PARROT Ireland: Placental growth factor in Assessment of women with suspected pre-eclampsia to reduce maternal morbidity: a Stepped Wedge Cluster Randomised Control Trial Research Study Protocol

Deirdre Hayes-Ryan,¹,² Karla Hemming,³ Fionnuala Breathnach,⁴ Amanda Cotter,⁵ Declan Devane,⁶,⁷ Alyson Hunter,⁸ Fionnuala M McAuliffe,⁹ John J Morrison,¹⁰ Deirdre J Murphy,¹¹ Ali Khashan,¹²,¹³ Brendan McElroy,¹⁴ Aileen Murphy,¹⁴ Eugene Dempsey,¹³,¹⁵ Keelin O’Donoghue,²,¹³ Louise C Kenny¹³,¹⁶

ABSTRACT

Introduction Women presenting with suspected pre-eclampsia are currently triaged on the basis of hypertension and dipstick proteinuria. This may result in significant false positive and negative diagnoses resulting in increased morbidity or unnecessary intervention. Recent data suggest that placental growth factor testing may be a useful adjunct in the management of women presenting with preterm pre-eclampsia. The primary objective of this trial is to determine if the addition of placental growth factor testing to the current clinical assessment of women with suspected preterm pre-eclampsia, is beneficial for both mothers and babies.

Methods and analysis This is a multicentre, stepped wedge cluster, randomised trial aiming to recruit 4000 women presenting with symptoms suggestive of preterm pre-eclampsia between 20 and 36+6 weeks’ gestation. The intervention of an unblinded point of care test, performed at enrolment, will quantify maternal levels of circulating plasma placental growth factor. The intervention will be rolled out sequentially, based on randomisation, in the seven largest maternity units on the island of Ireland. Primary outcome is a composite outcome of maternal morbidity (derived from the modified fullPIERS model). To ensure we are not reducing maternal morbidity at the expense of earlier delivery and worse neonatal outcomes, we have established a co-primary outcome which will examine the effect of the intervention on neonatal morbidity, assessed using a composite neonatal score. Secondary analyses will examine further clinical outcomes (such as mode of delivery, antenatal detection of growth restriction and use of antihypertensive agents) as well as a health economic analysis, of incorporation of placental growth factor testing into routine care.

Ethics and dissemination Ethical approval has been granted from each of the seven maternity hospitals involved in the trial. The results of the trial will be presented both nationally and internationally at conference and published in an international peer-reviewed journal.

Strengths and limitations of this study

- Randomised trial.
- Multiple sites with wide geographic distribution.
- Stepped wedge design.
- Placental growth factor testing only in the intervention arm.

Trial registration number NCT02881073.

BACKGROUND

Pre-eclampsia is characterised by hypertension and proteinuria, complicates 2%–8% of pregnancies, and is associated with significant maternal and neonatal morbidity and mortality.¹ Currently, women who present with suspected pre-eclampsia are triaged on the basis of hypertension and dipstick proteinuria. Both of these clinical end points are subject to observer error and poor test accuracy, with false positive and negative diagnoses of pre-eclampsia occurring in clinical practice.²⁻⁵ Current biochemical tests are imperfect at stratifying women for more intensive surveillance as they only identify advanced disease where there is already marked end-organ damage.⁶ While biomarkers and imaging techniques have been evaluated for improving detection, none has adequate sensitivity and/or specificity for the diagnosis of pre-eclampsia.⁷

Placental growth factor (PIGF) belongs to the vascular endothelial growth factor (VEGF) family and represents a key regulator
of angiogenic events in pathological conditions. PIGF exerts its biological function through the binding and activation of the receptor Flt-1. In pre-eclampsia, it is thought that endothelial dysfunction leads to an increased level of a circulating decoy receptor, known as soluble Flt-1 (sFlt-1), a soluble receptor for both vascular endothelial growth factor type A (VEGF-A) and PIGF. Circulating levels of sFlt-1 are increased in pre-eclampsia and particularly in the early onset form of the disease, resulting in reduced levels of free VEGF-A and PIGF in the maternal circulation. Thus, the endothelial dysfunction observed in pre-eclampsia may be due to excess neutralisation of VEGF-A and PIGF by circulating sFlt-1. Levine et al showed that in normal pregnancy, PIGF levels track the development of the placenta, peaking at about 32 weeks’ gestation when the placenta is developed fully and then declining until delivery. However, in pre-eclampsia, this rise and fall is considerably lower throughout pregnancy, and levels are strikingly lower when the condition presents clinically.

The PELICAN study (Plasma Placental Growth Factor in the Diagnosis of Women with Pre-Eclampsia Requiring Delivery Within 14 Days) was the first and largest prospective evaluation of PIGF in women presenting with suspected pre-eclampsia. This blinded observational cohort study was conducted in seven consultant-led maternity units in the UK and Ireland between January 2011 and February 2012. It enrolled women being investigated for suspected pre-eclampsia, quantified their plasma PIGF using a point of care device, the Alere Triage PIGF test, but did not reveal the result to their clinician. The study found that a PIGF value <100 pg/mL, in women presenting prior to 35 completed weeks’ gestation had a negative predictive value of 98% (95% CI 93 to 99.5) and a positive predictive value of 44% (95% CI 36 to 52) in determining those that would require delivery for a confirmed diagnosis of pre-eclampsia within the next 14 days. The study reported a PIGF <100 pg/mL to be a better predictor than all other current commonly used predictive tests of pre-eclampsia, either singly or in combination (blood pressure, urinalysis or biochemical markers) with an area under the receiver operating characteristic curve for low PIGF of 0.87 compared with 0.76 for the next best predictor.

The PROGNOSIS study (Predictive value of the sFlt-1:PIGF ratio in women with suspected preeclampsia) was a prospective, multicentre, blinded, observational study conducted in 14 countries from 2011 to 2014. Its aim was to derive and validate a ratio of serum sFlt-1 to PIGF that would be predictive of the absence or presence of pre-eclampsia in the short term. It included women with singleton pregnancies from 24 weeks to 36+6 weeks’ gestation in whom a clinical suspicion of pre-eclampsia existed. The Elecsys immunoassay was used to quantify levels of PIGF and sFlt-1. The development cohort of over 500 participants identified a sFlt-1:PIGF ratio of 38 as having an important predictive value. The subsequent validation cohort, again with over 500 participants, reported a negative predictive value of 99.3% (95% CI 97.9 to 99.9) for ruling out pre-eclampsia within 1 week. Interestingly, the same cut-off of 38 was predictive of the absence of fetal adverse outcomes within 1 week; negative predictive value of 99.3% (95% CI 97.9 to 99.9). The study showed that an sFlt-1:PIGF ratio of 38 or lower can be used to predict the short-term absence of pre-eclampsia and adverse fetal events in women in whom the syndrome is suspected clinically. The positive predictive value, a diagnosis of pre-eclampsia, eclampsia or the HELLP syndrome (haemolysis, elevated liver enzymes and a low platelet count) within 4 weeks, was 36.7% (95% CI 28.4 to 45.7) using the same sFlt-1:PIGF ratio of 38. Post hoc analysis however showed this was still an improvement in prediction compared with the use of clinical variables such as blood pressure and urinalysis alone.

The National Institute for Health and Clinical Excellence, UK (NICE) has recently published guidance on incorporation of PIGF testing, in addition to clinical assessment, in women presenting with suspected pre-eclampsia from 20 to 34+6 weeks’ gestation. It advises that the Triage PIGF test or Elecsys immunoassay sFlt-1/PIGF ratio test may be used, in combination with clinical assessment, to ‘rule-out’ pre-eclampsia in this group of women. However, it advises that these tests should not yet be used to diagnose pre-eclampsia until further research is available, specifically on how an abnormal PIGF result would affect management decisions regarding timing and gestation of delivery and the outcomes associated with this.

The objective of this randomised trial is to evaluate the impact of knowledge of PIGF measurement on clinically relevant outcomes. We hypothesise that adding PIGF measurement to current clinical assessment of women with suspected pre-eclampsia prior to 37 weeks’ gestation will reduce associated maternal morbidity through improved risk stratification, earlier diagnosis and targeted management of women with the disease. Any intervention in late pregnancy may have an impact on the fetus. On the one hand, earlier diagnosis of pre-eclampsia may precipitate earlier delivery and lead to an increase in neonatal morbidity and mortality secondary to iatrogenic prematurity. Conversely, improved identification of those neonates at highest risk of imminent placental dysfunction may reduce neonatal morbidity by allowing for timely intervention. It is therefore imperative that full evaluation of both potential benefit and harm is conducted before PIGF testing is implemented routinely into clinical practice. If this trial demonstrates a beneficial impact on maternal morbidity and/or neonatal morbidity, alongside a favourable health economic assessment, then there would be a strong case for incorporating PIGF testing into routine clinical investigations for women presenting with suspected pre-eclampsia before 37 weeks’ gestation in a wide variety of healthcare settings.
Methods and design

Study design

PARROT Ireland is a multicentre, stepped wedge cluster-controlled trial of PlGF measurement in women presenting with suspected pre-eclampsia from 20 weeks and prior to 37 weeks’ gestation. As implementation of a diagnostic test may alter physician management, a cluster design was chosen rather than individual randomisation. This allows for a change in management to occur at a hospital rather than at an individual woman level, which is preferable in trials involving a diagnostic test and allows the clinical influence of the additional test to be evaluated in a pragmatic fashion.17 Each maternity hospital acts as a cluster. All clusters commenced the trial in the control arm and in turn, each cluster transitions at random from the control to the intervention at prespecified time points. Once a cluster has changed over to the intervention, it continues as such for the remainder of the trial so that by the end of the trial all clusters will be in the intervention arm (figure 1). A stepped wedge design was chosen so as to increase the social acceptability of the trial to the seven hospitals (the stakeholders/decision makers in all of the hospitals expressed a desire to participate in a trial in which they were guaranteed to get the intervention); and because a trial with just seven clusters risks baseline imbalance in a parallel design.

The trial will continue for a period of 22 months, and with seven clusters the interval between transitions is approximately 3 months in duration. A restricted method of randomisation was used to provide a balance in total (expected) number of observations across intervention and control periods (details below).18–20 There is a short transition period of 1 week whenever a new cluster transitions from control to the intervention. Data collected during this transition period will not be included in any analysis of outcomes. Recruitment will stop on a prespecified fixed date in late April 2019 and the study will end when the last recruited participant and neonate are discharged and all outcome data collected.

Setting and participants

The trial is being conducted within the Health Research Board Mother and Baby Clinical Trial Network Collaborative. The Coombe Women and Infants University Hospital Dublin, Cork University Maternity Hospital, University Maternity Hospital Limerick, The Royal Jubilee Maternity Hospital Belfast, University College Hospital Galway, The National Maternity Hospital Dublin and The Rotunda Maternity Hospital Dublin are the seven largest consultant-led maternity units on the island of Ireland. Combined, they have an annual birth rate of over 44 000, representing over half of the country’s total annual births. Women attending these maternity units who present with suspected preterm pre-eclampsia are eligible for inclusion in this trial. Detailed inclusion and exclusion criteria are described in boxes 1 and 2.

Randomisation

The trial statisticians for the study developed a randomisation sequence for site transition from control to intervention; however, the order of site transitioning is concealed from sites and principal investigators until 12 weeks prior to the sites transition date. An allocation sequence was randomly selected (ie, a cross-over order for the seven clusters) from a set of random sequences constrained so that the sum of the total cluster sizes in
the intervention status was similar to the total sum of the cluster sizes in the control status. Similar was defined to be a difference in the total sums exposed to intervention and control statuses being no different than the expected middle 25th percentile range of differences. To implement this, 10,000 simulations of possible (unique) allocation sequences were performed. From this, the difference in number exposed to intervention and control for each sequence was determined. An allocation sequence was then selected at random from those falling within the middle 25th percentile range of differences.17–19

Control

Eligible women are approached and provided with detailed information about the trial, both verbally and written, by a trained researcher. Eligibility is determined by review of symptoms and signs at the time of presentation to the maternity hospital by the local researcher. Participants are not aware of their maternity hospitals current randomisation prior to their enrolment on the trial. Informed consent is obtained in accordance with ICH—GCP guidelines.21 Once an eligible woman has given written informed consent for inclusion in the study, her maternity hospitals current group allocation is revealed (figure 2). Participants enrolled in the control arm receive usual hospital care as per National guidelines; these are Health Service Executive/Institute of Obstetrics and Gynaecology Irish guidelines for those in the Republic or the NICE guidelines for those in Northern Ireland (figure 3A,B).22,23 Eligible women who are approached but who decline to participate in the trial will continue to receive usual hospital care.

Intervention

Participants enrolled in the intervention arm have their plasma PlGF quantified in addition to routine hospital investigations. The PlGF result is made immediately available to the participants clinical team and documented clearly in the participant’s medical notes. A suggested further management algorithm is provided to the clinician based on both the degree of hypertension present and the PlGF result (figure 4). This algorithm advocates increased frequency of review for those participants identified as having an abnormal PlGF result. The final decision regarding frequency of review remains with the treating clinician. If 4 weeks or more pass and the participant represents with symptoms suggestive of pre-eclampsia, a repeat PlGF quantification may be performed as long as the inclusion/exclusion criteria are still satisfied. In certain sites the option of plasma Biobanking will be available. Participants will be consented separately for this. For those who give consent, a portion of the specimen taken will be used to measure the level of PlGF in the plasma and the remainder of the sample will be stored in University College Cork Biobanking facility.

Box 1 Inclusion criteria

- Pregnant women between 20+0 and 36+6 weeks of gestation (inclusive) with a:
  - Singleton pregnancy.
  - Aged 18 years or over.
  - Able to give informed consent.
  - Presenting with suspected pre-eclampsia: (one or more of the following):
    - Hypertension;
    - Dipstick proteinuria;
    - Headache;
    - Visual disturbances;
    - Epigastric or right upper quadrant pain;
    - Increasing oedema;
    - Suspected fetal growth restriction;
    - If the healthcare provider deems that the woman requires further evaluation for possible pre-eclampsia.

Box 2 Exclusion criteria

- Confirmed pre-eclampsia at point of enrolment: ‘sustained hypertension with systolic blood pressure (BP)≥140 or diastolic BP≥90 mm Hg on at least two occasions (at least 4 hours apart) with significant quantified proteinuria (>300 mg protein on 24 hour collection or urine protein creatinine ratio>30 mg/mmol) or abnormal pre-eclampsia bloods’.
- ≥37 weeks’ gestation.
- Multiple pregnancy.
- Abnormal pre-eclampsia bloods (new-onset reduced number of platelets or deranged liver function/renal function tests, identified during routine care prior to enrolment and not attributable to anything other than pre-eclampsia).
- Decision regarding imminent delivery already made.
- Lethal fetal abnormality present.
- Previous participation in PELICAN trial in a prior pregnancy.
- Participation in a conflicting trial at the same time as PARROT Ireland.
- Plan to use off protocol placental growth factor testing.

PlGF quantification

Maternal plasma PlGF quantification is performed on an EDTA venous blood sample obtained in the standard fashion. Plasma is obtained through centrifugation and the sample is then processed immediately using a CE marked validated point of care platform; the automated Triage Meterpro (ALERE San Diego, California, USA). Each hospital has the necessary equipment in situ and appropriately trained researchers in place, to perform this test as per manufacturer’s guidelines. The PlGF measurement is reported as the absolute value in pg/mL within 30 min of commencing processing of the sample. All samples taken will be analysed without delay by the researcher after venepuncture has occurred and in accordance with manufacturer’s instructions. The Triage PlGF test platform and consumables necessary to perform testing are brought to the cluster just at the point of transition to intervention. It is therefore not available at site for use while the site is in the control arm.
Patient and public involvement

Patients/public were not involved in the development of this trial.

Outcome measure

Primary outcome measure

To evaluate if the intervention is beneficial to both women and their babies and more importantly to ensure it is not harmful to either, the study has two equally important co-primary outcome measures. These are maternal morbidity and neonatal morbidity. For maternal morbidity assessment, an adaption of the full-PIERS score is used (box 3). The definition of hepatic dysfunction is based on alanine aminotransferase rather than International Normalised Ratio (INR), requirement for intensive care unit admission is included as well as the presence of severe hypertension. Severe systolic hypertension is an independent risk factor for stroke in pregnancy and in high-resource settings uncontrolled hypertension is the main cause of death in women with pre-eclampsia. The interval from diagnosis of pre-eclampsia to delivery is not a suitable outcome measure to use, as we are aware that knowledge of PI GF result may alter clinician management and expedite delivery. For neonatal morbidity assessment, babies are dichotomised into having or not having identified neonatal morbidity by means of a composite neonatal score (box 4). In order to avoid subjectivity in the diagnosis of morbidity, the majority of components of the neonatal composite score are objective measures; pH<7.2, positive cultures, admission to NICU. We acknowledge that some subjectivity can arise with staging of disease hence why all stages of each disease will be captured and will comprise the composite outcome; NEC stage 1–3, IVH grade 1–4 and ROP stage 1–5. Neonatal outcomes and morbidity will be captured from local case note review, as documented by the treating neonatologist. In cases where any uncertainty is present, the researcher will discuss the case with the local Principal Investigator (PI) and or the trial clinical fellow and a consensus will be reached.

Secondary outcome measure

Secondary outcomes include each component of the primary outcome reported individually as well as further maternal and neonatal assessments such as mode of delivery and use of antihypertensive agents (boxes 5 and 6). Fetal growth restriction, identified on antenatal ultrasound, has been included as a secondary outcome measure of neonatal morbidity. As PI GF correlates well with placental dysfunction, it may be able to differentiate between those babies with pathological growth restriction rather than constitutional growth restriction and hence improve neonatal outcomes. A separate health economic evaluation is assessing the intervention’s economic impact. This is achieved through the use of participant quality of life (QoL) questionnaires (EuroQol 5D and Short-Form 36), a specially designed study-specific participant costing questionnaire and by assessment of costs to the health service.
of community-based/inpatient/day case care, through chart review at discharge.30–32

Data collection

Trial data captured locally at site by researchers are transmitted securely using an electronic clinical record form (eCRF) to a specific database developed by MedSciNet. Baseline demographic data, QoL questionnaires and the PlGF result are entered live to the eCRF at point of recruitment. The full eCRF is completed after discharge from the maternity hospital postdelivery, and includes neonatal and maternal medical outcome, costing questionnaire and repeat QoL questionnaires. All data entered to the eCRF is pseudo-anonymised with each participant identified by a unique study number. The identifier key is kept separately locally at site in a secure location. The data system is built to the same security and confidentiality standards as those of hospital electronic health records. The data at each participating centre are handled in accordance with local regulatory legislation and ethics committee approval. A detailed description of schedule and timing of data collection is provided in table 1.
Sample size
The sample size was fixed by the number of sites and the study duration. It is anticipated that the total sample size will be in the region of 4000 participants; split across seven clusters and the eight time periods in the design (equivalent to a cluster-period size of about 71). With a sample size of 4000 and using a two-sided type I error rate of 0.025 (to allow for two co-primary outcomes), we determined the power to detect a 7% reduction in maternal morbidity (relative risk reduction of 20%) from 35% to 28% in the intervention, that is, ‘active’ group (based on a reported rate of adverse maternal outcome in the region of 35% in the PELICAN trial). This is assuming an intraclass correlation (ICC) in the region of 0.01; and considering sensitivity to a range of intraclass correlation (ICC) values between 0.005 and 0.05. The second co-primary outcome is adverse neonatal outcomes. Due to scarcity of information on the ICC, the same ICC as for the maternal outcome is assumed. Current rates of adverse events are around 10%. We determine power to detect an absolute change in neonatal adverse outcomes of 6%.

To allow for the longitudinal nature of the trial, where correlations may differ between observations in the same management, the following considerations are made:

**Box 3 Components of the maternal morbidity composite score**
- Confirmed placental abruption.
- Intensive care admission.
- Central nervous system compromise:
  - Generalised tonic clonic seizure due to eclampsia, Glasgow Coma Scale (GCS)<13, cerebral haemorrhage/infarction, cortical blindness, retinal detachment, transient ischaemic attack, reversible ischaemic neurological deficit.
- Cardiorespiratory compromise:
  - Myocardial ischaemia/infarction, SpO₂<90%, >50% FiO₂ for >1 hour, intubation (other than for caesarean section), pulmonary oedema, need for positive inotrope support.
- Haematological compromise:
  - Transfusion of any blood product, platelet count <100×10⁹/L.
- Liver compromise:
  - Hepatic dysfunction (alanine aminotransferase or aspartate aminotransferase>70 IU/L, haematoma, rupture.
- Kidney compromise:
  - Acute renal insufficiency (creatinine>150 μmol/L); haemodialysis.
- Severe hypertension:
  - Systolic blood pressure>160 mm Hg on at least one occasion.

**Box 4 Components of the neonatal morbidity composite score**
- Perinatal death or death before hospital discharge.
- Neonatal intensive care unit admission for >48 hours.
- Birth weight ≤5th customised centile*.
- Apgar score <7 at 5 min.
- Umbilical artery acidosis at birth (cord pH <7.2).
- Admission to neonatal unit.
- Respiratory distress syndrome.
- Intraventricular haemorrhage (IVH).
- Retinopathy of prematurity (ROP).
- Confirmed infection (confirmed on blood or cerebrospinal fluid cultures).
- Necrotising enterocolitis (NEC).

*Customised birth weight at delivery is calculated using the Gestation Related Optimal Weight (GROW) centile.
### Box 5 Secondary outcomes—maternal
- Final diagnosis of hypertensive disorder of pregnancy (chronic hypertension (HTN), gestational HTN or pre-eclampsia).
- Gestation at diagnosis of pre-eclampsia.
- Use of one or more antihypertensive drugs.
- Instrumental delivery (ventouse or forceps).
- Severe HTN (systolic blood pressure (BP) ≥160 mm Hg on at least one occasion).
- Maternal morbidity by fullPIERS model:
  - Confirmed placental abruption;
  - Intensive care admission;
  - Central nervous system compromise;
  - Cardiorespiratory compromise;
  - Haematological compromise;
  - Liver compromise;
  - Kidney compromise.
- Progression to severe pre-eclampsia as defined by the American College of Obstetricians and Gynecologists practice bulletin:
  - Systolic BP ≥160 mm Hg or diastolic BP ≥110 mm Hg on two occasions at least 4 hours apart while the patient is on bed rest (unless antihypertensive therapy is initiated before this time);
  - Thrombocytopenia (platelet count <100×10^9/L);
  - Impaired liver function as indicated by abnormally elevated blood concentrations of liver enzymes (to twice normal concentration), severe persistent right upper quadrant or epigastric pain unresponsive to medication and not accounted for by an alternative diagnoses, or both;
  - Progressive renal insufficiency (serum creatinine concentration >1.1 mg/dL (150 μmol/L) or a doubling of the serum creatinine concentration in the absence of other renal disease);
  - Pulmonary oedema;
  - New-onset cerebral or visual disturbances.
- Elective delivery: induction of labour or caesarean section.
- Caesarean section: emergency and elective.

### Box 6 Secondary outcomes—neonatal
- Fetal growth restriction identified on antenatal ultrasound* (estimated fetal weight and/or abdominal circumference <10th customised centile, abnormality in umbilical artery Doppler velocity or reduced level of amniotic fluid).
- Gestation at delivery.
- Perinatal death or death before hospital discharge.
- Admission to neonatal intensive care unit (NICU).
- NICU admission for ≥ 48 hours.
- Birth weight ≤ 5th customised centile.
- Apgar score <7 at 5 min.
- Umbilical artery acidosis at birth (arterial cord pH<7.2).
- Respiratory distress syndrome.
- Interventricular haemorrhage.
- Retinopathy of prematurity.
- Confirmed infection (confirmed on blood or cerebrospinal fluid cultures).
- Necrotising enterocolitis.

*Antenatal detection of fetal growth restriction is based on formal ultrasound assessment of fetal biometry using the Hadlock formula.

Cluster-period; and those measured in different cluster periods, we incorporate cluster-autocorrelations (CAC). There is little information to support likely values for the CAC, so we are guided by values in the literature and explore sensitivity across a range of values (0.64, 0.80 and 0.96).^{34,35}

The power has been estimated using an online RShiny App.^{36,37} We have not included transition periods in the calculation but given the transition periods are just 1 week in length, this is not expected to significantly affect power. There has been no allowance for varying cluster sizes as this is currently not something which is technically possible in a stepped wedge study. Sample size calculations were performed assuming linear mixed models with categorical effects for time; random cluster and random cluster by period effects.^{38} Under these assumptions, we constructed power curves, which reveal that under most anticipated scenarios the trial will have in the region of 80% power (figures 5 and 6).^{35,39}

### Data analysis
#### Clinical outcome
The primary aim of the study is to evaluate whether there is a difference in the two composite outcomes before and after exposure to the intervention. There will be no double counting of outcomes, individuals not events will be presented for the composite. Mixed effects regression models will be used to allow for the clustering within sites. Calendar time will also be adjusted for since the intervention is sequentially rolled-out both by including fixed categorical time effects and random cluster by categorical time effects.^{40}

The primary estimate of the treatment effects will therefore be cluster and time adjusted. Time adjustment is essential, as it is a stepped wedge trial. Log Poisson regression models with robust variance estimation (to allow for misspecification of binomial errors) will be used so as to allow estimates of relative risks; to estimate risk differences corresponding Binomial models with log links will be fitted. Secondary analysis will adjust for individual and cluster level covariates. In the first instance, comparative estimates of differences between groups will be adjusted for variables used in the randomisation procedure (eg, site, time and hospital size). Furthermore, more fully adjusted analyses will also be performed. These more fully adjusted analyses will adjust for gestational age at recruitment, maternal age, smoking status, maternal body mass index, public versus private obstetric care and maternal comorbidities such as chronic renal disease, systemic lupus erythematosus (SLE), antiphospholipid syndrome (APS) and diabetes. It will also adjust for hospital size (<5000 or >5000 deliveries/annum). Categorised continuous variables (eg, age) will be treated as continuous variables in this adjustment. If covariate adjustment is not practical, unadjusted estimates will be produced and it will be made clear in the output why this occurred (eg, not possible due to low event rate lack of model convergence). Null hypotheses and analyses for secondary outcomes take a similar form to that for the primary outcome, and where outcomes are not binary, analysis will be using the generalised linear mixed model.
Transformations will be performed where data are markedly not normally distributed. For the analysis adjusted for covariates and for the secondary outcomes (unadjusted), multiple imputation methods will be used if the proportion of missing data is >5%, and this multiple imputation will also allow for the clustered and temporal nature of the trial. It is not expected that there will be any missing data in the primary outcome, as it will be assumed that if the outcome is present then it will be recorded and if it is not recorded we will assume it is absent. This is a standard and realistic assumption. Results will be presented as adjusted risk ratios with CIs and risk differences to allow full appreciation of clinical effect. To allow for the two primary outcomes, we will follow good practice and adjust for this multiplicity using a Bonferroni correction and so report 97.5% CIs.

For secondary continuous outcomes mean differences will be reported and 99% CIs. We will report latent ICCs for all outcomes, along with 95% CIs. Prespecified subgroup analysis will be undertaken on the primary outcome based on women presenting <35 weeks’ gestation vs >35 weeks’ gestation; size of unit and final confirmed diagnosis. The stepped wedge trial design will also allow investigation of treatment effect heterogeneity across clusters and time. These exploratory analyses will be reported using 99% CIs.

Analysis will be conducted by intention to treat and sites will be considered exposed to the intervention postrandomised cross-over date.

Health economic outcome

The economic evaluation will be informed by a decision analytical model, which will be designed and constructed for the study to reflect the maternal and fetal pathway and health states. Employing a decision analytical model allows for the extrapolation of existing data and the opportunity

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**Table 1**

| Standard Protocol Items: Recommendations for Interventional Trials flow diagram for schedule of events in PARROT Ireland |
|---|---|---|
| **On presentation with suspected PET Between 20+0 and 36+6 weeks** | **From enrolment to discharge postdelivery** | **Discharge postdelivery** |
| | In-person visit | Chart | In-person visit | Chart | In-person completed |
| Randomisation-institutional level | X | | | | |
| Inclusion/exclusion | | | | | |
| Informed consent | X | | | | |
| Demographics | | X* | | | |
| History, comorbidities | | X* | | | |
| Con medications | | X* | X | | |
| Physical measurements | | X* | | | |
| Clinical readings | | X* | | | |
| PlGF† measurement | X | | | X‡ | |
| Biobank sample§ | X | | | | |
| Fetal assessments | | | X | | |
| Prenatal admissions | | | X | | |
| Maternal PET bloods | | | X | | |
| Newborn data | | | X | | |
| Neonatal outcome | | | X | | |
| Maternal outcome | | | X | | |
| Complications | | | X | | |
| Postnatal admissions | | | X | | |
| Clinical management | | | X | | |
| Final outcomes | | | X | | |
| EQ-5D, SF-36 | X | | | X | |
| Costing questionnaire | | | X | | |
| In-person visits | X | | X‡ | | |

*May be captured in chart review or in consultation with participant at any time following enrolment.
†PlGF testing depends on institutional randomisation allocation.
‡PlGF testing will be repeated if readmission for suspected pre-eclampsia. May be repeated more than once. No more often than four weekly.
§Only at biobanking sites.

EQ-5D, EuroQol 5D; PlGF, placental growth factor; SF-36, Short-Form 36.
to systematically synthesise evidence from various sources. Primary data on maternal health outcomes will be available from the study with the distribution of 5-level EQ-5D version and SF-36 questionnaires which will inform the estimation of quality-adjusted life years (QALYs). Neonatal outcomes will be informed by secondary sources. A systematic literature review will be conducted to identify QOL/utilities (or proxies for same) associated with neonate outcomes, which will be incorporated into the decision analytical model to estimate QALYs. Primary data on resource utilisation will be collected using the costing questionnaire. The costs and effects of the intervention and comparator will be compared with an incremental cost-effectiveness ratio in a cost utility analysis. To address parameter and structural uncertainties, a probabilistic sensitivity analysis will be performed.

**Trial management**

Day-to-day running of the trial will be coordinated by the Trial Management Group (TMG). The TMG consist of the lead site investigator plus the project manager and the clinical fellow. The TMG will act on behalf of the sponsor and will be responsible to the Trial Steering

**Figure 5** Power curve for PARROT Ireland for maternal adverse outcomes. CAC, cluster-autocorrelation; ICC, intracluster correlation.

**Figure 6** Power curve for PARROT Ireland for neonatal adverse outcomes. CAC, cluster-autocorrelation; ICC, intracluster correlation.
Committee (TSC) to ensure that all sponsors’ responsibilities are carried out. The TSC comprises all PIs as well as the TMG, sponsor, Health Research Board (HRB) and representatives from statistics, economics, neonatology, laboratory and a lay person. The role of the TSC is to provide overall supervision of the trial. In particular, the TSC will concentrate on the progress of the trial, adherence to the protocol, participant safety and consideration of new information.

Data monitoring
To provide protection for study participants an independent data monitoring committee (DMC) has been appointed for this trial. The DMC comprises four members who are not involved with any other aspect of the trial. They include an obstetrician, a neonatologist, a statistician and a midwife. The DMC met and ratified their charter and have advised that all serious adverse events such as stillbirth/neonatal death or profound maternal morbidity in the Intervention arm of the study be reported to them immediately. The DMC will receive regular updates on the progress of the trial every quarter from the TMG. The purpose of these updates is for the DMC to: 1) ensure the quality of data collection; 2) ensure that the intervention is being rolled out according to the randomisation plan; 3) monitor balance between arms to monitor for potential selection biases and 4) ensure PIGF testing is not overwhelmingly better or worse than no PIGF testing with respect to maternal morbidity with neonatal morbidity. Once 1500 outcomes are available an interim analysis will be conducted and reviewed by the DMC. The interim analysis will report on the co-primary outcomes, follow the same methods as those of the primary analysis and examine if there is proof beyond reasonable doubt that one particular intervention is definitely indicated or definitely contraindicated in terms of a net difference of a major end point. There will be no formal stopping criteria put in place, but the DMC will be guided by the knowledge that proof beyond reasonable doubt cannot be specified precisely, but a difference of at least three SD in an interim analysis of the primary outcome would be consistent with strong level of evidence. No allowance for this interim analysis has been made in power calculations.

There will be no stopping of the trial for futility as the study will be underpowered to detect small effects.

DISCUSSION
Based on previous experience during the PELICAN study, an analysis of success criteria and barriers to our proposed study was conducted. Potential barriers include the over-estimation of (i) identification of eligible women by the research team, (ii) primary outcome event rate (iii) and retention/attrition, that is, gaining outcomes data on all women included.

A recruitment feasibility audit conducted in Cork University Maternity Hospital (CUMH) over the course of a typical week in July 2016 identified 21 women who would be eligible for inclusion in the PARROT Ireland study. This would equate to almost 1100 women per annum in CUMH, approximately 13% of its annual delivery rate. This is in keeping with the quoted 10% incidence of hypertensive disorders of pregnancy in the population. It is anticipated that over the 22-month duration of the study across the 7 hospitals approximately 10,486 women will meet the study inclusion criteria (13% of the combined annual delivery rate), and of these 4000 will be recruited into this trial (approximately 38% of those eligible). As inclusion in the trial will be optional and require informed consent from participants, not all eligible women in each unit will be included. Projected inclusion rates will be apparent via a dedicated MedSciNet database pre-programmed, available online and contemporaneously updated, allowing prompt action to intervene when not optimal. A conservative requirement of <50% of all eligible women to be recruited in order to reach targets has deliberately been chosen and successful recruitment of the same population in the PELICAN study is reassuring. As with any study we may get a higher or lower incidence of the primary outcome of interest than anticipated. We should get an early indication of this at the interim analysis.

As participation in the trial does not require any extra attendances/input from the participant for the remainder of the pregnancy, it is likely that retention of participants will not be an issue. Similarly, the data outcome to assess for maternal and neonatal morbidity can be readily obtained postdelivery following discharge of the participant from their stored medical records locally at each unit. However, in order to fully examine the health economic outcomes there exists a reliance on the return of completed questionnaires by the participant postdelivery. To minimise attrition rates, the researcher at each site will endeavour to meet with each participant postdelivery prior to their discharge and encourage them to complete the health economic questionnaires. In the PELICAN study, only 1% of the cohort were lost to follow-up. The risk of incomplete data collection of outcomes in studies such as this is more relevant if women deliver in a different unit to that which they are recruited into the trial. However, all seven clusters in our trial are large tertiary referral units and patient transfer during pregnancy is rare. We are therefore confident that the likely rate of loss to follow-up will be similar and in the order of 1%.

There are a number of advantages with the use of stepped wedge design. It allows a phased implementation of the intervention, which is preferable when commencement in all clusters simultaneously would be challenging. As all clusters ultimately receive the intervention, it increases willingness of the clusters to partake in the trial. We acknowledge that seven clusters is a small number of clusters and this is an important limitation of the study. Mostly, this is a limitation because it will mean that the findings have questionable generalisability. But, if these clusters are representative then the findings may still be
generalisable in part. The other limitation that seven clusters brings about is questionable internal reliability. However, because all of the clusters receive both the intervention and control condition, the clusters serve as their own controls. This lessens the impact of chance imbalance and it increases the power of the study (particularly so when the ICC is large, as is the case here). The study does only have in the region of 80% power and should parameters such as the ICC be very different to that which we have assumed, then it is correct that the study might be underpowered. To ensure that this is properly accounted for at the analysis stage, we will report appropriate CIs around all point estimates, so the impact of any impression is properly reported.

Another potential limitation worth noting is the slightly different management algorithm for one cluster, Belfast, in the control arm. The Belfast control arm algorithm is taken directly from the NICE Hypertension in Pregnancy guidelines. All other clusters are using an algorithm taken from the HSE guidelines for Hypertension in Pregnancy. The two are essentially the same except the HSE algorithm also includes a recommendation for a fetal ultrasound in cases where the participant is <34 weeks’ gestation. It is not anticipated that the difference in these algorithms should have any bearing on the overall trial results. We will conduct a sensitivity analysis with the Belfast site removed and see if the result remains consistent.

Ideally, PlGF testing should be performed for all participants enrolled in the study, with blinding of the result for those in the control arm. This would allow for test performance statistics to be performed. Unfortunately, testing of control participants will not be conducted in our trial, which is a notable limitation of the study.

The primary aim of the PARROT Ireland trial is to establish the effectiveness of revealed plasma PlGF measurement in reducing maternal morbidity (with assessment of neonatal safety in parallel) in women presenting with suspected pre-eclampsia prior to 37 weeks’ gestation. Should the trial show a reduction in maternal morbidity without an increase in neonatal morbidity, or indeed a reduction in neonatal morbidity with no change in maternal morbidity, it would provide a strong argument for its incorporation into routine obstetric practice. The long-term aim of the trial is to demonstrate if PlGF measurement enables appropriate antenatal stratification of women presenting with suspected pre-eclampsia.

Avoiding unnecessary hospital admission would be both clinically and economically beneficial. In contrast, those at increased risk of imminent adverse events, identified by an abnormal PlGF result, would have hospital resources redirected to them. We anticipate that this trial will provide a definitive result on the benefits of PlGF testing, which will act to influence international clinical practice.

A separate randomised controlled trial (RCT), also entitled ‘PARROT’, has completed recruitment in the UK since the end of 2017. Although recruiting a similar population of women and using the same PlGF platform, the primary outcome measure for the two RCTs is different, with the UK PARROT trial focusing on time from enrolment to diagnosis. Both studies are using the same electronic clinical record forms developed by MedSciNet and thus will have a large cross-over of data. The advantage of having these two similar RCTs conducted almost simultaneously is that robust information on the impact of incorporation of PIGF into clinical care will be generated. In addition, the potential exists for a collaborative project such as an individual participant data meta-analyses in the future.

DECLARATIONS

Ethics approval and consent to participate

The trial is being conducted in accordance with ethical principles that have their origin in the Declaration of Helsinki and are consistent with Good Clinical Practice (GCP) and applicable regulatory requirements. The local ethics committee at each participating site has reviewed the trial protocol, including the patient information and informed consent form, and full ethical approval granted. Each eligible woman identified is required to give written informed consent prior to her inclusion in the trial. A GCP trained researcher at the local site obtains this consent.

Clinical Research Ethics Committee Cork: ECM 3 (h) 08/11/16.
University College Hospital Galway EC: Ref 50/12.
Coombe Womens & Infants University Hospital EC: Study No 20–2016.
University Hospital Limerick EC: Ref: 68/16.
Health Research Authority (Belfast): 16/WM/0484.
Rotunda Hospital EC: REC-2016–020.

Dissemination

The success of the trial will be dependent entirely on the collaboration of clinicians in the participating hospitals and those who hold key responsibility for the trial. Hence, the credit for the study will be assigned to the key collaborator(s) from a participating site as it is crucial that those taking credit for the work have actually carried it out. The results of the trial will be reported first to trial collaborators. The results from the PARROT Ireland trial will be published in an established peer-reviewed journal. At least one publication of the main results will be made. Links to the publication will be provided in all applicable trial registers. Dissemination of results to participants will take place via the media, trial website and relevant participant organisations. Collaborating investigators will play a vital role in disseminating the results to colleagues and participants.

Author affiliations

1The Irish Centre for Fetal and Neonatal Translational Research (INFANT), UCC, Cork, Ireland
2University College Cork, Department of Obstetrics and Gynaecology, Cork, Ireland
3Public Health, University of Birmingham, Birmingham, UK
4Rotunda Hospital, Royal College of Surgeons in Ireland, Dublin, Ireland
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Contributors All authors contributed to the overall study design and specific methodologies. LK conceived and designed the study with DD. LK and DHR

Patient consent for publication Not required.

Ethics approval Ethical approval has been granted from each of the seven maternity hospitals involved in the trial. The results of the trial will be presented both nationally and internationally at conference and published in an international peer-reviewed journal.

Provenance and peer review Not commissioned; externally peer reviewed.

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