Case Report Form
CONFIDENTIAL

Maternal and Offspring outcomes after Treatment of HyperEmesis by Refeeding

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Please read CRF instructions carefully!
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CRF MOTHER TRIAL for RANDOMIZED patients, Version: 2.0 final, 09-10-2015
CRF INSTRUCTIONS

1. Fill out CRF using a blue or black pen
2. Write clearly, if possible using capitals
3. Answer all questions. Only use ‘unknown’ when this option is given
4. Selection of buttons should be: ☑ or ☒ or ☐ (if applicable)
5. Radio buttons ‘Ο’ are used when only one answer is possible
6. Check boxes ‘☐’ are used when multiple answers are possible
7. Open Combs, e.g. ‘|__|’ are used for numerical answers
8. For missing numeric values, write ‘-1’. Otherwise, select ‘unknown’
9. Write dates as follows: dd-mm-jj e.g. 01-01-2001
10. When a date is (partly) missing, use ‘99’ for missing part, e.g. when the day is missing: 99-01-2001. When day and month are missing: 99-99-2001. When complete date is missing: 99-99-99
11. Never hide corrections (using Tipp-Ex)
12. When you want to correct something, strike through false answer with a single line, followed by the right answer. Sign and date this correction
13. After CRF completion, the local PI has to sign the CRF to declare complete and truthful filling out. After signing, corrections can no longer be made to the CRF.
1. General information

Case number | __ | __ | __ | __ | __ | __ | __ | __ | (clinic - case)

Date of birth | __ | __ | __ | __ | __ | __ | __ | __ | (dd-mm-yyyy)

Patient initials | __ | __ | (initials first name and female family name)

2. Randomization

Date of randomization | __ | __ | __ | __ | __ | __ | __ | __ | (dd-mm-yyyy)

Treatment allocation
- Tube feeding + standard care
- Standard care

Randomized on day of hospital admission for hyperemesis gravidarum (HG)*

| __ | __ | __ | __ | __ | __ | __ | __ | (dd-mm-yyyy)

Current admission is first admission for HG

| __ | __ | __ | __ | __ | __ | __ | __ | (dd-mm-yyyy)

Number of previous admissions for HG | __ | __ | (0-10)

*Hyperemesis gravidarum will further be abbreviated as HG
3. Medical history

**Disease(s)**
- Hypothyroidism: O no, O yes
- Hyperthyroidism: O no, O yes
- Pre-existing diabetes (type I/type II): O no, O yes
- Pre-existing hypertension: O no, O yes
- Peptic ulcer: O no, O yes
- Depressive disorder: O no, O yes
- Anxiety disorder: O no, O yes
- Eating disorder: O no, O yes
- Other: ________________________________

*History of disease or ongoing disease
†Including postnatal depression
§Including posttraumatic stress disorder (PTSS)

4. Obstetric history

**Gravidity**
- |__|__| (0-15)
**Parity**
- |__|__| (0-10)
**Miscarriage**
- |__|__| (0-10)
**EUG**
- |__|__| (0-10)
**Termination of pregnancy** (APLA)
- |__|__| (0-10)
**Progeniture** (children alive)
- |__|__| (0-10)

**Previous pregnancy**
- O no, O yes
  - HG in a previous pregnancy
    - O no, O yes, O unknown
  - Requiring hospital admission
    - O no, O yes, O unknown
5. Current pregnancy

5.1 Baseline characteristics

**Estimated date of delivery**  
Based on ultrasound or embryo transfer  
(dd-mm-yy)

**Type of pregnancy**  
- Singleton  
- Twins / Higher order multiple

**Height**  
unknown: -1  
(cm 140-210)

**Smoking**  
- No/Quit before pregnancy  
- Yes  
- Unknown

**Drug use**  
- No/Quit before pregnancy  
- Yes  
- Unknown  
- Cannabis  
- Other

**Folic acid use**  
- No  
- Yes  
- Unknown

**Antiemetics started before admission** (medicatie tegen misselijkheid)  
- No  
- Yes  
- Unknown
### 5.2 Urinary ketones at admission (during which patient was randomized)

**Date first measurement of urinary ketones during admission process**

| __ | __ | - | __ | __ | - | __ | __ | (dd-mm-yy)  
|---|---|---|---|---|---|---|---|

*Unknown: 99-99-99
†May or may not be the same as admission date

**Fill out corresponding value for urinary ketones**

<table>
<thead>
<tr>
<th>O Negative</th>
<th>O Positive</th>
<th>O Unknown</th>
</tr>
</thead>
<tbody>
<tr>
<td>O</td>
<td>+</td>
<td>++</td>
</tr>
<tr>
<td>O</td>
<td>+++</td>
<td>++++</td>
</tr>
<tr>
<td>O</td>
<td>Unknown</td>
<td></td>
</tr>
</tbody>
</table>

### 5.3 Blood testing at admission (during which patient was randomized)

**Date first blood test during admission process**

| __ | __ | - | __ | __ | - | __ | __ | (dd-mm-yy)  
|---|---|---|---|---|---|---|---|

*Unknown: 99-99-99
†May or may not be the same as admission date

**Fill out corresponding blood values for**

- **Hb**  | __ | __ | | mmol/l |
- **Ht**  | __ | __ | | l/l   |
- **TSH** | __ | __ | | mU/l |
- **Na**  | __ | __ | | mmol/l |
- **K**   | __ | __ | | mmol/l |
- **ASAT**| __ | __ | | U/l   |
- **ALAT**| __ | __ | | U/l   |
- **Urea**| __ | __ | | mmol/l |
- **Creatinin** | __ | __ | | mmol/l |

*Unknown: -1

**Have magnesium (Mg) and phosphate (P) been measured during admission**

<table>
<thead>
<tr>
<th>O No</th>
<th>O Yes</th>
<th>O Unknown</th>
</tr>
</thead>
</table>
| Date of first measurement**| __ | __ | - | __ | __ | - | __ | __ | (dd-mm-yy)  
|---|---|---|---|---|---|---|---|

*Unknown: 99-99-99
†May or may not be the same as first blood test date. In some hospitals, Mg and P are only tested in patients receiving tube feeding

**Fill out corresponding blood values for**

- **Mg (magnesium)**  | __ | __ | | mmol/l |
- **P (fosfaat/fosfor)** | __ | __ | | mmol/l |

*Unknown: -1
5.4 Medication during first admission (during which patient was randomized)

**Antiemetics** (medicatie tegen misselijkheid)

<table>
<thead>
<tr>
<th></th>
<th>No</th>
<th>Yes</th>
<th>Unknown</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Emesafene p.o. / supp</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Primperan p.o. / supp</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Primperan i.v.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ondansetron (Zofran) p.o. / supp</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ondansetron (Zofran) i.v.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Steroids*</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Other</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Prescribed as antiemetic, not because of fetal lung ripening

**Vitamins**

<table>
<thead>
<tr>
<th></th>
<th>No</th>
<th>Yes</th>
<th>Unknown</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>B1 (thiamin)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>B6 (pyridoxin)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>B12 (cobalamin)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>B complex</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Other</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

CRF MOTHER TRIAL for RANDOMIZED patients, Version: 2.0 final, 09-10-2015
6. Intervention (started during admission in which patient was randomized)

6.1 Intravenous (i.v.) drip

Note: For exact intravenous rehydration regimen, check both medical and nursing record to help you fill out this section

Did patient receive an i.v. drip*

- No
- Yes
- Unknown

| Patient declined
| Doctor advised against
| Other; _________________________

*As standard treatment or in combination with tube feeding

If i.v. drip:

Was i.v. drip placed on day of randomization

- No
- Yes
- Unknown

| Date of i.v. drip placement* (dd-mm-yy)

*Unknown: 99-99-99

i.v. drip solution contained*

- NaCl
- Glucose
- KCl
- Other; _________________________

*If in medical or nursing record only: ‘i.v. drip according to protocol’, fill out what is prescribed according to local protocol for HG

Did patient experience any side effect(s) of i.v. drip

- No
- Yes*
- Unknown

| Phlebitis (pain/redness/swelling at insertion area)
| Allergic reaction
| Other; _________________________

* See paragraph 8.3 whether SAE form needs to be filled out
| Case Number |  |  |  |

**Was i.v. drip removed because of side effects**

<table>
<thead>
<tr>
<th>O No</th>
<th>O Yes</th>
<th>O Unknown</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>O Patients request</td>
<td>O Doctors' advice</td>
<td>O Other; _________________</td>
</tr>
</tbody>
</table>

**Date of i.v. drip removal**

*Unknown: 99-99-99

†All reasons for i.v. drip removal

**If i.v. drip was removed, was it replaced**

<table>
<thead>
<tr>
<th>O No</th>
<th>O Yes</th>
<th>O Unknown</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Date of i.v. drip replacement**

*Unknown: 99-99-99

†Replacement during this admission, not placement of i.v. drip during readmission. In some hospitals patients receive a venflon and return the next day again for rehydration. This is not considered i.v. drip replacement but continuation of care until venflon is removed

**Date of replaced i.v. drip removal**

*Unknown: 99-99-99

†All reasons for i.v. drip removal
### 6.2 Tube feeding

**Note:** The dietician in your hospital will fill out a weekly registration form for all patients in this trial that have received tube feeding. When you fill out a CRF, please collect these form(s) for the specific patient, to help you fill out this section. These forms also contain complementing information on tube feeding regimen and caloric intake (not asked in this CRF).

Please attach the dietician registration form(s) to this CRF.

**If randomized for standard care:** Did patient receive a tube

- No
- Yes
- Unknown

- Patients request
- Doctors'/dieticians' advice
- Other: _________________________

**If randomized for tube feeding + standard care:** Did patient receive a tube

- No
- Yes
- Unknown

- Patient refused tube placement
- Tube not in stock
- Other: _________________________

**If tube:**

- **Was tube placed on day of randomization**
  - No
  - Yes
  - Unknown

- **Date of tube placement**:
  - (dd-mm-yy)
  - Unknown: 99-99-99

**Did patient experience any side effect(s) of tube**

- No
- Yes

- Nose/throat irritation
- Continuation of vomiting
- Tube obstruction
- Tube dislocation
- Aspiration
- Intestinal bleeding
- Intestinal perforation
- Other: _________________________

* See paragraph 8.3 whether SAE form needs to be filled out
### Case Number |__|__| __|  - |__| __| __|

**CRF MOTHER TRIAL for RANDOMIZED patients, Version: 2.0 final, 09-10-2015**

### If tube was dislocated, how often did this occur*  
|__|__| times (0-10)

* Unknown: -1

### Was tube removed because of side effects
- O No
- O Yes
- O Unknown
- O Patients request
- O Doctors' advice
- O Other; _________________

### Date of tube removal†§  
|__|__| - |__|__| - |__|__| (dd-mm-yy)

*Unknown: 99-99-99  
† All reasons for tube removal  
§ Patients are normally discharged with tube in situ. Date may be later than date of discharge.

### If the tube was removed, was it replaced
- O No
- O Yes
- O Unknown

### Date of tube replacement†  
|__|__| - |__|__| - |__|__| (dd-mm-yy)

*Unknown: 99-99-99  
† Tube replacement directly following tube removal (within 48 hours), not tube placement during readmission

### Date of replaced tube removal†§  
|__|__| - |__|__| - |__|__| (dd-mm-yy)

*Unknown: 99-99-99  
† All reasons for tube removal  
§ Patients are normally discharged with tube in situ. Date may be later than date of discharge.

### Was a duodenal or jejunal tube placed at any point*
- O No
- O Yes
- O Unknown

### Reason(s) for duodenal or jejunal tube placement
- O Nasogastric tube dislocation
- O Continuation of vomiting
- O Other; _________________

---
7. **Level of care**

Did patient return/go to primary care after remission (or stabilization) of HG during the course of pregnancy

<table>
<thead>
<tr>
<th>O No</th>
<th>O Yes</th>
<th>O Unknown</th>
</tr>
</thead>
</table>

*Date of return to primary care* |

*Unknown: 99-99-99*

Was patient afterwards referred to secondary/tertiary care

<table>
<thead>
<tr>
<th>O No</th>
<th>O Yes</th>
<th>O Unknown</th>
</tr>
</thead>
</table>

*Date of referral* |

*Unknown: 99-99-99*

Was there a medical indication(s) for (continuation of) antenatal/perinatal care after 20 weeks gestation in a secondary/tertiary center

<table>
<thead>
<tr>
<th>O No*</th>
<th>O Yes</th>
<th>O Unknown</th>
</tr>
</thead>
</table>

*Continuation of HG

| PE/HELLP

| IUGR

| Diabetes (gravidarum/type I/type II)

| Premature rupture of membranes

| Threatening preterm labour

| Placenta previa (marginalis/totalis)

| Vaginal bleeding 2nd/3rd trimester

| Placental abruption (partial/total)

| Breech/transverse position

| Other

* No secondary/tertiary care, or patients wish for secondary/tertiary care without medical indication
7.1 Routine pregnancy check-ups

Was maternal weight measured during routine pregnancy check-ups (primary/secondary/tertiary care)

| 0 No | 0 Yes | 0 Unknown |

<table>
<thead>
<tr>
<th>Measurement</th>
<th>Weight:* kg (35-250)</th>
<th>Date of measurement:* (dd-mm-yy)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

7.2 Hospital readmissions for HG

Number of readmissions for HG* | _ _ _ _ (0-10)

*After admission in which patient was randomized

Fill out dates of readmission(s) because of HG

<table>
<thead>
<tr>
<th>Readmission*</th>
<th>Hospital admission (dd-mm-yy)</th>
<th>Discharge home (dd-mm-yy)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*If more than 5 readmissions, please print/copy this page another time to fill out dates of subsequent readmissions
## Fill out details on each readmission because of HG*

*If more than 1 readmission, please print/copy this page another time to fill out details on subsequent readmission(s)

<table>
<thead>
<tr>
<th>Readmission No</th>
<th>___</th>
<th>___</th>
<th>(0-10)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weight at readmission*</td>
<td>___</td>
<td>___</td>
<td>___ kg (35-250)</td>
</tr>
<tr>
<td>Urinary ketones*</td>
<td>___</td>
<td>___</td>
<td>___</td>
</tr>
<tr>
<td>Intravenous drip</td>
<td>___</td>
<td>___</td>
<td>___</td>
</tr>
<tr>
<td>Was i.v. drip placed on day of readmission</td>
<td>___</td>
<td>___</td>
<td>___</td>
</tr>
<tr>
<td>Date of placement*</td>
<td>___</td>
<td>___</td>
<td>___ (dd-mm-yy)</td>
</tr>
<tr>
<td>Was i.v. drip removed on day of discharge</td>
<td>___</td>
<td>___</td>
<td>___</td>
</tr>
<tr>
<td>Date of removal*</td>
<td>___</td>
<td>___</td>
<td>___ (dd-mm-yy)</td>
</tr>
<tr>
<td>Tube feeding</td>
<td>___</td>
<td>___</td>
<td>___</td>
</tr>
<tr>
<td>Was tube placed on day of readmission</td>
<td>___</td>
<td>___</td>
<td>___</td>
</tr>
<tr>
<td>Date of placement*</td>
<td>___</td>
<td>___</td>
<td>___ (dd-mm-yy)</td>
</tr>
<tr>
<td>Was tube removed on day of discharge</td>
<td>___</td>
<td>___</td>
<td>___</td>
</tr>
<tr>
<td>Date of removal*</td>
<td>___</td>
<td>___</td>
<td>___ (dd-mm-yy)</td>
</tr>
<tr>
<td>Antiemetics</td>
<td>___</td>
<td>___</td>
<td>___</td>
</tr>
<tr>
<td>Vitamins</td>
<td>___</td>
<td>___</td>
<td>___</td>
</tr>
</tbody>
</table>

*Unknown: 99-99-99

*If in medical or nursing record only: 'i.v. drip according to protocol', fill out what is prescribed according to local protocol for HG

*Prescribed as antiemetic, not because of fetal lung ripening

*Patients are normally discharged with tube in situ. Date may be later than date of discharge

---

CRF MOTHER TRIAL for RANDOMIZED patients, Version: 2.0 final, 09-10-2015
8. Delivery

Onset of labour
Ο Spontaneously
Ο Primary caesarean section
Ο Induction

Induction of labour or primary caesarean section
Ο No ○ Yes ○ Unknown
□ Maternal indication*
□ Fetal indication†
□ Elective§

*Such as hypertension
†Such as growth restriction or breech position
§Non-medical reason such as patients wish or ‘DDD’ (discrepancy draagkracht draaglast)

Place of birth
Ο Home delivery, primary care
Ο Hospital delivery, primary care
Ο Hospital delivery, secondary/tertiary care

Analgesics during labour*
Ο No ○ Yes ○ Unknown

*Not during surgical intervention, e.g. caesarean section

Route of delivery
Ο vaginally
Ο caesarean section

Delivery of placenta
Ο Spontaneously
Ο Manual removal
Ο Manual removal during caesarean section

Placental weight measured
Ο No ○ Yes ○ Unknown

*Placental weight* ○ | | | | grams (100-2000)

*Unknown: -1
| Case Number | __|__|__| - |__|__|__| |

**Total haemorrhage** | __|__|__|__| ml (50-9999) |

**Maternal death**

O No
O Yes

| Date of death | __| __| - |__|__| - |__|__| (dd-mm-yy) |

*Within 6 weeks of delivery
†Fill out SAE form
9. **Neonatal birth data**

**Multiple pregnancy**
- O No
- O Yes

**Child no.** *|__|__| 0-3*

*If multiple, please print/copy this page another time to fill out details for subsequent child(ren)*

**Date of birth** |__|__| - |__|__| - |__|__| (dd-mm-yy)

**Live birth**
- O Yes
- O No, deceased during delivery < 24 hours postpartum*
- O No, deceased before delivery*

**Estimated date of death** |__|__| - |__|__| - |__|__| (dd-mm-yy)

*Fill out SAE form

**Apgar score (5 min)** |__|__| (00-10)

**Umbilical cord pHs measured**
- O No
- O Yes
- O Unknown

- Arterial pH |__|__| - |__|__| (6.00-7.70)
- Venous pH |__|__| - |__|__| (6.00-7.70)

**Sex**
- O Boy
- O Girl

**Birth weight** |__|__|__|__| grams (300-6500)

**Neonatal death ≥ 24 hours postpartum**
- O No
- O Yes*
- O Unknown

**Date of death** |__|__| - |__|__| - |__|__| (dd-mm-yy)

*Fill out SAE form
10. Hospital admission postpartum

10.1 Maternal hospital admission

<table>
<thead>
<tr>
<th>Maternal hospital admission &lt; 12 hours postpartum</th>
</tr>
</thead>
<tbody>
<tr>
<td>O No</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>□ Maternal indication</td>
</tr>
<tr>
<td>□ Neonatal indication</td>
</tr>
</tbody>
</table>

*Fill out SAE form

If hospital admission on maternal indication: indication(s) for admission

- Suspected infection
- Tromboembolic complication
- Hypertensive disorder
- Postpartum haemorrhage
- Post-caesarean
- Eclampsia/HELLP
- Other

Date of hospital admission equal to date of delivery

<table>
<thead>
<tr>
<th>O No</th>
<th>O Yes</th>
<th>O Unknown</th>
</tr>
</thead>
</table>

Date of hospital admission | __ | __ | - | __ | __ | - | __ | __ | (dd-mm-yy)

Date of discharge home | __ | __ | - | __ | __ | - | __ | __ | (dd-mm-yy)
10.2 Neonatal hospital admission

Multiple pregnancy
☐ No ☑ Yes

Child no. * | ___ | (0-3)
*If multiple, please print/copy this page another time to fill out details for subsequent child(ren)

Neonatal hospital admission <6 weeks postpartum
☐ No ☑ Yes*
☐ Maternal indication
☐ Neonatal indication
*Fill out SAE form

If hospital admission on neonatal indication: indication(s) for admission*
☐ Small for gestational age (defined as birth weight < P10)
☐ Large for gestational age (defined as birth weight > P 90)
☐ Congenital anomaly or suspicion for abnormality
☐ Hypoglycaemia (i.v. treatment needed)
☐ Hyperbilirubinemia (phototherapy or transfusion needed)
☐ Infection/ Sepsis (suspected or proven positive culture)
☐ Convulsions
☐ Other
*As reported in the final discharge letter of pediatrician

Date of hospital admission equal to date of birth
☐ No ☑ Yes ☐ Unknown

Date of hospital admission | ___ | ___ | ___ | ___ | ___ | ___ | (dd-mm-yy)
Date of discharge home | ___ | ___ | ___ | ___ | ___ | ___ | (dd-mm-yy)

Note: please attach an anonymized copy of the final pediatric discharge letter; mark this letter with maternal case number on every page
11. **(Serious) Adverse Events**

**In this pregnancy, has there been an adverse event(s) (AE)**

<table>
<thead>
<tr>
<th></th>
<th>O No</th>
<th>O Yes</th>
<th>O Unknown</th>
</tr>
</thead>
<tbody>
<tr>
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Please specify; ___________________________

*Any undesirable event during the course of the study whether or not considered related to the intervention, spontaneously reported by patient, or observed by caregiver/research staff, but not leading to hospital admission (see SAE). Examples of AE’s are: broken leg, fall from stairs, brain contusion etc.

**In this pregnancy, has there been a Serious Adverse Event(s) (SAE)**

<table>
<thead>
<tr>
<th></th>
<th>O No</th>
<th>O Yes</th>
<th>O Unknown</th>
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- Miscarriage
- Hospital admission(s) after initial admission at study entry†
- Significant prolongation of hospital admission
- Maternal (pregnancy) complications§
- Complications due to tube placement (e.g. aspiration/intestinal bleeding/intestinal perforation)
- Maternal death
- Neonatal hospital admission¥
- Birth defect/congenital anomaly neonate
- Perinatal death
- Other; _______________________________

*Any of the above mentioned situations, or AE causing significant disability to the patient, or AE requiring interventions to prevent a SAE from happening.
†Any reason, excluding hospital admissions for HG (these are already reported in CRF)
§Think of ICU admission (e.g. due to massive postpartum haemorrhage), uterine rupture, placental abruption etc.
¥Any reason

**Note: in all cases of SAE fill out SAE form (see study website → documents)**

---

CRF MOTHER TRIAL for RANDOMIZED patients, Version 2.0 final, 09-10-2015
12. End Study Form

I have filled out this form accurately and truthfully

Name: ________________________________

Date: ___ ___ ___ ___ ___ ___ ___ ___ ___ ___ ___ ___ ___ (dd-mm-yyyy)

Signature: ______________________________

12.1 Study stop

Has patient reached the endpoint of the study (childbirth)
O No  O Yes  O Lost to follow up

Has patient (partially) withdrawn from study
O No  O Yes

☐ Withdrawn from biobank
☐ Withdrawn from filling out diaries
☐ Completely withdrawn from study participation*

*Patient has explicitly expressed wish that patient information may no longer be used for this study

Reason for (partial) study withdrawal
O Unknown
O Adverse event
O Other; __________________

---------------------------------------------------------------
END OF Case Report Form
THANK YOU FOR FILLING OUT THIS FORM
---------------------------------------------------------------
13. **Supplement for patients included before 01-10-2014**

*Note: Is the patient included before 01-10-2014? Please try to answer the following questions.*

For patients included from 01-10-2014 onwards, the following questions have been asked in questionnaire A (at inclusion) and filling out this supplement is not necessary.

### 13.1 Maternal background

#### Ethnicity
- Unknown
- Dutch
- European / North American / Oceanian
- South American
- Turkish
- Surinamese
- Antillean / Aruban / Cape Verdian
- African (Sub-Sahara)
- Indonesian / Moluccan / Japanese
- Indian / Pakistani
- Other; __________________________

#### Highest finished education
- Unknown
- Primary school
- Secondary school
- Lower professional school (VMBO)
- Medium professional school (MBO)
- Higher professional school (HBO)
- University (WO)
- Other; __________________________

#### Relationship
- Unknown
- Single
- Living together with partner
### 13.2 Current pregnancy

**Mode of conception**
- Unknown
- Spontaneous (natural route)
- IUI (intrauterine insemination)
- Ovulation-induction (medication to effectuate menstrual cycle)
- IVF (in vitro fertilization)
- ICSI (intracytoplasmic sperm injection)
- Other: __________________________

<table>
<thead>
<tr>
<th>Nausea reported since gestational age</th>
<th>__</th>
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<th>weeks (0-20)</th>
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<table>
<thead>
<tr>
<th>Vomiting reported since gestational age</th>
<th>__</th>
<th>__</th>
<th>weeks (0-20)</th>
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