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University College Cork, Ireland Coláiste na hOllscoile Corcaigh

1	A prospective study of placental growth factor in twin
2	pregnancy and development of a dichorionic twin pregnancy
3	specific reference range
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21	Running Title: Placental growth factor in twin pregnancy
22	

1 Abstract

Objective: The aim of this study was twofold; to develop a dichorionic twin pregnancy 2 specific reference range for placental growth factor, and to compare gestational 3 4 specific placental growth factor levels in twin pregnancies later complicated by preeclampsia, hypertensive disorder of pregnancy or fetal growth restriction to controls 5 6 Design: Prospective observational study 7 8 9 Setting: Single large tertiary maternity unit in Ireland 10 *Population or Sample:* Women with a twin pregnancy 11 12 *Methods:* Consenting pregnant women, across a variety of gestations, had a single 13 blood sample taken at one time point only during their pregnancy. The plasma was 14 initially biobanked and PIGF was measured later in batches using the point of care 15 Triage® PIGF test 16 17 Main Outcome Measures: Development of preeclampsia, hypertensive disorder of 18 pregnancy or fetal growth restriction 19 20 *Results:* PIGF levels in uncomplicated dichorionic twin pregnancies were significantly 21 lower in the women who later developed preeclampsia than in the controls at all 22 gestational intervals. In those that later developed any hypertensive disorder of 23 pregnancy median PIGF was lower only in those recruited before 24 weeks' gestation 24

while in infants with a customised birthweight below the 3rd centile, PIGF was lower
only in those sampled after 24 weeks' gestation.

3

Conclusions: PIGF levels in twin pregnancy differ significantly between those women
with a pregnancy that will later be complicated by preeclampsia and those that will not.
This difference is present many weeks before clinical signs or symptoms of disease
are present. Using cross sectional values from uncomplicated twin pregnancies, we
have developed a dichorionic twin pregnancy specific reference range for PIGF.

9

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14

Keywords: 15

- Hypertensive Disorders of Pregnancy
- Placental Growth Factor
- 18 Preeclampsia
- 19 Reference Range
- Twin Pregnancy

21

Tweetable Abstract: PIGF levels in twin pregnancy differ significantly between
 women that will later develop preeclampsia and those that will not.

1 Introduction

2 Preeclampsia is a common complication of pregnancy characterised by new onset hypertension and either proteinuria or other maternal organ dysfunction after 20 3 weeks' gestation (1). Along with other hypertensive disorders of pregnancy (HDP), it 4 is a major contributor to maternal and neonatal morbidity and mortality (2, 3). 5 6 Potentially serious maternal morbidity may arise in the form of seizures, cerebral 7 haemorrhage, renal failure, liver rupture and disseminated intravascular coagulation 8 (4). The only definitive treatment for preeclampsia is removal of the placenta, often 9 resulting in iatrogenic pre-term delivery and subsequent fetal morbidity (5). Women with a twin pregnancy are at a two to three fold increased risk of developing 10 preeclampsia, possibly due to a combination of larger placental mass and use of 11 assisted reproductive therapy (ART), especially use of non-autologous gametes (6-8). 12 Rates of twin pregnancy have risen over the last number of decades globally (9-12). 13

14 Although the exact aetiology of preeclampsia is not fully understood, a growing body of evidence suggests that an imbalance of angiogenic factors of placental origin play 15 a crucial role in its development (13-17). Placental growth factor (PIGF) is an 16 angiogenic protein and a member of the vascular endothelial growth factor family (18). 17 Studies in singleton pregnancies have shown lowered levels of PIGF and increased 18 levels of its soluble receptor sFIt-1 in maternal plasma, weeks prior to the clinical onset 19 of preeclampsia (19, 20). The UK National Institute for Clinical Excellence (NICE) 20 advocates PIGF testing, combined with routine clinical care, to help rule out pre-term 21 preeclampsia in singleton pregnancies (21). A number of international randomised 22 control trials (RCTs) are currently on-going, investigating the clinical impact of the 23 integration of PIGF into clinical care pathways (22). The international INSPIRE trial 24 evaluated the use of sFIt-1/PIGF ratio in women presenting with suspected 25

preeclampsia and showed that use of the ratio, in conjunction with standard clinical
practice, significantly improved clinical precision without changing admission rates
(23). The UK PARROT study, demonstrated a reduction in time taken to diagnosis
preeclampsia and reduced maternal morbidity when PIGF is integrated into clinical
care algorithms (24).

6 Few studies to date have evaluated the levels of circulating angiogenic factors during 7 twin pregnancy. In those that have been described, huge variations exist in; the primary outcome (i.e preeclampsia, fetal growth restriction or other adverse clinical 8 9 outcome); the definition/classification of the primary outcome; the gestational age at time of sampling; and the immunoassay used for guantification. (25-33). The aim of 10 this study was twofold; to develop a dichorionic twin pregnancy specific reference 11 range for PIGF and secondly to compare gestational specific PIGF levels in twin 12 pregnancies complicated by preeclampsia, any hypertensive disorder of pregnancy 13 (HDP) or fetal growth restriction to controls. 14

15

16 Materials & Methods

17 Setting and Design

This study was conducted in a single maternity hospital in Ireland with over 8000 deliveries per annum. The study was a prospective cross-sectional cohort study of PIGF in twin pregnancy. From the start of July 2015 to the end of December 2017, women attending the hospital's dedicated twin pregnancy clinic were approached to participate in the study. Any woman with an uncomplicated twin pregnancy from 12+0-36+6 weeks' gestation inclusive, without signs/symptoms or a diagnosis of preeclampsia was eligible for inclusion. Those with complications such as a known

congenital anomaly in either baby, severe early onset growth restriction or twin-to-twin 1 transfusion syndrome (TTTS) were excluded from recruitment. Following informed 2 patient consent, a 3ml ethylenediaminetetraacetic acid (EDTA) blood sample was 3 taken, centrifuged, divided into aliquots and the plasma biobanked at -80C within 3 4 hours of sampling. All sampling, processing and biobanking was carried out within the 5 same building according to previously published Standard Operating Procedures 6 7 (SOPs) (34). Women had venepuncture performed at one random gestational time point only. Clinically relevant outcome data such as the diagnosis of any HDP (chronic 8 9 hypertension, gestational hypertension, preeclampsia or superimposed preeclampsia) and infant birthweights were taken from medical notes following delivery. Anonymised 10 clinical and demographic data pertaining to the participant and their offspring were 11 recorded in the study database. For our study, the NICE definitions of hypertensive 12 disorders of pregnancy were utilised (35). Fetal growth restriction was calculated based 13 on actual birthweight, gestation at birth, fetal gender and maternal ethnicity, parity and 14 BMI using the Gestation Related Optimal Weight (GROW) centile calculator (36). 15 Patients were not involved in the development of this research study and a core 16 outcome set was not used. 17

18 Placental Growth Factor Immunoassay

Biobanked plasma samples were analysed in batches for circulating levels of PIGF using a point of care immunoassay; the Triage® PIGF test (Quidel Inc., San Diego). This test is not routinely available in the hospital for clinical use. It was purchased by our research centre for the purpose of this study. The test manufacturers had no part in the study design, conduct, analysis or manuscript development. The immunoassay was performed as per manufacturer's instructions, in a single freeze thaw cycle to minimise protein denaturation. The results are displayed on the meter screen in

approximately 15 minutes and have a measurable range from 12-3000 pg/ml. The
Triage® has a reported measurable range from 12-3000 pg/ml. The manufacturers
report total precision on plasma controls at concentrations of 85.2 and 1300 pg/mL as
12.8% and 13.2% respectively. For the purposes of this study, any result obtained <12
pg/ml was allocated the value of 10 pg/ml.

6 <u>Statistics</u>

7 SPSS Version 23 and Stata 15 were used to analyse the data.

8 Part 1: Descriptive statistics were employed to examine the baseline maternal demographics, clinical outcomes and the PIGF distribution in the cohort. When 9 developing a reference range for PIGF, all cases where a stillbirth was diagnosed in 10 11 either of the twins as well as cases where any form of HDP later developed were removed. Secondly in order to facilitate development of a reference range for PIGF in 12 an uncomplicated dichorionic twin pregnancy all monochorionic twin pregnancies were 13 removed. Lastly, cases that developed fetal growth restriction resulting in both twins 14 having a customised birthweight of less than the 3rd centile were then removed. The 15 remaining women were divided according to gestational age at recruitment and PIGF 16 ranges calculated for each gestational week were calculated. A heterogeneous 17 regression of log(PIGF) on gestational age of was used to develop a reference range 18 19 for PIGF in uncomplicated dichorionic twin pregnancies.. The mean of log(PIGF) was modelled using fractional polynomials and the variance was modelled as proportional 20 to some power of gestational age (37). The residuals were assumed to follow a Normal 21 22 distribution at each gestational age. Centiles of log(PIGF) were exponentiated to yield centiles 23

Part 2: To examine the effect of hypertensive disorders and placental dysfunction on PIGF, the entire cohort including abnormal cases, was divided into 2 groups based on the woman's gestational age at time of her enrolment to the study and hence sampling of maternal plasma PIGF; <24 weeks' gestation and \geq 24 weeks' gestation. This gestational cut-off was employed as pregnancy related hypertensive complications are unusual prior to this timepoint and also it equated well with the median of the cohort.

7

8 <u>Results</u>

In total, 275 women with a twin pregnancy were recruited. There were no withdrawals 9 or losses to follow up. Three women (1% of the cohort) had a stillbirth occur in one of 10 11 the twins while in 4.7% (n=12) of women, an anomaly of one or both twins was diagnosed. Given with twin pregnancy there is differing placental volumes present 12 dependent on chorionicity, circulating levels of PIGF may also vary in line with 13 chorionicity. We found that PIGF was lower in monochorionic twin pregnancy (data not 14 shown) but given the high incidence of complications as well as the small numbers 15 present (n=40) in this subgroup, further analysis was not possible. We limited our 16 analysis to dichorionic cases only for development of the reference range. 17

18

19 <u>Part 1:</u>

20 Reference Range Demographics

Removal of those with an abnormal pregnancy outcome (preeclampsia or HDP in the mother, stillbirth of either twin or where both twins had a customised birthweight of $<3^{rd}$ centile at delivery), or a monochorionic pregnancy left 173 women with an uneventful

dichorionic twin pregnancy for inclusion in the reference range analysis (Supplemental 1 Material Figure 1). Median maternal age was 34 years, booking BMI was <30Kg/m² 2 for the majority (81.5%; n=141) and most were Caucasian (93.6%; n=162). Over half 3 of the group were multiparous (56.1%; n=97), just over a third (35.1.1%; n=60) had 4 conceived the twin pregnancy with use of ART. All women with pre-existing renal 5 disease or essential hypertension developed superimposed preeclampsia in their 6 7 pregnancies and hence were not included in the reference range cohort (Supplemental Material Table 1). Comparison of participant characteristics between each gestational 8 9 group showed no significant difference in enrolment characteristics (Supplemental Material Table 2). 10

11 Reference Range Development

The 3rd-97th centiles of PIGF as a function of gestational age concentrations were 12 calculated (Table 1). With progressing gestational age the median PIGF was seen to 13 rise, simultaneously to the development and maturation of the placentae, peaking at 14 25 weeks gestation, and then steadily decreased towards term. The lowest acceptable 15 PIGF value for each gestational week is presented (Figure 1). Removal of cases where 16 women developed any form of HDP or where both twins had a customised birthweight 17 <3rd centile did not alter the reference range significantly. These data provide a valid 18 19 reference range for PIGF in a normal dichorionic twin pregnancy (Figure 2).

20

21 <u>Part 2:</u>

22 Comparison of Gestational PIGF

The second aim of this study was to compare gestational PIGF in twin pregnancies complicated by preeclampsia, HDP or customised birthweight of both twins $<3^{rd}$ centile, to controls. To this end, the entire cohort (n=275) was divided into 2 groups

based on the woman's gestational age at time of her enrolment to the study and hence gestational age at time of sampling of maternal plasma PIGF; <24 weeks' gestation and \geq 24 weeks' gestation. Just under half the cohort (43.6%; n=120) were recruited at <24 weeks' gestation with the remainder (56.4%, n=155) recruited at \geq 24 gestational weeks. The groups were then stratified by presence of preeclampsia, HDP or customised birthweight <3rd centile for both infants.

7

8 Demographics of Entire Cohort

9 The maternal age of the study group ranged from 20 to 50 years, with 134 women (48.7%) aged >35 years at booking. The majority of the cohort had a Body Mass Index 10 (BMI) of <30Kg/m² at booking (78.9%; n=217) and were of Caucasian ethnicity 11 (93.8%; n=258). Just under half the cohort were nulliparous (46.9%; n=129). The 12 majority of the group were dichorionic twin pregnancies (81.5%; n=224) and 13 approximately two thirds (65.5%; n=180) of the population studied had conceived the 14 twin pregnancy spontaneously. Where assisted reproductive therapy (ART) was 15 utilised, almost a fifth (17.1%; n=47) had conceived through the assistance of In Vitro 16 Fertilisation (IVF) and a large proportion of these using a donor oocyte (12%; n=33). 17 There was a small number of women with pre-existing renal disease or hypertension 18 (1.8%; n=5). The two gestational groups were well matched, with no differences seen 19 20 in BMI <30, ethnicity, parity or chorionicity. However, there were significantly more women with ART assisted pregnancies (40.2%; n=48 v 27.8%; n=42, p=0.04), oocyte 21 donation (18.5%; n=22 v 7.3%; n=11, p=0.009) and those with a maternal age >35 22 23 years (57.5%; n=69 v 41.9%; n=65, p=0.01) sampled in the <24 weeks' gestational group compared to the \geq 24 weeks group (Supplemental Material Table 3). 24

1 Clinical Outcomes

Overall, the incidence of a subsequent diagnosis of HDP was 15.3% (n=42) and of 2 these 11.3% (n=31) developed preeclampsia (Supplemental Material Table 4). Of the 3 532 infants with maternal BMI information available, 11.8% (n=65) had a customised 4 birthweight <3rd centile with both twins <3rd customised birthweight in twelve cases. 5 Gestation at delivery ranged from 23 to 38 weeks' gestation, with two thirds of the 6 7 cohort delivered via Caesarean section (66.5%, n=183). Pre-term delivery at <35 weeks occurred in almost a fifth of the cohort (17.8%, n=49) and in over half of cases 8 9 was iatrogenic (59.2%, n=20). Preterm delivery at <32 weeks was less common (6.9%, n=19), and again half of cases were iatrogenic (47.4%, n=9). There were no significant 10 differences between the two gestational groups in terms of incidence of HDP or 11 preeclampsia, nor were there any differences in pre-term delivery or mode of delivery. 12 13

14 Comparison of PIGF

The median PIGF was 230.5 pg/mL when sampling occurred at <24 weeks and 276 15 pg/mL when sampling was ≥24 weeks. The cohort was then stratified by subsequent 16 diagnosis of preeclampsia, HDP or customised birthweight <3rd centile in both twins. 17 PIGF levels were 0.6 times higher in the controls than in the women who later 18 developed preeclampsia at <24 weeks gestation (247 pg/ml vs 153 pg/ml) and 2.0 19 times higher at >24 weeks gestation (304 pg/ml vs 99.8 pg/ml.)(Table 2). In those that 20 subsequently developed any form of HDP, PIGF was 0.7 times higher in the >24 weeks 21 group (250 pg/ml vs 150 pg/ml) (Table 3). In those that subsequently had either twin 22 born at a birthweight <3rd customised centile, PIGF was 0.8 times higher in the group 23 recruited <24 weeks (170 pg/ml vs 304pg/ml)(Table 4). 24

1 Discussion:

2 Main Findings

3 This study demonstrates that maternal plasma PIGF in twin pregnancy follows the same gestational pattern as described in singletons (38, 39); a steady rise 4 5 corresponding with development of the placenta, peaking slightly earlier at approximately 25 weeks' gestation, and then declining thereafter. It also shows that 6 7 maternal plasma PIGF is significantly lower in twin pregnancies that will later develop 8 preeclampsia but not other HDP, independent of gestational age at time of sampling of PIGF, compared to controls. PIGF was also noted to be lower in those babies with 9 a customised birthweight <3rd centile when maternal sampling occurred after 24 10 11 weeks gestation.

12 Interpretation

13 To our knowledge, this is the largest prospective study of PIGF in twin pregnancy from a single site. This allows us to describe the twin pregnancy specific distribution of 14 gestational PIGF, as well as develop a dichorionic specific reference range for PIGF 15 16 in twin pregnancy, which has not been previously described. This is also the only study to date examining PIGF in twin pregnancies specifically using the Triage® PIGF test. 17 The Triage® PIGF test is currently the only point of care test on the market for 18 measuring PIGF, is CE marked and has been endorsed by NICE for use in further 19 research (21). 20

Previous studies of angiogenic factors in twin pregnancy have had limited numbers of participants, varied gestations at quantification, varied outcome measures and often involve pooled results from a number of sites or countries across a variety of time periods (40-42). Often these studies require shipment of specimens to laboratories in

other countries, which may affect the quality of samples. In contrast, all of the
laboratory analysis in our study was performed on site, by a single researcher, in a
single freeze thaw cycle, to minimise the chance of protein denaturation.

A Spanish study in 2011 examined first trimester levels of circulating angiogenic factors in 61 women with a twin pregnancy (40). Using a R&D systems immunoassay, they reported higher serum concentrations of both PIGF and sFlt-1 in twins compared to matched singletons. They also reported maternal serum sFlt-1 levels were higher in twin pregnancies conceived through ART compared to spontaneous twin conceptions, supporting the well-accepted concept that ART pregnancies are at increased risk of preeclampsia development.

11 A study from Boston in 2012 (41) described 79 women with a twin pregnancy 12 presenting with suspected preeclampsia in the third trimester. Serum PIGF and sFIt-1 from the women was quantified using the Roche Elecsys immunoassay Ratio test. 13 14 The outcome measure utilised was the diagnosis of an adverse clinical event in the subsequent fortnight, of which 52 women met the criteria. The authors reported 15 median PIGF was significantly reduced, while median sFIt-1 was elevated in those that 16 did develop an adverse event indicating that these angiogenic factors have potential 17 utility as prognostic indicators in twin pregnancies with suspected preeclampsia. 18

A German group in 2014 published on a small cohort of 49 women with a twin pregnancy, 18 of which developed preeclampsia. Maternal serum PIGF and sFIt-1 was quantified again using the Roche Elecsys immunoassay Ratio test. The researchers reported PIGF levels were decreased and sFIt-1 levels increased in the preeclampsia cases at time of presentation with preeclampsia symptoms compared to the twin

controls, indicating the potential for integration of angiogenic factors into clinical care
 pathways for investigation of suspected preeclampsia in twin pregnancy (42).

Clearly, potential exists for use of PIGF and sFIt-1 as biomarkers for prediction of 3 preeclampsia in twin pregnancies. However, before these biomarkers are introduced 4 into clinical use for twins, it is important that relevant cut-offs are developed and 5 6 validated specifically for this group and specific to each PIGF platform. Differences in 7 PIGF results may arise between commercially available platforms owing to measurement of different PIGF isoforms by each assay gene (43). The Triage 8 9 immunoassay predominantly measures PIGF Isoform-1, the R&D System assay detects PIGF-2 and PIGF-3 isoforms in addition to PIGF-1 while the Roche is a ratio 10 of PIGF to sFLT-1 (44, 45). 11

12 Several large prospective observational studies have published on clinically relevant cut-offs for use in singletons. The PROGNOSIS study, using the Roche Elecsys 13 14 immunoassay Ratio test in 550 women with suspected pre-eclampsia, reported a sFIt-1:PIGF ratio of ≤38 as having a negative predictive value for preeclampsia in 15 singletons in the subsequent 7 days of 99.3% (46). The PELICAN study, using the 16 Triage® PIGF test in 625 women with suspected pre-eclampsia, reported a PIGF of 17 >100 pg/ml as having a 98% negative predictive value for preeclampsia in the 18 subsequent 14 days in singletons presenting at < 35 weeks' gestation (47). 19

A 2018 Dutch study compared PIGF and sFlt-1 levels in normotensive and preeclamptic singleton and twin pregnancies using the Roche Elecsys immunoassay Ratio test (48). Numbers were small, with only 22 twin pregnancies included. Again, differences in serum sFlt-1 and PIGF levels were noted in the normotensive twins compared to the matched singletons and in the pre-eclamptic twin cases compared to

1 the twin controls. Importantly, they demonstrated that the previously defined sFIt-1/PIGF ratio cut-off of ≤38 for predicting short-term absence of preeclampsia in 2 singleton pregnancies is not applicable to twin pregnancies. Importantly this 3 demonstrates that established reference ranges for PIGF/sFIt-1 in singletons are not 4 transferrable to twin or higher order multiple pregnancies. This highlights the need for 5 quality prospective observational studies of women with twin pregnancy presenting 6 7 with suspected preeclampsia, in order to develop and validate clinically useful cut-offs for PIGF/sFlt-1 in twins. 8

9 It appears there may be a role for PIGF in identification of pathologically related growth 10 restriction in twin pregnancy, given the lower PIGF levels we observed in the maternal 11 plasma of those women that later went on to deliver babies <3rd CBW centile. Studies 12 in singleton pregnancies have demonstrated the utility of PIGF in discriminating 13 pathological growth restriction from constitutional smallness and in the prediction of 14 adverse perinatal outcomes (49, 50).

15 Strengths and Limitations.

We recognise there are limitations to our study specifically the use of a customised 16 birthweight centile not specific to twin pregnancy and the exclusion of cases where 17 only both twins were <3rd customised centile. This choice was pragmatic given our 18 numbers however we recognise that reduced placental volume in either twin may 19 affect the circulating maternal PIGF levels. Normal twin growth patterns are the subject 20 of much debate with differing opinion as to which is the most appropriate growth curve 21 22 to use in clinical practice (51, 52). Concerns exist that twin specific growth charts, adjusted to reflect the smallness of twins compared to singletons, may not identify 23 growth restricted twins with underlying placental pathology, thereby resulting in 24

increased perinatal morbidity (53). There is no consensus as to whether fetal growth
charts should be customised by factors such as ethnicity, height, weight and parity or
not and there is also no agreement regarding which is the most appropriate growth
calculator to use (54-59).

A second limitation of the study was single sampling of participants. Serial sampling 5 6 of maternal PIGF may have provided a much more robust, informative account of PIGF 7 distribution. However repeated phlebotomy, solely for the purposes of a research study, may have deterred many women from taken part and given that participation 8 9 was truly altruistic, a single timepoint only approach was adopted. Our population is largely homogenous; white Caucasian and non-obese which potentially limits 10 extrapolation to minority ethnic groups however this is representative of the twin 11 pregnancy population that attends our unit. Although a large number of women with a 12 twin pregnancy were enrolled, we do not have sufficient power at present to develop 13 a monochorionic twin pregnancy specific reference range, although it would be 14 possible to expand on the study and add to our numbers in the future to achieve this. 15 An additional limitation of our study is the use of only one automated commercial 16 platform for quantification of PIGF alone and not s-FLT1, rather than on multiple 17 commercially available platforms such as the DELFIA Xpress PIGF 1-2-3 test, Brahms 18 Kryptor and the Roche Elecsys ratio test, as advocated by NICE (21). Comparative 19 studies performed in singleton pregnancies have shown similar performance of all 20 three platforms in ability to rule out preeclampsia (60). As sufficient plasma remains 21 biobanked in our site, this is an area for potential future research subject to funding 22 and ethical approval. 23

24 Conclusion

We have shown that PIGF levels in twin pregnancy differ between those pregnancies that later will be complicated by preeclampsia and those that will not. This difference is present many weeks before clinical signs or symptoms of disease are present. We provide a valid overall reference range for PIGF in a normal twin pregnancy and specifically in a normal dichorionic twin pregnancy. With further research, PIGF has potential as an adjunct to clinical care as a predictor of evolving preeclampsia and/or adverse clinical outcomes in twin pregnancy.

8

9 Financial Disclosure

10 All authors report no financial disclosures

11

12 **Conflict of Interest**

13 All authors report no conflicts of interest

14

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18

19 **Contribution to Authorship**

All authors contributed to the overall study design and specific methodologies. KOD and LK conceived and designed the study with DHR. DHR conducted the data collection with assistance from CN and EOM. DHR conducted the analysis with assistance from SM and AF. DHR drafted the manuscript with assistance from SM and 1 KOD. All authors have critically read, contributed with inputs and revisions and 2 approved the final manuscript.

3

4 **Details of Ethics Approval**

5 Ethical approval for the study was granted from the Cork Research Ethics committee
6 (ECM 3 (PPP) 19/05/15).

7

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13

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Figure Legend

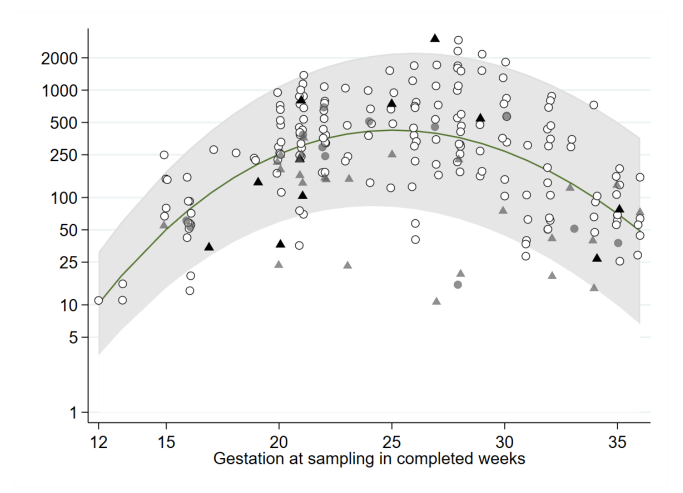


Figure 1: Scatter plot of gestational PIGF. Shaded area represents the reference range from the 5th to 95th percentiles (n=222). ○ uncomplicated dichorionic twin pregnancy cohort, ▲ HDP and PET present, ▲ HDP present, ● both neonates <3rd CBW

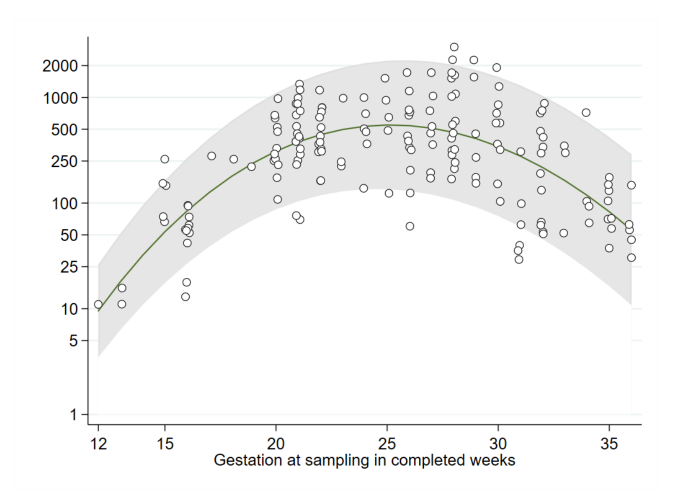


Figure 2: Scatter plot of gestational PIGF for the uncomplicated dichorionic twin pregnancy cohort. Shaded area represents the reference range from the 5th to 95th percentiles (n=173).

Gestational age at sampling (weeks)	Percentile of PIGF (pg/ml)								
	3rd	5th	10th	25th	50 th	75th	90th	95th	97th
12	3.0	3.5	4.3	6.3	9.5	14.5	21.0	26.2	30.3
13	5.5	6.3	8.0	11.8	18.1	27.8	40.9	51.5	59.9
14	9.3	10.9	13.8	20.6	32.1	50.0	74.6	94.7	110.7
15	15.0	17.6	22.5	33.9	53.6	84.7	127.9	163.7	192.1
16	22.7	26.8	34.5	52.8	84.5	135.4	206.9	266.6	314.4
17	32.8	38.8	50.3	77.8	126.2	204.8	316.5	410.7	486.5
18	44.8	53.3	69.7	108.9	178.9	293.8	459.3	600.0	713.7
19	58.5	69.8	91.8	145.1	241.1	400.8	633.3	832.7	994.7
20	72.7	87.1	115.3	184.1	309.6	520.7	831.3	1100.0	1319.4
21	86.2	103.8	138.2	223.0	379.3	645.2	1040.7	1385.5	1668.4
22	97.9	118.3	158.5	258.2	444.0	763.7	1244.1	1666.1	2014.0
23	106.4	129.1	173.9	286.0	497.2	864.4	1422.0	1915.3	2324.1
24	110.8	134.9	182.8	303.5	533.3	936.8	1555.7	2107.4	2566.6
25	110.7	135.3	184.3	308.9	548.2	973.0	1630.8	2221.4	2715.2
26	106.2	130.2	178.3	301.6	540.7	969.5	1639.6	2245.4	2754.2
27	97.8	120.4	165.8	282.9	512.1	927.3	1582.1	2178.3	2681.0
28	86.7	107.0	148.1	255.0	466.1	852.1	1466.5	2029.5	2506.3
29	73.9	91.5	127.3	221.0	407.9	752.7	1306.5	1817.3	2251.7
30	60.6	75.4	105.3	184.4	343.4	639.6	1119.5	1565.0	1945.3
31	47.9	59.7	83.9	148.1	278.3	523.1	923.1	1296.8	1617.1
32	36.5	45.6	64.4	114.6	217.3	412.0	732.9	1034.6	1294.1
33	26.8	33.6	47.6	85.4	163.4	312.6	560.6	795.0	997.5
34	19.0	23.9	34.0	61.4	118.5	228.7	413.2	588.7	740.9
35	12.9	16.3	23.4	42.6	82.9	161.3	293.6	420.3	530.5
36	8.5	10.8	15.5	28.5	55.9	109.7	201.3	289.4	366.3

 Table 1: Normal Reference Range percentiles of PIGF by gestational age interval quantified using the Triage® PIGF test (n=173)

Centiles were calculated based on a Normal distribution for log(PIGFR) where $\mu = exp (-19.8165 + 6.9434 \times \sqrt{G} - 0.01375 \times G^2)$ $\sigma = exp (-1.5851 + 0.4422 \times log(G))$ and G = gestational age (weeks)

Table 2: PIGF by gestational group at enrolment in twin pregnancies complicated by ^A Preeclampsia (de novo or superimposed) compared to those that were not, quantified using the Triage® PIGF test (n=275)

Gestation at recruitment (weeks)	Median (IQR) PIGF pg/mL (n=275)	Median (IQR) PIGF PE ^A present pg/mL (n=31)	Median (IQR) PIGF PE not present pg/mL (n=244)
<24	230.5 (79.4-437.8)	153 (54-224)	247 (81-489)
≥24	276 (71.6-577)	99.8 (24-273)	304 (73-652)

Table 3: PIGF by gestational group at enrolment in twin pregnancies complicated by ^B Hypertensive Disorder of Pregnancy compared to those that were not, quantified using the Triage® PIGF test (n=275)

Gestation at recruitment (weeks)	Median (IQR) PIGF pg/mL (n=275)	Median (IQR) PIGF HDP ^B present pg/mL (n=42)	Median (IQR) PIGF HDP not present pg/mL (n=233)
<24	230.5 (79.4-437.8)	150 (45-229)	250 (84-490)
≥24	276 (71.6-577)	123 (32-425)	304 (73-598)

Table 4: PIGF by gestational group at enrolment in offspring of twin pregnancies complicated by ^C birthweight <3rd customised centile compared to those that were not, quantified using the Triage® PIGF test (n=532)

Gestation at recruitment (weeks)	Median (IQR) PIGF pg/mL (n=532)*	Median (IQR) PIGF pg/mL CBW <3rd ^c (n=109)	Median (IQR) PIGF pg/mL CBW not <3rd ^c (n=423) 231 (105-413)	
<24	230.5 (79.4-437.8)	234 (54.2-460.5)		
≥24	276 (71.6-577)	170 (42.7-462)	304 (73-652)	

*BMI not available for 9 women so CWB not available for 18 offspring