

Device Deficiency

Record ID

(REDCap auto generated)

*Phagenesis Ltd, who manufacture the Phagenyx® PES system, indemnify their equipment.



Pharyngeal Electrical stimulation for Acute Stroke dysphagia Trial (PhEAST)

UK ISRCTN 98886991

UK IRAS306761

UK CPMS 50913

WHO UTN U1111-1273-9942

Device deficiency form v1.3

► Please check consent form obtained.

Section A: Participant details

A1. Centre name:

(Centre)

A2. Participant ID :

(Participant ID)

A3. Participant initials (e.g. ABC or A-C) :

(3 uppercase letters, or 2 separated by a hyphen (-))

Section B: Device deficiency Phagenyx® is CE marked

B1. Date and time of device deficiency

(Date & time (DD-MM-YYYY HH:MM:SS))

B2. Timing of deficiency

- Before starting patient treatment, i.e. before first PES treatment
- During patient treatment, i.e. between first and last PES treatments
- After treatment, i.e. after last PES treatment
- Unknown

B3. Device component

Base station Catheter

B4a. Were there any equipment/device problems?

- Cable could not connect to basestation
- Cable could not connect to catheter
- Cable broke/snapped
- Catheter broke
- Base station screen not working
- Base station software freezes
- Safety button used to stop PES
- Device calculated stimulation differs from calculation by more than 2mA
- Other

(Choose one answer)

B4b. If "Other", please give a short description of the problem.

B4c. Was there an associated safety event?

- None
- ADE
- SADE
- UADE
- USADE

B5. Please provide as much information as possible on the equipment failure/problem.

This may be shared with the manufacturer so no identifying information please.

(This may be shared with the manufacturer so no identifying information please.)

Photo of defective device, catheter or basestation

B6. You may upload an image of the device, catheter or basestation showing the problem.

( Photo (but no faces, name badges or identifiers please).)

B7. Description of event

B8a. Deficiency related to

- Labeling
- Quality
- Durability
- Reliability
- Performance
- Safety
- Other

(Choose one answer)

B8b. If "Other", please explain the deficiency

B9. Deficiency associated with an (serious) adverse event?

If yes, please also complete a SAE form even if after 7 days post-randomisation.

Yes No
(If yes, please also complete a SAE form)

B10. Device component(s) returned, or intention to return, to manufacturer (Phagenesis)?

Yes No

B11. Safety button used to stop PES (Urgent safety measure)

Yes No

Section C: Catheter and device details.

C1. Catheter Lot number

(([[pes_catheter_lotno_1]]) ([[pes_catheter_lot_no_2]])
([[pes_catheter_lot_no_3]]) ([[pes_catheter_lot_no_4]])
([[pes_catheter_lot_no_5]]) ([[pes_catheter_lotno_2]]))

C2. Base station serial number

(([[pes_baselstation_serialno_1]])
([[pes_baselstation_serialno_2]]))

Section D: Assessor information.

D1. Please enter your name

D2a. What is your professional role?

- Doctor
- Research coordinator
- Nurse, clinical
- Research nurse
- Physiotherapist
- Occupation therapist
- Speech & Language therapist
- Other

(Choose one answer)

D2b. If "Other", please specify your role

D3. Does your role involve working on stroke wards?

- Yes
- No

D4. Please enter your name if you did not collect the information

D5. Please sign the form

 (Signature)

Comments and full explanation for missing data

Are any values missing due to tests not done (or measures not taken), or because data are unknown and every effort has been made to find the data - i.e. 'Not done' / 'Not known'?

- Yes
- No

If any values are missing, please provide a full explanation Comments