



Children Growing Up in Liverpool Longitudinal Birth Cohort

We are inviting pregnant women/people and their partners to take part in a research study. Before you decide whether or not to take part it is important for you to understand why the research is being performed and what it will involve. Please take the time to read the following information carefully and discuss it with others if you wish. A member of our research team will go through this information sheet with you and answer any questions you may have. Please take time to decide whether or not you wish to take part.

Thank you for reading this.

Why are we doing the study?

Children Growing Up in Liverpool (C-GULL) is an exciting research study focused on improving the health and wellbeing of children and their families in the Liverpool City Region. It is the first new longitudinal birth cohort to be funded in the UK for almost 20 years. Longitudinal birth cohorts are large studies of babies born around the same time and followed up for many years. They are important as they can help us understand the connection between poverty, pregnancy and childhood, and the later development of illness. The C-GULL Study will collect information, including samples and data, from 10,000 children and their families, starting in pregnancy and continuing throughout life.

One of the main areas of focus of the C-GULL Study is mental health. We know that the first months of a baby's life are extremely important for how the brain develops, and that this can have a long-lasting impact on a person's mental health. There is also evidence suggesting that the bacteria in our gut play an important role in brain development and mental health. By collecting samples such as breast milk and stool (poo) from mothers/birthing parents and babies, the study aims to find out more about how these things are linked.

Who should take part?

We are inviting all women/people aged 16 years and older who are pregnant and planning for their care to be at the Liverpool Women's NHS Foundation Trust to take part in this study. We will also ask partners (if applicable) if they would like to take part.

We want to involve families from different ethnic, cultural, social and family backgrounds so that we can understand how to improve the health and wellbeing of all children in the Liverpool City Region.

Do I have to take part?

No, you do not have to take part. You can choose whether to take part or not. Even if you agree now, you can choose to stop at any time. If you decide not to take part, your healthcare will not be affected.

If you decide to take part in the study, you will be able to choose which aspects you would like to be part of. It is up to you to decide what you feel happy to be involved with. Before making any decisions, you must take time to read this information sheet carefully and discuss any questions you may have with the research team.

What should I do if I want to take part?

If you would like to take part in the C-GULL study you can speak to the research team located in either the Antenatal or Ultrasound (scanning) Department when you are next visiting the Liverpool Women's Hospital. Alternatively, you can register your interest online and the research team will contact you by telephone. A member of the team will talk to you about the study, answer any questions you may have and ask you if you would like to take part. You can take as









must time as you need to decide. If you are happy to take part in the study, you will be asked to make an appointment for your first assessment with one of our Research Midwives. You can arrange this appointment at a time that is convenient for you. Before taking part in the study you will need to complete and sign the study consent form. The research team will be available to support you with this process, if needed.

What will happen to me if I take part?

If you agree to take part in the C-GULL Study, we will ask you to:

- Visit the research centre at Liverpool Women's Hospital or one of our community centres twice while you are pregnant, once in early and once in late pregnancy. At the first visit you will be asked to answer some questions, complete some questionnaires, and give a small amount of blood, urine, and hair. At the second visit you will be asked to complete some questionnaires, give a small amount of blood and take a kit home to collect a small sample of your poo. At this visit you will also be given some kits to collect samples from yourself and your baby at home after your baby is born.
- Provide some samples. Around the time of birth we will ask you to donate some samples, including your placenta (afterbirth), some blood left in the umbilical cord (cord blood) and a small amount of colostrum (first breast milk). Please be reassured, taking blood from the umbilical cord will not cause any pain or distress to your baby. After your baby is born, we will ask you to collect some samples of your breast milk and your baby's poo using the kits provided at your second visit to the research centre. For more detailed information about the samples we will be collecting and how they will be used, please refer to our Supplementary Participant Information: Taking Part in the Study / Microbiome
 - Shortly after birth, we will also ask if we can measure your baby's body composition (the amount of fat, bone, water, and muscle in the body) using a piece of equipment called a 'PEA POD'. Your baby will need to lie in a special incubator for a short time whilst measurements are being taken, however this should not cause any pain or discomfort. For more detailed information about the PEA POD and what will happen to your baby, please refer to our Supplementary Participant Information: PEA POD.
- Answer some questions on your and your baby's health, when your baby is around the ages of 3 months and 12 months. We will do this by email, text or by sending you a letter, whichever you prefer.
- Visit one of our community centres when your baby is two years old so that we can see how they are growing and developing. You will also be able to ask any questions you may have about your health or your baby's health and development.
- Agree to be contacted again in the future. We will contact you again in the future to invite you and your baby
 to take part in follow up assessments (answer more questions and / or attend more visits). It is your choice
 whether to take part in each of these. You will be provided with full information for each and given time to
 decide if you would like to be involved. If you decide not to take part, the care you or your family receives will
 not be affected in anyway.
- Give permission for us to request information that the NHS, government departments and other organisations already hold about you and your baby. We would like to look at this information so that we can understand how health and other life events are related. This will only be done with your permission. You will need to agree to us getting some information from your NHS, healthcare and birth records, but anything more than this will be optional.

For more detailed information about what will happen at each study visit, please refer to our Supplementary Participant Information: Taking Part in the Study.









During your first assessment, we will ask your permission to contact your partner (if applicable) about taking part in the study. If they agree, we will provide a link to our website where they can find more information. They will be able to register their interest online and shortly after, they will be able to consent and fill in the study questionnaires via our online systems. If they prefer to do this face to face, we can arrange for this also. Partners will also be asked if they would like to provide a blood or saliva sample. If they agree, we will arrange for them to come into the Research Centre where blood can be drawn or a saliva sample collected by a member of our Research Midwife team. If preferred however, they can take the saliva sample themselves using a home collection kit. If your partner does not wish to take part in the study, that is completely fine. For more detailed information about partners taking part in the study, please refer to our Partner Information Sheet.

Will my taking part in this study be kept confidential?

Yes, we will follow ethical and legal guidance and all information collected about you and your baby will be kept confidential and secure. Any personal information you give us will only be used by the research team during the course of the study. All samples and data will be coded (given a unique identification number) and stored separately from your personal details (name, date of birth, contact details etc.) to protect your identity. Only members of the research team who need to contact you will know your name and contact details. To prevent unauthorised access to your personal details we will use things like encryption, password protected computers systems, key card controlled rooms and lockable storage cabinets to make sure your information is stored securely.

How will you use my data?

We will need to use information from you and from your and your baby's medical records for this study. This information will include identifiable data, such as name, initials, date of birth, NHS number and contact details (telephone number, email address, house address and postcode). We will keep any identifiable data collected for at least 10 years after the study has finished.

In addition to identifiable data, the C-GULL Study will also collect demographic, health (hospital, GP, midwife / health visitor community), lifestyle, nutrition, housing, education, employment, social care and economic (income) information. A lot of this information will be obtained from the questions you will answer / questionnaires you will complete (self-reported data). However, we would also like to access health, education, social care and economic information that the NHS and government departments (Education, Health and Social Care, Work and Pensions) already hold about you and your baby. This is called linked data and we would only do this with your permission.

Data linkage in a longitudinal study such as the C-GULL Study means collecting information from different sources and using it to track the health and development of a person throughout their life. To link all this information together, identifiable information like their name and date of birth is used. This helps researchers to identify patterns in the data and understand what factors might contribute to them. Data linkage in a longitudinal study usually starts from when the person was born.

To ensure that your data is handled with the utmost care and respect, the C-GULL Study starts data linkage from the point of your consent. This means that your NHS number and date of birth will only be used to link information collected after you have agreed to participate in the study. By doing so, we can help protect your privacy and ensure that your data is handled securely.

Dynamic consent is a process that enables individuals to have control over their involvement in research studies. In the C-GULL Study, we would like to stay in touch with you after your child reaches the age of two years to obtain your consent for continued involvement in the study. This means that you can choose to opt out of any further data collection or analysis if you prefer. We believe that by using dynamic consent, we can establish a dialogue with you and ensure









that you're informed about the study's progress and how your data is being used. This approach allows us to maintain transparency and trust with our participants and respects their privacy.

You can find out more about how we use your information

- at www.hra.nhs.uk/information-about-patients/
- from the leaflet available at www.hra.nhs.uk/patientdataandresearch
- by asking one of the research team
- by sending an email to <u>legalservices@liverpool.ac.uk</u>, or
- by ringing us on 0151 795 0523.

For more detailed information about the data we are collecting and how it will be used, please refer to our Supplementary Participant Information: Data.

What are the possible benefits of taking part?

Taking part in the C-GULL Study will allow the development of your child to be tracked throughout their childhood and teenage years, to adulthood and beyond. The samples and data collected from you and your child will allow us to better understand the factors that can improve the lives and health outcomes of the children (and their families) within our city.

What are the possible risks of taking part?

The risks involved in the study have been carefully assessed and the main objective of the research team is to maintain yours and your baby's safety at all times. Taking part in the study will not put you or your baby at any risk of harm. The samples collected have been used with pregnant women/people before and are not known to have any risks. The questions / questionnaires you will be asked to answer should not make you feel uncomfortable and you can decide whether to answer them or not. However, if you should experience any discomfort at any time during the study, please tell a member of the research team immediately. If the team member has any serious concerns about your health, wellbeing or safety, they will ensure that the appropriate support is provided to you via the Liverpool Women's NHS Foundation Trust safeguarding pathway. There are also contact details provided at the end of this information sheet for organisations who can offer support should you need it.

What will happen if I do not want to continue in the study?

You are free to withdraw at any time throughout the course of the study, without explanation. The care you or your family receives will not be affected in anyway. If you withdraw, you can ask the C-GULL Study for:

- No further contact: we would not contact you again, but would have your permission to keep and use information and samples that you have already given and to request further information from your health and other records in the future.
- No further access: we would not contact you again and would not request information from your health and other records in the future. We would still however have your permission to keep and use the information and samples you have already given us.
- **No further use:** we would no longer contact you and would not request information from your health and other records in the future AND any information and samples already collected would no longer be given to researchers.

Can I take a break from the study?

Yes, if you feel you need a break you can opt out for a while. During this period we will not contact you, but would have your permission to continue to collect information from your health and other records. Once the break is over, we will get back in touch to check that you are happy to take part in the study again.





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What will happen to any samples I give?

Sample collection for the C-GULL Study is optional. You can choose which samples you would and would not like to give. Any samples you do give will be collected specifically for the study. For most of the samples you provide, there will be no tests performed immediately. They will be stored so that in the future researchers will be able to use them to better understand the link between early-life and later poor health. We will use some of the breast milk and poo samples you and your baby provide straight away. We will use these samples to look at the bacteria that live in the gut and the nutrients in milk to find out more about the role they play in brain development and mental health.

All samples will be coded and stored separately to any personal details (name, date of birth, contact details etc.) you provide to us. We will make sure your samples are handled properly and ethically and stored securely throughout the course of the study.

If you agree, your samples will be used by approved researchers in specialised laboratories in the UK and laboratories abroad with specific ethical approval in place. As a result, some of your samples and data may be transferred out of the UK. We will ensure that they are sent in a fully anonymised format so that you and your baby can not be identified.

If you give permission, we will extract your DNA and / or your RNA from some of your samples for use in human genetic studies. If you agree to gift (donate) your samples, they will be used in other ethically approved research in the future.

For more detailed information about the samples we are collecting and how they will be used, please refer to our Supplementary Participant Information: Taking Part in the Study / Microbiome.

What will happen to the results of the research study?

It is intended that the results generated from the study will be published as research papers in medical journals. We will also engage with national and charitable organisations. Summaries of the study results will also be made available on our website and social media.

Where can I get further information or discuss any problems?

If you have any questions or concerns about any aspect of this study, please contact a member of the C-GULL Study research team on 0151 795 6700. If your concerns are not resolved, you can contact the Patient Advisory Liaison Services (PALS) on 0151 702 4353. You can also visit PALS by asking at the Liverpool Women's Hospital main reception.

Who is organising and funding the research?

Wellcome Trust is funding the study (Reference: 217067/Z/19/Z) and Professor Louise Kenny is the study Chief Investigator. The study is sponsored by The University of Liverpool and managed by the Harris Research Centre, University of Liverpool.

Who has reviewed the study?

All research in the NHS is looked at by an independent group of people, called a Research Ethics Committee, to protect your interests. This study has been reviewed for ethical considerations and given a favorable opinion by members of the North East - Newcastle and North Tyneside 1 Research Ethics Committee.









Where can I find further information?

Should you need any further information about the study, please contact us on cgullstudy@liverpool.ac.uk or 0151 795 6700. You can also visit our website cgullstudy.com or follow us on social media @ChildrenGrowing.

You can register your interest in the study by visiting our website using the link above or the QR code below and completing the registration form within the 'Taking Part' section.



Organisations who offer support should you need it:

- Mersey Care (urgent mental health support): https://www.merseycare.nhs.uk/urgent-help
- Anxiety UK: https://www.anxietyuk.org.uk/anxiety-type/stress/
- Merseyside Domestic Violence Service: https://www.mdvs.org/
- Pandas (postnatal depression support): https://pandasfoundation.org.uk/
- Bambis (breastfeeding support): https://www.liverpoolbambis.co.uk

Thank you for taking the time to read and consider this information sheet. Should you decide to take part in the study, you will be given a copy of the information sheet and a signed consent form to keep.



