# Empowering parents of obese children: development and controlled evaluation of an obesity-specific parenting skills training

Submission date 12/10/2007	<b>Recruitment status</b> No longer recruiting	<ul> <li>Prospectively registered</li> <li>Protocol</li> </ul>
<b>Registration date</b> 06/03/2008	<b>Overall study status</b> Completed	 [_] Statistical analysis plan [X] Results
Last Edited 16/11/2016	<b>Condition category</b> Nutritional, Metabolic, Endocrine	[_] 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

EudraCT/CTIS number

**IRAS number** 

ClinicalTrials.gov number

Secondary identifying numbers

### Study information

#### Scientific Title

Empowering Parents of Obese Children: development and controlled evaluation of an obesityspecific parenting skills training

#### Acronym

EPOC

#### **Study objectives**

A compact parent training enhancing "obesity-specific" parenting skills combined with written information and a training video contemporaneous with the inpatient training of the children leads to a greater reduction of percentage overweight (BMI-SDS) in the obese children and an improvement of children and parents' psychosocial variables one year after the end of intervention compared to a parent group only receiving written information.

#### Ethics approval required

Old ethics approval format

**Ethics approval(s)** Ethics Committee of the University of Potsdam, 19/05/2006

**Study design** Prospective randomised controlled study

#### **Primary study design** Interventional

**Secondary study design** Randomised controlled trial

**Study setting(s)** Other

**Study type(s)** Treatment

#### Participant information sheet

Not available in web format, please use the contact details to request a patient information sheet

Health condition(s) or problem(s) studied Obesity

#### **Interventions** Parent training course (intervention group) vs written information only (control group)

Parent training course:

Key activity: Cognitive-behavioral training, focusing on how parents can support their children at home in critical behavioural tasks (increasing physical activity, dietary changes).

Materials and techniques: Group discussions, role play, work sheets, introduction of critical situations (video), take-home video and take-home written information.

The participants will be taught by health professionals of the participating clinics (psychologists, pedagogues, nutritionists, dieticians, etc.)

Duration of the course: 2 days (in most instances weekend-course), 10 sessions, on average 50 minutes per session

Topics/contents of the written information provided to the control group:

- 1. Etiology of obesity
- 2. "Traffic light" diet
- 3. Advice on increasing activity (physical exercise and daily motor activity)
- 4. Dietary changes

#### Intervention Type

Other

Phase Not Specified

#### Primary outcome measure

BMI-SDS in children, assessed at T1, T2, T4, T5 and T6 (assessed by a physician, an interview and a questionnaire)

Timepoints:

- T1: Start of the child's inpatient stay
- T2: End of the child's inpatient stay
- T3 (Applicable only for the intervention group; timepoint when a telephone booster session takes place) : One month after the child's inpatient stay
- T4: Three months after the child's inpatient stay
- T5: Six months after the child's inpatient stay
- T6: Twelve months after the child's inpatient stay

#### Secondary outcome measures

- 1. Psychosocial variables of parents:
- 1.1. Self-efficacy, assessed at T1, T2, T4, T5 and T6 by questionnaires and an interview

1.2. Psychosocial strain, assessed at T1 and T6 by the Short Form (SF)-12 State of Health Questionnaire

1.3. Familial support, assessed at T1, T4, T5 and T6 by questionnaires and an interview 1.4. Feeding style, assessed at T1, T4, T5 and T6 by the Child Feeding Questionnaire (CFQ), Caregiver's Feeding Styles Questionnaire (CFSQ) and an interview

2. Psychosocial variables of children:

2.1. General quality of life, assessed at T1, T4, T5 and T6 by a children's quality of life questionnaire (KINDL)

2.2. Weight-related quality of life, assessed at T1, T2, T4, T5 and T6 by a weight-related quality of life questionnaire (GW-LQ-KJ) and an interview

- 2.3. General self-efficacy, assessed at T1, T2, T4, T5 and T6 by questionnaires
- 2.4. Weight-related self-efficacy, assessed at T1, T4, T5 and T6 by a weight-related self-efficacy

questionnaire (GW-SW-KJ)

2.5. Eating behaviour, assessed at T1, T4, T5 and T6 by the Eating Behaviour Questionnaire (FKE-KJ)

2.6. Activity behaviour, assessed at T1, T4, T5 and T6 by the German Health Interview and Examination Survey for Children and Adolescents questionnaire

2.4. Familial support, assessed at T1, T4, T5 and T6 by a questionnaire

2.5. Psychosocial strain, assessed at T1 and T6 by the Strengths & Difficulties Questionnaire (SDQ)

Timepoints:

T1: Start of the child's inpatient stay

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T3 (Applicable only for the intervention group; timepoint when a telephone booster session takes place): One month after the child's inpatient stay

T4: Three months after the child's inpatient stay

T5: Six months after the child's inpatient stay

T6: Twelve months after the child's inpatient stay

#### Overall study start date

01/04/2007

#### **Completion date**

31/12/2012

## Eligibility

#### Key inclusion criteria

Current inclusion criterria as of 16/05/2012:

1. Children aged 7-13 years

2. Duration of inpatient stay at least four weeks

3. BMI-SDS exceeding the 97th percentile

4. Informed consent to participate in the study and to participate in the parent training if randomised into the intervention group

Previous inclusion criteria:

1. Children aged 7-12 years

2. Duration of inpatient stay at least four weeks

3. BMI-SDS exceeding the 97th percentile

4. Informed consent to participate in the study and to participate in the parent training if randomised into the intervention group

Participant type(s) Patient

**Age group** Child

**Lower age limit** 7 Years

Upper age limit

13 Years

**Sex** Both

**Target number of participants** 500

#### Key exclusion criteria

- 1. Inadequate German language skills of the parent
- 2. Secondary obesity as a consequence of other diseases
- 3. Major comorbid diseases
- 4. Major behavioural or psychiatric disorder of the child
- 5. Psychiatric disorder of the parent

6. Previous or concomitant parent training by other sources or accompaniment by the parent during the child's inpatient stay

#### Date of first enrolment

01/04/2007

### Date of final enrolment

31/12/2012

### Locations

**Countries of recruitment** Germany

**Study participating centre University of Potsdam** Potsdam OT Golm Germany 14476

### Sponsor information

**Organisation** German Research Foundation (Deutsche Forschungsgemeinschaft) (Germany)

**Sponsor details** Kennedyallee 40 Bonn Germany 53175 +49 (0)228 8851 postmaster@dfg.de

**Sponsor type** Government

Website http://www.dfg.de

ROR https://ror.org/018mejw64

### Funder(s)

**Funder type** Government

**Funder Name** Deutsche Forschungsgemeinschaft

**Alternative Name(s)** German Research Association, German Research Foundation, DFG

Funding Body Type Government organisation

Funding Body Subtype National government

**Location** Germany

### **Results and Publications**

**Publication and dissemination plan** Not provided at time of registration

Intention to publish date

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 01/08/2016 Results article Yes No results Results article 14/11/2016 Yes No