# Insulin sensitivity in preterm appropriate-forgestational-age and small-for-gestational-age infants

| Submission date   | Recruitment status       | [X] Prospectively registered                  |
|-------------------|--------------------------|---|
| 26/02/2007        | No longer recruiting     | Protocol                                      |
| Registration date | Overall study status     | Statistical analysis plan                     |
| 26/02/2007        | Completed                | Results                                       |
| Last Edited       | Condition category       | Individual participant data                   |
| 26/08/2021        | Pregnancy and Childbirth | <ul><li>Record updated in last year</li></ul> |

### **Plain English Summary**

Not provided at time of registration

## Contact information

## Type(s)

Scientific

#### Contact name

Prof H P Sauerwein

### Contact details

Academic Medical Centre (AMC)
Department of Endocrinology and Metabolism, F5-170
P.O. Box 22660
Amsterdam
Netherlands
1100 DD
+31 (0)20 566 3061
h.p.sauerwein@amc.uva.nl

## Additional identifiers

## EudraCT/CTIS number

Nil known

### IRAS number

## ClinicalTrials.gov number

Nil known

### Secondary identifying numbers

NL874 (NTR888)

## Study information

### Scientific Title

Insulin sensitivity in preterm appropriate-for-gestational-age and small-for-gestational-age infants

### Study hypothesis

Insulin sensitivity is already reduced at birth in preterm Small-for-Gestational-Age (SGA) infants, compared to preterm Appropriate-for-Gestational-Age (AGA) infants.

## Ethics approval required

Old ethics approval format

### Ethics approval(s)

Approval received from the Central Committee on Research inv. Human Subjects on the 30th January 2006 (ref: P05.1488C).

### Study design

Observational study

### Primary study design

Observational

### Secondary study design

Multi-centre

## Study setting(s)

Not specified

## Study type(s)

Other

### Participant information sheet

### Condition

Small for gestational age, prematurity, insulin sensitivity

### **Interventions**

Methods used:

- 1. Glucose concentration: this will be measured with the glucose oxidase method using a Beckman Glucose Analyzer 2 (Beckman, Fullerton, CA)
- 2. Insulin: this will be determined with a chemiluminescent immunometric assay (Immulite 2000, Diagnostic Products Corporation, Los Angeles, USA)
- 3. Free Fatty Acid (FFA) concentration: this will be determined with an enzymatic colorimetric method (NEFA-C test kit, Wako Chemicals GmbH, Neuss, Germany)
- 4. Cortisol: this will be determined with a chemiluminiscent immunoassay (Immulite 2000,

Diagnostic Products Corporation, Los Angeles, USA)

- 5. Adiponectin: this will be determined by a radioimmunoassay (Linco, St. Charles, USA)
- 6. Stable isotope measurements: Newborns are infused with [U-13C] glucose and [2-13C] glycerol. Isotope dilution and label incorporation will be determined by gas chromatography mass spectrometry (GCMS) and mass isotopomer distribution analysis (MIDA) in glucose, isolated from plasma

### Calculations:

- 1. Rate of appearance (Ra) of glucose during steady state is calculated by the isotope dilution technique from the [U-13C] enrichment of glucose, using calculations for steady state kinetics, adapted for the use of stable isotopes:  $Ra = (Ei/Ep) \times I$ , where Ei and Ep are the enrichments of infusate and plasma respectively, and I is the infusion rate of [U-13C] glucose
- 2. Rate of disappearance (Rd): rate of exogenous glucose infusion plus the rate of endogenous glucose production
- 3. Endogenous glucose production: Rate of appearance minus rate of exogenous glucose infusion
- 4. Absolute gluconeogenesis: fractional gluconeogenesis (measured by MIDA) times rate of appearance
- 5. Glycogenolysis: Endogenous glucose production minus absolute gluconeogenesis

### Intervention Type

Other

#### Phase

**Not Specified** 

### Primary outcome measure

Rate of appearance and disappearance of glucose during insulin infusion

## Secondary outcome measures

- 1. Rate of gluconeogenesis and glycogenolysis
- 2. Plasma Free Fatty Acid (FFA) concentrations
- 3. Plasma concentrations of insulin, cortisol and adiponectin

### Overall study start date

01/04/2007

### Overall study end date

01/04/2008

## Eligibility

### Participant inclusion criteria

- 1. Premature infants 28 to 32 weeks gestational age
- 2. Presence of a (central) venous and arterial catheter for clinical reasons
- 3. For preterm SGA infants: growth retardation caused by placental insufficiency, assessed by maternal history (pregnancy induced hypertension, preeclampsia), and confirmed by Doppler flow measurements of the umbilical arteries (Pulsatility Index [PI] more than +2 Standard Deviation [SD] for gestational age, measured on two occasions)

### Participant type(s)

### **Patient**

### Age group

Neonate

#### Sex

**Not Specified** 

### Target number of participants

16

### Participant exclusion criteria

- 1. For preterm SGA infants: growth retardation based on other causes (e.g. congenital infections, congenital malformations)
- 2. Major congenital malformations
- 3. Severe perinatal asphyxia defined as five minute Apgar score less than seven
- 4. Severe disturbances of glucose metabolism (glucose intake less than 4 or more than 8 mg.kg-1. min-1, or need for insulin therapy to maintain the glucose concentration between 2.6 and 8 mmol /l)
- 5. Severe respiratory distress. Mild ventilatory support is allowed:
- a. nasal Continuous Positive Airway Pressure (nCPAP) with maximum Fraction of Inspired Oxygen (FiO2) of 0.40, maximum Positive End Expiratory Pressure (PEEP) 6 cm Water (H2O)
- b. Synchronised Intermittent Mandatory Ventilation (SIMV) with maximum inspiratory peak pressure of 18 cm H2O and maximum FiO2 of 0.40
- c. High Frequency Oscillatory Ventilation (HFOV) with maximum continuous distending pressure of 12 cm H2O and maximum FiO2 of 0.30
- 6. Need of vasopressor support for hypotension
- 7. Treatment with systemic corticosteroids
- 8. Clinical or laboratory evidence of sepsis: lethargy or irritability, hypo- or hyperthermia, temperature instability, tachypnea, apnea, bradycardia, hypotension, gastric retention, abdominal distension, pallor, elevated C- Reactive Protein (CRP)-level, leukocytosis or leukocytopenia and increased number of band neutrophils
- 9. Low haemoglobin level at the study days with need for a blood transfusion
- 10. Positive family history for type two diabetes in first degree relatives
- 11. No informed consent from parents or legal guardians

#### Recruitment start date

01/04/2007

### Recruitment end date

01/04/2008

## Locations

### Countries of recruitment

Netherlands

### Study participating centre

## Academic Medical Centre (AMC)

Amsterdam Netherlands 1100 DD

## Sponsor information

### Organisation

Diabetes Fonds Nederland (The Netherlands)

## Sponsor details

Stationsplein 139 Amersfoort Netherlands 3818 LE

\_

info@diabetesfonds.nl

### Sponsor type

Research organisation

### Website

http://www.diabetesfonds.nl/

### **ROR**

https://ror.org/04ch2g225

## Funder(s)

### Funder type

Hospital/treatment centre

### **Funder Name**

Academic Medical Centre (AMC) (The Netherlands)

## **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