

Effect of botulinum toxin A injections and specific intensive rehabilitation therapy in children with hemiparetic cerebral palsy on upper limb functions and skills

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Registration date 31/03/2008	Overall study status Completed	<input type="checkbox"/> Protocol
Last Edited 20/08/2015	Condition category Nervous System Diseases	<input type="checkbox"/> Statistical analysis plan
		<input checked="" type="checkbox"/> Results
		<input type="checkbox"/> Individual participant data

Plain English summary of protocol
Not provided at time of registration

Contact information

Type(s)
Scientific

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Additional identifiers

Protocol serial number
NL12005.096.06

Study information

Scientific Title

Effect of botulinum toxin A injections and specific intensive rehabilitation therapy in children with hemiparetic cerebral palsy on upper limb functions and skills

Acronym

BoBiVa (Botuline toxine Bimanuele Vaardigheden)

Study objectives

Research question:

What is the effect of botulinum toxin A (btA) injections (B), an intensive physical and occupational therapy program aimed at improving arm function and skills (C), or a combination of both (A), on arm function, bimanual skills and use of the affected arm, in children with hemiparetic cerebral palsy, relative to the course in such children who receive usual care (D)?

Ethics approval required

Old ethics approval format

Ethics approval(s)

1. Medical Ethics Committee of the Meuse Hospital (Medisch Ethische Toetsingscommissie Atrium MC-Maaslandziekenhuis), 27/07/2006
2. This trial is also registered at the Centrale Commissie Mensgebonden Onderzoek (CCMO) Central Committee for Research Involving Human Subjects (<https://toetsingonline.ccmo.nl>) (ref: NL12005.096.06)

Study design

Randomised controlled trial

Primary study design

Interventional

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Cerebral palsy

Interventions

This study will take place in three hospitals in the Netherlands: University Hospital Maastricht (Franciscusoord Valkenburg), Maartenskliniek Nijmegen Hospital and Free University Medical Centre (Vrije Universiteit Medisch Centrum [VUMC]) Amsterdam.

Interventions:

Group A: BtA injections (Dysport®) prior to therapy programme and intensive physical and occupational therapy programme

Group B: BtA injections only

Group C: Intensive physical and occupational therapy programme

Group D: Usual care

BtA injections:

The most spastic muscles hampering function will be injected. Dysport® dilution: 25 U/0.1 ml, dose 6 - 9 U/kg body weight muscles above elbow, 3 - 6 U/kg body weight muscles in forearm,

limited to no more than 150 units (0.6 ml) at any one injection site. In the intrinsic thumb muscles the maximum dose will be 25 U per muscle. A maximum Dysport® dose of 1,000 U per child in total per session will be used.

Intensive physical and occupational therapy programme:

Participants will receive one hour of occupational therapy and 30 minutes of physical therapy, twice a week for 12 weeks.

Intervention Type

Drug

Phase

Not Applicable

Drug/device/biological/vaccine name(s)

Botulinum toxin A (Dysport®)

Primary outcome(s)

1. Assisted Hand Assessment (AHA): original test kit for children 2.5 - 6 years and board game for children 7 - 12 years (T2, T4, T6)
2. A measure of manual ability for children with upper limb impairments (ABILHAND)-Kids questionnaire (T1 - T6)
3. Canadian Occupational Performance Measure (COPM): establishing treatment goals; Goal Attainment Scaling (GAS) of the most important bimanual treatment goal (T1, T4, T6)
4. Video recording of two fine motor tasks (children 7 - 12 years: buttering and cutting bread, screw construction task; children 2.5 - 6 years: building with 'poppons', threading beads) and one gross motor task (children 2.5 - 6 years: building blocks; children 7 - 12 years: stacking cylinders). These videos will be scored with newly developed and reliability tested Video Observation (VO) criteria (T2, T6).

T1 and T2: Baseline

T3: 6 weeks after btA and start of the therapy program

T4: 12 weeks, end of therapy program

T5: 18 weeks

T6: 24 weeks

Key secondary outcome(s)

1. Wrist and elbow tone and Tardieu Scale or Spasticity Test (SPAT): supine and sitting (T1 - T6)
2. Active and passive range of motion (ROM) of wrist (with fist and with extended fingers), and of elbow and thumb (T1 - T6)
3. Grip strength: E-link (biometrics®) and functional grip strength (T1 - T6)

T1 and T2: Baseline

T3: 6 weeks after btA and start of the therapy program

T4: 12 weeks, end of therapy program

T5: 18 weeks

T6: 24 weeks

Completion date

01/01/2012

Eligibility

Key inclusion criteria

1. Aged 2.5 - 12 years, either sex
2. Cerebral palsy
3. Hagberg diagnosis: spastic hemiparesis or extreme asymmetric diplegia
4. Hand function impairment Zancolli grade I with evident problems in thumb extension and supination, Zancolli grade IIA and IIB
5. Mentally able to comprehend and perform tasks
6. Children and their parents should be able to cope with the intensive rehabilitation therapy programme and the measurement sessions
7. Children and the parents/caregivers should comprehend and speak Dutch
8. Children and their parents indicate the necessity for improvement of the children's abilities

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Child

Lower age limit

2.5 years

Upper age limit

12 years

Sex

All

Key exclusion criteria

1. Severe structural contractures of the muscles at the extremity to be treated:
 - 1.1. Passive elbow extension maximum 160 degrees or less
 - 1.2. Supination maximum 30 degrees or less from neutral position
 - 1.3. Wrist dorsal flexion maximum 20 degrees or less in children aged 2.5 - 6 years, or 45 degrees or less in age group 7 - 12 years
2. Severe impairment of hand function: no active hand function is expected after treatment (Zancolli III)
3. Hand surgery or phenolisation or btA injections in the arm less than nine months ago
4. Contraindication for botulinum toxin (muscular diseases such as myasthenia gravis, tetanus vaccination less than three months before the injection, use of aminoglycoside antibiotics or spectinomycine and known hypersensitivity for human albumin)
5. Contraindication for anaesthesia
6. Children who cannot bear touching the affected arm and hand

Date of first enrolment

01/01/2008

Date of final enrolment

01/01/2012

Locations

Countries of recruitment

Netherlands

Study participating centre**Franciscusoord Child Rehabilitation**

Valkenburg

Netherlands

6301 KA

Sponsor information

Organisation

Ipsen Biopharm Ltd (UK)

ROR

<https://ror.org/00gmnqd91>

Funder(s)

Funder type

Research organisation

Funder Name

Rotterdam Foundation Child Rehabilitation Fund 'Adriaan Fund' (Stichting Rotterdams Kinderrevalidatie Fonds Adriaanstichting) (The Netherlands)

Funder Name

Johanna Children's Fund (Johanna Kinderfonds) (The Netherlands) - <http://www.johannakinderfonds.nl>

Funder Name

Phelps Foundation for Spasticity (Phelps Stichting voor Spastici) (The Netherlands) - <http://www.phelps-stichting.nl>

Funder Name

Profile Fund of the University Hospital Maastricht (Profileringsfonds azM) (The Netherlands)

Funder Name

Foundation for Children's Illness (Stichting het gebrekkige Kind) (The Netherlands)

Funder Name

Ipsen Biopharm Ltd (UK) - provided Dysport®

Results and Publications

Individual participant data (IPD) sharing plan

IPD sharing plan summary

Not provided at time of registration

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article	results	19/08/2015		Yes	No
Study website	Study website	11/11/2025	11/11/2025	No	Yes