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Published in: Placenta

DOI (link to publication from Publisher): 10.1016/j.placenta.2018.06.309

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Publication date: 2018

Document Version Accepted author manuscript, peer reviewed version

Link to publication from Aalborg University

Citation for published version (APA):

Thunbo, M. Ø., Sinding, M., Bogaard, P., Korsager, A. S., Frøkjær, J. B., Østergaard, L. R., Petersen, A., & Sørensen, A. (2018). Postpartum placental CT angiography in normal pregnancies and in those complicated by diabetes mellitus. *Placenta*, *69*, 20-25. https://doi.org/10.1016/j.placenta.2018.06.309

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Accepted Manuscript

Postpartum placental CT angiography in normal pregnancies and in those complicated by diabetes mellitus

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PII: S0143-4004(18)30183-8

DOI: 10.1016/j.placenta.2018.06.309

Reference: YPLAC 3844

To appear in: *Placenta*

Received Date: 24 April 2018

Revised Date: 11 June 2018

Accepted Date: 28 June 2018

Please cite this article as: Thunbo MetteØ, Sinding M, Bogaard P, Korsager AS, Frøkjær JensBrø, Østergaard LR, Petersen A, Sørensen A, Postpartum placental CT angiography in normal pregnancies and in those complicated by diabetes mellitus, *Placenta* (2018), doi: 10.1016/j.placenta.2018.06.309.

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- ¹ Postpartum placental CT angiography in normal
- ² pregnancies and in those complicated by
- 3 diabetes mellitus

4	Running headline: Placental CT angiography in diabetic pregnancies
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26	Word count, abstract: 250
27	Word count, main text: 2498
28	
29	Declarations of interest: None
30	

31 Abstract

32 Introduction

- 33 Pregnancy complicated by diabetes mellitus (DM) is a central obstetric problem often complicated by fetal
- 34 macrosomia and increased risk of intrapartum asphyxia. This risk might be explained by fetoplacental
- 35 vascular abnormalities. This study aimed to investigate the fetoplacental vascular volume by placental CT
- 36 angiography in normal pregnancies and in pregnancies complicated by type 1 DM (T1DM), diet controlled
- 37 gestational DM (GDMd), and insulin treated gestational DM (GDMi).

38 Methods

- 39 Postpartum, barium contrast enhanced placental CT angiography was performed in 27 normal pregnancies
- 40 and 25 DM pregnancies (8 T1DM, 8 GDMd, and 9 GDMi). The fetoplacental vascular volume/placenta
- 41 weight (FVV/PW)-ratio and fetoplacental vascular volume/birth weight (FVV/BW)-ratio of each diabetic
- 42 group were compared to the normal group with multiple regression analysis adjusted for GA. In all
- 43 pregnancies a standardized histopathological placental examination was performed postpartum.

44 Results

- 45 In normal pregnancies, the fetoplacental vascular volume increased with GA (*p*<0.001), placental weight
- 46 (p<0.001), and birth weight (p<0.001). In T1DM and GDMi pregnancies, the gestational age adjusted
- 47 placental weight and the birth weight were increased when compared to normal pregnancies (*p*<0.05). The
- 48 FVV/BW-ratio was significantly reduced in both T1DM and GDMi pregnancies when compared to normal
- 49 pregnancies (*p*=0.003 and *p*=0.009, respectively).

50 Discussion

- 51 This study demonstrates, that in insulin treated DM pregnancies the fetus as well as the placenta is larger
- 52 than normal. However, despite a large placenta, a relatively smaller fetoplacental vascular volume supplies
- 53 the macrosomic fetus. This finding might explain why fetuses from insulin treated DM pregnancies have
- 54 high vulnerability to intrauterine and intrapartum asphyxia.

55 Keywords

56 Diabetes mellitus, placenta, CT angiography, vasculature, fetoplacental vascular volume, gestational
 57 diabetes

58 Abbreviations

- 59 Computed tomography angiography (CTA), diabetes mellitus (DM), diet controlled gestational diabetes
- 60 mellitus (GDMd), fetoplacental vascular volume/birth weight-ratio (FVV/BW-ratio), fetoplacental vascular
- 61 volume/placenta weight-ratio (FVV/PW-ratio), gestational age (GA), insulin treated gestational diabetes
- 62 mellitus (GDMi), magnetic resonance angiography (MRA), and type 1 diabetes mellitus (T1DM).

63 Introduction

Pregnancies complicated by diabetes mellitus (DM), both type 1 DM (T1DM) and gestational DM (GDM),
 are a central obstetric challenge, as the fetal and maternal morbidity and perinatal mortality is high[1]. It is

- 66 well described that DM pregnancies are associated with neonatal complications such as fetal macrosomia,
- perinatal asphyxia, and metabolic syndrome in later life[1]. The increased risk of intrauterine and
 intrapartum asphyxia in pregnancies complicated by DM may partly relay on the increased metabolic
- 69 demand of the macrosomic diabetic fetus and a decreased transplacental oxygen transfer capacity due to
- altered oxygen binding capacity of hemoglobin[2]. However fetoplacental vascular abnormalities related to
- 71 DM may also contribute to the increased risk[3–5].
- It is known that DM pregnancies are associated with increased placental weight and birth weight and
 an increased birth weight/placental weight-ratio[6]. Current knowledge on the fetoplacental vasculature in
 DM pregnancies is based on macroscopic examinations[7], histomorphometry[3,8–16], stereology[4,17–
- 75 20], x-ray angiograms[21], and measurements of the placental residual blood volume after birth[22]. In
- 76 T1DM pregnancies conflicting results are demonstrated as some studies describe an increased
- fetoplacental vascular volume, surface area, and capillary length compared to normal[3,4,8,9,17–19,22],
- 78 while others describe decreased vessel diameter and number of vessels[7,10–15,21]. Also in GDM
- 79 pregnancies, existing knowledge on the fetoplacental vasculature demonstrates conflicting results with
- 80 studies reporting increased vascular volume[15], surface area[15], and number of vessels[12,23] as well as
- 81 decreased number of vessels[20,24]. The inconsistent findings in the literature may be explained by
- differences in glycemic control, treatment regime, and lack of methods to demonstrate vascularpathology[25].
- Imaging technologies such as placental computed tomography angiography (CTA) and magnetic resonance angiography (MRA) have a great potential to investigate the fetoplacental vasculature in threedimensions (3D). By using these methods the fetoplacental vasculature has been investigated in normal pregnancies[26–29], however to the best of our knowledge CTA has never been performed in pregnancies complicated by different types of DM.
- To improve our understanding of the perinatal risk of asphyxia associated with DM, a better knowledge of the fetoplacental vasculature is essential. Therefore this study aimed to investigate the fetoplacental vascular volume by using postpartum 3D placental CTA in normal pregnancies in comparison to pregnancies complicated by DM (T1DM, diet controlled GDM (GDMd), and insulin treated GDM (GDMi)).
- 93 Methods
- 94Twenty-five placentas (35-41 weeks' gestation) from singleton pregnancies complicated by DM (8 T1DM95placentas, 8 GDMd placentas, and 9 GDMi placentas) were included in the study[30,31]. 32 placentas (30-
- 96 42 weeks' gestation) from normal singleton pregnancies constituted the control group. We excluded
- stillbirths, abnormal fetal karyotype or congenital malformations, and pregnancies with clinical signs of
 placental insufficiency (umbilical artery Doppler flow Pulsatility index (PI) Z-score ≥ 2[32], cerebroplacental
- 98 placental insufficiency (umbilical artery Doppler flow Pulsatility index (PI) Z-score \geq 2[32], cerebroplacental 99 Doppler ratio Z-score \leq -2[33] and birth weight \leq - 22%[34]. All placentas were collected at Aalborg
- 100 University Hospital, Denmark, between July 1st, 2015 and December 1st, 2016. The Danish National Ethics
- 101 Committee (N-20150018) and the Danish Data Protection Agency (2008-58-0028) approved the study, and
- 102 all participants gave oral and written informed consent. Maternal and pregnancy characteristics are

presented in Table 1. Data were collected from medical records and the electronic ultrasound databaseAstraia version 1.24.7 (Astraia Software Gmbh, Munich, Germany).

105 Just after delivery, the placentas were stored at -5°C, and on the day of CTA the placenta was thawed 106 in a warm water bath (37°C). The umbilical cord vessels were cannulated 5cm from the umbilical cord 107 insertion using 3 venous cannulas size 1.3x32mm (BD Venflon Pro, Helsingborg, Sweden). The placenta was 108 flushed with a saline 9mg NaCl/ml and Heparin 4.5IE/ml (Leo Pharma A/S, Ballerup, Denmark) solution until 109 the venous efflux was clear. Hereafter a heated (<40°C) contrast mixture of gelatin 0.05g/ml (Urtegaarden 110 Djursland, Allingåbro, Denmark), barium sulphate 0.17g/ml (E-Z Em Inc, Westbury, NY, USA), and saline 111 9mg NaCl/ml was injected with a hand syringe. When the contrast mixture appeared in the venous efflux, 112 the vein was plugged, and injection was continued until resistance was felt. Hereafter the placenta was 113 cooled on ice to set the gelatin solution (Figure 1 (A and C)).

114 CTA was performed on a 128-slice Siemens SOMATOM Definition Flash scanner (Siemens Healthcare 115 GmbH, Erlangen, Germany) with software version VA48A and the flowing parameters: 0.6mm slice 116 thickness, 0.4mm increment, 1° pitch, 140kV, effective 200mAs, and 1sec rotation time. Post processing 117 analysis was performed using the commercial software AW Server version 3.0 (GE Healthcare, Little 118 Chalfont, Great Britain) to calculate the fetoplacental vascular volume by computing the volume of all 119 voxels above 550HU (Figure 1 (B, D, and E)). All 3D reconstructions of the fetoplacental vasculature were 120 visually inspected. Five placentas were excluded from the normal group due to insufficient contrast filling 121 of the fetoplacental vessels. No placentas were excluded from the DM group.

After the CTA, a standardized postnatal histopathological examination according to the Amsterdam consensus guideline[35] was performed by experienced placental pathologists (PB and AP), who were blinded to the CTA vascular outcome, but not the clinical information. Selected diabetic histopathological findings are reported in Table 3 using the following references for placental weight[36], delayed villous maturation[35], and the umbilical cord[37].

127 In normal pregnancies, the association between the fetoplacental vascular volume and the following 128 variables; gestational age at birth (GA), placental weight, and birth weight, was investigated by linear 129 regression analysis. In each of the diabetic groups, the fetoplacental vascular volume, the fetoplacental 130 vascular volume/placental weight (FVV/PW)-ratio and the fetoplacental vascular volume/birth weight 131 (FVV/BW)-ratio was compared to the normal group by multiple linear regression adjusting for GA. *p*<0.05 132 was considered significant. All analyses were performed in SPSS Statistics version 25.0 (IBM, North Castle, 133 New York, USA).

134 **Results**

135 As demonstrated in Table 1, the placental weight and birth weight (given as Z-scores and hence corrected 136 for GA) were increased in pregnancies complicated by T1DM (p=0.026 and p<0.001, respectively) and in 137 GDMi (p=0.002 and p=0.003, respectively) pregnancies. In addition, the T1DM and GDMi groups had a 138 higher HbA_{1c} when compared to GDMd pregnancies, indicating poorer glycemic control in these diabetic 139 groups. Given the small number of patients in this study, the rare event of umbilical cord pH < 7 and Apgar 140 score < 7 five minutes postpartum was not apparent. However, there was a trend towards more caesarian 141 sections (elective and acute) among the patients with insulin dependent diabetes. 142 As illustrated in Figure 1, the 3D reconstruction of the segmented fetoplacental vascular volume

- 143 included both the chorionic vessels on the placental surface and the stem villi vessels that bend
- 144 perpendicularly to the placental surface, which further branches into intermediate villi vessels (Figure 1E).

The smallest vessels of the fetoplacental vascular tree (capillaries) were not included as a part of thecomputed fetoplacental vascular volume.

147 In normal pregnancies at term (GA 40+0) the fetoplacental vascular volume was 172.2ml (95% CI:

148 154.2-189.9 ml/kg), and we demonstrated a positive linear association between the fetoplacental vascular 149 volume and GA (r^2 =0.585, p<0.001), placental weight (r^2 =0.405, p<0.001), and birth weight (r^2 =0.499,

p < 0.001 (Figure 2). In the DM groups, the fetoplacental vascular volume did not differ from that in normal

151 pregnancies at equivalent GA.

In normal pregnancies at term (GA 40+0) the FVV/BW-ratio was 48.9 ml/kg (95% CI: 44.5-56.7ml/kg).
In all DM groups the FVV/BW-ratio was lower, however this difference was only significant in the insulin
dependent DM groups; T1DM (-16.2ml/kg, *p*=0.003), GDMi (-12.1ml/kg, *p*=0.009), and GDMd (-7.8ml/kg, *p*=0.198).

In normal pregnancies at term (GA 40+0) the FVV/PW-ratio was 33.9ml/kg (95% CI: 32.9-37.9ml/kg),
and this ratio was reduced in all DM groups, but the difference was only significant for pregnancies
complicated by GDMi (-81.5ml/kg, *p*=0.012) and with a strong trend in T1DM (-66.0ml/kg, *p*=0.068) (Table
2).

The placental histopathological examination is presented in Table 3. Characteristic diabetic
 abnormalities are seen predominantly in the insulin treated diabetic pregnancies, as two or more diabetic
 findings are seen in 12.5% of pregnancies complicated by T1DM and in 44.4% of pregnancies complicated

163 by GDMi.

164 **Discussion**

This study demonstrated that in normal pregnancies the fetoplacental vascular volume increased with GA, placental weight, and birth weight. In pregnancies complicated by T1DM and GDMi, the placental weight and birth weight was higher than normal. However, in these pregnancies the relative fetoplacental vascular volume was reduced, as demonstrated by a lower FVV/PW-ratio and FVV/BW-ratio. These findings indicate, that the large placenta in pregnancies complicated by insulin dependent diabetes has relatively fewer fetoplacental vessels, and therefore a relatively smaller fetoplacental volume supplies the macrosomic diabetic fetus. This finding might explain one of the underlying mechanisms why fetuses from insulin

172 dependent diabetic pregnancies are more vulnerable to asphyxia during pregnancy and labor.

173 This study had some methodological limitations. Proper placental preparation is crucial to obtain 174 reliable fetoplacental vascular assessments. To ensure sufficient contrast filling of the entire fetoplacental 175 vasculature without presence of intravascular blood clots, the placentas in this study were frozen 176 postpartum, which is known to reduce clotting without adverse effects on the fetoplacental 177 vasculature[38–40]. Furthermore, the thawed placentas were flushed with heparinized saline to remove 178 vascular clots prior to contrast injection. The fetoplacental vascular volume may have varied according to 179 the degree of vascular contrast filling injected by a hand syringe. However in this study, one person (MØT) 180 performed the contrast injection, and in each placenta contrast injection was performed uniformly until a 181 specific resistance was felt. To avoid contrast drainage we added gelatin to the contrast solution, and the 182 contrast-perfused placenta was cooled on ice packs to set the contrast solution.

Strength of our study was the ability to demonstrate the entire fetoplacental vasculature except for the smallest vessels (capillaries). The majority of previous studies is based on placental biopsies[3,9,14,15], which might not reflect the placental pathology of the entire placenta. It is well known that the normal placenta has a heterogeneous vessel maturation[41], and also the diabetic placenta is known to have focal

- 187 pathology such as dysmature villous structures[25]. Another strength of the present method was the
- 188 demonstration of the fetoplacental vasculature in 3D. 3D reconstruction provides information of spatial

vessel architecture such as the fetoplacental vascular volume, which is not available by 2D imaging[42]

- 190 which is the basis of the majority of previous studies [3,7,9,14,15,20,21]. Lastly, angiography allows for
- histopathological placental examination to be performed following the vascular investigation, which is not
- available with comparable methods such as corrosion casting.

A limitation of the study design was the relatively small study population. GDM pregnancies were grouped according to the White Classification of Diabetes in Pregnancy[43], but because of the small number of T1DM pregnancies in this study, this group was not further subdivided.

Strength of the study was the placental histopathological examination that confirmed, that placental pathology is predominantly seen in pregnancies complicated by insulin dependent diabetes. Another strength was the adjustment for GA in the analysis when comparing the fetoplacental vasculature between groups. This is of major importance, as the fetoplacental vascular volume is known to increase dramatically as pregnancy advances[44,45], which is also in accordance with the findings of our study.

201 In normal pregnancies, this study demonstrated that the normal fetoplacental vascular volume at 202 term is 172.2ml (95% CI: 154.2-189.9 ml/kg). As demonstrated by images of the 3D segmentation, the 203 vascular volume includes the chorionic vessels, the stem villi vessels, and the intermediate villi vessels. This 204 finding is very much in line with the previous published literature, where the normal fetoplacental vascular 205 volume is estimated to be 5-159ml based on residual placental blood-volume[22,46] or 12-124ml based on 206 MR-angiography[27], The normal fetoplacental vascular volume in this study is reassuring, as it indicates 207 that under-segmentation was not a major limitation of this study. Furthermore, a positive linear association 208 was found between the fetoplacental vascular volume and GA. This finding is in accordance with a previous 209 study based on stereological analysis of placental biopsies demonstrating a linear increase of stem and 210 intermediate villi vessel volume throughout pregnancy[47]. Furthermore, our study demonstrated a linear 211 correlation between the fetoplacental vascular volume and placental weight and birth weight. This finding 212 is in accordance with a previous study on placental MRA in normal pregnancies at term by Rasmussen et 213 al[27]. In contrast to our study, Rasmussen et al. were not able to demonstrate an association between 214 placental vascular volume and placental weight. However, this discrepancy may be explained by a different 215 range of GA included in the two studies, as our study included placentas at a wide range of GA (30-42 216 weeks' gestation) as compared to Rasmussen et al. who included term placentas.

We found an increased placental weight and birth weight in pregnancies complicated by insulin dependent DM, which is in line with previous literature[6]. However there is evidence to support that this pathology should also be found among diet dependent diabetes[48]. In this study, the GDMd group had a lower HbA_{1c} suggesting a good glycemic control, which may explain why this group did not demonstrate such pathology in our study.

Furthermore, we found that in all three DM groups the fetoplacental vascular volume did not differ from that in normal pregnancies. In previous studies, the fetoplacental vascular volume has been investigated by different methods, which could explain the rather conflicting results. By estimating the placental residual blood volume, Klebe et al[22] found an increased fetoplacental vascular volume in DM pregnancies when compared to normal pregnancies. However, another study by Singer et al[14] found no difference in the vascular volume and surface areas of the chorionic vessels and stem villi vessels when estimated by placental histological examination.

- In this study, intrauterine and intrapartum asphyxia, defined as umbilical blood pH < 7 and Apgar score < 7 after 5 minutes, was not present. This might be due to a very small number of patients included in the study, appropriate antenatal monitoring, and adequate obstetric intervention of the included diabetic pregnancies. In Table 1, it should be noted, that there is a trend towards a higher proportion of elective and acute cesarean section among the DM pregnancies.
- 234 The disparities in FVV/BW-ratio among insulin dependent and insulin independent DM pregnancies 235 in this study might be caused by differences in the placental pathophysiology. As fetoplacental vascular 236 abnormalities are hypothesized to result from hypoxia caused by fetal and maternal hyperglycemia[1], the 237 observed differences might indicate differences in time of onset and severity of hyperglycemia between 238 the different DM types and differences in preexisting maternal factors[49]. It is known that when DM is 239 well controlled during pregnancy, the fetoplacental vasculature is normally developed and the neonatal 240 outcome improved[25]. The measured HbA_{1c} values in this study is highest in T1DM both pre-gestational 241 and during pregnancy (Table 1), but time for onset of hyperglycemia in the GDM pregnancies cannot be 242 withdrawn. Future research is recommended to consider this.
- In conclusion, we have demonstrated that the placental weight and birth weight is increased in
 insulin dependent diabetic pregnancies, however the fetoplacental vasculature is not correspondingly
 increased. This study highlights that insulin dependent pregnancies are subject to relative placental
 insufficiency, as in these pregnancies the macrosomic fetus is supplied by a relatively smaller fetoplacental
 vascular volume. This finding might explain one of the underlying mechanisms, why fetuses from insulin
 dependent diabetic pregnancies are more vulnerable to asphyxia during pregnancy and labor.
- Acknowledgements: The authors are grateful to Pernille Veiss-Pedersen, radiologist, for competent
 assistance in performing the CTA. The author thanks Anne Staub Rasmussen, MD, for fruitful discussions
 regarding the placental preparation and the angiography method.
- Funding: This work was supported by The Faculty of Medicine at Aalborg University and by Aalborg
 University Hospital at The North Denmark Region.
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257 **Conflicts of interest:** The authors declare no conflict of interest.

Author's contribution: MØT, MS, and AS planned the research project. MØT collected all placentas and
 prepared it for scanning. JBF took part of planning the CT angiography. ASK, LRØ, and JBF participated in
 cooperation with MØT, MS, and AS in extracting the fetoplacental vascular volume based on the scannings.
 PB and AP performed all the placental pathological examinations and assessed the pathological data. MØT,
 MS, and AS performed the statistical analysis and data assessment and wrote the manuscript. All authors
 read and approved the final manuscript.

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396 Figure legends397

Figure 1: Images of a placenta from a normal pregnancy GA 37+4 (A and B) and a placenta from a pregnancy complicated by type 1
 diabetes mellitus, GA 37+1 (C and D). Macroscopic photography (A and C) and a 3D reconstruction of the fetoplacental blood
 vessels (B and D).

401

402 Figure 2: Association between fetoplacental vascular volume (ml) and A) gestational age at birth (weeks), B) placental weight (g),

403 and C) birth weight (g). The solid lines indicate ordinary least squares fit. The dashed lines indicate 95% confidence interval for the

404 normal placentas. Normal placentas (open circle), type 1 diabetes mellitus (T1DM) (squares), diet controlled gestational diabetes

 $405 \qquad {\rm mellitus} \ {\rm (GDMd)} \ {\rm (triangles), insulin treated gestational diabetes mellitus} \ {\rm (GDMi)} \ {\rm (pentagons).}$

Characteristics	Normal (N=27)	T1DM (N=8)		GDMd (N=8)		GDMi (N=9)	
			p-value		p-value		p-value
Maternal age at birth, yr	29.0 (27.0 – 34.0)	27.5 (26.5 – 31.0)	0.433	29.0 (27.0 - 34.8)	0.884	31.0 (27.0 – 36.5)	0.443
Pre-gestational maternal	24.1 (20.7 – 30.1)	29.1 (26.8 – 33.5)	0.027	26.6 (20.4-31.9)	0.550	28.2 (25.9 – 34.6)	0.086
BMI, kg/m ²					<i>Y</i>		
Nulliparity	48.1 (13/29)	37.5 (3/8)	0.700	50.0 (4/8)	1.000	33.3 (3/9)	0.700
Maternal smoking	14.8 (4/29)	0.0 (0/8)	0.553	37.5 (3/8)	0.312	22.2 (2/9)	0.627
Umbilical artery Doppler, Z- score[32]	0.3 (-0.6 – 0.9) (N=22)	0.2 (-0.3 – 1.6)	0.185	-0.1 (-0.6 – 0.4)	0.530	0.4 (-0.3 – 0.9)	0.362
Cerebroplacental-ratio, Z- score[33]	-0.5 (-0.7 – 0.2) (N=7)	-1.3 (-3.70.7) (N=4)	0.065	-1.1 (-1.3) (N=2)	0.283	0.0 (-1.5) (N=3)	0.922
Gestational age, weeks	38.9 (34.3 - 40.6)	36.9 (35.9 – 37.1)	0.192	40.6 (39.4 - 41.1)	0.104	37.6 (36.7 - 38.2)	0.295
Birth weight, Z-score[34]	-0.4 (-0.7 – 0.5)	2.2 (1.5–3.2)	<0.001	0.0 (-0.1 – 2.2)	0.286	1.1 (0.2 – 3.2)	0.003
Placental weight, Z-score	-0.1 (-0.4 - 0.4)	0.5 (0.0 - 1.9)	0.026	-0.2 (-0.9 – 0.8)	0.784	1.5 (0.4 – 2.3)	0.002
Umbilical venous vessel pH	0.0 (0/27)	0.0 (0/8)	- \	0.0 (0/8)	-	0.0 (0/9)	-
< 7.00							
Apgar score < 7 after 5 min	0.0 (0/27)	0.0 (0/8)	-	0.0 (0/8)	-	0.0 (0/9)	-
Delivery mode)				
 Vaginal birth 	• 70.4 (19/27)	• 37.5 (3/8)	0.116	• 75.0 (6/8)	1.000	• 33.3 (3/9)	0.111
 Elective cesarean section 	• 7.4 (2/27)	• 25.0 (2/8)	0.218	• 12.5 (1/8)	0.553	• 33.3 (3/9)	0.088
Acute cesarean section	• 22.2 (6/27)	• 37.5 (3/8)	0.396	• 12.5 (1/8)	1.000	• 33.3 (3/9)	0.660
Shoulder dystocia	0.0 (0/27)	0.0 (0/8)	-	0.0 (0/8)	-	0.0 (0/9)	-
Vacuum delivery	11.1 (3/27)	0.0 (0/8)	1.000	0.0 (0/8)	1.000	0.0 (0/9)	0.558
Postpartum bleeding	14.8 (4/27)	0.0 (0/8)	0.559	12.5% (1/8)	1.000	0.0 (1/9)	0.553
>500ml							
Maternal age at DM debut,	-)	12.5 (3.0 – 17.8)	-	29.0 (27.0 - 34.8)	-	31.0 (27.0 – 36.5)	-
yr							
Gestational age at debut,	-	-	-	30.9 (28.8 – 33.2)	-	28.6 (21.0 - 30.6)	-
weeks							
HbA _{1c} , mmol/mol	-		-		-		-

406 Table 1: Maternal, placental, and neonatal characteristics of included patients.

 Pre-gestational 		• 58.0 (51.25 – 90.0)		• 31.5 (28.8-35.8)		• 39.0 (36.0 – 45.5)	
(T1DM) or pre-							
treatment (GDM)							
•1 st trimester		• 55.0 (46.8 – 67.0)		• -	_	• -	
•2 nd trimester		• 50.0 (42.5 – 60.3)		• 37.0 (N=1)		• 37.0 (33.8 – 38.0)	
					Y	(N=4)	
•3 rd trimester		• 52.0 (49.5 – 57.8)		• 32.0 (29.5 – 38.5)		• 41.0 (34.0 – 46.5)	
White classification[43],	-	C: 50.0 (4/8)	-	A1: 8/8	-	A2: 9/9	-
class		D: 25.0 (2/8)					
		R: 25.0 (2/8)					

407 Data are given as median (IQR) or % (n/N). All p-values indicate comparison to normal pregnancy. The continuous data are analyzed with independent samples Student t-test or Mann-

Inal pregnaile, data are analyzed withs. 408 Whitney U-test based on the appearance of normal distribution, and categorical data are analyzed with Fishers exact test. GDMd = diet controlled gestational diabetes mellitus, GDMi

409 = insulin treated gestational diabetes mellitus, T1DM = type 1 diabetes mellitus.

410

411

Table 2: Summary of multiple regression analysis. The normal pregnancies are the reference.

	Unstandardi	zed coefficients	<i>p</i> -value	95% confidence interval for B	
	В	Std. Error		Lower bound	Upper bound
Fetoplace	ental vascular	volume (ml)			
T1DM	-14.3	15.2	0.356	-45.3	16.8
GDMd	-18.6	19.9	0.357	-59.3	22.0
GDMi	-10.9	13.9	0.440	-39.2	17.4
Fetoplace	ental vascular	volume / placent	al weight rat	io (ml/kg)	
T1DM	-66.0	34.9	0.068	-137.2	5.2
GDMd	-48.8	40.0	0.231	-130.2	32.6
GDMi	-81.5	30.6	0.012	-143.7	-19.3
Fetoplace	ental vascular	volume / birth wo	eight ratio (n	nl/kg)	·
T1DM	-16.2	49.5	0.003	-26.3	-6.1
GDMd	-7.8	5.9	0.198	-19.8	4.3
GDMi	-12.1	4.4	0.009	-21.0	-3.3

GDMd = diet controlled gestational diabetes mellitus, GDMi = insulin treated gestational diabetes mellitus, T1DM = type 1 diabetes mellitus.

Table 3: Diabetic placental histopathological findings.

Placenta pathology \ Group	Normal	T1DM	GDMd	GDMi
Large placental weight, p>90 [36]	14.8 (4/27)	37.5 (3/8)	12.5 (1/8)	55.6 (5/9)
Delayed villous maturation [35]	0.0 (0/27)	25.0 (2/8)	0.0 (0/8)	33.3 (3/9)
Long umbilical cord [37]	22.2 (6/27)	12.5 (1/8)	12.5 (1/8)	33.3 (3/9)
≥ 2 diabetic findings	7.4 (2/27)	12.5 (1/8)	0.0 (0/8)	44.4 (4/9)

Data are given as % (n/N). GDMd = diet controlled gestational diabetes mellitus, GDMi = insulin treated gestational diabetes mellitus, T1DM = type 1 diabetes mellitus.





Highlights

- Pregnancies complicated by diabetes (DM) are at high risk of intrapartum asphyxia.
- This risk may be related to fetoplacental vascular abnormalities.
- This study examines the fetoplacental vascular volume by placental CT angiography.
- In insulin dependent DM the fetoplacental vascular volume/birthweight-ratio is low.
- This finding might partly explain the higher vulnerability to fetal asphyxia.

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