

**Brain Injury  
MedTech Co-operative**

  
**National Institute for  
Health Research**

# **Register for Healthcare Involvement and Technology Evaluation**

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# Areas of public involvement

A horizontal bar composed of several colored segments: green, dark green, orange, purple, red, dark blue, and light blue.

## Informing of latest updates, progress and innovation

Newsletters

Tailored information

## Involvement in research evaluation and prioritisation

Workshop participation

Survey completion

Roadmapping

## Research participation

Feasibility studies and pilots

Clinical trials

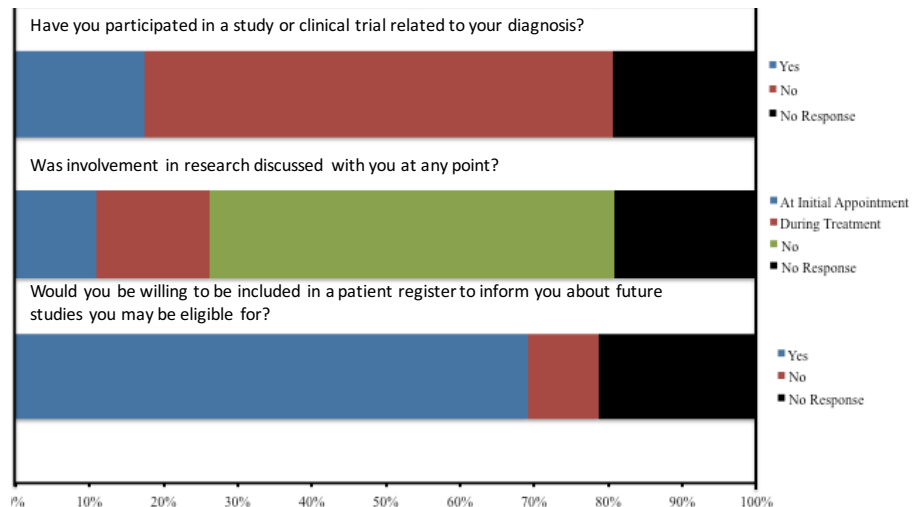
# Unmet clinical need

## A neuro-oncology patient/carer perspective



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Question	Positive Response	Neutral Response	Negative Response
At your first appointment how well were your symptoms addressed	65%	17%	18%
At your first appointment, how much opportunity did you get to ask questions	71%	11%	18%
Before seeing a specialist, how well were you update/kept informed	29%	13%	47%
At the first appointment, how well did you understand the information provided	41%	34%	25%
How useful would a structured questionnaire recording your symptoms and concerns at your first appointment be	74%	11%	15%



# Register for Healthcare Involvement and Technology Evaluation (RHITE)



National Institute for Health Research

The screenshot shows the registration form for the NIHR Brain Injury Healthcare Technology Co-operative (HTC). It includes the logos for the University of Cambridge and the National Institute for Health Research. The form title is "Register for Healthcare Involvement and Technology Evaluation". The text explains that the HTC has developed a volunteer register for patients, carers, and other interested persons to assist in the development and advance of healthcare technologies. It provides the website [www.brainhtc.org](http://www.brainhtc.org) for more information. A note states that the patient and carer register is delivered in partnership with the Outcome Registry Intervention and Operation Network (ORION), hosted within the University of Cambridge. The registration fields include: Title (dropdown), Forename, Surname, Gender (radio buttons for Male, Female, Other), Date of birth (DDMMYY), Email, and Confirm email. There are also checkboxes for "How else would you prefer us to contact you?" with options for "by post" and "by telephone".

## Role

- Patient
- Carer
- Professional

## Condition

- Head injury
- Stroke
- Tumour
- Hydrocephalus
- Neurodegeneration

## Activity

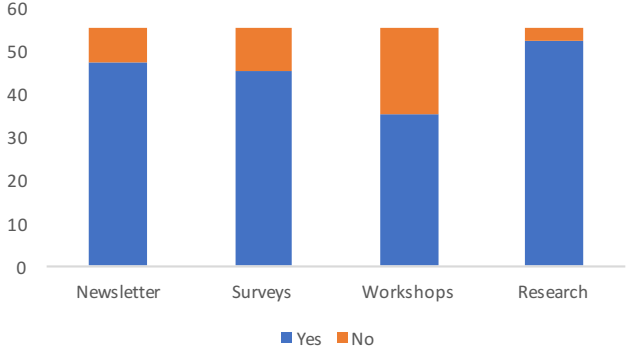
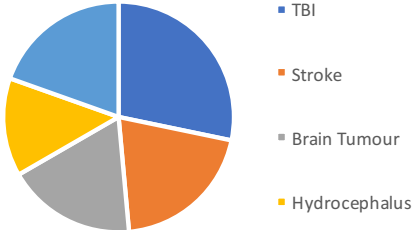
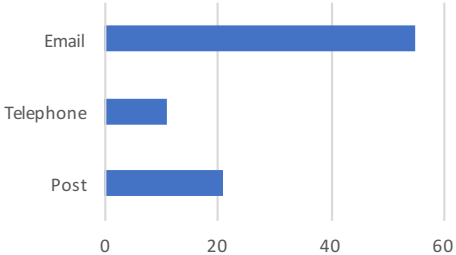
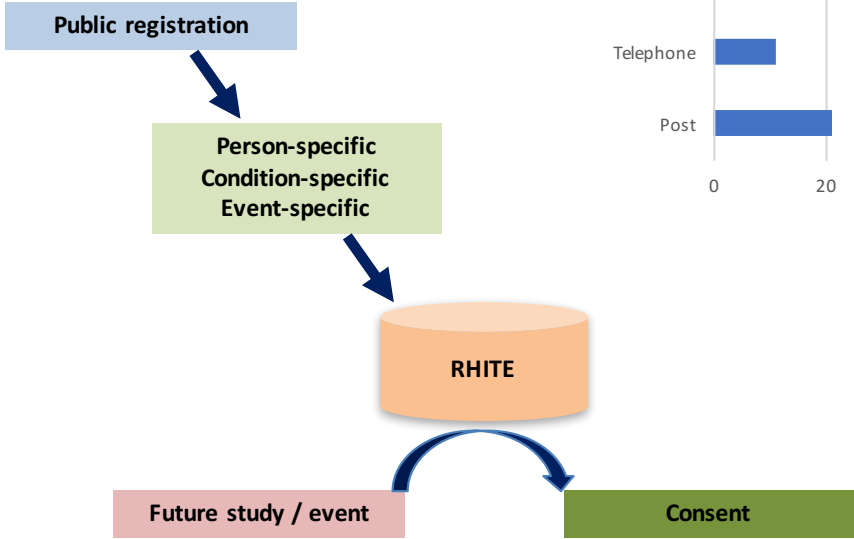
- Newsletter
- Surveys
- Workshops
- Research



# RHITE - initial evaluation



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# RHITE - next steps

