Standardised, Concentrated and Additional Macronutrients in neonatal Parenteral nutrition study

Submission date Recruitment status [X] Prospectively registered 23/04/2009 No longer recruiting [X] Protocol Statistical analysis plan Overall study status Registration date 28/05/2009 Completed [X] Results [] Individual participant data Last Edited Condition category **Neonatal Diseases** 27/11/2023

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s)

Scientific

Contact name

Mrs Sarah Azurdia

Contact details

Lead Research Nurse
Department of Neonatology
Liverpool Women's Hospital NHS Foundation Trust
Crown Street
Liverpool
United Kingdom
L8 7SS
+44 (0)151 7089988 x 4382
sarah.azurdia@lwh.nhs.uk

Additional identifiers

Clinical Trials Information System (CTIS)

Nil known

ClinicalTrials.gov (NCT)

Nil known

Protocol serial number

LWH 0776

Study information

Scientific Title

Standardised, Concentrated, Additional Macronutrients, Parenteral (SCAMP) nutrition in very preterm infants: a phase IV randomised, controlled study of macronutrient intake, growth and other aspects of neonatal care

Acronym

SCAMP

Study objectives

Maximising the macronutrient content of the standardised, concentrated neonatal parenteral nutrition formulation will improve head growth in very preterm infants.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Central Manchester Research Ethics Committee, North West Strategic Health Authority. Approval expected on 11/05/2009 (ref: 09/H1008/91).

Study design

Phase IV randomized controlled trial

Primary study design

Interventional

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Head growth in very preterm infants

Interventions

Informed consent and randomisation to the trial. Intravenous administration of either scNPN (standard) or scNPNmax (study) solution. Daily collection of routine biochemical monitoring data and all oral and intravenous fluid/drug administration on the case report form. All infants will be measured for weight, lower leg length, occipito-frontal circumference and length on day 0 (baseline) day 7,14,21,28. The total duration of interventions will be until the infant is 28 days old. The follow up will consist of weekly growth measurements until the infant reaches 36/40 post conceptual age and then neuo-developmental follow-up at 18 months corrected gestational age.

Intervention Type

Drug

Phase

Drug/device/biological/vaccine name(s)

Neonatal parenteral nutrition formulation

Primary outcome(s)

To compare early head growth in premature infants receiving a standardised formulation of parenteral nutrition(scNPN) with a standardised, concentrated formulation of parenteral nutrition containing additional protein, fat and carbohydrate (scNPNmax). Timepoints: when the infant is 28 days old (having already had growth measurements on days 0, 7, 14 and 21).

Key secondary outcome(s))

- 1. Weight gain and linear growth
- 2. Monitoring supplementary infusion requirements such as insulin and electrolytes
- 3. Routine biochemical measures of PN tolerance
- 4. Cost-benefit analysis
- 5. Efficiency and safety of prescribing and administration
- 6. Neurodevelopmental outcome (Bayley III)

Timepoints:

For outcomes 1 - 5: At 28 days and 36 weeks post conceptual age For outcome 6: At 18 months corrected for gestational age

Completion date

01/05/2013

Eligibility

Key inclusion criteria

All infants (both males ane females) born less than 29 weeks gestation and weighing <1,200 grams born at Liverpool Women's Hospital (LWH) will be eligible. All infants meeting these criteria born outside LWH will be eligible if transfer takes place to LWH within 48 hours.

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Neonate

Sex

Αll

Key exclusion criteria

- 1. Infants born in poor condition in the first 72 hours and are unlikely to survive the first week after birth
- 2. Infants diagnosed with major congenital and chromosomal abnormalities know to affect head growth or gastrointestinal function

Date of first enrolment

01/06/2009

Date of final enrolment

01/05/2013

Locations

Countries of recruitment

United Kingdom

England

Study participating centre Lead Research Nurse

Liverpool United Kingdom L8 7SS

Sponsor information

Organisation

Liverpool Women's Hospital NHS Foundation Trust (UK)

ROR

https://ror.org/04q5r0746

Funder(s)

Funder type

Charity

Funder Name

BLISS (UK) - Innovations in Care Programme

Results and Publications

Individual participant data (IPD) sharing plan

IPD sharing plan summaryNot provided at time of registration

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article	results	01/06/2013		Yes	No
Results article	results	01/01/2014		Yes	No
Results article	results	01/05/2015		Yes	No
Results article		10/11/2023	27/11/2023	Yes	No
<u>Protocol article</u>	protocol	10/06/2011		Yes	No
HRA research summary			28/06/2023	No	No
Participant information sheet	Participant information sheet	11/11/2025	11/11/2025	No	Yes