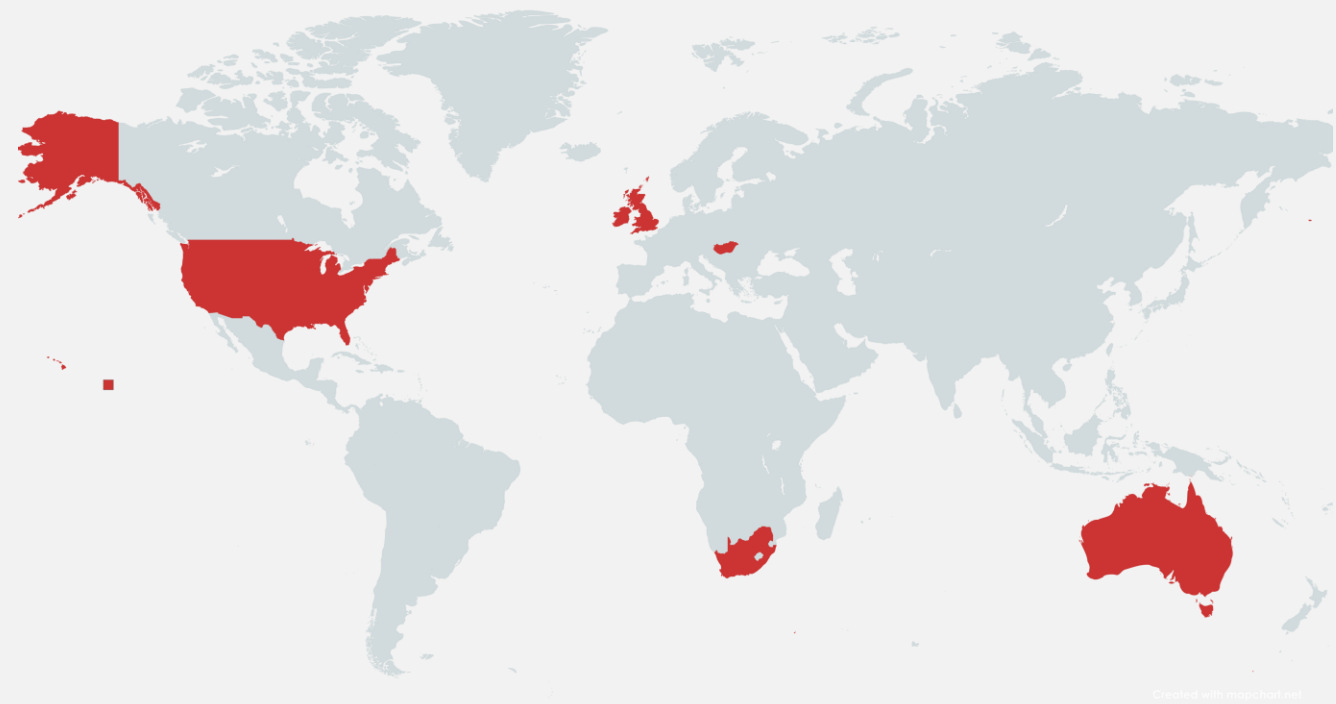
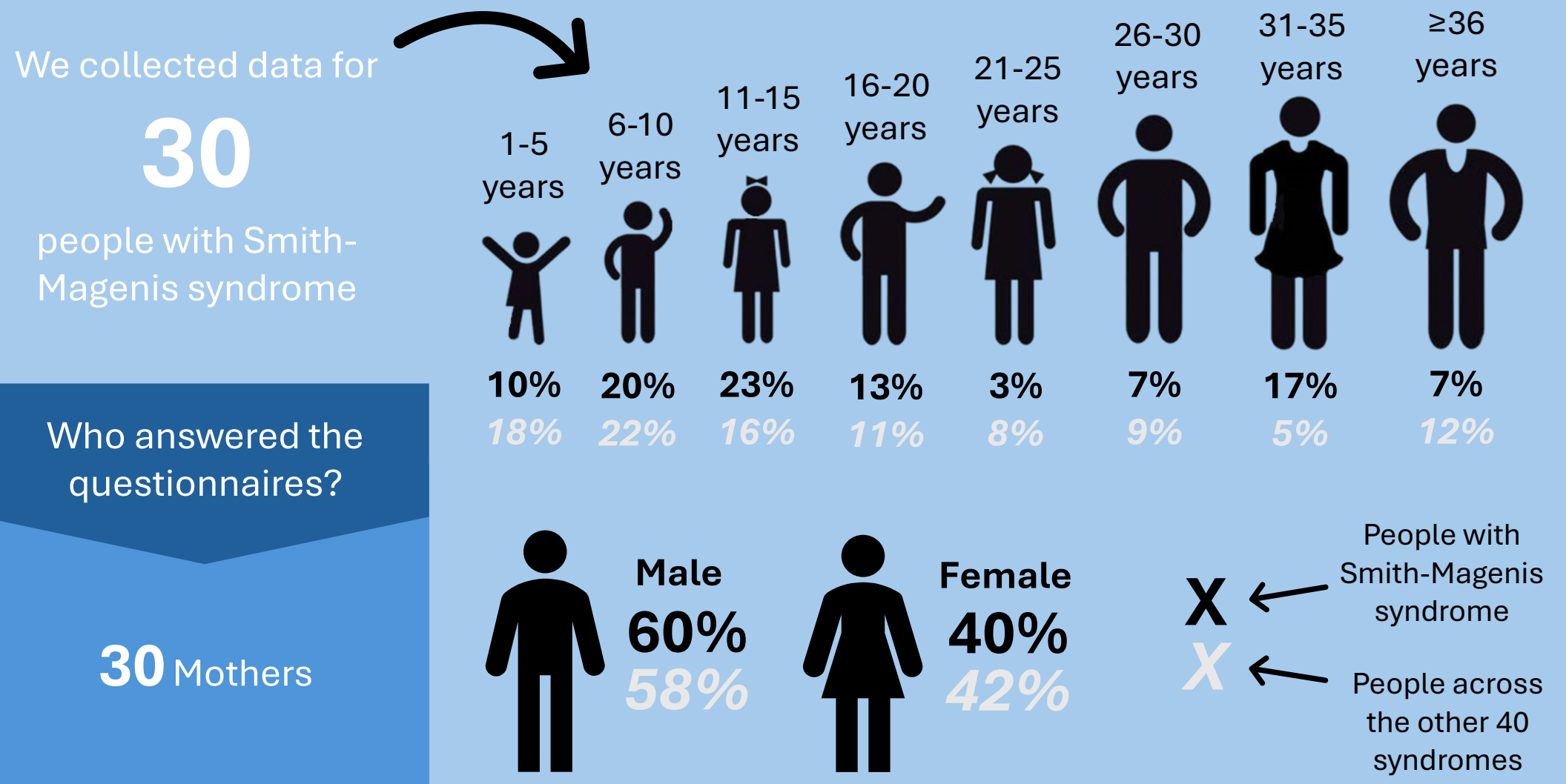


Behavioural and Emotional Outcomes in Neurodevelopmental Disorders (BEOND)

Study summary report



For the BEOND study, we recruited **627** parents and caregivers of people with rare genetic syndromes. These parents and caregivers represented **41** syndrome groups, one of which was Smith-Magenis syndrome. We asked parents and caregivers to complete questionnaires about the person they care for. Below are some facts about the study.



- Where were families from?
- 24 United Kingdom
 - 2 United States of America
 - 1 Australia
 - 1 Hungary
 - 1 Ireland
 - 1 South Africa

We collected data between Nov 2022 – Jan 2024



Questionnaires completed online: **24**



Questionnaires completed via mail: **6**

30 personalised feedback reports were sent to families summarising their data

The results of the study were shared at **15** conferences, syndrome-support events, and invited talks

The data you provided helped us to...

- 1 Develop a new questionnaire to assess anxiety in people with intellectual disabilities.
- 2 Investigate the importance of sleep for behaviour, health, and psychological outcomes in people with intellectual disabilities.
- 3 Further understand parent-carer wellbeing and how this relates to the support received from family doctors.

The BEOND study will open again to families in 2026!

Find out everything you need to know below.



Meet the team!



Prof Caroline Richards



Dr Hayley Crawford



Dr Joanna Moss



Dr Jane Waite



Dr Rory Devine



Dr Rory O'Sullivan

With BEOND 2026, we aim to:

Create a new measure of everyday functioning for people with rare genetic syndromes and intellectual disabilities.

Explore quality of life in people with rare genetic syndromes.

Investigate changes over time in sleep, anxiety, autism characteristics, and behaviour for people with rare genetic syndromes.

In response to families' feedback, we've made a few changes to BEOND 2026.



A 'skip' button has been added in case some questions are tricky to answer.

1/2

The survey is now 50% shorter.



We streamlined our feedback report processes to ensure these are sent to families quicker.

What you need to know before taking part...

Who can take part?

All families with Smith-Magenis syndrome are welcome to take part! We welcome families who previously took part in BEOND as well as new families.

What do I get for taking part?

All families that take part will receive a personalised feedback report which summarises their data. For families who took part in BEOND last time, we will link their past data with their current data.

Families will also be offered a £20 voucher upon completion of the survey.

What does taking part involve?

Parents/carers will complete a survey, either online or via a posted paper copy. The survey includes questions about your child/person you care for. A range of topics are covered such as health, behaviour, wellbeing, and cognition.

This information will remain confidential.

When will I be able to take part?

We intend to open the study to families in early 2026. There will be further updates towards the end of 2025.

Where can I find more information about BEOND 2026?

You can find out more about BEOND 2026 on the Cerebra Network for Neurodevelopmental Disorders website: <https://www.cerebranetwork.com/beond>

You can also email any queries to Dr Rory O'Sullivan at r.osullivan@bham.ac.uk