Utero-placental vascularisation in normal and preeclamptic and intra-uterine growth restriction pregnancies: third trimester quantification using 3D power Doppler with comparison to placental vascular morphology (EVUPA): a prospective controlled study

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ABSTRACT

Introduction: Preeclampsia (PE) and intra-uterine growth restriction (IUGR) are two major pregnancy complications related to chronic utero-placental hypoperfusion. Three-dimensional power Doppler (3DPD) angiography has been used for the evaluation of utero-placental vascularisation and three vascular indices have been calculated: the vascularisation index (VI), flow index (FI) and vascularisation-FI (VFI). However, several technical endpoints hinder the clinical use of 3DPD as physical characteristics and machine settings may affect 3DPD indices, and so its clinical significance is not yet clear.

Objectives: The primary objective is to better understand the clinical significance of 3DPD indices by evaluating the relationship between these indices and morphological indices. Secondary objectives are (i) to determine the impact of machine settings and physical characteristics on 3DPD indices, and (ii) to evaluate physio-pathological placental vascularisation patterns.

Methods and analysis: This is a prospective controlled study. We expect to include 112 women: 84 with normal pregnancies and 28 with PE and/or IUGR (based on our former cohort study on 3DPD indices for PE and/or IUGR prediction (unpublished data)). Within 72 h before planned or semi-urgent caesarean section, utero-placental 3DPD images with five different machine settings will be acquired. Placentas will be collected and examined after surgery and stereological indices (volume density, surface density, length density) will be calculated. The 3DPD indices (VI, FI and VFI) of the placenta and adjacent myometrium will be calculated. Correlation between Doppler and morphological indices will be evaluated by Pearson or Spearman tests. Agreement between 3DPD indices and morphological indices will be assessed by Bland and Altman plots. The impact of Doppler settings and maternal characteristics on 3DPD indices will be evaluated with a multivariate linear regression model.

Ethics: The study and related consent forms have been approved by the French Ethics Committee (CPP, Comité de Protection des Personnes) Est III on 4 March 2014.
Major advances in the understanding of utero-placental physiological were made by Ramsey and Donner during the 20th century. The changes due to pregnancy in the uterine spiral arteries and blood flow in the intervillous space are now well characterised. Evaluation of utero-placental vascular modification during pregnancy using non-invasive methods such as ultrasound recently became possible. After implantation, trophoblast cells induce vascular remodelling, which begins first in the endometrium and then in adjacent myometrium, and can be detected by Doppler imaging. A rich and dense vascular network develops inside the myometrium under the placental basal plate, which coincides with changes in the terminal parts of the spiral arteries induced by invasion by extra-villous trophoblast cells. A functional anatomical arteriovenous shunt was also identified in the sub-placental myometrium from the first trimester until the end of pregnancy, and plays a role in gas exchange.

Unlike the traditional ‘in series’ vascular communication model, the intervillous space is connected ‘in parallel’ with the uterine circulation. This type of circulation would offset the maternal haemodynamic changes to protect placental villi, and may also provide a dissolved oxygen reservoir to feed the intervillous space when blood flow is temporarily reduced or modified. Therefore, the development of this rich anatomical vascular network in the myometrium is a crucial step for normal pregnancy development and fetal growth.

Nowadays, the available in vivo methods for the assessment of organ vascularisation include ultrasound with or without contrast agent injection, CT scanning with contrast agent injection, MR angiography with contrast agent injection and functional MRI. However, in pregnant women, traditional radioactive examination or the use of contrast agents is either forbidden or discouraged. Consequently, ultrasound is recommended for non-invasive, in vivo assessment of placental vascularisation.

Until recently, the most promising method to screen for PE/IUGR was uterine artery Doppler velocimetry by 2D pulsed Doppler. This approach, however, has some major disadvantages: the blood flowing perpendicular to the axis of the ultrasonic beam cannot be studied, and sensitivity is too poor for the study of slow flows. Furthermore, movement (organ movement and adherent tissue movement) reduces the accuracy and reproducibility of 2D pulsed Doppler acquisition, especially for small vessels and slow flows.

Published data suggest that impaired placental perfusion, which is associated with development of PE/IUGR, could be reflected in an increased uterine artery pulsatility index (PI). Measurement of the uterine artery PI is influenced by sonographer experience and by gestational age, maternal weight, racial origin and pre-existing diabetes mellitus. Using the uterine artery PI, the detection rate of PE before 34 gestational weeks is estimated to be 59% with a 5% false positive rate or 75% with a 10% false positive rate. The clinical value of spectral analysis of the uterine artery is still uncertain. Besides, the uterine artery, as the main afferent vessel of the utero-placental unit, only partially reflects the true haemodynamics of the placenta. Therefore, non-invasive and direct quantification of intervillous space perfusion might provide more valuable information about placental function.

Since 2004, it has been possible to quantify placental and myometrium vascularisation by 3D power Doppler angiography (3DPD). This method allows the vascularisation of an organ of interest to be studied and quantified in a non-invasive manner and without the use of contrast agent. Quantification is based on calculation of the ratios of voxels with Doppler signals to the intensity of Doppler signals in the voxels. It provides a new approach to quantify the vascular signals from an organ of interest. Three typical indices of a volume of interest were calculated by this method: the vascularisation index (VI), flow index (FI) and vascularisation-FI (VFI).

The feasibility and reproducibility of Doppler signal quantification by calculating VI, FI and VFI were found to be satisfactory in vitro and in vivo. In a previous study by our group, a good correlation between measured 3DPD indices and actual blood flow has also been shown in vivo in a sheep model. A direct relationship between the number of fetal capillary vessels per villus during the first trimester and placental vascular indices (VI, FI and VFI) has been found in first trimester chorionic villus samples.

A significant decrease in placental VI, FI and VFI was observed in preeclamptic patients during the first trimester as well as the second and third trimesters. The decrease in 3D placental indices appeared before anomalies in uterine artery PI were seen. The 3D placental indices seem to be superior for predicting PE compared with the PI of the uterine artery on 2D Doppler at the end of the first trimester. In a larger population (4325 pregnant women), the myometrial VI during the first trimester had the best performance of all types of PE prediction compared with uterine artery PI and maternal biomarkers. The 3D Doppler measurement has been shown its value and superiority for the detection of maternal hypertensive disorders.

In a nitric oxide treated IUGR rabbit model, the placental indices (VI, FI and VFI) were significantly reduced at 28 days of gestation and accompanied important placental morphological alterations. In humans, a reduction in placental VI and VFI was always observed in IUGR during the second and third trimesters. During the first trimester, all placental 3DPD indices in the case of IUGR were similar to those measured in normal pregnancies in three studies. However, a positive correlation was confirmed between 3DPD indices and IUGR severity.

Although 3DPD appears to be a very promising clinical approach for pathological placental perfusion assessment, several technical endpoints still need to be evaluated for optimal use. The main problem with this technique is that the calculated 3DPD indices are...
dependent on different technical and physiological parameters, such as flow rate, distance between the organ of interest and the probe, machine settings, age of the ultrasound system and maternal characteristics.15 29 To reduce the influence of personal characteristics on the 3DPD indices (including body mass index (BMI), and position and depth of the placenta), a new approach was described, known as ‘sub-noise gain’ (SNG), whereby gain is adjusted to the maximum value for examination without artefacts.30 The inter- and intra-observer intraclass gain is adjusted to the maximum value for examination described, known as 3DPD indices (including body mass index (BMI), and reduce the in

The main objective of our study is to better understand the clinical signification of 3DPD indices by evaluating the relationship between the indices and placental morphometry in normal and impaired pregnancies.

Secondary objectives

▸ To evaluate the impact of different machine settings (including the SNG setting) on 3DPD indices and the relationship between the 3DPD indices and placental morphometry. This should permit assessment of Doppler sensitivity to different vessels and allow future setting to be optimised for clinical use.

▸ To concomitantly assess in vivo and in vitro normal and pathological utero-placental vascularisation.

METHODS

Trial design

The Etude de la Vascularisation UtéroPlacentaire par echoAngiographie (EVUPA) protocol is a monocentric (Maternité Régionale du CHRU de Nancy, France), prospective controlled study.

Study population

To avoid the effect of uterine contractions during delivery on Doppler signals and ensure the integrity of the placenta, pregnant women with a planned or semi-urgent caesarean section will be recruited into this study. Included women will be divided into two groups (normal and pathological) according to current pregnancy issues. Caesarean sections for normal pregnancies are based on national health authority (HAS, Haute Autorité de Santé) recommendations.32 Women with normal pregnancies will be hospitalised on the day before the programmed caesarean section in the maternity unit according to routine protocol. In pathological pregnancies, a caesarean section might be scheduled in semi-urgent situations after benefit-risk evaluation.

The pathological group will include patients with small for gestational age (SGA) fetuses or IUGR and/or PE diagnosed based on pregnancy monitoring and tests. PE is identified as systolic blood pressure (SBP) ≥140 mm Hg and/or diastolic blood pressure (DBP) ≥90 mm Hg associated with proteinuria >300 mg/24 h after 20 weeks of gestation. Early-onset PE (<34 weeks’ of gestation) and late-onset PE (≥34 weeks’ of gestation) will be identified according to gestational age. IUGR is identified as fetal weight <10th percentile as estimated by ultrasound and based on the personal fetal growth estimation curve (AUDIPOG, France) with an umbilical artery PI >95th percentile, in the absence of malformation, infection or a chromosomal defect. SGA is identified as a birth weight <10th percentile based on the personal fetal growth estimation curve, in the absence of malformation, infection or a chromosomal defect.

The inclusion and exclusion criteria are reported in table 1. All eligible women will receive an information sheet about this study and explanations by investigators. The informed consent form will be signed and collected if the patient is willing to participate. Patients’ medical history including BMI, smoking history, family history, and gynaecological and obstetric history will be collected. Medical information on the current pregnancy will also be collected.

Doppler acquisition

Doppler images will be acquired for all participating women within 72 h before their caesarean section using a Voluson E8 system and 4–8 MHz probe (General Electric Healthcare). All images will be identified by subject code following the same coding rules (initials

Table 1: Study inclusion and exclusion criteria

<table>
<thead>
<tr>
<th>Inclusion criteria</th>
<th>Exclusion criteria</th>
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<tbody>
<tr>
<td>Age ≥18 years and &lt;65 years</td>
<td>Fetal anomalies</td>
</tr>
<tr>
<td>Gestational age ≥30 gestational weeks</td>
<td>Maternal or fetal vital urgency</td>
</tr>
<tr>
<td>and &lt;42 gestational weeks</td>
<td>Multiple pregnancy</td>
</tr>
<tr>
<td>With social insurance</td>
<td>Non-placental origin IUGR</td>
</tr>
<tr>
<td>With written consent for participation</td>
<td>Language barrier</td>
</tr>
<tr>
<td>With medical examination</td>
<td>Patient under legal protection</td>
</tr>
<tr>
<td>Hospitalised pregnant women with</td>
<td></td>
</tr>
<tr>
<td>scheduled or semi-urgent</td>
<td></td>
</tr>
<tr>
<td>caesarean section according to national HAS recommendations or for pathological management (PE and/or IUGR)</td>
<td></td>
</tr>
<tr>
<td>Normal pregnancy issues or with PE and/or IUGR (control or pathological group)</td>
<td></td>
</tr>
<tr>
<td>Normal fetal morphology</td>
<td></td>
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</table>

| HAS, Haute Autorité de Santé; IUGR, intra-uterine growth restriction; PE, preeclampsia. | |
followed by inclusion order). 3DPD acquisition of utero-placental vascularisation will be carried out first. Basic 3D acquisition settings are pre-programmed in the machine. As it is difficult to acquire all placental volumes during the third trimester, we will use the widest 3D scan angle to obtain as much of the placental volume as possible. Placental position will be noted. To evaluate the impact of different machine settings on 3DPD indices, five settings will be used with modification each time of pulse repetition frequency (PRF), wall motion filter (WMF) and gain (table 2). A SNG setting will be used as recommended in the literature. This setting is obtained by increasing the gain until artefactual noise is present, and then slowly reducing it until the noise artefact has just disappeared. To evaluate the effect of attenuation on 3DPD indices, the patient’s BMI and placental location will be recorded.

2D colour Doppler indices of fetal and maternal major vessels (such as the PI and the resistive index of the umbilical artery, middle cerebral artery, ductus venosus and uterine arteries) will be acquired afterwards following ISUOG guidelines. JD and A-CC-L have been trained on the ultrasound system and will perform all Doppler acquisitions.

Image analysis
All Doppler images will be transferred and stored in CIC-IT (Centre d’Investigation Clinique—Innovation Technologique), Nancy. Each 3D utero-placental volume will be analysed using VOCAL software. The placenta and its entire adjacent myometrium will be each traced with 30° rotation, and the 3DPD vascular indices (VI, FI and VFI) calculated automatically (figure 1). The 2D colour Doppler waveform parameters of each vessel will be noted.

Placenta collection and analysis
After delivery, the umbilical cord will be immediately clamped and cut, and each placenta will be collected and transferred to the laboratory of fetal and placental pathology for systematic examination. The macroscopic features of each placenta to be noted are: umbilical cord and membrane insertion, placental size and thickness, and putative presence of thrombosis, infarcts, cysts and haemorrhage. The placenta will be weighed after the removal of membranes and the remaining length of umbilical cord. The placenta will then be fully immersed in water. The initial and final water level after placenta immersion will be noted and the volume of placenta calculated as the final water level minus the initial water level.

The entire placenta will be fixed in 4% formaldehyde for about 3 weeks. Placental volume and weight will be measured again after fixation using the same technique as described above. A systematic uniform random sampling (SURS) design will be used and six cubes (≈1.5×1.5×1.5 cm³) containing whole placenta thickness will be sampled for each placenta. Each tissue cube will be dehydrated in step-increased concentrations of ethyl alcohol. Once the tissue block is embedded in paraffin wax, a 6µm vertical uniform random (VUR) section containing the entire thickness of the placenta will be taken and mounted on a Superfrost Plus glass slide for staining with H&E (figure 2).

Images will be taken at different magnifications. Certified pathologists will read slides in order to establish a histopathological diagnosis. Stereological parameters such as volume density (Vv), surface density (Sv) and length density (Lv) will be calculated using STEPanizer software. Structures to be measured are summarised in table 3. Correlation coefficients between 3DPD indices and morphological indices will be estimated by Pearson or Spearman tests as appropriate.

Outcomes
Validation of the 3DPD technique as a reliable indicator of utero-placental vascularisation and a better understanding of utero-placental physio-pathological mechanisms are expected. If a high degree of correlation between 3DPD indices and placental morphometry is confirmed, 3D Doppler angiography could have an important role in future strategies for placental vascular pathology screening and diagnosis. Earlier screening (during the first trimester) for placental vascular pathologies could be established and corresponding medical care instituted, which might reduce maternal and fetal morbidity and mortality. An adapted 3DPD setting for clinics will be proposed at the end of this study.

Participant timeline
The enrolment of women started in February 2014. In view of the caesarean rate in our hospital, recruitment should be achieved by February 2017. The flow chart of patient participation is presented in figure 3.

Premature ending of patient participation
Participants will be excluded from the study in the following situations:

- Lack of 3DPD acquisition
- No placental morphological analysis
- Patient has an undesirable event related to the study
- Withdrawal of consent before the end of the study

<table>
<thead>
<tr>
<th>Table 2 3D power Doppler settings</th>
<th>PRF (KHz)</th>
<th>WMF</th>
<th>Gain (dB)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Setting 0</td>
<td>0.9</td>
<td>Mid 1</td>
<td>−7.2</td>
</tr>
<tr>
<td>Setting 1</td>
<td>1.8</td>
<td>Mid 1</td>
<td>−7.2</td>
</tr>
<tr>
<td>Setting 2</td>
<td>0.9</td>
<td>Max 2</td>
<td>−7.2</td>
</tr>
<tr>
<td>Setting 3</td>
<td>0.9</td>
<td>Low 1</td>
<td>−7.2</td>
</tr>
<tr>
<td>Setting 4</td>
<td>0.9</td>
<td>Mid 1</td>
<td>SNG</td>
</tr>
</tbody>
</table>

SNG was set by increasing the gain until artefactual noise was present then slowly reducing it to where the noise artefact has just disappeared.

PRF, pulse repetition frequency; SNG, sub-noise gain; WMF, wall motion filter.

Patients will be immediately excluded from the study and replaced with other new participants. Any decision to withdraw consent should not affect the patient’s routine medical care. In the case of an undesirable event related to the study, the patient should be informed and excluded. Revised medical care should be offered.

Follow-up
No specific follow-up has been planned for participants except for routine healthcare after caesarean section. Information on newborns will be collected after delivery, such as gender, birth weight, Apgar score, umbilical cord arterial pH, base excess and lactate value, and neonatal intensive care. Any undesirable event after caesarean section such as maternal or neonatal death, transfusion, hysterectomy, hospitalisation in the intensive care unit, and significant post-partum haemorrhage will be noted and declared.

Sample size consideration
As this, as far as we know, is the first study to evaluate concordance between 3DPD indices and placental morphometry, there is no available reference for sample size determination. Therefore, the ability to detect differences in 3DPD indices between normal and pathological pregnancies was used for sample size estimation. Based on our previous cohort study on 3DPD indices for PE and/or IUGR prediction (unpublished data of Morel et al), in order to detect a difference in placental FI (the best index in our study) between normal pregnancy and PE (PE prevalence of 3%), with an $\alpha$ of 5% and a power of 80%, a population of 112 pregnant women (28 with PE and/or IUGR and 84 with normal pregnancies) have to be included. As the main objective of this study is to evaluate the correlation between Doppler and morphometric indices, different pathologies (such as early onset and late onset PE, and IUGR) will be combined during analysis.

Data collection and management
Participant consent will be obtained after detailed explanation to patients during recruitment. An electronic case report form (e-CRF) will then be created for each patient on a secure website dedicated to clinical research (2008 Clinsight). All information related to patient and pregnancy outcomes mentioned above will
be noted. Each subject will be anonymized as initials followed by inclusion order. Each investigator will have an individual account allowing them to fill in the e-CRF.

Confirmation of participation in this study will be noted in patients’ medical records by the investigators. Patients’ identities will be kept confidential. The subject code will be used for all related data (case report form, ultrasound and morphological images and undesirable invent declaration form) using the same coding rules (initials followed by inclusion order). One list only with patients’ identities and corresponding subject codes will be stored in the principle investigator’s file in a secure place, together with patient consent forms.

Data management will be carried out by CIC-IT, Nancy. Queries will be answered in order to solve questions regarding missing data, unreadable text, and coherence. Study monitors will be responsible for obtaining written and signed answers by investigators to all questions. Before the database is frozen, a data review will be conducted in order to identify errors during recruiting and data input. The clinical study coordinator, study monitor and data manager will jointly conduct the data review and approve the list of omitted data.

When the queries process has ended, the data manager will declare database closure so that the dataset can be sent to the statistician. The database will be frozen once all corrections required by the investigators have been implemented and all data reviewed.

All ultrasound and morphological images will be anonymized and archived in the ARCHIMED database, which has been declared to CNIL (National Commission for Data Protection and Liberties; declaration number: 1410005). All data (numeric and paper) will be archived for 15 years after the end of the study.

### Statistical analysis

Statistical analysis will be performed using R software (R Foundation for Statistical Computing, Vienna, Austria). Quantitative results will be expressed as the mean±SD. Correlation coefficients between 3DPD indices and morphological indices will be estimated by Pearson or Spearman tests as appropriate. Agreement between 3DPD indices and morphological indices will be assessed

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**Table 3** Stereological indices

<table>
<thead>
<tr>
<th>Structure</th>
<th>Parameter</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intervillous space</td>
<td>Vv</td>
</tr>
<tr>
<td>Villi</td>
<td>Vv, Sv, Lv</td>
</tr>
<tr>
<td>Capillary</td>
<td>Vv, Sv, Lv</td>
</tr>
<tr>
<td>Stroma of villi</td>
<td>Vv</td>
</tr>
<tr>
<td>Trophoblast cells</td>
<td>Vv</td>
</tr>
<tr>
<td>Trophoblast knots</td>
<td>Vv, Lv</td>
</tr>
</tbody>
</table>

Lv, length density (μm²); Sv, surface density (μm); Vv, volume density.
QUALITY CONTROL
Right of access to data and source documents
The Centre Hospitalier Régional Universitaire (CHRU) of Nancy is responsible for obtaining the agreement of all parties involved in the study so as to guarantee direct access to all study sites, source data, source documents and reports so that the sponsor can control data quality and perform an audit.

Investigators will make available the documents and individual data strictly required for monitoring, quality control and audit of the biomedical study to persons allowed access to these, in accordance with the statutory and regulatory provisions in place (Articles L.1121-3 and R.5121-13 of the French Public Health Code).

Any original document or object that allows the existence or accuracy of a data point or information recorded during the study to be proved is defined as a source document.

In accordance with the statutory provisions in place (Articles L.1121-3 and R.5121-13 of the French Public Health Code), the persons having direct access to source data will take every precaution necessary to ensure the confidentiality of information relating to investigational medicinal products, studies, participants, particularly concerning their identity, as well as the results obtained. These persons, as the investigators themselves, are subject to professional confidentiality.

The CHRU of Nancy will ensure that each participant has given her written consent for access to her personal data that is strictly required for study quality control.

Study monitoring
The method of data monitoring has been established by DRI (Department of Clinical Research and Innovation) of CHRU, Nancy. A clinical research associate (CRA) will regularly visit the Nancy regional university maternity unit of the CHRU for the quality control of reported data entered in the e-CRF. The CRA will verify that the study is being conducted according to protocol and ensure that the data form contains all requested information.

During monitoring, the source documents of each patient should be provided (medical files) and must be consistent with the e-CRF. The CRA will have access to the e-CRF, medical files and all other related documents. The CRA is bound by a duty of confidentiality regarding the reviewed patients’ information.

The risk of the study has been classified as category B by the CHRU of Nancy according to the GT5 work group of the national assembly of the Delegation of Clinical Research and Innovation (DRCI), which is validated by the primary healthcare department (Direction Générale de l’Offre de Soins, DGOS). A proper monitoring plan including verification of partial patient’ consent forms, inclusion and exclusion criteria, data entry and security, principle endpoint, and undesirable event reports has been established.

POTENTIAL RISKS RELATED TO THE STUDY
The research team does not foresee any special medical risks for participating women or their unborn children. This study follows good clinical practice as defined by the French ministry of health. The only difference related to the study is the addition of utero-placental 3DPD acquisition. The duration of this acquisition does not exceed 5 min. The ultrasound machine is EC (European Community) marked and routinely used in clinic service. Patients with a maternal or fetal vital emergency will not be included in this study.

ETHICAL PERMIT
The CHRU of Nancy and investigators commit that this research will be conducted in accordance with legislative regulation no. 2004–806 of 9 August 2004, as well as in agreement with Good Clinical Practices (ICH V.4 of 1 May 1996 and Decision of 24 November 2006) and the Helsinki Declaration (Ethical Principles for Medical Research Involving Human Subjects, Tokyo 2004). In order to initiate the research, the CHRU of Nancy as its sponsor has submitted an authorisation request to the competent authority, ANSM (French National Agency for Medicines and Health Products Safety). The competent authority, as defined in Article L. 1123-12, has provided its decision with regard to the safety of individuals who consent to biomedical research, by especially taking into consideration the safety and quality of products used during the research in accordance with, where appropriate, existing repositories, their condition of use and the safety of persons with regard to acts performed and the methods used as well as intended procedures for patient follow-up. Approval of ANSM was obtained on 14 February 2014 and the reference number of the study is 203-A01049-36.

In accordance with Article L. 1123-6 of the Public Health Code, the research protocol has been submitted by the sponsor to the Committee for the Protection of Persons (CPP, Comité de Protection des Personnes). The study and related consent forms have been approved by CPP Est III on 4 March 2014 and the CPP reference number of this study is 13.09.02.

PROTOCOL AMENDMENT
A substantial change is a change that is liable, in one way or another, to modify the assurances made to
participants who consent to the study (modification of an inclusion criterion, extending the inclusion period, participation of new centres, etc).

Once the research has begun, any substantial modification thereof on the initiative of the sponsor must obtain, prior to its implementation, approval by the committee and authorisation from the competent authority. In this case, if necessary, the committee ensures that a new consent form is obtained from individuals participating in research.

Any substantial change requires authorisation from ANSM and CPP in accordance with legislative regulation no. 2004–806 of 9 August 2004.

FINAL RESEARCH REPORT

The coordinator and the mandated biostatistician will collaboratively write the final research report. This report will be submitted to each of the investigators for review. Once consensus has been reached, the final version must be endorsed with the signature of each of the investigators and sent to the sponsor as early as possible after the effective end of the research. A report prepared according to the reference plan of the competent authority must be forwarded to the competent authority and the CPP within a year after the end of the research.

DISCUSSION

This study aims to evaluate concordance between 3DPD indices and placental morphology. The reliability of 3DPD for the evaluation of utero-placental vascularisation will be established if a high correlation between 3DPD indices and morphological indices is found. Stereological analysis, which can provide three-dimensional unbiased and quantitative data from two-dimensional cross-sections, has been widely used in placental morphological analysis. When defining our 3DPD indices, we found similarities with stereological indices such as volume density, surface density and length density. Therefore, we will use stereology for our primary morphological analysis for relationship assessment; traditional morphological analysis will also be used for further physio-pathological interpretation. As far as we know, this is the first study to evaluate the relationship between utero-placental 3DPD and morphology in pregnant women.

Furthermore, the clinical use of 3DPD will also be assessed in this study. The impact of different Doppler settings, including the new SNG setting, on 3DPD indices in women with normal and pathological pregnancies will be evaluated. To our knowledge, such evaluations have only been carried out in phantoms and animal models. The homogeneity of 3DPD indices in a specific population, as well as the relationship between 3DPD and morphology with one particular setting, will be considered regarding future suggestions for clinical settings. We anticipate that our study data will inform recommendations about machine settings to provide the maximum information about utero-placental physiology and pathophysiology while minimising ultrasound exposure for the pregnant women and elucidate the impact of maternal physical characteristics such as BMI and placental position on Doppler signals.

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Contributors JD participated in study design and is carrying out or will carry out recruitment, Doppler acquisition, placental examination, data analysis and manuscript writing. A-CC-L is carrying out or will carry out recruitment, Doppler acquisition and data analysis, EP-G and OM are major investigators for clinical assessment, Doppler acquisition and study design. CC is in charge of placental morphological analysis. AC is project manager. GH is in charge of statistical analysis. All authors read and approved the final manuscript.

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Competing interests None declared.

Patient consent Obtained.

Ethics approval ANSM (the French National Agency for Medicines and Health Products Safety) and the Committee for the Protection of Persons (Comité de Protection des Personnes, CPP) approved this study.

Data sharing statement All data generated during the project will be made freely available via CIC-IT, Nancy. Data obtained from this study will be deposited at CIC-IT Nancy where they will be maintained for a minimum of 15 years. There are no security, licensing or ethical issues related to the expected data, and all data used in the project will be generated directly as a result of the project, without any pre-existing data being used.

Trial status This is an ongoing trial. Recruitment began in February 2014. We expect to complete recruitment by February 2017. We plan to publish final results in 2017.

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REFERENCES