Sugar or Salt (SOS) Trial: comparing two current treatments for patients with a brain injury

Submission date	Recruitment status No longer recruiting	[X] Prospectively registered		
09/04/2019		[X] Protocol		
Registration date	Overall study status Ongoing	Statistical analysis plan		
16/04/2019		Results		
Last Edited 13/12/2024	Condition category Injury, Occupational Diseases, Poisoning	Individual participant data		
		[X] Record updated in last year		

Plain English summary of protocol

Background and study aims

Over one million people a year suffer injuries to their heads which require them to go to hospital. The most severe injuries often result in significant brain swelling. If left untreated, this swelling causes the pressure inside the head to increase, compressing the brain and causing further brain damage. The main treatments used for severe brain swelling involve placing the patient into an artificial coma (to rest the brain), giving drugs (to reduce brain swelling) or brain surgery (to release the pressure). Even with current treatments delivered in intensive care, over half of people with severe brain injury die or are left with severe brain damage. To improve outcomes for patients, doctors need to know the best treatments for severe brain swelling after head injuries. The two main drugs that are currently used to treat brain swelling are hypertonic saline (a strong salt solution) and mannitol (a sugary solution). Both of these drugs work by reducing brain swelling which helps to reduce pressure on the brain. Currently, it is not known which drug is the most effective treatment. Both drugs have undesirable side effects (hypertonic saline causes an imbalance of salts in the blood and mannitol can cause kidney failure). To deliver the best treatment doctors need to know which is most the safest and most effective. This study aims to work out which is the safest and most effective drug to treat the swelling of the brain that occurs after severe trauma to the head.

Who can participate?

Patients aged 16 or over admitted to an intensive care unit with a traumatic brain injury (an injury to the brain which occurs after trauma to the head)

What does the study involve?

Participants are randomly allocated to receive either the salty solution (hypertonic saline) or the sugary solution (mannitol). The study compares how effective the different drugs are at reducing the pressure on the brain. It also assesses which was better at helping the patient to recovery and what the side effects of treatment were. The study team keeps in contact with patients for 12 months after the study to check on how well they have recovered over time. Researchers also calculate how much each treatment costs and compare this to how beneficial they were.

What are the possible benefits and risks of participating?

Doctors do not know which of the two treatments is best, and that is why we are conducting this research. The researchers therefore cannot promise any direct benefits as a result of taking part in this study. However, it is hoped that the research will provide benefit to future patients who have a severe brain injury, as it will help doctors to know which is the best treatment to give. The risk of physical harm from taking part in the study is not considered to be any higher than the risks of standard clinical care, because the study is testing two existing treatments rather than a new treatment. Because the study involves completing questionnaires, there is a risk that participants may find it upsetting to answer some questions about their recovery. Trained research staff are available to talk to participants about any such feelings and can offer to put them in contact with professional services if this would be helpful.

Where is the study being run from? Queen Elizabeth Hospital - University Hospitals Birmingham NHS Foundation Trust (UK)

When is the study starting and how long is it expected to run for? June 2019 to February 2026

Who is funding the study? National Institute for Health Research (NIHR) Health Technology Assessment (HTA) Programme (UK)

Who is the main contact? University of Warwick study team sostrial@warwick.ac.uk

Contact information

Type(s)

Scientific

Contact name

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Type(s)

Public

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Additional identifiers

Clinical Trials Information System (CTIS)

2019-001688-66

Integrated Research Application System (IRAS)

260350

ClinicalTrials.gov (NCT)

Nil known

Protocol serial number

17/120/01, IRAS 260350

Study information

Scientific Title

Sugar or Salt (SOS) Trial: hyperosmolar therapy in traumatic brain injury

Acronym

SOS

Study objectives

The primary hypothesis is that hypertonic saline is more effective than mannitol in the management of raised ICP after severe TBI through improving clinical outcomes and cost-effectiveness.

Ethics approval required

Ethics approval required

Ethics approval(s)

approved 09/09/2019, East of England – Essex Research Ethics Committee (The Old Chapel, Royal Standard Place, Nottingham, NG1 6FS, United Kingdom; +44 (0)207 104 8115; essex. rec@hra.nhs.uk), ref: 19/EE/0228

Study design

Multicentre open-label randomized controlled clinical and cost-effectiveness trial with an internal pilot

Primary study design

Interventional

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Traumatic brain injury

Interventions

Current interventions as of 10/06/2019:

A simple and secure, web-based and allocation concealed randomisation system will be used. Randomisation will be stratified by site and predicted probability of 6-month unfavourable outcome. This predicted probability will be calculated using age, pupillary response and documented Glasgow Coma Scale (GCS) motor score at intubation using the IMPACT calculator (Steyerberg et al, 2008).

Patients will be randomized in a 1:1 ratio to receive intravenous boluses of either 2 ml/kg 20% mannitol or 2 ml/kg hypertonic saline (or equivalent osmolar dose using concentration used locally by participating study centres).

If intracranial pressure (ICP) remains higher than 20mmHg, boluses of each treatment can be repeated until serum sodium is >155 mmol/L. If there is a second spike in ICP over 20 mmHg then the allocated IMP should continue to be used.

Trial treatment will continue until therapeutic targets have been met. The total duration of follow-up for both treatment arms will be 12 months.

Previous interventions:

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If intracranial pressure (ICP) remains high, boluses of each treatment can be repeated until either ICP is less than 20 mmHg or serum sodium is >155 mmol/L or osmolarity is >320 mosmol/L. If there is a second spike in ICP over 20 mmHg then the allocated IMP should continue to be used.

Trial treatment will continue until therapeutic targets have been met. The total duration of follow-up for both treatment arms will be 12 months.

Intervention Type

Drug

Phase

Phase III

Drug/device/biological/vaccine name(s)

Mannitol, hypertonic saline

Primary outcome(s)

Neurological outcome measured by patient/relative/clinician completion of the Extended Glasgow Outcome Scale (GOS-E) questionnaire at 6 months

Key secondary outcome(s))

- 1. Intracranial pressure (ICP) control recorded continuously or at regular intervals from ICP bolt readings during the period of monitoring on ICU
- 2. Progression to stage 3 therapies (i.e. any use of additional treatments e.g. barbiturate coma, decompressive craniectomy, hypothermia, CSF drainage) recorded from the patient's medical records during their ICU stay
- 3. Which stage 3 therapies were required, recorded from the patient's medical notes during their ICU stay
- 4. Organ support requirements during ICU recorded from the patient's medical records, or through data linkage, according to the Critical Care Minimum Data Set definitions
- 5. ICU length of stay obtained from hospital records and through data linkage
- 6. Hospital length of stay obtained from hospital records and through data linkage
- 7. Discharge location obtained from hospital records and through data linkage
- 8. Longer term neurological outcomes measured using the modified Oxford Handicap Score (mOHS) completed by the research or clinical team at hospital discharge, and the Extended Glasgow Outcome Scale (GOS-E) completed by the patient/relative/clinician at 12 months
- 9. Survival measured from the patient's medical records at hospital discharge, 3 months, 6 months and 12 months
- 10. Health-related quality of life measured using the EQ-5D-5L at hospital discharge, 3 months, 6 months and 12 months post-TBI, completed by the patient/relative/clinician
- 11. Resource use collected from hospital records and through data linkage for the patient's duration of hospital stay and up to 12 months post-TBI
- 12. Serious adverse events recorded from the time that the patient is randomised through and including 28 calendar days after the last administration of IMP

Completion date

28/02/2026

Eligibility

Key inclusion criteria

- 1. Age 16 years or over
- 2. Admission to ICU following traumatic brain injury
- 3. ICP > 20mmHg for more than 5 minutes despite stage 1 procedures
- 4. <10 days from initial head injury
- 5. Abnormal CT scan consistent with traumatic brain injury

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Lower age limit

16 years

Sex

All

Key exclusion criteria

Current participant exclusion criteria as of 21/12/2023:

- 1. Devastating brain injury with withdrawal of treatment anticipated in the next 24 hours
- 2. Pregnancy
- 3. Severe hypernatraemia (Na > 155 mmol/L)
- 4. 2 or more prior doses of hyperosmolar therapy given on ICU

Previous participant exclusion criteria as of 06/08/2020:

- 1. Devastating brain injury with withdrawal of treatment anticipated in the next 24 hours
- 2. Pregnancy
- 3. Severe hypernatraemia (Na > 155 mmol/L)

Previous participant exclusion criteria from 10/06/2019 to 06/08/2020:

- 1. Devastating brain injury with withdrawal of treatment anticipated in the next 24 hours
- 2. Pregnancy
- 3. Severe hypernatraemia (Na > 160 mmol/L)

Original participant exclusion criteria:

- 1. Unsurvivable injuries
- 2. Pregnancy
- 3. Severe hypernatraemia (Na > 160 mmol/L)

Date of first enrolment

01/12/2019

Date of final enrolment

28/02/2025

Locations

Countries of recruitment

United Kingdom

England

Northern Ireland

Scotland

Wales

Study participating centre

Queen Elizabeth Hospital - University Hospitals Birmingham NHS Foundation Trust

Heritage Building Mindelsohn Way Edgbaston Birmingham United Kingdom B15 2TH

Study participating centre John Radcliffe Hospital

Oxford University Hospitals NHS Foundation Trust Headley Way Headington Oxford United Kingdom OX3 9DU

Study participating centre Salford Royal Hospital

Salford Royal NHS Foundation Trust Stott Lane Salford United Kingdom M6 8HD

Study participating centre Derriford Hospital

University Hospitals Plymouth NHS Foundation Trust Derriford Rd Plymouth United Kingdom PL6 8DH

Study participating centre The Walton Centre NHS Foundation Trust

Lower Lane Fazakerley Liverpool United Kingdom L9 7LJ

Study participating centre Southampton General Hospital

University Hospital Southampton NHS Foundation Trust Tremona Road Southampton United Kingdom SO16 6YD

Study participating centre Royal Victoria Hospital

Belfast Health & Social Care Trust Grosvenor Road Belfast United Kingdom BT12 6BA

Study participating centre

King's College Hospital King's College Hospital

King's College Hospital NHS Foundation Trust Denmark Hill London United Kingdom SE5 9RS

Study participating centre Royal Infirmary of Edinburgh

NHS Lothian Little France Cres Edinburgh United Kingdom EH16 4SA

Study participating centre Addenbrookes Hospital

Cambridge University Hospitals NHS Foundation Trust Hills Road Cambridge United Kingdom CB2 0QQ

Study participating centre

Lancashire Teaching Hospitals NHS Foundation Trust

Royal Preston Hospital Sharoe Green Lane Fulwood Preston United Kingdom PR2 9HT

Study participating centre University Hospital of Wales

Heath Park Cardiff United Kingdom CF14 4XW

Study participating centre The Royal Victoria Infirmary

Queen Victoria Road Newcastle upon Tyne United Kingdom TS1 4LP

Study participating centre South Tees Hospitals NHS Foundation Trust

James Cook University Hospital Marton Road Middlesbrough United Kingdom TS4 3BW

Study participating centre Royal Hallamshire Hospital

Glossop Road Sheffield United Kingdom S10 2JF

Study participating centre
Nottingham University Hospitals NHS Trust - City Campus
Nottingham City Hospital
Hucknall Road

Nottingham United Kingdom NG5 1PB

Study participating centre Aberdeen Royal Infirmary

Foresterhill Road Aberdeen United Kingdom AB25 2ZN

Study participating centre Queen Elizabeth University Hospital

1345 Govan Road Glasgow United Kingdom G51 4TF

Study participating centre University Hospital of North Staffordshire

Princes Road Stoke-on-trent United Kingdom ST4 7LN

Study participating centre Leeds General Infirmary

Great George Street Leeds United Kingdom LS1 3EX

Study participating centre Imperial College Healthcare NHS Trust

The Bays St Marys Hospital South Wharf Road London United Kingdom W2 1BL

Study participating centre St George's University Hospitals NHS Foundation Trust

St George's Hospital Blackshaw Road Tooting London United Kingdom SW17 0QT

Study participating centre Barts Health NHS Trust

The Royal London Hospital 80 Newark Street London United Kingdom E1 2ES

Sponsor information

Organisation

University Hospitals Birmingham NHS Foundation Trust

ROR

https://ror.org/014ja3n03

Organisation

University of Warwick

Funder(s)

Funder type

Government

Funder Name

Health Technology Assessment Programme

Alternative Name(s)

NIHR Health Technology Assessment Programme, Health Technology Assessment (HTA), HTA

Funding Body Type

Government organisation

Funding Body Subtype

National government

Location

United Kingdom

Results and Publications

Individual participant data (IPD) sharing plan

The data sharing plans for the current study are unknown and will be made available at a later date.

IPD sharing plan summary

Data sharing statement to be made available at a later date

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<u>Protocol article</u>	protocol	25/02/2020	06/08/2020	Yes	No
HRA research summary			28/06/2023		No
Participant information sheet	Participant information sheet	11/11/2025	11/11/2025	No	Yes