

# Does repeat placental growth factor blood sample testing reduce harm from pre-eclampsia to babies?

<b>Submission date</b> 10/10/2019	<b>Recruitment status</b> No longer recruiting	<input checked="" type="checkbox"/> Prospectively registered <input checked="" type="checkbox"/> Protocol
<b>Registration date</b> 25/11/2019	<b>Overall study status</b> Completed	<input type="checkbox"/> Statistical analysis plan <input checked="" type="checkbox"/> Results
<b>Last Edited</b> 12/01/2026	<b>Condition category</b> Pregnancy and Childbirth	<input type="checkbox"/> Individual participant data

## Plain English summary of protocol

### Background and study aims

Pre-eclampsia is a condition occurring only in pregnancy, thought to be caused by the way the placenta implants into the wall of the womb. Women with pre-eclampsia can suffer from high blood pressure, problems with their kidneys, liver and blood clotting. The problems with the placenta can mean that the baby's growth is affected. In some cases the baby can be stillborn. Once diagnosed, the only cure is to deliver the baby. If pre-eclampsia occurs before 37 weeks of pregnancy, women may need to be admitted to hospital to have treatment and monitoring for complications, whilst planning for safe delivery of the baby. Some women become unwell very quickly and need to have their babies delivered; others have long stays in hospital. It can be difficult to identify women at high-risk of severe complications of pre-eclampsia. This study looks at the levels of a protein produced by the placenta called Placenta Growth Factor (PLGF). Women with suspected pre-eclampsia can have a simple blood test for this protein. Studies have shown that women with very low PLGF levels are at greater risk of severe pre-eclampsia and stillbirth. However, it is not known how PLGF levels change over time. When pre-eclampsia is suspected, it is difficult to predict how severely a woman and her baby will be affected. The aim of this study is to find out whether using repeated blood samples can help to reduce severe complications for babies, and for women.

### Who can participate?

Pregnant women aged 18 or over suspected of having pre-eclampsia

### What does the study involve?

If a woman agrees to take part, she will be asked to sign a consent form, and details about her and her pregnancy will be put into a secure computer database. This will also be noted in her hospital maternity records. The study computer will then select for her to have repeat PLGF-based blood tests with the results known or not known to her, her doctors and her midwives while she is pregnant. There will be a 50:50 chance of being in either study group. She will then have routine bloods to test for pre-eclampsia that will include her first PLGF-based test. Her doctor will then use the results of these tests to guide her care following their hospital standard practice. Depending on the result of the blood tests (including the PLGF-based test), the doctors

and midwives will decide if a woman needs to be admitted to hospital or how often they will need to see her again in her pregnancy to make sure she and the baby are okay. When she is asked to have repeat blood tests for routine follow up care, we will ask her for an extra 10 ml or two teaspoons of blood for a PlGF-based test. Depending on which study group she is in, the doctors and midwives will be given or not given the result of the PlGF-based test. If the doctors and midwives are given the PlGF-based test result, they can use this to guide ongoing care, in addition to following their hospital standard practice. If the doctors and midwives are not given the PlGF-based test results ongoing antenatal care will be exactly the same as if women were not taking part in the study. Women will only be asked to provide an extra blood sample for this study once per week or once every two weeks (depending on the result of the first test) and only for a maximum of four times during their pregnancy.

What are the possible benefits and risks of participating?

The first PARROT study showed that some women may spend less time in hospital, and if they have an abnormal result they may benefit from their doctors and midwives having more information about their pre-eclampsia condition. There are no expected serious side effects to having the blood tests and participants will be having blood tests as part of their normal clinical care.

Where is the study run from?

1. Guy's and St Thomas' NHS Foundation Trust (UK)
2. Manchester University NHS Foundation Trust (UK)
3. Leeds Teaching Hospitals NHS Trust (UK)
4. Liverpool Women's NHS Foundation Trust (UK)
5. Royal United Hospitals Bath NHS Foundation Trust (UK)
6. Bradford Teaching Hospitals NHS Foundation Trust (UK)
7. St George's University Hospitals NHS Foundation Trust (UK)
8. Kingston Hospital NHS Foundation Trust (UK)
9. Chelsea and Westminster Hospital NHS Foundation Trust (UK)
10. North Bristol NHS Trust (UK)
11. University Hospitals Bristol NHS Foundation Trust (UK)
12. Imperial College Healthcare NHS Trust (UK)
13. NHS Lothian (UK)
14. Ashford and St Peter's Hospitals NHS Foundation Trust (UK)
15. Croydon Health Services NHS Trust (UK)
16. Norfolk and Norwich University Hospitals NHS Foundation Trust (UK)
17. University College London Hospitals NHS Foundation Trust (UK)
18. Nottingham University Hospitals NHS Trust (UK)
19. Warrington and Halton Hospitals NHS Foundation Trust (UK)

When is the study starting and how long is it expected to run for?

April 2019 to March 2023

Who is funding the study?

1. Moulton Charitable Trust (UK)
2. Tommy's Baby Charity (UK)

Who is the main contact?

Dr Louise Webster

[louise.m.webster@kcl.ac.uk](mailto:louise.m.webster@kcl.ac.uk)

# Contact information

## Type(s)

Public

## Contact name

Dr Louise Webster

## Contact details

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# Additional identifiers

## Protocol serial number

265824; CPMS: 43092

# Study information

## Scientific Title

Placental growth fActor Repeat sampling for Reduction of adverse perinatal Outcomes in women with suspecTed pre-eclampsia

## Acronym

PARROT-2

## Study objectives

Repeat PlGF-based testing, in women presenting with suspected preterm pre-eclampsia, reduces adverse perinatal outcomes (perinatal death/neonatal unit admission).

## Ethics approval required

Old ethics approval format

## Ethics approval(s)

Approved 11/11/2019, East of England - Cambridge East Research Ethics Committee (The Old Chapel, Royal Standard Place, Nottingham, NG1 6FS, UK; Tel: +44 (0)207 104 8101; Email: NRESCcommittee.EastofEngland-CambridgeEast@nhs.net), ref: 19/EE/0322

## Study design

Multi-centre randomized controlled trial

## Primary study design

Interventional

## **Study type(s)**

Diagnostic

## **Health condition(s) or problem(s) studied**

Pre-eclampsia

## **Interventions**

Multi-centre RCT of revealed versus concealed repeat PlGF-based testing in women presenting with suspected pre-eclampsia between 22+0 and 35+6 weeks' gestation. The trial will be conducted in at least 15 maternity units across England. Recruitment will run for approximately 18 months in total.

All women participating in the trial will have an initial revealed PlGF-based test, allowing the clinician to formulate an individualised management plan.

Randomisation will be via a secure web-based platform (MedSciNet) with 50% of women assigned to revealed repeat PlGF-based testing and 50% of women assigned concealed repeat PlGF-based testing.

For the trial, the women will be asked to provide one extra tube of blood (as far as possible at the same time as clinical blood samples) up to four times during the rest of their pregnancy as per the schedule below. The results of the repeat PlGF-based test will be revealed to the health care professionals and the women in the intervention arm and used in addition to the other clinical features to inform ongoing management plan integrated with the NICE Hypertension in Pregnancy Guideline. The results of the repeat tests will be concealed in the usual care arm. It is recognised that some women will only provide one sample; from previous studies it is anticipated that most women will provide two samples as the majority of women will be delivered within that time interval.

For both the revealed repeat testing and concealed repeat testing groups, the repeat sampling strategy will be based on the first PlGF test result as follows:

1. If PlGF  $\leq 100$  pg/ml (including women  $< 12$  pg/ml) or sFlt-1/PlGF ratio  $> 38$ , i.e. at higher risk, sampling will be weekly whilst attending for clinical review.
2. If PlGF  $> 100$  pg/ml or sFlt-1/PlGF ratio  $\leq 38$  (lower risk) and asymptomatic of pre-eclampsia, sampling will be every two weeks (+/- 7 days) whilst attending for routine antenatal checks. If a woman presents  $\geq 7$  days from last sample and is symptomatic, an additional sample can be taken and reported.

Women will only be asked to provide repeat samples while they are still pregnant and pregnancy outcome data for mother and baby will be collected by the site research teams from care records without the need for further review in person of the participants.

## **Intervention Type**

Other

## **Primary outcome(s)**

Composite of:

1. Stillbirth defined as death of a fetus after 24 weeks' gestation and before birth collected by 6 weeks post birth
2. Early neonatal death defined as death occurring within the first 7 days of life collected by 6 weeks post birth

3. Neonatal unit admission defined as admission of the neonate to the neonatal unit and captured within the first 6 weeks from birth

### **Key secondary outcome(s)**

Additional fetal and neonatal outcomes:

1. Late neonatal death defined as neonatal death occurring between 7 and 28 days after birth and captured from hospital records by 6 weeks post birth
2. Need for respiratory support on Neonatal Unit defined as the need for CPAP/high flow /endotracheal ventilation and recorded by 6 weeks post birth
3. Gestational age at delivery measured in days and recorded by 6 weeks post birth
4. Birthweight centile <10th calculated using recorded birth weight and using the Intergrowth 21 birthweight centile calculator and calculated by the trial statistician prior to data analysis

Added 07/07/2022:

5. Survival to discharge without severe morbidity: defined as survival to neonatal discharge without any of the following: bronchopulmonary dysplasia, retinopathy of prematurity, severe necrotising enterocolitis, brain injury, late-onset sepsis

Maternal secondary outcomes (between enrollment and delivery):

1. Proportion of women diagnosed with pre-eclampsia defined using the ISSHP definition and captured by 6 weeks post birth
2. Severe adverse maternal outcome composite defined by the fullPIERS consensus and captured by 6 weeks post birth
3. Systolic blood pressure  $\geq 160$  mmHg measured during routine blood pressure readings captured in maternity records and occurring on at least one occasion between study enrolment and birth of the baby
4. Concealed first repeat PlGF-based test performance (with comparison against currently utilised tests) for clinically indicated delivery for diagnosed pre-eclampsia within 14 days measured in peripheral blood samples at 1 to 3 weeks post study enrollment and analysed following completion of the trial

Health economic outcomes:

1. Perinatal: intensive care, high dependency and special care unit days measured as total of these days and captured by 6 weeks post birth
2. Maternal: antenatal outpatient attendances and inpatient days; intensive care unit use measured as total numbers of each of these antenatal care episodes and captured by 6 weeks post birth

### **Completion date**

31/03/2023

## **Eligibility**

### **Key inclusion criteria**

1. Women aged 18 years or more between 22+0 and 35+6 weeks' gestation with clinical suspicion of pre-eclampsia
2. Viable singleton pregnancy
3. Able to give written informed consent

### **Participant type(s)**

Patient

**Healthy volunteers allowed**

No

**Age group**

Adult

**Lower age limit**

18 years

**Upper age limit**

55 years

**Sex**

Female

**Total final enrolment**

1252

**Key exclusion criteria**

Confirmed preterm pre-eclampsia at presentation

**Date of first enrolment**

06/12/2019

**Date of final enrolment**

30/09/2022

**Locations****Countries of recruitment**

United Kingdom

England

Scotland

**Study participating centre**

**Guy's and St Thomas' NHS Foundation Trust**

St. Thomas's Hospital

249 Westminster Bridge Road

London

England

SE1 7EH

**Study participating centre**

**Manchester University NHS Foundation Trust**

Cobbett House  
Oxford Road  
Manchester Greater  
Manchester  
England  
M13 9WL

**Study participating centre**

**Leeds Teaching Hospitals NHS Trust**

Beckett Street  
Leeds  
England  
LS9 7TF

**Study participating centre**

**Liverpool Women's NHS Foundation Trust**

Crown Street  
Liverpool  
England  
L8 7SS

**Study participating centre**

**Royal United Hospitals Bath NHS Foundation Trust**

Combe Park  
Bath  
England  
BA1 3NG

**Study participating centre**

**Bradford Teaching Hospitals NHS Foundation Trust**

Duckworth Lane  
Bradford  
England  
BD9 6RJ

**Study participating centre**

**St George's University Hospitals NHS Foundation Trust**

Blackshaw Road  
Tooting  
London

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SW17 0QT

**Study participating centre**  
**Kingston Hospital NHS Foundation Trust**  
Galsworthy Road  
London  
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KT2 7QB

**Study participating centre**  
**Chelsea and Westminster Hospital NHS Foundation Trust**  
369 Fulham Road  
London  
England  
SW10 9NH

**Study participating centre**  
**North Bristol NHS Trust**  
Southmead Road  
Westbury-on-Trym  
Bristol  
England  
BS10 5NB

**Study participating centre**  
**University Hospitals Bristol NHS Foundation Trust**  
Marlborough Street  
Bristol  
England  
BS13NU

**Study participating centre**  
**Imperial College Healthcare NHS Trust**  
St. Mary's Hospital  
Praed Street  
London  
England  
W2 1NY

**Study participating centre**

**NHS Lothian**

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2-4 Waterloo Place  
Edinburgh  
Scotland  
EH1 3EG

**Study participating centre**

**Ashford and St Peter's Hospitals NHS Foundation Trust**

Guildford Road  
Surrey  
England  
KT160PZ

**Study participating centre**

**Croydon Heath Services NHS Trust**

London Road  
Thornton Heath  
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England  
CR7 7YE

**Study participating centre**

**Norfolk and Norwich University Hospitals NHS Foundation Trust**

Colney Lane  
Colney  
Norwich  
England  
NR4 7UY

**Study participating centre**

**University College London Hospitals NHS Foundation Trust**

250 Euston Road  
London  
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NW1 2PG

**Study participating centre**

**Nottingham University Hospitals NHS Trust**  
Derby Road  
Nottingham  
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NG7 2UH

**Study participating centre**  
**Warrington and Halton Hospitals NHS Foundation Trust**  
Lovely Lane  
Warrington  
England  
WA5 1QG

## **Sponsor information**

**Organisation**  
King's College London

**ROR**  
<https://ror.org/0220mzb33>

**Organisation**  
Guy's and St Thomas' NHS Foundation Trust

## **Funder(s)**

**Funder type**  
Charity

**Funder Name**  
Moulton Charitable Trust

**Funder Name**  
Tommy's Baby Charity

**Alternative Name(s)**

**Funding Body Type**

Private sector organisation

## Funding Body Subtype

Other non-profit organizations

## Location

United Kingdom

# Results and Publications

## Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study are/will be available upon request from Louise Webster (louise.m.webster@kcl.ac.uk) and Lucy Chappell (lucy.chappell@kcl.ac.uk).

Type of data: quantitative

When the data will become available and for how long: 01/12/2022

By what access criteria data will be shared including with whom: to be determined at a later date

For what types of analyses, and by what mechanism: to be determined at a later date

Whether consent from participants was obtained: not applicable

Comments on data anonymisation: Data would only be supplied in fully anonymised format

Any ethical or legal restrictions: not aware of any

Any other comments: no

## IPD sharing plan summary

Available on request

## Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<a href="#">Results article</a>		08/02/2024	12/02/2024	Yes	No
<a href="#">Results article</a>		06/05/2024	07/05/2024	Yes	No
<a href="#">Results article</a>	Health economic assessment	06/01/2026	12/01/2026	Yes	No
<a href="#">Protocol article</a>		02/09/2022	05/09/2022	Yes	No
<a href="#">HRA research summary</a>			28/06/2023	No	No