Fetal growth restriction (FGR), conventionally defined as an ultrasound estimated fetal weight less than the 10th centile, is associated with an increased risk of perinatal morbidity and mortality as well as an increased risk of adult diseases. Many fetuses identified as less than the 10th centile, however, are not pathologically small, but rather constitutionally small for gestational age (SGA). Additionally, due to the inherent error of ultrasound estimation of fetal weight, which may be as great as 25%, fetuses that are appropriate for gestational age (AGA) can be misclassified as FGR. Constitutionally SGA and misclassified AGA fetuses are not at higher risk of antenatal or postnatal complications. They are, however, subject to the cost and morbidity of invasive testing, antenatal monitoring, and iatrogenic preterm delivery. Given this, identifying potential biomarkers that can distinguish these groups of fetuses would have clinical and cost benefits.

Transposable elements are a class of mobile genetic elements that have been estimated to comprise half of the human genome. Retroelements (REs), a subset of transposable elements, originate from retroviruses, integrate into the germline and are thus transmitted to all the cells of the host. REs can cause insertional mutagenesis or other adverse effects and are...
often suppressed in somatic tissues by epigenetic modifications, including DNA methylation.4

Interestingly, REs are often hypomethylated and highly expressed in the placenta. Furthermore, some REs have been co-opted to perform essential functions in the placenta.5,6 For example, Syncytin-1, encoded by ERVV-W-1 (SYN1), and Syncytin-2, encoded by ERVRD-1 (SYN2), have intact env genes that have evolved to mediate cell-to-cell fusion in the placenta to form the syncytiotrophoblast.7 Mice that lack expression of Syncytin-A (the murine orthologue of Syncytin-1), die between 11.5 and 13.5 days of gestation due to failure of the syncytiotrophoblast layer to form.8,9 Limited studies from human pregnancy suggest these gene products are also important in the human placenta. Altered expression and methylation patterns are associated with growth discordance in twin pregnancies and FGR or other placental syndromes in singleton pregnancies.10–14 As a mutable epigenetic mark, methylation patterns may be of particular interest because they are more likely than fixed genetic marks to reflect the environmental circumstances that may predispose to FGR and other placental syndromes.

Given the importance of Syn1 and Syn2 to placental function, our primary objective was to determine if methylation and expression patterns of SYN1 and SYN2 differed from FGR placentas compared with SGA placentas. We hypothesized that expression or methylation differences in Syncytin-1 and Syncytin-2, would plausibly distinguish pathologic FGR from constitutional SGA.

Materials and Methods

Placental Biopsies

Samples were obtained from a placental biopsy biobank that is maintained at the Magee-Womens Research Institute. A trained research nurse obtained at least two placental biopsy samples immediately after delivery. One sample was snap frozen in liquid nitrogen and the other was placed in RNAlater (Qiagen, Hilden, Germany) and stored at –80°C. A chart abstraction was performed at the time of collection and entered into a de-identified database linked to the samples. The University of Pittsburgh Institutional Review Board under project number PRO08050177 approved specimen collection.

Subject Selection Criteria

Using the Magee Biobank database, placental biopsy samples were selected from singleton pregnancies delivered after 36 weeks gestation. Women with diabetes mellitus or those carrying fetuses with suspected anomalies or aneuploidy were excluded. For analysis, subjects were divided into AGA (n = 10), SGA (n = 9), and FGR (n = 7) groups. AGA was defined by birth weight >10th centile and <90th centile using the Alexander growth reference.15,16 Subjects in both the SGA and FGR groups had birth weight ≤10th centile. Subjects that were categorized in the FGR group had antenatal evidence of uteroplacental insufficiency, defined as oligohydramnios, decreased fetal movement, or abnormalities in the biophysical profile, nonstress testing, contraction stress testing, or umbilical artery Doppler waveform.

Identifying Candidate Genes

The main target genes for this study were SYN1 and SYN2. To explore how SYN1 and SYN2 compare with other REs present in the placenta, we identified other REs for analysis that are expressed in the placenta or have placental-specific RE-derived regulatory regions. REs in the former category included endogenous retrovirus group 3 (ERV-3), paternally expressed 10 (PEG10) and retrotransposon-like 1 (RTL1), while the latter included leptin (LEP), endothelin receptor B (EDNRB), aromatase (CYP19A1), early placenta insulin-like peptide (INSL4), midline-1 (MID1), and pleiotrophin (PTN).5 Expression analysis was performed in all of these and methylation assessment performed on SYN1, SYN2, PEG10, and PTN.

Real-time Quantitative Polymerase Chain Reaction

Each sample was mechanically homogenized and digested in TRIzol followed by chloroform extraction and 100% ethanol precipitation. RNA was transferred to silica spin-columns (Epoch Biotechnologies, Missoula, MT) for column RNase-free DNase treatment (Qiagen, Hilden, Germany) and washing. The RNA pellet was suspended in RNAse-free water. Quantification and purity testing of the eluted RNA was performed by spectrophotometric analysis at OD260 and OD280 with the NanoDrop 1000 and by gel electrophoresis. Samples of 8 AGA, 6 SGA, and 4 FGR had high quality RNA for expression analysis. Complementary DNA was prepared using Applied Biosystems’ high-capacity RNA-to-cDNA™ kit (Thermo Scientific, Waltham, MA) per the manufacturer’s instructions. Primers were identified using the Massachusetts General Hospital Primer Bank (https://pga.mgh.harvard.edu/primerbank/index.html) and checked for specificity using the National Center for Biotechnology Information’s Primer BLAST (http://www.ncbi.nlm.nih.gov/tools/primer-blast/index.cgi?LINK_LOC=BlastHome). The primer sequences used are listed in Table 1. RT-qPCR was performed in triplicate using SYBR® green PCR Master Mix (Thermo Scientific, Waltham, MA) and the Viia™ 7 Real-Time PCR System (Thermo Scientific, Waltham, MA). A template control was run for each primer set and samples were analyzed using the DDCt method (delta-delta cycle threshold). YWHAZ was used as the internal control.

DNA Extraction

Each sample was mechanically homogenized and placed in DNA digest buffer with Proteinase K at 50°C for 3 hours. RNase A was added and a 1:1 phenol/chloroform extraction subsequently performed. The samples were washed with chloroform and DNA was precipitated using 100% ethanol with a 70% ethanol wash. The resulting pellet was suspended in TE buffer. Quantification and purity testing of DNA was performed with spectrophotometric analysis at OD260 and OD280 with the NanoDrop 1000 (Thermo Scientific, Waltham, MA).

DNA Methylation

The region of interest for methylation assessment was identified based on previous studies showing methylation changes in regulatory regions for each gene.14,16–20 The final determined base positions, based on the Genome Reference Consortium build 38, were 92477267–92478260 on chromosome 7 for SYN1, 11111566–11112154 on chromosome 6
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Table 1 RT-qPCR primers used for expression studies

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer pair sequences</th>
<th>Amplicon size (base pairs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SYN1</td>
<td>f GAAGGCCCCCTTACATAACCAATGA</td>
<td>83</td>
</tr>
<tr>
<td></td>
<td>r GATATTTGGTACAGGAGGTGATGTC</td>
<td></td>
</tr>
<tr>
<td>SYN2</td>
<td>f TACACCCCAACCAATTCGCC</td>
<td>93</td>
</tr>
<tr>
<td></td>
<td>r CCGGCTGATATTATCTAGCAAG</td>
<td></td>
</tr>
<tr>
<td>ERV3</td>
<td>f TGTTTCCTGACTCCTCCCTATCC</td>
<td>86</td>
</tr>
<tr>
<td></td>
<td>r GTCGCCGACCAGATG</td>
<td></td>
</tr>
<tr>
<td>PEG10</td>
<td>f AACGCAAGATCAGACGCCTG</td>
<td>75</td>
</tr>
<tr>
<td></td>
<td>r GGGCAATCTCGGAAAGCAT</td>
<td></td>
</tr>
<tr>
<td>RTL1</td>
<td>f GTCTGCAAGGGGTTCACACC</td>
<td>86</td>
</tr>
<tr>
<td></td>
<td>r CCGATGGTGTCTGATG</td>
<td></td>
</tr>
<tr>
<td>LEP</td>
<td>f GACACTGGCAGCTTCAAACAGA</td>
<td>97</td>
</tr>
<tr>
<td></td>
<td>r GTGAGAGAAGATGCCGGAGTT</td>
<td></td>
</tr>
<tr>
<td>CYP19A1</td>
<td>f CCACAGCTGAGAAACTGGAAGA</td>
<td>78</td>
</tr>
<tr>
<td></td>
<td>r TCCCTCGCTTCCACGCTCT</td>
<td></td>
</tr>
<tr>
<td>EDNRB</td>
<td>f GGGAAGGAACCTGTTACTGG</td>
<td>110</td>
</tr>
<tr>
<td></td>
<td>r ACTTGAGGGCGGTGCAAT</td>
<td></td>
</tr>
<tr>
<td>INSL4</td>
<td>f AGCTCTGTCGCTTTATCT</td>
<td>211</td>
</tr>
<tr>
<td></td>
<td>r GTGAGGTTGACCGACACCATTC</td>
<td></td>
</tr>
<tr>
<td>MID1</td>
<td>f CTGCACCTGGCCTTTTAGTCT</td>
<td>107</td>
</tr>
<tr>
<td></td>
<td>r GCACATGTGTCATGACTGGAGATG</td>
<td></td>
</tr>
<tr>
<td>PTN</td>
<td>f GGAGCTGAGTGCGAAGCAAC</td>
<td>157</td>
</tr>
<tr>
<td></td>
<td>r CTGCTTCAGACTCTCCAGTTC</td>
<td></td>
</tr>
</tbody>
</table>

for SYN2, 94656185–94656702 on chromosome 7 for PEG10, and 137268077–137268597 on chromosome 7 for PTN. Genomic DNA methylation patterns were determined by Epityper application (Agena Bioscience, San Diego, CA) as previously described21 (Roswell Park Cancer Institute Genomics Shared Resource with Core grant NCI P30CA16056, Buffalo, NY). Three amplicons were needed for SYN1 and SYN2 and two amplicons for PEG10 and PTN. Samples were run in duplicate. Each amplicon was analyzed separately using mean CpG methylation. Only differentially methylated regions are presented in the results. These regions are amplicon 2 for SYN1, amplicon 1 for SYN2, and amplicon 1 for PEG10. The primer sequences used are listed in Table 2.

Statistical Analysis

Given the distribution of the data, Kruskal–Wallis test was used to compare median expression fold change and methylation levels between all three groups. If significant, two-way comparisons were then performed with the Wilcoxon rank-sum test and reported in the results. The summation of SYN1 and SYN2 methylation percentages in differentially methylated amplicons was then analyzed as a possible predictive test. Receiver operating characteristic curves were generated and sensitivity and specificity were calculated. Categorical baseline data were analyzed by chi-square testing. Statistical significance was defined by p-value <0.05 in all analyses.

Results

Patient Characteristics

A total of 26 specimens were identified from the placenta biobank–10 AGA, 9 SGA, and 7 FGR. As expected, these groups differed significantly by gestational age at birth, placental and birth weights, mode of delivery, and performance of umbilical artery cord gas (Table 3). Specifically, FGR babies weighed 470 g less than SGA and 690 g less than AGA babies at birth (p < 0.001). Only 43% of FGR babies were delivered vaginally, compared with 100% of SGA babies and 90% of AGA babies (p = 0.01).

SYN1 Expression is Significantly Increased in Both FGR and SGA Placentas

Placental SYN1 expression was significantly increased in both FGR and SGA samples compared to AGA samples (p = 0.027 and p = 0.005, respectively). There was, however, no significant difference between SYN1 expression in SGA and FGR placentas. There was no difference in SYN2 expression between the three groups (Fig. 1A and B).

SYN1 and SYN2 Methylation is Decreased Uniquely in FGR Placentas

Methylation of SYN1 was decreased in FGR samples [23.5% CpG methylation (IQR 21.5, 26.5)] compared with SGA [29.6% CpG methylation (IQR 24.0, 32.1); p = 0.044] and AGA [28.9% CpG methylation (IQR 26.6, 33.5); p = 0.006]. Interestingly, despite the lack of change in expression patterns, SYN2 methylation was also decreased in FGR samples [16.5% CpG methylation (IQR 14.8, 19.8)] compared with SGA [21.9% CpG methylation (IQR 19.8, 22.3); p = 0.008] and AGA [22.9% CpG methylation (IQR 21.7, 24.0); p = 0.011] (Fig. 1C and D).

SYN1 and SYN2 Methylation Accurately Identifies FGR Placentas

Methylation of SYN1 was decreased in FGR samples [23.5% CpG methylation (IQR 21.5, 26.5)] compared with SGA [29.6% CpG methylation (IQR 24.0, 32.1); p = 0.044] and AGA [28.9% CpG methylation (IQR 26.6, 33.5); p = 0.006]. Interestingly, despite the lack of change in expression patterns, SYN2 methylation was also decreased in FGR samples [16.5% CpG methylation (IQR 14.8, 19.8)] compared with SGA [21.9% CpG methylation (IQR 19.8, 22.3); p = 0.008] and AGA [22.9% CpG methylation (IQR 21.7, 24.0); p = 0.011] (Fig. 1C and D).

The Methylation Pattern in SYN1 and SYN2 is Unique to These Retroelements

There was no statistical difference between the three groups in expression in other REs that are highly expressed in the placenta including ERV3, PEG10, RTL1, LEP, EDNRB, CYP19A1, INSL4, MID1 or PTN. There was a trend toward increased expression in FGR samples in PEG10 [1.15 fold change for
AGA (IQR 0.95, 1.36) vs. 1.59 fold change for SGA (IQR 1.26, 2.85) vs. 2.31 fold change for FGR (IQR 1.14, 3.33) and PTN [0.66 fold change for AGA (IQR 0.38, 0.83) vs. 1.20 fold change for SGA (IQR 0.81, 1.76) vs. 1.02 fold change for FGR (IQR 0.58, 1.38)], and thus methylation analyses for these genes were performed. In contrast to SYN1 and SYN2, the differentially methylated region of PEG10 showed higher methylation in FGR [50.2% CpG methylation (IQR 43.1, 60.2)] compared with AGA [21.0% CpG methylation (IQR 11.1, 23.8); \( p = 0.005 \)], but not to SGA samples [41.8% CpG methylation (IQR 29.0, 42.4)]. Including this with SYN1 and SYN2 methylation did not improve the area under the curve of the generated ROC curves, so was not considered in the final predictive model above.

**Discussion**

The current study demonstrates significantly lower placental methylation of the regulatory regions of SYN1 and SYN2 in
FGR compared with SGA pregnancies. These differences could be used to distinguish pathologic FGR from constitutional SGA with reasonable predictive accuracy in this cohort. The methylation differences corresponded with a biologically consistent increase in expression of SYN1, though not SYN2. While DNA methylation regulates expression of both SYN1 and SYN2, expression patterns do not always follow that predicted by changes in DNA methylation. This suggests that other mechanisms regulate the expression of these critical gene products.

The methylation differences described in this study are consistent with previously published data showing differences in expression and methylation of these genes in growth discordant twins as well as pregnancies complicated by SGA and other placental syndromes. To our knowledge, however, this is the first study to attempt to assess differences in placental expression and methylation of SYN1 and SYN2 in pathologic FGR compared with physiologic SGA. This is of particular interest because it is biologically plausible that epigenetic marks such as DNA methylation are modifiable by environmental differences, such as hypoxia, that would lead to FGR versus SGA. An additional strength of our study is that all samples were obtained from deliveries done at 36 weeks or greater. This minimized the impact of gestational age on our results.

There are also limitations with the current study that deserve comment. First, the FGR and SGA groups were not identified prospectively and thus these groups may have
some overlap. Nevertheless, the FGR group clearly represents a sicker population. Birth weights are smaller, there is a trend toward lower cord pH and a markedly increased rate of cesarean delivery compared with the SGA group. An additional concern is that our sample size in each group was small. To be more confident about the significance and magnitude of differences, larger studies replicating these results would be important. Additionally, by limiting our samples to term or near term, we may have missed cases with more severe FGR, as these would have been more likely to be delivered significantly preterm. As true differences would likely be exaggerated in more severely affected pregnancies, this decision should support the null hypothesis in the current study, but we cannot exclude the possibility of a more complicated relationship between SYN1 and SYN2 methylation and severity of placental dysfunction.

Despite these limitations, our data suggest that SYN1 and SYN2 may be useful biomarkers for distinguishing FGR from SGA. An important next step is a prospective study. To make these data clinically useful a source of placental DNA must be available prior to delivery. Thus, we propose to replicate our results in prospectively obtained cell-free fetal DNA as the majority of this is suspected to be placental in origin.23 Studies have already been performed that illustrate the feasibility of assessing differential CpG methylation in cell-free fetal DNA and maternal DNA,24 supporting the potential of this approach.

In conclusion, we identified significant differences in methylation patterns of SYN1 and SYN2 that distinguished FGR from SGA. This work adds to a growing effort to define FGR biologically, rather than by a threshold centile on a growth chart.25 If these methylation differences are replicated in cell-free DNA, this approach has the potential to provide noninvasive information about placental function that could be used clinically.

Presentation Information
Findings presented at the 36th Annual Pregnancy Meeting, Society for Maternal Fetal Medicine, Atlanta, Georgia, February 1–6, 2016

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Conflict of Interest and Financial Disclosure
The authors have no financial disclosures or potential conflict of interest.

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