; Hackman & Farah, 2009; Noble et al., 2012). However, the majority of these studies rely heavily on correlational analyses in later childhood rather than studying development at an early age to fully understand the mechanistic processes driving these differences (for a review of studies that have looked at general differences in brain volume and poverty in infancy see Hurt & Betancourt, 2016). While the development of new methodology has increased our understanding of brain and cognitive development in infants and young children over the last decade, this research has been largely restricted to financially mobile participants within high-income countries (Henrich *et al.*, 2010). Studies that examine the impact of extreme poverty (defined by the World Bank as living on less than \$1.9 per household member per day) on brain development are extremely scarce. Furthermore, it remains unclear whether poverty-associated risk factors influence brain development more severely during sensitive periods of development (*i.e.* fetal life/early infancy/early childhood), or whether their impact builds over time depending on the chronicity, pervasiveness, severity and/ number of incidences (Berens *et al.*, 2017; Jensen *et al.*, 2017). Furthering our understanding of the developmental impact of early environmental adversity in both the short- and long-term is thus of high priority, particularly during the understudied period of the first 1000 days of life.

A further consideration is that the majority of child development research conducted in LMICs has been limited to the use of behavioural assessments of cognitive development to measure the effect of exposure to early adversity. Such measures are often undertaken later in childhood rather than at the time that vulnerability to exposure is most critical *i.e.* during prenatal and early postnatal life (Sabanathan et al., 2015). Furthermore, it is important to note that developmental trajectories of perceptual, motor and language domains have different timescales and cascading effects on one another. Therefore, they may have different key periods of sensitivity to insults (for an example of a key sensitive period for visual cortex see Hensch, 2005). Thus, it is imperative that neurocognitive development is studied from as close to birth as possible ideally with a longitudinal framework to track age-related changes - taking contemporaneous measurements of brain function and behaviour in parallel with measurements of exposure to environmental challenges. Furthermore, such research could offer new pathways for the provision of widely applicable, objective paradigms and methods that can assess early brain development in hard-to-reach populations (Isaacs, 2013). This would be synergistic with current large scale global health initiatives to optimise behavioural measures of early cognitive development (Murray-Kolb et al., 2014; Richter et al., 2019) such as the Global Scales of Early Development (GSED) (Cavallera et al., 2023). Furthermore the introduction of objective brain imaging paradigms to global health research could address some of the current challenges associated with behavioural measures of child development (Isaacs et al., 2008; Perkins et al., 2017). The optimisation of tools for measuring neurocognitive development will in turn support the development of early intervention strategies from the first days and months of life offering the potential for large lifetime cost savings (i.e. "1001 Critical Days" cross party manifesto, UK; Sure Start (Cattan et al., 2021)).

Over the last five to ten years, several research collaborations have been established to bring new neurocognitive technology (*i.e.* eye-tracking and neuroimaging tools) to the field of infant and early child brain and cognitive development research in LMICs: these include within The Gambia (Begus *et al.*, 2016; Katus *et al.*, 2019; Katus *et al.*, 2020; Katus *et al.*, 2023; Katus *et al.*, 2022b; Lloyd-Fox *et al.*, 2014; Lloyd-Fox *et al.*, 2017; Lloyd-Fox *et al.*, 2019); Cote D'Ivoire

(Jasinska & Guei, 2018); Guinea Bissau (Roberts et al., 2017); Malawi (Forssman et al., 2017; Pyykkö et al., 2019; Pyykkö et al., 2020); Bangladesh (Jensen et al., 2019; Perdue et al., 2019; Turesky et al., 2019; Xie et al., 2018); India (Wijeakumar et al., 2019); South Africa (Wedderburn et al., 2020) and Brazil (Alarcão et al., 2021). These global infant and child brain development studies cover age ranges from 0 - 65 months of age, reporting on areas as wide ranging as social information processing (Lloyd-Fox et al., 2017; Perdue et al., 2019; Xie et al., 2019), early brain anatomical and connectivity development (Collins-Jones et al., 2021; Fishell et al., 2020; Turesky et al., 2020; Turesky et al., 2019), the development of brain networks associated with visual working memory (Wijeakumar et al., 2019) and the development of attentional/neural markers of habituation and novelty detection (Katus et al., 2020; Katus et al., 2023; Katus et al., 2022b; Lloyd-Fox et al., 2019). Finally, a recent step change in research has been to begin to use measures of brain and cognitive development to understand the impact of interventions within global health studies; as evidenced by the recent work by Alarcão and colleagues in Brazil to measure the efficacy of a home-visiting program for adolescent mothers for enhancing early infant brain development and behaviour (Alarcão et al., 2021).

In this paper we describe the BRIGHT project, a follow-on from a pilot study, which ran from 2012-2014, in which we demonstrated the feasibility of combined neuroimaging, behavioural assessments and growth measures in longitudinal and crosssectional studies from birth to 24 months of age in rural Gambia (Begus et al., 2016; Lloyd-Fox et al., 2017; Lloyd-Fox et al., 2014; Papademetriou et al., 2013). Importantly we showed that fNIRS, can be easily implemented in rural contexts such as in The Gambia and used from the first weeks of life to provide quantitative and objective markers of neurocognitive function. We identified testing paradigms that elicit reliable brain responses that can be used to chart development as a function of age, and which aligned with findings in age-matched groups of infants from studies conducted in the UK. As part of our pilot study, we also successfully performed a quality control assessment of the adaptation and administration of a behavioural assessment, the Mullen Scales of Early Learning (MSEL), for use in rural Gambia (Milosavljevic et al., 2019). The BRIGHT project has been established to extend this pilot phase to a larger longitudinal observation cohort study of infant and early child development from birth to two years of age, with an additional follow-up at pre-school age (3-5 years).

### The BRIGHT project design overview

The BRIGHT project (Phase I), which ran from 2015 - 2020, established two prospective cohorts of families in the UK and The Gambia using a longitudinal multi-methods approach. Families were recruited during pregnancy and, following delivery, longitudinal measures of infant brain and cognitive development were conducted from 0–24 months of age across 10 data collection phases: antenatal – recruitment and 32–36 weeks' gestation, and postnatal – 1–3, 7–14 days, 1, 5, 8, 12, 18 and 24 months of age. Additional data on diet and health were collected continuously across this time period. The project

implemented brain imaging measures (fNIRS and electroencephalography [EEG]), neurocognitive behavioural measures (utilising eye-tracking methods), population-specific cognitive developmental measures (MSEL and the Communicative Development Inventory [CDI]), family-caregiving assessments (Family Care Indicators [FCI]), caregiver-infant interaction videos and questionnaires) and home environment measures (Language Environment Analysis [LENA]) alongside regular collection of biological, socioeconomic, parental health and nutritional measures at both sites. This data collection framework was implemented to allow the modelling of longitudinal changes in brain function, cognitive development, and growth within the rural Gambian population. Further, the collection of parallel behavioural and environmental data was designed to enable the identification of critical developmental moderators, mediators, and markers of risk and resilience. The purpose of the BRIGHT project is to firstly establish longitudinal trajectories across populations, and secondly to provide a framework for in-depth investigations of interindividual differences within the Gambian cohort. Given that neuroimaging data provided the backbone of this project, it was essential that a UK cohort was also established to measure the longitudinal developmental trajectories of the different fNIRS and EEG paradigms across different populations as several of these had not been studied across this longitudinal time span before in any population internationally. This was chosen to broadly match the context of previously acquired developmental neuroimaging data, given that to date the vast majority of research of this kind has been undertaken in high income countries (HICs).

The target cohort sizes (The Gambia n=200, UK n=60,) were based on previous infant fNIRS and EEG studies conducted in the UK, which indicated that sample sizes from 20 (moderate effect size) to 42 (small effect size) were sufficient to determine regions of significant cortical brain activation in response to stimuli. The Gambian cohort was designed to be larger to allow within-cohort sub-group comparisons and individual differences analyses; for example, grouping by growth trajectories (mild, medium and severe markers of undernutrition) on the assumption that approximately 25–30% of the cohort would be stunted (z-score of length-for-age < 2 standard deviations below the WHO reference) by two years of age (Nabwera *et al.*, 2017).

# BRIGHT Kids (BRIGHT project phase II)

Previous research highlights a marked impact of exposure to early adversity and neurobehavioral outcomes at preschool age. With this in mind, in 2023 we conducted a follow-up assessment at preschool age in the Gambian cohort of the BRIGHT project at 3–5 years. This cross-sectional follow up will allow us to examine additional questions, regarding the long-term stability of our infant neural markers to predict long-term outcomes.

# Objectives

The primary objectives of the BRIGHT project are to:

(1) develop brain and neurocognitive function-for-age curves from birth to 24 months of age using prospective

longitudinal datasets from the UK and The Gambia. These reference curves will be used to enable age-adjusted group comparisons of differences in average trajectories, group-wise differences in variability, and for characterizing the range of individual developmental trajectories within each cohort.

- (2) establish the association between context-associated moderators, including poverty-associated risk factors (*i.e.*, undernutrition and consequent growth faltering), and developmental trajectories across the first two years of life in The Gambia.
- (3) establish the association between context-associated moderators and developmental trajectories across the first two years of life in The Gambia and pre-school outcomes at three to five years.

Secondary aims of interest are to:

- 1. Assess whether infants with similar trajectories of growth have the potential to reach the same developmental milestones within the first 24 months of life.
- 2. Establish whether neuroimaging markers of brain function are more robust indicators of development within individual infants across age, as compared to behavioural measures.
- 3. Assess the capability of fNIRS, EEG and eyetracking methods to deliver specific and early biomarkers of altered developmental pathways.

Here we describe the formation of a common BRIGHT study protocol across the two sites (The Gambia and the UK) and, where appropriate, site-specific additional measures, particularly those focused on family context, nutrition, diet, and biological samples are presented for The Gambia. We briefly outline how we recruited participants at each site; selected and implemented experimental neurocognitive and behavioural measures at each site; how we standardised lab practices across sites to ensure comparability; and how we developed analytical pipelines for the different datasets. We also describe the demographic and socioeconomic distribution of our cohorts.

#### Methods

### The Gambian site and population

The Gambia is situated on the West coast of Africa, bordering Senegal. The majority (60%) of the roughly 2.4 million inhabitants of The Gambia live in the coastal regions surrounding the capital, Banjul, while the remainder of the population live rurally, often supporting themselves through subsistence farming (Hennig *et al.*, 2017) living in extended, multi-generational households (Brotherton *et al.*, 2021; Kea, 2013; Sear & Mace, 2009). The Gambia is one of the lowest ranking countries with regard to gross national income, years of schooling, and life expectancy, with over half of adults never having received formal education (Hennig *et al.*, 2017). School attendance has risen rapidly over the last decades thanks to the introduction of free universal education,

and 97% of children now attend to primary level relative to 66.7% in the early 1970s (The Gambia Government National Education Statistics, 2018; CEICdata.com). Preschool education has also become increasingly available across the timeframe of the BRIGHT project (Blimpo et al., 2019). Childcare is viewed as a shared responsibility among family members, with grandmothers and older sisters having the biggest role in supporting parents (Brotherton et al., 2021; Sear & Mace, 2009). Islam is the predominant religion and raising children in accordance with religious and community values is of high importance (Sosseh et al., 2023). Marriages are commonly polygamous with over half of married women living with one or two co-wives (Hennig et al., 2017), though gendered hierarchies and intra-household relations are dynamic and subject to change (Kea, 2013; Kea, 2020). Furthermore, over the past decades infant and child mortality has decreased, birth spacing has increased, and overall family size has reduced (Nabwera et al., 2017).

The Gambian arm of the BRIGHT project was hosted at a rural site of the Medical Research Council The Gambia Unit at the London School of Hygiene and Tropical Medicine (MRCG@ LSHTM; www.mrc.gm). The UK Medical Research Council (MRC) has a long-standing research partnership with The Gambia, established in the late 1940's. Currently, research conducted within MRCG@LSHTM is focused on three broad themes centred around major public health priorities, specifically Vaccines and Immunity, Disease Control and Elimination, and Nutrition and Planetary Health; the latter of which the BRIGHT project is situated within.

The BRIGHT project was undertaken at the Keneba Field Station of MRCG@LSHTM, situated in the rural West Kiang region, 145 km inland from the capital. Seasonality has an impact on nutrient availability for the population living here as weather patterns alternate between four months of heavy rainfall (July-October) and eight months of extreme dryness directly affecting the availability of key nutrients (Moore et al., 1997). In 2015 at the onset of the project, the Keneba Field Station was relatively isolated, accessed via unmade roads and required to independently maintain all facilities necessary for research and clinical care (e.g., generator powered electricity, bore hole water supply, satellite communication). However, over the course of the study, the country and local region have been witnesses to several changes. At the local level several infrastructure improvements have been made, including the road being tarmacked, and therefore allowing greater access to urban resources, and the field station and local community now have nationally sourced electrical power. BRIGHT project participants were drawn from Keneba and surrounding villages within a 20km radius of the field station.

All women of reproductive age (18–45 years) who were reported to be pregnant within the West Kiang Demographic Surveillance System between June 2016 and March 2018, spoke Mandinka as their primary language, and were expected to reside in West Kiang for the duration of the project were invited to participate (see Figure 1 for recruitment pathway in The Gambia). Further eligibility criteria for study participation pertaining to the pregnant women included: carrying a singleton pregnancy, < 36 weeks' gestation on presentation to the first antenatal study visit and being medically fit to participate, as determined by the study midwife. The project was designed to recruit participants so that deliveries were spread evenly throughout the recruitment period, aiming for around 10-15 deliveries per month. This was to ensure that workload was achievable and as consistent as possible, ensuring timely scheduling of follow-up visits. For this reason, an additional exclusion criterion of 'gestational age incompatible with study requirements' was introduced. From an ultrasound scan at the first antenatal study visit, gestational age was measured, and expected delivery date calculated. If a participant was due to deliver in a month that was already at full capacity, they were excluded at this point. Postnatally, mother-infant dyads were excluded from the project if the infant was diagnosed with a developmental disability e.g., Down's Syndrome or cerebral palsy. Participants were free to withdraw from participation at any point in the study.

# The UK site and population

In the UK, participants were recruited from the city of Cambridge and surrounding villages. Demographically, the population in Cambridgeshire is representative of that across the UK with regard to ethnicity, employment rates and family structure (Cambridge County Council, 2011). The area however differs from the rest of the UK with regard to levels of education within the population, with twice as many inhabitants holding a higher education degree (Cambridge County Council, 2011). The research involved in the UK arm of the BRIGHT project was conducted at dedicated facilities either within the Evelyne Perinatal Imaging Unit at the Rosie Hospital, Cambridge University Hospitals NHS Foundation Trust or within the Department of Psychology, University of Cambridge. Once per week during the recruitment phase, families who attended an antenatal clinic at the Rosie Maternity Unit at Cambridge University Hospitals between June 2016 and January 2017, with a healthy singleton pregnancy less than 36 weeks gestational age, were approached and given information about the project (see Figure 2 for UK recruitment pathway).

#### Ethical considerations

Protocols were approved by the relevant committee at each site. In The Gambia, ethical approval was given by the joint Gambia Government - MRC Ethics Committee (SCC 1351) and the Scientific Coordinating Committee at the MRC Unit The Gambia. Additional approval was granted for the BRIGHT Kids follow up (Project reference 22737). Informed consent was obtained in writing, or via thumbprint if individuals were unable to write, from all parents/carers prior to participation. In the UK, the study was approved by the National Research Ethics Service East of England Committee, NHS Health Research Authority (REC reference 13/EE/0200), and informed written consent was obtained from parents of infants to participate. The project is guided by a consistent set of principles which ensured that the infants' and child's wellbeing is always prioritised. Infants/children are always with their caregiver. The protocols were designed to be engaging and interesting to the infants and children, and the setup comfortable. Caregivers were made aware that the study can be interrupted, rescheduled,

Intenatal Screening (research lab visit)         Pregnancy Dating         Eligibility confirmed         Questionnaires: Pregnancy history, medical history         24-36 weeks' gestation (research lab visit)         Questionnaires: EPDS, PRAS, PSS         Biological Samples: Maternal: 5mL fasted sample of venous blood and a 10mL spot urine sample         Physical and behavioural assessments: maternal anthropometry         Sinth – 5 days (home visit)         Delivery details         Physical and behavioural assessments: MIPH, anthropometry         7-14 days after birth (home visit)         Infant physical and behavioural assessments: Anthropometric measures, NBAS.         Questionnaires: Demographics, SES.         I month (research lab visit)         Infant neurocognitive measures: fNIRS (SOC, HaND, FC), EEG/ERP         Infant physical and behavioural assessments: PCI video recording, anthropometry.         Questionnaires: EPDS, PANAS, PSS, Sleep Diary         Biological sample: Infant: 0.5ml Capillary blood sample         Simonths (research lab visit)         Infant neurocognitive measures: fNIRS (SOC, HaND, FC), EEG/ERP, eye tracking tasks.         Infant physical and behavioural assessments: PCI video recording, MSEL, anthropometry.         Questionnaires: Infant: 0.5ml Capillary blood sample         Simonths (research lab visit)         Infant physical and behavioura	
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<ul> <li>34-36 weeks' gestation (research lab visit)</li> <li>Questionnaires: EPDS, PRAS, PSS</li> <li>Biological Samples: Maternal: 5mL fasted sample of venous blood and a 10mL spot urine sample</li> <li>Physical and behavioural assessments: maternal anthropometry</li> <li>36rth - 5 days (home visit)</li> <li>Delivery details</li> <li>Physical and behavioural assessments: MIPH, anthropometry</li> <li>7-14 days after birth (home visit)</li> <li>Infant physical and behavioural assessments: Anthropometric measures, NBAS.</li> <li>Questionnaires: Demographics, SES.</li> <li>I month (research lab visit)</li> <li>Infant neurocognitive measures: fNIRS (SOC, HaND, FC), EEG/ERP</li> <li>Infant physical and behavioural assessments: PCI video recording, anthropometry.</li> <li>Questionnaires: EPDS, PANAS, PSS, Sleep Diary</li> <li>Biological sample: Infant: 0.5ml Capillary blood sample</li> <li>imonths (research lab visit)</li> <li>Infant neurocognitive measures: fNIRS (SOC, HaND, FC), EEG/ERP, eye tracking tasks.</li> <li>Infant neurocognitive measures: fNIRS (SOC, HaND, FC), EEG/ERP, eye tracking tasks.</li> <li>Infant neurocognitive measures: fNIRS (SOC, HaND, FC), EEG/ERP, eye tracking tasks.</li> <li>Infant neurocognitive measures: fNIRS (SOC, HaND, FC), EEG/ERP, eye tracking tasks.</li> <li>Infant neurocognitive measures: fNIRS (SOC, HaND, FC), EEG/ERP, eye tracking tasks.</li> <li>Infant physical and behavioural assessments: PCI video recording, MSEL, anthropometry.</li> <li>Questionnaires: EPDS, PANAS, PSS, Milestones, Sleep Diary,</li> <li>Biological sample: Infant: 3mL venous blood, spot urine sample; Maternal: 10mL breast milk (5mL each breast)</li> </ul>	
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B months (research lab visit)	
Infant neurocognitive measures: fNIRS (SOC, HaND, FC, WM), eye tracking tasks.	
<ul> <li>Infant physical and behavioural assessments: PCI video recording, MSEL, deferred imitation task, anthropometry</li> </ul>	/
Questionnaires: Demographics, SES, Milestones, Sleep Diary	
Biological sample: Infant: 0.5ml Capillary blood sample	
2 months (research lab visit and home visit*)	
Infant neurocognitive measures: fNIRS (SOC, HaND, FC, WM), eye tracking tasks.	
Infant physical and behavioural assessments: PCI video recording, MSEL, deferred imitation task, anthropometer	try,
LENA* Questionnaires: EPDS, PANAS, PSS, Milestones, Sleep Diary, CDI, FCI, family caregiving interview*.	
8 months (research lab visit and home visit*). Infant neurocognitive measures: fNIRS (SOC, HaND, FC, WM), EEG/ERP, eye tracking tasks.	
<ul> <li>Infant heurocognitive measures: Niks (SOC, Hand), PC, Will, ECG/ERP, eye tracking tasks.</li> <li>Infant physical and behavioural assessments: PCI video recording, MSEL, anthropometry, LENA*.</li> </ul>	
<u>Questionnaires</u> : Demographics, SES, Milestones, Sleep Diary, CDI, FCI, family caregiving interview <sup>*</sup> .	
Biological sample: Infant: Capillary blood sample	
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24 months (research lab visit and home visit*)	
Infant neurocognitive measures: fNIRS (SOC, HaND, FC, WM), eye tracking tasks.	
Infant physical and behavioural assessments: PCI video recording, MSEL, anthropometry, LENA*	
Questionnaires: Milestones, Sleep Diary, CDI, FCI, family caregiving interview* Biological samples: Infant: 3mL venous blood sample, spot urine sample	
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Digoing assessments throughout postnatal phase of study <u>     Food diary/questionnaires</u>	

**Figure 1. Summary of study visits: The Gambia.** EPDS, Edinburgh Postnatal Depression Scale; PRAS, Pregnancy-related Anxiety Scale; PSS, Perceived Stress Scale; PANAS, Positive and Negative Affect Scale; NBAS, Neonatal Behavioural Assessment Scale; SES, Socioeconomic Status; MIPH, Maternal and Infant Physical Health; fNIRS, functional near infrared spectroscopy; EEG/ERP, electroencephalography/event related potentials; SOC, Social versus Non-Social Response; HaND, Habituation and Novelty Detection; FC, Functional Connectivity Networks; WM, Working Memory; DI, Deferred Imitation task; PCI, Parent-Child Interaction; MSEL, Mullen Scales of Early Learning; LENA, Language Environment Analysis; FCI, Family Care Indicators; CDI, Communicative Development Inventory; \* indicates assessments undertaken in the family's home at the later time points.

#### Recruitment at 36 weeks' gestation

- Eligibility women identified via research midwife at the hospital
- · Detailed explanation of the study: (verbal and written)
- Informed consent (or information taken home for consideration)

36-40 weeks' gestation (home visit)

- Informed consent (if not already given)
- <u>Questionnaires</u>: EPDS, PRAS, PSS, Family details, Demographics, SES,
- <u>Physical and behavioural assessments</u>: maternal anthropometry

#### Birth

- Update on birth midwife monitors birth records at hospital / parents contact research team / researchers reach out via
   email to check for an update
- Birth details telephone/email contact to update D.O.B. and arrange first home visit

7-14 days after birth (home visit)

- Infant physical and behavioural assessments: anthropometry, NBAS
- Questionnaires: MIPH, Healthy Pregnancy, Demographics, SES

1 month (research lab visit)

- Infant neurocognitive measures: fNIRS (SOC, HaND, FC), EEG/ERP
- Infant physical and behavioural assessments: PCI video recording, anthropometry.
- Questionnaires: EPDS, PANAS, PSS, Sleep Diary, Infant Feeding questionnaire issued

5 months (research lab visit)

- Infant neurocognitive measures: fNIRS (SOC, HaND, FC), EEG/ERP, eye tracking tasks.
- Infant physical and behavioural assessments: PCI video recording, MSEL, anthropometry.
- Questionnaires: EPDS, PANAS, PSS, Milestones, Sleep Diary,

8 months (research lab visit)

- Infant neurocognitive measures: fNIRS (SOC, HaND, FC, WM, DI), eye tracking tasks.
- Infant physical and behavioural assessments: PCI video recording, MSEL, anthropometry
- Questionnaires: Demographics, SES, Milestones, Sleep Diary

12 months (research lab visit and home visit\*)

- Infant neurocognitive measures: fNIRS (SOC, HaND, FC, WM, DI), eye tracking tasks.
- Infant physical and behavioural assessments: PCI video recording, MSEL, anthropometry, LENA\*
- Questionnaires: EPDS, PANAS, PSS, Milestones, Sleep Diary, CDI, FCI, family caregiving interview\*.

18 months (research lab visit and home visit\*).

- Infant neurocognitive measures: fNIRS (SOC, HaND, FC, WM), EEG/ERP, eye tracking tasks.
- Infant physical and behavioural assessments: PCI video recording, MSEL, anthropometry, LENA\*.
- Questionnaires: Demographics, SES, Milestones, Sleep Diary, CDI, FCI, family caregiving interview\*.

24 months (research lab visit and home visit\*)

- Infant neurocognitive measures: fNIRS (SOC, HaND, FC, WM), eye tracking tasks.
- Infant physical and behavioural assessments: PCI video recording, MSEL, anthropometry, LENA\*
- <u>Questionnaires:</u> Milestones, Sleep Diary, CDI, FCI, family caregiving interview\*

Ongoing assessments throughout postnatal phase of study

Food diary/questionnaires

**Figure 2. Summary of study visits: The UK.** Abbreviations: EPDS, Edinburgh Postnatal Depression Scale; PRAS, Pregnancyrelated Anxiety Scale; PSS, Perceived Stress Scale; PANAS, Positive and Negative Affect Scale; NBAS, Neonatal Behavioural Assessment Scale; SES, Socioeconomic Status; MIPH, Maternal and Infant Physical Health; fNIRS, functional near infrared spectroscopy; EEG/ERP, electroencephalography/event related potentials; SOC, Social versus Non-Social Response; HaND, Habituation and Novelty Detection; FC, Functional Connectivity Networks; WM, Working Memory; DI, Deferred Imitation task; PCI, Parent-Child Interaction; MSEL, Mullen Scales of Early Learning; LENA, Language Environment Analysis; FCI, Family Care Indicators; CDI, Communicative Development Inventory; \* indicates assessments undertaken in the family's home at the later time points. or stopped at any time if the infant/child became fussy or tired, or, if the caregiver expressed a wish to end the study. Each research team includes researchers fluent in Mandinka and/or English, as relevant. Data protection and confidentiality shape our approach to data sharing within the BRIGHT research team and externally, and is detailed in the *Standardisation of protocol across sites* section.

# Study protocol

The full study protocol is shown in Table 1 and outlined below. The draft protocol was developed through the integration of expertise from our international multi-disciplinary (psychology, neuroscience, medical physics and bioengineering, maternal and infant health and nutrition, global health) research leadership team. Following this, all measures were

**Table 1. Summary of BRIGHT protocol.** Note on abbreviations: \* The UK only; • The Gambia only; ^ Recorded every two weeks from 2 weeks to 24 months of age; L,W, H, HC,MUAC, KHL (L – length, W – weight, H – height, HC – head circumference, MUAC – mid to upper arm circumference, KHL – knee to heel length).

Study Measure	Time point:	Antenatal	Birth	7–14 d	1 mo	5 mo	8 mo	12 mo	18 mo	24 mo
	Neu	roimaging m	neasures							
fNIRS: Social/1	Non-social				x	x	x	x	х	х
fNIRS: Habituation and	Novelty Detection				х	x	x	х	х	х
fNIRS: Functiona	l connectivity				x	x	х	x	x	х
fNIRS: Workin	g memory						x	x	x	x
fNIRS*/Behavioural: [	Deferred imitation						х	x		
EEG: Auditory	y Oddball				х	x			x	
	Behavioura	l/Neurocogr	nitive mea	sures						
Neonatal Behavioural Ass	essment Scale (NBAS)			х						
Eye-tracking: Cog	nitive control					x	x	x	х	х
Eye-tracking: H	labituation					x	x	х		
Eye-tracking: G	Eye-tracking: Gap/Overlap					x	x	x	х	х
Eye-tracking: Non-sc	ocial contingency					x	x	х		
Eye-tracking: Fa	ace popout					x	x	x	х	х
Eye-tracking: Dyr	namic scenes					x	x	x	х	х
Eye-tracking: Word-p	picture-matching								х	х
Mullen Scales of Early	y Learning (MSEL)					x	x	х	х	х
Parent-Child I	nteraction				х	x	x	x	х	х
LENA language asse	essment in home							х	х	х
LENA in	PCI				<b>X</b> △	<b>X</b> ^	<b>X</b> ^	<b>X</b> ^	х	х
Tablet-based Cognit	ive Assessment*								x *	x *
	Questionnaires/Interviews – Infant/Child									
Communication Develop	ment Inventory (CDI)							х	х	х
Adapted Oxford	l Sleep Diary				x	x	x	x	x	x
Food Frequency	Q (FFQ) △ ^		x	x	х	х	x	x	x	x
Infant feeding onl	ine Q (IFQ) * ^				x	x				
Food diary (Inta	ake24UK) *						x	х	х	х
Early Childhood Deve										

Study Measure Time point:	Antenatal	Birth	7-14 d	1 mo	5 mo	8 mo	12 mo	18 mo	24 mo
Question	naires/Inter	views – Fa	mily						
Edinburgh Postnatal Depression Scale (EDPS) – Maternal	x			x	x		x		x
Edinburgh Postnatal Depression Scale (EDPS) – Paternal *	x			x	x		x		x
Pregnancy Related Anxiety form (PRAS) – Maternal	x								
Pregnancy Related Anxiety form (PRAS) – Paternal*	x								
Pregnancy Specific Anxiety (PSA) – Maternal	x								
Pregnancy Specific Anxiety (PSA) – Paternal*	x								
Positive Negative Affect Schedule (PANAS) – Maternal				x	x		х		x
Positive Negative Affect Schedule (PANAS) – Paternal*				x	х		х		х
Perceived Stress Scale (PSS) – Maternal	x			x			x		x
Perceived Stress Scale (PSS) – Paternal*	x			х			х		x
Socioeconomic Status (SES)	x *		<b>X</b> $ riangle$			х *		х	
Demographic and Family Information	x *		<b>X</b> ^			x *		x	x *
Family details (from DSS – Gambia or antenatal call UK)	x					х *			x *
Family Caregiving Questionnaire (FCQ)							х	x	x
Family Care Indicators (FCI) 🗅							х	x	x
Clinical r	neasures / M	edical det	ails						
Healthy Pregnancy Questionnaire *			x						
Antenatal Medical form *	x								
Delivery Information and Baby check ${\scriptscriptstyle  riangle}$		x							
Anthropometric measures – infant (L,W,HC,MUAC, KHL)		<b>x</b> (L,W,HC)	<b>x</b> (L,W,HC)	x	x	x	x	x	x
Anthropometric measures – mother (W, H)									
Maternal blood sample 🗠	x								
Maternal urine sample 🗠	x								
Maternal breast milk sample 🗠					х				
Infant blood sample 🗠				x	x	x	x	x	x
Infant urine sample 🗠					х		х		x

reviewed during several multi-site web-based meetings to identify the necessary adaptations and translations required to ensure each paradigm and assessment was appropriate for the population and culture of the cohort. Furthermore, some field testing and adaptation had already been undertaken in previous pilot phases within our research group (Lloyd-Fox *et al.*, 2014; Milosavljevic *et al.*, 2019).

During the protocol development phase, for paradigms that used images, videos, or audio that included people, actors representative of the ethnicity and language of the participants were used. For paradigms that included toys or objects (either real or in image form), appropriate representatives of the contextual environment of each cohort were identified. For The Gambia only, when appropriate, questionnaires were translated and administered in Mandinka (e.g., all mental health questionnaires, and the Mullen Scales of Early Learning [MSEL]). A full adaptation process with forward and backward translation by the authors and the BRIGHT Project team in Keneba was undertaken for all questionnaires and assessments (for an example see Milosavljevic et al., 2019). As Mandinka is not a written language and literacy rates among caregivers were low, the questionnaires were converted into interview versions and conducted by trained field assistants. Of note, where translations were undertaken for standardised measures that were not open-source and free to use (i.e., Mullen Scales of Early Learning [MSEL]) we purchased the equivalent number of copies of the original forms from the publisher that would be required to administer the measure at each age point. Neurocognitive measures were selected, where possible, on the basis of test re-test reliability and previous evidence indicating that they showed robust data quality (i.e., fNIRS (Blasi et al., 2014); Eye-tracking (Jones et al., 2019); EEG (Dzhelyova et al., 2019; Räikkönen et al., 2003)). Within each battery of measures, we selected a combination of well tested and robust paradigms, and, when necessary to allow us to target particular cognitive domains or informative metrics, paradigms with novel designs were developed by the BRIGHT research group (i.e., fNIRS tasks to assess working memory, delayed imitation, habituation, repetition suppression and novelty detection). Tasks were administered in a pre-determined order across the study visit where possible (i.e., anthropometrics were taken at the end of the session to optimise infants' attention and energy for experimental tasks), and also within a testing modality (i.e., in the fNIRS session infants viewed paradigms in a set order according to the stimulus presentation scripting framework). On occasion, when infants tired before completing the full session, families were invited to return on a separate day to complete the tasks, but where possible infants were encouraged to continue after a nap and/or feed within the same visit. We found that a second visit was required more often in The Gambian cohort than in the UK. Details of the session and completion of tasks were recorded in a Session Log Form at each visit.

#### Neuroimaging measures

#### Electroencephalography (EEG)

Electroencephalography (EEG) has a long-standing tradition in neurodevelopmental research. It provides a direct measure of infants' neural responses to stimuli without requiring them to overtly respond or to follow task instructions. Through the use of innovative, wireless EEG hardware, it is now possible to implement EEG tasks in remote rural contexts and in the absence of standardised lab settings (Katus et al., 2019). The EEG task implemented in the BRIGHT project assessed auditory habituation and novelty detection, at 1, 5 and 18 months of infant age (for a description of the full protocol see Katus et al., 2020). Due to the nature of the sounds (pure tones, bursts of white noise etc.), no adaptations had to be undertaken allowing for identical protocols at both project sites. Infants were presented with auditory stimuli for approximately 15 minutes while asleep (at 1 month) or awake (at 5 and 18 months): during the latter an experimenter quietly entertained the infant

with bubbles or silent toys to maintain calm attention during the task.

#### Functional near infrared spectroscopy (fNIRS)

Functional near infrared spectroscopy (fNIRS) is a relatively recent addition to the battery of neuroimaging measures available to neurodevelopmental research. fNIRS measures the haemodynamic response to the neural activation measured by EEG. It has become the technique of choice for many studies given its ease of use with infants and young children, improved spatial resolution (relative to EEG) and low cost (relative to MRI) (Gervain *et al.*, 2023; Lloyd-Fox *et al.*, 2010). In addition, fNIRS is relatively portable, opening a pathway for implementation in the remote and/or out-of-lab settings often associated with global health research contexts (Blasi *et al.*, 2019; Katus *et al.*, 2019).

The fNIRS paradigms implemented in the BRIGHT project assessed a range of cognitive functions and domains, namely social cognition (Lloyd-Fox et al., 2017), habituation and novelty detection (Lloyd-Fox et al., 2019), working memory, deferred imitation and functional connectivity. Paradigms were included, at age-appropriate time points, across the 1, 5, 8, 12, 18 and 24 months, as well as in BRIGHT Kids at 3-5 years of age. Paradigms contained auditory and/or visual stimuli and were presented while infants were asleep at 1 month of age and while infants were awake and alert at all other time points. Audio and visual stimuli were adapted with site-relevant content (see the Preliminary Results section and Katus et al., 2019). The full fNIRS battery lasted 24 min for the shortest sessions at 1 and 5 months to up to 35 min for the longest at 8 and 12 months (where we included a live behavioural Deferred Imitation task). When accounting for preparation time such as settling the infant, taking head measurements, capping, and photographing headgear, the total assessment time was approximately 45 minutes. This multi-domain battery was designed to interrogate whether global health risk factors impact on development to result in global/cross-domain differences in brain activity or localized/domain-specific differences or altered function.

#### Behavioural/neurocognitive assessments Neonatal Behavioural Assessment Scale (NBAS)

The NBAS is a structured clinical assessment of infant neurology and behaviour, which can be performed within the first few days of life. The NBAS is regarded as the most comprehensive examination of newborn behaviour available. It has been used across multiple cultures and in different LMICs (Zambia: Brazelton *et al.*, 1976, Chile: Ayala *et al.*, 2021, Mexico: Soler-Limón *et al.*, 2019, Kenya: Super & Harkness, 2020). The NBAS is a standard protocol which requires an initial training period culminating in assessment and certification as described in *The Neonatal Behavioral Assessment Scale* Manual (Brazelton & Nugent, 1995). Prior to the BRIGHT Project, we conducted a qualitative pilot study to assess the cultural acceptability and feasibility of using the NBAS within the rural low resource settings of families living in the West Kiang region of The Gambia. To this end, fifteen infants were assessed with the NBAS, and their families' feedback showed that the NBAS was acceptable to parents in this population (Bartram, 2018). In line with feedback from parents from other countries and populations, a few aspects of the assessment (specifically, shining a light over closed eyes while sleeping, covering eyes with cloth while awake and undressing the infant) were questioned or perceived negatively. While no items were altered or removed from the assessment, the more controversial items were introduced with special care during all study visits. In both the UK and The Gambia, the NBAS was performed between 7–14 days after birth during a pre-arranged home visit. Administration times for the NBAS ranged from 20 to 45 minutes, depending on the infants' state of alertness.

#### Eye-tracking neurocognitive battery

Eye-tracking is a non-invasive and well-tolerated measure in infant neurodevelopmental research (Jones *et al.*, 2019). The tasks in our eye-tracking battery were selected to provide broad coverage of several key domains of neurocognitive functioning, including working memory (Elsabbagh *et al.*, 2009; Elsabbagh *et al.*, 2013a; Johnson, 1995; Scerif *et al.*, 2005), visual attention (Kaldy *et al.*, 2011), habituation (Webb *et al.*, 2010), reversal learning (Wass, 2015), social *versus* non-social visual preference (Elsabbagh *et al.*, 2013b) and language learning (Fernald *et al.*, 2008). Table 1 displays the specific tasks used per age point. Many of the tasks, and all of the fixation stimuli that preceded a task, were gaze-contingent, that is they rely on the child's gaze to proceed through the battery. The duration of the battery therefore varied slightly between participants but averaged 20 minutes.

The majority of the tasks focused on the use of visual stimuli (accompanied by simple alerting sounds) with no language modification required for use in The Gambia. Therefore, the eye-tracking battery was run with an existing stimulus package used in other longitudinal large scale cohort studies (developed by researchers at the Centre for Brain and Cognitive Development, Birkbeck, University of London) to aid future comparative analyses. An additional task (word-picture matching task) was developed specifically for the BRIGHT project to assess language comprehension of participants at the ages of 18 and 24 months. The task measured processing efficiency (speed and accuracy) in terms of infants' ability to direct their gaze to one of two visual stimuli to match to a spoken target noun. This task was adapted from the behavioural Looking-while-Listening eye movement methodology used by Fernald and colleagues (2008), which had been recently adapted for a research project conducted in neighbouring Senegal (Weber et al., 2017). During the design phase of the paradigm, common items were photographed and audio recordings of sentences relating to the photographed items were made in the appropriate language at each site. A series of pilot studies were then completed in both the UK and The Gambia to identify population- and age- appropriate word-picture stimulus pairs.

#### Mullen Scales of Early Learning (MSEL)

The MSEL measures cognitive ability and motor development using five scales: Gross Motor, Visual Reception, Fine Motor, Expressive Language, and Receptive Language. In both the UK and The Gambia, the MSEL was performed at visits to the research lab conducted at 5, 8, 12, 18 and 24 months of age, as well as at 3-5 years of age in The Gambia. During each visit, the MSEL was conducted using the standardized protocol appropriate for the age of the participant, as detailed in the Mullen Scales of Early Learning Manual and the Item Administration Book (Mullen, 1995) and the MSEL training DVD. During a pilot phase to optimize and adapt the MSEL for use in The Gambia, n = 171 infants were tested across the age ranges described above (Milosavljevic et al., 2019). The MSEL is broadly similar to the Bayley Scales of Infant and Toddler Development, both of which have been adapted for use across multiple countries, but originate in the U.S.A. and are normed to this population. A further restriction of this measure is that it is not open source and must be purchased, both the toolkit, manual and the assessment forms, i.e. a fee per participant is required to be paid.

#### Parent-child interaction videos

We assessed parent-child interaction styles in both the UK and The Gambia at 1, 5, 8, 12, 18 and 24 months of age. The parent and child engaged in a video recorded, 10-minute free-play session, which consisted of five minutes of play without toys and five minutes with a standardised collection of toys provided by the research team. The parent and child were seated on a mat in front of a mirror, to ensure that both of their faces were visible on the recording. At the younger ages, the infants were placed on a baby mat facing the parent, but, as they became more mobile, they were allowed to move around the room. The parent was instructed to play with their child as they normally would at home. For the younger time points the parent-infant dyad were left alone in the room to encourage a more relaxed environment, however, at later time points the researcher remained in the room to be able to move the camera around as the child became more mobile. These videos can be coded to assess multiple aspects of parental and child behaviour and engagement. Parental interactive characteristics have been shown to associate with child neural and cognitive development across a range of cultural contexts (i.e. Bozicevic et al., 2016; Sethna et al., 2017).

#### Language Environment Analysis system (LENA)

The Language Environment Analysis system (LENA) provides automated counts of the linguistic environment. A digital language processor (DLP) is worn by the participant in the front pocket of a specially designed vest and able to record the audio environment within 1- to 3- metres. In BRIGHT, this was used to record the acoustic environment of the participants during a typical day in the home. Our standard protocol assessed the auditory home environment using LENA at the 12, 18 and 24 months of age. The recordings took place during two consecutive days, with seven hours of recording per day. During these recordings, parents were asked to complete a Family Caregiving Questionnaire or Interview. In the UK this was a logbook describing the main activities performed by the toddler, the locations in which the recording had taken place. who was around during the recording as well as technical details regarding the usage of the device. In The Gambia, this was conducted as an interview by a field assistant at the end of each recording day.

Via the LENA software, we extracted, 1) Adult word counts (AWC) defined as the number of adult words the key child hears - these adult words may be or may not be directed to the key child; 2) Key child vocalization counts (CVC), defined as words or prelinguistic babbling produced by the key child (crying and laughing sounds are not included in this category); 3) Conversational turn taking (CTT), identified as those instances when the key child and an adult speak one at a time in alternating turns. The reliability of these estimates has been shown in multiple languages; including English (Gilkerson & Richards, 2008; Xu et al., 2009), French (Canault et al., 2016), Dutch (Busch et al., 2018), Spanish (Weisleder & Fernald, 2013), Vietnamese (Ganek & Eriks-Brophy, 2018) and Korean (Pae et al., 2016). In line with previous research, field assistants, with Mandinka as their first language, transcribed a subset of recordings to determine reliability estimates for the Mandinka language. In addition to the home visits, within The Gambia a sub-sample (N = 40) of infants were followed more intensively from 1- 24 months of age with LENA recording included during the PCI sessions where the context is restricted to one parent and the target child.

#### Tablet-basedased cognitive assessment

The Babyscreen software application V1.83 (Hello Games, Guilford, UK) was used to measure key domains of neurodevelopment, including selective attention, working memory, and general learning ability. The task consists of 18 items and provides two performance variables: number of items accurately completed and speed of item completion (Twomey *et al.*, 2018). The task was administered on an Apple iPad (6<sup>th</sup> generation, 9.7" screen). Given the young age of the BRIGHT participants and varying exposure to touch screen technology, two free play tasks were administered at the start of the testing session to familiarise the participants with the tablet, these involved drawing on the screen and moving shapes around.

In the UK, the task was administered to participants at 18 and 24 months of age (Macrae *et al.*, 2022). In The Gambia, several challenges arose, and the task was removed from the protocol: whereas pilots of the task were well-received by infants and parents, the density of assessment during the study visit meant that infants often were too fatigued to complete this task. Additionally, participants were more reluctant to touch or play with the tablet in the lab setting, even after encouragement from examiners and mothers. Attempts were made to complete the task at separate home visits. However, this had disadvantages as it increased the burden on the testing team and introduced wider variance in testing conditions. Therefore, the task was excluded from the main BRIGHT protocol in The Gambia.

#### *Questionnaires/interviews – infant/child* Communicative development inventory (CDI)

The McArthur-Bates CDI (Fenson *et al.*, 2007) was used to assess language development at the 12, 18 and 24 month time points. In the UK, the full English version was used, which consists of a vocabulary checklist that asks parents to report how many words their child can understand and how many they

can understand and say. The questionnaire also asks about the child's use of grammar and gestures.

An adaptation of the CDI was developed for use in the Mandinka language, following guidelines outlined by the MacArthur-Bates CDI Advisory Board. To construct an inventory of words for use in the Mandinka adaptation, a list of 200-250 words was compiled; these were taken from the standard CDI, the Malawian CDI and the Senegalese CDI. Mandinka-speaking field staff translated these words into Mandinka and suggested alternatives if words were not applicable in the West Kiang district (i.e., baby buggy/stroller) or more affected by seasonality (i.e. some food items). The inventory probed specific categories from the MacArthur-Bates CDI, such as animals, food and drink, and clothing. It also asked whether the child had started to combine words and use more complex sentences. Subsequently, this inventory underwent pilot testing in two phases, with a total of 60 mothers of children aged 24 to 48 months. During the first phase, 30 women were interviewed, and the list was revised, removing words that were not frequently endorsed and adding new words that had been suggested by the pilot participants themselves. Subsequently, a second phase of pilot testing was conducted, where a further 30 women were interviewed. From these interviews, the list was reduced further by (1) eliminating all words which less than 10% of mothers said their child knew; (2) selecting 54 words of moderate difficulty (known by 40 to 70 % of the children); (3) selecting 18 easy words known by 70 - 100% of the children; and (4) selecting 18 more difficult words for which 10-40% of the children knew. This adaptation received full approval by the CDI committee as an official adaptation into the Mandinka language.

# Parental report sleep diary (adapted from the brief infant sleep questionnaire)

A daily sleep log was administered over three consecutive days in the week prior to each lab visit and averaged over the monitored period at 1, 5, 8, 12, 18 and 24 months of age. This diary was adapted from the Brief Infant Sleep Questionnaire (Sadeh, 2004) and methods for parental reporting (Sadeh, 1996) into a three-day diary differentially for each site. In the UK the diary was sent to caregivers by post, and completed by caregivers, with an option to fill out an online version if preferred. The diary asked for a record of all periods of sleep (time and location) over a 72-hour period including information on anything that made the day/night unusual relative to their regular routine (i.e., illness/activity). In The Gambia the questionnaire was adapted in several ways following advice from the local ethics committees and through the formation of a consultation group comprised of local research staff who live and work in the West Kiang region of The Gambia and who had young children (i.e., to provide guidance on the range of locations parents might use for daytime naps). Firstly, the local population in West Kiang do not adhere to strict observance of equinoctial hours, nor do they necessarily possess a time piece in each home. Therefore, the diary was adapted so that caregivers could answer questions about sleep based around the sections of the day that are divided by prayer calls from the local mosques (i.e., morning, afternoon, evening, last prayer time). The prayer time calls differed

slightly by season, which will be accounted for in analyses. Therefore, while fragmentation of sleep, location and number of daytime naps could be accurately recorded, length of sleep was restricted to an approximation. As with other questionnaires, the sleep diary was administered as an interview at the lab visit with caregivers asked to recall the previous three days and nights. While it would have been more accurate to interview caregivers day by day, the research team did not have the capacity for this many home visits. Families were, however, reminded to attend to their child's sleeping patterns for three days prior to the study visit when they were notified about their visit date the week before the scheduled research visit.

# Questionnaires/interviews – Family Family Care Indicators (FCI)

In The Gambia, families were asked to complete the FCI questionnaire when the infants were 12, 18 and 24 months of age. The development of this set of indicators was initiated by UNICEF to provide measures of family care practises and resources with globally relevant application (Kariger et al., 2012). The items measure the support provided by caregivers for a stimulating environment for infants to learn from, and the caregiving resources available within the home. For example, caregivers were asked about number of books and play items in the home, who was engaging with the child at home and how many different types of stimulating activities the child was encouraged to do. The questionnaire was developed by an international panel of experts who reviewed existing surveys used in low- and high-income countries (i.e., Home Observation for Measure of the Environment Inventory; Caldwell & Bradley, 1984) and field-tested new candidate questionnaire items across populations in five low-income countries before finalising this set of indicators for use in global health and epidemiological studies.

# Parental mental health

Parental mental health was assessed using a range of questionnaires, starting at the antenatal visit and followed up until the 24-month time point. In the UK, the original English versions of the questionnaires were given to both parents (where applicable) to complete in their own time. In The Gambia, questionnaires were translated into Mandinka and administered in interview format (see below). Since mothers always accompanied infants to visits and fathers were often working away from home, we could only collect data on maternal mental health.

The Edinburgh Postnatal Depression Scale (EPDS) (Cox *et al.*, 1987) was administered at the antenatal, 1, 5, 12 and 24 month visits. This is a 10-item self-report questionnaire that asks participants to rate how frequently they have experienced a range of depressive symptoms in the last seven days. Items are scored on scale of 0-3 ("No, not at all" to "Yes, most of the time") and possible scores range from 0 to 40. A cut-off of 10 is considered to indicate elevated levels of depression. The EPDS is a validated tool used to screen for postnatal depression and has previously been used to assess

maternal mental health in The Gambia, as well as other LMICs (Coleman *et al.*, 2006; Nabwera *et al.*, 2018).

The Positive and Negative Affect Schedule (PANAS) (Watson *et al.*, 1988) was administered at the 1, 5, 12 and 24 month visits. This 20-item self-report questionnaire asks participants to rate how frequently they have experienced a range of positive and negative emotions in the past few hours. Items are scored on a range of 1-5 ("Very slightly or not at all" to "Extremely"). There are six items that correspond to the Positive Affect (PA) and the Negative Affect (NA) scales, which are summed to compute scores for each scale, with a possible maximum of 30 for each scale.

The Percieved Stress Scale (PSS) (Cohen *et al.*, 1983) is a 10-item self-report questionnaire that asks participants to rate how often they have experienced a series of stress-related feelings in the last month. Items are scored on a scale of 0-4 ("Never" to "Very often"), with a possible total score of 40. The PSS has been shown to have robust psychometric properties across diverse low- and middle-income settings (Katus *et al.*, 2022a) and across different modes of assessment (Murray *et al.*, 2023). The PSS was administered at the antenatal, 1, 5, 12 and 24 month visits.

The Pregnancy Related Anxiety Scale (PRAS) (Rini et al., 1999) and the Pregnancy Specific Anxiety scale (Roesch et al., 2004) were used as measures of anxiety related to pregnancy at the antenatal visit only. The PRAS is a 10-item scale that asks respondents to rate how frequently they have experienced a range of concerns related to their pregnancy in the last few months. The scale is rated on a scale of 1-4 ("Not at all" to "Very Much" or "Never" to "A lot of the time"). The total score is computed by summing scores on all items, with a maximum score of 40 possible for the scale. The PSA asks participants to rate how often they have felt a range of emotions in the last week. Scores range from 1-5 ("Never" - "Always"). A total score is generated by summing the scores on four items that are specific to anxiety (anxious, concerned, afraid, panicky). Total scores can range from 4-20. The paternal versions of these questionnaires ask fathers to rate their feelings in reference to their partner's pregnancy.

Over a period of eight months (2015–2016) each questionnaire was adapted following World Health Organization guidelines (World Health Organization, 2013), the questionnaire developers, and procedures described in other studies using these measures in LMICs (Hanlon *et al.*, 2008; Kohrt *et al.*, 2016; Nabwera *et al.*, 2018; Tesfaye *et al.*, 2010; Weobong *et al.*, 2015). The adaptation protocol was the same for all measures and involved a core team of five researchers, as well as an additional nine staff members who supported this intermittently where needed. an initial translation from English to Mandinka by a panel of three Gambian research staff, the local PI (MD), who were all native speakers, and two researchers from the UK who were experienced in mental health data collection. Following best practice guidelines outlined by (Peña, 2007), we attempted to align the translated items as closely as

possible to the original English, while taking into account cultural equivalence. This involved replacing English idioms and medical terminology with phrases that would be comprehensible in Mandinka, and changing the structure of statements into questions (Kohrt et al., 2016). The translation process involved several rounds of back translation by Mandinka-speaking staff who were blind to the original questionnaires and the translation process, as well as several meetings with local clinical staff (i.e. midwives) and visits to families to discuss the language of the questions with mothers and pregnant women. Where discrepancies were noted between the original English and the back-translation, or regional variations in wording identified, the panel made necessary adjustments. The translated questionnaires were pilot tested with N=12 volunteers from West Kiang, to assess their understanding of the measures, corrections were made where issues with comprehension emerged.

While every attempt was made to ensure equivalence between the original English and the Mandinka translations, one item on the EPDS (item 10) that asked about suicidal ideation/behaviours was changed because of the highly sensitive nature of the question in this culture and local population. (Nabwera et al., 2018) noted that, due to the highly communal way of life in this community, the desire to be isolated from others was seen as a sign that the individual may be suffering from a mental health problem. Therefore, this item was changed to ask participants whether they wanted to be isolated or alone. Participants who scored above clinical cut-off (a score of 10) on the EPDS were given the opportunity to be referred to the MRC clinic for support. Furthermore, some of the English words used, in the PANAS in particular, were not differentiable in Mandinka, and so the number of items were reduced in The Gambian version relative to the one administered in the UK.

To simplify administration and reduce recall of response options, mothers were first asked whether they had experienced the issue described in the question (yes/no) and, only if they responded with a yes, would the interviewer elaborate with the frequency options (Hanlon *et al.*, 2008). To help mothers remember the time period that each questionnaire was referring to, they were administered in order of timeframe, from shortest to longest, and the timeframe was reiterated with each question. Finally, to reduce the impacts of stigma, mothers were reminded that all participants were being asked the same questions.

Field staff responsible for administering the interviews received extensive training on understanding the conceptual framework of each measure. Subsequently, they were trained in administration using vignettes and role play scenarios, practicing administration and managing different types of potential responses.

# Socioeconomic status (SES), demographic and family information

Families were asked to complete a questionnaire (UK) or interview (The Gambia) regarding their family demographics and socioeconomic circumstances. These were conducted as a series of questionnaires/interviews spanning from the first antenatal visit across the postnatal sessions, tailored to ask questions relevant to each time point, and with reduced time burden at each session, given that questions became an update on whether circumstances had changed. At both sites information was gathered on biological parents, biological grandparents (ethnicity, date, and place of birth) and any other applicable caregivers of the key participating infant/child. The caregiver information gathered included ethnicity, age, caregiving role, employment status, highest level of education, languages spoken. Wider family information was collected including parity of parents, size, and composition of household. Finally, housing information was collected pertaining to indication of wealth, i.e., number of bedrooms (UK), wall/floor materials, access to water, durable assets (The Gambia). In The Gambia this information was gathered through a combination of observation by the field assistant during home visits and reports from the participant during interview.

#### Pregnancy, birth and family health information

In the UK and The Gambia mothers were asked to complete a questionnaire (UK) or interview (The Gambia) regarding their pregnancy, birth, and family medical history. Information gathered included: (i) antenatal information on maternal obstetric and medical history; (ii) fetal ultrasound information including gestational age and anthropometric foetal measurements (The Gambia only); (iii) birth and delivery information including neonatal anthropometric measurements (The Gambia only) and a maternal health check.

# Growth and diet measures

# Anthropometric measures

Anthropometric measurements were made by research assistants in the UK and by field assistants in The Gambia. In both sites, measurements were taken in triplicate, following standard protocols and all staff underwent training. In the Gambian sample, maternal height and weight were measured in late pregnancy and infant length, weight and head circumference were measured at birth. In addition, infant length, weight, head circumference, mid-upper-arm-circumference (MUAC) and knee-heel length were measured at both sites at 7-14 days, 1, 5, 8, 12, 18 and 24 months of age. At birth, infant length was measured using a flexible length mat, and a fixed length board (SECA 417) was used thereafter. Infant weight was measured using a calibrated electronic baby scale (SECA 336), with a precision of 10g. Mid-upper-arm circumference and head circumference were both measured using a SECA 201 head and body measuring tape, precise to 1mm. Knee-heel length was measured using a calliper, also precise to 1mm.

### **Dietary data**

In The Gambia, infant feeding data was collected every two weeks from birth to 24 months of age. The questionnaire was administered verbally by a field assistant at the participant's home or by telephone if a home visit was not possible. The mother was asked to report on the infant's diet in the two weeks prior to the questionnaire. Details included whether the infant received breastmilk feeds, and/or other liquids, semi-solid or solid foods. The questionnaire included common examples of local weaning foods, as well as free text space for additional items. The mother was also asked to report the frequency (never, once, more than once, most days) at which the infant received each food or drink.

In the UK, similar feeding questionnaires were completed monthly by parents online, from 1-7 months of age. In addition, parents completed a detailed food diary reporting all food and drink consumed by their infant, for four consecutive days prior to each study visit from 8 months onwards (8, 12, 18, 24 months). This data was then coded on a food composition database (DINO; (Fitt *et al.*, 2015)). Mothers also completed online 24-hour dietary recall questionnaires reporting on their own diets, in pregnancy and at 6 and 12 months postnatally, using the Intake24 UK platform.

#### **Biological samples**

In The Gambia, to investigate nutritional factors in more detail, samples of breast milk, blood and urine were collected. Maternal venous blood and urine were collected in late pregnancy (34–36 weeks' gestation) and breast milk was collected from the mother at the five-month visit. Infant urine was collected at 5, 12 and 24 months of age. Samples were stored at  $-70^{\circ}$ C for subsequent analysis. In addition, infant blood samples were collected at all infant visits, alternating between a 0.5mL capillary sample (at 1, 8 and18 months) and a 3mL venous sample (5, 12 and 24 months). On each sample, a full blood count was run using a Medonic analyser and the remaining sample was centrifuged. Plasma and cell pellets were separated and stored at  $-70^{\circ}$ C for subsequent nutritional analysis and DNA extraction, respectively.

#### Standardisation of protocol across sites

All measures were collected using Standardised Operating Procedures (available upon request). Longitudinal infant and toddler testing requires standardisation of the (i) equipment (ii) environment in which the measures are administered, (iii) experimental protocol, and (iv) behaviour of researchers during administration of measures. The site in The Gambia had not previously undertaken research of this kind until the pilot phase of the BRIGHT project (Lloyd-Fox et al., 2014; Milosavljevic et al, 2019). Therefore, all equipment and testing materials had to be purchased prior to the start of the study sessions. To reduce site differences due to hardware, an identical set of equipment was purchased for both sites. While it is challenging outside of a research lab context to replicate the environment that the testing is undertaken in, where possible, we replicated the UK room setups at the Gambian site. In the UK (both at the hospital and university sites) testing rooms for visits from 1 - 24 months of age were sound proofed and windowless with temperature and lighting control. In The Gambia, the rooms were air conditioned to control the temperature, and the neurocognitive testing (fNIRS, EEG, eye-tracking) room was windowless with some light control. However, none of the rooms were sound-proofed. Therefore, environmental noise was more inconsistent across data collection within the Gambian sample as external

sounds could sometimes be heard within the testing rooms. At both sites testing at 7 - 14 days of age was done at the family's home, therefore environmental noise differed between the cohorts (for example family size was generally larger in The Gambia and houses often had open windows and doors- see Table 3). To address this, researchers at both sites optimised data quality where possible by discussing the needs of each measure with the family who were present during data collection (*i.e.*, they discussed with the family that there would be times when they needed the room to be quiet during a measure of attention to sound or light, or when they might need their help in eliciting a smile from their baby)).

Broadly, the protocol was identical across sites. As described above, some measures were site-specific either because a measure was found to be unsuccessful at one site (*i.e.*, the tablet task) or because the measure is only relevant at one site (*i.e.*, FCI). For the neurocognitive testing we adopted the use of the TaskEngine framework, which was developed for a separate multi-site neurocognitive study, Eurosibs (Jones *et al.*, 2019), to optimise data quality and standardisation of acquisition. This framework allows the presentation of the paradigms to be identical across sites and produces identical data outputs for cross-site quality control reports and analysis.

As described above, the order of testing was kept consistent across sites, however we have adopted the practice of standardization with flexibility to be responsive to the needs of the individual infant/toddler. Before the onset of the project, project coordinators for both sites were trained in the UK for a period of two months. Following this, training continued in The Gambia for one - two months (depending on the measure) and the first study sessions were conducted under supervision. The BRIGHT project team were committed to building long-term capacity for neurodevelopmental research at MRC Keneba and therefore across the duration of the project, trained and supported local researchers to conduct and co-ordinate all aspects of the research in The Gambia. To facilitate harmonization across sites, staff were trained in the practical detail of data collection and administration as well as researcher responsiveness to infant behaviour.

Working in partnership with local researchers during their training phase, we discussed the observed infant responses and behaviours elicited by each paradigm and used this experience to devise a standardized strategy for responding to the needs of the infant and caregiver. This strategy was agreed at both sites and included in the Standard Operating Procedures (SOPs). These steps took into account the infant's state of alertness and fussiness, as well as the caregivers needs, to ensure optimal data collection and participant comfort. Significant fussiness typically leads to inadequate data quality across both the neurocognitive and behavioral measures. Fussiness was defined as excessive motion (i.e., infant wriggling on lap, or toddler walking away from task), inattention to the task (i.e., looking away) or negative affect (i.e., crying, negative vocalizations). Furthermore, particularly in The Gambia, a further category of inattentiveness existed as infants,

**Table 2. Participant characteristics (age/sex) and retention rate of cohort at each time point.** Note: If infant became tired or fussy before session was complete a call back was arranged for a second visit to complete testing; DOV = date of visit; N = sample size; SD = standard deviation of mean.

The Gambia (N = 214)							
Sex (female/male)	103/111						
Maternal age at birth Mean, SD (min-max)	29.76 (6.61)	), 18.2 – 44.7					
Timepoint	7–14 days	1 months	5 months	8 months	12 months	18 months	24 months
Enrolled at DOV (N)	205	204	200	193	192	191	185
Attended visit (N)	157	185	198	188	188	177	161
Mean age in days (SD)	12.3 (3.87)	36.0 (5.64)	159.9 (10.14)	247.2 (11.39)	372.7 (14.05)	558.6 (17.13)	745.0 (27.94)
Range in days	5 - 44	29 - 65	148 - 208	211 - 314	353 - 428	533-641	722-896
% attended visit	76.5	90.7	99.0	97.4	97.9	92.7	87.0
% attended two visits per age point*	n/a	3.2	22.2	23.9	25.9	37.9	16.8
UK (N = 62)							
Sex (female/male)	31/31						
Maternal age at birth Mean, SD (min-max)	32.96 (2.93)	), 28.4 – 40.8					
Timepoint	7–14 days	1 months	5 months	8 months	12 months	18 months	24 months
Enrolled at DOV (N)	62	61	60	60	60	57	50
Attended visit (N)	58	60	58	57	59	55	50
Mean age in days (SD)	12.2 (3.33)	33.2 (5.53)	155.8 (6.54)	251.7 (9.89)	375.7 (12.51)	557.1 (15.02)	736.9 (15.79)
Range (days)	7 - 23	22 - 56	144 - 184	235 - 279	353 - 411	536 - 603	700 - 784
% attended visit	93.6	98.4	96.7	95	98.3	96.5	100
% attended two visits per age point*	n/a	n/a	n/a	n/a	n/a	27.3	20

on occasion, become drowsy or fell asleep during the task, likely an effect of the climate. We agreed upon a hierarchy of responses, tailored to the task, to ensure maximum participation. For example, during screen-based tasks (i.e., eye-tracking, fNIRS) to address possible boredom, when the infant began to fidget and look away, "attention-grabbers" (non-social short sounds) were employed to re-orient the participant to the screen. These were automatically recorded by the task presentation framework. If this was ineffective the researcher moved through the following strategies (also applicable to other measures); parents were asked to speak reassuringly and hold hands, the infant was given something to hold or at older age points infants/toddlers are offered a snack (i.e. rice cake/rusk), a short break was offered before re-engaging with the task, or if none of these strategies were successful families were given a longer break for a nap or feed. If the parent was happy to resume the session continued. If a participant was unable to complete the full session within a day, families were asked if they were happy to return on a subsequent visit. All "manual" strategies of engagement and breaks for naps and food are recorded in the Session Log Form.

#### Data analysis plan

## **Quality control**

Frequent refresher training sessions and quality control have been implemented across the duration of the BRIGHT project. At the Gambian site, due to the number of participants, time frame of testing and number of age points measured, testing load was very high at the peak of the project – with up to four infants tested per day, seven days per week, with staff on rotating schedules - for over two years. To maintain high data quality, research staff were trained to be highly specialised in a subset of measures rather than every measure. A researcher might oversee the neurocognitive tasks (EEG, fNIRS, eye-tracking), behavioural measures (MSEL, NBAS), mental health parent interviews or anthropometric measures.

**Table 3. Demographic characteristics and socioeconomic status, The Gambia.** This socioeconomic information is derived from that reported or observed at the 7–14-day home visit. *n*=172; IQR = inter quartile ratio; NA = not applicable.

	Mothers (%)	Fathers (%)	
Education			
No formal education	59.4	55.0	
Some primary education	12.9	5.9	
Complete primary education	3.5	4.7	
Some secondary education	18.2	7.1	
Complete Secondary education	5.9	27.2	
Household characteristics			
Number of children, Median (IQR) min-max	5(4), 1-10	6 (7), 1-23	
Number of wives	NA	1 (1), 0-4	
Assets			
	Household (n min-		
Number of people in household	11 (8)	, 3-36	
Diet			
Meals per week containing meat	1 (1)	, 0-5	
Meals per week containing fish	6 (1)	, 0-7	
Meals per week containing fish	7 (0), 2-12		
or meat	/ (0//		
or meat Housing Attribute		olds (%)	
Housing Attribute		olds (%)	
Housing Attribute Primary Water Source	Househ	<b>olds (%)</b> 25	
Housing Attribute Primary Water Source Open Public Well	Househ	<b>olds (%)</b> 25 37	
Housing Attribute Primary Water Source Open Public Well Protected Public Well	Househ 2.2 7.8	<b>olds (%)</b> 25 37 2.6	
Housing Attribute Primary Water Source Open Public Well Protected Public Well Public Tap	Househ 2.: 7.8	<b>olds (%)</b> 25 37 2.6	
Housing Attribute Primary Water Source Open Public Well Protected Public Well Public Tap Piped water in compound	Househ 2.: 7.8	olds (%) 25 37 26 25	
Housing Attribute Primary Water Source Open Public Well Protected Public Well Public Tap Piped water in compound Cooking Fuel	Househ	olds (%) 25 37 2.6 25 5.1	
Housing Attribute Primary Water Source Open Public Well Protected Public Well Public Tap Piped water in compound Cooking Fuel Firewood	Househ	olds (%) 25 37 2.6 25 5.1	
Housing Attribute Primary Water Source Open Public Well Protected Public Well Public Tap Piped water in compound Cooking Fuel Firewood Charcoal	Househ	olds (%) 25 37 26 25 3.1 95	
Housing Attribute Primary Water Source Open Public Well Protected Public Well Public Tap Piped water in compound Cooking Fuel Firewood Charcoal Toilet Facilities	Househ	olds (%) 25 37 26 25 3.1 95	
Housing Attribute Primary Water Source Open Public Well Protected Public Well Public Tap Piped water in compound Cooking Fuel Firewood Charcoal Toilet Facilities Pit Latrine	Househ	olds (%) 25 37 26 25 3.1 95 07 69	
Housing Attribute Primary Water Source Open Public Well Protected Public Well Public Tap Piped water in compound Cooking Fuel Firewood Charcoal Toilet Facilities Pit Latrine Improved Pit latrine	Househ	olds (%) 25 37 26 25 3.1 95 07 69	
Housing Attribute Primary Water Source Open Public Well Protected Public Well Public Tap Piped water in compound Cooking Fuel Firewood Charcoal Toilet Facilities Pit Latrine Improved Pit latrine Flush Toilet	Househ	olds (%) 25 37 26 25 3.1 95 07 69	
Housing Attribute Primary Water Source Open Public Well Protected Public Well Public Tap Piped water in compound Cooking Fuel Firewood Charcoal Toilet Facilities Pit Latrine Improved Pit latrine Flush Toilet Flooring	Househ	olds (%) 25 37 26 25 37 25 3.1 3.1 35 3.1 35 3.1 35 3.1 35 35 37 37 37 37 37 37 37 37 37 37 37 37 37	
Housing Attribute Primary Water Source Open Public Well Protected Public Well Public Tap Piped water in compound Cooking Fuel Firewood Charcoal Toilet Facilities Pit Latrine Improved Pit latrine Flush Toilet Flooring Earth, Sand, Mud	Househ	olds (%) 25 37 26 25 3.1 25 07 69 25 25 25 25 25	
Housing Attribute Primary Water Source Open Public Well Protected Public Well Public Tap Piped water in compound Cooking Fuel Firewood Charcoal Toilet Facilities Pit Latrine Improved Pit latrine Flush Toilet Flooring Earth, Sand, Mud Cement	Househ	olds (%) 25 37 26 25 25 3.1 25 07 69 25 25 25 25 25 24 2.4 2.2 26	

Housing Attribute	Households (%)
Roofing	
Corrugate	100
Walls	
Mud	12.9
Earth Bricks	3.93
Cement/ Burnt Bricks	82.0
White Lime	1.12
Household Assets	
Electricity	2.81
Television	16.3
Refrigerator	5.06
Bicycle	73.0
Motorbike	14.0
Other vehicle	7.87

For example, for fNIRS studies, to minimize data loss, designated researchers were trained to monitor the system's performance, detect potential problems with the acquisition and instructed to implement basic repairs (Blasi et al., 2019). Designated researchers with expertise in a particular measure were also supervised by a senior team member with specialist knowledge of the measure or paradigm. The majority of the senior team members were located in the UK. From the start of the project, web-based multi-site meetings were routinely held. For the duration of the project, these consisted of fortnightly group meetings (which include senior team members and all designated researchers at the sites of data collection) and fortnightly quality control meetings (which included at minimum one senior team members and designated researchers). At these meetings recruitment, data quality, testing practices, outputs and other issues were discussed. Furthermore, during the quality control meetings training sessions were also conducted, including researcher responsiveness, as and when the need arose. For example, during the life course of the project, training in the MSEL had to be administered as each new age point was reached to ensure the data was being collected in a standardised way across sites. During the BRIGHT-Kids phase of the project, training on all measures was complete, therefore, team meetings were reduced to fortnightly quality control meetings until completion of data collection.

After data collection, each data set required extensive quality control assessments.

For example, EEG and NIRS data needed to be assessed for artefact from motion and infant inattentiveness, and segments of data removed as appropriate. Session level data from the "Session Log Form" was also reviewed so that contextual variables such as infant state (*e.g.*, fussiness), interference from

researcher/caregiver (*e.g.* talking during task), experimenter error (*e.g.* video not recorded), technical issues (*e.g.* computer program crashed, headgear misaligned for NIRS/EEG), or position of participant (*e.g.* infant facing wrong way, or parent out of view during interaction video) could be taken into account. Data was marked for validity at each stage of this process to monitor data attrition during the stages of the processing stream. Data quality metrics were extracted for all paradigms at each site and age point. Specific guidance for conducting EEG and fNIRS studies in global health contexts to maintain high quality control measures can be found in two methods papers concerning the BRIGHT project (Blasi *et al.*, 2019; Katus *et al.*, 2019)

#### Data storage and handling

While it would be preferable for data pre-processing and analysis to be conducted at the site of data acquisition, such skills are highly specialised and usually attributed to post graduate education in neuroimaging and such expertise among local researchers in The Gambia is, as of 2023, very limited. This is likely because developmental neuroscience is a very new field of research in The Gambia, and as such, opportunities to learn the required analytic skills have previously been lacking. This has been identified as an important focus for capacity building in the long term. However, to ensure timely data quality control during the BRIGHT project, much of the data analysis was conducted within the UK and a reliable data transfer method had to be established to enable this. A dual protocol was designed to ensure the integrity of the data transfer.

As a standard procedure, initially, the data was stored and backed-up locally on site in a separate location to the research data acquisition computers. Personal data (i.e., contact details, DOB) was stored securely at each site within locked cabinets as well as within a password encrypted electronic database isolated from the research data. Data from the anthropometric measures, parent-report questionnaires, and behavioural assessments (i.e., NBAS, MSEL) was pseudonymised and housed in password-protected encrypted databases locally. All data obtained using paper forms was double entered on the local databases to ensure reliability across staff. Experimental data (*i.e.*, eye-tracking, neuroimaging measures, parent-infant videos) was pseudonymised and stored on password encrypted storage hard drives at each local site. Following this, a Secured File Transfer Protocol (SFTP) server functioned as a bridge between sites. This system allowed the transfer of data in both directions to account for (i) planning of protocol updates, (ii) software and stimuli transfer from the development site (London, UK) to cohort sites (Cambridge, UK and Keneba, The Gambia), (iii) data transfer from the cohort site and (iv) feedback on pilot data and quality control checks (see Blasi et al., 2019 for further detail). A second full copy of all research data was transferred to a secure password encrypted server in one of the participating UK centres (at the Centre for Brain and Cognitive Development, Birkbeck, University of London).

One consistent challenge was the unreliable internet and/or poor bandwidth, which lead to slow transfer times for large files. The fNIRS tasks of the BRIGHT project, for example, involved acquiring over 160 files, equating to over 3GB of data, in total across all time points for each participant (Blasi *et al.*, 2019). Furthermore, during the life course of the project over 25,000 data files were collected for the infant/ child measures alone (this excluded datasets for anthropometric data, biological data and family questionnaire data). Therefore, when the rate of data transfer became restricted, we prioritized the transfer of specific data types based on the data quality control checks and analysis pipelines required for each.

Data transfers, quality control checks and inventories were carried out at regular intervals. Access was fully audited, and to ensure data security, access is governed by a management team. A web-based interface enables internal BRIGHT researchers to access the database using personalized login details to search, filter, and download data. Data access is overseen by the data management team in the UK and The Gambia, and access is granted for internally pre-registered projects (see Data.

#### Data analysis and statistical plan

Overall, we aim to identify developmental brain and neurocognitive function-for-age curves across the first two years of life, establish which context associated moderators (*i.e.*, under nutrition, family income, caregiving/family support) impact significantly on infant development and how these associate with pre-school outcomes at three to five years. Further to the plans outlined for each aim below, tests of normality and sensitivity analyses (comparing observed values and imputed missing values) will be conducted. Non-linear tests of significance and interpolation approaches will be applied where appropriate.

To address Aim 1, brain and neurocognitive function-for-age curves from birth to 24 months of age will be generated across 1, 5, 8, 12, 18 and 24 months of age. These will be derived from the fNIRS, EEG, MSEL and neurocognitive eye-tracking batteries of tasks. These reference curves will be used to enable age-adjusted group comparisons of differences in average trajectories, group-wise differences in variability, and for characterizing the range of individual developmental trajectories within each cohort. To generate appropriate metrics for each dataset we will explore which type of derivative measures and the level of complexity required to meaningfully capture brain and cognitive change across this developmental window. For the neuroimaging data we will explore these derivative measures (i.e., localisation versus globalisation of brain response, latency of response, profiles of change in haemoglobin) by running time varying parameter models to explore variation across and within these values over our varying age points. Following this, several approaches will be undertaken, including longitudinal growth modelling to explore relationships between measures, and regularisation methods such as latent function connectivity modelling, to explore whether we can derive a common measure across the different brain function tasks which predict rate of change in other measures such as MSEL and the neurocognitive battery.

To address Aim 2, a combination of quantitative analyses will be undertaken to establish the association between contextassociated moderators (i.e., undernutrition in mother and infant and consequent growth faltering, parental mental health, SES factors) and developmental trajectories across the first two years of life in The Gambia. To generate appropriate metrics for each dataset we will explore which type of derivative measures most meaningfully capture measures of poverty (i.e. maternal iron status versus infant iron levels, physical growth at birth, versus change in growth measures over time, SES measures of income, household size, household assets). Structural equation modelling (SEM) and hypothesis-driven regressions will explore how these latent pre- and post-natal variables associate with latent outcomes of infant development and regression analyses conducted to understand the directional relationships between outcome variables.

To address Aim 3, outputs from Aims 1 and 2 will be nominated using lasso regression coefficients in relation to longitudinal brain and cognitive development trajectories (*i.e.* across language, motor, sensory, attentional correlates, brain connectivity, ) across 0–24 months of life using SEM, applying full information maximum likelihood to account for missingness and to identify developmental-hypothesis driven clusters to explore how context-associated moderators (including risk factors of poverty) of altered infant development, and the impact of these in turn on pre-school outcome measures of early child development.

#### **Preliminary results**

Recruitment and retention of participants: The Gambia Figure 3 illustrates the recruitment and retention of participants in The Gambian cohort. In total 280 families were recruited and consented into this project. A total of 58 families were excluded from the study prior to delivery, as outlined in Figure 3, leaving a total of 222 families enrolled at delivery. Eight infants were stillborn, leaving a total postnatal cohort of 214 mother-infant dyads. A further seven infants were lost to neonatal death during the first two weeks of life and two families chose to withdraw from the study prior to the first postnatal home visit at 7–14 days of infant age, leaving 205 mother infant dyads within the study. At the 24-month time point, 185 families remained enrolled within the study.

#### Recruitment and retention of participants, UK

Figure 4 illustrates the initial recruitment process in the UK, and the retention of participants throughout the study. In total, 62 families were recruited and consented into this study, undertook the antenatal assessment and took part in their first postnatal home visit when the infants were 7–14 days of age. Participating families live either in the centre of Cambridge (N = 22) or in surrounding urban or rural communities within a 20-mile radius (N = 40).

#### Participant attendance to study visits

This data was derived from the first data collection point and includes all available data for N = 214 (The Gambia) and N = 62 (UK). At each age point a preferred testing window was established; +/- 1 week at 7-14 days, +/- 2 weeks at 1 - 8months, +/-1 month at 12 - 24 months. To minimise data loss, we allowed data collection up to + 1 month at 5 - 8 months. and + 2 months from 12 months upwards. This became necessary on occasion, particularly if both parents returned to work or when families had travelled outside of the region and were unavailable at the time of test. The total number of participants who attended each visit is given in Table 2. Due to the COVID-19 pandemic, testing of the full 24-month time point was suspended in The Gambia during their first country-wide lockdown. Following a prolonged period of suspension of research, a decision was made to end data collection for the BRIGHT project, therefore 20 participants could not be invited for their final time point at 24 months of age. For those families enrolled at each time point (i.e., excluding those who had been withdrawn for health reasons or because they moved away) we have experienced a high retention rate for the majority of the completed time points (> 90% of enrolled cohort attended visits). In The Gambia the exception has been at the 7 - 14 days visit, where the proportion of families attending the visit fell to 76.6%. This occurred as a result of mothers who gave birth away from their village and would therefore stay with other family members during the first weeks of their infant's life. Consequently, it was sometimes challenging to identify that these women had given birth and arrange their first visits. Furthermore, during late 2016 - early 2017 the country experienced political unrest during the general elections, which impacted on our ability to schedule infants at 7-14 days and 1 month of age for their visit. In the UK, retention rates have been high in the first time points, for example 91.9% of participants were seen at all five of their visits during their first year of life. While the majority of those families were still enrolled in the study at 18 and 24 months of age and attended the visit, we experienced higher rates of self-withdrawal in the UK as the testing burden on some participating families became too high (the most common reasons for withdrawal were either that both parents had returned to work or that the mother was expecting/had given birth to a further child).

# Demographic characteristics and socioeconomic status (SES), The Gambia.

A summary of the family demographic and SES distribution of The Gambian cohort is given in Table 3. Families live in multigenerational households with up to 36 members per compound. Polygamy was common within the cohort with 38.8% of fathers having more than one wife. Consequently, while mothers had on average 4.4 children, including the infant enrolled in the study, fathers had on average 6.9, with a range of 1 - 23 children attributed to a single father.

For the generation of parents within our cohort, formal schooling was readily available when they themselves were children, therefore, on average, mothers and fathers within the study had completed three and four years of schooling respectively.

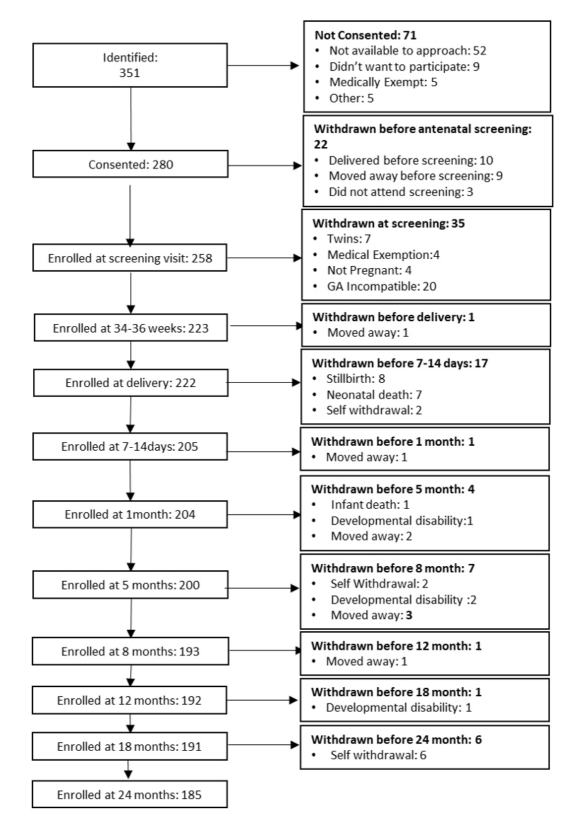


Figure 3. Recruitment and retention of participants, The Gambia.

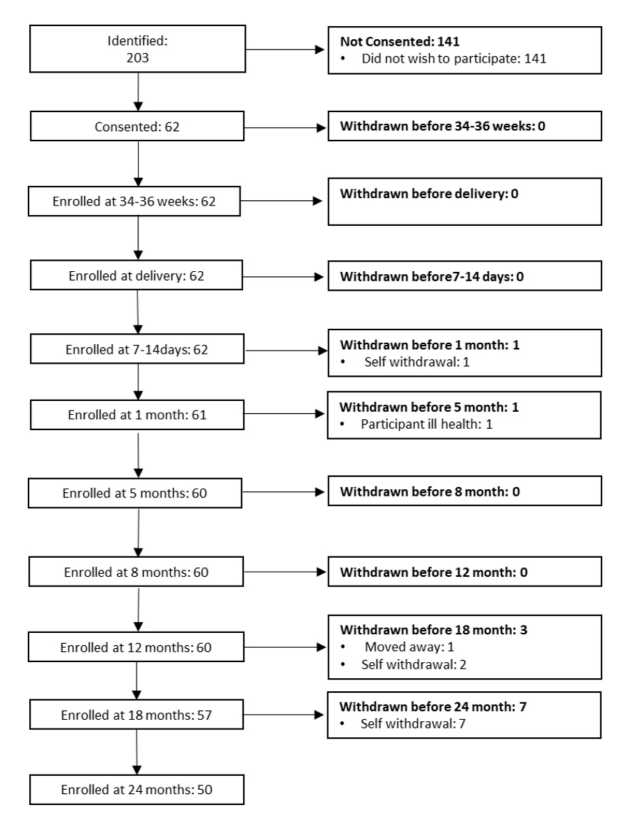


Figure 4. Recruitment and retention of participants, UK.

Over five times more fathers (26.6%) than mothers (5.62%) had completed high school (grade 12). Subsistence farming was the most common profession among both mothers (64.4%) and fathers (43.4%). In terms of livestock, fathers owned more livestock (sheep, goats, donkeys, or cows) than mothers. Goats were the most commonly owned livestock among both mothers and fathers, with 45.5% mothers and 46.6% fathers owning at least one goat. Cows are the most valuable livestock, and ownership was heavily skewed in favour of the fathers with 29.7% of fathers compared to 6.7% mothers owning at least one cow. Around one third (34%) of farming fathers sold a proportion of their produce, however the mothers' farm produce was largely consumed by the household, with only a small minority (6%) of mothers selling any of their harvest. Durable asset ownership was low, while 73.3% of households reported owning a bicycle, only 14% owned a motorbike, 7.9% owned a vehicle, 16.3% a television and 5.1% a refrigerator.

Within the cohort, the mother was reported to be the primary caregiver for all infants at 7–14 days of age, and for all but one child, where the primary caregiver was reported as the aunt, at 18 months of age. As can be seen in Figure 5, at 7–14 days of age, the most common secondary caregiver was the grandmother (41.1%), whereas by 18 months almost half (45.0%) of all participants reported an elder sibling to be the secondary caregiver. Fathers were reported as secondary or tertiary caregivers in 28.2% of families at 7–14 days, and in 32.2% of families when infants were 18 months old.

All primary caregivers reported that their first language was Mandinka. In addition, 16.7% of primary caregivers reported that they spoke a second language and 3.5% of primary caregivers spoke three languages. Mandinka was also the most common first language among secondary and tertiary caregivers, the only exceptions being three participants for whom their tertiary (n=2) or secondary and tertiary (n=1) caregivers' first language was an alternative local language. In addition, 25% of secondary caregivers and 13% of tertiary caregivers spoke a second language and a small minority spoke a

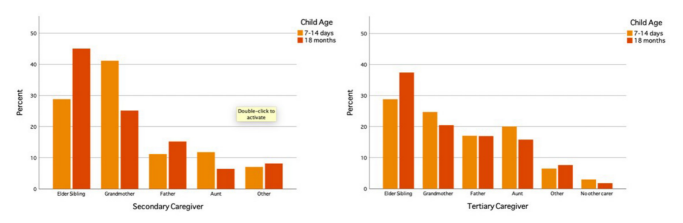
third language (2.3% and 4.1% of secondary and tertiary caregivers respectively) (Figure 6). Other than Mandinka, languages spoken included local languages such as Fula, Wolof and Jola, as well as English and Arabic.

# Demographic characteristics and socioeconomic status (SES): UK

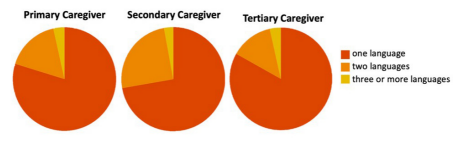
A summary of the family demographic and SES distribution of the UK cohort is given in Table 4.

As can be seen in Figure 7, at 8 months of age, caregiving in the UK cohort was primarily provided by the mother and/or father. Other reported caregivers at this age included grandparents and a childminder. Similarly, at 18 months of age, caregiving was primarily provided by the mother or jointly by the mother and father, with the exception of one family who reported the grandmother as the primary caregiver. Nursery workers, childminders and grandparents were also frequently reported as additional caregivers at this age (Figure 8).

In the UK we followed a recruitment strategy that encompassed natural population variance within the region of recruitment. A large proportion of the population living in the city of Cambridge and surrounding areas is multi-lingual. Consequently, a significant proportion of our recruited infants were exposed to multiple languages (Figure 8). As highlighted in Figure 9, exposure to languages both within, and outside of, the home was recorded antenatally and then at 8, 18 and 24 months of age. Within the home, the proportion of households exposed to English-only varied from 55 - 72.9% across age points. A further 13.6 - 27.5% of households had one additional language, 10 - 15.4% had two additional languages and 0 - 7.5%of infants were not exposed to English within the home at all. The languages that the infant was exposed to outside of the home differed with 75.6 - 85.7% of families reporting that infants were exposed only to the English language outside of home across 8 - 24 months of age. A further 11.4 - 14.7% reported that their infants heard English and one additional language outside of the home, one further family reported that their infants heard two other languages in addition to



**Figure 5. Additional caregivers at 7–14 days and 18 months of age, The Gambia.** The Figures display the make-up of secondary and tertiary caregivers at 7–14 days and 18 months of age in The Gambian cohort. Secondary caregiver; a person who looks after the infant a substantial proportion of the time. Tertiary caregiver; a person who looks after the infant 'sometimes'. A. n=170, B. n=165, C. n=171, D. n=168.



**Figure 6. Number of languages spoken by caregivers, The Gambia.** The Figures show the proportion of primary (n=172), secondary (n=171) and tertiary (n=167) caregivers at 18 months of age, who spoke one, two or three languages.

	Mother (%)	Father (%)			
Education					
Secondary	3.2	6.5			
Tertiary	3.2	4.8			
Undergraduate	33.9	33.9			
Postgraduate	54.8	48.4			
Family Ethnicity					
Caucasian	87.1	83.9			
Asian	3.2	6.5			
Black	1.6	1.6			
Mixed/Other	3.2	3.2			
Parental SOC Classification					
Higher managerial or professional	69.8	76.8			
Intermediate	22.6	19.6			
Routine/semi-routine	3.8	3.6			
Unemployed > 6 months	3.8	0			
	Household (mean	(SD), min-max)			
No of children (incl. key child)	1.19 (0.4	), 0-3)			
Annual Household Income	Percent of Hou	ıseholds (%)			
£20,000 - £29,999	1.6				
£30,000 - £39,999	1.6				
£40,000 - £59,999	25.8				
£60,000 - £79,999	37.1				
£80,000 - £99,999	21.0				
£100,000 - £149,999	6.5				
>£149,999	1.6				
Do not wish to answer	3.2				

Table 4. Demographic characteristics and socioeconomic status, UK.Note that this summary is compiled from the 18 month visit; SD = standarddeviation of the mean.

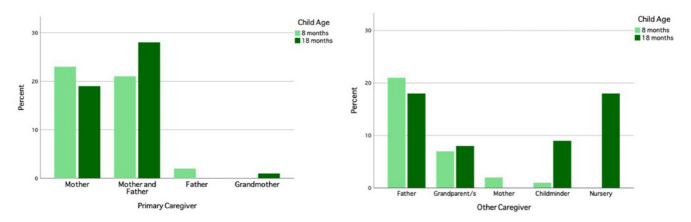


Figure 7. Caregivers reported at 8 and 18 months of age, UK. The Figure shows the primary caregivers and other reported caregivers at 8 and 18 months of age for the UK cohort. 8 months of age n=46, 18 months of age n=48.

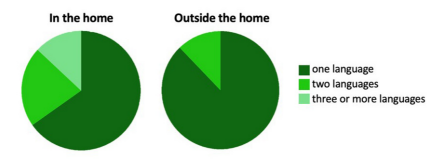
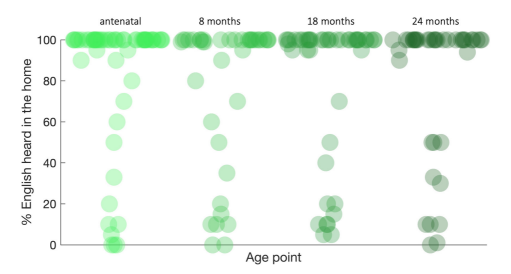


Figure 8. Number of languages heard in the home by infants at 18 months.



**Figure 9. Percent of English heard by UK infants across age points.** Note: While the majority of families completed this questionnaire antenatally (95.2%), the proportion of families who provided this information at 8 – 24 months dropped significantly to 62.9 – 74.2% complete datasets from the full cohort. This was due either to families withdrawing from the study at 18 – 24 months or missing data within the demographic questionnaire.

English, and one family reported that their infant did not hear English outside of their home at all. At this stage we wish to avoid premature consideration of cultural and linguistic factors that could affect the interpretation of our results within the UK, given that this was not a primary focus of our project. However, this factor will be taken into account during our planned analyses.

# BRIGHT Kids: Implementation, challenges, and successes

The protocol for BRIGHT Kids is outlined below and summarised in Table 5.

### Aims of BRIGHT Kids

While a growing body of research has begun to identify early markers of risk and resilience, the consequences of adversity often do not become fully manifest until later in childhood. Preschool age is a period where children experience rapid progress in a number of developmental domains (*e.g.*, language, executive functions), as well as an increase in external demands from caregivers. Consequently, this is a period where delays in development start to become outwardly observable and to interfere with everyday functioning. Thus, to fully understand the consequences of early exposure to context-associated moderators, it was necessary to continue tracking the development of children in The Gambian cohort beyond infancy and into the preschool period.

The preschool age follow-up (hereafter "BRIGHT Kids") has several key aims: (i) to assess a broader, age-appropriate set of developmental outcomes; (ii) to further elucidate developmental trajectories of the measures that had been administered since infancy; and (iii) to examine associations between biomarkers of risk in infancy and outcomes at preschool age.

To examine these aims at the BRIGHT Kids follow-up, we implemented a combination of existing and new measures, described below. Prior to inviting the BRIGHT sample for the follow-up, we recruited a separate pilot cohort of 24 participants to evaluate the implementation of the new measures into the protocol.

Participant retention, sample size and age distribution In 2021, all participants that had attended the 24-month visit were invited to take part in the BRIGHT Kids follow-up. Of these, 181 provided informed consent for participation. Two families were traveling and could not be reached and two declined to participate. A further three families withdrew from BRIGHT Kids after consenting and six remained in the study but did not attend the scheduled visits. Finally, one child was withdrawn from the study due to evidence of developmental delay (which interfered with their ability to complete any of the assessments). This left a final sample of 171 (49.7% female) participants that took part in BRIGHT Kids.

**Table 5. Protocol for BRIGHT-Kids.** Summary of abbreviations; fNIRS, functional near infrared spectroscopy; EEG/ERP, electroencephalography/event related potentials; L – length, W – weight, H – height, HC – head circumference, MUAC – mid to upper arm circumference, KHL – knee to heel length. \* Indicates a new measure introduced for BRIGHT-Kids.

Measure					
Neuroimaging measures	Questionnaires/Interviews – Infant/Child				
fNIRS: Social/Non-social	Food Insecurity Q (FFQ) *				
fNIRS: Habituation and Novelty Detection	Dietary Diversity Q (IFQ) *				
fNIRS: Functional connectivity	Early Childhood Development Index *				
EEG: Auditory Oddball	Questionnaires/Interviews – Family				
Behavioural/Neurocognitive measures	Generalised Anxiety Disorder – Maternal *				
Eye-tracking: Cognitive control	Patient Health Questionnaire – Maternal *				
Eye-tracking: Gap/Overlap	Perceived Stress Scale (PSS) – Maternal				
Eye-tracking: Non-social contingency	Impact of COVID-19 *				
Eye-tracking: Face popout	Family Care Indicators (FCI)				
Eye-tracking: Dynamic scenes	Clinical measures/ Medical details				
Eye-tracking: Word-picture-matching	Anthropometric measures – infant (L,W,HC,MUAC, KHL)				
Mullen Scales of Early Learning (MSEL)					
Tablet-based Early Years toolbox *					
Test of Gross Motor Development *					

As a result of over recruitment in the early stages of the BRIGHT project and interruptions to testing caused by a political crisis in December 2016-January 2017 (which resulted in the short-term closure of MRC Keneba), a recruitment pause was needed in the summer of 2017. This recruitment pause gave the number of infants born into the study per month a bimodal distribution. Furthermore, the initial BRIGHT-Kids phase of the BRIGHT project was conducted over a compressed time window relative to the earlier longitudinal time points of the BRIGHT project, due to funding and pandemic related restrictions during 2021-2022. Children were assessed starting with the oldest (first recruited) and ending with the voungest (last recruited). Therefore, the ages of children at follow-up reflected the same bimodal distribution as the number of infants born per month. The age range of participants in Phase I of BRIGHT-Kids was 45-63 months (M=52.8, SD=5.06); see Figure 10.

With the support of additional funding, due to this bimodal distribution of ages, a Phase II data collection of BRIGHT-Kids, has now been established to collect data during 2023. This additional follow-up is designed to assess the younger half of the age distribution again when they are age matched with the older participants from Phase I. The same BRIGHT-Kids protocol is being utilised to allow for us to assess potential outcome and exposure effects by collecting data from all participants when they reach 4 - 5 years.

#### Protocol for neuroimaging, ET, and MSEL

A number of paradigms that had previously been used in the BRIGHT protocol were retained in BRIGHT Kids. These included fNIRS tasks (social cognition, habituation and novelty detection, functional connectivity), EEG, ET tasks and the MSEL. The aforementioned assessments were either developed using stimuli that was suitable for broader age ranges throughout early childhood (fNIRS, EEG, ET) or had specific items suitable for preschool-aged children (MSEL). Using consistent paradigms enabled further tracking of developmental trajectories, which is informative because it tells us whether a particular biomarker is relevant at only one developmental/age period, or if it continually predicts outcomes throughout early childhood (Loth *et al.*, 2017). While the neuroimaging paradigms were suitable for the age group tested, several practical considerations were required to ensure suitability of the headgear.

**fNIRS headgear.** Data acquired at this visit were collected using a headgear based on an EasyCap, rather than the custom-built headband that were used at prior visits. These caps are made of soft fabric that covers the whole head. The optodes were held in place with purpose-made holders for a secure fit on the head and the design was changed to have a pointy, rather than flat, end to better penetrate hair in older children. Both of these changes allowed for better data acquisition, particularly considering the variety of hair types at this age.

**Practical considerations.** One challenge of undertaking fNIRS and EEG with older children is that they have more hair than infants, and children (girls in particular) often have their hair tied or in braids (Katus *et al.*, 2019). Several steps were taken to maintain data quality and account for a variety of hair types. If participants had large braids, mothers were asked to undo these before the study visit. When fitting the fNIRS or EEG caps, research staff brushed aside stray hair to ensure that the optodes/electrodes made contact with the scalp. To acknowledge this additional burden on participating families, and to thank them for their time, we hired an assistant, who was local in West Kiang, to act as a "hairdresser" for the participants. At the end of the assessments, she washed out the EEG gel from the children's hair and re-braided the hair of every child (with caregiver consent).

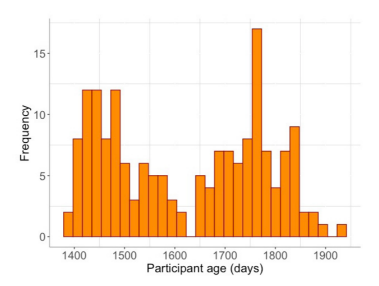


Figure 10. Distribution of participant age (in days) in the BRIGHT Kids follow up.

### *Protocol for additional behavioural measures* Gross motor skills

The *Test of Gross Motor Development* 3<sup>rd</sup> *Edition* TGMD-3 (Ulrich, 2019) was used to assess gross motor (GM) skills. The MSEL does not provide age-appropriate test items, nor does it describe developmental GM milestones, for children older than 39 months (Mullen, 1995). The TGMD-3 is suitable for children aged 3–11 years and assesses GM skills across two domains: locomotor and ball skills. The locomotor subscale tests skills that require coordinated and fluid body movements (*e.g.*, running, skipping, hopping). The ball skills subscale evaluates proficiency in throwing, catching and striking movements. Age-normed scores can be computed for both scales, which are then combined to form the Gross Motor composite score.

#### Tablet-based executive functioning assessments

Tablet based assessments from the Early Years Toolbox (Howard & Melhuish, 2017) were used to assess executive functions (EFs) across three domains: working memory ("Mr Ant" task), Inhibitory Control (the Go/No-Go task) and Cognitive Flexibility (Card Sorting task). These tasks were previously used among preschool aged children (3–5 years) in South Africa, who demonstrated good EF abilities and, on some scales, outperformed their Australian counterparts (Howard *et al.*, 2020). Task instructions were presented as an audio playing from the tablet. These had been translated into Mandinka and recorded by field staff and embedded into the app. While the tablet task was challenging at the earlier time point, we found this tablet task to be successful in The Gambia with our cohort at 3 – 5 years (Milosavljevic *et al.*, 2023).

#### **Parent-report measures**

We continued to use the Family Care Indicators (FCI, described above) to assess play materials, enrichment activities and caregiving. In addition to this, a number of new questionnaires were implemented to assess child development, caregiver wellbeing and the home environment.

### **Child development**

The *Early Childhood Development Index (ECDI2030)* is a caregiver report questionnaire that assesses the achievement of fundamental developmental milestones among children aged 24–59 months. The measure was developed by UNICEF with the aim of creating a tool that could be both nationally representative and also provide internationally comparable data on child development (for full details, see the UNICEF ECDI resources). The questionnaire consists of 20 items that assess a range of skills related to a learning, health, and psychosocial wellbeing. Since a Mandinka version was not available, we translated the tool following the customisation and translation guidelines outlined by UNICEF.

#### Maternal mental health and caregiving

Maternal wellbeing has important implications for child wellbeing and development beyond infancy and into school age (Bennett *et al.*, 2016; Kingston & Tough, 2014). Therefore, we continued to assess maternal mental health at the BRIGHT Kids visits. The Perceived Stress Scale (PSS, described above) continued to be used, as this assessment is not specific to pregnancy and was, thus, deemed suitable for this time point. In addition to this, two new assessments of depression and anxiety that were designed to assess these mental health constructs more generally and were not specific to pregnancy, were introduced. The Generalised Anxiety Disorder-7 (Spitzer et al., 2006) and the Patient Health Questionnaire-9 (PHQ-9; Kroenke et al., 2001) were implemented to assess anxiety and depression, respectively. These measures were translated and adapted following the same method as the previously used mental health assessments. For the adaptation of these two measures, we collaborated with the PRECISE-DYAD study, who were assessing maternal health in a different area of The Gambia (for further details see PRECISE-DYAD website) to create a Mandinka version of these measures that could be used in multiple sites and studies. As with the previous measures, these were administered as interviews. The two measures are described in more detail below.

The GAD-7 is a seven-item self-report questionnaire that asks respondents to report how often they experienced a range of anxiety-related symptoms in the last two weeks. Responses are rated on a Likert scale ranging from 0–3 (not at all-nearly every day). The scale generates a total score, which can range from 0–21. There is also one item assessing the severity of symptoms, where respondents are asked to report how much the symptoms that they reported add difficulty to their daily lives on a scale of 0–3 (not at all to very difficult). The GAD-7 has previously been used in Kenya (Nyongesa *et al.*, 2020), where it was reported to have good psychometric properties and the same factor structure as the original English. For our translations, all of the items were deemed to be appropriate for the local setting by the field staff working on the translations.

The PHQ-9 is a nine-item self-report questionnaire that asks respondents to rate how often they experienced a range of depression-related symptoms in the last two weeks. Responses are rated on a Likert scale ranging from 0-3 (not at all-nearly every day). Scores can range from 0-27 and different ranges indicate varying severity of depression (0-4 - none; 5-9 - mild; 10-14 - moderate; 15-19 - moderately severe; and 20-27 severe). Similar to the GAD-7, the PHQ-9 has one item assessing severity of symptoms, where respondents are asked to rate how much difficulty they experience in their daily lives as a result of reported symptoms on a scale of 0-3 (not at all to very difficult). Mothers who report elevated levels of depression were offered an opportunity for referral to the on-site clinic, as was done at previous visits. The field team found most of the items on this scale suitable for the local population. However, as with the EPDS, there were concerns about asking participants to report on suicidal thoughts. Thus, as for the EPDS, this item was changed to asking participants how often they wanted to be alone or isolated.

#### **Impact of COVID-19**

The BRIGHT Kids follow up was conducted during the COVID-19 pandemic and, thus, it was vital to better understand how the pandemic impacted on each family's wellbeing, financial situation and access to healthcare. The questionnaire was adapted from a measure that was being used in a UK study at the time to assess the impact of the pandemic on families with young children (Aydin *et al.*, 2022). Items that were deemed by field staff to be irrelevant or unsuitable for the West Kiang community were removed. The final measure consisted of a series of questions that asked the caregiver to report on how their living and financial situation had changed due to the pandemic, whether they had family members who got sick with COVID-19, their ability to access healthcare, and their general feelings and concerns related to the situation.

#### Nutrition and child growth

Measures of child physical size, namely height, weight, and head circumference, continued to be measured at this visit (see above for details). In addition to this, measures of family food insecurity and the child's dietary diversity were also implemented.

The Dietary Diversity Questionnaire (Swindale & Bilinsky, 2006)) is a caregiver report questionnaire that asks respondents to indicate what the child eats on a typical day, from 12 food groups. There is also an option for "miscellaneous". The *Food Insecurity* questionnaire asks two sets of questions. The first comprises of a single item that asks respondents if anybody in the family has missed a meal due to food shortage in the last seven days. Subsequently, there are seven items asking respondents to indicate any food shortages in the household over the previous month.

### Discussion

## Good scientific practice

We recognise the importance of maximising outputs from the data collected in the BRIGHT project, both by serving the participants and communities that have agreed to partake in this research and the wider scientific community by providing access to the collected data for further analysis. Access to any data collected during or generated by the BRIGHT project is fully audited, and to ensure data security, is overseen by the data management team in the UK and The Gambia. While data sharing is critically important to maximising the benefit of research, we must also consider the need to protect the confidentiality of this sensitive group (particularly the infants within the mother-infant dyads, who as minors do not consent for themselves). Furthermore, to generate maximum value from this dataset we must link data points together (i.e. NIRS/ EEG data with outcome data or contextual factor data). Due to the nature of the data being collected (i.e. collected from a specific geographical location, longitudinal dataset of several datapoints) the majority of the data cannot be fully de-identified under the guidance included in the European General Data Protection Regulation (GDPR). Furthermore, some BRIGHT measures include photo, audio and video material, and therefore is inherently identifiable and requires even stricter governance. In line with other EU based studies of this nature (i.e. EUROSIBS, (Jones et al., 2019)), currently, access to the data sets for the wider scientific community is governed by the data management team to ensure that users comply with all relevant data

protection laws and have appropriate ethical permissions. Access operates via a project approval form, which is reviewed by a committee consisting of representatives from each cohort site, and implements relevant data sharing agreements. Collaborations are encouraged, and projects are evaluated primarily on their consistency with the ethical principles and aims of the project that the families signed up to when partaking in this study. All planned analyses (both internal to the BRIGHT team and external) are pre-specified either on an internal database monitored by our management committee or via web-based pre-registration platforms. These procedures continue to be evaluated annually and updated to optimise the BRIGHT Project's value to the scientific community and public priorities.

Dissemination of key findings has, and will continue, to take place, through presentation of findings by the BRIGHT research team at national and international conferences, internal reports and peer reviewed publications hosted on our website, and through direct interaction with key stakeholders and public engagement events.

#### Lessons learned

During the life course of the BRIGHT Project (and the pilot phases that preceded this) we have encountered many challenges and opportunities that provide us with important guidance for future work. Firstly, the importance for harmonisation of goals, sense of purpose and motivation across the research team and study sites should not be underestimated. Only through our dedicated research team, which includes the 46 authors of this paper, and countless more research midwives, clinical staff, lab assistants, village community assistants, drivers, mechanics, (even a hairdresser for the children's hair) and research centre staff have we managed to succeed in achieving what we have thus far in the BRIGHT Project. In particular in The Gambia, staff coordination was critical while we managed the study workload across the time course of the study with the research team working on rolling interleaved working hours so that the participants could be tested seven days a week during the peak period of longitudinal timepoints. Furthermore, a strength of our study came in the establishment of rigorous SOPs - adapted for each site, age point and measure obtained - and overseen by the quality control and data handling procedures and regular meetings outlined above. The pilot phase of our project enabled us to identify areas where sites differed and discuss with the research team at each site how to adapt best to the contextual factors that were relevant. An important area was to operationalise our management of infant behaviour to ensure successful collection of neuroimaging and behavioural infant measures. We broadly harmonised these approaches and used a series of sequential "attention getters" within task, and calming strategies (such as a snack, break or comforting social strategies) to maintain successful data collection, with levels of attrition being broadly similar across sites and measures. However, there were some differences noted across sites. In The Gambia for example, the climate required us to more carefully monitor environmental conditions, a significant rise in temperature could lead the infant to become drowsy

and fall asleep during the session at a more frequent rate at some age points relative to the UK. In the UK, in contrast, we found that age related changes in the behaviour of individuals led to some losing attention more frequently and moving around or leaving their parent's lap at older ages relative to younger. While in The Gambia this age-related behavioural shift was less evident. It is possible that some of these site differences exist due to fluctuations in the testing practice of the research team, or differences in caregiving approaches or infant motivational behaviours that led to differences in performance during the sessions, or differences in other contextual factors of the sample at each site (for example the proportion of infants and families having a stressful or long journey to the research site). It is hard to tease apart these differences systematically, but our approach to data analysis will include provision for these contextual factors. It should also be noted that heterogeneity can be a strength, and individual variance within sites as well as across sites can be used to further our understanding of infant and early childhood development. As noted by the EUROSIBS consortium, "multisite studies need to consider balancing the resources put into standardisation with the desirability of testing whether metrics are robust across natural variation between sites" (Jones et al., 2019).

Building on this, a strength of our project comes in our approach to incorporating contextual information in the design, development and adaptation of measures to different contexts. We were faced with a need to adapt measures to be broadly applicable across different cultural and linguistic contexts, while also ensuring we designed site specific measures when appropriate. While many of these measures had been used extensively in the UK, and other HICs, not all of these had been used across all the age points of the BRIGHT protocol, so piloting work focused on ensuring that the questionnaires and paradigms were relevant across the developmental time frame that we are studying at both sites. Furthermore, in the Gambia, our approach was to ensure Gambian research staff co-produced the necessary adaptation of measures and assessed the appropriateness of tools in a context that they had not been used previously to take advantage of the local expertise at each site. Furthermore, staff in The Gambia are also involved in data processing and quality control to ensure that, when possible, our training strategies incorporate not just data collection knowledge transfer but also steps towards transferring knowledge on data analysis and dissemination. The latter is certainly an area for future development, as this area of research (developmental psychology/neuroscience) is relatively new to the country and so capacity building is a key priority for our group. While we strive to be sensitive to cultural context, a limitation of some of the measures used in this project is that they are drawn from a pool of existing developmental measures of cognitive outcomes which have largely been derived from tasks developed for high-income settings. While we went to lengths to adapt these measures, we cannot fully eliminate possibility of cultural bias. Equally, these measures likely overlook some skills that are relevant in a rural, farming community as many measures have been designed in urban contexts, therefore we cannot be certain that we are measuring all of the crucial predictors of SES (Hermida *et al.*, 2019). For example, some studies have defined preschool attendance and access to services as a better predictor of cognitive outcomes than income. In focussing on poverty initially, we also overlook many of the strengths of this community and culture, which are highly relevant. While we strive to understand potential protective factors in this environment, we may not have fully captured the complex family dynamics that exist, and will focus on furthering our understanding in future work.

A final strength of the BRIGHT project comes from the diversity of measures incorporated in the cohort design and the density of time points. The spacing of study visits and the range of cognitive, brain and contextual measures employed means that we are able to more accurately pinpoint when certain skills emerge and when they start to become relevant for later outcomes. Likewise, we are also able to investigate whether a particular marker is relevant throughout childhood or only during a specific age/time in development.

#### Future efforts

The ultimate objective of the next phase of the BRIGHT project is the identification and validation of a marker, or "fingerprint" combination of brain function markers, that predict the contribution of exposure phenotypes (i.e., undernutrition / caregiving context) to the substantive variation in developmental outcomes seen in infants born into a low-income setting, such as rural Gambia. Innovative modelling and analytic approaches will be applied to allow us to address the following aims: (1) To employ feature extraction and time varying parameter modelling to identify the most reliable "neural fingerprinting" biomarkers (from fNIRS and EEG) of longitudinal developmental cortical specialization; (2) To create a profile of metrics across developmental outcomes in later childhood (executive function, language, general cognitive development, adaptive skills) at the group level and identify latent classes that sub-group individuals into developmental profiles of pre-academic skills; (3) To expand our predictive models of longitudinal fNIRS and EEG trajectories of early developing brain networks (0-2 years) to determine which age points and "fingerprint" of biomarkers are the most consistent and reliable predictors of developmental outcomes of language, pre-academic skills and functional brain specialization in later childhood (3-5 years); (4) To model exposure phenotypes and establish health, social and environmental risk and resilience determinants of developmental trajectories from birth to preschool which associate with early biomarkers of development, to identify primary targets of intervention.

## Conclusions

The BRIGHT project is a comprehensive, and multi-method study of development, during the first two years of life in the UK and the first five years in The Gambia. The combination of neuroimaging, behavioural, parent-report measures, and biological samples provides a unique opportunity to study a variety of context associated moderators, which may, or may not be related to poverty, as well as the mechanistic processes underlying associations between markers and outcomes. On one hand, we hope that our work will be an asset to global health research, where the study of neurocognitive development in early infancy, particularly with the use of neuroimaging tools, is still emergent. On the other hand, this work has broader value for developmental research in general. Due to the logistical and financial constraints of longitudinal research, our project is among the few to assess early development, across a large number of study visits, and that incorporates such a variety of methods. Thus, the generated results will enable us to identify critical windows for developmental vulnerability and act as rationale to guide future interventions which aim to protect and enrich the developing brain within context associated risk contexts. We propose that our project provides a roadmap for other researchers interested in conducting studies of neurocognitive development in LMICs with similar contextual factors.

### **Data availability**

Underlying data No data are associated with this article.

### Extended data

We recognise the importance of maximising outputs from the data collected in the BRIGHT project, both by serving the participants and communities that have agreed to partake in this research and the wider scientific community by providing access to the collected data for further analysis. Access to any data collected during or generated by the BRIGHT project is fully audited, and to ensure data security, is overseen by the data management team in the UK and The Gambia. While data sharing is critically important to maximising the benefit of research, we must also consider the need to protect the confidentiality of this sensitive group (particularly the infants within the motherinfant dyads, who as minors do not consent for themselves). Furthermore, to generate maximum value from this dataset we must link data points together (i.e. NIRS/EEG data with outcome data or contextual factor data). Due to the nature of the data being collected (i.e. collected from a specific geographical location, longitudinal dataset of several datapoints) the majority of the data cannot be fully de-identified under the guidance included in the European General Data Protection Regulation (GDPR).

The data used to support this study are stored in the Brain Imaging for Global Health Data Repository. The conditions of our ethics approval do not allow public archiving of <u>pseudonymised</u> study data. The data cannot be fully anonymized due to the nature of combined sources of information, such as neuroimaging, sociodemographic, geographic and health measures, making it possible to attribute data to specific individuals, and hence, falling under personal information, the release of which would not be compliant with GDPR guide-lines unless additional participant consent forms are completed. Our data sharing procedures were created in consultation with stakeholders and external consultation (Begum-Ali *et al.*, 2023).

Collaborations are encouraged, and projects are evaluated primarily on their consistency with the ethical principles and aims of the project that the families signed up to when partaking in this study. All planned analyses (both internal to the BRIGHT team and external) are pre-specified either on an internal database monitored by our management committee or via web-based pre-registration platforms. These procedures continue to be evaluated annually and updated to optimise the BRIGHT Project's value to the scientific community and public priorities. To access the data, interested readers should contact the BRIGHT coordinator on the Contact page of our website. Access will be granted to named individuals following ethical procedures governing the reuse of sensitive data. Specifically, requestors must pre-register their proposal, and clearly explain the purpose of the analysis so as to ensure that the purpose and nature of the research is consistent with that to which participating families originally consented. Additionally, requestors must complete and sign a data sharing agreement to ensure data is stored securely. Approved projects would need to adhere to the BRIGHT project's policies on Ethics, Data Sharing, Authorship and Publication.

Legal copyright restrictions do not permit us to publicly archive the full set of behavioural tests and task paradigms used in this experiment. Readers seeking access to these tests are advised to contact the lead author or the reference list.

No part of the study procedures or analysis plans was preregistered prior to the research being conducted.

Contributor Role	Role Definition	Authors
Conceptualization	Ideas; formulation or evolution of overarching research goals and aims.	Main Authors: SLF, BM, SEM, CEE BRIGHT Study Team: TA, MDH, AP
Data Curation	Management activities to annotate (produce metadata), scrub data and maintain research data (including software code, where it is necessary for interpreting the data itself) for initial use and later reuse.	Main Authors: BM, SMC, LK, AB, MCL, GG, LM, MP, CB, EM, ET BRIGHT Study Team:
Formal Analysis	Application of statistical, mathematical, computational, or other formal techniques to analyze or synthesize study data.	Main Authors: SLF, BM, SMC, LK, AB, MCL, GG, LM, CB, MS, ON BRIGHT Study Team:

Contributor Role	Role Definition	Authors
Funding Acquisition	Acquisition of the financial support for the project leading to this publication.	Main Authors: SLF, BM, LK, SEM, CEE BRIGHT Study Team: TA, MDH, AP
Investigation	Conducting a research and investigation process, specifically performing the experiments, or data/evidence collection.	Main Authors: SMC, BM, LK, SLF, AB, MCL, GG, TF, EM, FN, ON, MP, MR, FS, MS, ET BRIGHT Study Team: LA, ABa, CBT, SB, DC, MC, YD, NH, ED, SDa, SD, AJ, SJ, BJ, MJ, OK, KK, JL, LS, LSt
Methodology	Development or design of methodology; creation of models.	Main Authors: SLF, BM, LK, AB, MCL, LM, SEM, MR
		BRIGHT Study Team: CBT, MDH, JL
Project Administration	Management and coordination responsibility for the research activity planning and execution.	Main Authors: SLF, BM, SMC, MCL, EM, MP, MR, ET, SEM, CEE
		BRIGHT Study Team: LA, TA, SB, NH, SD, PN
Resources	Provision of study materials, reagents, materials, patients,	Main Authors: SLF, CEE, SM,
	laboratory samples, animals, instrumentation, computing resources, or other analysis tools.	BRIGHT Study Team: TA, EC, AF
Software	Programming, software development; designing computer programs; implementation of the computer code and supporting	Main Authors: AB, LM, CB, BM, SMC, LK
	algorithms; testing of existing code components.	BRIGHT Study Team:
Supervision	Oversight and leadership responsibility for the research activity planning and execution, including mentorship external to the core	Main Authors: SLF, BM, SMC, LK, AB, CB, MCL, EM, LM, MR, SEM, CEE
	team.	BRIGHT Study Team: LA, CBT, SB, MDH, SD
Validation	Verification, whether as a part of the activity or separate, of the overall replication/reproducibility of results/experiments and other	Main Authors: SLF, AB, BM, MCL, CB, SCB,
	research outputs.	BRIGHT Study Team:
Visualization	Preparation, creation and/or presentation of the published work,	Main Authors: SLF, BM, SMC, LK
	specifically visualization/data presentation.	BRIGHT Study Team:
Writing – Original Draft	Creation and/or presentation of the published work, specifically	Main Authors: SLF, BM, SMC, LK
Preparation	writing the initial draft (including substantive translation).	BRIGHT Study Team:
Writing – Review and Editing	Preparation, creation and/or presentation of the published work by those from the original research group, specifically critical review, commentary or revision – including pre- or post-publication stages.	Main Authors: SLF, BM, SMC, LK, AB, CB, MCL, GG, TJ, EM, LM, FN, ON, MP, MR, FS, MS, ET, SEM, CEE
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#### References

Alam S: Majority World: Challenging the West's Rhetoric of Democracy. Amerasia Journal. 2008; 34(1): 87–98. **Publisher Full Text** 

Alarção FSP, Shephard E, Fatori D, et al.: Promoting mother-infant relationships and underlying neural correlates: Results from a randomized controlled trial of a home-visiting program for adolescent mothers in Brazil. Dev Sci. 2021; 24(6): e13113. PubMed Abstract | Publisher Full Text | Free Full Text

Alderman H, Behrman JR, Grantham-McGregor S, et al.: Economic perspectives **on integrating early child stimulation with nutritional interventions.** *Ann N Y Acad Sci.* 2014; **1308**(1): 129–138.

#### PubMed Abstract | Publisher Full Text

Avala A Christensson K Christensson F et al: Newborn infants who received skin-to-skin contact with fathers after Caesarean sections showed stable physiological patterns. Acta Paediatr. 2021; 110(5): 1461-1467 PubMed Abstract | Publisher Full Text | Free Full Text

Aydin E, Weiss SM, Glasgow KA, et al.: COVID-19 in the context of pregnancy, infancy and parenting (CoCoPIP) study: Protocol for a longitudinal study of parental mental health, social interactions, physical growth and cognitive development of infants during the pandemic. BMJ Open. 2022; 12(6): e053800.

#### PubMed Abstract | Publisher Full Text | Free Full Text

Barch D, Pagliaccio D, Belden A, et al.: Effect of Hippocampal and Amygdala Connectivity on the Relationship Between Preschool Poverty and School-Age Depression. Am J Psychiatry. 2016; **173**(6): 625–634. PubMed Abstract | Publisher Full Text | Free Full Text

Bartram SC: Early infant social behaviour and maternal perinatal depression in Kiang West, The Gambia. [Phd, University of Warwick], 2018. **Reference Source** 

Begum-Ali JB, Holman R, Goodwin A, et al.: Parent attitudes towards data sharing in developmental science. PsyArXiv, 2023. Publisher Full Text

Begus K, Lloyd-Fox S, Halliday D, et al.: Using fNIRS to Study Working Memory PubMed Abstract | Publisher Full Text

Bennett IM, Schott W, Krutikova S, et al.: Maternal mental health, and child growth and development, in four low-income and middle-income countries. J Epidemiol Community Health. 2016; **70**(2): 168–173. PubMed Abstract | Publisher Full Text | Free Full Text

Berens AE, Jensen SKG, Nelson CA 3rd: Biological embedding of childhood adversity: From physiological mechanisms to clinical implications. BMC Med. 2017; 15(1): 135

PubMed Abstract | Publisher Full Text | Free Full Text

Berkes J, Raikes A, Bouguen A, et al.: Joint roles of parenting and nutritional status for child development: Evidence from rural Cambodia. Dev Sci. 2019; 22(5): e12874. PubMed Abstract | Publisher Full Text

Blasi A, Lloyd-Fox S, Johnson MH, et al.: Test-retest reliability of functional near infrared spectroscopy in infants. *Neurophotonics*. 2014; 1(2): 025005. PubMed Abstract | Publisher Full Text | Free Full Text

Blasi A, Lloyd-Fox S, Katus L, et al.: FNIRS for Tracking Brain Development in the Context of Global Health Projects. Photonics. 2019; 6(3): 89. PubMed Abstract | Publisher Full Text | Free Full Text

Blimpo M, Carneiro P, Jervis P, et al.: Improving Access and Quality in Early Childhood Development Programs: Experimental Evidence from the Gambia. SSRN Electronic Journal. 2019.

#### **Publisher Full Text**

Bozicevic L, De Pascalis L, Schuitmaker N, et al.: Longitudinal Association between Child Emotion Regulation and Aggression, and the Role of Parenting: A Comparison of Three Cultures. Psychopathology. 2016; 49(4): 228-235

PubMed Abstract | Publisher Full Text | Free Full Text

Brazelton TB, Koslowski B, Tronick E: Neonatal behavior among urban Zambians and Americans. J Am Acad Child Psychiatry. 1976; 15(1): 97. PubMed Abstract | Publisher Full Text

Brazelton TB, Nugent JK: Neonatal behavioral assessment scale. Cambridge University Press, 1995.

Brotherton H, Daly M, Johm P, et al.: "We All Join Hands": Perceptions of the Kangaroo Method Among Female Relatives of Newborns in The Gambia. Qual Health Res. 2021; **31**(4): 665–676.

PubMed Abstract | Publisher Full Text | Free Full Text

Busch T, Sangen A, Vanpoucke F, et al.: Correlation and agreement between Language ENvironment Analysis (lena™) and manual transcription for Dutch natural language recordings. Behav Res Methods. 2018; 50(5): 1921-1932

PubMed Abstract | Publisher Full Text

Caldwell BM, Bradley RH: Home Observation for Measurement of the Environment (HOME)-revised edition. University of Arkansas, Little Rock, 1984

#### **Reference Source**

Cambridge County Council: Cambridge City annual demographic and socioeconomic report. 2011.

**Reference Source** 

Canault M, Le Normand MT, Foudil S, *et al*.: **Reliability of the Language** ENvironment Analysis system (LENA<sup>™</sup>) in European French. *Behav Res* Methods. 2016; 48(3): 1109-1124 PubMed Abstract | Publisher Full Text

Cattan S, Conti G, Farquharson C, et al.: The Health Effects of Universal Early Childhood Interventions: Evidence from Sure Start.2021; 82. **Reference Source** 

Cavallera V, Lancaster G, Gladstone M, et al.: Protocol for validation of the Global Scales for Early Development (GSED) for children under 3 years of age in seven countries. *BMJ Open.* 2023; **13**(1): e062562. PubMed Abstract | Publisher Full Text | Free Full Text

#### CEIC data: CEICdata.Com. 2018. Reference Source

Cho A: Economics Nobel honors trio taking an experimental approach to fighting poverty. Science. 2019. **Publisher Full Text** 

Cohen S, Kamarck T, Mermelstein R: A GLOBAL MEASURE OF PERCEIVED STRESS. J Health Soc Behav. 1983; 24(4): 385-96. PubMed Abstract

Coleman R, Morison L, Paine K, et al.: Women's reproductive health and depression: a community survey in the Gambia, West Africa. Soc Psychiatry Psychiatr Epidemiol. 2006; 41(9): 720–727.

#### PubMed Abstract | Publisher Full Text

Collins-Jones LH, Cooper RJ, Bulgarelli C, *et al.*: **Longitudinal infant fNIRS channel-space analyses are robust to variability parameters at the** group-level: An image reconstruction investigation. NeuroImage. 2021; 237: 118068. PubMed Abstract | Publisher Full Text | Free Full Text

Cox JL, Holden JM, Sagovsky R: Detection of Postnatal Depression. Development of the 10-item Edinburgh Postnatal Depression Scale. Br J Psychiatry. 1987; **150**(6): 782–786. PubMed Abstract | Publisher Full Text

Development Initiatives: 2018 Global Nutrition Report: Shining a light to spur action on nutrition. Bristol. 2018. **Reference Source** 

Draper CE, Barnett LM, Cook CJ, *et al.*: **Publishing child development research from around the world: An unfair playing field resulting in most of the** world's child population under-represented in research. Infant Child Dev. 2022. **Publisher Full Text** 

Draper CE, Klingberg S, Wrottesley SV, et al.: Interventions to promote development in the next 1000 days: A mapping review. Child Care Health Dev. 2023; 49(4): 617-629.

#### PubMed Abstract | Publisher Full Text

Dzhelyova M, Jacques C, Dormal G, et al.: High test-retest reliability of a neural index of rapid automatic discrimination of unfamiliar individual faces. Vis Cogn. 2019; 27(2): 127-141. **Publisher Full Text** 

East P, Lozoff B, Blanco E, et al.: Infant Iron Deficiency, Child Affect, and Maternal Unresponsiveness: Testing the Long-Term Effects of Functional Isolation. *Dev Psychol.* 2017; **53**(12): 2233–2244.

PubMed Abstract | Publisher Full Text | Free Full Text

Elsabbagh M, Fernandes J, Webb SJ, et al.: Disengagement of visual attention in infancy is associated with emerging autism in toddlerhood. Biol Psychiatry. 2013a; 74(3): 189-194.

PubMed Abstract | Publisher Full Text | Free Full Text

Elsabbagh M, Gliga T, Pickles A, et al.: The development of face orienting mechanisms in infants at-risk for autism. Behav Brain Res. 2013b; 251: 147-154

#### PubMed Abstract | Publisher Full Text | Free Full Text

Elsabbagh M, Volein A, Holmboe K, et al.: Visual orienting in the early broader autism phenotype: Disengagement and facilitation. J Child Psychol Psychiatry. 2009; 50(5): 637–642.

PubMed Abstract | Publisher Full Text | Free Full Text

Farah MI: The Neuroscience of Socioeconomic Status: Correlates. Causes. and Consequences. Neuron. 2017; 96(1): 56-71. PubMed Abstract | Publisher Full Text

Farah MJ, Shera DM, Savage JH, et al.: Childhood poverty: Specific associations with neurocognitive development. Brain Res. 2006; 1110(1): 166-174

PubMed Abstract | Publisher Full Text

Fenson L, Marchman VA, Thal DJ, et al.: MacArthur-Bates Communicative Development Inventories: User's guide and technical manual (2nd ed.). Brookes, 2007.

#### Reference Source

Fernald A, Zangl R, Portillo AL, *et al.*: Looking while listening: Using eye movements to monitor spoken language comprehension by infants and young children. In: *Developmental Psycholinguistics: On-line Methods in Children's* Language Processing. John Benjamins Publishing Company, 2008; 97–135. **Publisher Full Text** 

Fishell AK, Arbeláez AM, Valdés CP, et al.: Portable, field-based neuroimaging using high-density diffuse optical tomography. NeuroImage. 2020; 215: 116541.

PubMed Abstract | Publisher Full Text

Fitt E, Cole D, Ziauddeen N, et al.: DINO (Diet In Nutrients Out) - an integrated dietary assessment system. Public Health Nutr. 2015; 18(2): 234–241. PubMed Abstract | Publisher Full Text | Free Full Text

Forssman L, Ashorn P, Ashorn U, et al.: Eye-tracking-based assessment of cognitive function in low-resource settings. Arch Dis Child. 2017; 102(4): 301-302

PubMed Abstract | Publisher Full Text | Free Full Text

Ganek HV, Eriks-Brophy A: A Concise Protocol for the Validation of Language ENvironment Analysis (LENA) Conversational Turn Counts in Vietnamese. Comm Disord Q. 2018; 39(2): 371-380.

Publisher Full Text

Gervain J, Minagawa Y, Emberson L, et al.: Using functional near-infrared spectroscopy to study the early developing brain: Future directions and new challenges. *Neurophotonics*. 2023; **10**(2): 023519. PubMed Abstract | Publisher Full Text | Free Full Text

Gilkerson J, Richards JA: **The LENA Natural Language Study.** (LENA Technical Report LTR-02-2). LENA Foundation, 2008; 1–26. **Reference Source** 

Giovanelli A, Reynolds AJ, Mondi CF, et al.: Adverse Childhood Experiences and Adult Well-Being in a Low-income, Urban Cohort. Pediatrics. 2016; 137(4): e20154016.

PubMed Abstract | Publisher Full Text | Free Full Text

Hackman DA. Farah MI: Socioeconomic status and the developing brain. Trends Cogn Sci. 2009; 13(2): 65-73

PubMed Abstract | Publisher Full Text | Free Full Text

Hair NL, Hanson JL, Wolfe BL, et al.: Association of Child Poverty, Brain Development, and Academic Achievement. JAMA Pediatr. 2015; 169(9): 822-829

PubMed Abstract | Publisher Full Text | Free Full Text

Hanlon C, Medhin G, Alem A, et al.: Detecting perinatal common mental disorders in Ethiopia: Validation of the self-reporting questionnaire and Edinburgh Postnatal Depression Scale. J Affect Disord. 2008; **108**(3): 251–262. PubMed Abstract | Publisher Full Text

Hennig BJ, Unger SA, Dondeh BL, et al.: Cohort Profile: The Kiang West Longitudinal Population Study (KWLPS)-a platform for integrated research and health care provision in rural Gambia. Int J Epidemiol. 2017; 46(2): e13. PubMed Abstract | Publisher Full Text | Free Full Text

Henrich J, Heine SJ, Norenzayan A: Beyond WEIRD: Towards a Broad-based Behavioral Science. Behav Brain Sci. 2010; 33(2-3): 111-135. **Publisher Full Text** 

Hensch TK: Critical period mechanisms in developing visual cortex. Curr Top Dev Biol. 2005; 69: 215-237.

PubMed Abstract | Publisher Full Text

Hermida MJ, Shalom DE, Segretin MS, et al.: Risks for Child Cognitive Development in Rural Contexts. Front Psychol. 2019; 9: 2735 PubMed Abstract | Publisher Full Text | Free Full Text

Howard SJ, Cook CJ, Everts L, et al.: Challenging socioeconomic status: A

cross-cultural comparison of early executive function. Dev Sci. 2020; 23(1): e12854

PubMed Abstract | Publisher Full Text

Howard SJ, Melhuish E: An Early Years Toolbox for Assessing Early Executive Function, Language, Self-Regulation, and Social Development: Validity, Reliability, and Preliminary Norms. J Psychoeduc Assess. 2017; 35(3): 255–275. PubMed Abstract | Publisher Full Text | Free Full Text

Hurt H, Betancourt LM: Effect of socioeconomic status disparity on child language and neural outcome: How early is early? Pediatr Res. 2016; 79(1-2): 148-158

#### PubMed Abstract | Publisher Full Text

Isaacs EB: Neuroimaging, a new tool for investigating the effects of early diet on cognitive and brain development. Front Hum Neurosci. 2013; 7: 445. PubMed Abstract | Publisher Full Text | Free Full Text

Isaacs E, Oates J, ILSI Europe a.i.s.b.l: Nutrition and cognition: Assessing cognitive abilities in children and young people. Eur J Nutr. 2008; 47 Suppl 3(3): 4-24.

#### PubMed Abstract | Publisher Full Text

Jasińska KK, Guei S: Neuroimaging Field Methods Using Functional Near Infrared Spectroscopy (NIRS) Neuroimaging to Study Global Child Development: Rural Sub-Saharan Africa. J Vis Exp. 2018; (132): 57165. PubMed Abstract | Publisher Full Text | Free Full Text

Jensen SKG, Berens AE, Nelson CA: Effects of poverty on interacting biological systems underlying child development. Lancet Child Adolesc Health. 2017; 1(3): 225–239. PubMed Abstract | Publisher Full Text

Jensen SKG, Tofail F, Haque R, et al.: Child development in the context of biological and psychosocial hazards among poor families in Bangladesh. PLoS One. 2019; 14(5): e0215304. PubMed Abstract | Publisher Full Text | Free Full Text

Johnson MH: The inhibition of automatic saccades in early infancy. Dev Psychobiol. 1995; **28**(5): 281–291. PubMed Abstract | Publisher Full Text

Jones EJH, Mason L, Begum Ali J, et al.: Eurosibs: Towards robust measurement of infant neurocognitive predictors of autism across Europe. Infant Behav Dev. 2019; 57: 101316.

PubMed Abstract | Publisher Full Text | Free Full Text

Kaldy Z, Kraper C, Carter AS, et al.: Toddlers with Autism Spectrum Disorder are more successful at visual search than typically developing toddlers. Dev Sci. 2011; 14(5): 980-988.

PubMed Abstract | Publisher Full Text | Free Full Text

Kariger P, Frongillo EA, Engle P, et al.: Indicators of Family Care for Development for Use in Multicountry Surveys. J Health Popul Nutr. 2012; 30(4): 472-486

PubMed Abstract | Publisher Full Text | Free Full Text

Katus L, Blasi A, McCann S, et al.: Longitudinal fNIRS and EEG metrics of habituation and novelty detection are correlated in 1-18-month-old infants. Neuroimage. 2023; 274: 120153. PubMed Abstract | Publisher Full Text | Free Full Text

Katus L, Foley S, Murray AL, et al.: Perceived stress during the prenatal period: Assessing measurement invariance of the Perceived Stress Scale (PSS-10) across cultures and birth parity. Arch Womens Ment Health. 2022a; 25(3): 633-640

PubMed Abstract | Publisher Full Text | Free Full Text

Katus L, Hayes NJ, Mason L, et al.: Implementing neuroimaging and eye tracking methods to assess neurocognitive development of young infants in low- and middle-income countries [version 2; peer review: 2 approved]. Gates Open Res. 2019; 3: 1113.

PubMed Abstract | Publisher Full Text | Free Full Text

Katus L, Mason L, Milosavljevic B, et al.: ERP markers are associated with neurodevelopmental outcomes in 1-5 month old infants in rural Africa and the UK. Neuroimage. 2020; 210: 116591.

PubMed Abstract | Publisher Full Text | Free Full Text

Katus L, Milosavljevic B, Rozhko M, et al.: Neural Marker of Habituation at 5 Months of Age Associated with Deferred Imitation Performance at 12 Months: A Longitudinal Study in the UK and The Gambia. Children (Basel). 2022b; 9(7): 988. PubMed Abstract | Publisher Full Text | Free Full Text

Kea P: 'The complexity of an enduring relationship': Gender, generation, and the moral economy of the Gambian Mandinka household. J R Anthropol Inst. 2013: 19(1): 102-119. **Publisher Full Text** 

Kea P: Gambian educational migration, care and the persistence of the domestic moral economy. Child Geogr. 2020; 18(6): 642-653. **Publisher Full Text** 

Kingston D, Tough S: Prenatal and Postnatal Maternal Mental Health and School-Age Child Development: A Systematic Review. Matern Child Health J. 2014; 18(7): 1728-1741.

PubMed Abstract | Publisher Full Text

Kohrt BA, Luitel NP, Acharya P, et al.: Detection of depression in low resource settings: Validation of the Patient Health Questionnaire (PHQ-9) and cultural concepts of distress in Nepal. BMC Psychiatry. 2016; 16(1): 58. PubMed Abstract | Publisher Full Text | Free Full Text

Kroenke K, Spitzer RL, Williams JBW: **The PHQ-9: Validity of a brief depression** severity measure. *J Gen Intern Med.* 2001; **16**(9): 606–613. PubMed Abstract | Publisher Full Text | Free Full Text

Lloyd-Fox S, Begus K, Halliday D, et al.: Cortical specialisation to social stimuli from the first days to the second year of life: A rural Gambian cohort. Dev Cogn Neurosci. 2017; 25: 92–104.

PubMed Abstract | Publisher Full Text | Free Full Text

Lloyd-Fox S, Blasi A, Elwell CE: Illuminating the developing brain: The past, present and future of functional near infrared spectroscopy. Neurosci Biobehav Rev. 2010; **34**(3): 269–284.

PubMed Abstract | Publisher Full Text

Lloyd-Fox S, Blasi A, McCann S, et al.: Habituation and novelty detection fNIRS brain responses in 5- and 8-month-old infants: The Gambia and UK. Dev Sci. 2019; 22(5): e12817.

PubMed Abstract | Publisher Full Text | Free Full Text

Lloyd-Fox S, Papademetriou M, Darboe MK, et al.: Functional near infrared spectroscopy (fNIRS) to assess cognitive function in infants in rural Africa. Sci Rep. 2014; 4: 4740.

PubMed Abstract | Publisher Full Text | Free Full Text

Loth E, Charman T, Mason L, et al.: The EU-AIMS Longitudinal European Autism Project (LEAP): Design and methodologies to identify and validate stratification biomarkers for autism spectrum disorders. Mol Autism. 2017; 8(1):24

#### PubMed Abstract | Publisher Full Text | Free Full Text

Luby J, Belden A, Botteron K, *et al.*: **The Effects of Poverty on Childhood Brain Development: The Mediating Effect of Caregiving and Stressful Life Events.** *JAMA Pediatr.* 2013; **167**(12): 1135–1142.

PubMed Abstract | Publisher Full Text | Free Full Text

Macrae E. Milosavlievic B. Mason L. et al.: Executive functions in infancy: Measurement using a novel tablet task and exploration of longitudinal attentional and cognitive predictors. [Preprint]. PsyArXiv. 2022. **Publisher Full Text** 

Martorell R, Horta BL, Adair LS, et al.: Weight Gain in the First Two Years of Life Is an Important Predictor of Schooling Outcomes in Pooled Analyses from Five Birth Cohorts from Low- and Middle-Income Countries. / Nutr. 2010; 140(2): 348-354

#### PubMed Abstract | Publisher Full Text | Free Full Text

McCoy DC, Peet ED, Ezzati M, et al.: Early Childhood Developmental Status in Low- and Middle-Income Countries: National, Regional, and Global Prevalence Estimates Using Predictive Modeling. PLoS Med. 2016; 13(6): e1002034

#### PubMed Abstract | Publisher Full Text | Free Full Text

Milosavljevic B, Cook CJ, Fadera T, et al.: Executive functioning skills and their environmental predictors among pre-school aged children in South Africa and The Gambia. *Dev Sci.* 2023; e13407.

PubMed Abstract | Publisher Full Text

Milosavljevic B, Vellekoop P, Maris H, et al.: Adaptation of the Mullen Scales of Early Learning for use among infants aged 5- to 24-months in rural Gambia. Dev Sci. 2019; 22(5): e12808

PubMed Abstract | Publisher Full Text | Free Full Text

Moore SE, Cole TJ, Poskitt EME, et al.: Season of birth predicts mortality in rural Gambia. Nature. 1997; 388(6641): 434.

PubMed Abstract | Publisher Full Text

Moriguchi Y: Beyond bias to Western participants, authors, and editors in developmental science. Infant and Child Development. 2022; 31(1). Publisher Full Text

Mullen EM: Mullen scales of early learning (AGS ed.). Circle Pines, MN: American Guidance Service Inc. 1995

Murray AL, Xiao Z, Zhu X, et al.: Psychometric evaluation of an adapted version of the perceived stress scale for ecological momentary assessment research. Stress Health. 2023.

PubMed Abstract | Publisher Full Text

Murray-Kolb LE, Rasmussen ZA, Scharf RJ, et al.: The MAL-ED Cohort Study: Methods and Lessons Learned When Assessing Early Child Development and Caregiving Mediators in Infants and Young Children in 8 Low- and Middle-Income Countries. *Clin Infect Dis.* 2014; **59**(suppl\_ 4): S261–S272. PubMed Abstract | Publisher Full Text | Free Full Text

Nabwera HM, Fulford AJ, Moore SE, et al.: Growth faltering in rural Gambian children after four decades of interventions: A retrospective cohort study. Lancet Glob Health. 2017; 5(2): e208-e216.

PubMed Abstract | Publisher Full Text | Free Full Text

Nabwera HM, Moore SE, Mwangome MK, et al.: The influence of maternal psychosocial circumstances and physical environment on the risk of severe wasting in rural Gambian infants: A mixed methods approach. BMC Public Health. 2018; 18: 109.

## PubMed Abstract | Publisher Full Text | Free Full Text

Nielsen M, Haun D, Kärtner J, et al.: The persistent sampling bias in developmental psychology: A call to action. J Exp Child Psychol. 2017; 162: 31-38

#### PubMed Abstract | Publisher Full Text

Noble KG, Houston SM, Brito NH, et al.: Family income, parental education and brain structure in children and adolescents. Nat Neurosci. 2015; 18(5):

#### 773-778

#### PubMed Abstract | Publisher Full Text | Free Full Text

Noble KG, Houston SM, Kan E, et al.: Neural correlates of socioeconomic status in the developing human brain. Dev Sci. 2012; 15(4): 516-527. PubMed Abstract | Publisher Full Text | Free Full Text

Nyongesa MK, Mwangi P, Koot HM, et al.: The reliability, validity and factorial structure of the Swahili version of the 7-item generalized anxiety disorder scale (GAD-7) among adults living with HIV from Kilifi, Kenya. Ann Gen Psychiatry. 2020; **19**(1): 62.

#### PubMed Abstract | Publisher Full Text | Free Full Text

Pae S, Yoon H, Seol A, et al.: Effects of feedback on parent-child language with infants and toddlers in Korea. First Language. 2016; 36(6): 549–569. **Publisher Full Text** 

Papademetriou MD, Richards J, Correia T, *et al.*: **Cortical Mapping of 3D Optical Topography in Infants.** In: S. Van Huffel, G. Naulaers, A. Caicedo, D. F. Bruley, and D. K. Harrison (Eds.): Oxygen Transport to Tissue XXXV. 2013; 789: 455-461

#### **Publisher Full Text**

Peña ED: Lost in translation: Methodological considerations in crosscultural research. Child Dev. 2007; 78(4): 1255-1264. PubMed Abstract | Publisher Full Text

Perdue KL, Jensen SKG, Kumar S, et al.: Using functional near-infrared spectroscopy to assess social information processing in poor urban Bangladeshi infants and toddlers. Dev Sci. 2019; 22(5): e12839. PubMed Abstract | Publisher Full Text | Free Full Text

Perkins JM, Kim R, Krishna A, et al.: Understanding the association between stunting and child development in low- and middle-income countries: Next steps for research and intervention. Soc Sci Med. 2017; 193: 101–109. PubMed Abstract | Publisher Full Text

Pyykkö J, Ashorn U, Chilora E, et al.: Associations between individual variations in visual attention at 9 months and behavioral competencies at 18 months in rural Malawi. *PLoS One.* 2020; **15**(10): e0239613. PubMed Abstract | Publisher Full Text | Free Full Text

Pyykkö J, Forssman L, Maleta K, et al.: Early development of visual attention in infants in rural Malawi. Dev Sci. 2019; 22(5): e12761. PubMed Abstract | Publisher Full Text

Räikkönen K, Birkás E, Horváth J, et al.: Test-retest reliability of auditory ERP components in healthy 6-year-old children. *Neuroreport*. 2003; 14(16): 2121-2125

PubMed Abstract | Publisher Full Text

Richter L, Black M, Britto P, et al.: Early childhood development: An imperative for action and measurement at scale. BMJ Global Health. 2019; 4(Suppl 4): e001302.

PubMed Abstract | Publisher Full Text | Free Full Text

Rini CK, Dunkel-Schetter C, Wadhwa PD, et al.: Psychological adaptation and birth outcomes: The role of personal resources, stress, and sociocultural context in pregnancy. *Health Psychol.* 1999; **18**(4): 333–345. PubMed Abstract | Publisher Full Text

Roberts SB, Franceschini MA, Krauss A, et al.: A Pilot Randomized Controlled Trial of a New Supplementary Food Designed to Enhance Cognitive Performance during Prevention and Treatment of Malnutrition in Childhood. Curr Dev Nutr. 2017; 1(11): e000885. PubMed Abstract | Publisher Full Text | Free Full Text

Roesch SC, Schetter CD, Woo G, et al.: Modeling the types and timing of stress in pregnancy. ANXIETY STRESS COPIN. 2004; 17(1): 87–102.

**Publisher Full Text** Sabanathan S, Wills B, Gladstone M: Child development assessment tools in low-income and middle-income countries: How can we use them more appropriately? Arch Dis Child 2015; 100(5): 482-8.

PubMed Abstract | Publisher Full Text | Free Full Text

Sadeh A: Evaluating Night Wakings in Sleep-Disturbed Infants: A Methodological Study of Parental Reports and Actigraphy. Sleep. 1996; **19**(10): 757-762.

#### PubMed Abstract | Publisher Full Text

Sadeh A: A Brief Screening Questionnaire for Infant Sleep Problems: Validation and Findings for an Internet Sample. Pediatrics. 2004; 113(6): e570-e577. PubMed Abstract | Publisher Full Text

Scerif G, Karmiloff-Smith A, Campos R, et al.: To look or not to look? Typical and atypical development of oculomotor control. J Cogn Neurosci. 2005; 17(4): 591-604.

#### PubMed Abstract | Publisher Full Text

Sear R, Mace R: Family Matters Kin, Demography and Child Health in a Rural Gambian Population. In: G. Bentley and R. Mace (Eds.), *Substitute* Parents. Berghahn Books, 2009; 50-76. **Publisher Full Text** 

Sethna V, Pote I, Wang S, et al.: Mother-infant interactions and regional brain volumes in infancy: An MRI study. Brain Struct Funct. 2017; 222(5): 2379-2388.

#### PubMed Abstract | Publisher Full Text | Free Full Text

Smith T, Noble M, Noble S, et al.: The English Indices of Deprivation 2015:

Technical report. Department for Communities and Local Government, 2015. **Reference Source** 

Soler-Limón KM, Rivera-González R, Velázquez-Moctezuma J, et al.: Neurobehavior of mexican newborns in conditions of low perinatal risk. Acta Pediátrica de México. 2019; 40(3): 113–121. **Reference Source** 

Sosseh SAL, Barrow A, Lu ZJ: Cultural beliefs, attitudes and perceptions of lactating mothers on exclusive breastfeeding in The Gambia: An ethnographic study. BMC Womens Health. 2023; 23(1): 18. PubMed Abstract | Publisher Full Text | Free Full Text

Spitzer RL, Kroenke K, Williams JBW, et al.: A Brief Measure for Assessing Generalized Anxiety Disorder: The GAD-7. Arch Intern Med. 2006; 166(10): 1092-7

#### PubMed Abstract | Publisher Full Text

Super CM, Harkness S: Culture and the perceived organization of newborn behavior: A comparative study in Kenya and the United States. *New Dir* Child Adolesc Dev. 2020; 2020(172): 11-24. PubMed Abstract | Publisher Full Text

Swindale A, Bilinsky P: Household Dietary Diversity Score (HDDS) for Measurement of Household Food Access: Indicator Guide. FHI 360/FANTA, 2006.

#### **Reference Source**

Tesfaye M, Hanlon C, Wondimagegn D, et al.: Detecting postnatal common mental disorders in Addis Ababa, Ethiopia: Validation of the Edinburgh Postnatal Depression Scale and Kessler Scales. J Affect Disord. 2010; 122(1-2): 102-108

#### PubMed Abstract | Publisher Full Text

Turesky TK, Jensen SKG, Yu X, et al.: The relationship between biological and psychosocial risk factors and resting-state functional connectivity in 2month-old Bangladeshi infants: A feasibility and pilot study. Dev Sci. 2019; 22(5): e12841.

#### PubMed Abstract | Publisher Full Text | Free Full Text

Turesky T, Xie W, Kumar S, et al.: Relating anthropometric indicators to brain structure in 2-month-old Bangladeshi infants growing up in poverty: A pilot study. Neuroimage. 2020; 210: 116540. PubMed Abstract | Publisher Full Text | Free Full Text

Twomey DM, Wrigley C, Ahearne C, et al.: Feasibility of using touch screen technology for early cognitive assessment in children. Arch Dis Child. 2018; 103(9): 853-858

#### PubMed Abstract | Publisher Full Text

Ulrich DA: Test of Gross Motor Development. (3rd ed.). Pro-Ed, 2019. **Reference Source** 

Victora CG, Adair L, Fall C, et al.: Maternal and child undernutrition: Consequences for adult health and human capital. Lancet. 2008; 371(9609): 340-357

PubMed Abstract | Publisher Full Text | Free Full Text

Wass SV: Applying cognitive training to target executive functions during early development. Child Neuropsychol. 2015; 21(2): 150-166. PubMed Abstract | Publisher Full Text | Free Full Text

Watson D, Clark LA, Tellegen A: Development and validation of brief measures of positive and negative affect: The PANAS scales. J Pers Soc Psychol. 1988; 54(6): 1063-1070. PubMed Abstract | Publisher Full Text

Webb SJ, Jones EJH, Merkle K, et al.: Toddlers with elevated autism symptoms show slowed habituation to faces. Child Neuropsychol. 2010; 16(3):

PubMed Abstract | Publisher Full Text | Free Full Text

Weber A. Fernald A. Diop Y: When Cultural Norms Discourage Talking to Babies: Effectiveness of a Parenting Program in Rural Senegal. Child Dev. 2017; 88(5): 1513-1526.

#### PubMed Abstract | Publisher Full Text

Wedderburn CJ, Subramoney S, Yeung S, et al.: Neuroimaging young children and associations with neurocognitive development in a South African birth cohort study. NeuroImage. 2020; 219: 116846.

PubMed Abstract | Publisher Full Text | Free Full Text

Weisleder A, Fernald A: Talking to Children Matters: Early Language Experience Strengthens Processing and Builds Vocabulary. Psychol Sci. 2013; 24(11): 2143-2152

PubMed Abstract | Publisher Full Text | Free Full Text

Weobong B, Ten Asbroek AH, Soremekun S, et al.: Determinants of postnatal depression in rural ghana: Findings from the don population based cohort study. Depress Anxiety. 2015; 32(2): 108–119. PubMed Abstract | Publisher Full Text

Wijeakumar S, Kumar A, Reyes LMD, et al.: Early adversity in rural India impacts the brain networks underlying visual working memory. Dev Sci. 2019: 22(5): e12822

PubMed Abstract | Publisher Full Text | Free Full Text

Worku BN, Abessa TG, Wondafrash Kibebew M, et al.: Effects of home-based play-assisted stimulation on developmental performances of children living in extreme poverty: A randomized single-blind controlled trial. BMC Pediatr. 2018; 18(1): 29.

PubMed Abstract | Publisher Full Text | Free Full Text

World Health Organization: Process of translation and adaptation of instruments. 2013. **Reference Source** 

Xie W. Jensen S. Wade M. et al.: Child Growth Predicts Brain Functional Connectivity and Future Cognitive Outcomes in Urban Bangladeshi Children Exposed to Early Adversities. bioRxiv. 2018. **Publisher Full Text** 

Xie W, Kumar S, Kakon SH, et al.: Chronic inflammation is associated with neural responses to faces in bangladeshi children. NeuroImage. 2019; 202: 116110

#### PubMed Abstract | Publisher Full Text | Free Full Text

Xu D, Richards JA, Gilkerson J, et al.: Automatic Childhood Autism Detection by Vocalization Decomposition with Phone-like Units. Proceedings of the 2Nd Workshop on Child, Computer and Interaction. 2009; 5: 1-7. **Publisher Full Text** 

## **Open Peer Review**

### Current Peer Review Status: 🥇 🗸

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Reviewer Report 27 February 2024

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### Rihui Li 匝

University of Macau, Taipa, Macao

This study protocol introduces the detailed design and implementation of the Brain Imaging for Global Health (BRIGHT) project, which primarily aims to study the longitudinal change of brain function, cognitive development, and growth within the rural Gambian population. The protocol also aims to find out the critical developmental moderators, mediators, and markers of risk and resilience by comparing the Gambian cohort with the UK cohort. I think this is an exciting project that makes use of a variety of brain imaging techniques and behavioral assessments to collect multi-dimensional data. I have several minor suggestions to be considered.

It may be beneficial to list the specifications of the EEG, fNIRS system as well as the eye-tracking device used in this project, such as channel number, sampling rate, and whether short channels are used. Such information could help to justify the reliability and feasibility of the protocol.

A few neuroimaging tasks (e.g., working memory, habituation) are used in this project, please provide more information about these tasks.

Psychiatric disorders, such as Autism or anxiety disorder, may be potential factors affecting brain and cognitive development. I wonder whether the research team is capable of screening and excluding those participants.

Since the Phase 1 of BRIGHT is completed 3-4 years ago (ran from 2015-2020), I recommend the authors add a section or/and table to briefly describe relevant publications and main findings in order to highlight the value of the BRIGHT project.

Is the rationale for, and objectives of, the study clearly described?

Yes

Is the study design appropriate for the research question?

Yes

### Are sufficient details of the methods provided to allow replication by others?

Yes

# Are the datasets clearly presented in a useable and accessible format? $\ensuremath{\mathsf{Yes}}$

Competing Interests: No competing interests were disclosed.

*Reviewer Expertise:* neuroimaging, neuropsychiatry, social interaction, biomedical engineering

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Author Response 12 May 2024

#### Sarah Lloyd-Fox

We thank you for your helpful and constructive comments. We have addressed all comments in our revision and believe this has strengthened the manuscript. Below we provide detailed responses to the individual points raised (your points in bold, ours in standard or italicised.

This study protocol introduces the detailed design and implementation of the Brain Imaging for Global Health (BRIGHT) project, which primarily aims to study the longitudinal change of brain function, cognitive development, and growth within the rural Gambian population. The protocol also aims to find out the critical developmental moderators, mediators, and markers of risk and resilience by comparing the Gambian cohort with the UK cohort. I think this is an exciting project that makes use of a variety of brain imaging techniques and behavioral assessments to collect multi-dimensional data. I have several minor suggestions to be considered.

1. It may be beneficial to list the specifications of the EEG, fNIRS system as well as the eye-tracking device used in this project, such as channel number, sampling rate, and whether short channels are used. Such information could help to justify the reliability and feasibility of the protocol.

**Response:** We agree that these details would help contextualise our protocol and findings further. As outlined below, we have added additional detail to the relevant sections: fNIRS:

This multi-domain battery was designed to interrogate whether global health risk factors impact on development to result in global/cross-domain differences in brain activity or localized/domainspecific differences or altered function. Data were collected using the NTS optical imaging system (Gowerlabs Ltd. London, Everdell et al., 2005), which emits near-infrared light at the 780nm and 850nm wavelengths with a sampling rate of 10Hz. The total array consisted of 18 channels at 1 month, 34 channels at 5, 8, 12, 18 and 24 months. The same 34-channel layout was used in BRIGHT kids. Channel separation was kept constant at 20mm across age points. Co-registration information from age and head size matched MRIs was used to adjust the placement guidelines of the headgear to ensure the channels were aligned over the frontal and temporal regions at each age point.

EEG:

Due to the nature of the sounds (pure tones, bursts of white noise etc.), no adaptations had to be undertaken allowing for identical protocols at both project sites. Infants were presented with auditory stimuli for approximately 15 minutes while asleep (at 1 month) or awake (at 5 and 18 months): during the latter an experimenter quietly entertained the infant with bubbles or silent toys to maintain calm attention during the task. We used a low-density (8 electrodes) gel based Neurolectrics Enobio EEG system with passive electrodes with a sampling rate of 500Hz. Eye tracking:

A series of pilot studies were then completed in both the UK and The Gambia to identify population- and age- appropriate word-picture stimulus pairs. Data were collected using a Tobii TX-300 (Tobii AB, Sweden) remote eye tracker, set to a sampling rate of 60Hz.

# 2. A few neuroimaging tasks (e.g., working memory, habituation) are used in this project, please provide more information about these tasks.

**Response:** We agree that such detail would be helpful to others wishing to implement similar task protocols. To not overload the manuscript with detailed methodological descriptions for all components of this extensive protocol, we have opted to more clearly signpost prior work (both our own and other groups') to provide easily accessible links to the specific tasks used.

- EEG: (for a description of the EEG task and full protocol see Katus et al., 2020)
- fNIRS: The fNIRS paradigms implemented in the BRIGHT project assessed a range of cognitive functions and domains, namely social cognition (see Lloyd-Fox et al., 2017 for details of task), habituation and novelty detection (see Lloyd-Fox et al., 2019), working memory (using an adapted version of the task described in Begus et al., (2016)), deferred imitation (see Katus et al., (2022) for task description) and functional connectivity (see Bulgarelli et al., (2024)).
- Eye-tracking: The tasks used assessed cognitive control (Wass et al., 2011), habituation (Katus et al., 2019), attentional disengagement using the Gap/Overlap task (Holmboe et al., 2010), non-social contingency (Delligianni, 2011), preference for faces (Gliga et al., 2009), and spontaneous looking (de Urabain et al., 2017).

# 3 .Psychiatric disorders, such as Autism or anxiety disorder, may be potential factors affecting brain and cognitive development. I wonder whether the research team is capable of screening and excluding those participants.

**Response:** We agree that the conditions highlighted by the reviewer might affect the responses measured in this project. Infants who had suspected developmental delay or disability were referred for medical review and, as outlined in our exclusion criteria, those diagnosed with a developmental disability were excluded from the study. However, because autism and anxiety are typically not diagnosed until later in childhood, and due to the lack of appropriate medical expertise and follow up care, we did not screen for these conditions. The following is included in the main text:

Infants with suspected developmental delay or disability were referred for medical review and, as per the exclusion criteria, infants/ children who were diagnosed with developmental disabilities (e.g. Down's Syndrome and cerebral palsy) were excluded from the study. Previously collected data relating to these participants was removed from the dataset. Specific screening for further psychiatric disorders including autism and anxiety disorder was not completed as (i) these

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conditions are usually diagnosed later in childhood and (ii) appropriate necessary medical expertise and follow up care was not available. Further screening will be taken into consideration if the cohort is followed up in the future.

4. Since the Phase 1 of BRIGHT is completed 3-4 years ago (ran from 2015-2020), I recommend the authors add a section or/and table to briefly describe relevant publications and main findings in order to highlight the value of the BRIGHT project.

**Response:** We added this information in the Table below, to provide a more accessible summary of key findings obtained to date:

We have multiple publications that describe both the methods development and pilot testing that was conducted prior to the start of the project, as well as publications from the data collected in BRIGHT (please see Table 6 for list and summary of BRIGHT publications to date).

Table 6 overview for your benefit in this review reply (is formatted properly in the resubmission file).

Info given in the following order:

Citation Title Data examined Main findings

Lloyd-Fox et al., 2019

Habituation and novelty detection fNIRS brain responses in 5- and 8-month-old infants: The Gambia and UK.

fNIRS habituation and novelty detection at 5 and 8 months in The Gambia and the UK Differential profiles of habituation and novelty detection across the two cohorts.

Katus et al., 2019

Implementing neuroimaging and eye tracking methods to assess neurocognitive development of young infants in low- and middle-income countries NA

fNIRS, EEG and eye tracking show robustness for use in a less well-controlled lab setting in The Gambia. We address common challenges related to implementing a multi-method protocol longitudinally.

Blasi et al., 2019

fNIRS for Tracking Brain Development in the Context of Global Health Projects NA

Review of the use of fNIRS in the context of global health studies. We discuss the implementation of fNIRS studies in LMICs with a particular emphasis on the Brain Imaging for Global Health (BRIGHT) project, and we consider its potential in this emerging field.

Katus et al., 2020

ERP markers predict neurodevelopmental outcomes in young infants in rural Africa and the UK.

EEG auditory oddball data at 1 and 5 months, MSEL data at 5 months in UK and Gambia UK and Gambian cohort both show immature novelty response at 1 month of age. A novelty

response emerges at group level in UK, but not in the Gambian cohort. The emergence of a novelty response is associated with concurrently MSEL scores at 5 months of age.

Collins-Jones et al., 2021

Longitudinal infant fNIRS channel-space analyses are robust to variability parameters at the group-level: An image reconstruction investigation.

fNIRS Social paradigm at 5, 8 and 12 months in The Gambia

Traditionally, fNIRS analysis is done in the channel space, where data from equivalent channels across participants is combined, assuming that head size and source and detector positions are constant across participants. We conclude that channel -space analysis of longitudinal fNIRS data is robust to assumptions about head size and array position given the variability in these parameters in our dataset.

#### Katus et al., 2022

Neural Marker of Habituation at 5 Months of Age Associated with Deferred Imitation Performance at 12 Months: A Longitudinal Study in the UK and The Gambia.

EEG auditory oddball data at 1 and 5 months, deferred imitation data at 8 and 12 months of age in UK and Gambia

EEG habituation indices at 5 months of age were associated with imitation responses at 12 months of age in both cohorts. In the Gambian cohort, EEG novelty responses at 5 months were also associated with imitation responses as 12 months of age.

#### Katus et al., 2023

Longitudinal fNIRS and EEG metrics of habituation and novelty detection are correlated in 1-18-month-old infants.

EEG auditory oddball data and fNIRS habituation and novelty detection data at 1, 5 and 18 months in Gambian cohort only.

fNIRS and EEG responses were correlated for habituation at 1 and 5 months of age and for novelty detection at 5 and 18 months of age.

#### Milosavljevic et al., 2023

# Executive functioning skills and their environmental predictors among pre-school aged children in South Africa and The Gambia.

Executive Functioning (EF; working memory, inhibitory control, and cognitive flexibility) collected at 3-5 years in the Gambia, and a South African sample of the same age. Performance on these tasks was examined in relation to task norms, and family enrichment factors (enrichment activities, diversity of caregivers) and socioeconomic status (SES), and caregiver educational attainment.

The Gambian and South African participants showed normative performance across EF tasks, with evidence of heightened performance on the measure of cognitive flexibility. However, there were no associations between EF performance and measures of caregiver enrichment, SES or caregiver education in either site except an association between enrichment activities and working memory in the Gambian sample.

#### McCann et al., 2023

Iron status in early infancy is associated with cognitive development up to pre-school age in rural Gambia.

Iron status in early infancy and trajectories of cognitice development using MSEL and the Eye-Tracking Gap Overlap task.

Iron status (measured by soluble transferrin receptor -sTfR) at 5 months of age is associated with MSEL cognitive score at 5 months and trajectory of cognitive development from 5months to 5 years. sTfR at 5 months was also associated with an eye tracking measure of visual attention in cross sectional analysis, however this relationship was diminished by 5 years of age.

Katus\*, Crespo-Llado\* et al., 2024

It takes a village: caregiver diversity and language contingency in the UK and rural Gambia LENA data at 12, 18 and 24 months for UK and Gambia.

In the Gambia, earlier LENA measures of turn taking were associated with subsequent child vocalisations. In the UK, there was a trend toward such associations. In the Gambian cohort, we explored the role of caregiving: variability in caregiver numbers from one day to the next was associated with less turn taking. Total number of caregivers showed an inverted u-shaped relationship with turn taking: a medium number of caregivers was associated with highest turn taking frequencies.

Macrae et al., (2024)

Cognitive control in infancy: attentional predictors using a tablet-based measure

Tablet-based measure of cognitive skills, attentional disengagement (Gap-overlap task) and general cognitive skills (MSEL) at 5-24 months in the UK.

Participants showed significant improvements (at 24 months compared to 18 months) in cognitive control. General cognitive skills (measured using the MSEL) were not significantly associated with cognitive control, while attentional disengagement at 8 and 18 months was a significant predictor.

Pilot phase of the BRIGHT Project (2013 - 2014)

#### Milosavljevic et al., 2019

### Adaptation of the Mullen Scales of Early Learning for use among infants aged 5- to 24months in rural Gambia.

Pilot data collected using the MSEL, and measures of physical growth (height-for-age [HAZ], weight-for-height [WHZ], Head Circumference [HCZ]), collected at 9-24 months in The Gambia.

Primarily, we report on the method adapting the MSEL for use in the rural Gambian context. We also find that infants show normative MSEL performance early in infancy (at 5-9 months), however from 10-14 months, they show poorer performance compared to normed scores. Additionally, both HAZ and WHZ were significantly associated with MSEL performance.

#### Lloyd-Fox et al., (2017)

# Cortical specialisation to social stimuli from the first days to the second year of life: A rural Gambian cohort.

Pilot fNIRS and anthropometric data collected in first wave of longitudinal and crosssectional analyses of infants aged 1 – 24 months.

#### Lloyd-Fox et al., (2014)

# Functional Near Infrared Spectroscopy (fNIRS) to assess cognitive function in infants in rural Africa.

Pilot fNIRS data collected during first study in The Gambia – 4-8 month olds watched videos of a auditory and visual social cognition paradigm; a task showing Gambian adults performing social movements (i.e. Peekaboo) or vocalisations vs non-social stimuli Distinct regions of the posterior superior temporal and inferior frontal cortex evidenced either visual-social activation or vocally selective activation (vocal > non-vocal). The patterns replicated those observed within similar aged infants in previous studies in the UK. These are the first reported data on the measurement of localized functional brain activity in young infants in Africa

#### Begus et al., (2016)

#### Using fNIRS to study working memory in infants in rural Africa.

Pilot fNIRS data collected in cross-sectional analyses - 12-16 month olds watched videos of an *object permanence* paradigm; a task testing the ability to create and hold a mental schema of an object in mind, when it is no longer visually accessible Differential neural activity was seen when infants observed objects being hidden for 3 (posterior superior temporal activity) compared to 6 seconds (additional regions of inferior frontal and anterior superior temporal activity). This cortical activation could potentially reflect working memory and executive functioning.

#### Papademetriou et al, (2014) Optical imaging for brain activation in Gambian infants.

Optical topography of fNIRS data from Lloyd-Fox et al., 2014 A finite element model to model the propagation of light through tissue and create optical topography images of brain activation

Blasi et al., (2014) Test–retest reliability of functional near infrared spectroscopy in infants

Test re-test reliability of fNIRS signals from longitudinal data collection at 4 – 8 and 12 – 16 months of age

At the group level, good spatial overlap of significant responses and signal reliability was seen At participant level, spatial overlap was acceptable although results highlighted that signal reliability varied between participants.

*Competing Interests:* No competing interests were disclosed.

Reviewer Report 20 November 2023

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### ? Prerna Aneja 匝

University of East Anglia, Norwich, England, UK

The article makes an important contribution by highlighting the need to understand developmental trajectories in context of adversity. The paper proposes a comprehensive project through a longitudinal design and by considering various socio-cultural (risk and protective) factors. The key aims of the research include developing brain and neurocognitive function-for-age curves, establishing associations between poverty-related risk factors and developmental trajectories, and examining associations between context variables and preschool outcomes (long term outcomes). To this end, the authors outline the project design to investigate early brain and cognitive development in both The Gambia and UK (with the former being the primary focus of the project) as well as the developmental outcomes at 3-5 years of age in The Gambia.

### Methods:

#### NBAS:

The authors mention introducing controversial items with "special care". A short follow up with some concrete information could help the reader understand better what the special care would entail for work with this community. The authors mention that some aspects of task have been negatively perceived in other countries (I interpret that this falls outside of their own pilot). A citation for the research carried out in other nations would be helpful for readers who would like to do further readings on evaluations of the NBAS. Last, given the statement of "other countries" is not very concrete (e.g., do "other countries" include western nations?), it was not clear if added care was also necessary when using the clinical assessment in the UK. Clarifying this point would be helpful.

### Parent-child interaction videos:

The authors provide a good and clear general overview of the value and function of addressing PCI. To further enhance clarity, it would be helpful if the authors elaborated on points such as "videos can be coded to assess multiple aspects of parental and child behaviour and engagement". Are there specific key measures that the authors are planning to focus on or whether they are taking a more open and exploratory approach?

### **Sleep Diary:**

The Sleep diary section mentions that the diary was provided to caregivers. However, given the variability in caregiving figures across ages (as noted in the demographic characteristics section), I wondered whether the sleep diary was initially provided to mothers specifically (due to potentially being expected to act as primary caregivers), to other family members (e.g., grandparents), or if it was shared with the whole family for anyone to complete. Similarly, I wondered whether the person completing the sleep journal changed over time, given that secondary and tertiary caregivers shifted as children grew older.

#### PANAS:

The authors note that some of the English words used in the PANAS were not differentiable in

Mandinka, so the number of items were reduced in The Gambian version relative to the one administered in the UK. What was the total number of items for the positive and negative affect scores in the Gambia PANAS measure? It also may be worth it to address some information about which specific positive and negative affect words had to be reduced. For instance, the negative affect in the PANAS includes measures linked to emotions of fear (e.g., afraid and scared) as well as items conceptually linked to anger (e.g., hostile). Given that the emotions have some distinct characteristics (despite being grouped as negative), it could be helpful for readers to understand which specific dimensions of negative affect were affected most by the loss of items (e.g., did all items linked to a specific emotion, let's say feelings of anger, get removed from the scale? Did all emotional dimensions lose some items but keep others?).

#### <u>Results:</u>

#### BRIGHT Kids & Motor Skills:

The authors use an adaptation of the TGMD. It may be useful for the authors to draw from research that shows food reliability and consistency of the scale across a few countries to support their use of the measure (e.g., Rey *et al.*, 2020<sup>1</sup>).

#### Parent-report measures:

Given the nuanced approach to cultural and contextual sensitivity in the project and considering the complex influences that the COVID pandemic exerted across different communities, I wondered whether the Impact of COVID-19 was completed as a questionnaire by family members on their own, through an interview (e.g., by a field assistant, or through telephone calls, as noted in regards to the dietary data). Adding a bit more concrete information on the measure (e.g., example of items or scales options for responses as done in the Parental Mental Health section) would be helpful.

#### References

1. Rey E, Carballo-Fazanes A, Varela-Casal C, Abelairas-Gómez C, et al.: Reliability of the test of gross motor development: A systematic review.*PLoS One*. 2020; **15** (7): e0236070 PubMed Abstract | Publisher Full Text

### Is the rationale for, and objectives of, the study clearly described?

Yes

#### Is the study design appropriate for the research question?

Yes

# Are sufficient details of the methods provided to allow replication by others? Partly

### Are the datasets clearly presented in a useable and accessible format?

Yes

*Competing Interests:* No competing interests were disclosed.

Reviewer Expertise: Infant cognitive development, visual attention, cultural

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

#### Author Response 12 May 2024

#### Sarah Lloyd-Fox

We thank you for your helpful and constructive comments. We have addressed all comments in our revision and believe this has strengthened the manuscript. Below we provide detailed responses to the individual points raised (your points in bold, ours in standard or italicised.

The article makes an important contribution by highlighting the need to understand developmental trajectories in context of adversity. The paper proposes a comprehensive project through a longitudinal design and by considering various socio-cultural (risk and protective) factors. The key aims of the research include developing brain and neurocognitive function-for- age curves, establishing associations between poverty-related risk factors and developmental trajectories, and examining associations between context variables and preschool outcomes (long term outcomes). To this end, the authors outline the project design to investigate early brain and cognitive development in both The Gambia and UK (with the former being the primary focus of the project) as well as the developmental outcomes at 3-5 years of age in The Gambia.

Methods:

1. NBAS:

The authors mention introducing controversial items with "special care". A short follow up with some concrete information could help the reader understand better what the special care would entail for work with this community. The authors mention that some aspects of task have been negatively perceived in other countries (I interpret that this falls outside of their own pilot). A citation for the research carried out in other nations would be helpful for readers who would like to do further readings on evaluations of the NBAS. Last, given the statement of "other countries" is not very concrete (e.g., do "other countries" include western nations?), it was not clear if added care was also necessary when using the clinical assessment in the UK. Clarifying this point would be helpful.

**Response:** We agree that this added information would be useful to the reader, and have thus provided the below information:

In line with feedback from parents from other countries (across low-, middle- and high-income settings, see Katus et al., 20204) and populations, a few aspects of the assessment (specifically, shining a light over closed eyes while sleeping, covering eyes with cloth while awake and undressing the infant) were questioned or perceived negatively (Super et al., 2020). While no items were altered or removed from the assessment, the more controversial items were introduced with special care during all study visits. Adjustment to the protocol included to either i) provide additional explanation on the rationale for an item (e.g., for habituation to light, or

covering the infant's eyes) to check whether parents approved of the item's administration (e.g., undressing); or ii) allow examiners to skip an item if faced with an unsuitable environment (e.g., no suitable surface to assess reflexes). Overall, the interviews emphasised the need to communicate with caregivers prior to and during the assessment, and in addition to observe the infant and be aware of cues from parents throughout. Another recurrent theme was that parents allowed certain actions to be performed specifically because they were part of a research study. This emphasised the ethical responsibility of making sure all items administered were relevant and provided a benefit to the participant or the wider community. In both the UK and The Gambia, the NBAS was performed between 7-14 days after birth during a pre-arranged home visit. Administration times for the NBAS ranged from 20 to 45 minutes, depending on the infants' state of alertness.

1. Parent-child interaction videos:

The authors provide a good and clear general overview of the value and function of addressing PCI. To further enhance clarity, it would be helpful if the authors elaborated on points such as "videos can be coded to assess multiple aspects of parental and child behaviour and engagement". Are there specific key measures that the authors are planning to focus on or whether they are taking a more open and exploratory approach?

**Response:** Thank you for this suggestion, the following has been added to the main text to address this:

Coding parent-infant interactive styles is an ongoing process, with different approaches undertaken at different age points. At the 1- and 5-month visits, we implemented an adapted version of the Global Rating Scales (Murray et al., 1996) that had previously been used with rural and/or low-income communities in South Africa and India (Bozicevic et al., 2016; Cooper et al., 2002; Holla et al., 2021; Tomlinson et al., 2005). The GRS assesses maternal (contingency, responsiveness, affect), infant (active communication, attentiveness) and interactive (level and intensity of dyadic engagement) characteristics. The GRS have been designed to be comparable across cultures by ensuring that key dimensions are assessed independently of modality. Thus, the main adaptation involved removing items that relied heavily on speech content, as prior work demonstrated that, at this young age, speech content does not influence infant behaviour (Bozicevic et al., 2016; Stern et al., 2016). However, as infants get older, both language and cultural norms play an increasing role in social interactions. Thus, it is our aim to develop objective, culturally tailored assessments, of parent-infant interactions, rather than relying solely on schemes developed for other cultures. Our approach in developing these new coding schemes thus far have been to run focus groups with Gambian research staff. These involve an iterative process whereby we begin by asking for reflections on caregiving and infant/child behaviours (across different ages) within this population; then ask open questions within the group while viewing videos to describe behaviours seen and whether they are frequently used by families living in this cultural setting; then introduce the group to commonly used coding schemes; and finally co-create a coding scheme that combines the advantages of standardised frameworks with the specificity required for a given context.

1. Sleep Diary:

The Sleep diary section mentions that the diary was provided to caregivers. However, given the variability in caregiving figures across ages (as noted in the demographic characteristics section), I wondered whether the sleep diary was initially provided to

mothers specifically (due to potentially being expected to act as primary caregivers), to other family members (e.g., grandparents), or if it was shared with the whole family for anyone to complete. Similarly, I wondered whether the person completing the sleep journal changed over time, given that secondary and tertiary caregivers shifted as children grew older.

**Response:** We have added the following clarification to the section describing the sleep diary to address this point:

As was the case with the maternal mental health questionnaires, the sleep diary was administered as an interview by field staff with the primary caregiver who accompanied the child to the visit. This was most commonly the mother across study visits.

1. **PANAS:** 

The authors note that some of the English words used in the PANAS were not differentiable in

Mandinka, so the number of items were reduced in The Gambian version relative to the one

administered in the UK. What was the total number of items for the positive and negative affect scores in the Gambia PANAS measure? It also may be worth it to address some information about which specific positive and negative affect words had to be reduced. For instance, the negative affect in the PANAS includes measures linked to emotions of fear (e.g., afraid and scared) as well as items conceptually linked to anger (e.g., hostile). Given that the emotions have some distinct characteristics (despite being grouped as negative), it could be helpful for readers to understand which specific dimensions of negative affect were affected most by the loss of items (e.g., did all items linked to a specific emotion, let's say feelings of anger, get removed from the scale? Did all emotional dimensions lose some items but keep others?).

**Response:** We have added the following to the description of the PANAS to address this query:

*Furthermore, some of the English words used, in the PANAS, were not differentiable in Mandinka, and so the number of items were reduced in The Gambian version relative to the one administered in the UK. The Mandinka version of the PANAS included 6 items relating to PA (Curious, Happy, Strong, Proud, Determined, Active) and 6 items relating to NA (Distressed, Regretful, Scared, Aggressive, Ashamed, Uncomfortable).* **Results:** 

1. BRIGHT Kids & Motor Skills:

The authors use an adaptation of the TGMD. It may be useful for the authors to draw from

research that shows food reliability and consistency of the scale across a few countries to support their use of the measure (e.g., Rey et al., 20201).

**Response:** We thank the reviewer for this comment and suggested reference, we include the below in the main text to address this:

The TGMD (in its various editions) has been used to assess GM skills in multiple countries with good reliability and consistency across settings (Rey et al., 2020). Of note is that it has successfully been implemented in other rural African communities to assess pre-school and

school-aged children (Cook et al., 2019; Tomaz et al., 2019; Tsikata et al., 2021). 1. **Parent-report measures:** 

Given the nuanced approach to cultural and contextual sensitivity in the project and considering the complex influences that the COVID pandemic exerted across different communities, I wondered whether the Impact of COVID-19 was completed as a questionnaire by family members on their own, through an interview (e.g., by a field assistant, or through telephone calls, as noted in regards to the dietary data). Adding a bit more concrete information on the measure (e.g., example of items or scales options for responses as done in the Parental Mental Health section) would be helpful.

**Response:** We have included the below in the description of the Covid-19 questionnaire to address this.

Impact of COVID-19

The BRIGHT Kids follow up was conducted during the COVID-19 pandemic and, thus, it was vital to better understand how the pandemic impacted on each family's wellbeing, financial situation and access to healthcare. The questionnaire was adapted from a measure that was being used in a UK study at the time to assess the impact of the pandemic on families with young children (Aydin et al., 2022). Items that were deemed by field staff to be irrelevant or unsuitable for the West Kiang community were removed. The final measure consisted of a series of questions that asked the caregiver to report on how their living and financial situation had changed due to the pandemic (e.g., "Has COVID-19 had an impact on your ability to get enough food for your household?"), whether they had family members who got sick with COVID-19 (e.g., "Do you think someone in your household has had, or currently has COVID-19?"), their ability to access healthcare (e.g., "Has COVID-19 impacted your ability to see a doctor or go to the hospital when you or a family member needed to?"), and their general feelings and concerns related to the situation (e.g., "I am afraid of COVID-19" – not at all, a little, a lot). As was the case with the maternal mental health questionnaires, the questionnaire was administered as an interview by field staff with the primary caregiver who accompanied the child to the visit. Once again, we thank the two reviewers for their thorough, thoughtful, and constructive feedback, which will undoubtedly strengthen this manuscript. We hope that we have addressed in sufficient detail.

Competing Interests: No competing interests were disclosed.