Early first trimester uteroplacental flow and the progressive disintegration of spiral artery plugs: New insights from contrast-enhanced ultrasound and tissue histopathology.

Running title: First trimester uteroplacental flow

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Abstract

Study question: Does the use of a vascular contrast agent facilitate earlier detection of maternal flow to the placental intervillous space in the first trimester of pregnancy?

Summary answer: Microvascular filling of the intervillous space was demonstrated by contrast-enhanced ultrasound from 6 weeks of gestation onwards.

What is known already: During placental establishment and remodeling of maternal spiral arteries, endovascular trophoblast cells invade and accumulate in the lumen of these vessels to form ‘trophoblast plugs’. Prior evidence from morphological and Doppler ultrasound studies has been conflicting as to whether the spiral arteries are completely plugged, preventing maternal blood flow to the intervillous space until late in the first trimester.

Study design, size, duration: Uteroplacental flow was examined across the first trimester in human subjects given an intravenous infusion of lipid-shelled octofluoropropane microbubbles with ultrasound measurement of destruction and replenishment kinetics. We also performed a comprehensive histopathological correlation using two separately archived uteroplacental tissue collections to evaluate the degree of spiral artery plugging and evaluate remodeling of the upstream myometrial radial and arcurate arteries.

Participants/materials, setting, methods: Pregnant women (n=34) were recruited in the first trimester (range: 6+3 – 13+6 weeks gestation) for contrast-enhanced ultrasound studies with destruction-replenishment analysis of signal intensity for assessment of microvascular flux rate. Histological samples from archived in situ (Boyd Collection, n=11) and fresh first, second, and third trimester decidual and post-hysterectomy uterine specimens (n=16) were evaluated by immunohistochemistry and ultrastructural analysis.
Main results and the role of chance: Contrast agent entry into the IVS was visualized as early as 6+3 weeks of gestation with some variability in microvascular flux rate noted in the 6 to 7+6 week samples. Spiral artery plug canalization was observed from 7 weeks with progressive disintegration thereafter. Of note, microvascular flux rate did not progressively increase until 13 weeks which suggests that resistance to maternal flow in the early placenta may be mediated more proximally by myometrial radial arteries that begin remodeling at the end of the first trimester.

Limitations, reasons for caution: Gestational age was determined by crown rump length measurements obtained by transvaginal ultrasound on the day of contrast-enhanced imaging studies, which may explain the variability in the earliest gestational age samples due to the margin of error in this type of measurement.

Wider implications of the findings: Our comprehensive in situ histological analysis, in combination with the use of an in vivo imaging modality that has the sensitivity to permit visualization of microvascular filling has allowed us to reveal new evidence in support of increasing blood flow to the intervillous space from 6 weeks of gestation. Histologic review suggested the mechanism may be blood flow through capillary sized channels that form through the loosely cohesive ‘plugs’ by 7 weeks gestation. However, spiral artery remodeling on its own did not appear to explain why there is significantly more blood flow at 13 weeks gestation. Histologic studies suggest it may be related to radial artery remodeling, which begins at the end of the first trimester.
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Keywords: Spiral artery, intervillous blood flow, Contrast-enhanced ultrasound; placental perfusion; in vivo imaging, trophoblast plugs
Introduction

Many factors contribute to pregnancy success but arguably the most critical one is the establishment of the uteroplacental vasculature. Without the appropriate interplay between the maternal and fetal circulations, inadequate maternal-fetal exchange will inevitably result in compromised fetal growth and development. Remodeling of the maternal spiral arteries from tightly coiled vessels in the non-pregnant state, to wide-bore, low resistance conduits in pregnancy is fundamental to the establishment of the uteroplacental circulation. This complex conversion involves multiple processes that require the invasion of placental extravillous trophoblasts (EVTs) into the lumen of decidual spiral arteries (For review, (Pijnenborg, et al., 2006, Whitley and Cartwright, 2010)). The EVT accumulate in the lumen and form what has been referred to as ‘trophoblast plugs’. These collections of EVT, or ‘plugs’, are postulated to obstruct maternal arterial blood flow into the intervillous space (IVS) before the end of the first trimester; yet, there is conflicting support for and against this hypothesis (Burton, et al., 1999, Burton, et al., 2002, Carbillon, et al., 2001, Coppens, et al., 1996, Jauniaux, et al., 2001, Meekins, et al., 1997, Valentin, et al., 1996). Moreover, little is known about the breakdown of these ‘plugs’, how this process is initiated, regulated, and the relationship between disintegrating plugs and blood flow.

The primary challenge in addressing these questions is in situ access to the spiral arteries in relation to the placenta. Curettage specimens from miscarriage and elective pregnancy terminations may provide partial snap-shots, but they are limited by confounders such as tissue damage associated with collection and sample processing. In addition, current in vivo assessment of placental perfusion in early pregnancy is hindered by limitations in assessing perfusion through low flow and small diameter vessels like mid-first trimester decidual spiral arteries. In
this study, we sought to overcome these obstacles using two approaches; firstly, using a vascular
contrast agent to visualize vascular filling of the IVS by ultrasound in early pregnancy in a
cohort of women with non-continuing pregnancies, and secondly, by histological examination of
two separately archived uteroplacental tissue resources.

Contrast-enhanced ultrasound (CEUS) relies on the acoustic detection of gas-filled, lipid-encapsulated microbubbles to visualize and quantify microvascular perfusion (Kaufmann and
Lindner, 2007, Kaufmann, et al., 2007). The use of a contrast agent facilitates assessment of
perfusion in small capillary networks that are difficult to assess solely with the use of Doppler ultrasound. Microbubbles have similar rheology to red blood cells and do not interfere with
hemodynamics (Lindner, et al., 2002). Work in nonhuman primate models has demonstrated the
feasibility of using contrast agents to enhance detection of blood flow during implantation and
early pregnancy (Keator, et al., 2011, Simpson, et al., 1997). Similarly, we recently implemented
CEUS to visualize spiral arteries and vascular filling in the human placenta at 11–13 weeks of
gestation (Roberts, et al., 2016). Utilizing this technology, we are able to calculate the
microvascular flux rate constant which is a measure of in-flow velocity. Although this is not an
absolute measure of ‘flow’ (volume/time), we will refer to the presence of contrast agent in the
placenta as placental blood flow.

To compare uteroplacental arterial remodeling changes with changes in uteroplacental
blood flow, we utilized the in situ placental hysterectomy collection at Cambridge University
(Boyd Collection). This previously published tissue resource is a collection of hysterectomies
during pregnancy performed for reasons of uterine bleeding, trauma, pregnancy termination, and
fetal demise (Burton, et al., 1999). Maternal age and parity are not available, but gestational age
is estimated by embryonic and fetal crown rump length (CRL). The advantage of the Boyd
Collection is the placenta and uterus have been sectioned into large blocks and embedded in paraffin, providing complete serial sections through the entire uteroplacental interface. Employment of the Boyd Collection, in combination with archived hysterectomy and decidual biopsies from our own institution, allowed for a comprehensive assessment of vascular remodeling, EVT plug immunophenotype, and ultrastructural analysis of the disintegrating plugs in the first trimester.

Since it has been previously suggested that the spiral artery plugs are only loosely cohesive and may even develop vascular channels communicating with the IVS (Burton, et al., 1999), the primary objective of this study was to utilize CEUS to examine maternal blood flow to the human placenta in the first trimester, and to correlate these data with an independent histological review of spiral artery ‘plugs’ in early pregnancy.
Materials and Methods

Contrast-Enhanced Ultrasound

Women scheduled for elective termination of pregnancy (n=34) underwent ultrasound studies at 6+3-13+6 weeks’ gestation. Patient demographics are given in Table I. Gestational age was determined by CRL on transvaginal ultrasound performed day-of the procedure. CEUS was performed using a Sequoia system (Siemens Medical Systems, Mountain View, CA) equipped with a 4C1-S transducer. Lipid-shelled octofluoropropane microbubble reagent (Definity®, Lantheus Medical Imaging, Billerica, MA) was intravenously infused (rate: 60 ml/hour) for visualization of uteroplacental perfusion as previously described (Roberts, et al., 2016).

Microbubble re-entry in the IVS was recorded in three replicates for each study. Replenishment kinetic curves were generated and flux rate (β) calculated as the rate of refilling of the vascular space until signal saturation (Figure 1A). Vascular filling of the entire placenta was visualized and captured in a single field of view within two visual planes: coronal and sagittal. The coronal plane was achieved by identifying bilateral uterine artery sources immediately after branching from the internal iliac artery. The sagittal plane was determined using the vesicocervical junction as a landmark with visualization of the internal iliac artery branching into the uterine artery.

In Situ Placental Histopathology: The Boyd Collection

The majority of the in situ placental specimens of the Boyd Collection (Figure 2A, Table II) were fixed by immersion of the intact uterus in formalin. Excised blocks were secondarily fixed in formalin or Bouin’s (e.g. case H916). One specimen (case H653) was perfused by the uterine artery with india ink, which may have affected spiral artery plug morphology and/or the presence or absence of maternal RBCs in the IVS. The width of decidua basalis viewed ranged
from 1.5 cm–7.0 cm (gestational ages 6-13 weeks based on CRL). These paraffin blocks had been previously serially sectioned, numbered and stained with hematoxylin and eosin, or Masson’s trichrome. All histological assessments were performed by one pathologist (TKM). By reviewing the entire sequence of serial sections through the placental bed, spiral arteries were traced from their openings distally through the decidua basalis proximally and then into their upstream radial and arcuate arteries. Artery segments (spiral, radial, arcuate) were scored for the presence of placental trophoblast invasion, obstruction of the vascular lumen (plug), degree of plugging (complete, channels, partial, absence of plugs), and the relative dilation of the lumen (+/- 2-fold increase in diameter compared with unremodeled vessels in a nulligravida hysterectomy specimens).

Spiral Artery Plug Ultrastructural and Immunohistochemical Analysis

Three nulligravida post-hysterectomy and five additional hysterectomies (two primigravida with placentas in situ and three multiparous postmenopausal cases) were obtained from the archives of the Department of Pathology, (OHSU, Table II). The OHSU archives also provided six first trimester decidua basalis specimens from primigravida women (Age: 22-33 years) having an elective termination at 6-8 weeks gestation (confirmed by CRL). Histologic sections of these OHSU specimens were similarly scored, although decidua basalis from terminations only provided cross-sections of spiral arteries.

Decidua basalis specimens from elective terminations (OHSU) were used to identify completely plugged and cannulated plugged spiral artery cross-sections. Serial sections were obtained for immunohistochemical studies; these spiral artery targets were then mapped onto the
accompanying paraffin block and cored using a 14-gauge needle. The cores were deparaffinized, fixed in 2.5% glutaraldehyde and processed by the Electron Microscopy facility (OHSU). Complete plugs were analyzed for the presence or absence of tight junctions (endothelial cells served as positive controls), desmosomes, apoptosis (nuclear membrane integrity), and necrosis (cytoplasmic vacuolization). Serial sections were stained for pancytokeratin (epithelial marker), E-cadherin (cell adhesion marker), Ki-67 (proliferation marker), CD31 (endothelial marker), CD3 (T-cell marker) and CD56 (cell adhesion marker [also stains uterine natural killer cells]). Slides were scored for the presence or absence of staining compared with internal tissue controls within each decidual basalis histologic section. Patterns were assessed for reproducibility within and between multiple sections per case and between cases.

Statistical Analysis

CEUS parameters were compared across gestation using an ANOVA with a Dunnett’s multiple comparison post hoc test, and within data acquisition replicates by linear regression (PRISM software, version 7.01; GraphPad). A value of p<0.05 was considered significant.

Ethical Approval

The CEUS protocol was approved by the OHSU Institutional Review Board (IRB#10744) with written consent obtained.
Results

Placental Perfusion in the First Trimester

Using CEUS we were able to assess maternal blood flow into the IVS in samples from 6+3-13+6 weeks of gestation. Figure 1B shows the flux rate, a measure of vascular impedance, across this gestational age range. These data demonstrate maternal perfusion through the spiral arteries as early as 6 weeks and more clearly from 8 weeks onwards (Video clip link). Within the 6+3-7+6 week age range, we observed wide variability in flux rates from 0.041 to 0.125 msec⁻¹.

Surprisingly, we do not demonstrate a progressive increase in microvascular flux rate with increasing gestational age across 6 to 13 weeks (Figure 1B). However, when comparing the change in microvascular flux rate relative to 6 weeks, there is a significant increase at 13 weeks (1 vs. 1.88 p<0.05, one way ANOVA with Dunnett’s post hoc test). We analyzed the variability within data acquisition replicates in one field of view (Figure 1C) and between whole placenta perfusion data acquired from coronal and sagittal orientations (Figure 1D). Replicates were highly correlated (p<0.0001).

Channels in Spiral Artery Trophoblastic Plugs

Our objective was to evaluate the Boyd Collection (6-13 weeks gestation, Table II) within the context provided by CEUS data suggesting intervillous flow beginning as early as 6 weeks of gestation. Similar to a thrombus being cannulated by endothelial cells, we observed well-demarcated channels forming within the spiral artery trophoblastic plugs (Figure 2B). The single 6 week specimen had spiral arteries apparently plugged by loosely cohesive EVTs, but maternal RBCs were also seen intermixed with these ‘plugs’. Vascular channels through the EVTs plugs could be traced to the IVS by 7-8 weeks and these channels became more apparent.
with larger luminal diameters as gestational age increased. Additionally, increasing numbers of maternal RBCs were apparent in the IVS after 7 weeks, consistent with blood flow. Although maternal RBCs could have leaked into the IVS secondary to trauma (reason for hysterectomy), preparation artifacts, or possible retrograde flow through venous connections during handling, the presence of well-formed channels within the plugs suggest they are physiologic. After 8 weeks most of the spiral arteries examined showed only partial EVT obstruction (Figure 2). These histological findings suggest that the collection of EVTs, termed ‘plugs’, do not completely obstruct flow to the IVS.

To further characterize the EVT plug channels we immunostained histologic sections and performed ultrastructural analysis. Unlike the cytotrophoblast cells rimming chorionic villi, the EVTs within spiral artery plugs lose E-cadherin expression and gain NCAM (neural cell adhesion molecule, CD56) (Figure 2E). Interestingly, CD56 is also a marker of uterine natural killer cells, which are thought to contribute to spiral artery remodeling. Importantly, the EVTs in the ‘plugs’ lack mitotic activity and are negative for Ki-67 immunostaining (Figure 2F). This finding contrasts with the trophoblasts in the anchoring villi which have numerous conspicuous mitotic figures. Instead, EVT plugging cells appear to be undergoing necrosis as gestation progresses in a distal to proximal fashion (highlighted by the red to blue color change in trichrome stained sections, Figure 2B). Apoptag staining of DNA strand breaks was negative suggesting that these cells are not undergoing apoptosis (data not shown).

To further investigate the cell adhesion characteristics of the EVTs in the plugs, we performed ultrastructural studies of decidua basalis samples at 6-8 weeks. These studies reveal that EVT plugs are loosely held together by desmosomes, not tight junctions (Figure 3). Many of the trophoblasts in these plugs also showed evidence of necrosis with cytoplasmic vacuolization,
but no nuclear membrane degradation (further data against apoptosis). Taken together, our
histological observations of the progressive disintegration of EVT plugs and the formation of
vascular channels into the IVS provide a correlation between tissue architecture and our CEUS
data suggesting flow into the IVS in the first trimester. However, we do not observe a
progressive increase in microvascular flux rate coincident with the progressive disintegration of
the spiral artery plugs which provided rationale for examining proximal regulation of maternal
blood flow to the IVS as a potential regulator of perfusion to the early placenta.

_Pregnancy-Induced Radial and Arcuate Artery Remodeling_

Previous reports of the Boyd Collection did not address the radial and arcuate arteries in
the myometrium upstream from the decidual spiral arteries (Burton, _et al._, 1999). Here, we
scored these vessels for the presence or absence of luminal dilation compared with nulligravida
hysterectomy controls and earlier gestational age specimens (Figure 4). Our observations
suggest that the distal radial arteries intersecting with the decidua basalis may begin dilating
around 9 weeks (case H1029) with placental cells tracking along the loose perivascular
adventitia by 15 weeks (Figure 5). The placental cells did not invade the radial artery muscular
walls, which were instead infused with CD3 positive T-cells. By 33 weeks the radial arteries are
even more dilated than the second trimester cases in the Boyd Collection (Figure 4C). This
suggests a potential temporal sequence in radial vascular remodeling. Moreover, it appears the
remodeling changes persist after the completion of pregnancy, because radial remodeling
changes are still present in postmenopausal uterine specimens (Figure 4D) and absent in the
nulligravida cases. Similarly, the arcuate arteries upstream from the radial arteries remodel
during pregnancy (Figure 4E compared with 4F). Although we see clear placental invasion of the
radial artery perivascular adventitia in the 15 week case, the arcuate arteries are negative for
perivascular trophoblasts in the 33 week specimen. A mechanism other than medial invasion and smooth muscle destruction may regulate radial and arcuate artery pregnancy-induced remodeling.
Discussion

Our comprehensive *in situ* histological analysis, in combination with the use of an *in vivo* imaging modality that has the sensitivity, and higher resolution, to facilitate visualization of microvascular filling of the IVS has allowed us to reveal new evidence of first trimester flow earlier in pregnancy than previously reported. Our novel CEUS data unequivocally demonstrate maternal blood flow in the IVS from 6-7 weeks of gestation, and are supported by morphological data that demonstrate the progressive disintegration of trophoblast plugs from 7 weeks onwards.

Maternal blood flow to the IVS has been addressed in several prior imaging studies mainly utilizing Doppler ultrasound with some evidence of blood flow as early as 5-7 weeks (Kurjak and Kupesic, 1998, Makikallio, *et al.*, 2004, Merce, *et al.*, 1996), others suggesting absence of flow prior to 8 (Carbillon, *et al.*, 2005, Makikallio, *et al.*, 2004, Makikallio, *et al.*, 1999, Valentin, *et al.*, 1996) or 10 weeks (Jauniaux, *et al.*, 2005) and general agreement that continuous flow is established by 12 weeks of gestation (Coppens, *et al.*, 1996, Hustin, *et al.*, 1988, Jauniaux, *et al.*, 1991, Jauniaux, *et al.*, 1992). Yet uncertainty remains, and while flow has been demonstrated early, at best it is conceded that this is limited to slow, non-continuous flow prior to the end of 8 weeks of gestation (Burton, *et al.*, 1999). Additionally, it has been suggested that early flow is indicative of pregnancy failure, as correlations have been made that support this notion (Jaffe, 2001, Jauniaux, *et al.*, 2003, Merce, *et al.*, 2009). However, the sensitivity and resolution of the ultrasound system must be considered when interpreting these conflicting data sets. Pregnancies that are destined for spontaneous demise may indeed have higher flow but it is possible that the increased flow may be within the limits of detection of current Doppler capabilities, but does not confirm that early flow is not present in successful pregnancies which may have lower, undetectable flow rates. Specifically, previous Doppler studies emphasizing no
perfusion to the early placenta may have been impaired by limitations and error generated when attempting to quantify low velocity perfusion in small vessels. Using a contrast agent enhanced our ability to track vascular filling of the placenta. Importantly in our study, we were able to carefully observe microbubble replenishment post destruction and do not have evidence to suggest impeded spiral artery refilling, or flow to the IVS. Flow appears to be continuous and evenly distributed as opposed to slowly meandering around obstructive plugs. Using the CEUS analysis parameters, it is possible to calculate an estimate of flow within a vessel (Keator, et al., 2011), however we have limited our data presentation to microvascular flux rate as the placenta has a complex vascular network, and absolute blood volume and flow calculations are derived using equations and assumptions that are dependent on a standard capillary network. However, the entry of a contrast agent with similar rheology to red blood cells, as detected by replenishment kinetic curves, is in itself strong evidence of the presence of flow at earlier gestational ages than previously assumed.

Despite inconsistencies in previously reported Doppler ultrasound studies attempting to determine if blood flow to the early human placenta was present, previous analysis of the Boyd Collection suggested the presence of channels in the spiral artery plugs as early as 7 weeks (Burton, et al., 1999). Of interest, by the eighth week of pregnancy these channels were formed within the spiral artery plugs and overlying anchoring villi that communicated with the IVS and were measured to be 10-20µm in diameter and by 9-10 weeks there were well defined channels (~100µm in diameter) into the IVS, clearly suggesting that blood flow to the early placenta could be possible. The unequivocal demonstration of contrast agent in the IVS at 6-7 weeks of gestation identified by CEUS motivated our re-examination of the Boyd Collection. Our review of this Collection independently supported the observation of vascular channels. In an effort to
characterize the temporal sequence of the formation of these channels, we extended these observations with immunohistochemistry and ultrastructural analysis.

Our histologic investigation of in situ preserved specimens from the Boyd Collection demonstrated that the trophoblasts in the loosely cohesive clusters of EVTs, so-called ‘plugs’, are not mitotically active, they are loosely held together by desmosomes, and they provide at least capillary-sized intercellular channels that would permit blood flow by 7 weeks. In contrast to the trophoblasts in the anchoring columns which proliferate and express E-cadherin, the EVTs do not proliferate and change cell surface adhesion markers from E-cadherin to CD56 (Kam, et al., 1999). If the identified channels from 6-7 weeks provide capillary sized pathways for intervillous blood flow, they may explain our microbubble reappearance kinetic data. These channels are reproducibly evident by 7-8 weeks gestation and we are confident they represent an in vivo differentiation rather than an artifact, because the Boyd Collection provides placentas fixed in situ. This approach avoids potential disruptions that may arise in elective termination curettage specimens.

Significantly, we do not observe a progressive increase in microvascular flux rate into the IVS with increasing gestational age, as would be predicted based on our histological assessment of the progressive disintegration of spiral artery plugs. Therefore, we suspect that the significant increase in flux rate at 13 weeks is more likely related to radial artery luminal diameter increases at the end of the first trimester, rather than loss of spiral artery plugs. This idea is supported by our histological analysis of a small cohort from the Boyd Collection, and we suggest that pregnancy-induced remodeling of the upstream vessels in the myometrium may be an important and under-appreciated component.
Spiral artery remodeling has been the most studied pregnancy-induced change in the uterine vascular network probably because of the accessibility of the tissue and the assumption that the spiral artery lumen must be the narrowest in the uterine vascular network. The rate of flow in a blood vessel is proportional to the fourth power of the radius of the vessel. If the spiral arteries are the narrowest lumen in the vascular tree, then dilation and opening of these vessels would be the key to regulating flow rate. In the Boyd Collection, it is clear that spiral arteries are more dilated and attenuated than the radial arteries with the exception of the trophoblastic plugs.

The impedance in the conduit progressively disintegrates from 7-12 weeks. If flow was entirely dependent on the diameter of the channels within these plugs, we would expect a linear relationship between flow and gestational age. This was not the case. Therefore, we suspect the radial artery luminal diameter may be key to regulating uteroplacental blood flow in the first and early second trimester, although it is accepted that these observations are based on immersion, rather than perfusion-fixed material.

The uterine vascular network is complex and the spiral arteries are only a short segment in the distal most aspect of this network. The uterine artery leads to the arcuate arteries that track in parallel with the serosal surface. The arcuates then penetrate the myometrium via perpendicular radial arteries, which feed the endometrial/decidual spiral arteries. Spiral arteries are angiogenic and grow and are shed each month with the menstrual cycle. During pregnancy the decidual spiral arteries are the first to dilate in a distal to proximal fashion from the IVS towards the radial arteries. The Boyd Collection illustrates this process clearly and shows dilated spiral arteries intersecting with more narrow unremodeled radial arteries. Others have described a progressive transformation of the uterine vascular network over time (Harris and Ramsey, 1966, Pijnenborg, et al., 1980), and our CEUS data suggest to us the upstream radial artery
remodeling may be the key to understanding early second trimester changes in uteroplacental flow resistance. In fact, new data from mice suggest radial artery diameter, not spiral artery diameter, may be the key resistance regulatory point for uteroplacental blood flow (Rennie, et al., 2016), consistent with this hypothesis. Moreover, our findings suggest that like mice, radial/arcuate remodeling by mechanisms other than direct placental invasion of the vascular wall may be significant contributors to blood flow regulation in early human pregnancy. These findings should spur investigations into proximal vascular remodeling mechanisms and the relationship with common obstetric complications like pre-eclampsia (Ong, et al., 2005).

Myometrial vascular remodeling may help explain why complications like pre-eclampsia are more common in a woman’s first pregnancy. It is important to emphasize that only the decidual lining is lost after pregnancy and with it, the decidual spiral arteries. The myometrial radial and arcuate arteries remain. The precise timeline of complete radial and arcuate remodeling during pregnancy is yet to be established, but there are certainly changes that continue throughout pregnancy (Burchell, 1967, Harris and Ramsey, 1966) and changes that seem to remain embedded in the muscular walls of these myometrial blood vessels post-pregnancy. In our analysis, the OHSU post-hysterectomy cases provided four nulligravida and four multigravida postmenopausal uterine specimens for review: there were clear differences between radial and arcuate arteries from women who had been pregnant compared with nulligravidas.

Future studies need to investigate the mechanisms regulating myometrial artery remodeling. Similar to decidual spiral artery remodeling in cases of ectopic pregnancies (Craven, et al., 1998), our histologic review of the radial arteries and the arcuate arteries suggests to us that the maternal inflammatory T-cell response may play an important role in this process.
Although the second trimester sample size is small, we did not observe placental trophoblast invasion of the vascular media in any of the radial or arcuate artery cross-sections. These hypotheses warrant further study by immunohistochemical analysis of placental in situ hysterectomy specimens and high resolution blood flow imaging analyses.

Our CEUS cohort included four participants in the 6+3 to 6+6 week gestational age range with low flux rates measured in two women and higher flux in the other two. This variability may indicate that this is a critical time period in the establishment of maternal flow, although this interpretation is speculative at this time and requires further investigation. It is also important to note that gestational dating was determined by fetal biometry and therefore the margin of error in such measurements could be a contributing factor to the observed differences at this gestational age range. A possible future approach would be to implement super resolution ultrasound imaging in combination with microbubble use (Errico, et al., 2015). This technology is still under development for placental imaging, but has the potential to provide invaluable flow data both for understanding normal physiology, and for the improved clinical identification and management of pregnancies at-risk for vascular compromise. Specifically, advances in imaging technology may aid our understanding of early fetal development and the pathophysiology of common obstetric problems, which in turn will inform the search for biomarker discovery.

In summary, here we demonstrate maternal blood flow to the placental IVS at 6 weeks of gestation. Although we cannot comment on the resistance to flow prior to 6 weeks, our data support the onset of maternal flow into the IVS earlier than previously suspected. Our data are correlated with a comprehensive morphological and phenotypic review of spiral artery plugs and the uterine vascular network. The so-called ‘plugs’ are only loosely cohesive at 6 weeks and begin forming clear capillary-sized channels into the IVS by 7 weeks. We suggest there may be
a two-step process that involves progressive remodeling of EVT cell clusters, or ‘plugs’, with reduced resistance to intervillous flow, and that this is protected hemodynamically by persistence of radial artery resistance until the end of the first trimester when the radial arteries start to remodel and develop more dilated lumens. We propose that the upstream radial and arcuate arteries may be significant regulators of uteroplacental blood flow, especially by the end of the first trimester.
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Figure Legends
**Figure 1: Contrast-Enhanced Ultrasound flux rate and variance.**

A) Example replenishment kinetic curve demonstrating calculation of flux rate (\(\beta\)) from the slope of the curve (y). B) Flux rate (\(\beta\) value in sec\(^{-1}\)) plotted over 6 to 13 weeks of gestation (mean+SEM). The number of participants is represented by the number overlay in each bar. The variance between measures of flux rate within C) individual data acquisition replicates and D) in two different anatomical orientations.

**Figure 2: Characterization of channels within the spiral artery plugs.**

A) *En bloc* sections from the Boyd Collection (gross example is from 10 weeks’ gestation) were available for serial section analysis revealing well-formed channels (*) through loosely cohesive endovascular trophoblast cells (B), which trichrome staining suggested may be related to cell death at the maternal arterial blood interface (arrow). C) Elective termination specimens were employed for immunophenotyping. D) Extravillous trophoblast cells expressed less E-cadherin than the strong staining seen in villous cytotrophoblast cells (top left). E) The endovascular plugs had their own specific CD56 staining pattern, which was absent in all other types of trophoblast. F) Although the overlying anchoring villi were proliferating, the cells making up the spiral artery plugs were Ki-67 negative. Figure 2B was photographed using a 20x objective (~200x); C-F were photographed using a 10x objective (~100x magnification).

**Figure 3: Analysis of vascular channels through spiral artery plugs.**

A) By 8 weeks gestation the channels were well formed and clearly communicated with the intervillous space. B) They were lined by trophoblast cells negative for CD31 (*). C-F) Electron microscopy showed loosely cohesive cells connected by desmosomes (red arrow) that were undergoing necrotic degeneration forming fibrin-lined channels. A and B were photographed
using a 20x objective (~200x magnification). C and D 3000x (note RBCs); E 6000x (note RBCs); F 60,000x.

**Figure 4**: Pregnancy-induced remodeling of myometrial radial and arcuate arteries.

A) The radial arteries appeared to begin remodeling at the end of the first trimester associated with perivascular lymphocytes and few extravillous trophoblast cells that track along the adventitia, but they did not invade the media. B) By 15 weeks the upstream proximal radial arteries showed signs of remodeling that was clearly evident in the 33 week hysterectomy specimen (C) and left permanent fibrin changes seen in multigravida postmenopausal specimens (D). Compared with nulligravida uteri (E), the arcuate arteries also dilated and attenuated (F); this occurred in the absence of trophoblast. Photographs taken using a 5x objective (~50x magnification).

**Figure 5**: Radial artery remodeling involves perivascular trophoblasts and accumulation of medial CD3 positive and CD3 negative lymphocytes.

A) Remodeling radial artery at 15 weeks gestation by hematoxylin and eosin stain showed medial muscular and extracellular matrix changes appreciated best in complete cross-sections (dashed circle). B) PAS stain highlighted perivascular adventitia that was invaded by cytokeratin positive extravillous trophoblast cells (C) that do not appear to invade the muscular media. D) Instead, there are numerous lymphocytes, including CD3 positive T-cells in the walls of these remodeling arteries. Photographs taken using a 5x objective (~50x magnification).
Video legend

Video 1: Visualization of uteroplacental vascular filling in a human subject at gestational age 8 weeks and 4 days. Microbubbles are destroyed by a 2 second increase in mechanical index and vascular refilling is observed until signal saturation is reached. Digital recording obtained at 1 frame/75 msec.
Authors’ Roles

VHJR assisted with acquisition, and performed all analysis of the CEUS data, and wrote the manuscript.

TKM conceived of and designed the study, performed all histological review, and co-wrote the manuscript.

PB facilitated and assisted with the CEUS studies, and reviewed the manuscript.

MM assisted with data acquisition, histological sample processing and analysis, and reviewed the manuscript.

GJB facilitated the histological studies and edited the manuscript.

JOL acquired the CEUS data and edited the manuscript.

AEF conceived of and designed the study, acquired the CEUS data and edited the manuscript.

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Funding

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Conflict of Interest There are no conflicts to report
Table 1: Contrast-Enhanced Ultrasound study participant demographics

<table>
<thead>
<tr>
<th>Baseline Characteristics</th>
<th>Gestational Age (weeks)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>6 (4)</td>
</tr>
<tr>
<td>Maternal age (y)</td>
<td>28</td>
</tr>
<tr>
<td>Nulliparous (%)</td>
<td>25</td>
</tr>
<tr>
<td>Race/Ethnicity (%)</td>
<td></td>
</tr>
<tr>
<td>Caucasian</td>
<td>100</td>
</tr>
<tr>
<td>African American</td>
<td>-</td>
</tr>
<tr>
<td>Hispanic</td>
<td>-</td>
</tr>
<tr>
<td>Other</td>
<td>-</td>
</tr>
<tr>
<td>Body Mass Index (kg/m²)</td>
<td>22</td>
</tr>
<tr>
<td>Smoking (%)</td>
<td></td>
</tr>
<tr>
<td>None</td>
<td>50</td>
</tr>
<tr>
<td>Current</td>
<td>25</td>
</tr>
<tr>
<td>Former</td>
<td>25</td>
</tr>
<tr>
<td>Specimens</td>
<td>Gestational Age (CRL)</td>
</tr>
<tr>
<td>-------------</td>
<td>------------------------</td>
</tr>
<tr>
<td><strong>Boyd Collection</strong></td>
<td></td>
</tr>
<tr>
<td>H710</td>
<td>6 weeks (4.0 mm)</td>
</tr>
<tr>
<td>H750</td>
<td>7 weeks (10.0 mm)</td>
</tr>
<tr>
<td>H937</td>
<td>8 weeks (15.0 mm)</td>
</tr>
<tr>
<td>H673</td>
<td>8 weeks (15.0 mm)</td>
</tr>
<tr>
<td>H916</td>
<td>9 weeks (20.0 mm)</td>
</tr>
<tr>
<td>H1029</td>
<td>9 weeks (20.0 mm)</td>
</tr>
<tr>
<td>H630</td>
<td>10 weeks (30.0 mm)</td>
</tr>
<tr>
<td>H653</td>
<td>11 weeks (46.0 mm)</td>
</tr>
<tr>
<td>H870</td>
<td>12 weeks (55.0 mm)</td>
</tr>
<tr>
<td>H691</td>
<td>12.5 weeks (60.0 mm)</td>
</tr>
<tr>
<td>H1094</td>
<td>13 weeks (73.0 mm)</td>
</tr>
<tr>
<td><strong>OHSU Collection</strong></td>
<td></td>
</tr>
<tr>
<td>Case 1</td>
<td>6.5 weeks (6.0 mm)</td>
</tr>
<tr>
<td>Case 2</td>
<td>7 weeks (11.0 mm)</td>
</tr>
<tr>
<td>Case 3</td>
<td>7 weeks (10.0 mm)</td>
</tr>
<tr>
<td>Case 4</td>
<td>7.5 weeks (13.0 mm)</td>
</tr>
<tr>
<td>Case 5</td>
<td>8 weeks (14.0 mm)</td>
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<tr>
<td>Case 6</td>
<td>8 weeks (16.0 mm)</td>
</tr>
<tr>
<td>Case 7</td>
<td>15 weeks (85.0 mm)</td>
</tr>
<tr>
<td>Case 8</td>
<td>32 weeks (260.0 mm)</td>
</tr>
<tr>
<td>Cases 9-12</td>
<td>Postmenopausal G0</td>
</tr>
<tr>
<td>Case 13-16</td>
<td>Postmenopausal G2-5</td>
</tr>
</tbody>
</table>

Boyd Collection is housed within the *Center for Trophoblast Research*, Cambridge, UK; it is a collection of serial sections through paraffin blocks taken from the entire uteroplacental interface from a series of hysterectomies performed in the 1950s. Archived decidua basalis paraffin blocks from primigravida elective terminations and four hysterectomy specimens were obtained from Oregon Health & Science University (OHSU). EVT: extravillous trophoblasts; EM: electron microscopy; IHC: immunohistochemistry.
Figure 4

A

B

C

D

E

F