





CARER/FAMILY MEMBER INFORMATION SHEET & INFORMED CONSENT FORM

SCARF-BT

Social Cognition Assessment and Rehabilitation for Families Living with Brain Tumour (SCARF-BT) - Stage 2

You are being invited to take part in a research study. Before deciding whether to take part, you need to understand why we need to do this research and what it involves for you. We'd like you to take time to read the following information carefully and talk to others about the study if you wish. Please ask us if anything is not clear or if you would like more information. Please take time to decide whether or not you wish to take part.

Section 1 tells you the purpose of this study and what will happen to you if you take part.

Section 2 gives you more detailed information about how the study is carried out.

Section 1: Purpose of the study and what will happen

1. What is the purpose of the study?

Brain tumours can affect how the brain works. This can make interacting with others difficult, including friends and family, by the effect they can have on how the brain works. Detecting emotions in friends and family is a key part of relationships and communication. One way we recognise emotions in others is by looking at facial expressions. The effects of a brain tumour can make this recognition difficult, making it harder to engage with others. This in turn can leave patients with brain tumours, and their carers feeling isolated and alone.

Similar problems can occur in patients with head injuries and strokes. A computer-based rehabilitation method has been shown to improve the ability to recognise emotions for these patients. We think such rehabilitation may help with similar difficulties in patients with brain tumours. In this study we want to see if this computer-based rehabilitation, called 'FACES' can be used by patients with brain tumours. In the future, we hope to help patients re-learn how to recognise emotions in others and improve their quality of life by enjoying easier interactions with friends and family.

2. What is the treatment being tested?

We would like to test 'FACES', a computer-based rehabilitation package. We know it provides benefit for patients with head injury and strokes who also have difficulties with recognising emotions. We will provide 'FACES' or a general brain training rehabilitation computer package for patients taking part. Patients will be shown how to use the computer package with the help of a trained assistant psychologist, so that they can use it at home for 3 one hourly sessions per week for 3 weeks.

We will see if patients can use 'FACES' or a general brain training rehabilitation computer package after having surgery for a brain tumour. We also want to see if there are any improvements in the ability to recognise emotions and day-to-day function after using the



This study is funded by the National Institute for Health and Care Research (NIHR) [RfPB Programme (NIHR200495]). The views expressed are those of the author(s) and not necessarily those of the NIHR or the Department of Health and Social Care.





computer packages. This will be assessed with tests of the ability to recognise emotions before and after using the computer packages and also by interviews with patients, their carers and family members to see what effect, if any, the computer-based rehabilitation had on their day-to-day interactions.

Where necessary we will provide patients with a computer, for the duration of the study, to allow them to do the interventions.

3. Why have I been invited?

You have been invited to take part in this study because a family member/someone you care for has a brain tumour and we believe computer based rehabilitation may offer relief to patients with symptoms, especially those who find it more difficult to interact with others (including friends and family) since having the diagnosis of a brain tumour.

We are inviting patients that have been involved in the CogENT and/or SIND studies.

4. Do I have to take part?

No. Taking part in this study is completely voluntary. If you decide to take part you will be asked to sign an Informed Consent Form, however you are still free to change your mind and leave the study at any time without giving a reason.

5. What will happen to me if I take part?

If you agree to take part in the study, you will sign the Informed Consent Form at the end of this document and be given a copy of this to take away and refer to later.

Once you have signed a consent form to join this study, we will ask you to complete a quality of life questionnaire. The purpose of this questionnaire is to help us understand the impact of a brain tumour on the carer/family. We will ask your family member / person you care for to complete some tests to see if they have any difficulties recognising emotions.

Then our trained assistant psychologist will arrange to help your family member / person you care use 'FACES' or a general brain training computer package at home, when they are discharged from hospital to recover after surgery. They will use computer based rehabilitation 3 times a week for 3 weeks, each session lasting one hour. Whether your family member / person you care for gets to use 'FACES' or a general brain training rehabilitation will be randomly chosen, by means of a computer program, in order to keep the comparisons between the two as fair as possible. Your family member / person you care for will have a 50% chance of receiving 'FACES' or general brain training rehabilitation. Half the patients taking part in this study will receive the FACES intervention and half will receive general brain training.

After these 3 weeks of recovering at home and using the computer packages, we will repeat the tests for difficulties in recognising emotions and the quality of life questionnaires to find out if computer based rehabilitation has helped. We will also ask you to complete another quality of life questionnaire.

Following this, we may contact you to arrange some interviews and/or focus groups, where we would discuss your experience of your family member / person you care for, using the computer rehabilitation and any effect it had on their day to day interactions. Participation in interviews and focus groups is optional and if you are interested in taking part we will provide a separate information sheet. We will need to give your contact details to the researcher conducting the interviews (an NHS Clinical Psychologist). This







researcher will then contact you to answer any questions, ask you to sign the separate consent form and arrange the interviews and/or focus groups.

At the end of this study, you and your family member/person you care for will be offered a final feedback session to discuss any problems you are having and possible methods of addressing your problems.

This study will not require any invasive procedures or changes in drugs and routine treatment for your family member/ person you care for. The only difference is the use of this computer package during a time when patients are not routinely seen by hospital staff as they are recovering from surgery at home.

As this is not a study of a drug or invasive treatment, we do not expect your family member/person you care for to feel unwell or any side-effects from using the 'FACES' or general brain training computer-based rehabilitation.

You should tell the study team if your family member/person you care for feels unwell or different in any way. If you have any major concerns please contact the study doctor immediately using the contact numbers at the end of this information sheet.

6. What are the possible disadvantages and risks of taking part?

Although we do not expect any additional hospital visits for your family member / person you care for, these visits may be longer. This is due to the completion of quality of life questionnaires by both you and your family member/ person you care for. There will be some extra tests to look for any difficulties recognising emotions.

Quality of life questionnaires will be completed before the clinic visit so that any possible distress caused by the topics raised can be addressed by the clinical research team during the clinic visit.

If during the interviews with you and the patient about their experience of using 'FACES' or the general brain training, other incidental concerns are raised, we will find the appropriate support to address them.

Attending a hospital during the Pandemic

Although there is a risk of hospital COVID-19 transmission, we do not think the risk is significant. Local policies are in place for patients attending a hospital for planned care to minimise the risk of disease transmission.

7. What are the possible benefits of taking part?

There is no guarantee that your family member / person you care for will benefit from taking part in this study. Although we hope they may experience relief in their symptoms or improvement in their experience of interactions with family and friends, this study is not designed to show any possible benefit. However, information collected as part of your taking part in this study may benefit patients with brain tumours in the future.

Previous research has shown that participants find completing quality of life questionnaires helpful in getting them to think about their problems and thinking about what they wanted to discuss in outpatient clinics.

We are offering all participants (patients and their family member/ carer) the opportunity to attend a final feedback session. Participants in previous research have found this helpful.



This study is funded by the National Institute for Health and Care Research (NIHR) [RfPB Programme (NIHR200495]). The views expressed are those of the author(s) and not necessarily those of the NIHR or the Department of Health and Social Care.





8. What are the alternatives for treatment?

Currently there are no other alternative or standard treatments for patients with brain tumours who have difficulties with recognising emotions.

9. What happens when the study stops?

Your family member/ person you care for will return to normal standard of care once all the study assessments have been completed.

10. Expenses & Payment?

You will not receive payment for taking part in this study. We aim to perform the tests required during routine hospital visits, however some of these visits may be longer, we will reimburse any additional parking costs incurred.

This completes Section 1 of the information sheet.

If you are considering taking part in the study, please continue to read the additional information in Part 2 before making any decisions.







Section 2: Study Conduct

Please bear in mind that the following is information that we must provide to all patients and carers involved in all research studies.

11. What if new information becomes available?

Sometimes during the course of a study, new information becomes available which might affect your decision to continue taking part in this study. The study doctor will contact you to discuss the new information and whether you wish to continue taking part in the study. If you still wish to continue on the study, you will be asked to sign a new Informed Consent Form.

The study sponsor, the regulatory authority or the study doctor may decide to stop the study at any time. If that happens we will tell you why the study has been stopped.

12. What if I decide I no longer wish to take part in the study?

You are free to come off this study at any time without giving a reason. If you decide not to take part any further, you will no longer be asked to complete questionnaires or invited for interview. Any data (which is not identifiable) already collected will continue to be used in the study analysis.

The study doctor may also choose to withdraw you from the study if they feel it is in your best interests or if you have been unable to comply with the requirements of the study. A reason for study withdrawal could be you are unable to complete the visits or study documentation as required

13. What if there is a problem?

Any complaint about the way you have been dealt with during the study or any possible harm you might suffer will be addressed. If you have any concerns about any aspect of this study you should speak to the study doctor who will do their best to answer your questions.

In the event that something does go wrong and you are harmed by taking part in the research and this is due to someone's negligence then you may have grounds for a legal action for compensation against Cambridge University Hospitals NHS Foundation Trust or the University of Cambridge. The normal National Health Service complaints mechanisms will still be available to you (if appropriate). The University has obtained insurance which provides no-fault compensation i.e. for non-negligent harm, you may be entitled to make a claim for this

If you wish to complain or have any concerns about any aspect of the way you have been approached or treated during this study, you can do this through the NHS complaints procedure. In the first instance it may be helpful to contact the *Patient Advice* and *Liaison Service (PALS)*.

14. Will my taking part in this study be kept confidential?

Cambridge University Hospitals NHS Foundation Trust (CUH) and the University of Cambridge are the Sponsors for this clinical study based in the United Kingdom. They will be using information from you in order to undertake this study and will act as the data controller for this study. This means that they are responsible for looking after your information and using it properly. The Sponsor organisations will keep identifiable information about you for up to 15 years after the study has finished to ensure your safety and allow the study to be reviewed by the authorities after it is finished.







Your rights to access, change or move your information are limited, as the Sponsor organisation(s) need to manage your information in specific ways in order for the research to be reliable and accurate. To safeguard your rights, we will use the minimum personally-identifiable information possible.

You can find out more about how the Sponsors use your information using the information below:

- For Cambridge University Hospitals NHS Foundation Trust, please visit: https://www.cuh.nhs.uk/corporate-information/about-us/our-responsibilities/looking-after-your-information, or email the Data Protection Officer at: gdpr.enguiries@addenbrookes.nhs.uk
- For University of Cambridge, please visit: https://www.medschl.cam.ac.uk/research/information-governance/, or email the Information Governance team at: researchgovernance@medschl.cam.ac.uk

Cambridge University Hospitals will collect your name and contact details to contact you about this study, and make sure that relevant information about the study is recorded and to oversee the quality of the study. Individuals from the Sponsors and regulatory organisations may look at your research records to check the accuracy of this study. Cambridge University Hospitals will pass these details to the Sponsors along with the information collected from you. The only people in the Sponsor organisations who will have access to information that identifies you will be people who need to contact you in relation to this study and to audit the data collection process. Cambridge University Hospitals will keep identifiable information about you from this study for up to 15years after the study has finished.

All information collected about you as a result of your taking part in the study will be kept strictly confidential. Your personal information (e.g. age, gender assigned at birth, ethnicity, relationship to the person with brain tumour, length of time caring) will be kept in a secured file and be treated in the strictest confidence.

Once you have agreed to take part in this study you will be allocated a Study ID Number that links you to your family member/ person you care for. This study number will be used on all your study documentation.

The people who analyse the information will not be able to identify you and will not be able to find out your name or contact details. Only anonymous study data, without any personal information will be published at the end of the study.

When you agree to take part in this study, the information directly related to you taking part in this study may be provided to researchers running other research studies in this organisation and in other organisations. These organisations may be universities, NHS organisations or companies involved in health and care research in this country or abroad. Your information will only be used by organisations and researchers to conduct research in accordance with the UK Policy Framework for Health and Social Care Research.

Your information could be used for research in any aspect of health or care, and could be combined with information about you from other sources held by researchers, the NHS or government. Where this information could identify you, the information will be held securely with strict arrangements about who can access the information. The







information will only be used for the purpose of health and care research. It will not be used to make decisions about future services available to you, such as insurance. Where there is a risk that you can be identified your data will only be used in research that has been independently reviewed by an ethics committee.

15. What will happen to the results of the study?

The results of the study will be anonymous and you will not be able to be identified from any of the data produced. When the results of this study are available they may be published in peer reviewed medical journals and used for medical presentations and conferences. They will also be published a Clinical Studies Register website, a central registry for all conducted clinical studies.

Anonymous datasets from the study may also be made available to other researchers in line with national and international data transparency initiatives.

If you would like to obtain a copy of the published results please contact your study doctor directly who will be able to arrange this for you. We are planning to feedback your results to you after you complete the study.

16. Who is funding the study?

The study is being funded by the National Institute for Healthcare Research under their remit of 'Research for Patient Benefit' (Grant Reference: NIHR200495).

The study is managed by the Cambridge Clinical Trials Unit - Cancer Theme and sponsored by the Cambridge University Hospitals NHS Foundation Trust and the University of Cambridge.

17. Who has reviewed this study?

All research within the NHS is reviewed by an independent group of people called a Research Ethics Committee, to protect your interests. This study has been reviewed and given favourable opinion by Cambridge South Research Ethics Committee.

18. Further information and contact details

If you require any further information or have any concerns before/while taking part in this study, please contact a member of the study team during normal office hours (9:00am – 5:00 pm):

CNS Study Research Nurse.

Name: XXXXXX Tel: XXXXXX

Study Doctor/Principal Investigator

Name: XXXXXX Tel: XXXXXX

24-hour contact

Hospital/Ward: XXXXXX Tel: XXXXXX







Alternatively, if you or your relatives have any questions about this study, you may wish to contact one of the following organisations that are independent of the hospital:

Brainstrust is a UK based charity and community which supports and helps thousands of patients and carers who are affected by a brain tumour diagnosis. They work to help patients and carers regain control and be confident that they are working towards the best outcome for their situation. They do this by sharing understanding of their local services, by providing our own services that include 24/7 email and telephone support, practical resources such as our brainbox and personalised resources that help our community feel a little less lonely. For more information have a look at www.brainstrust.org.uk or email hello@brainstrust.org.uk

The Brain Tumour Charity (originally called the Samantha Dickson Brain Tumour Trust) was set up in 1996. It is the UK's largest dedicated brain tumour charity, funding scientific and clinical research into brain tumours. It also offers support and information to brain tumour patients, whilst raising awareness and influencing policy. You can contact the organisation for advice, information or support on 0808 800 0004. You can access the information leaflets, discussion forum and support group details on the website at http://www.thebraintumourcharity.org/

MACMILLAN CANCER SUPPORT: a registered charity providing information about all aspects of cancer for cancer patients and their families. They have published several useful booklets on different types of cancer, chemotherapy, radiotherapy, and clinical studys in general. These booklets may be requested from Macmillan Cancer Support, 89 Albert Embankment, London SE1 7UQ. Alternatively, you may view the contents of these booklets on their website (www.macmillan.org).

In addition, MACMILLAN CANCER SUPPORT also provide advice from specialist cancer nurses on: FREEPHONE 0808 808 0000 (9am to 8pm, Monday to Friday, excluding Bank Holidays).

Contact details for Patient Liaison and Advice Service (or equivalent)

Addenbrooke's Patient Advice & Liaison Service

Tel: 01223 216756

Email: pals@addenbrookes.nhs.uk

Address: Box 53, Cambridge University Hospitals, Cambridge Biomedical Campus,

Hills Road, Cambridge, CB2 0QQ







CARER/FAMILY MEMBER INFORMED CONSENT FORM

Study Title: Social Cognition Assessment and Rehabilitation for Families Living with Brain Tumour (SCARF-BT) - Stage 2

_	agree with each sentence below, please initial the box	INITIAL
1	I have read and understood the Carer / Family member	
	Information Sheet version 2.0 dated 25 th April 2022, for the above	
	study and I confirm that the study procedures and information	
	have been explained to me. I have had the opportunity to ask	
	questions and I am satisfied with the answers and explanations	
	provided.	
2	I understand that my taking part in this study is voluntary and that	
	I am free to withdraw at any time, without giving a reason and	
2	without my legal rights being affected.	
3	I understand that personal information about me will be collected and used in accordance with this information sheet. This	
	information will be kept in the strictest confidence and none of my	
4	personal data will be published. I understand that information related directly to my taking part in	
4	this study may be looked at by responsible individuals from the	
	sponsors, regulatory authorities and research personnel where it	
	is relevant to my taking part in research and that they will keep	
	my personal information confidential. I give permission for these	
	individuals to have access to my records.	
5	I have read and understood the compensation arrangements for	
•	this study as specified in the Carer / Family Member Information	
	Sheet.	
6	I understand that the doctors in charge of this study may close	
	the study, or stop my taking part in it at any time without my	
	consent.	
7	I have read and understood my responsibilities for the study.	
8	I understand that the data and information collected about me will	
	be used to support other research in the future and may be	
	shared anonymously with other academic and commercial	
	researchers external to the project, within the UK and beyond.	
	ptional Consent	
ξ	I am interested in taking part in interviews/focus groups and	Yes / No
	agree to my contact details being shared with the researcher	
	conducting interviews/focus groups	
gre	e to take part in this study:	
•	·	
o of	carer Signature	 Date
ie oi	Care Signature	Date
ne of	person taking consent Signature	Date
e of (Consent (24hr clock):	
	ne carer, 1 copy for the study team, 1 copy to be retained in the hospital notes.	
y iOi ti		

This study is funded by the National Institute for Health and Care Research (NIHR) [RfPB Programme (NIHR200495]). The views expressed are those of the author(s) and not necessarily those of the NIHR or the Department of Health and Social Care.