PIGF-based testing to help diagnose suspected pre-eclampsia (Triage PIGF test, Elecsys immunoassay sFlt-1/PIGF ratio, DELFIA Xpress PIGF 1-2-3 test, and BRAHMS sFlt-1 Kryptor/BRAHMS PIGF plus Kryptor PE ratio)

Diagnostics guidance
Published: 11 May 2016
nice.org.uk/guidance/dg23
Your responsibility

This guidance represents the view of NICE, arrived at after careful consideration of the evidence available. When exercising their judgement, healthcare professionals are expected to take this guidance fully into account. However, the guidance does not override the individual responsibility of healthcare professionals to make decisions appropriate to the circumstances of the individual patient, in consultation with the patient and/or guardian or carer.

Commissioners and/or providers have a responsibility to implement the guidance, in their local context, in light of their duties to have due regard to the need to eliminate unlawful discrimination, advance equality of opportunity, and foster good relations. Nothing in this guidance should be interpreted in a way that would be inconsistent with compliance with those duties.

Commissioners and providers have a responsibility to promote an environmentally sustainable health and care system and should assess and reduce the environmental impact of implementing NICE recommendations wherever possible.

PlGF-based testing to help diagnose suspected pre-eclampsia (Triage PlGF test, Elecsys immunoassay sFlt-1/PlGF ratio, DELFIA Xpress PlGF 1-2-3 test, and BRAHMS sFlt-1 Kryptor/BRAHMS PlGF plus Kryptor PE ratio) (DG23)
## Recommendations

1.1 The Triage PlGF test and the Elecsys immunoassay sFlt-1/PlGF ratio, used with standard clinical assessment and subsequent clinical follow-up, are recommended to help rule-out pre-eclampsia in women presenting with suspected pre-eclampsia between 20 weeks and 34 weeks plus 6 days of gestation.

- When pre-eclampsia is not ruled-out using a PlGF-based test result, the result should not be used to diagnose (rule-in) pre-eclampsia (see text box).

1.2 The Triage PlGF test and the Elecsys immunoassay sFlt-1/PlGF ratio, used with standard clinical assessment and subsequent clinical follow-up, show promise in helping to diagnose (rule-in) pre-eclampsia in women presenting with suspected pre-eclampsia between 20 weeks and 34 weeks plus 6 days of gestation. However, there is currently insufficient evidence to recommend their routine adoption for diagnosing pre-eclampsia in the NHS (see text box). Further research is recommended on using these tests in women with suspected pre-eclampsia to rule-in pre-eclampsia (see section 6.2).

1.3 The DELFIA Xpress PlGF 1-2-3 test and BRAHMS sFlt-1 Kryptor/BRAHMS PlGF plus Kryptor PE ratio are not recommended for routine adoption in the NHS. Further research by the companies is needed to show the clinical effectiveness of these tests, including diagnostic accuracy and analytical validity.

This guidance only considers using PlGF-based testing to help diagnose suspected pre-eclampsia. NICE is aware of ongoing research linking low PlGF levels and high sFlt-1/PlGF ratios (positive test results) with placental disease, but placental disease is beyond the scope of this guidance. Therefore, the recommendations in this guidance do not consider using PlGF-based testing for conditions other than suspected pre-eclampsia and this guidance is not intended to give advice on diagnosing or managing placental disease. If placental disease is suspected, additional clinical surveillance may be needed (see section 5.10).
Clinical need and practice

The problem addressed

2.1 Placental growth factor (PlGF)-based tests are intended to be used with clinical judgement and other diagnostic tests, to help diagnose suspected pre-eclampsia. This assessment focuses on diagnosing pre-eclampsia in the second and third trimesters of pregnancy. Using PlGF-based tests in addition to standard clinical assessment could result in a faster and more accurate diagnosis of pre-eclampsia, and better risk assessment for adverse outcomes in women with suspected pre-eclampsia. It could also allow women in whom pre-eclampsia has been ruled out with a PlGF-based test to return to community care instead of being admitted to hospital for observation.

2.2 PlGF-based tests measure the amount of PlGF in blood plasma or serum. PlGF is a protein involved in placental angiogenesis (the development of new blood vessels). In pre-eclampsia, levels of PlGF can be abnormally low. In normal pregnancy, PlGF levels rise and peak at 26–30 weeks, so when PlGF levels do not rise during pregnancy there may be placental dysfunction.

2.3 In addition, some PlGF-based tests measure soluble FMS-like tyrosine kinase-1 (sFlt-1), a protein that is thought to disable other proteins associated with blood vessel formation, such as PlGF. In women who develop pre-eclampsia, the levels of sFlt-1 are higher than those seen in normal pregnancy.

2.4 Four PlGF-based tests were identified during scoping as relevant to this assessment: the Triage PlGF test (Alere International); the Elecsys immunoassay sFlt-1/PlGF ratio (Roche Diagnostics); the DELFIA Xpress PlGF 1-2-3 test (Perkin Elmer); and the BRAHMS sFlt-1 Kryptor/BRAHMS PlGF plus Kryptor PE ratio (Thermo Fisher Scientific).

The condition

2.5 Pre-eclampsia is a potentially serious complication of some pregnancies, which when identified, needs referral to a specialist and hospital admission for both maternal and fetal monitoring. It is thought to be related to problems with the development of the placenta. Pre-eclampsia is characterised by high blood pressure (hypertension) and proteinuria, which occurs when the kidneys leak PIGF-based testing to help diagnose suspected pre-eclampsia (Triage PlGF test, Elecsys immunoassay sFlt-1/PIGF ratio, DELFIA Xpress PlGF 1-2-3 test, and BRAHMS sFlt-1 Kryptor/BRAHMS PlGF plus Kryptor PE ratio) (DG23)
protein into the urine. The presence of either hypertension or proteinuria alone during pregnancy can also indicate a risk of developing pre-eclampsia. Other symptoms include headache, visual disturbances, right upper quadrant abdominal (epigastric) pain, oedema (swelling of the hands, face or feet) and oliguria (low output of urine).

2.6 If pre-eclampsia is not diagnosed and closely monitored, it can lead to potentially life-threatening complications including eclampsia, HELLP syndrome (haemolysis, elevated liver enzymes and low platelets), disseminated intravascular coagulation, stroke, or organ dysfunction. Women who have hypertension or pre-eclampsia during pregnancy may have a higher risk of placental abruption. Women who develop pre-eclampsia during pregnancy may also be at greater risk of cardiovascular disease in later life.

2.7 Gestational hypertension and pre-eclampsia may also affect the fetus, placing it at increased risk of intrauterine growth restriction, prematurity and intrauterine death.

The diagnostic and care pathways

2.8 The NICE guideline on antenatal care recommends measuring blood pressure and testing urine for proteinuria to screen for pre-eclampsia at each routine antenatal visit.

2.9 The NICE pathway on pre-eclampsia describes the assessment and treatment of women at risk of pre-eclampsia or with pre-eclampsia. The NICE guideline on hypertension in pregnancy was used to create the pathway.

Identifying and managing the risk of developing pre-eclampsia

2.10 The NICE guideline on hypertension in pregnancy states that women who are classified as being at high risk of pre-eclampsia are those who have any of the following risk factors identified during the booking appointment:

- hypertensive disease during a previous pregnancy
- chronic kidney disease
autoimmune disease such as systemic lupus erythematosus or antiphospholipid syndrome

- type 1 or type 2 diabetes

- chronic hypertension.

2.11 Women who are classified as being at moderate risk of pre-eclampsia are those who have any of the following risk factors identified during the booking appointment:

- first pregnancy
- age 40 years or older
- pregnancy interval of more than 10 years
- BMI of 35 kg/m\(^2\) or more at first visit
- family history of pre-eclampsia
- multiple pregnancy.

2.12 Women with either 1 high risk factor, or more than 1 moderate risk factor for pre-eclampsia, are advised to take 75 mg of aspirin daily from 12 weeks' gestation until the birth of the baby. They are also considered for more frequent blood pressure monitoring, and assessment for proteinuria. Women who have significant hypertension (diastolic pressure of 90–110 mmHg) or a proteinuria result of 1+ on urinalysis reagent strips need increased surveillance.

Management of pregnancy with gestational hypertension

2.13 The NICE guideline on hypertension in pregnancy defines gestational hypertension as new hypertension presenting after 20 weeks' gestation without significant proteinuria. Increased surveillance is needed to confirm a diagnosis of gestational hypertension, because some women may present with transient hypertension. Women with gestational hypertension are recommended to have assessment for proteinuria at each visit to detect the onset of suspected pre-eclampsia (see table 1).
Table 1 Management of pregnancy with gestational hypertension

<table>
<thead>
<tr>
<th>Degree of hypertension</th>
<th>Mild (140/90 mmHg to 149/99 mmHg)</th>
<th>Moderate (150/100 mmHg to 159/109 mmHg)</th>
<th>Severe (160/110 mmHg or higher)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Admit to hospital</td>
<td>No</td>
<td>No</td>
<td>Yes (until blood pressure is 159/109 mmHg or lower)</td>
</tr>
<tr>
<td>Treat</td>
<td>No</td>
<td>With oral labetalol as first-line treatment</td>
<td>With oral labetalol as first-line treatment</td>
</tr>
<tr>
<td>Measure blood pressure</td>
<td>Not more than once a week</td>
<td>At least twice a week</td>
<td>At least 4 times a day</td>
</tr>
<tr>
<td>Test for proteinuria</td>
<td>At each visit</td>
<td>At each visit</td>
<td>Daily</td>
</tr>
<tr>
<td>Blood tests</td>
<td>Only those for routine antenatal care</td>
<td>Test kidney function, electrolytes, full blood count, transaminases, bilirubin</td>
<td>Test at presentation and then monitor weekly: kidney function, electrolytes, full blood count, transaminases, bilirubin</td>
</tr>
</tbody>
</table>

Birth before 37 weeks should not be offered to women with gestational hypertension whose blood pressure is lower than 160/110 mmHg with or without antihypertensive treatment.

Management of pregnancy with pre-eclampsia

2.14 The NICE guideline on hypertension in pregnancy defines pre-eclampsia as new hypertension with significant proteinuria after 20 weeks' gestation. Women diagnosed with pre-eclampsia should be assessed at each consultation by a healthcare professional trained in the management of hypertensive disorders of pregnancy and offered an integrated package of care that includes admission, testing and treatment that relates to the severity of hypertension (see table 2).
Table 2 Management of pregnancy with pre-eclampsia

<table>
<thead>
<tr>
<th>Degree of hypertension</th>
<th>Mild (140/90 mmHg to 149/99 mmHg)</th>
<th>Moderate (150/100 mmHg to 159/109 mmHg)</th>
<th>Severe (160/110 mmHg or higher)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Admit to hospital</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Treat</td>
<td>No</td>
<td>With oral labetalol as first-line treatment</td>
<td>With oral labetalol as first-line treatment</td>
</tr>
<tr>
<td>Measure blood pressure</td>
<td>At least 4 times a day</td>
<td>At least 4 times a day</td>
<td>More than 4 times a day</td>
</tr>
<tr>
<td>Test for proteinuria</td>
<td>Do not repeat quantification of proteinuria</td>
<td>Do not repeat quantification of proteinuria</td>
<td>Do not repeat quantification of proteinuria</td>
</tr>
<tr>
<td>Blood tests</td>
<td>Monitor the following twice a week: kidney function, bilirubin, electrolytes, full blood count, transaminases</td>
<td>Monitor the following 3 times a week: kidney function, bilirubin, electrolytes, full blood count, transaminases</td>
<td>Monitor the following 3 times a week: kidney function, bilirubin, electrolytes, full blood count, transaminases</td>
</tr>
</tbody>
</table>
3 The diagnostic tests

The interventions

Triage PlGF test

3.1 The Triage PlGF test (Alere International) is a CE-marked, single-use, fluorescence immunoassay device, which is used with the Triage MeterPro point-of-care analyser for the quantitative determination of placental growth factor (PlGF) in blood plasma samples. The test is intended for use with clinical judgement and other diagnostic tests, to help diagnose suspected pre-eclampsia and the level of risk for delivery arising from pre-eclampsia within 14 days of testing. It is recommended for use in pregnant women between 20 weeks and 34 weeks plus 6 days of gestation.

3.2 The test has a limit of detection of 9 picograms/ml and a measurable range of 12 to 3,000 picograms/ml. The test turnaround time is about 15 minutes. The test cut-off values recommended by the company are shown in table 3.

Table 3 Recommended cut-off values for the Triage PlGF test

<table>
<thead>
<tr>
<th>Result</th>
<th>Classification</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>PIGF &lt;12 pg/ml</td>
<td>Test positive – highly abnormal</td>
<td>Highly abnormal and suggestive of patients with severe placental dysfunction and at increased risk for preterm delivery</td>
</tr>
<tr>
<td>PIGF ≥12 pg/ml and &lt;100 pg/ml</td>
<td>Test positive – abnormal</td>
<td>Abnormal and suggestive of patients with placental dysfunction and at increased risk for preterm delivery</td>
</tr>
<tr>
<td>PIGF ≥100 pg/ml</td>
<td>Test negative – normal</td>
<td>Normal and suggestive of patients without placental dysfunction and unlikely to progress to delivery within 14 days of the test</td>
</tr>
</tbody>
</table>

Abbreviations: PIGF, placental growth factor; pg/ml, picograms per millilitre.

Elecsys immunoassay sFlt-1/PIGF ratio

3.3 The Elecsys immunoassay sFlt-1/PIGF ratio (Roche Diagnostics) measures the amounts of PIGF relative to soluble FMS-like tyrosine kinase-1 (sFlt-1; also
known as VEGFR1) in serum samples from women with suspected pre-eclampsia. The ratio is formed by combining the results from 2 CE-marked sandwich electrochemiluminescence immunoassays (Elecsys PIGF and Elecsys sFlt-1 assays), which are compatible with both the Roche Elecsys and the Cobas e automated analysers. The laboratory information system calculates and reports the sFlt-1/PIGF ratio and the individual assay values. The Elecsys immunoassay sFlt-1/PIGF ratio is intended to be used with clinical judgement and other diagnostic tests to diagnose pre-eclampsia. The ratio may also be used to help predict pre-eclampsia, eclampsia and HELLP syndrome in the short term. It may be used for testing pregnant women from 20 weeks' gestation up until the time of delivery.

3.4 The Elecsys sFlt-1 assay has a limit of detection of 10 picograms/ml (measuring range 10 to 85,000 picograms/ml) and a limit of quantitation of 15 picograms/ml. The Elecsys PIGF assay has a limit of detection of 3 picograms/ml (measuring range 3 to 10,000 picograms/ml) and a limit of quantitation of 10 picograms/ml. The turnaround time of the Elecsys immunoassay sFlt-1/PIGF ratio is about 18 minutes. The test cut-off values recommended by the company are shown in table 4.

### Table 4 Recommended cut-off values for the Elecsys immunoassay sFlt-1/PIGF ratio

<table>
<thead>
<tr>
<th>Aid in diagnosis of pre-eclampsia</th>
<th>20 weeks to 33 weeks plus 6 days</th>
<th>Rule-out cut-off</th>
<th>33</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Rule-in cut-off</td>
<td>85</td>
</tr>
<tr>
<td>34 weeks to delivery</td>
<td></td>
<td>Rule-out cut-off</td>
<td>33</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Rule-in cut-off</td>
<td>110</td>
</tr>
<tr>
<td>Short-term prediction of pre-eclampsia (24 weeks to 36 weeks plus 6 days)</td>
<td>Rule-out*</td>
<td>&lt;38</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Rule-in**</td>
<td>&gt;38</td>
<td></td>
</tr>
</tbody>
</table>
**Rule-out pre-eclampsia for 1 week.**

**Rule-in pre-eclampsia within 4 weeks.**

**DELFIA Xpress PIGF 1-2-3 test**

3.5 The DELFIA Xpress PIGF 1-2-3 test (Perkin Elmer) is a CE-marked solid-phase, 2-site fluoroimmunometric sandwich assay for the quantitative determination of PIGF in serum samples. The assay is compatible with the 6000 DELFIA Xpress random access analyser. The test is intended to help diagnose pre-eclampsia during the second and third trimesters of pregnancy, and is used with clinical assessment.

3.6 The assay has a limit of detection of 1.9 picograms/ml (measuring range 1.9 to 4,000 picograms/ml) and a limit of quantitation of 3.3 picograms/ml. The test has a turnaround time of about 30 minutes. The cut-off values for PIGF measurements obtained during the second trimester are highly dependent on gestational day and the company suggests that cut-off values should be established by individual laboratories. In the third trimester, in addition to laboratory-calculated cut-off values based on gestational day, the company suggests that a fixed cut-off value of 184 picograms/ml can be used. Levels of PIGF lower than 184 picograms/ml indicate an increased probability of pre-eclampsia developing.

**BRAHMS sFlt-1 Kryptor/BRAHMS PIGF plus Kryptor PE ratio**

3.7 The BRAHMS sFlt-1 Kryptor/BRAHMS PIGF plus Kryptor PE ratio (Thermo Fisher Scientific) is formed by combining the results from 2 automated immunofluorescent sandwich assays, the BRAHMS sFlt-1 Kryptor assay and the BRAHMS PIGF plus Kryptor assay. They are indicated for the quantitative determination of sFlt-1 and PIGF in serum samples and are compatible with the BRAHMS Kryptor compact plus analyser. The assays are intended to be run at the same time, with the analyser reporting both the concentrations for each assay and the sFlt-1/PIGF ratio. The BRAHMS sFlt-1 Kryptor/BRAHMS PIGF plus KRYPTOR PE ratio is intended to be used with clinical assessment to help diagnose pre-eclampsia.

3.8 The BRAHMS sFlt-1 Kryptor assay has a limit of detection of 22 picograms/ml (measuring range 22 to 90,000 picograms/ml) and a limit of quantitation of
34 picograms/ml. The BRAHMS PIGF plus Kryptor assay has a limit of detection of 3.6 picograms/ml (measuring range 3.6 to 7,000 picograms/ml) and a limit of quantitation of 6.9 picograms/ml. Reference ranges for each of the assays and the sFlt-1/PIGF ratio are given in the company instructions, and the company recommends that individual laboratories should validate these ranges or establish their own reference ranges before use. The turnaround time for the BRAHMS sFlt-1 Kryptor assay is 9 minutes and the turnaround time for the BRAHMS PIGF plus Kryptor assay is 29 minutes.

The comparator

3.9 The comparator used in this assessment is standard clinical assessment to help diagnose suspected pre-eclampsia, guided by a combination of the following clinical information:

- maternal hypertension (categorised as mild, moderate or severe)
- quantitative proteinuria test
- clinical symptoms suggestive of pre-eclampsia (for example, headache, oedema, visual disturbances)
- fetal growth restriction.
4  Outcomes

The diagnostics advisory committee (section 9) considered evidence from a number of sources (section 10). Full details of all the evidence are in the committee papers.

How outcomes were assessed

4.1  The assessment consisted of:

- A systematic review of the evidence on the diagnostic accuracy of the 4 index tests for the assessment of suspected pre-eclampsia in the second and third trimesters of pregnancy.

- A review of cost-effectiveness evidence on the 4 index tests for the assessment of suspected pre-eclampsia in the second and third trimesters of pregnancy.

- A de novo economic model designed to assess the cost effectiveness of placental growth factor (PlGF)-based tests when used with standard clinical assessment compared with standard clinical assessment alone for the assessment of suspected pre-eclampsia in the second and third trimesters of pregnancy.

Assessment of test accuracy

4.2  Studies were included in the systematic review if they contained information on:

- Women with suspected pre-eclampsia between 20 weeks and 36 weeks plus 6 days of pregnancy who had blood pressure assessment and qualitative assessment of proteinuria.

- Triage PIGF test; Elecsys immunoassay sFlt-1/PIGF ratio; DELFIA Xpress PIGF 1-2-3 test; or BRAHMS sFlt-1 Kryptor/BRAHMS PlGF plus Kryptor PE ratio, with standard clinical assessment, or with standard clinical assessment excluding quantitative determination of proteinuria.

- A reference standard of clinical assessment guided by maternal hypertension, proteinuria, symptoms suggestive of pre-eclampsia, and ultrasound fetal growth measurements.

- Test performance outcomes, including diagnostic and prognostic test accuracy (sensitivity, specificity, incidence and related outcome measures) for pre-eclampsia.
Overview of included studies

4.3 After searches and inclusion screening, 12 publications of 4 studies were included in the review. Two of these studies were on the Triage PlGF test and 2 studies were on the Elecsys immunoassay sFlt-1/PIGF ratio. None of the studies included more than 1 test; so no head-to-head comparisons of the index tests were available. None of the included studies were on the Perkin Elmer DELFIA Xpress PlGF 1-2-3 test or the Thermo Fisher Scientific BRAHMS sFlt-1 Kryptor/BRAHMS PlGF plus Kryptor ratio.

4.4 The PETRA study was a multicentre study of the Triage PlGF test. The details of this study are academic in confidence at the time of writing this diagnostics guidance.

4.5 The PELICAN study (2013) was a prospective, single cohort study of the Triage PlGF test done in 7 centres in the UK and Ireland. It included 287 women with suspected pre-eclampsia between 20 weeks and 34 weeks plus 6 days of gestation, and 137 women with suspected pre-eclampsia between 35 weeks and 36 weeks plus 6 days of gestation.

4.6 The PROGNOSIS study (2016) was a prospective, international multicentre study of the Elecsys immunoassay sFlt-1/PIGF ratio, with 1 study centre in the UK. The study had 2 cohorts: a 'feasibility' cohort with 500 patients to derive a cut-off value based prediction model for the sFlt-1/PIGF ratio, and a 'validation' cohort with 550 patients to test the model. It included women with suspected pre-eclampsia between 24 weeks and 36 weeks and 6 days of gestation.

4.7 The study by Alvarez-Fernandez et al. (2014) was a retrospective study of the Elecsys immunoassay sFlt-1/PIGF ratio in a single cohort of patients. The study was done in Spain and included 62 women with suspected pre-eclampsia between 20 weeks and 34 weeks of gestation.

4.8 The PELICAN study (2013) and the PROGNOSIS study (2016) defined hypertensive disorders according to the American College of Obstetrics and Gynecology practice bulletin (2002). This gives a broader definition of pre-eclampsia than the NICE guideline on hypertension in pregnancy, because it includes superimposed pre-eclampsia and atypical pre-eclampsia. The definition of pre-eclampsia used in the PETRA study is academic in confidence at the time.
of writing this diagnostics guidance. The study by Alvarez-Fernandez et al. (2014) used a simple definition of pre-eclampsia, which expands on the definition in the NICE guideline on hypertension in pregnancy by including pre-existing proteinuria with superimposed pre-eclampsia.

4.9 The PELICAN study (2013), the PROGNOSIS study (2016) and the study by Alvarez-Fernandez et al. (2014) were judged to be at low risk of bias using the Cochrane Collaboration adaptation of the QUADAS tool. However, all 3 studies were judged to be at high risk of clinical review bias because the diagnosis of pre-eclampsia was based solely on whether index-test results were above or below the cut-off value, whereas in clinical practice index-test results would likely be interpreted alongside clinical signs and symptoms, such as information about hypertension and proteinuria. The PETRA study is academic in confidence at the time of writing this diagnostics guidance.

Diagnostic-accuracy results for the Triage PIGF test

4.10 Diagnostic-accuracy results for the Alere Triage PIGF test are available for 3 test cut-off values: 100 picograms/ml, the fifth centile of PIGF concentration for gestational age, and 12 picograms/ml. PIGF concentrations above 100 picograms/ml are considered normal and would be used to identify women unlikely to develop pre-eclampsia needing delivery within 14 days. As such, a result of 100 picograms/ml or greater is used to rule-out pre-eclampsia.

4.11 Results from the PELICAN study (2013) show that the Triage PIGF test at cut-off values of 100 picograms/ml and the fifth centile of PIGF concentration for gestational age gave high sensitivity with good precision for identifying women likely to develop pre-eclampsia needing delivery within 14 days of testing, when presenting with suspected pre-eclampsia before 35 weeks’ gestation. The cut-off value of 12 picograms/ml yielded lower sensitivity for identifying women likely to develop pre-eclampsia needing delivery within 14 days of testing, when presenting with suspected pre-eclampsia between 20 weeks and 34 weeks plus 6 days of gestation (see table 5).
### Table 5 PELICAN study results – Triage PlGF test accuracy for predicting pre-eclampsia needing delivery within 14 days for women presenting between 20 weeks and 34 weeks plus 6 days of gestation

<table>
<thead>
<tr>
<th>Test cut-off</th>
<th>Sensitivity (95% CI)</th>
<th>Specificity (95% CI)</th>
<th>PPV (95% CI)</th>
<th>NPV (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;100 pg/ml</td>
<td>0.96 (0.89 to 0.99)</td>
<td>0.56 (0.49 to 0.63)</td>
<td>0.44 (0.36 to 0.52)</td>
<td>0.98 (0.93 to 1.00)</td>
</tr>
<tr>
<td>≥100 pg/ml</td>
<td>0.96 (0.89 to 0.99)</td>
<td>0.56 (0.49 to 0.63)</td>
<td>0.43 (0.36 to 0.51)</td>
<td>0.98 (0.93 to 1.00)</td>
</tr>
<tr>
<td>&lt;fifth centile</td>
<td>0.96 (0.89 to 0.99)</td>
<td>0.55 (0.48 to 0.61)</td>
<td>0.43 (0.36 to 0.51)</td>
<td>0.98 (0.93 to 1.00)</td>
</tr>
<tr>
<td>&lt;12 pg/ml</td>
<td>0.63 (0.51 to 0.74)</td>
<td>0.90 (0.85 to 0.94)</td>
<td>0.70 (0.57 to 0.80)</td>
<td>0.87 (0.82 to 0.91)</td>
</tr>
</tbody>
</table>

Abbreviations: CI, confidence interval; NPV, negative predictive value; PPV, positive predictive value; pg/ml, picograms per millilitre.

### Table 6 PELICAN study results – Triage PlGF test accuracy for predicting pre-eclampsia needing delivery within 14 days for women presenting between 35 weeks and 36 weeks plus 6 days of gestation

<table>
<thead>
<tr>
<th>Test cut-off</th>
<th>Sensitivity (95% CI)</th>
<th>Specificity (95% CI)</th>
<th>PPV (95% CI)</th>
<th>NPV (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;fifth centile</td>
<td>0.70 (0.58 to 0.81)</td>
<td>0.64 (0.52 to 0.75)</td>
<td>0.65 (0.53 to 0.76)</td>
<td>0.69 (0.57 to 0.80)</td>
</tr>
<tr>
<td>&lt;12 pg/ml</td>
<td>0.22 (0.13 to 0.34)</td>
<td>0.91 (0.82 to 0.97)</td>
<td>0.71 (0.48 to 0.89)</td>
<td>0.55 (0.46 to 0.64)</td>
</tr>
</tbody>
</table>

4.12 Results from the PELICAN study (2013) also show that for the cut-off values for the fifth centile of PlGF for gestational age and 12 picograms/ml, the Triage PlGF test had poor diagnostic accuracy for predicting pre-eclampsia needing delivery within 14 days in women presenting with suspected pre-eclampsia between 35 weeks and 36 weeks plus 6 days of gestation (see table 6).
4.13 The PELICAN study (2013) also reported that the Triage PlGF test at a cut-off value of 100 picograms/ml had high sensitivity for predicting preterm pre-eclampsia and delivery within 14 days of testing independent of the pre-eclampsia diagnosis. A cut-off value of 12 picograms/ml had poor sensitivity but good specificity for predicting preterm delivery independent of the pre-eclampsia diagnosis (see table 7).

**Table 7 PELICAN study results – Triage PlGF test accuracy for other outcomes for women presenting between 20 weeks and 34 weeks plus 6 days of gestation**

<table>
<thead>
<tr>
<th>Test cut-off</th>
<th>Sensitivity (95% CI)</th>
<th>Specificity (95% CI)</th>
<th>PPV (95% CI)</th>
<th>NPV (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Preterm pre-eclampsia</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;100 pg/ml</td>
<td>0.90 (0.83 to 0.95)</td>
<td>0.65 (0.58 to 0.73)</td>
<td>0.65 (0.57 to 0.72)</td>
<td>0.90 (0.83 to 0.95)</td>
</tr>
<tr>
<td>≥100 pg/ml</td>
<td>0.94 (0.87 to 0.98)</td>
<td>0.57 (0.50 to 0.64)</td>
<td>0.47 (0.39 to 0.55)</td>
<td>0.96 (0.91 to 0.99)</td>
</tr>
<tr>
<td><strong>Delivery within 14 days of testing</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;100 pg/ml</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≥100 pg/ml</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Preterm delivery</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;12 pg/ml</td>
<td>0.44 (0.36 to 0.52)</td>
<td>0.97 (0.93 to 0.99)</td>
<td>0.94 (0.86 to 0.98)</td>
<td>0.62 (0.55 to 0.68)</td>
</tr>
</tbody>
</table>

Abbreviations: CI, confidence interval; NPV, negative predictive value; PPV, positive predictive value; pg/ml, picograms per millilitre.

4.14 The PROGNOSIS study (2016) derived a cut-off value of 38; values below 38 were considered negative and were used to rule-out pre-eclampsia within 1 week; values above 38 were considered positive and used to rule-in pre-eclampsia within 4 weeks. Results from the combined cohort show that for women with suspected pre-eclampsia between 24 weeks and 36 weeks and...
6 days of gestation, sensitivity and specificity for ruling out pre-eclampsia within 1 week was relatively high. Sensitivity for ruling in pre-eclampsia within 4 weeks was lower than for ruling out pre-eclampsia within 1 week, but specificity was relatively high (see table 8).

Table 8 PROGNOSIS study results – Elecsys immunoassay sFlt-1/PIGF ratio accuracy for women presenting between 24 weeks and 36 weeks plus 6 days of gestation; cut-off value 38

<table>
<thead>
<tr>
<th></th>
<th>Sensitivity (95% CI)</th>
<th>Specificity (95% CI)</th>
<th>PPV (95% CI)</th>
<th>NPV (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rule-out of pre-eclampsia within 1 week</td>
<td>0.86 (0.73 to 0.94)</td>
<td>0.79 (0.77 to 0.82)</td>
<td>0.17 (0.12 to 0.22)</td>
<td>0.99 (0.98 to 1.00)</td>
</tr>
<tr>
<td>Rule-in of pre-eclampsia within 4 weeks</td>
<td>0.70 (0.62 to 0.78)</td>
<td>0.83 (0.81 to 0.86)</td>
<td>0.39 (0.33 to 0.45)</td>
<td>0.95 (0.93 to 0.96)</td>
</tr>
</tbody>
</table>

Abbreviations: CI, confidence interval; NPV, negative predictive value; PPV, positive predictive value; pg/ml, picograms per millilitre.

The study by Alvarez-Fernandez et al. (2014) analysed the Elecsys immunoassay sFlt-1/PIGF ratio at cut-off values of 23 and 85 (see table 9). Results show that for women presenting with suspected pre-eclampsia between 20 weeks and 34 weeks of gestation, the cut-off value of 23 had higher sensitivity than the cut-off value of 85 for rule-out of pre-eclampsia within 3 weeks (92% compared with 56%). Specificity was lower for the cut-off value of 23 than the cut-off value of 85 for the rule-out of pre-eclampsia within 3 weeks (81% compared with 97% respectively).

Table 9 Alvarez-Fernandez et al. (2014) study results – Elecsys immunoassay sFlt-1/PIGF ratio accuracy for rule-out of pre-eclampsia within 3 weeks for women presenting between 20 weeks and 34 weeks of gestation

<table>
<thead>
<tr>
<th>Test cut-off</th>
<th>Sensitivity (95% CI)</th>
<th>Specificity (95% CI)</th>
<th>PPV (95% CI)</th>
<th>NPV (95% CI)</th>
</tr>
</thead>
</table>

The study by Alvarez-Fernandez et al. (2014) analysed the Elecsys immunoassay sFlt-1/PIGF ratio at cut-off values of 23 and 85 (see table 9). Results show that for women presenting with suspected pre-eclampsia between 20 weeks and 34 weeks of gestation, the cut-off value of 23 had higher sensitivity than the cut-off value of 85 for rule-out of pre-eclampsia within 3 weeks (92% compared with 56%). Specificity was lower for the cut-off value of 23 than the cut-off value of 85 for the rule-out of pre-eclampsia within 3 weeks (81% compared with 97% respectively).
Abbreviations: CI, confidence interval; NPV, negative predictive value; PPV, positive predictive value.

### Review of cost-effectiveness evidence

4.16 Searches were done to identify existing studies investigating the cost effectiveness of Triage PIGF test, Elecsys immunoassay sFlt-1/PIGF ratio, DELFIA Xpress PIGF 1-2-3 test and BRAHMS sFlt-1 Kryptor/BRAHMS PIGF plus Kryptor PE ratio for diagnosing pre-eclampsia in the second and third trimesters of pregnancy.

4.17 Four studies described in 4 full text articles (1 unpublished) were included in the review. All studies were cost analyses; that is, they focused on potential savings and did not formally evaluate health outcomes of the mother or baby.

4.18 Two studies by Hadker et al. (2010 and 2013) used the same model, populated with identical clinical inputs, to address UK and German healthcare payer perspectives. The study population was all women assessed for pre-eclampsia after 20 weeks’ gestation. The intervention used was standard assessment plus Elecsys immunoassay sFlt-1/PIGF ratio (cut-off value of 85) compared with standard assessment alone. Base-case results from a UK healthcare payer perspective show that there was an overall reduction in cost of £945 per patient from £2,726 to £1,781 associated with the use of the Elecsys immunoassay sFlt-1/PIGF ratio. From a German healthcare payer perspective, base-case results show an overall cost reduction of €637 per patient from €1,579 to €942 associated with the use of the Elecsys immunoassay sFlt-1/PIGF ratio.

4.19 A study by Schnettler et al. (2013) included women before 34 weeks’ gestation with suspected pre-eclampsia. The intervention used was standard clinical assessment plus Elecsys immunoassay sFlt-1/PIGF ratio (cut-off value of 85) compared with standard clinical assessment alone. The perspective was that of a US healthcare payer. Base-case results show an overall cost reduction of
$1,215 per patient, from $3,022 to $1,807, associated with the use of the Elecsys immunoassay sFlt-1/PIGF ratio.

4.20 The fourth study was by Hunter et al. (2013) and is unpublished and commercial in confidence.

**Economic evaluation**

4.21 A decision tree model was developed to assess the cost effectiveness of PIGF-based tests used with standard clinical assessment compared with standard clinical assessment alone in women with suspected pre-eclampsia in 2 groups:

- those presenting between 20 weeks and 33 weeks plus 6 days of gestation
- those presenting between 34 weeks and 36 weeks plus 6 days of gestation.

**Model structure**

4.22 The model used a linked evidence approach in which maternal, fetal and neonatal outcomes were modelled from diagnostic test accuracy and prevalence of pre-eclampsia data. The model had 4 components: risk stratification, management, maternal outcomes, and fetal and neonatal outcomes.

4.23 Women with suspected pre-eclampsia were classified as being at high, intermediate or low risk of pre-eclampsia. This was based on clinical signs, symptoms or findings with or without the addition of a PIGF-based test. The probability of pre-eclampsia in women with suspected pre-eclampsia was based on the prevalence of pre-eclampsia and the reported sensitivity and specificity of each diagnostic strategy.

4.24 Suspected pre-eclampsia could be managed using expectant management or immediate delivery, dependent on the risk of pre-eclampsia (high, intermediate or low) and the number of weeks’ gestation. Expectant management involves monitoring clinical signs, symptoms and findings, active management of conditions such as hypertension, and planned delivery at 37 weeks of gestation. Immediate delivery involves delivery much sooner, irrespective of gestational age because of clinical findings indicating severe risk to a pregnant woman or
fetus. A low risk of pre-eclampsia is managed on the gestational hypertension pathway (expectant management). An intermediate risk of pre-eclampsia is managed on a modified version of the gestational hypertension pathway, which has an increased frequency of surveillance (expectant monitoring). A high risk of pre-eclampsia presenting before 35 weeks' gestation is managed using expectant monitoring when there are no signs of increased risk for the mother or fetus. A high risk of pre-eclampsia presenting from 35 weeks' gestation is managed by immediate delivery when there are signs of increased risk for the mother or fetus. These assumptions are in line with the NICE guideline on hypertension in pregnancy.

4.25 Maternal and fetal outcomes in the model are assumed to be related to the presence or absence of pre-eclampsia. As a result, the outcome components of the model are preceded by an evaluation of true disease status. This is the probability of pre-eclampsia in women in each of the risk categories (high, intermediate, low) assigned in the first stage of the model.

4.26 The maternal outcome component begins with delivery, resulting either from spontaneous labour, induced labour, or planned caesarean section. Each of these modes of delivery may be associated with a risk of conversion to assisted or instrumental vaginal delivery, or to emergency caesarean section. Each mode of delivery is associated with a risk of a severe adverse event associated with the progression of severity of pre-eclampsia during the delivery, which can result in convulsions. These adverse events may result in admission to an intensive or high-dependency care unit and the need for anti-convulsive therapy. The model assumes that women who do not have convulsions are transferred to the ward after delivery and those who do not have any further adverse events have a normal length of stay for the given mode of delivery.

4.27 The fetal and neonatal outcome component of the model first establishes whether the labour results in a live birth or stillbirth. After a live birth, a neonate may or may not need to be admitted to a neonatal intensive care unit or high dependency unit, the probability of which is related to gestational age, presence or absence of pre-eclampsia, principal cause of early delivery (maternal condition or fetal distress), mode of delivery, and the presence or absence of complications during delivery. The neonate may then survive or die.
Test accuracy data for the Triage PlGF test were taken from the PELICAN study (2013) and test accuracy data for the Elecsys immunoassay sFlt-1/PlGF ratio were taken from the PROGNOSIS study (2016). Test accuracy data for standard clinical assessment were taken from a study included in the systematic review of economic evaluations (Schnettler et al. 2013). The data on prevalence of pre-eclampsia were taken from the PELICAN study, which was done in the UK. Other clinical and resource use inputs were taken from a variety of published studies.

Test costs used in the base-case model were taken from economic models produced by the companies. These were slightly different from the list price test costs submitted by the companies. The list price for a single Triage PlGF test is £40. The list price for a single Elecsys immunoassay sFlt-1/PlGF ratio is £57.23. Other costs were taken from NHS reference costs, the British national formulary and published literature.

Utility values were taken from the published literature, however, many values had to be mapped from SF-36 to EQ-5D (see table 10).

### Table 10 Utility values used in the economic model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline QALYs from (vaginal) delivery to 6 months post-partum</td>
<td>Birth to 3 weeks post-partum</td>
<td>0.0389</td>
</tr>
<tr>
<td></td>
<td>3–12 weeks post-partum</td>
<td>0.1496</td>
</tr>
<tr>
<td></td>
<td>12–6 months post-partum</td>
<td>0.2171</td>
</tr>
<tr>
<td>Decrement for caesarean delivery (birth to 3 weeks post-partum)</td>
<td>Non-emergency caesarean section</td>
<td>0.0050</td>
</tr>
<tr>
<td></td>
<td>Emergency caesarean section</td>
<td>0.0092</td>
</tr>
</tbody>
</table>
Decrement for non-spontaneous delivery (induced) | 3–6 months post-partum | 0.0084 | Petrou et al. 2009

Abbreviation: QALYs, quality-adjusted life years.

**Base-case results**

4.31 Key assumptions made in the model include:

- UK guidelines for management of suspected pre-eclampsia, gestational hypertension and pre-eclampsia are followed.

- The 2 different outcomes, pre-eclampsia needing delivery within 14 days (Triage PIGF test) and pre-eclampsia within 4 weeks irrespective of delivery time (Elecsys immunoassay sFlt-1/PIGF ratio) are compared as if they were the same.

- Costs of neonatal intensive care unit stay capture the effects of neonatal morbidity for deliveries occurring between 35 and 37 weeks’ gestation.

- Tests are done in a central laboratory.

- The unit costs associated with birth are not dependent on whether the mother has hypertension or pre-eclampsia.

- Utility scores for birth are assumed to last for 3 weeks.

4.32 For women with suspected pre-eclampsia presenting before 35 weeks’ gestation, in the base case, total costs varied between £6,048 for the Triage PIGF test to £8,945 for standard clinical assessment. Both the Triage PIGF test and the Elecsys immunoassay sFlt-1/PIGF ratio were cost saving compared with standard clinical assessment. Cost reductions per patient compared with standard clinical assessment were £2,896 for the Triage PIGF test and £2,488 for the Elecsys immunoassay sFlt-1/PIGF ratio. Total quality-adjusted life years (QALYs) for each diagnostic strategy were similar, with no more than 0.00076 QALYs separating the most clinically-effective diagnostic strategy and the least clinically-effective diagnostic strategy (see table 11). Although the PIGF-based tests dominated standard clinical assessment, the differences in QALYs were so small that they could be considered comparable and so the calculation of incremental cost-effectiveness ratios (ICERs) was not appropriate.
For women with suspected pre-eclampsia presenting between 35 and 37 weeks' gestation, the cost differences were much smaller than in women with suspected pre-eclampsia presenting before 35 weeks. In the base case, total costs varied between £3,393 for the Triage PlGF test and £3,758 for standard clinical assessment. Both strategies including PIGF-based tests were cost saving compared with standard clinical assessment. Cost reductions per patient compared with standard assessment were £365 for the Triage PlGF test and £174 for the Elecsys immunoassay sFlt-1/PlGF ratio. There was no difference in QALYs between any of the strategies (see table 12). Therefore, ICERs could not be calculated in this analysis.

### Table 11 Base-case results for women presenting with suspected pre-eclampsia before 35 weeks' gestation

<table>
<thead>
<tr>
<th>Strategy</th>
<th>Costs</th>
<th>QALYs</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total</td>
<td>Increment compared with standard clinical assessment</td>
</tr>
<tr>
<td>Triage PlGF test</td>
<td>£6,048</td>
<td>−£2,896</td>
</tr>
<tr>
<td>Elecsys immunoassay sFlt-1/PlGF ratio</td>
<td>£6,456</td>
<td>−£2,488</td>
</tr>
<tr>
<td>Standard clinical assessment</td>
<td>£8,945</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviation: QALYs, quality-adjusted life years.

### Table 12 Base-case results for women presenting with suspected pre-eclampsia between 35 and 37 weeks' gestation

<table>
<thead>
<tr>
<th>Strategy</th>
<th>Costs</th>
<th>QALYs</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total</td>
<td>Increment compared with standard clinical assessment</td>
</tr>
<tr>
<td>Triage PlGF test</td>
<td>£3,393</td>
<td>−£365</td>
</tr>
</tbody>
</table>
4.34 A scenario analysis looking at using PIGF-based tests as an alternative to quantitative proteinuria testing was done. Quantitative proteinuria testing may delay the diagnostic assessment of women with suspected pre-eclampsia, leading to some being unnecessarily admitted for overnight stays to await results of the quantitative proteinuria test. No evidence was found relating to this question, so a simple cost-based analysis was done. Results suggested that cost savings could increase slightly if PIGF-based tests replace quantitative proteinuria.

4.35 A scenario analysis looking at the effect on costs of doing the Triage PIGF test in a near-patient setting (a midwifery day unit) rather than in a hospital laboratory was also done. The same unit cost of the test was assumed as in the base case. It was also assumed that because of the adoption of near-patient testing in the midwifery day unit, no women need to be admitted overnight while waiting for test results. For the Elecsys immunoassay sFlt-1/PIGF ratio, it was assumed that 10% of women being assessed for suspected pre-eclampsia need an overnight stay while waiting for test results, and for standard clinical assessment a range of 10–50% was assumed. Results suggested that cost savings could increase slightly as a result of doing near-patient testing rather than hospital laboratory testing.

4.36 A scenario analysis was done that considered using PIGF-based tests to rule-out (and not rule-in) pre-eclampsia. Results suggested that when PIGF-based tests are used to rule-out (and not rule-in) pre-eclampsia, the total costs increase compared with the base case. For women with suspected pre-eclampsia presenting before 35 weeks' gestation, the total costs for the Triage PIGF test increased by £1,939 and the total costs for the Elecsys immunoassay sFlt-1/PIGF ratio increased by £294. However, the Triage PIGF test and the Elecsys

## Scenario and sensitivity analyses

<table>
<thead>
<tr>
<th></th>
<th>£3,584</th>
<th>−£174</th>
<th>0.3954</th>
<th>0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Elecsys immunoassay sFlt-1/PIGF ratio</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Standard clinical assessment</td>
<td>£3,758</td>
<td></td>
<td>0.3954</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviation: QALYs, quality-adjusted life years.
immunoassay sFlt-1/PIGF ratio remain cost-effective compared with standard clinical assessment for women presenting with suspected pre-eclampsia between 20 weeks and 34 weeks plus 6 days of gestation (see table 13).

### Table 13 Rule-out only scenario analysis results for women presenting with suspected pre-eclampsia before 35 weeks’ gestation

<table>
<thead>
<tr>
<th>Strategy</th>
<th>Costs</th>
<th>QALYs</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total</td>
<td>Increment compared with standard clinical assessment</td>
</tr>
<tr>
<td>Elecsys immunoassay sFlt-1/PIGF ratio</td>
<td>£6,750</td>
<td>−£2,195</td>
</tr>
<tr>
<td>Triage PlGF test</td>
<td>£7,987</td>
<td>−£958</td>
</tr>
<tr>
<td>Standard clinical assessment</td>
<td>£8,945</td>
<td>0.39368</td>
</tr>
</tbody>
</table>

Abbreviation: QALYs, quality-adjusted life years.

4.37 Deterministic sensitivity analyses were done on the following model inputs:

- test sensitivity and specificity
- prevalence of pre-eclampsia in women suspected of having pre-eclampsia
- test cost
- probability of admission and length of stay in neonatal intensive care
- distribution of hypertension across women included in the model.

PIGF-based test strategies remained cost saving compared with standard clinical assessment in all sensitivity analyses. Changing the assumptions around neonatal intensive care had the biggest impact on costs, but strategies involving a PIGF-based test remained cost saving compared with standard clinical assessment.
Analysis of index tests not included in the base-case model

4.38 The BRAHMS sFlt-1 Kryptor/BRAHMS PIGF plus Kryptor PE ratio and the DELFIA Xpress 1-2-3 PIGF test were included in the scope for the assessment but were not included in the base-case economic analysis because there were insufficient data. One study comparing the diagnostic accuracy of the BRAHMS sFlt-1 Kryptor/BRAHMS PIGF plus Kryptor PE ratio with the Elecsys immunoassay sFlt-1/PIGF ratio was identified, but it did not meet the inclusion criteria for the systematic review of diagnostic test accuracy (Anderson et al. 2015). However, a threshold cost analysis was done for the BRAHMS sFlt-1 Kryptor/BRAHMS PIGF plus Kryptor PE ratio by assuming that it has equivalent diagnostic accuracy to the Elecsys immunoassay sFlt-1/PIGF ratio.

4.39 The study by Andersen et al. (2015) compared the diagnostic accuracy of the BRAHMS sFlt-1 Kryptor/BRAHMS PIGF plus Kryptor PE ratio with the Elecsys immunoassay PIGF/sFlt-1 ratio. It had 39 patients with confirmed pre-eclampsia and 76 patients with normotensive pregnancies. The study was not done in a population of patients with suspected pre-eclampsia, so sensitivity and specificity may be exaggerated. The BRAHMS sFlt-1 Kryptor/BRAHMS PIGF plus Kryptor PE ratio had a cost based on throughput, therefore the threshold analysis used regression analysis to estimate the point at which the BRAHMS sFlt-1 Kryptor/BRAHMS PIGF plus Kryptor PE ratio would cost the same as the Elecsys immunoassay sFlt-1/PIGF ratio, assuming equivalent sensitivity and specificity. The results of this analysis are commercial in confidence.

4.40 No relevant information on the diagnostic accuracy of the DELFIA Xpress 1-2-3 PIGF test was available, so no cost-analysis was done.
5 Considerations

5.1 The committee considered the potential effects of pre-eclampsia on a pregnant woman's life and on the life of her partner. The committee heard from a patient expert that women who have had pre-eclampsia fear that they will get pre-eclampsia again if they have another baby. It heard further that a few women decide not to complete their family because of the experience of pre-eclampsia in a previous pregnancy. The committee also heard that there are currently no tests that can be used to confidently rule-out the presence of pre-eclampsia. Therefore pregnant women with suspected pre-eclampsia often need increased monitoring or admission to hospital, which can be inconvenient and can cause anxiety. The committee concluded that tests that could help to assess suspected pre-eclampsia could help to reduce anxiety and unnecessary monitoring, in women with suspected pre-eclampsia or pre-eclampsia.

5.2 The committee considered the role of placental growth factor (PlGF)-based tests and noted that PlGF is also a biomarker for placental disease. It heard from clinical experts on the committee that the occurrence of placental disease does not mean that pre-eclampsia is present. The committee also heard that the requirements for the surveillance and management of placental disease are not well defined and noted that surveillance and management of this disease were beyond the scope of this guidance.

5.3 The Committee considered the definitions of pre-eclampsia used in the studies compared with the definition of pre-eclampsia used in the NICE guideline on hypertension in pregnancy. It noted that the definition in the NICE guideline is narrower than the definitions used in the studies. It heard from clinical experts on the committee that the decision on whether to deliver the baby in a woman with pre-eclampsia is based on clinical symptoms suggesting risk to the mother or baby, which aligns with the expanded definitions of pre-eclampsia used in the studies. The committee concluded that the expanded definitions of pre-eclampsia used in the studies do not affect the generalisability of the studies to clinical practice in the NHS.

5.4 The committee further considered the generalisability of the studies included in the systematic review of diagnostic accuracy to clinical practice in the NHS. The committee noted that 3 of the 4 studies were done mainly outside the UK. It heard from clinical experts on the committee that in the NHS, women with
suspected pre-eclampsia would be referred from community care to a day unit for further assessment before a decision on whether to admit them to hospital is made. It heard further that unlike in the NHS, some countries do not have day units so suspected pre-eclampsia would be managed differently. In countries where there are no day units, women with suspected pre-eclampsia would be referred from community care directly to hospital for further assessment. This results in suspected pre-eclampsia being managed for longer in the community than in the NHS, but if symptoms of suspected pre-eclampsia become worse, admission to hospital would happen earlier. The committee concluded that this may affect the generalisability of some of the studies to clinical practice in the NHS.

5.5 The committee considered the test accuracy reported in the studies. It noted that test accuracy of the Triage PlGF test was better in women presenting with suspected pre-eclampsia between 20 weeks and 34 weeks plus 6 days of gestation than in women presenting with suspected pre-eclampsia between 35 weeks and 36 weeks plus 6 days of gestation. It heard from clinical experts on the committee that decisions on managing suspected pre-eclampsia are more difficult in women at earlier gestations. For example, in women with pre-eclampsia between 35 weeks and 36 weeks plus 6 days of gestation, the average time to delivery is about 4–5 days from diagnosis of pre-eclampsia. However, in women at earlier gestations the risk to the baby from premature delivery is higher than in women at later gestation, so the average time to delivery of a woman at 28 weeks’ gestation is about 13 days. The committee heard from a clinical expert that the reason for delaying delivery at early gestations is to reduce the risk of adverse neonatal outcomes. For a woman at 26 weeks’ gestation, every day that the pregnancy can be extended leads to a 3% improvement in neonatal outcomes. The committee concluded that PlGF-based tests would be more clinically useful in women presenting before 35 weeks’ gestation.

5.6 The committee considered the 2 tests that were not included in the base-case economic evaluation (the DELFIA Xpress PlGF 1-2-3 test and the BRAHMS sFlt-1 Kryptor/BRAHMS PlGF plus Kryptor PE ratio). It noted that the DELFIA Xpress PlGF 1-2-3 test does not have validated cut-off values, and neither test has diagnostic accuracy data for a population of women presenting with suspected pre-eclampsia between 20 weeks and 36 weeks and 6 days of gestation. The committee concluded that the diagnostic accuracy of the DELFIA
Xpress PIGF 1-2-3 test and the BRAHMS sFlt-1 Kryptor/BRAHMS PIGF plus Kryptor PE ratio could not be assumed to be equivalent to the diagnostic accuracy of the Triage PIGF test and the Elecsys immunoassay sFlt-1/PIGF ratio.

5.7 The committee considered the use of PIGF-based tests to rule-out pre-eclampsia. The committee noted that the negative predictive value for the Triage PIGF test for the rule-out of pre-eclampsia needing delivery within 14 days in women presenting with suspected pre-eclampsia between 20 weeks and 34 weeks plus 6 days of gestation, using a cut-off value of 100 picograms/ml, was 98% (PELICAN study 2013). It also noted that the negative predictive value for the Elecsys immunoassay sFlt-1/PIGF ratio for the rule-out of pre-eclampsia within 1 week in women presenting with suspected pre-eclampsia between 24 weeks and 36 weeks plus 6 days of gestation, using a cut-off value of 38, was 99% (PROGNOSIS study 2016). The committee heard from clinical experts that PIGF-based tests indicate that the placenta is functioning correctly, which gives them confidence to return a woman with a negative PIGF-based test result (Triage PIGF test result of 100 picograms/ml or more; Elecsys immunoassay sFlt 1/PIGF ratio of less than 38) to community care. It heard further that all women who return to community care, including the small number of women with false negative PIGF-based test results, would still be monitored, so if symptoms of pre-eclampsia reappeared they would be picked up by community midwives and the women would be referred back to the day unit for further assessment. The committee therefore concluded that the use of PIGF-based tests for the rule-out of pre-eclampsia in women presenting with suspected pre-eclampsia between 20 weeks and 36 weeks plus 6 days of gestation was safe and would help to avoid unnecessary hospital admissions. The committee noted its conclusion that PIGF-based tests would be more clinically useful in women presenting before 35 weeks' gestation (see section 5.5). It concluded further that PIGF-based tests for the rule-out of pre-eclampsia should be used for women presenting with suspected pre-eclampsia between 20 weeks and 34 weeks plus 6 days of gestation.

5.8 The committee considered the end point used to determine the accuracy data for the rule-in of pre-eclampsia using the Triage PIGF test. It noted that the external assessment group used the end point of pre-eclampsia needing delivery within 14 days, and considered if an alternative end point could be preterm delivery. The committee heard from clinical experts that the end point of preterm delivery is not equivalent to the end point of pre-eclampsia needing
preterm delivery, because preterm delivery can be for various reasons, not just pre-eclampsia. The committee concluded that the correct end point had been used in the assessment for the rule-in of pre-eclampsia.

5.9 The committee considered the use of PIGF-based tests for the rule-in of pre-eclampsia. The committee noted that the positive predictive value for the Triage PIGF test for the rule-in of pre-eclampsia needing delivery within 14 days in women presenting with suspected pre-eclampsia between 20 weeks and 34 weeks plus 6 days of gestation, using a cut-off value of 12 picograms/ml, was 70% (PELICAN study 2013). It also noted that the positive predictive value for the Elecsys immunoassay sFlt-1/PIGF ratio for the rule-in of pre-eclampsia within 4 weeks in women presenting with suspected pre-eclampsia between 24 weeks and 36 weeks plus 6 days of gestation, using a cut-off value of 38, was 39% (PROGNOSIS study 2016). It heard from clinical experts on the committee that although the PIGF-based tests are helpful for ruling-in pre-eclampsia because they indicate the presence of placental disease, they do not help with the decision on when to deliver the baby. The decision on when to deliver is based on clinical symptoms that indicate risk to the mother or baby, rather than the presence of pre-eclampsia alone. The committee was concerned that in women with suspected pre-eclampsia and a positive PIGF-based test result (Triage PIGF test result of 12 picograms/ml or less; Elecsys immunoassay sFlt 1/PIGF ratio of greater than 38), a decision may be made to deliver the baby sooner on the basis of the PIGF-based test result alone, rather than on clinical symptoms indicating risk to the mother or baby. The committee considered the unpublished PreOS study of the Elecsys immunoassay sFlt-1/PIGF ratio in women with suspected pre-eclampsia. The committee noted that the study was done in Germany and Austria, and heard from a clinical expert on the committee that although the care pathways on admission and surveillance are likely to be slightly different to those in the UK, the results are probably generalisable to clinical practice in the NHS. The results of this study are academic in confidence at the time of writing this diagnostics guidance. The committee concluded that the use of PIGF-based tests to diagnose (rule-in) pre-eclampsia in women presenting with suspected pre-eclampsia between 20 weeks and 36 weeks plus 6 days of gestation could lead to more unnecessary medical intervention resulting in a greater number of premature babies being delivered. The committee noted its conclusion that PIGF-based tests would be more clinically useful in women presenting before 35 weeks' gestation (see section 5.5). It concluded that further research on PIGF-based tests to diagnose (rule-in)
5.10 The committee considered how the results of PIGF-based tests would be used in clinical practice. It was concerned that although PIGF-based tests could be used to safely rule-out pre-eclampsia in women presenting with suspected pre-eclampsia (see section 5.7), the interpretation of positive test results to rule-in pre-eclampsia is more difficult. The committee was concerned that if a PIGF-based test result were available and the result were positive (Triage PIGF test result of 12 picograms/ml or less; Elecsys immunoassay sFlt 1/PIGF ratio of greater than 38) to rule-in pre-eclampsia, too much emphasis might be placed on this result, and not enough emphasis on clinical assessment, which could result in the unnecessary early delivery of the baby (see section 5.9). The committee heard from clinical experts that low levels of PIGF probably indicate placental disease and this is normally managed by close clinical surveillance of the woman and the baby. The committee emphasised the importance of clarity of laboratory reporting on PIGF-based tests, which should include the relevant negative predictive values of the tests and explain to clinicians that a positive test result (Triage PIGF test result of 12 picograms/ml or less; Elecsys immunoassay sFlt 1/PIGF ratio of greater than 38) when using PIGF testing to rule-out pre-eclampsia, does not mean that pre-eclampsia should be ruled-in. The committee concluded that careful laboratory reporting of PIGF-based test results combined with targeted medical education for midwives, obstetricians and laboratory staff is important and should prevent diagnostic drift, in which the results of a test designed to make a rule-out decision start to be used to make a rule-in decision. The committee concluded that PIGF-based testing could be used for the rule-out of pre-eclampsia in NHS clinical practice but could not be used to rule-in pre-eclampsia because further evidence is needed.

5.11 The committee considered the results of the economic analysis and noted that the model included short-term outcomes. It also noted that there was very little difference in quality-adjusted life years (QALYs) for PIGF-based test strategies compared with standard clinical assessment in women presenting with suspected pre-eclampsia between 20 weeks and 34 weeks plus 6 days of gestation. Further, there was no difference in QALYs for PIGF-based strategies compared with standard clinical assessment in women presenting with suspected pre-eclampsia between 35 weeks and 36 weeks plus 6 days of gestation. The committee concluded that PIGF-based testing strategies could
have comparable short-term clinical benefit to standard clinical assessment, and cost-saving benefits (see section 5.13).

5.12 The committee considered long-term outcomes after premature birth. It heard from a clinical expert on the committee that babies born very prematurely are at higher risk of long-term adverse outcomes, such as cerebral palsy. It heard further that these adverse outcomes are very rare, but would have a significant impact on the quality of life of both the child and the parents. Clinicians would take this into consideration when making decisions on when to deliver a fetus in a woman with pre-eclampsia at less than 34 weeks’ gestation. The committee noted that long-term adverse outcomes for the baby were not included in the economic model and concluded that the QALYs in the economic model were therefore likely to have been underestimated. However, the committee concluded further that any underestimation was likely to be small because of the rarity of these adverse events.

5.13 The committee considered the cost savings in the economic analysis. It heard from the external assessment group that the high negative predictive value of the PIGF-based tests to rule-out pre-eclampsia in women with suspected pre-eclampsia was driving the cost savings. This happens because using the PIGF-based tests to assess pre-eclampsia results in fewer false positive results compared with standard clinical assessment, so fewer women are admitted to hospital unnecessarily. Immediate delivery of the fetus is also reduced, which results in fewer premature babies needing time in a neonatal intensive care unit. The committee noted that most of the cost savings came from using the PIGF-based tests to rule-out pre-eclampsia, through monitoring women in a community care setting rather than admitting them to hospital unnecessarily. The committee concluded that these cost savings are plausible if the PIGF-based test results are implemented correctly in clinical practice.

5.14 The committee considered the difference in cost savings between the different gestational age groups. It noted that for women presenting with suspected pre-eclampsia between 20 weeks and 34 weeks plus 6 days of gestation, cost savings of PIGF-based test strategies compared with standard clinical assessment ranged between £2,488 and £2,896. However, for women presenting with suspected pre-eclampsia between 35 weeks and 36 weeks plus 6 days of gestation, cost savings of PIGF-based test strategies compared with standard clinical assessment ranged between £174 and £365. The committee
further noted its conclusion on the clinical utility of PlGF-based tests at different gestation ages (see section 5.5). The committee concluded that the tests show most promise in those women presenting with suspected pre-eclampsia between 20 weeks and 34 weeks plus 6 days of gestation because, on the basis of limited evidence, both clinical utility and cost savings appear likely to be greater in this group compared with women presenting with suspected pre-eclampsia between 35 weeks and 36 weeks plus 6 days of gestation.

5.15 The committee considered whether repeat testing would be done in clinical practice. It noted that the economic evaluation did not include repeat testing, and that no diagnostic accuracy data were available on repeat testing. However, the economic evaluation did include sensitivity analyses doubling and tripling the cost of the test. Results showed that PlGF-based testing remained cost saving compared with standard clinical assessment. The committee concluded that the costs of PlGF-based tests are not prohibitive to repeat testing, but the effect on clinical outcomes is unknown. The committee heard from a clinical expert that repeat testing is often not indicated because a woman who presented with suspected pre-eclampsia, but had a negative PlGF-based test result (Triage PlGF test result of 100 picograms/ml or more; Elecsys immunoassay sFlt-1/PIGF ratio of less than 38), may have no symptoms of pre-eclampsia a week later based on standard clinical assessment, and therefore would not need a PlGF-based test. In other cases, repeat testing is indicated and would normally be done 2 weeks later unless a woman presents again with suspected pre-eclampsia before 2 weeks after the previous test. The committee concluded that research on when repeat testing should be done, and the diagnostic accuracy of repeat testing would be useful.

5.16 The committee considered whether PlGF-based testing could replace the need for quantitative proteinuria testing in the assessment of pre-eclampsia. It noted that the assessment did not identify any evidence on this question and an exploratory cost analysis suggested that cost savings may only increase slightly if PlGF-based testing replaced quantitative proteinuria compared with using both tests together. The committee heard from clinical experts that PlGF-based testing could potentially replace quantitative proteinuria testing in the future.

5.17 The committee considered the advantages and disadvantages of near-patient testing compared with laboratory-based testing. It noted that laboratory-based
testing was used in the base-case economic evaluation, and a scenario cost analysis was done on near-patient testing. The committee heard from a clinical expert that the advantages of near-patient testing include an instant result to inform patient care, which may result in a reduction in waiting times for patients and a reduction in unnecessary admittance to hospital while waiting for a result. It also heard that the disadvantages of near-patient care include: midwife time to do the test which may take them away from patient care; the risk of lack of compliance with the standard operating procedure; the risk of inadequate training in doing the test, and interpreting the results. The committee noted further that an exploratory cost analysis on near-patient testing suggested there may be a slight increase in costs savings compared with laboratory-based testing. The committee concluded that each hospital should make their own decision on whether to implement laboratory-based testing or near-patient testing.

Research considerations

5.18 The committee considered the ongoing and planned research on PIGF-based tests. It heard from an expert on the committee that PIGF-based testing is being used with data collection at Liverpool Women's NHS Foundation Trust and that these data may be available soon. It heard further that a small study was due to start in January 2016: Placental growth factor to assess and diagnose hypertensive pregnant women (PARROT). This is a study of PIGF-based testing for the assessment of suspected pre-eclampsia, which will be done across 6 centres in England and is expected to report in 2018. The aim of the study is to evaluate the implementation of PIGF-based tests by investigating whether PIGF measurement in women presenting with suspected pre-eclampsia between 20 weeks and 37 weeks' gestation decreases the time to a diagnosis of pre-eclampsia. The study will also collect data on clinical outcomes, but is not powered to show a significant difference in outcomes. The committee also noted that the INSPIRE study, a randomised controlled study, is currently running at the Oxford University Hospitals NHS Foundation Trust. This study aims to determine whether the Elecsys immunoassay sFlt-1/PIGF ratio will reduce unnecessary hospital admissions and monitoring for patients with suspected pre-eclampsia by ruling-out pre-eclampsia within 1 week. The committee also heard about the ASPRE study, which includes testing with the DELFIA Xpress PIGF 1-2-3 test, and noted that this study is a screening study to identify women at high risk of pre-eclampsia, rather than to help diagnose
pre-eclampsia in women presenting with suspected pre-eclampsia. The committee concluded that women with suspected pre-eclampsia who are eligible to enrol in the PARROT and INSPIRE studies should be encouraged to do so. The committee concluded further that it is difficult to carry out research into adverse outcomes arising from early delivery because the outcomes are rare and therefore a study would need to be very large.
Recommendations for further research

6.1 Further research is recommended on the use of repeat PIGF-based testing (Triage PIGF test and Elecsys immunoassay sFlt-1/PIGF ratio), with standard clinical assessment, in women presenting with suspected pre-eclampsia between 20 weeks and 34 weeks plus 6 days of gestation, who have had a negative PIGF-based test result (Triage PIGF test result of 100 picograms/ml or more; Elecsys immunoassay sFlt 1/PIGF ratio of less than 38) that was used to rule-out pre-eclampsia (see section 5.15). This should include:

- exploration of the different scenarios in which repeat testing may be indicated
- the appropriate intervals between PIGF-based tests
- the diagnostic accuracy of PIGF-based testing in women with suspected pre-eclampsia who have previously had 1 or more negative PIGF-based test results.

6.2 Further research is recommended on the use of the Triage PIGF test and Elecsys immunoassay sFlt-1/PIGF ratio, with standard clinical assessment, to rule-in pre-eclampsia in women presenting with suspected pre-eclampsia between 20 weeks and 34 weeks plus 6 days of gestation (see section 5.9). This should specifically investigate how a positive PIGF-based test result (Triage PIGF test result of 12 picograms/ml or less; Elecsys immunoassay sFlt 1/PIGF ratio of greater than 38) used to rule-in pre-eclampsia would affect management decisions on time to delivery and the outcomes associated with this.
7 Implementation

NICE has developed tools, in association with relevant stakeholders, to help organisations put this guidance into practice.

- Adoption support resource
- Resource impact report
- Resource impact template

NICE will also support this guidance through a range of activities to promote the recommendations for further research. The research proposed will be considered by the NICE Medical Technologies Evaluation Programme research facilitation team for the development of specific research study protocols as appropriate. NICE will also incorporate the research recommendations in section 6 into its guidance research recommendations database (available on the NICE website) and highlight these recommendations to public research bodies.
8 Review

NICE updates the literature search at least every 3 years to ensure that relevant new evidence is identified. NICE will contact product sponsors and other stakeholders about issues that may affect the value of the diagnostic technology. NICE may review and update the guidance at any time if significant new evidence becomes available.

Andrew Dillon
Chief Executive
May 2016
Diagnostics advisory committee members and NICE project team

Diagnostics advisory committee

The diagnostics advisory committee is an independent committee consisting of 22 standing members and additional specialist members. A list of the committee members who participated in this assessment appears below.

Standing committee members

Professor Adrian Newland
Chair, diagnostics advisory committee and Professor of Haematology, Barts Health NHS Trust

Dr Mark Kroese
Vice Chair, diagnostics advisory committee and Consultant in Public Health Medicine, PHG Foundation, Cambridge and UK Genetic Testing Network

Professor Ron Akehurst
Professor in Health Economics, School of Health and Related Research (ScHARR), University of Sheffield

Dr Phil Chambers
Research Fellow, Leeds Institute of Cancer and Pathology, University of Leeds

Dr Sue Crawford
GP Principal, Ch illington Health Centre

Professor Erika Denton
National Clinical Director for Diagnostics, NHS England, Honorary Professor of Radiology, University of East Anglia and Norfolk and Norwich University Hospital

Dr Steve Edwards
Head of Health Technology Assessment, BMJ Evidence Centre

Mr David Evans
Lay member
Dr Simon Fleming  
Consultant in Clinical Biochemistry and Metabolic Medicine, Royal Cornwall Hospital

Mr John Hitchman  
Lay member

Professor Chris Hyde  
Professor of Public Health and Clinical Epidemiology, Peninsula Technology Assessment Group (PenTAG)

Mr Matthew Lowry  
Director of Finance and Infrastructure, Doncaster and Bassetlaw Hospitals NHS Foundation Trust

Dr Michael Messenger  
Deputy Director and Scientific Manager National Institute for Health Research Diagnostic Evidence Co-operative, Leeds

Dr Peter Naylor  
GP, Chair Wirral Health Commissioning Consortia

Dr Dermot Neely  
Consultant in Clinical Biochemistry and Metabolic Medicine, Newcastle upon Tyne NHS Trust

Ms Gail Norbury  
Consultant Clinical Scientist, Guy's Hospital

Dr Deirdre Ryan  
Consultant Cellular Pathologist, Royal London Hospital

Dr Steve Thomas  
Consultant Vascular and Cardiac Radiologist, Sheffield Teaching Hospitals Foundation Trust

Mr Paul Weinberger  
Chief Executive Officer, DiaSolve Ltd, London

Professor Anthony Wierzbicki  
Consultant in Metabolic Medicine and Chemical Pathology, St Thomas' Hospital

PIGF-based testing to help diagnose suspected pre-eclampsia (Triage PIGF test, Elecsys immunoassay sFlt-1/PIGF ratio, DELFIA Xpress PIGF 1-2-3 test, and BRAHMS sFlt-1 Kryptor/BRAHMS PIGF plus Kryptor PE ratio) (DG23)
PIGF-based testing to help diagnose suspected pre-eclampsia (Triage PIGF test, Elecsys immunoassay sFlt-1/PIGF ratio, DELFIA Xpress PIGF 1-2-3 test, and BRAHMS sFlt-1 Kryptor/BRAHMS PIGF plus Kryptor PE ratio) (DG23)

Specialist committee members

Mrs Ann Marie Barnard
Lay member

Dr Jenny Myers
Senior Lecturer and Consultant Obstetrician, Maternal and Fetal Health Research Centre, Central Manchester Foundation Trust

Mr Nigel Simpson
Senior Lecturer in Obstetrics and Gynaecology, and Honorary Consultant, Division of Women and Children's Health, University of Leeds

Professor Jimmy Walker
Professor of Obstetrics and Gynaecology, St James's University Hospital Trust

Mr David Wells
Pathology General Manager, Viapath, St Thomas' Hospital

NICE project team

Each diagnostics assessment is assigned to a team consisting of a Technical Analyst (who acts as the topic lead), a Technical Adviser and a Project Manager.

Frances Nixon
Topic Lead

Sarah Byron
Technical Adviser

Robert Fernley
Project Manager
10 Sources of evidence considered by the committee

The diagnostics assessment report was prepared by Southampton Health Technology Assessments Centre.


Registered stakeholders

The following organisations accepted the invitation to participate in this assessment as registered stakeholders. They were invited to attend the scoping workshop and to comment on the diagnostics assessment report and the diagnostics consultation documents.

Companies/sponsors:

- Alere International Ltd
- Perkin Elmer LAS UK
- Roche Diagnostics Ltd
- Siemens
- Thermo Fisher Scientific

Professional/specialist and patient/carer groups:

- Action on Pre-eclampsia (APEC)
- Birth Trauma Association
- British Maternal and Fetal Medicine Society
- Institute of Biomedical Science
- National Childbirth Trust (NCT)
- Royal College of Nursing
- Royal College of Physicians
PIGF-based testing to help diagnose suspected pre-eclampsia (Triage PIGF test, Elecsys immunoassay sFlt-1/PIGF ratio, DELFIA Xpress PIGF 1-2-3 test, and BRAHMS sFlt-1 Kryptor/BRAHMS PIGF plus Kryptor PE ratio) (DG23)

- Sands (Stillbirth and neonatal death charity)
- The Multiple Births Foundation
- Royal College of Pathologists

Others:

- Department of Health
- Healthcare Improvement Scotland
- NHS England
- Welsh Government
About this guidance

NICE diagnostics technologies guidance is designed to help the NHS adopt efficient and cost-effective medical diagnostic technologies more rapidly and consistently.

The programme concentrates on pathological tests, imaging, endoscopy and physiological measurement, since these represent most of the investigations performed on patients. The types of products that might be included are medical diagnostic technologies that give greater independence to patients, and diagnostic devices or tests used to detect or monitor medical conditions. Diagnostic technologies may be used for various purposes: diagnosis, clinical monitoring, screening, treatment triage, assessing stages of disease progression, and risk stratification.

This guidance was developed using the NICE diagnostic technologies guidance process.

We have produced a summary for patients and carers. Tools to help you put the guidance into practice and information about the evidence it is based on are also available.

Your responsibility

This guidance represents the view of NICE, which was arrived at after careful consideration of the evidence available. Healthcare professionals are expected to take it fully into account when exercising their clinical judgement. However, the guidance does not override the individual responsibility of healthcare professionals to make decisions appropriate to the circumstances of the individual patient, in consultation with the patient and/or guardian or carer.

Implementation of this guidance is the responsibility of local commissioners and/or providers. Commissioners and providers are reminded that it is their responsibility to implement the guidance, in their local context, in light of their duties to have due regard to the need to eliminate unlawful discrimination, advance equality of opportunity, and foster good relations. Nothing in this guidance should be interpreted in a way which would be inconsistent with compliance with those duties.

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