Gestational Dietary Protein Is Associated with Sex Specific Decrease in Blood Flow, Fetal Heart Growth and Post-Natal Blood Pressure of Progeny

Juan H. Hernandez-Medrano, Katrina J. Copping, Andrew Hoare, Wendela Wapanaar, Rosalie Grivell, Tim Kuchel, Giuliana Miguel-Pacheco, I. Caroline McMillen, Raymond J. Rodgers, Viv E. A. Perry

1 School of Veterinary and Medical Science, University of Nottingham, Sutton Bonington, United Kingdom, 2 School of Paediatrics and Reproductive Health Robinson Institute, University of Adelaide, Adelaide, Australia, 3 South East Vets, Mt Gambier, South Australia, Australia, 4 The Chancellory, University of Newcastle, Newcastle, New South Wales, Australia, 5 South Australian Health and Medical Research Institute, Adelaide, Australia, 6 Department of Perinatal Medicine, The Womens and Childrens Hospital, North Adelaide, Australia

Current address: Facultad de Medicina Veterinaria y Zootecnia, Universidad Nacional Autonoma de Mexico, Mexico D.F., Mexico

These authors contributed equally to this work.

This author is the senior author on this work.

viv.perry@nottingham.ac.uk

Abstract

Study Overview

The incidence of adverse pregnancy outcomes is higher in pregnancies where the fetus is male. Sex specific differences in feto-placental perfusion indices identified by Doppler assessment have recently been associated with placental insufficiency and fetal growth restriction. This study aims to investigate sex specific differences in placental perfusion and to correlate these changes with fetal growth. It represents the largest comprehensive study under field conditions of uterine hemodynamics in a monotocous species, with a similar long gestation period to the human. Primiparous 14mo heifers in Australia (n=360) and UK (n=180) were either individually or group fed, respectively, diets with differing protein content (18, 14, 10 or 7% crude protein (CP)) from 60d prior to 98 days post conception (dpc). Fetuses and placentae were excised at 98dpc (n = 48). Fetal development an median uterine artery blood flow were assessed monthly from 36dpc until term using B-mode and Doppler ultrasonography. MUA blood flow to the male feto-placental unit increased in early pregnancy associated with increased fetal growth. Protein restriction before and shortly after conception (-60d up to 23dpc) increased MUA diameter and indices of velocity during late pregnancy, reduced fetal heart weight in the female fetus and increased heart rate at birth, but decreased systolic blood pressure at six months of age.

Conclusion and Significance

Sex specific differences both in feto-placental Doppler perfusion indices and response of these indices to dietary perturbations were observed. Further, maternal diet affected
development of fetal cardiovascular system associated with altered fetal haemodynamics in utero, with such effects having a sex bias. The results from this study provide further insight into the gender specific circulatory differences present in the fetal period and developing cardiovascular system.

**Introduction**

It has been established that the male fetus is more susceptible to placental insufficiency and adverse pregnancy outcomes [1,2]. Increased median uterine artery resistance indices have been used in human medicine to identify placental insufficiency. These techniques have been adapted for use in the bovine [3,4] where it was shown that characteristic variations in the median uterine artery (MUA) blood flow could be observed over pregnancy. Moreover, these blood flow volume (BFV) measures were correlated with birth weight of calves.

Dietary restriction in early gestation affects fetal growth from 36dpc to the end of the first trimester [5–7] and may affect postnatal reproduction and production traits in cattle progeny [8–13]. Critical windows exist during oocyte, embryo and fetal development where decreased maternal dietary protein may reset early cell lineages [14]. In sheep maternal nutrition affects oocytes number, quality [15] and embryo survival [16]. In cattle sex specific differences are seen in embryo glucose uptake [17] DNA methylation pattern [18], expression of key developmental genes [19] and fetal and placental perfusion [1,16]. Intriguingly, neonatal and fetal death in cattle is increased if the fetus is male [20,21], as reported in other mammalian species [22,23].

The aim of this study is to further evaluate the critical windows during development where dietary protein may alter fetal development and subsequently impact upon lifetime health and production. This paper focuses on the effects of protein restriction during specific windows of development known to affect growth of the fetus; preconception [24] periconception (including implantation) [7] and first trimester [20], upon blood flow to the fetus and the developing cardiovascular system. We hypothesise that low maternal protein during the peri-conception period and the first trimester will increase fetal blood flow and cardiovascular development in a gender dependent manner.

**Materials and Methods**

**Ethics statement**

This project was approved by the University of South Australia IMVS Animal Ethics Committee (Approval number 18/11; Experiment 1) and the United Kingdom’s Animals (Scientific Procedures) Act 1986 (Experiment 2), respectively.

**Experiment 1 (Expt 1) Peri-conception and Post-implantation**

The purpose of this experiment was to evaluate the impact of dietary protein during the peri-conception (PERI; day -60 to day 23 (implantation being 18-22dpc); insemination day = conception, day 0) and post-implantation (POST; day 23 to day 98) periods in heifers. This study was carried out at Tungali, South Australia (Lat. 34°30'9.2" South, Long. 139°18'8.3" East) and full details of materials and methods have been published previously [7]. Briefly, primiparous *Bos taurus* x *Bos indicus* heifers (n = 360) were selected and acclimatized for 60 days prior to the start of the experiment. Heifers that did not eat well in individual stalls were...
removed from the study. After acclimatisation, heifers were split into 4 groups based on a two-by-two crossover factorial design with conception stage (PERI) or post-implantation (POST) and dietary crude protein (CP) content (14% or 7%CP; reflecting pasture conditions in Australia) as main factors. Heifers were randomly assigned according to bodyweight (BW) to either high (H; 14%CP, n = 177; 345.5 ±26.9kg BW (mean ±S.E.M.)) or low CP diet (L; 7%CP, n = 173; 343.1 ±25.5kg BW) sixty days prior to conception. At 23 days post-conception (dpc), L and H groups were split into two further groups based on %CP content (post-implantation H and L groups) to give the crossover design resulting in 4 groups as follows: HH group (PERI 14%CP and POST 14%CP; n = 88); HL group (PERI 14%CP and POST 7%CP; n = 89); LH group (PERI 7%CP and POST 14%CP; n = 87) and LL group (PERI 7% and POST 7%; n = 86). Diets used in the experiment varied in crude protein content but were otherwise isocaloric and supplemented with a commercial vitamin and mineral preparation. The pellet portion of the diet was fed individually to the heifers daily in stalls with straw ad libitum in pens. At the end of the first trimester of gestation (98 dpc), all heifers were fed a similar maintenance diet (12% CP) until parturition. Detailed composition of the heifer rations was previously published [7] and a brief nutrient content is presented in Table 1.

### Synchronization Protocol

Oestrous cycles were synchronized 50 days after initiation of diet treatment using intravaginal progesterone releasing devices (n = 2; progesterone 1.56g, CUE-MATE, Bioniche Animal Health, Bayer Australia Ltd, Pymble, Australia) inserted sequentially on day -10, relative to artificial insemination (AI; day 0), with simultaneous intramuscular (i.m.) injection of 2mg oestradiol benzoate (Bomerol, Bayer Australia Ltd, Pymble, Australia). On day -2 intravaginal devices were removed and heifers received i.m. injections of prostaglandin 2 alpha (2ml; Ovuprost) and equine chorionic gonadotrophin (300 IU; eCG; Pregnecol). Twenty-four hours after device removal, heifers were injected i.m. with 1mg oestradiol benzoate and inseminated.

![Table 1. Nutrient content of heifer rations used in Experiment 1 (Australian n = 360; PERIconception and POSTimplantation) and Experiment 2 (UK, n = 188; PREconception and POSTconception) based on a dry matter basis.](https://doi.org/10.1371/journal.pone.0125694.t001)
24 hours later (day 0 = conception day) with frozen semen from a single bull (*Bos indicus x Bos taurus*) with known estimated breeding values (EBVs) for growth and low birth weight.

**Experiment 2 (Expt 2) Pre- and post-conception**

The aim of this experiment was to further study the effects of dietary protein during differing intervention periods and at levels representative of UK commercial beef cattle protein intake. Dietary intervention occurred from day -60 to conception (AI day = conception, day 0) or from conception to 90dpc. Eighty-eight primiparous, 10–14 month-old *Bos taurus* heifers in two locations (North Yorkshire: Lat. 54°23'41.3" North, Long. 1°18'11.6" West; and Buckinghamshire: Lat. 51°34'15.7" North, Long. 0°38'32.2" West) were selected and allocated to two regimens during the preconception period (PRE, n = 88) from -60d to conception of either high crude protein (18%CP) or low dietary protein (10%CP). A further group (n = 100, POST; Northumberland, UK: 55°22'11.6" North, 1°45'04.8" West) were allocated to two dietary regimens from conception to 90dpc fed either 10%CP (Low) or 14% CP (High). This resulted in 4 independent groups: pre-conception high (PRE H, n = 44; 347.2 ±9.4kg BW), pre-conception low (PRE L, n = 44; 355.5 ±10.8kg BW), post-conception high (POST H, n = 51; 392.6 ±3.8kg BW) and post-conception low (POST L, n = 49; 393.9 ±3.6kg BW). Heifers in this experiment were group-fed the corresponding diets. After the end of the dietary intervention, all animals were run together on each individual farm under similar diet and management practices for the remainder of the pregnancy.

**Synchronization Protocol**

Oestrous synchronisation was carried out by a commercial breeding company following a 5-day synchronization protocol for beef heifers reported previously by Bridges *et al.*[25]. Similarly to experiment 1, following synchronisation heifers were inseminated using semen from a single bull with known EBVs for growth and low birth weight on each farm.

**Heifer and Fetal Measurements**

Fetal growth was measured using trans-rectal ultrasound (probe 5–12.5; Sonosite M-Turbo, Sonosite Inc., Bothell, Washington, USA) at 35 and 60 dpc. Simultaneously, fetal crown-rump length (CRL; 35dpc), biparietal diameter (BPD; 60dpc) and sex (60dpc) were recorded for all pregnant heifers. Crown-rump length (CRL) was measured from a lateral view of the fetus from the tip of the nose to the base of the tail, while biparietal diameter (BPD) was measured from a dorso-ventral view of the cranium and perpendicular to the sagittal crest as the widest span between the most lateral parts of the parietal bone [5]. After 60dpc, empty heifers were withdrawn from the experiment.

Transrectal pulsed B mode Doppler sonography (Sonosite M-Turbo, Sonosite Inc., Bothell, Washington, USA) of the MUA (as described previously [4]) was performed at 120, 150, 180 and 210 dpc. Briefly, in longitudinal color Doppler section of the artery, blood-flow waveforms were measured at an interrogation angle between Doppler ultrasound beam and flow direction from 45° to 60°, with the volume sample mark located in the middle of the vessel. Evaluations of the MUA diameter were done in a cross-section of the vessel just forward of the external iliac vein [26]. Doppler indices, such as resistance index (RI), pulsatility index (PI) and timed average maximum velocity (TAMV) were recorded. Total blood flow volume (BFV) to the pregnant uterus was estimated following the equations presented by Herzog *et al.*[4]. Time constraints considering the number of heifers to be scanned dictated that only the ipsilateral MUA was scanned over two days. The relationship between the two arteries in blood flow parameters of resistance, velocity and volume have previously been reported [26], with resistance...
decreasing and velocity, volume and diameter increasing in the ipsilateral side at an increased rate from the 4th month of pregnancy.

At 98dpc, a subset of heifers (HH = 12; HL = 15; LH = 10; LL = 9) in Expt 1 were killed at a commercial abattoir. The pregnant uterus from each heifer was collected and the fetus excised, blood sampled, measured (CRL and BPD, as previously described; [7]) and weighed. Fetal sex was recorded and tissue samples (i.e. heart) weighed and collected. The remaining heifers in Exp 1 were allowed to calve and postnatal arterial blood pressure measured as describe in the following section.

Postnatal Arterial Blood Pressure

In Experiment 1, arterial blood pressure was measured to determine the long-term effects of maternal under-nutrition during gestation on the cardiovascular system in the offspring. This was carried out using the tail-cuff system previously described and validated for calves [13] at 6, 8 and 10 months of age (200, 250 and 350d, respectively), using a non-invasive blood pressure monitor (Cardell Veterinary Monitor 9401BP, SHARN Veterinary). Briefly, the measurement was taken by the same operator on each occasion whilst the calf was restrained quietly in a crush. At least five consecutive measurements were taken for each calf. Only measures where three of the readings ranged by < 10 mmHg were used to obtain a mean [13]. Those with greater variation were discarded.

Statistical Analysis

All data was analysed using STATA software (STATA/IC for Windows; StataCorp LP, College Station, Texas, USA). Data was checked for normality and transformed if required. Data regarding fetal measurements (CRL and BPD) was analysed using factorial ANOVA with diet (L or H), conception stage (PRE, PERI- POSTconception and POSTimplantation) and fetal sex as main factors, with interactions between these being explored. Experiment 1 and 2 were analysed separately as dietary treatment and intervention periods differed. Additionally, for experiment 2 a farm factor was used as a blocking factor. As heifer is considered the experimental unit, repeated measurements ANOVA with Greenhouse-Geisser epsilon adjustment for the within-subject variance was used when comparing Doppler measurements. Data is presented as the mean values of the raw data ± standard error of the mean (S.E.M) unless stated otherwise. After comparison of means (hypothesis tests), a p-value ≤ 0.05 was considered as statistically significant and a P-value ≤ 0.10 considered as a tendency.

Results

Uterine Artery Blood Flow

Blood flow volume (BFV; Fig 1) increased over gestation in heifers in all experimental treatments (P = 0.01) in association with reduced PI (P < 0.001) and increased MUA diameter (P = 0.01; Fig 2). In Expt 1, BFV increased over gestation (P < 0.001), this effect being influenced by sex (P = 0.01) and PERIconception diet (P = 0.005; Fig 1A). Those heifers with a male feto-placental unit at 120dpc had significantly greater BFV (P = 0.03) compared to those with a female. This effect on BFV was associated with low PERIconception protein reducing PI in all heifers at 120dpc compared to those on the high PERIconception diet (P = 0.02; Fig 3). Moreover heifers in the low PERIconception diet carrying male fetuses had an increased PI (P = 0.05) and a reduced BFV at 150dpc (P = 0.02), compared to those that received the high PERIconception diet.
Fig 1. Blood Flow Volume according to sex and PERI and PREconception diet. Mean (±SEM) estimated blood flow volume (cm$^3$) in heifers carrying male (M) or female (F) fetuses by treatment diet in A) Experiment 1 (Low 7%CP, M n = 20 and F n = 8; and High 14%CP, M n = 25 and F n = 11, from 60d prior to conception to 23dpc) and in B) Experiment 2 (Low 10%CP, M n = 13 and F n = 3; and High 18%CP, M n = 7 and F n = 11 from -60d to conception); ** denotes $P \leq 0.01$. 

doi:10.1371/journal.pone.0125694.g001
Similarly TAMV was affected by sex (P = 0.03) and diet (P = 0.05). At 120dpc TAMV was increased in those heifers carrying a male rather than a female fetus (P = 0.03; Fig 4). At 150dpc, TAMV increased in heifers receiving the low compared to high PERIconception protein diet (P = 0.05). At 210dpc, heifers fed the low PERIconception or POSTimplantation protein diet had increased TAMV (P = 0.03; Fig 4A) compared to those fed the high diet during these periods.

Maximal velocity (Vmax) was affected similarly to TAMV. At 150dpc, heifers with a male fetus fed the low protein PERIconception diet had a lower Vmax than heifers fed the high diet (P = 0.01; Fig 5). At 210dpc, heifers with a male fetus fed the low PERI and POSTimplantation diet had a higher Vmax measurement than heifers that received the high PERI and POSTimplantation diet (P = 0.03).

In combination PERIconception low protein diet increased blood flow to fetus over gestation by reducing PI (resistance) and increasing TAMV (velocity; Fig 6).

In Expt 2, there were lower numbers of heifers per treatment group (n = 40 in the PREconception dietary treatments and n = 50 in the POSTconception treatments). There was a trend for the low PERIconception protein diet to increase BFV (P = 0.06; Fig 1B) and TAMV (p = 0.06) at both 36 and 60dpc and at 180 and 210dpc (P = 0.08; Fig 4B). At 210dpc MUA diameter was increased in those heifers that received the low PREconception diet (P = 0.01; Fig 2). The low protein POSTconception diet increased MUA diameter (P = 0.04) at 60dpc associated with a trend to increased BFV (P = 0.09).

**Blood Flow and Birth Weight**

Birth weight was not affected by dietary treatment in Expt 1 but in Expt 2 the low PREconception diet (10 vs 18% CP) increased birth weight in male calves (P = 0.03). Birth weight was positively correlated with MUA blood flow in both experiments under either PRE or PERIconception dietary regimen (P = 0.01; Fig 7). This effect however, was greater in male calves on the low PERIconception diet (r = 0.42, P = 0.004) with the opposite dietary effect in the female; that is, high PERIconception diet in the female increased the correlation between MUA blood flow and birth weight.
flow and birth weight ($r = 0.47$, $P = 0.03$). In Expt 2 again it was the high PREconception diet that caused the significant correlation with birth weight in females ($r = 0.57$, $P = 0.03$) with a trend to significance in the low PREconception diet males ($r = 0.52$, $P = 0.07$). There was no significant correlation when males were subjected to a high PREconception diet or females to the low PREconception diet.

**Fetal Heart Weight (Expt 1)**

In the excised 98dpc fetus low protein diet (7%CP) compared to high (14% CP) during the PERIconception period reduced fetal heart weight in female fetuses ($P < 0.05$; Fig 8).

**Fetal Growth (Expt 2)**

Protein intake during the PREconception period in Experiment 2 had no effect upon fetal growth at 36 or 60dpc. Low protein (10%CP) during the first trimester (POSTconception diet)
was associated with a transient increase in fetal growth as measured by CRL at 36dpc compared to the high (14%CP) diet, but this effect was not present at 60dpc (Fig 9).

Neonatal Heart Rate

Low PERI conception dietary protein (7%CP) increased heart rate at birth in female offspring in Expt 1 compared to the high diet (14%CP; P < 0.05). There was no effect on heart rate in the animals in Expt 2 (10% vs 18%CP; Fig 10).
Low protein during the PERIconception period (7% CP) decreased systolic blood pressure (P < 0.05) in female progeny at six months of age with no effect at eight or ten months (Fig 11).

Blood Pressure in Progeny

Low protein during the PERIconception period (7% CP) decreased systolic blood pressure (P < 0.05) in female progeny at six months of age with no effect at eight or ten months (Fig 11).
Discussion

This study is the first to show sexually dimorphic effects of transient protein restriction during peri-conception upon blood flow to the placenta and the developing fetal cardiovascular system. This research represents the largest comprehensive study of uterine hemodynamics in a monotocous species, with a similar long gestation period to the human completed under field conditions.

We have recently published the first report of the enhanced sensitivity of the male fetal calf to dietary perturbation in early pregnancy, affecting growth [7] and placental function [20]. In the heifers from Expt 1, the PERIconception low protein diet at 7%CP reduced fetal growth.
compared with 14%CP diet with the effect being greater in the male [7]. Interestingly this effect upon growth was no longer apparent at birth. In contrast in the data presented here from Expt 2, the high protein diet (14%CP) during the first trimester of pregnancy transiently decreased fetal growth compared to the low (10%CP) diet. Furthermore, the low protein PREconception diet of 10% increased birth weight compared to the high PREconception diet of 14%.

We consider that these differing effects may be influenced by the different levels of protein and the different dietary intervention periods in the two studies. In Expt 1, the PERI conception diet did not cease until 23dpc (after implantation of the embryo) whereas in Expt 2 the PREconception diet ended at conception (i.e. AI) and the POSTconception diet started at conception. Both POSTconception diets, however, in Expt 1 and 2, continued until the end of the first trimester. We could expect this PERI conception diet, therefore, to exert a larger effect as it encapsulated both the development of the oocyte, the embryo and initial fixation to the uterus. The protein levels in Expt 1 were also more restrictive than those in Expt 2. The low protein diet in Expt 1 allowed a daily gain of 0.2kg per day compared 0.5kg when heifers were fed the high diet (data not shown). In the UK (Expt 2), however, both the low and high protein diets allowed a gain of at least 0.7kg i.e. there was no significant difference in the weight gain of the two groups (data not shown). The low protein PREconception diet, however, yielded an increase in birth weight. This may reflect an increase in placental development, as blood flow to the placenta (at 210dpc) in these animals was also elevated compared to those on high PREconception diet with this blood flow strongly correlated with birth weight (P < 0.01). Previous reports have shown contradictory effects of low protein diets during the first trimester followed by increased protein after 2nd trimester on placental trophectoderm volume and cotyledon weight [27,28].

In accord with previous work in beef cattle [3] resistance indices in the MUA traced a characteristic pattern over gestation and further, similar sex specific venous flow rates were observed as in the human [1]. In Expt 1 (7% vs 14%CP) the male feto-placental unit received greater blood flow than the female at 120dpc, this being achieved by decreased resistance (PI), increased velocity (TAMV) and increased diameter of the MUA. Throughout gestation the low protein in both the PERI conception and PREconception diet enhanced blood flow to the fetus by reducing resistance and increasing volume flow in the MUA. This can be best illustrated in Fig 6 where the ratio of BFV over gestation compared to the initial measure is shown. This was
associated with significantly increased velocity (TAMV) and MUA diameter (P > 0.05) with no difference in resistance (PI) over the treatment groups from 150dpc.

The effects of PERIconception low protein diet upon MUA blood flow was greater in those heifers with a male fetus at both 120 and 150dpc. This is concomitant with the data presented on the correlation between blood flow and birth weight, that is when both sexes are grouped together this relationship appears consistent (P = 0.01), however when we consider this by sex the correlation between blood flow and birth weight in the male calf exists only if the heifer

Fig 11. Arterial Blood Pressure according to PERIconception diet. Arterial blood pressure (mean ± SEM) in male (left) and female (right) calves of heifers fed Low (7% CP □; n = 28) or High (14% CP ●; n = 36) PERIconception diet from 60 days prior conception to 23 days post conception in Experiment 1. * denotes P ≤ 0.05.

doi:10.1371/journal.pone.0125694.g011

Gestational Dietary Protein Uterine Blood Flow and Fetal Development
received a low protein diet either PREconception or PERIconception ($P = 0.004$). In the female calves this correlation only exists if their dams had received a high protein diet in the PERI and PREconception period ($P = 0.03$). This suggests that dietary perturbations may have different effects upon uterine hemodynamics and fetal development in utero dependent upon sex as previously reported [1,7,20].

Contrary to one recent report [29] nutrient restriction altered uterine blood flow from 120dpc to 210dpc. This apparent disparity may be caused by the larger numbers in this study, consideration here of the sex specific effects, the difference in dietary treatment (protein versus total nutrients), the use of only primiparous heifers in this study and the period of dietary perturbation. The effects of nutrient restriction upon fetal and placental growth in the primiparous compared to the mature female are significantly different [30] as the hormonal response of the mature female signals greater nutrients to the fetus than in the immature primiparous female.

The in utero effects upon fetal growth and blood supply in Expt 1 produced corresponding effects upon the cardiovascular system in the neonate and young offspring with neonatal heart rate being increased in the female neonate with six month old female progeny having decreased systolic blood pressure. This study supports previous work [13] that showed heart length and width was reduced by nutrient restriction in female fetus at 94dpc and that aortic diameter was increased [13]. This important study by Mossa et al. [13] showed that low protein postconception diet increased systolic pressure in the neonatal calves. PERIconception diet in the current study however decreased systolic pressure at six months of age. Interestingly, the low protein diet (7% vs 14%) significantly reduced fetal heart weight in the female compared to the male fetus at 98dpc.

The relationship between blood flow and resistance indices in the two arteries has previously been published in beef cows [26]. Due to the large number of animals in this study ($n = 360$ in Expt 1 and $n = 188$ in Expt 2), only the ipsilateral artery to the fetus was assessed by Doppler, as it is known to have a consistent relationship with the increased blood flow in the contralateral artery, although with an average four-fold increase in blood flow, compared to the contralateral artery during pregnancy [3].

In humans, when uterine artery resistance does not decrease during pregnancy there is an association with deficiency in nutrient supply [2]. This study shows that low protein diet decreased PI at 120dpc, presumably increasing blood supply and nutrients to the placenta and fetus. This was further enforced by the increased TAMV in these animals. In previous studies, we reported an increase in placental size and function (as measured by higher estrone sulfate (ES)) following a low protein diet during early pregnancy [27,31]. We believe that a similar effect of low protein diet may have occurred in the study reported herein as ES may be rapidly converted to estradiol which in turn may increase uterine blood flow via vasodilation [32]. However, this will need to be studied further in the serum samples collected during the current experiments.

Placental growth from early to midgestation in the cow is much greater than fetal growth and continues to increase exponentially until the last month of pregnancy, when growth slows down [33]. On the contrary, fetal growth during the last trimester is five times that of the placenta, therefore angiogenesis and vascularization must increase to enable sufficient nutrients to be supplied. This pattern of placental and fetal development is similar to the human but dissimilar to the sheep where placental growth plateaus in midgestation [34]. We have previously shown that vasculature in the placenta at term increased following protein restriction in the first trimester [27], similarly Vonnahme et al. [35] showed that the placental vasculature increased at term after midgestation restriction followed by re-alimentation.

It is important to note that after nutrient restriction in this study there was no attempt to re-aliment the heifers post dietary restriction. In Expt 1 heifers were individually fed for the entire...
gestation to restrict such possible variation. The use of only primiparous heifers of restricted breed type, mated to a single bull, reduced possible uterine variation caused by prior pregnancies and variations in fetal demand.

Conclusions

Protein restriction during the peri-conception period and first trimester altered uterine blood flow in primiparous heifers. Further, uterine blood flow was increased in early pregnancy if the heifer was pregnant to a male rather than a female fetus. Sex specific differences in feto-placental perfusion indices exist in the bovine as previously reported in the human [1] and such indices are susceptible to dietary perturbation. The identification of these effects adds to our knowledge of the physiology of male and female fetuses in utero. In accord with a previous report [13] the cardiovascular system of the female fetus was altered by the early in utero low protein diet. Fetal organ development has been shown to be affected by changes in maternal nutrition associated with changes in placental hormone production, with such effects having a sex bias. Altered fetal haemodynamics in utero may be part of the mechanism involved, making further investigation of the sex specific differences present during fetal life important. This study affords additional insight into the gender specific circulatory differences that exist through gestation.

Acknowledgments

We acknowledge the invaluable assistance of Ray Cranney in all animal husbandry procedures and Nicholas Hatzirodios, Katja Hummitzsch, Wendy Bonner in the collection of samples.

Author Contributions

Conceived and designed the experiments: VP ICM. Performed the experiments: VP AH WW KC TK RG JHM. Analyzed the data: GMP JHM KC VP. Contributed reagents/materials/analysis tools: VP RR TK. Wrote the paper: VP JHM GMP. Drafted the article or revised it critically for important intellectual content: KC RG VP JHM GMP. Gave final approval of the version to be published: ICM KC RG VP JHM GMP AH WW TK RR. Agreed to be accountable for all aspects of the work: ICM KC RG VP JHM GMP AH WW TK RR.

References

1. Prior T, Wild M, Mullins E, Bennett P, Kumar S (2013) Sex Specific Differences in Fetal Middle Cerebral Artery and Umbilical Venous Doppler. PLoS ONE 8: e56933. doi:10.1371/journal.pone.0056933 PMID: 23437275
2. Clifton VL (2010) Review: Sex and the human placenta: mediating differential strategies of fetal growth and survival. Placenta 31 Suppl: S33–39. doi: 10.1016/j.placenta.2009.11.010 PMID: 20004469
3. Bollwein H, Baumgartner U, Stolla R (2002) Transrectal Doppler sonography of uterine blood flow in cows during pregnancy. Theriogenology 57: 2053–2061. PMID: 12066865
4. Herzog K, Bollwein H (2007) Application of Doppler ultrasonography in cattle reproduction. Reprod Domest Anim 42 Suppl 2: 51–58. PMID: 17688602
5. Micke GC, Sullivan TM, Soares Magalhaes RJ, Rolls PJ, Norman ST, Perry VEA (2010) Heifer nutrition during early- and mid-pregnancy alters fetal growth trajectory and birth weight. Animal Reproduction Science 117: 1–10. doi: 10.1016/j.anireprosci.2009.03.010 PMID: 19394770
6. Long NM, Vonnahme KA, Hess BW, Nathanielisz PW, Ford SP (2009) Effects of early gestational undernutrition on fetal growth, organ development, and placental composition in the bovine. Journal of Animal Science 87: 1950–1958. doi: 10.2527/jas.2008-1672 PMID: 19213703
7. Copping KJ, Hoare A, Callaghan M, McMillen IC, Rodgers RJ, Perry VEA (2014) Fetal programming in 2-year-old calving heifers: peri-conception and first trimester protein restriction alters fetal growth in a gender-specific manner. Animal Production Science 54: 1333–1337.
8. Sullivan TM, Micke GC, Greer RM, Irving-Rodgers HF, Rodgers RJ, Perry VEA (2009) Dietary manipulation of Bos indicus x heifers during gestation affects the reproductive development of their heifer calves. Reproduction, Fertility and Development 21: 773–784. doi: 10.1071/RD09004 PMID: 19967220

9. Sullivan TM, Micke GC, Greer RM, Perry VEA (2010) Dietary manipulation of Bos indicus x heifers during gestation affects the prepubertal reproductive development of their bull calves. Animal Reproduction Science 118: 131–139. doi: 10.1016/j.anireprosci.2009.07.005 PMID: 19671489

10. Micke GC, Sullivan TM, Gatford KL, Owens JA, Perry VE (2010) Nutrient intake in the bovine during early and mid-gestation causes sex-specific changes in progesterone plasma IGF-I, liveweight, height and carcass traits. Anim Reprod Sci 121: 208–217. doi: 10.1016/j.anireprosci.2010.05.017 PMID: 20591585

11. Micke GC, Sullivan TM, McMillen IC, Gentili S, Perry VEA (2011) Heifer nutrient intake during early- and mid-gestation programs adult offspring adiposity and mRNA expression of growth-related genes in adipose depots. Reproduction 141: 697–706. doi: 10.1530/REP-10-0332 PMID: 21310814

12. Micke GC, Sullivan TM, McMillen IC, Gentili S, Perry VEA (2011) Protein intake during gestation affects postnatal bovine skeletal muscle growth and relative expression of IGFI, IGFIIR, IGFI and IGFIIR. Molecular and Cellular Endocrinology 332: 234–241. doi: 10.1016/j.mce.2010.10.018 PMID: 21056085

13. Mossa F, Carter F, Walsh SW, Kenny DA, Smith GW, Ireland JLH, et al. (2013) Maternal Undernutrition in Cows Impairs Ovarian and Cardiovascular Systems in Their Offspring. Biology of Reproduction 88: 92, 91–99. doi: 10.1095/biolreprod.110.107235 PMID: 23426432

14. Wu G, Bazer FW, Wallace JM, Spencer TE (2006) BOARD-INVITED REVIEW: Intrauterine growth retardation: Implications for the animal sciences. Journal of Animal Science 84: 2316–2337. PMID: 16908634

15. Ashworth CJ, Toma LM, Hunter MG (2009) Nutritional effects on oocyte and embryo development in mammals: implications for reproductive efficiency and environmental sustainability. Philosophical Transactions of the Royal Society B: Biological Sciences 364: 3351–3361. doi: 10.1098/rstb.2009.0184 PMID: 19833647

16. Borowczyk E, Caton JS, Redmer DA, Bliski JJ, Weigl RM, Vonnahme KA, et al. (2006) Effects of plane of nutrition on in vitro fertilization and early embryonic development in sheep. Journal of Animal Science 1599: 1599. PMID:16699117

17. Bermejo-Alvarez P, Rizos D, Lonergan P, Gutierrez-Adan A (2011) Transcriptional sexual dimorphism during preimplantation embryo development and its consequences for developmental competence and adult health and disease. Reproduction 141: 563–570. doi: 10.1530/REP-10-0482 PMID: 21339284

18. Dobbs KB, Rodriguez M, Sudano MJ, Hansen PJ (2013) Dynamics of DNA methylation during early development of the preimplantation bovine embryo. PLoS One 8: e66230. doi: 10.1371/journal.pone.2009.08.010 PMID: 21903264

19. Kwong WY, Wild AE, Roberts P, Willis AC, Flemming TP (2000) Maternal undernutrition during the preimplantation period of rat development causes blastocyst abnormalities and programming of postnatal hypertension. Development 127: 4195–4202. PMID: 10976051

20. Micke G, Hernandez J, Sullivan TM, Kennaway D, Perry VEA (2014) Maternal endocrine adaptation throughout pregnancy to nutrient manipulation: consequences for sexually dimorphic programming of thyroid hormones, growth and appetite of their progeny. Theriogenology accepted.

21. Micke GC, Sullivan TM, Rolls PJ, Hasell B, Greer RM, Norman ST, et al. (2010) Dystocia in 3-year-old beef heifers; Relationship to maternal nutrient intake during early- and mid-gestation, pelvic area and hormonal indicators of placental function. Animal Reproduction Science 118: 163–170. doi: 10.1016/j.anireprosci.2009.08.006 PMID: 19762178

22. O’Connell BA, Moritz KM, Walker DW, Dickinson H (2013) Sexually dimorphic placental development throughout gestation in the spiny mouse (Acomys cahirinus). Placenta. 34: 119–126. doi: 10.1016/j.placenta.2012.11.009 PMID: 23260227

23. Stark MJ, Hodyl NA, Wright IM, Clifton VL (2011) Influence of sex and glucocorticoid exposure on pre-term placental pro-oxidant-antioxidant balance. Placenta 32: 865–870. doi: 10.1016/j.placenta.2011.08.010 PMID: 21903264

24. MacLaughlin SM, Walker SK, Kleemann DO, Sibbons JP, Tosh DN, Gentili S, et al. (2007) Impact of Periconceptional Undernutrition on Adrenal Growth and Adrenal Insulin-Like Growth Factor and Steroidogenic Enzyme Expression in the Sheep Fetus during Early Pregnancy. Endocrinology 148: 1911–1920. PMID: 17194740

25. Bridges GA, Mussard ML, Helser LA, Day ML (2014) Comparison of follicular dynamics and hormone concentrations between the 7-day and 5-day CO-Synch + CIDR program in primiparous beef cows. Theriogenology 81: 632–638. doi: 10.1016/j.theriogenology.2013.11.020 PMID: 24388673
26. Panarace M, Garmil C, Marfil M, Jauregui G, Lagioia J, Luther E, et al. (2006) Transrectal Doppler sonography for evaluation of uterine blood flow throughout pregnancy in 13 cows. Theriogenology 66: 2113–2119. PMID: 16876855

27. Perry VEA, Norman ST, Owen JA, Daniel RCW, Phillips N (1999) Low dietary protein during early pregnancy alters bovine placental development. Animal Reproduction Science 55: 13–21. PMID: 10099675

28. Sullivan TM, Micke GC, Magalhaes RS, Phillips NJ, Perry VEA (2009) Dietary protein during gestation affects placental development in heifers. Theriogenology 72: 427–438. doi: 10.1016/j.theriogenology.2009.03.018 PMID: 19540576

29. Camacho LE, Lemley CO, Van Emon ML, Caton JS, Swanson KC, Vonnahme KA (2014) Effects of maternal nutrient restriction followed by realimentation during early and midgestation on beef cows. I. Maternal performance and organ weights at different stages of gestation. Journal of Animal Science 92: 520–529. doi: 10.2527/jas.2013-7017 PMID: 24664560

30. Aitken RP, Milne JS, Wallace JM (2007) Wide variations in gestational dietary intake differentially impact on pregnancy outcome in young adolescents and influence early postnatal offspring phenotype. Pediatric Research 62: 385(Abstract).

31. Sullivan TM, Micke GC, Magalhaes RS, Martin GB, Wallace CR, Green JA, et al. (2009) Dietary protein during gestation affects circulating indicators of placental function and fetal development in heifers. Placenta 30: 348–354. doi: 10.1016/j.placenta.2009.01.008 PMID: 19233467

32. Ferrell C (1991) Maternal and fetal influences on uterine and conceptus development in the cow: II. Blood flow and nutrient flux. Journal of Animal Science 69: 1954–1965. PMID: 2066304

33. Reynolds LP, Redmer DA (1995) Utero-placental vascular development and placental function. Journal of Animal Science 73: 1839–1851. PMID: 7545661

34. Redmer DA, Wallace JM, Reynolds LP (2004) Effect of nutrient intake during pregnancy on fetal and placental growth and vascular development. Domestic Animal Endocrinology 27: 199–217. PMID: 15451070

35. Vonnahme KA, Zhu MJ, Borowicz PP, Geary TW, Hess BW, Reynolds LP, et al. (2007) Effect of early gestational undernutrition on angiogenic factor expression and vascularity in the bovine placentome. J Anim Sci 85: 2464–2472. PMID: 17565057