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1 Study profile

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- 2 Cohort description: Measures of early-life behaviour and later psychopathology
- in the LifeCycle Project EU Child Cohort Network
- 5 Authors and Affiliations
- 6 Johanna L. Nader^{1*}, Mònica López-Vicente^{2,3}, Jordi Julvez^{4,2}, Monica Guxens^{2,3}, Tim Cadman⁵,
- 7 Ahmed Elhakeem⁵, Marjo-Riitta Järvelin⁶, Nina Rautio⁶, Jouko Miettunen^{6,7}, Hanan El Marroun^{3,8,9},
- 8 Maria Melchior¹⁰, Barbara Heude¹¹, Marie-Aline Charles^{11,12}, Tiffany C. Yang¹³, Rosemary R. C.
- 9 McEachan¹³, John Wright¹³, Kinga Polanska¹⁴, Jennie Carson¹⁵, Ashleigh Lin¹⁵, Sebastian Rauschert¹⁵,
- 10 Rae-Chi Huang¹⁵, Maja Popovic¹⁶, Lorenzo Richiardi¹⁶, Eva Corpeleijn¹⁷, Marloes Cardol¹⁷, Tuija M.
- 11 Mikkola^{18,19}, Johan G. Eriksson^{18,20,21,22,23}, Theodosia Salika²⁴, Hazel Inskip^{24,25}, Johan Lerbech
- 12 Vinther²⁶, Katrine Strandberg-Larsen²⁶, Kathrin Gürlich²⁷, Veit Grote²⁷, Berthold Koletzko²⁷, Marina
- Vafeiadi²⁸, Jordi Sunyer², Vincent W. V. Jaddoe^{8,9}, Jennifer R. Harris²⁹ for the LifeCycle Project
- 14 Group.

- 16 Department of Genetics and Bioinformatics, Division of Health Data and Digitalisation, Norwegian Institute of Public Health, Oslo, Norway
- 17 ²ISGlobal, Instituto de Salud Global de Barcelona, Barcelona, Spain
- 18 ³Department of Child and Adolescent Psychiatry, University Medical Center Rotterdam, Erasmus MC
- ⁴Institut d'Investigació Sanitària Pere Virgili, Hospital Universitari Sant Joan de Reus, 43204 Reus, Spain
- 20 5MRC Integrative Epidemiology Unit at University of Bristol, Population Health Sciences, Bristol Medical School, Bristol, UK
- 21 ⁶Center for Life Course Health Research, University of Oulu, Oulu, Finland
- ⁷Medical Research Center Oulu, Oulu University Hospital and University of Oulu, Oulu, Finland
- 23 *Department of Pediatrics, University Medical Center Rotterdam, Erasmus MC
- ⁹The Generation R Study Group, Erasmus MC, Rotterdam, CA, The Netherlands
- 25 ¹⁰Sorbonne Université, INSERM, Institut Pierre Louis d'Épidémiologie et de Santé Publique, IPLESP, 75012, Paris, France
- 26 11 Université de Paris, Centre for Research in Epidemiology and Statistics (CRESS), INSERM, INRAE, Paris, France
- 27 12Unité mixte Inserm-Ined-EFS Elfe, INED, Paris, France
- 28 13Bradford Institute for Health Research, Bradford Teaching Hospitals NHS Foundation Trust, Bradford, UK
- 29 ¹⁴Department of Hygiene and Epidemiology, Medical University of Lodz, Lodz, Poland
- 30 ¹⁵Telethon Kids Institute, University of Western Australia, Australia
- 31 ¹⁶Cancer Epidemiology Unit, Department of Medical Sciences, University of Turin and CPO Piemonte, Turin, Italy

32	¹⁷ Department of Epidemiology, University Medical Center Groningen, Groningen, The Netherlands
33	¹⁸ Folkhälsan Research Center, Helsinki, Finland
34	¹⁹ Clinicum, Faculty of Medicine, University of Helsinki, Finland
35	²⁰ Public Health Promotion Unit, National Institute for Health and Welfare, Helsinki, Finland
36	²¹ Department of General Practice and Primary Health Care, University of Helsinki and Helsinki University Hospital, Helsinki, Finland
37	²² Singapore Institute for Clinical Sciences, Agency for Science, Technology, and Research, Singapore
38	²³ Department of Obstetrics & Gynaecology, Yong Loo Lin School of Medicine, National University of Singapore, Singapore
39	²⁴ Medical Research Council Lifecourse Epidemiology Unit, Southampton General Hospital, University of Southampton, Southampton, UK.
40	²⁵ NIHR Southampton Biomedical Research Centre, University Hospital Southampton NHS Foundation Trust, University of Southampton,
41	Southampton, UK.
42	²⁶ Section of Epidemiology, Department of Public Health, University of Copenhagen, Copenhagen, Denmark
43	²⁷ Division of Metabolic and Nutritional Medicine, Department of Pediatrics, Dr. von Hauner Children's Hospital, University Hospital, LMU,
44	Munich, Germany
45	²⁸ Department of Social Medicine, Faculty of Medicine, University of Crete, Heraklion, Greece.
46	²⁹ Division of Health Data and Digitalization, Center for Fertility and Health and Department of Genetics and Bioinformatics, The Norwegian
47	Institute of Public Health, Oslo, Norway
48	
49	Corresponding author: Dr. Johanna Lucia Nader, Department of Genetics and Bioinformatics, Norwegian Institute of Public
50	Health, Sandakerveien 24 C, 0473 Oslo, Norway. E-mail: JohannaLuciaThorbjornsrud.Nader@fhi.no
51	
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Abstract

Background: The EU LifeCycle Project was launched in 2017 to combine, harmonise, and analyse data from more than 250,000 participants across Europe and Australia, involving cohorts participating in the EU-funded LifeCycle Project. The purpose of this cohort description is to provide a detailed overview over the major measures within mental health domains that are available in 17 European and Australian cohorts participating in the LifeCycle Project.

Methods: Data on cognitive, behavioural and psychological development has been collected on participants from birth until adulthood through questionnaire and medical data. We developed an inventory of the available data by mapping individual instruments, domain types, and age groups, providing the basis for statistical harmonization across mental health measures.

Results: The mental health data in LifeCycle contain longitudinal and cross-sectional data for ages 0-18+ years, covering domains across a wide range of behavioural and psychopathology indicators and outcomes (including executive function, depression, ADHD and cognition). These data span a unique combination of qualitative data collected through behavioural/cognitive/mental health questionnaires and examination, as well as data from biological samples and indices in the form of brain imaging (MRI, foetal ultrasound) and DNA methylation data. Harmonized variables on a subset of mental health domains have been developed, providing statistical equivalence of measures required for longitudinal meta-analyses across instruments and cohorts.

Conclusion: Mental health data harmonized through the LifeCycle project can be used to study life course trajectories and exposure-outcome models that examine early life risk factors for mental illness and develop predictive markers for later-life disease.

Keywords: Birth and pregnancy cohorts, Child behaviour and mental health, Population epidemiology, Child development, DataSHIELD

Background and Purpose

Effects of early life exposures on later life mental health are well known, but more research to understand and elucidate the pathways from stressors to outcomes is needed. The LifeCycle Project - EU Child Cohort Network, a Horizon 2020 project, is a pan-European and Australian initiative comprised of 19 pregnancy and birth cohorts, established to study exposure-to-outcome associations and trajectories across the life course (https://lifecycle-project.eu/). In general, studies in LifeCycle aim to construct developmental trajectories, develop risk assessment models, measure developmental adaptations and evaluate mediating epigenetic effects to better understand the consequences of early-life exposures to stressors for risk factors and diseases in adulthood. The large sample sizes achieved through this consortium facilitate high statistical power needed for increased accuracy of estimates and more robust findings.

Mental health is one of the main outcomes within the LifeCycle Project.¹ While mortality rates for many non-communicable diseases have steadily declined in some populations over the past few decades, such as coronary heart disease^{2,3} and chronic obstructive pulmonary disease,⁴ the global burden of mental illness is on the rise.⁵ The impact of mental illness on disability and socioeconomic prosperity is increasing around the world, and it is predicted that mental illness will contribute more to disability-adjusted life years (DALYs) than any other category of diseases by the year 2030.⁶ An understanding of how mental health impacts and mediates disease risk and prognosis for other conditions is also beginning to emerge, with recent meta-analyses revealing significantly higher risks for cardiovascular⁷ and metabolic⁸ diseases linked to severe mental illness.

This cohort description focuses on the extensive work done to catalogue and harmonise variables related to cognitive, behavioural, and psychological development within the broader LifeCycle consortium. It is well-recognised that experiences in early life play an important part in shaping later mental health and the data within the LifeCycle Project permit analyses of these associations. LifeCycle includes many pregnancy and birth cohorts that prospectively collected data on offspring from conception and across different ages of child, adolescent, and adult development. The

availability of data from multiple follow-up assessments is essential for probing questions about causality and linking early life stressors with later life mental health symptoms and outcomes.

The mental health studies in LifeCycle aim to investigate epidemiological interrelations between early life exposures, behaviour, and cognition, with later mental and physical health. Towards this end we have harmonised measures from 17 LifeCycle cohorts to enable studies that examine how environmental stressors *in utero* and in early childhood affect, or are associated with, psychological trajectories, behaviours, and mental outcomes throughout childhood, adolescence and adulthood. Additionally, we are examining the nature and degree of mediation of these associations through epigenetic changes and brain development (Figure 1). To our knowledge, the data compiled for these studies within LifeCycle represents the largest ongoing consolidation of childhood behaviour, psychopathology and cognition data to date, encompassing more than 200 multidimensional and multi-informant established mental health measures collected from at least 250,000 participants.

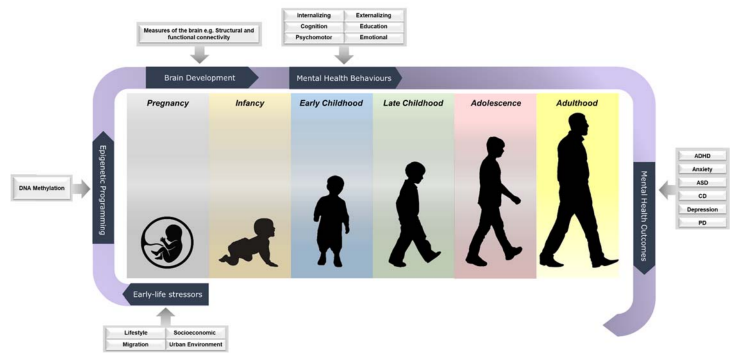


Figure 1. Available mental health outcomes from prenatal to adulthood in the LifeCycle mental health and cognitive

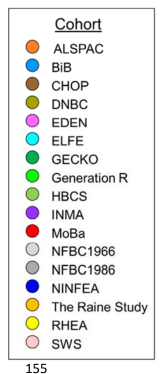
data

(ADHD: Attention deficit hyperactivity disorder; ASD: Autism spectrum disorders; CD: cognitive disorders; PD: psychiatric disorders)

Cohorts, participants and follow-up

A total of 17 child-parent cohorts based in 13 countries are contributing with mental health data: Avon Longitudinal Study of Parents and Children (ALSPAC, United Kingdom), Born in Bradford (BiB, United Kingdom), EU Childhood Obesity Programme (CHOP, Germany/Italy/Spain/Poland/Belgium), Danish National Birth Cohort (DNBC, Denmark), Etude des Déterminants du développement et de la santé de l'Enfant (EDEN, France), Etude Longitudinale Française depuis l'Enfance (ELFE, France), Groningen Expert Center for Kids with Obesity Drenthe cohort (GECKO Drenthe cohort, The Netherlands), the Generation R Study (Generation R, The Netherlands), Helsinki Birth Cohort Study (HBCS, Finland), Infancia y Medio Ambiente (INMA, Spain), The Norwegian Mother, Father and Child Cohort Study (MoBa, Norway), Northern Finland Birth Cohorts (NFBC1966/1986, Finland), Nascita e INFanzia: gli Effetti dell'Ambiente (NINFEA, Italy), The Raine Study (Australia), Rhea Mother & Child Cohort Study (RHEA, Greece), and the Southampton Women's Survey (SWS, United Kingdom).





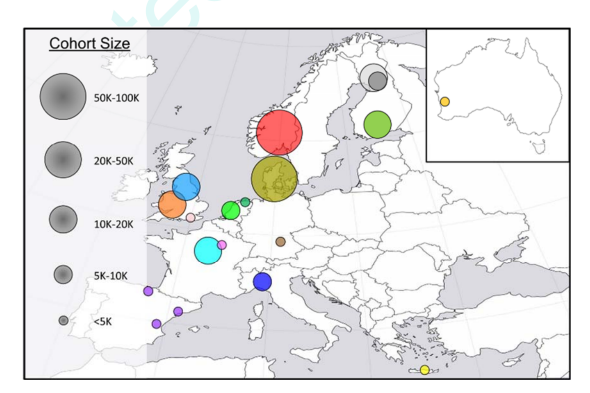


Figure 2. Geographic distribution and sample sizes of cohorts in LifeCycle contributing mental health

157 data

The geographic coverage is broad, spanning across much of northern, western, central and southern Europe as well as Western Australia (Figure 2). Mental health data from more than 250,000 children are available (as of June 2021), including either mother-child or mother-father-child cohorts, and the study population is diverse with respect to the age of the participants, cohort types, and data collection periods (Table 1). As described elsewhere for the LifeCycle consortium, most of the cohorts in the LifeCycle project (ALSPAC, CHOP, DNBC, EDEN, GECKO, HBCS, INMA, MoBa, NFBC1966/1986, NINFEEA, RHEA, and SWS) predominantly represent ethnic groups from the background population (more than 95% European/White), but certain cohorts like BiB, ELFE, The Generation R Study, and The Raine Study have significant representation of other ethnic groups as well. ¹⁰

170 Table 1. Summary characteristics of LifeCycle cohorts participating with mental health data

Cohort	Location of Coordinating Centre	Cohort Type	Data collection period	Recruitment	N (Live Births)	
ALSPAC ^{11,12}	Avon, United Kingdom	Population-based	1990-present	Pregnancy	14,953	
BiB ¹³	Bradford, United Kingdom	Population-based	Pregnancy	13,786		
CHOP ¹⁴	Belgium (Liege, Brussels), Germany (Munich, Nuremberg), Italy (Milano), Poland (Warsaw), Spain (Reus, Tarragona)	Mixed (Randomised controlled intervention trial (first year) with birth cohort)	2002-2015	First 8 weeks of life	1,678	
DNBC ¹⁵	Copenhagen, Denmark	Population-based	1996-present	Pregnancy	96,804	
EDEN ¹⁶	Nancy and Poitiers, France	Population-based	2003-2017	Pregnancy	1,907	
ELFE ¹⁷	Paris, France	Population-based	2011-present	Birth	18,329	
GECKO ¹⁸	Drenthe, The Netherlands	Population-based	2006-present	Pregnancy	2844	
The Generation R Study ^{19,20}	Rotterdam, The Netherlands	Population-based	2002-present	Pregnancy	9,749	
HBCS ²¹	Helsinki, Finland	Population-based	1934-present	Birth	13,345	
	Sabadell, Spain	Population-based	2004-present	Pregnancy	622	
INMA ²²	Valencia, Spain	Population-based	2003-present	Pregnancy	787	
	Gipuzkoa, Spain	Population-based	2006-present	Pregnancy	612	
MOBA ²³	Oslo, Norway	Population-based	1999-present	Pregnancy	113,564	
NFBC1966 ²⁴	Oulu, Finland	Population-based	1966-present	Pregnancy	12,058	
NFBC1986 ²⁵	Oulu, Finland	Population-based	1985/1986-present	Pregnancy	9,432	
NINFEA ²⁶	Torino, Italy	Population-based (Internet-based recruitment)	2005-present	Pregnancy (Internet- based recruitment)	6,816	
The Raine Study ²⁷	Perth, Australia	Population-based (Randomised assignment to multiple ultrasounds during pregnancy)	1989-present	Pregnancy	2,868	
RHEA ²⁸	Crete, Greece	Population-based	2007-present	Pregnancy	1,458	
SWS ²⁹	Southampton, United Kingdom	Population-based	1998-present	Pre-pregnancy	3,158	

The participating cohorts include child participants with follow-up data ranging from birth until adulthood (Table 2). Questionnaires, medical records, doctor diagnoses and registries were variably

used across the cohorts to collect data at different ages, but all of the cohorts collected baseline data during pregnancy or at birth, and included a follow-up data collection at least once by the time the child participant was 24 months of age. Although the regularity of follow-up differs substantially across cohorts, ranging from annually to many years apart, at least half of the cohorts performed some type of follow-up data collection for all incremental age groups up until 6 years of age. The overlapping age ranges enable comprehensive comparative analyses of mental health constructs between and within the populations to which these index children belong.

Table 2. Age ranges and sex (% male:female) of participants during assessment in LifeCycle cohorts

Cohort	Baseline (no. live births)	Age of child at assessment (years)														
		0 to <1	1 to <2	2 to <3	3 to <4	4 to <5	5 to <6	6 to <7	7 to <8	8 to <9	9 to <10	10 to <12	12 to <14	14 to <16	16 to <18	18+
ALSPAC ^a	14,953	11,466	11,097	9,993	9,779	9, 632	8,683	8,410	8,282	7,481	7,718	7,552	6,829	5,506	5,212	
Sex (% M:F)		51.6:48.4	51.7:48.3	51.8:48.2	51.7:48.3	51.8:48.2	51.6:48.4	51.4:48.6	50.7:49.3	49.8:50.2	49.3:50.7	49.4:50.6	49.1:50.9	47.1:52.0	43.6:56.4	
BiB Sex (% M:F)	13,786 51.6:48.4	1,436 49.6:50.4	3,484 50.3:49.7	2,911 50.1:49.9	1,167 47.9:52.1	2,505 49.9:50.1	79 51.9:48.1									
CHOP ^b Sex (% M:F)	1,678 50.7:49.3	1,175 49.0:51.0	1,067 48.1:51.9	934 48.2:51.8	747 46.6:53.4	674 47.2:52.8	655 47.2:52.8	1,028 48.5:51.5	594 49.0:51.0	589 47.0:53.0		719 46.5:53.5				
DNBC Sex (% M:F)	96,804 51.3:48.7	70,276 51.1:48.9	65,548 51.0:49.0				1,628° 52.0:48.0		57,156 51.2:48.8			46,345 ^d 49.7:50.3 48,579 ^c 48.2:51.8				35,558 ^f 41.6:58.4
EDENg	1,907		1,612	1,429	1,257	1,192	1,114					557				
Sex (% M:F)	10.220	16.515	52.8:47.2	52.2:47.8	52.4:47.6	51.3:48.7	52.7:47.3					51.3:48.7				
ELFE Sex (% M:F)	18,329 51.4:48.6	16,547 51.2;48.9	14,439 51.2:48.8	13,277 50.7:49.3	11,935 51.2:48.8											
GECKO	2.844	2,812	2.558	2,319	1,819	1,486	2,322					2,299				
Sex (% M:F)	2,0	50.3:49.7	50.3:49.7	50.1:49.9	51.2:48.8	51.4:48.6	50.3:49.7					49.8:50.2				
Generation R Sex (% M:F)	9,749 50.7:49.3	7,893 50.5:49.5					8,305 50.5:49.5				7,393 50.1:49.9		6,842 50.3:49.7			
HBCS Sex (% M:F)	13,345 52,3:47,7	13,345 52.3:47.7	13,342 52.3:47.7	13,342 52.3:47.7	8,947 52,0:48,0	7,252 51.7:48.3	9,947 52.6:47.4	10,055 52,7:47,3	10,046 52,6:47,4	10,033 52.7:47.3	9,985 52.8:47.2	9,902 52.8:47.2				13,345 52.3:47.7
INMA-Sabadell Sex (% M:F)	622		559 51.3:48.7			481 51.4:48.7		473 51.6:48.4			433 52.0:48.0		8 0 0 0			
INMA-Valencia Sex (% M:F)	787		694 52.6:47.4				530 51.7:48.3		469 50.8:49.3		429 50.6:49.4					
INMA-Gipuzkoa	612		556	506		394			397	382						
Sex (% M:F)	50.3:49.8		49.1:50.9	52.0:48.0		49.2:50.8			49.4:50.6	54.0:46.0						
MoBa Sex (% M:F)	113,564	87,801 51.0:49.0	74,750 51.0:49.0		58,835 51.0:49.0		41,617 50,9:49,1		53,517 51,3:48,7	43,609 50,9:49.1						
NFBC1966 Sex (% M:F)	12,058	31:0.49.0	10,729 50.8:49.2		31.0.49.0		30.9.49.1		31.3.46.7	30.3.43.1				10,927 50.4:49.6		9,517 51.3:48.7
NFBC1986 Sex (% M:F)	9,432		1,803 50.9:49.1						8,416 ^d 51.3:48.7 8,525 ^h 51.5:48.5					6,985 ^d 50.0:50.0 7,344 ^e 48.5:51.5 6,795 ⁱ 49.4:50.6		Data collection ongoing (2019- 2020)
NINFEA Sex (% M:F)	7,527 ^j	6,907 50.7:49.3	6,279			4,398 51.1:48.9			2,348 50.3:49.7			837 50.8:49.2				
The Raine Study Sex (% M:F)	2,868 50.7:49.3		2,430 50.9:49.1	1,974 52.1:47.9	2,260 50.9:49.1		2,236 51.6:48.4	0		2,140 51.4:48.6		2,048 51.7:48.3		1,864 51.4:48.6	1,693 49.9:50.1	1,462 48.9:51.1
RHEA Sex (% M:F)	1,458 50.1:49.9	1,257 50.2:49.8	569 54.5:45.5			904 52.3:47.7		626 55.1:44.9								
SWS Sex (% M:F)	3,158	2,959 51.7:48,3	2,875 51.9:48,1	2,779 51.8:48.2	2,625 52.1:47.9	1,182 51.9:48.1		2,034 51.3:48.7		1,214 49.4:50.6						

ALSPAC follow-up data is based on number of parents completing at least some of the questionnaire(s) on young person up to age 7 years, and number of children attending clinic from age 7 years and onwards

^bCHOP follow-up data is based on number of children with at least one anthropometric measurement at the considered age

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[°]DNBC follow-up data at 5 years based on a subsample, selected based on parental alcohol characteristics

^dParent-reported data

[°]Self-reported data

DNBC data collection for 18-year follow-up is currently ongoing geDEN follow-up data is based on number of children with at least one neurodevelopment assessment at the considered age

hTeacher-reported data

ⁱClinical data

^jNINFEA baseline data refers to no. pregnant women recruited

Main outcome measures

Psychological, motor and cognitive measures

Mental and cognitive disorders comprise some of the most frequently diagnosed conditions in children under 18 years of age. The combined data resource will contain information pertaining to the children from more than 200 mental health measures, covering eight clinical domains across 60 dimensions (Table S1). A majority of these measures assess domains under a broad banner of 'mental health', encompassing psychological functions, cognitive and executive functions and psychological development (67.0%; 136 of 203), covering dimensions such as neurodevelopmental disorders, internalising and externalising symptoms, temperament and mental diagnoses. Further domains include language skills (31.0%; 63 of 203), executive functions (29.1%; 59 of 203), memory (11.3%; 23 of 203) and general intelligence (8.4%; 17 of 203) (Table S1). There are many commonalities between mental health domain-types and significant overlap in the age groups with measures in specific domains (Figure 3). This makes it possible to harmonise the data. Most of the cohorts continuously follow up their participants, and the availability of harmonised data will tend to increase with time.

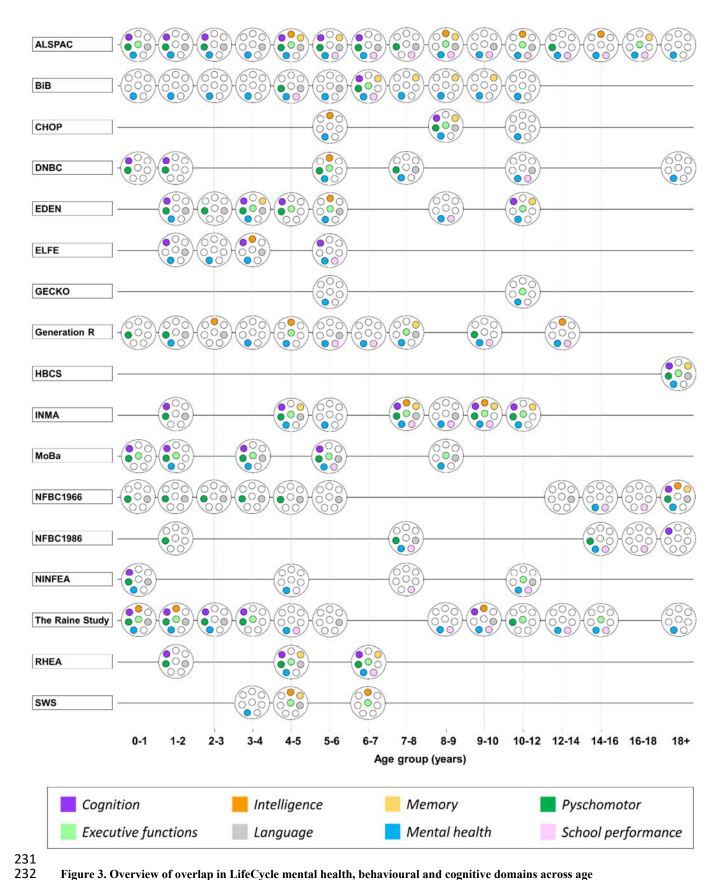


Figure 3. Overview of overlap in LifeCycle mental health, behavioural and cognitive domains across age

Basic illustration of the range of developmental domains and participant ages for data in the LifeCycle Project.

There are a number of approaches to harmonise data and several of these have been described and successfully implemented in large collaborations. 31,32,33,10 The LifeCycle Project has developed a protocol to generate harmonised variables across a selection of important cognitive and mental health domains. This harmonisation approach creates standardised scores and percentiles for important domains such as internalising and externalising symptoms, ADHD and ASD symptoms and diagnosis, and language and motor functions. Percentiles and standardised scores were used as they allow the pooling of mental health outcome data collected using different scales or instruments. One of the biggest harmonisation challenges this project faced was obtaining a thorough inventory of the available mental health data in individual cohorts, which was overcome by mapping the available data by instrument, measure, age group, and domain. A subset of cohorts has also employed items from the same mental health, cognitive and motor function measures, and these data can be pooled or coanalysed without the need for harmonisation (Figure 4). All of the measures harmonised thus far by age and cohort can be found in the LifeCycle online catalogue (https://catalogue.lifecycle-project.eu/).

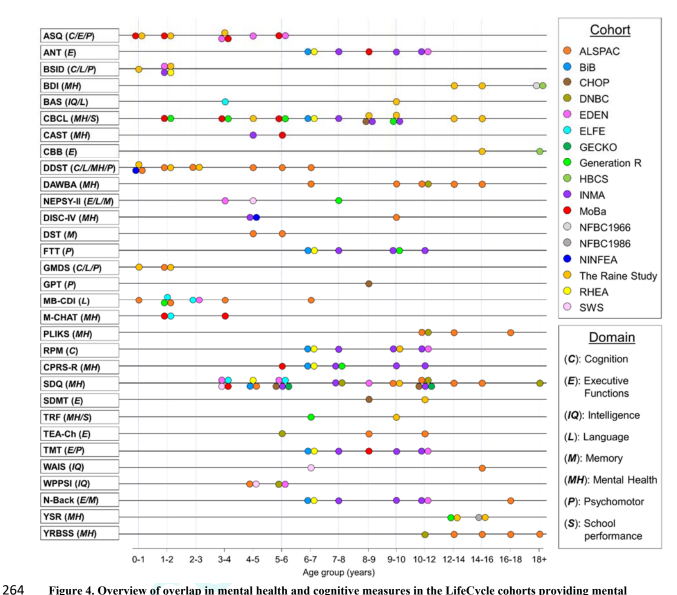


Figure 4. Overview of overlap in mental health and cognitive measures in the LifeCycle cohorts providing mental health data

Summary of overlapping measures and age ranges in participating cohorts. The full list of available measures (including non-overlapping) are described in supplementary table 1.

(ANT: Attention Network Task; ASQ: Ages and Stages Questionnaire; BAS: Behavioural Approach System; BDI: Becks Depression
Inventory; BRIEF: Behaviour Rating Inventory of Executive Function; BSID: Bayley Scales of Infant Development; CAST: Childhood
Asperger Syndrome Test; CBB: CogState Brief Battery; CBCL: Child Behaviour Checklist; CPRS-R: Revised Conners' Parent Rating Scale;
DAWBA: Development and Well-Being Assessment; DDST: Denver Developmental Screening Test; DISC-IV: Diagnostic Interview
Schedule for Children; DST: Digit Span Test; FTT: Finger Tapping Test; GMDS: Griffiths Mental Development scales; GPT: Grooved
Pegboard Test; M-CHAT: Modified Checklist for Autism in Toddlers; MB-CDI: MacArthur-Bates Communicative Development
Inventories; N-Back: Working Memory Test; NEPSY-II: Developmental NEuroPSYchological Assessment, Second Edition; PLIKS:
Psychosis-like symptoms measure; RPM: Raven's Progressive Matrices; SDMT: Symbol Digit Modalities Test; SDQ: Strengths and
Difficulties Questionnaire; TEA-ch: Test of Everyday Attention for Children; TMT: Trail Making Test; TRF: Teacher Report Form; WASI:
Wechsler Abbreviated Scale of Intelligence; WPPSI: Wechsler Preschool and Primary Scale of Intelligence; YRBSS: Youth Risk Behavior
Surveillance System; YSR: Youth Self-Report).

Early-life exposures – lifestyle, migration, socioeconomic, and urban environment

The LifeCycle online catalogue¹⁰ also contains information on harmonised data on diverse measures of exposures early in life. These will enable the analysis of risk models for mental health that assess the nature and impact of indirect and direct exposures experienced in early life, and comorbidities with adverse mental health symptoms and other health conditions. Comprehensive exposure-outcome analyses will also be used to develop predictive markers for mental health in children and adolescents, which may help shape the prediction of mental disorders, allowing for targeted early intervention.

Mediating Pathways - Brain Development

Early life is a particularly vulnerable time-window for brain development. The vital stages of neurogenesis, proliferation and migration occur almost exclusively during foetal development, and experience-dependent brain connectivity (i.e. myelination) is largely shaped and completed in early childhood.³⁴ Research-based evidence has repeatedly linked brain structure, volume, and connectivity indicators to a number of behavioural and cognitive outcomes. 35,36,37 However, study samples are often limited in size and population diversity, and only few longitudinal studies exist.³⁸ A subset of cohorts in LifeCycle have participant data on structural brain imaging (ALSPAC n=950; Generation R n≈4000;³⁹ NFBC1966 n=1000; NFBC1986 n=600), and will be contributing information on neuroanatomical markers such as total brain volume, cortical grey matter, white matter volume, ventricular volume, and volumes of subcortical brain structures including the hippocampus and amygdala. In addition, structural and functional connectivity metrics have been assessed. Data have been collected through neuroimaging techniques, such as foetal ultrasound and Magnetic Resonance Imaging (MRI) in childhood and /adulthood MRI. These data enable LifeCycle to describe changes in structural and functional development of the brain from foetal life and infancy, and to subsequently associate this brain development in early life with psychopathology outcomes in childhood, adolescence and adulthood.

Mediating Pathways - Epigenetics

An increasing number of studies are beginning to demonstrate the importance of epigenetic modification in mediating the risk of disease, including mental health outcomes. Epigeneticallymodified loci have been linked to a wide range of mental disorders such as schizophrenia, 40 as well as childhood onset disorders such as ADHD, 41 and ASD, 42 but conflicting and non-replicated associations mean that the causal relationships remain poorly understood. 43 LifeCycle mental health studies can currently analyse DNA methylation data on 14,368 offspring cohort participants (Figure 5), measured at birth (cord or placenta blood; N=7,783), childhood (0-12 years; N=3,055), adolescence (12-18 years; N=2,680), or adulthood (>18 years; N=850). Six of the thirteen contributing cohorts additionally contain longitudinal epigenetic data [ALSPAC, CHOP (multiple age groups in childhood), EDEN, Generation R, INMA, and RHEA]. The particular focus will be to identify epigenetic mechanisms that mediate the effect of early life exposures on behavioural and cognitive development, as well as mental health outcomes such as ASD, ADHD, depression and anxiety. This means it will be possible to track epigenetic changes in participants with behavioural and/or neurodevelopmental outcomes across time, and study causal relationships between environmental exposures in pregnancy or early life and later-life mental health outcomes mediated by DNA methylation.

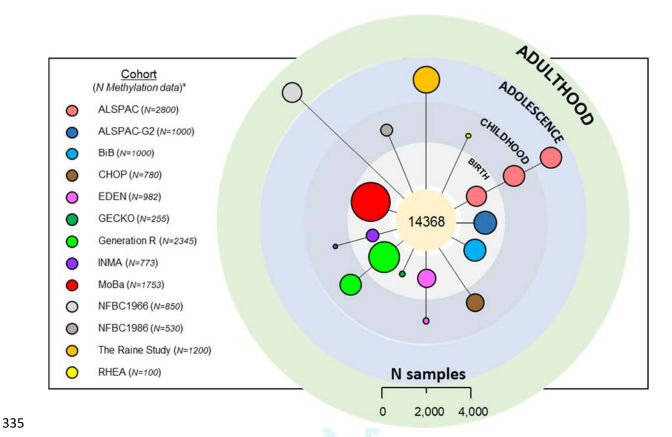


Figure 5. Overview of sample sizes for DNA methylation data in the offspring from birth to adulthood

Circle sizes are proportionate to the DNA methylation sample sizes as indicated in the scale at the bottom of the figure.

*Numbers relevant as of June, 2021 (sample processing and data collection is ongoing in several LifeCycle cohorts)

Framework for collaborative analyses

LifeCycle aims to perform most of the analyses through DataSHIELD. 44,45 With the recent launch of the platform and its analytical features for use with LifeCycle harmonised data, a number of novel collaborative studies have begun to form within the theme of mental health. Examples of planned and ongoing exposure-outcome analyses include infant feeding patterns and school-age externalising behaviours, maternal smoking in pregnancy and adverse child behaviours, associations between sleep, behaviour and cognition, sibling effects and prematurity, and socioeconomic inequalities and general mental health trajectories. Results from these studies are currently pending, but they have already shown that independent participant data resources have been successfully harmonised and can be coanalysed. The quantity and breadth of mental health and cognitive data available that have been mapped and harmonised by the LifeCycle mental health research group is a singular resource to enable developmental studies of mental health. These data will play an important role in replicating previous

findings with an enhanced statistical power, expanding upon previous associations through larger and more diverse samples, and in the development of novel models to describe how multi-faceted early-life exposures can shape and influence the landscape of mental health in later life.

Strengths and limitations

There are many strengths inherent in large consortia such as LifeCycle.¹ Key among these is that LifeCycle is building the EU Child Cohort Network, a sustainable research network that will enable continued exploitation of the LifeCycle data, metadata and collaborative progress beyond the usual timelines of a funded grant. Another important strength is the ability to study age differences and agerelated mental health and cognitive changes; this developmental aspect will help to understand the long- and short-term consequences of early life exposures, and how other factors such as epigenetic changes may mediate later health outcomes. Geographic diversity is also a key feature; it provides enhanced location coverage and generalizability of results, and also facilitates intra- and interpopulation comparisons. This makes it possible to make more reliable causal inferences due to different confounding structures.

The number of critical mental health domains covered is another strength, allowing for exposureoutcome research into many important and well-studied areas within this field. The availability of the
harmonisation protocols, coupled with the extensive overview of mental health measures, including
detailed information on the dimensions and age ranges across cohorts, provides users with an
integrated catalogue of psychological, cognitive, and psychomotor data in participating cohorts.
Furthermore, the use of DataSHIELD enables a flexible and data-secure approach that allows new
cohorts and centres to link into the analysis network and contribute with their own data, as well as the
addition of newly harmonised data as these are collected and updated. This open-source analysis
platform "takes the analysis to the data, not the data to the analysis", providing researchers with the
ability to remotely analyse data from multiple datasets without being able to access the data itself.^{45,46}
Removing the need to physically share data externally means participating cohorts bypass ethical
concerns related to the protection of privacy and other issues that arise when participant data are being

sent internationally to multiple users, and thus addresses some important ethico-legal considerations that are often associated with individual-level data sharing and analysis.

The heterogeneity of the psychological and cognitive measures available presents a potential limitation. Depending on the specific research question under investigation and measurement equivalence of constructs between different instruments, robust harmonisation 30,32 of certain measures may not be possible or may be limited to a small number of cohorts. This reduces the sample size or the range of participant ages that are possible to include. Within-country geographical bias of many of the cohorts may also present a weakness. Specifically, the urban-centric nature of many of the studies could mean that the generalizability of findings will be somewhat skewed, and the population-level inferences will need to take this bias into account. Furthermore, DNA methylation and brain imaging data are only available for less than 10% of total study participants. These smaller sample sizes may limit the number and strength of associations that can be found, as well as the distribution of participant ages and geographic and ethnic origins. However, the cohort studies are continuously expanding and adding new data on their participants, including phenotypic, genetic, epigenetic and biological data. The collaborative groundwork laid by LifeCycle will make it possible to continue building upon the analyses that have been performed, and help to mitigate some of the limitations that have been described.

Data Access

LifeCycle has developed an application procedure for data use proposals as described by Jaddoe et al.

It should be noted that approvals for data use and associated fees remain under the purview of the participating cohorts. This is the case regardless of whether one applies through LifeCycle or directly to the cohort, and these practices may vary across cohorts. The project strives to conduct as many analyses as possible within DataSHIELD. DataSHIELD is freely available to download and use (http://www.datashield.ac.uk/). This enables external cohorts to collaborate with LifeCycle and perform co-analyses. For more information, please visit the official website for the LifeCycle Project (https://lifecycle-project.eu/), or refer to the consortium design paper.

It is a described by Jaddoe et al.

It is hould be noted that approvals for data use and associated fees remain under the purview of the participant of the purview of the purview of the purview of the participant of the purview o

409	In some cases, data sharing and transfer agreements will need to be developed. These may vary due to
410	country-specific practices and restrictions as outlined by local General Data Protection Regulation
411	(GDPR) legislation. Application procedures directly to cohorts for data can be found at the following
412	websites:
413	
414	ALSPAC
415	http://www.bristol.ac.uk/alspac/researchers/access/
416	For more information on the ALSPAC cohort (including data dictionary, ethical considerations, and
417	funding), refer to Supplementary file 2.
418	BiB
419	https://borninbradford.nhs.uk/research/how-to-access-data/
420	СНОР
421	https://www.birthcohorts.net/birthcohorts/birthcohort/?id=137
422	DNBC
423	https://www.ssi.dk/English/RandD/Research%20areas/Epidemiology/DNBC/For%20researchers.aspx
424	EDEN
425	http://eden.vjf.inserm.fr/index.php/fr/contact
426	ELFE
427	https://www.elfe-france.fr/en/the-research/access-to-data-and-questionnaires/
428	GECKO
429	http://www.birthcohorts.net/birthcohorts/birthcohort/?id=138
430	The Generation R Study
431	https://www.generationr.nl/researchers/collaboration/
432	INMA
433	http://www.proyectoinma.org/presentacion-inma/politica-colaboracion/en_politica-colaboracion.html
434	HBCS
435	https://thl.fi/en/web/thlfi-en/research-and-expertwork/projects-and-programmes/helsinki-birth-cohort-
436	study-hbcs-idefix

437	MoBa
438	https://www.fhi.no/en/op/data-access-from-health-registries-health-studies-and-biobanks/data-from-
439	moba/research-and-data-access/
440	NFBC1966/1986
441	https://www.oulu.fi/nfbc/
442	NINFEA
443	https://www.progettoninfea.it/contact_us
444	The Raine Study
445	https://www.rainestudy.org.au/
446	RHEA
447	http://www.rhea.gr/en/research/data-access/
448	sws
449	https://www.mrc.soton.ac.uk/sws/
450	
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460	individual cohorts' funding information, refer to Supplementary File 3.
461	
462	Conflict of interest statement
463	The authors have no conflict of interest to declare.

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