Identification of placental nutrient transporters associated with intrauterine growth restriction and pre-eclampsia

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Abstract

Background: Gestational disorders such as intrauterine growth restriction (IUGR) and pre-eclampsia (PE) are main causes of poor perinatal outcomes worldwide. Both diseases are related with impaired materno-fetal nutrient transfer, but the crucial transport mechanisms underlying IUGR and PE are not fully elucidated. In this study, we aimed to identify membrane transporters highly associated with transplacental nutrient deficiencies in IUGR/PE.

Results: In silico analyses on the identification of differentially expressed nutrient transporters were conducted using seven eligible microarray datasets (from Gene Expression Omnibus), encompassing control and IUGR/PE placental samples. Thereby 46 out of 434 genes were identified as potentially interesting targets. They are involved in the fetal provision with amino acids, carbohydrates, lipids, vitamins and microelements. Targets of interest were clustered into a substrate-specific interaction network by using Search Tool for the Retrieval of Interacting Genes. The subsequent wet-lab validation was performed using quantitative RT-PCR on placentas from clinically well-characterized IUGR/PE patients (IUGR, n = 8; PE, n = 5; PE+IUGR, n = 10) and controls (term, n = 13; preterm, n = 7), followed by 2D-hierarchical heatmap generation. Statistical evaluation using Kruskal-Wallis tests was then applied to detect significantly different expression patterns, while scatter plot analysis indicated which transporters were predominantly influenced by IUGR or PE, or equally affected by both diseases. Identified by both methods, three overlapping targets, SLC7A7, SLC38A5 (amino acid transporters), and ABCA1 (cholesterol transporter), were further investigated at the protein level by western blotting. Protein analyses in total placental tissue lysates and membrane fractions isolated from disease and control placentas indicated an altered functional activity of those three nutrient transporters in IUGR/PE.

Conclusions: Combining bioinformatic analysis, molecular biological experiments and mathematical diagramming, this study has demonstrated systematic alterations of nutrient transporter expressions in IUGR/PE. Among 46 initially targeted transporters, three significantly regulated genes were further investigated based on the severity and the disease specificity for IUGR and PE. Confirmed by mRNA and protein expression, the amino acid transporters SLC7A7 and SLC38A5 showed marked differences between controls and IUGR/PE and were regulated by both diseases. In contrast, ABCA1 may play an exclusive role in the development of PE.

Keywords: Bioinformatics, Membrane transporters, Placenta, Pre-eclampsia, Intrauterine growth restriction
Background

Worldwide, more than one third of child deaths can be attributed to maternal and fetal undernutrition [1]. Along the course of gestation, the fetus receives organic and inorganic nutrients from the mother, such as amino acids, carbohydrates, lipids, vitamins and minerals, for adequate development and growth. To avoid direct materno-fetal blood contact, the placenta has distinctive anatomical barriers for regulating the substrate exchange [2, 3]. Various molecules originating from maternal blood enter the fetal circulation after crossing successively the apical microvillus plasma membrane (MVM), the cytoplasm, and the basal plasma membrane (BM) of the trophoblast layer [3, 4]. On the other side, the fetus eliminates waste products via efflux processes at the BM into the maternal veins. This highlights the crucial role of substrate specific membrane transporter proteins at these different locations. Indeed, for several membrane transporter proteins (e.g. Ca$^{2+}$-ATPase, ferroportin, system A amino acid transporters) an asymmetric distribution between MVM and BM has been reported. Although this asymmetry is a prerequisite for net transport across the placenta via transporter proteins, the mechanisms of protein trafficking that result in distribution to either MVM or BM in the polarized syncytiotrophoblast are largely unknown. It is evident, however, that any reduction of the quantity, density, and activity of membrane transporters at the placental microvilli could impact the rates of the nutrient supply, which may subsequently compromise the fetal development. It is well recognized that gestational diseases such as intrauterine growth restriction (IUGR) and pre-eclampsia (PE), which complicate a significant proportion of pregnancies leading to poor perinatal outcome, are closely related to impaired placental nutrient transfer [5, 6].

Transfer of amino acids across the placenta is mediated by substrate selective amino acid transport systems. So far, more than 15 different systems have been identified in the human placenta, which include Na$^+$-dependent systems A, N, ASC, β, $X_{AG}^-$, $X_C^-$, GLY and Na$^+$-independent systems L, γ, γL, T, b$_{0,+}^+$ [5, 7, 8]. Reduced activity of amino acid transporters e.g. system A in IUGR and PE has been reported [9–11], which might be due to reduced sodium-potassium ATPase (Na$^+$/$K^+$ATPase) activity [12] and a competitive inhibition by physiologically elevated maternal plasma homocysteine [13, 14]. The transplacental transport of glucose mainly occurs along a concentration gradient facilitated by the glucose transporter (GLUT) family, while lipids are mostly packed into circulating protein particles by virtue of active membrane transporters [15]. Current knowledge in this field revealed an increased concentration gradient from maternal to fetal sides in IUGR and PE to achieve higher glucose and lipid transfer [16, 17]. These significant elevations have also been considered as a risk factor for those gestational disorders [18, 19], which may be explained by an altered expression of membrane transporter [20, 21].

The fetal provision with vitamins and dietary elements is required, since they are essential to regulate the fetal metabolism as well as extra- and intra-cellular osmolality. Global studies under the auspices of the World Health Organization in 2008 proposed that deficiencies in micronutrients such as thiamine (Vitamin B1), riboflavin (Vitamin B2), folate (Vitamin B9), cobalamin (Vitamin B12), and ascorbic acid (Vitamin C), are highly prevalent and occur concurrently among pregnant women [1]. It has been reported that the elevated serum homocysteine levels in IUGR and PE were accompanied by decreasing serum levels of the aforementioned class B vitamins. The dietary treatment of pregnant women with diverse vitamin B supplements improved fetal weight [22–26]. However, data regarding the transplacental transport of vitamin B are scarce. In this context SLC19 family members were found to transport folate (SLC19A1) and thiamine (SLC19A2 and SLC19A3) [27]. Interestingly, in IUGR placentas, an increase in mRNA level of SLC19A1 and SLC19A2 has been reported [28], which was interpreted as a compensatory effect. Besides, deficiencies in serum calcium, magnesium and zinc might greatly contribute to the pathogenesis of PE, since it was suggested that the dietary supplementation of these elements can reduce the severity of PE [29–31]. Although calcium is probably the most studied element among all ions, publications investigating the role of transporters or exchangers in placental calcium homeostasis are negligible (<20). It was found that the expression of Ca$^{2+}$-selective receptor channels and Na$^+$/Ca$^{2+}$ exchangers were significantly increased in hypertensive placentae compared to normotensive tissues [32].

The studies mentioned above and numerous others have suggested that an altered transplacental nutrient transport might elicit IUGR/PE [5, 6, 33]. However, the detailed underlying mechanisms involving membrane transporters are still not fully elucidated. Therefore, we conducted the current study to gain a comprehensive knowledge on the importance of placental transporter proteins majorly involved in the fetal provision with amino acids, carbohydrates and derivatives, lipids, vitamins, and microelements. We hypothesized that the expression of placental membrane transporters was modified by IUGR/PE. To identify the key nutrient transporters in placenta, we have used a multifaceted approach including bioinformatic analysis, molecular biology, and mathematical diagramming (Fig. 1).
by the Markov cluster algorithm (MCL).

with a confidence score larger than 0.4), and clustered with medium protein-protein interactions (associated tool, we restricted the prediction to human placenta indirect (functional) targets. STRING-db.org with a focus on the direct (physical) and retrieval of Interacting Genes (STRING) 9.1 at http://

identified targets of interest to Search Tool for the Retention transporters from microarray data, we applied the biochemical approach using the limma package (R 3.1, Bioconductor 2.7). We analysed gene expression data of solute carriers (SLCs), ATP-binding cassette transporters (ABCs) and transient receptor potential channels (TRPs). Any SLC, ABC and TRP family member, which i) was represented in more than 3 studies, ii) was listed in top 12 up- or down-regulated genes, and iii) has an annotated a biological significance in placenta, was added to the candidate list.

Visualization of biomolecular interaction network
To better understand the network of the identified nutrient transporters from microarray data, we applied the identified targets of interest to Search Tool for the Retrieval of Interacting Genes (STRING) 9.1 at http://string-db.org with a focus on the direct (physical) and indirect (functional) targets’ connections. In the STRING tool, we restricted the prediction to human placenta with medium protein-protein interactions (associated with a confidence score larger than 0.4), and clustered by the Markov cluster algorithm (MCL).

Placental sample collection
The study was approved by the local ethic institutional review board of the Canton of Bern with an informed consent obtained from each participant prior to the caesarean section. Pregnant women were under obstetrical care at the Department of Obstetrics and Gynecology, University Hospital (Bern, Switzerland). They were assigned to five groups: term controls, preterm, PE, and PE combined with IUGR (PE + IUGR). Term controls consisted of the pregnancies without pathologies and terminated at term (≥ 37 gestational weeks) by elective primary caesarean section upon patients’ request or due to breech presentation. The preterm group was defined as the babies born prior to 37 weeks of pregnancy, but without further abnormal maternal or fetal symptoms. According to the definition from the International Society for the Study of Hypertension in Pregnancy, patients with the dramatic increase of systolic/diastolic blood pressure (> 140/90 mmHg) and urinary protein content in 24 h (> 0.30 g) were classified in PE related groups. When the fetal birth weight was below the 10th percentile for babies of the same gestational age, the baby was also referred to as small for gestational age due to the IUGR effects. As it is a severe complication when PE occurs concomitantly with IUGR, we also included and characterized patients suffering from PE combined with IUGR (PE + IUGR).

Placental tissue collection was performed and fully validated as previously described [21, 34]. Immediately after caesarean section, chorionic tissue was dissected from a standardized central location, and snap-frozen in liquid nitrogen after a thorough wash with Dulbecco’s phosphate buffered saline (Gibco). The collected tissue specimens were stored at -80 °C until further analysis.

Quantitative RT-PCR
Total RNA was isolated from placental tissue with a modified protocol, using a Trizol reagent kit (Invitrogen) [34]. Approx. 50 mg frozen placental tissues were homogenized in 1 ml cold Trizol on ice with a Polytron homogenizer (Kinematica AG), followed by phenol/chloroform extraction and precipitation by isopropanol, and final washing with 75% ethanol. After re-suspension in RNase free water, the extracted RNA was evaluated qualitatively (assessment of RNA integrity) by using Agilent 2100 Bioanalyzer (Agilent Technologies) and quantitatively (determination of RNA concentration) with NanoDrop 1000 (Thermo Scientific). All samples used in this study fulfilled the fixed selection criteria which were: RNA integration number (RIN) > 6, spectrophotometric measurements at 260/280 > 2.0 and 260/230 > 1.8, and the yield of RNA > 5 ng/μl.

Two microgram RNA was reverse transcribed into cDNA using GoScript™ Reverse Transcriptase System
GoTaq \textsuperscript{™} qPCR Master Mix (Promega) on the Viia\textsuperscript{TM} 7 Real-Time PCR System (Applied Biosystems). Based on the results of the transporter mRNA expression, the heatmap was generated using centered correlation (Cluster 3.0) and average linkage clustering analysis for similarity measurement and clustering (TreeView at http://diyhpl.us/~bryan/irc/protocol-online/protocolcache/EisenSoftware.htm). The consecutive statistical analysis and mathematical diagramming served for the identification of significant and disease-specific targets, respectively.

**Total protein and membrane fraction extraction**

To extract total protein from placenta, tissue samples were homogenized in ice-cold hypotonic buffer (10 mM Tris-HCl, 10 mM NaCl, 1.5 mM MgCl\textsubscript{2} and 0.1% Triton X-100, pH 7.4) supplemented with protease inhibitor cocktail (Sigma), and centrifuged at 1000 \texttimes g for 10 min at 4 °C. The supernatants were collected in fresh tubes and stored at -80 °C.

Membrane fractions from placenta tissues were extracted using membrane, nuclear and cytoplasmic protein extraction kit according to the manufacturer’s protocol (Bio Basic Inc.). The whole extraction procedure was carried out on ice in the presence of protease inhibitor, phosphatase inhibitor, dithiothreitol and phenylmethylsulfonyl fluoride (PMSF). The “identity” of the extracted cellular membrane fractions was validated by detecting the corresponding expression of the plasma membrane marker Novus Biologicals, the nuclear marker histone H3 (Novus Biologicals) and the cytoskeleton marker β-actin (Sigma).

The protein content of total lysates and membrane fractions was quantified by bicinchoninic acid assay (Pierce), using bovine serum albumin as standard.

**Western blotting**

90 μg of the total tissue lysate or 30 μg of membrane fraction was mixed with Laemmli sample buffer and separated by sodium dodecyl sulfate polyacrylamide gel electrophoresis (Bio-Rad) using 10% acrylamide gels. The immobilized bands were then semi-dry transferred to nitrocellulose membranes (GE Healthcare). Blots were blocked with 5% non-fat milk in Tris Buffered Saline with 0.1% Tween-20. The following antibodies were used: rabbit anti-ABCA1 (Novus Biologicals), rabbit anti-SLC7A7 (Novus Biologicals), and rabbit anti-SLC38A5 (Aviva Systems Biology), as well as mouse anti-β-actin or mouse anti-Na\textsuperscript{+}/K\textsuperscript{+} ATPase as internal controls. Primary antibodies were first incubated overnight at 4 °C followed by 4 times washing with Tris-buffered saline supplemented with 0.1% Tween 20 (TBST), and then incubation with DyLight 680 or 800 fluorescence conjugated secondary antibodies (Thermo Scientific). The immunoreactive bands were visualized with the Odyssey\textsuperscript{™} Sa Infrared Imaging System (LI-COR) to obtain relative densitometry values.

**Statistical analysis and mathematical diagramming**

The comparative threshold cycle (Ct) method was applied to analyze relative gene expression by fold change $= 2^{-\Delta(\Delta Ct)}$, where $\Delta Ct = Ct_{target \ gene} - Ct_{reference \ gene}$ and $\Delta(\Delta Ct) = \Delta(Ct_{test} - Ct_{control})$ [35].

Data were statistically evaluated using GraphPad Prism 6 Software. Kruskal-Wallis analysis followed by a Dunn’s multiple comparison test, were used to analyse statistical differences of the clinical records, the mRNA and protein expression data between the term controls and preterm, IUGR, PE and PE + IUGR groups. A p value of less than 0.05 was considered statistically significant.

In addition, scatter plotting was applied to estimate the influence of gestational diseases on transporter expression. Each point depicts a nutrient transporter, whose x-axis and y-axis represents the average log 2-fold changes (FC) in the IUGR group and PE group, respectively. To avoid possible bias (individual variation, experimental error, etc.), we selected only those targets, whose expression has been modulated with an absolute FC > 1.14 (i.e. >114% or <86%) over the expression in the term controls [36]. Subsequently, the ratio of y to x was considered as a rank variable. When the ratio is close to 0, it means that the predominant effect on transporter mRNA expression is caused by PE. Conversely, when it approaches infinity, the altered transporter is primarily influenced by IUGR. An intermediate state (i.e. a ratio equal to 1 or -1), indicates an equal (1) or an equal but reversed (-1) impact on the transporter expression level by IUGR and PE.

**Results**

**Microarray processing**

Six placental gene expression studies focusing on IUGR/PE plus one project investigating umbilical cords associated with low birth weight, were included in this meta-analysis (Table 1). These studies represented in total 258 samples, including 129 controls (term and preterm samples), 67 IUGR-related specimens and 62 PE placentas.

Based on the selection criteria for potential transporter candidates described in the section Methods, 46 of the 434 genes were chosen for consecutive analysis. These genes encode amino acids, carbohydrates and derivatives, lipids, vitamins, microelements, and other transport channels (Table 2). Among the selected targets, 32 genes exhibited increased expression levels in...
pathological pregnancies compared to controls, while 14 genes showed the opposite trend.

Integration of interaction between selected candidates
As one of the largest database for protein interactions, STRING aggregates the most available information (e.g. neighbourhood, gene fusion, co-occurrence, co-expression, experiments, databases, text mining and homology) derived from genomic context, high throughput experiments, co-expression and previous knowledge. Using the medium confidence score and clustered by MCL, 46 signature nutrient transporters were mapped into an interaction network (Fig. 2). The nodes visualize the proteins with known structure, and the solid lines represent the predicted functional associations with dashed inter-cluster edges. On grouping the network, it summarized the predicted associations for a particular group of proteins, for instance, amino acids associated transporters (blue cluster), vitamins associated transporters (red cluster), microelements and ions associated transporters (yellow cluster). The STRING view depicted that the selected targets were clearly clustered and strongly interconnected based on their substrate specificity, which reflected a high degree of functional association and a potential interplay among the nutrient transporters.

Identification of altered transporter gene expression in IUGR/PE
Characterization of patients
To validate the gene expression on a well characterized, clinically annotated tissue set, we have performed quantitative RT-PCR measurements on a number of accredited representative samples. Table 3 shows the demographic and clinical characteristics of the studied population. There were no major differences in the age, BMI, gravidity and parity numbers between the patients. The gestational age clearly isolated the term pregnancies from the rest, which was directly related with the placental weight and the birth weight.

In addition, as a PE related biomarker, placental leptin mRNA levels have been measured. A significantly increased expression of leptin has been shown in PE and PE + IUGR compared to both term and preterm control placentae, but not in isolated IUGR (data not shown).

Selection of reference genes
Prior to evaluating the nutrient transporter regulation in IUGR/PE, we first screened the mRNA abundance of commonly used reference genes: β2-microglobulin, mitochondrial ribosomal protein L19, ubiquitin (UBQ), and tyrosine 3-monooxygenase/tryptophan 5-monooxygenase activation protein, zeta polypeptide (YWHAZ). We found no differences between controls and the herein tested gestational diseases (data not shown). In accordance with our previous work [34], the mean value of UBQ and YWHAZ was used for quantitative RT-PCR normalization of target nutrient transporters.

Gene expression analysis
We evaluated the expression of the selected 46 targets in 43 well-characterized placenta samples using unsupervised hierarchical clustering. The heatmap shown in Fig. 3 illustrates an overview of the expression profiles of these transcripts based on the quantitative RT-PCR results. The generated expression profile data confirmed that the 43 well-characterized placenta samples used in this study highly reproduced the results obtained from microarray data of 258 samples all over the world.

Based on Kruskal-Wallis test, eight candidate transporters were identified showing significant changes on mRNA expression among gestational disorders: the amino acid transporters SLC7A7 (Fig. 4a), SLC38A2 (Fig. 4b), SLC38A5 (Fig. 4c), the glucose transporter SLC2A1 (Fig. 4d), the cholesterol transporter ABCA1 (Fig. 4e), the vitamin B transporters SLC19A3 (Fig. 4f)
| Substrate class                        | Predominant (predicted) substrate | Gene name | Accession number | Protein name | Log (FC) | Representative references |
|---------------------------------------|-----------------------------------|-----------|-----------------|--------------|----------|----------------------------|
| **Amino acids**                       | System L, y⁺L, xc-and asc [58, 59] | SLC3A2    | NM002394        | 4F2hc        | −0.42~0.28 | [60–63]                    |
|                                       | Cationic L- amino acids           | SLC7A1    | NM003045        | CAT1         | −0.52~0.89 | [64–67]                    |
|                                       | Large neutral L-amino acids       | SLC7A5    | NM003486        | LAT1         | −0.38~0.46 | [62, 63, 66, 68–71]        |
|                                       | Cationic amino acids, large       | SLC7A6    | NM003983        | y⁺LAT2       | −0.23~0.39 | [69, 72]                    |
|                                       | neutral L-amino acids             | SLC7A7    | NM001126105     | y⁻LAT1       | −0.44~0.89 | [72]                       |
|                                       | Neutral L – amino acids           | SLC7A8    | NM012244        | LAT2         | −0.12~0.25 | [73]                       |
|                                       | Cystine (anionic form), L-        | SLC7A1    | NM014331        | xCT          | −0.10~0.53 | –                          |
|                                       | glutamate                         |           |                 |              |          |                             |
|                                       | Gln, Ala, Asn, Cys, His, Ser     | SLC38A1   | NM030674        | SNAT1        | −0.28~1.24 | [68, 74, 75]                |
|                                       |                                  | SLC38A2   | NM018976        | SNAT2        | −0.11~0.11 | [50, 68, 75–78]             |
|                                       |                                  | SLC38A5   | NM033518        | SNAT5        | 0.01~0.21  | [49]                       |
|                                       | Polyamines, amino acids [79]     | SLC12A8   | NM024628        | CCC9         | −0.45~1.09 | –                          |
|                                       |                                  |           |                 |              |          |                             |
| **Vitamins**                          | Thiamine [27]                     | SLC19A2   | NM006996        | THTR1        | 0.28~0.66  | [80]                       |
|                                       |                                  | SLC19A3   | NM025243        | THTR2        | −1.17~−0.12 | [80, 81]                   |
|                                       | Folate                            | SLC19A1   | NM194255        | RFC          | −0.10~0.84 | [82–85]                    |
|                                       |                                  | SLC46A1   | NM080669        | PCFT         | −0.15~0.05 | [82, 86]                    |
|                                       | Biotin, pantothenic acid [87]     | SLC5A6    | NM021095        | SMVT         | 0.01~0.04  | [88, 89]                    |
|                                       | Cobalamin [90] (Pyridoxine) [91]  | LMBRD1    | NM018368        | LMBRD1       | −0.26~0.22 | –                          |
|                                       | L-ascorbid acid [92, 93]          | SLC23A1   | NM005847        | SVCT1        | 0.01~0.10  | [94, 95]                    |
|                                       |                                  | SLC23A2   | NM005116        | SVCT2        | −0.38~0.43 | [95–97]                    |
|                                       |                                  | SLC23A3   | NM144712        | SVCT3        | −0.21~0.33 | –                          |
|                                       |                                  |           |                 |              |          |                             |
| **Microelements and ions**            | Zn²⁺ [98, 99]                     | SLC30A1   | NM021194        | ZNT1         | −0.02~0.34 | [100–102]                  |
|                                       |                                  | SLC30A2   | NM032513        | ZNT2         | −0.06~0.17 | [100, 101, 103]            |
|                                       |                                  | SLC30A4   | NM013309        | ZNT4         | −0.21~0.05 | [100, 101]                 |
|                                       |                                  | SLC39A1   | NM014437        | ZIP1         | −0.13~0.04 | [100]                      |
|                                       |                                  | SLC39A8   | NM022154        | ZIP8         | 0.00~0.95  | [104]                      |
|                                       | Ca²⁺ [108] e.g. Cl⁻,[COO]⁻ [108]  | TRPV6     | NM018646        | TRPV6        | 0.02~0.43  | [32, 105–107]              |
|                                       |                                  | SLC2A15   | NM018420        | FLIPT1       | −0.14~0.10 | –                          |
|                                       |                                  | SLC23A1   | NM005847        | SVCT1        | 0.01~0.10  | [94, 95]                    |
|                                       |                                  | SLC23A2   | NM005116        | SVCT2        | −0.38~0.43 | [95–97]                    |
|                                       |                                  | SLC23A3   | NM144712        | SVCT3        | −0.21~0.33 | –                          |
|                                       |                                   |           |                 |              |          |                             |
| **Lipids**                            | Cholesterol, phospholipids [129]  | ABCA1     | NM005502        | ABCA1        | −0.19~0.71 | [21, 130, 132]             |
|                                       | Drugs, toxins [133]               | ABCG1     | NM207629        | ABCG1        | −0.20~0.33 | [130, 131]                 |
|                                       |                                  | ABCG2     | NM004827        | ABCG2        | −0.70~0.48 | [134–136]                  |
|                                       |                                  | SLC47A1   | NM018242        | MATE1        | −0.03~0.33 | [138, 139]                 |
|                                       |                                  | SLC27A2   | NM003645        | FATP2        | −0.58~1.37 | [141, 142]                 |
|                                       |                                  | SLC27A4   | NM005094        | FATP4        | −0.04~0.05 | [142–144]                  |
Table 2

Selected nutrient transporters according to their regulational signature and physiological significance (Continued)

| Substrate class | Predominant (predicted) substrate | Gene name | Accession number | Protein name | Log (FC) | Representative references |
|-----------------|----------------------------------|-----------|-----------------|-------------|---------|--------------------------|
| Carbohydrates (+ metabolic derivatives) | Glucose [145] | SLC2A1 | NM006156 | GLUT1 | 0.12~0.66 | [20, 40, 146] |
| | Lactate [147] | SLC16A1 | NM003051 | MCT1 | −0.36~1.35 | [148–150] |
| | | SLC16A3 | NM004207 | MCT4 | −0.12~0.48 | [148–151] |
| | | SLC16A4 | NM004696 | MCT5 | −0.50~0.47 | [149] |
| Neurotransmitter | Norepinephrine [152] | SLC6A2 | NM001043 | NET | −0.31~0.54 | [153–155] |

Abbreviations: SLC solute carrier, LCFA long chain fatty acid, VLCFA very long chain fatty acid

and SLC22A15 (Fig. 4g), as well as the sulphate transporter SLC26A2 (Fig. 4h). In order to directly compare the trend of gene regulation in IUGR or PE with the microarray analysis, we have presented the gene expression as the log (FC) to term controls. Almost all trends of alteration of transporters by IUGR/PE were comparable to those derived from microarray data, except the effects on ABCA1 and SLC38A2 by PE. The latter might be due to different criteria on PE sample selection, which result in greater variations between microarrays.

In addition, mathematical diagramming was applied to assess whether IUGR or PE is the major modulator of the determined alteration in gene expression. In the scatter plot shown in Fig. 5, each point represents the average expression level of one nutrient transporter. The log (FC)IUGR and log (FC)PE of independent transporters obtained from quantitative RT-PCR analysis, were defined as paired X-Y values. The absolute value on the X-axis or Y-axis reflects the severity of the gene expression changes by IUGR or PE, respectively. By ranking the absolute X or Y value, the top 3 genes dominantly influenced by IUGR were SLC23A1, SLC26A2, and SLC7A11 (yellow squares), while SLC16A4, ABCA1, and ABCG2 (blue squares) were regulated mainly by PE. On the
contrary, transporters such as SLC7A7, SLC38A5 and SLC7A8 (red squares), were equally affected by IUGR and PE since they locate on the y = x or y = -x line.

Taking into account the statistical analysis and the mathematical diagramming, the overlapping targets, namely ABCA1, SLC7A7, SLC38A5, and SLC26A2, were further subjected to protein quantification. However, due to the lack of a specific, validated antibody, the evaluation of SLC26A2 protein could not be further pursued.

Protein levels of the key membrane transporters

Using immunoblotting technique, we complemented the quantitative RT-PCR data with corresponding protein expression levels. As shown in Fig. 6, the total protein content of ABCA1 was significantly decreased by PE. In contrast, SLC7A7 transporter showed significant up-regulation in placenta associated with PE, while down-regulation in IUGR placenta was found. SLC38A5 expression in placenta tissue was only slightly elevated in IUGR and PE. The loss of significance may due to the limited sample size. Generally, the changes found on protein level for the three transporters ABCA1, SLC7A7 and SLC38A5, corresponded well with the transcriptional results.

As the target transporters modulate the fluxes of substrates through the plasma membrane to meet fetal requirements, we also isolated plasma membrane protein fractions from the placentae, and confirmed their high purity by enrichment of the marker protein Na+K+ATPase (data not shown). The latter was further used to normalize the transporter protein expression detected by Western blotting.

We have previously reported that the protein expression of the lipid transporter ABCA1 in extracted plasma membranes was significantly reduced only in PE placentae [21]. These findings were confirmed in the present study (data not shown). Interestingly, the opposite regulations for SLC7A7 protein between IUGR and PE-associated disease types that we detected in tissue lysates (Fig.6) were generally also reflected in extracted plasma membrane preparations (Fig. 7). Normalization to β-actin and Na+K+ATPase revealed a trend to decreased expression of SLC7A7 in IUGR tissues, and confirmed a significantly increased expression in PE tissues (Fig. 7a). SLC38A5 was not significantly altered in membrane proteins obtained from diseased and healthy (control) placentas (Fig. 7b). However, marked increases in IUGR and PE compared with the preterm group were found.
Discussion

Fetal growth is largely dependent on the availability of nutrients transported into the fetal circulation via the placenta. The transport of macro- and micronutrients across the placental barrier relies on the presence and activity of substrate specific membrane transporters. The altered expression of these transporters is thought to be implicated in diverse mechanisms of the fetal development. The current study focused on studying to what extent and which specific transporters are impaired by IUGR or PE. To this end, we screened the expression profiles of nutrient transporters whose substrates are amino acids, carbohydrates, lipids, vitamins and microelements in diseased and control placentas by analysing 7 publicly available GEO microarray datasets. The selected targets have been further validated on in-house well-characterized placenta tissues. A systematic alteration in mRNA expression of diverse transporters were identified in IUGR/PE. Among the 46 nutrient transporters, 8 were significantly regulated by those gestational disorders. In particular, the amino acid transporters SLC7A7 and SLC38A5 were strongly influenced by both IUGR and PE. The mRNA and protein expression of the cholesterol transporter ABCA1, by contrast, was only regulated under PE condition.

Microarray gene expression data provided a global overview on the nutrient transporter expression. However, the weakness of these data was that they were derived from studies using not exactly the same criteria for the clinical recruitment of patients. In this study, we have systematically reviewed our on-site collected samples in the following ways. First, the applied clinical and diagnostic data for each individual did meet the strictly defined clinical criteria for classification as IUGR or PE (Table 3). Second, the higher gene expression of leptin occurring only in the PE and PE + IUGR groups served as a placenta-derived marker of PE [37, 38]. Third, we have compared the regulated mRNA expressions that were generated on our 43 in-house samples with results from the cohort of character-matched 258 patients in seven microarray projects. It showed that only 26% of the targets exhibited inconsistent pathological effects
between our samples and the global tissue sets. Based on aforementioned evidences, we considered the in-house tissue samples applied in this study as adequate to reflect patterns occurring in a wider target population.

We further analysed the impact of IUGR/PE on the mRNA expression levels of the selected 46 transporters. Overall, our quantitative RT-PCR measurements revealed a differential expression pattern between control and diseased placentae, including 8 significantly regulated candidates, which might be critical for the fetal malnutrition associated with IUGR/PE. Within this list, SLC2A1 (GLUT1) has been repeatedly reported as significantly reduced at the mRNA level by PE [39, 40]. The studies in animals, such as rat and sheep, have shown that placental GLUT1 expression is significantly decreased in IUGR [41, 42], which is consistent with our findings. We also detected a mRNA down-regulation of the thiamine transporter SLC19A3 and the putative pyridoxine transporter SLC22A15 in IUGR and PE. To our knowledge, this is the first report demonstrating a reduction of placental SLC19A3 and SLC22A15 in IUGR and PE. Currently, there is a lack of specific clinical data on vitamin B1 or B6 deficiency during human pregnancy, except for B vitamin complex supplements. The
detected reduction of SLC19A3 and SLC22A15 mRNA in this study may raise our attention to serum thiamine and pyridoxine alterations during pregnancy, which could interrupt the structural and functional development of cerebella and hippocampus, respectively. Information on the sulfate transporter SLC26A2 in placenta is very scarce. Only Simmons et al. recently reported an abundant mRNA and CTB-specific expression of SLC26A2 in human placenta [43]. In the present study, we identified an up-regulation of SLC26A2 mRNA in IUGR and a down-regulation in PE, based on our in silico screening and the in house gene quantitation. Those opposite but significant changes between IUGR and PE may indicate a different role of SLC26A2 in these two diseases which should be further investigated.

The cholesterol transporter ABCA1, as well as the amino acid transporters SLC7A7 and SLC38A5 were consecutively chosen as the main targets for further investigation, as they popped up both in the statistical analysis and the mathematical diagramming procedure. For ABCA1, we have confirmed the previous finding of an exclusively decreased ABCA1 expression in PE [21]. The impaired ABCA1 function resulted in compromised cholesterol and phospholipid efflux, hence a concomitant accumulation of lipids was observed in PE placentas [44]. The amino acid transporter SLC7A7 encodes the y+LAT-1 protein, which is known to be located at the MVM and BM [7]. It forms a heterodimer complex with hF2hc (encoded by SLC3A2), mediating the Na+-independent cationic amino acids or large neutral L-amino acids transport [45]. Our finding regarding the decrease of SLC7A7 mRNA and protein in IUGR is in agreement with a previous study using Slc7a7-/- mice [46], which caused fetal growth retardation by downregulating insulin-like growth factor 1. In addition, we found an increased SLC7A7 expression in diseases associated with PE compared with gestational age-matched controls. As lysine, arginine and ornithine are substrates for SLC7A7, high SLC7A7 expression could cause reduced maternal lysine, arginine and ornithine levels. During pregnancy this may evoke similar symptoms as reported for lysinuric protein intolerance, which has been associated with serious gestational complications such as increased risk of developing gestational anaemia and PE [47].

SLC38A5, also known as SNAT5 protein, belongs to the Na+-coupled neutral amino acid transporter system N. It mediates the cotransport of glycine, asparagine, alanine, serine, glutamine and histidine with sodium [48]. Prior to the current study, SLC38A5 has only been proven to localize at the MVM and to participate in placental glutamine homeostasis [49]. Our work reports for the first time that SLC38A5 is differentially regulated (increased) at mRNA level in the clinically highly relevant pregnancy diseases IUGR and PE. The trend towards an elevated expression as compared to the control groups was also confirmed on protein level, but, supposedly due to the lower sample size and higher variation in the Western blotting experiments, the comparison between the groups in the ANOVA was not statistically different. In comparison, another sodium-coupled neural amino acid transporter, SLC38A2, which is localized at both the MVM and BM [50], was shown to be significantly decreased in IUGR and PE in the mRNA screening (Fig. 4). The differentially altered expression patterns of SLC38A5 and SLC38A2 in IUGR and PE (up- versus down-regulation) might not only result from their differing cellular localizations but

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**Fig. 6** Protein expression of the membrane transporters ABCA1, SLC7A7 and SLC38A5 in total tissue lysates. ABCA1 (a, 250 kD), SLC7A7 (b, 56 kD), and SLC38A5 (c, 51 kD) protein expressions in total lysates were quantified from control and diseased placentas. Western blotting (upper panels) was performed with 90 μg total lysates from term (n = 4), preterm (n = 5), IUGR (n = 6), PE + IUGR (n = 6), and PE (n = 5) (upper panels). After overnight exposure, an immunoreactive signal was observed in all lanes at the expected size. Relative protein expression levels from western blotting results were then calculated as the densitometric ratio of target versus β-actin (43 kD; lower panels). Data are expressed as median ± range. Significance at p < 0.05 was assessed by a Kruskal-Wallis test, followed by Dunn’s multiple comparison.
also from their distinct transport mechanisms. SLC38A2, as a member of system A, is a pH sensitive uniporter. However, SLC38A5 exhibits marked inhibition at low extracellular pH, because it mediates also the coupled countertransport of H+ with the substrate (e.g. glutamine) [51]. Therefore, the uptake or efflux of glutamine via SLC38A5 may strongly depend on the cellular metabolic status, which could be modified in gestational disorders such as IUGR and PE.

Conclusions

The current study reports a novel analytic strategy including bioinformatic analysis of publicly available microarray datasets, followed by molecular biological and biochemical quantification of mRNA and protein expression in biological tissue samples combined with mathematical diagramming. This approach was successfully applied, firstly to provide an overview on placental membrane transporters providing the fetus with optimal nutrients and microelements for adequate fetal growth, and secondly, to explore the significance of altered membrane transporter expressions in the clinically important pregnancy diseases IUGR and PE. A systematic change in nutrient transporter expression was detected, and 8 membrane transporters were found to be significantly regulated in IUGR/PE. Particularly, the amino acid transporters SLC7A7 and SLC38A5, were influenced by both IUGR and PE. In the present study, we demonstrated for the first time an enhanced mRNA expression of SLC38A5 in IUGR and PE placentae and the same trend on protein expression.
level. These findings are the basis for future investigations which should examine function of the selected transporters at different regions in placenta and elucidate the precise mechanisms regulating the observed changes in IUGR and/or PE.

Additional file

Additional file 1: Gene specific primer sequences used for quantitative RT-PCR. It presents a complete list of the primers designed for the selected candidate and control genes in this study. (DOCX 21 kb)

Abbreviations

ABC: ATP-binding cassette; BM: Basal plasma membrane; Ct: Cycle threshold; GEO: Gene expression omnibus; GLUT: Glucose transporter; IUGR: Intrauterine growth restriction; log (FC): Log 2 fold change; MCL: Markov cluster algorithm; MVM: Microvillous plasma membrane; Na⁺/K⁺ ATPase: Sodium potassium ATPase; PE: Pre-eclampsia; PE combined with IUGR: PE+IUGR; PE: Pre-eclampsia; Pre-eclampsia: PE; SCL: Solute carrier; STRING: Search tool for the retrieval of interacting genes; TRP: Transient receptor potential; UbQ: Ubiquitin; YWHAZ: Tyrosine 3-monooxygenase/tryptophan 5-monooxygenase activation protein, zeta polypeptide

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Availability of data and materials

The datasets analysed during the current study are mined from publicly available in the NCBI GEO repository under the following accession number: GSE 25861 [https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE25861], GSE 24129 [https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE24129], GSE 25906 [https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE25906], GSE 14722 [https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE14722], GSE 35574 [https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE35574], GSE 24818 [https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE24818]. Other datasets supporting the conclusions of this article are not publicly available due to privacy concerns, but are available from the corresponding author on request.

Authors’ contributions

ECO, XH and CA conceived and the study, PA and XH performed the bioinformatic analysis. MLB and DVS performed the clinical assessment of the participants and were involved in the interpretation of data. XH performed the for quantitative RT-PCR experiments, and wrote the manuscript. LH and XH evaluated the protein expressions. All authors read and approved the final manuscript.

Ethics approval and consent to participate

The study protocol was approved by the local ethic committees of the Canton of Bern, Switzerland (178/03). Witten informed consent was obtained from each participant for this research prior to the caesarean section. A copy of the consent form is available for review by the Editor of this journal.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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References

1. Black RE, Allen LH, Bhutta ZA, Caulfield LE, de Onis M, Ezzati M, Mathers C, Rivera J. Maternal, Child Undernutrition Study G. Maternal and child undernutrition: global and regional exposures and health consequences. Lancet. 2008;371(9608):243–60.
2. Wu G, Imhoff-Kunsch B, Girard AW. Biological mechanisms for nutritional regulation of maternal health and fetal development. Paediatr Perinatal Epidemiol. 2012;26(Suppl 1):4–26.
3. Pathology of the human placenta. https://www.springer.com/de/book/9783642239403
4. Manual of Pathology of the Human Placenta. https://link.springer.com/book/10.1007%2F978-1-4419-7494-5
5. Brett KE, Ferraro ZM, Yockell-Lelievre J, Gruslin A, Adamo KB. Maternal-fetal nutrient transport in pregnancy pathologies: the role of the placenta. Int J Mol Sci. 2014;15(9):16135–83.
6. Desorges M, Sibley CP. Placental nutrient supply and fetal growth. Int J Dev Biol. 2010;54(2-3):377–90.
7. Jansson T. Amino acid transporters in the human placenta. Pediatr Res. 2001;49(2):141–7.
8. Christensen HN. Role of amino acid transport and countertransport in nutrition and metabolism. Physiol Rev. 1990;70(1):43–77.
9. Mahendran D, Donnai P, Glazer JD, D’Souza SW, Boyd RD, Sibley CP. Amino acid (system a) transporter activity in microvillous membrane vesicles from the placentas of appropriate and small for gestational age babies. Pediatr Res. 1993;34(5):661–5.
10. Sibley CP, Turner MA, Cetin I, Ayuk P, Boyd CA, D’Souza SW, Glazier JD. Maternal-fetal nutrient transport in pregnancy pathologies: the role of the placenta. Pediatr Res. 2005;58(5):627–32.
11. Glazier JD, Cetin I, Perugini G, Ronzonesi S, Grey AM, Mahendran D, Marconi AM, Pardi G, Sibley CP. Association between the activity of the system a amino acid transporter in the microvillous plasma membrane of the human placenta and severity of fetal compromise in intrauterine growth restriction. Pediatr Res. 1997;42(4):514–9.
12. Johansson M, Karlsson L, Wenngren M, Jansson T, Powell TL. Activity and protein expression of Na+/K+-ATPase are reduced in microvillous syncytiotrophoblast plasma membranes isolated from pregnancies complicated by intrauterine growth restriction. J Clin Endocrinol Metab. 2003;88(6):2831–7.
13. Tsitsiou E, Sibley CP, D’Souza SW, Catanescu O, Jacobsen DW, Glazier JD. Homocysteine transport by systems L, a and y+L across the microvillous plasma membrane of human placenta. J Physiol. 2009;587(Pt 16):4001–13.
14. Tsitsiou E, Sibley CP, D’Souza SW, Catanescu O, Jacobsen DW, Glazier JD. Homocysteine is transported by the microvillous plasma membrane of human placenta. J Inherit Metab Dis. 2011;34(1):57–65.
15. Aye IL, Singh AT, Keelan JA. Transport of lipids by ABC proteins: interactions and implications for cellular toxicity, viability and function. Chem Biol Interact. 2009;180(1-3):327–39.
16. Marconi AM, Paolini C, Buscaglia M, Zerbe G, Battaglia FC, Pardi G. The impact of gestational age and fetal growth on the maternal-fetal glucose concentration difference. Obstet Gynecol. 1996;87(6):937–42.
17. Arkwright PD, Bademacher TW, Dwek RA, Redman CW. Pre-eclampsia is associated with an increase in trophoblast glycogen content and glycogen metabolism. J Clin Endocrinol Metab. 2003;88(6):2831–7.
61. Kanai Y, Segawa H, Miyamoto K, Uchino H, Takeda E, Endou H. Expression cloning and characterization of a transporter for large neutral amino acids activated by the heavy chain of 4F2 antigen (CD98). J Biol Chem. 1998;273(37):23629–32.

62. Aiko Y, Askew DJ, Arabaki S, Myoga M, Tomonaga C, Hachisuga T, Suga R, Kawamoto T, Tsuji M, Shiba M. Differential levels of amino acid transporters system L and ASC72, and the mTOR protein in placenta of preclampsia and IUGR. BM Pregnancy Childbirth. 2014;14:181.

63. Okamoto Y, Sakata M, Ogura K, Yamamoto T, Yamaguchi M, Tasaka K, Kurihara H, Tsurudome M, Murata Y. Expression and regulation of 4F2hc and hLAT1 in human trophoblasts. Am J Physiol Cell Physiol. 2002;282(1):C196–204.

64. Ayuk PT, Sibley CP, Donnai P, D’Souza S, Glazier JD. Development and expression and adaptive regulation of amino acid transporters and regulators in the human placenta. Am J Physiol Cell Physiol. 2000;278(6):C1162–71.

65. Speake PF, Glazier JD, Ayuk PT, Reade M, Sibley CP, D’Souza SW. L-Ariginine transport across the basal plasma membrane of the syncytiotrophoblast of the human placenta from normal and preeclamptic pregnancies. J Clin Endocrinol Metab. 2003;88(9):4287–92.

66. Noris M, Todeschini M, Cassi P, Pastaf F, Cappellini A, Bonazzola S, Macconi D, Maucci R, Porrait F, Benigni A, et al. L-Ariginine depletion in preclampsia orients nitric oxide synthase toward oxidant species. Hypertension. 2004;43(6):14–22.

67. Guzman-Gutierrez E, Westermeier F, Solomon C, Gonzalez M, Pardo F, Leiva A, Sobrevia L. Insulin-increased L-ariginine transport requires a(ZA) adenosine receptors activation in human umbilical vein endothelium. PLoS One. 2012;7(7):e41705.

68. Jones H, Tsurudome M, Murata Y. Expression and regulation of 4F2hc and hLAT1 in human trophoblasts. Am J Physiol Cell Physiol. 2002;282(1):C196–204.

69. Loubiere LS, Vasilopoulou E, Bulmer JN, Taylor PM, Stieger B, Verrey F, Araujo JR, Correia-Branco A, Ramalho C, Goncalves P, Pinho MJ, Keating E, Jones H, Crombleholme T, Habli M. Regulation of amino acid transporters system L and ASC72, and the mTOR protein in placenta of preclampsia and IUGR. BM Pregnancy Childbirth. 2014;14:181.
102. Andrews GK, Wang H, Dey SK, Palmer MD. Mouse zinc transporter 1 gene provides an essential function during early embryonic development. Genesis. 2004;40(2):74–81.

103. Seo YA, Lopez V, Kelleher SL. A histidine-rich motif mediates mitochondrial localization of ZnT2 to modulate mitochondrial function. Am J Physiol Cell Physiol. 2011;300(6):C1479–89.

104. Galvez-Peralta M, He L, Jorge-Nebert LF, Wang B, Miller ML, Eppert BL, Afton S, Nebert DW. ZIP zinc transporter: indispensable role for both multiple-organ organogenesis and hematopoiesis in utero. PLoS One. 2012;7(5):e36055.

105. Hache S, Takser L, LeBellego F, Fuster D, Forest JC, Giguere Y, Leduc L, Weiler H, Hache S, Takser L, LeBellego F, Weiler H, Leduc L, Forest JC, Giguere Y, Marsche G, Sadjak A, Lang U, Desoye G, et al. Human endothelial cells of the placental barrier efficiently deliver cholesterol to the fetal circulation via ABCA1 and ABCG1. Circ Res. 2009;105(4):600–8.

106. Bernucci L, Henriquez M, Diaz P, Riquelme G. Diverse calcium channel types are present in the human placental syncytiotrophoblast basolateral membrane. Placenta. 2006;27(11):1082–95.

107. Romero MF, Chen AP, Parker MD, Boron WF. The SLC4 family of bicarbonate (HCO3)(−) transporters. Mol Asp Med. 2013;34(2-3):159–82.

108. Powell TL, Lundquist C, Doughty IM, Glazer JD, Jansson T. Mechanisms of chloride transport across the syncytiotrophoblast basolateral membrane in the human placenta. Placenta. 1998;19(4):315–21.

109. Kondapalli KC, Kallay LM, Muszelik M, Rao R. Unconventional chemiosmotic coupling of NHA2, a mammalian Na+/H+ antiporter, to a plasma membrane H+ gradient. J Biol Chem. 2012;287(43):36239–50.

110. Donowitz M, Ming Tse C, Fuster D. SLC9/NHE gene family, a plasma membrane Na+/H+ exchanger. J Biol Chem. 2006;281(49):36239–50.

111. Alper SL, Sharma AK. The SLC26 gene family of anion transporters and channels. Mol Asp Med. 2013;34(2-3):494–515.

112. Dawson PA, Rakoczy J, Simmons DG. Placental, renal, and ileal sulfate transporter gene expression in mouse gestation. Biol Reprod. 2012;87(12):5686–8.

113. Mitchell AM, Manley SW, Morris JC, Powell KA, Bergert ER, Mortimer RH. Expression of Na+/H+ exchanger and Cl−/HCO3− anion exchanger mRNA expression in placenta using real-time QPCR. Placenta. 2005;26(1):93–8.

114. Donowitz M, Houston IE, Everett LA, Mori-Aoki A, Suzuki S, Nakamura K, Sakai T, Suzuki K, Royaux IE, Aye IL, Waddell BJ, Mark PJ, Keelan JA, Aye IL, Waddell BJ, Mark PJ, Keelan JA, Burns R, O’Herlihy C, Smyth PP. Regulation of iodide uptake in placental membranes and decidua and the influence of labour at term. Reprod Fertil Dev. 2008;20(2):328–34.

115. Yeboah D, Sun M, Kingdon J, Baczyn D, Lye SJ, Matthews SG, Gibb W. Expression and localization of Na/K ATPase beta 3-subunit expression in the human placenta gene expression in mouse gestation. Biol Trace Elem Res. 2013;154(3):338–47.

116. Yeboah D, Sun M, Kingdon J, Baczyn D, Lye SJ, Matthews SG, Gibb W. Expression of breast cancer susceptibility gene 1 (BRCA1) in human placenta gene expression in mouse gestation. Biol Trace Elem Res. 2013;154(3):338–47.

117. Lacey HA, Nolan T, Greenwood SL, Glazer JD, Sibbey CP. Gestational profile of Na+/H+ exchanger and Cl−/HCO3− anion exchanger mRNA expression in the placenta. J Biol Chem. 2005;280(21):20590–8.

118. Morris JC, Powell KA, Bergert ER, Mortimer RH. The sodium-iodide symporter and type 3 iodothyronine deiodinase gene expression in the placenta. J Clin Endocrinol Metab. 2002;87(2):938–43.

119. Forstner J, Eickhoff J, Czarnetzki BD, UH. Na+/I- symporter and type 3 iodothyronine deiodinase gene expression in the placenta using real-time QPCR. Placenta. 2005;26(1):93–8.

120. Stefulj J, Panzenboeck U, Becker T, Hirschmugl B, Schweizer C, Lang I, Marsche G, Sadjak A, Lang U, Desoye G, et al. Human endothelial cells of the placental barrier efficiently deliver cholesterol to the fetal circulation via ABCA1 and ABCG1. Circ Res. 2009;105(4):600–8.

121. Aye IL, Waddell BJ, Mark PJ, Keelan JA, Aye IL, Waddell BJ, Mark PJ, Keelan JA, Burns R, O’Herlihy C, Smyth PP. Regulation of iodide uptake in placental membranes and decidua and the influence of labour at term. Reprod Fertil Dev. 2008;20(2):328–34.

122. Yeboah D, Sun M, Kingdon J, Baczyn D, Lye SJ, Matthews SG, Gibb W. Expression of breast cancer resistance protein (BCRP) in human fetal membranes and decidua and the influence of labour at term. Reprod Fertil Dev. 2008;20(2):328–34.

123. Mitchell AM, Manley SW, Morris JC, Powell KA, Bergert ER, Mortimer RH. Expression of Na+/H+ exchanger and Cl−/HCO3− anion exchanger mRNA expression in placenta using real-time QPCR. Placenta. 2005;26(1):93–8.

124. Wright EM. Glucose transport families SLC5 and SLC50. Mol Asp Med. 2013;34(2-3):183–96.

125. Mitchell AM, Manley SW, Morris JC, Powell KA, Bergert ER, Mortimer RH. Sodium iodide symporter (NIS) gene expression in human placenta. Placenta. 2001;22(2):256–8.

126. Akhtar M, Won AC, Danisman N, Erkek S, Buyukkagaci U, Unlu E, Tazebay UH. Na+/H+ symporter and type 3 iodothyronine deiodinase gene expression in amniotic membrane and placenta and its relationship to maternal thyroid hormones. Biol Trace Elem Res. 2011;15(1):338–44.

127. Dimachkie MM, Ueda H, Resler D, Cerveny L, Ceckova M, Kacerovska M, Micuda S, Staud F. Organic cation transporter 3 (OCCT/CXCL22A3) and multidrug and toxic compound extrusion protein-1 (MATE1/SLC47A1) is a novel flavinoid transporter. J Agric Food Chem. 2016;64(20):9680–9.

128. Koepsell H, Lipti K, Volk C. Polyspecific organic cation transporters: structure, function, physiological roles, and biopharmaceutical implications. Pharm Res. 2007;24(7):1227–51.

129. Ahmadinaghdaram D, Zemankova L, Nachigal P, Dolezelova E, Neuvanov Z, Cerney L, Ceckova M, Kacerovska M, Micuda S, Staud F. Organic cation transporter 3 (OCCT/CXCL22A3) and multidrug and toxic compound extrusion 1 (MATE1/SLC47A1) transporter in the placental and fetal tissues expression profile and fetus protective role at different stages of gestation. Biol Reprod. 2013;88(3):55.

130. Andersson CM, Stahl A. SLC27 fatty acid transport proteins. Mol Asp Med. 2013;34(2-3):516–28.

131. Jadoon A, Cunningham P, McDermott LC. Regulation of fatty acid binding proteins by hypoxia inducible factors: 1alpha and 2alpha in the placenta: relevance to pre-eclampsia. Prostaglandins Leukot Essent Fatty Acids. 2015;93:25–9.

132. Schaff WT, Bildirici I, Cheong M, Chen PL, Nelson DM, Sadowsky Y. Peroxisome proliferator-activated receptor-gamma and retinoid X receptor signaling regulate fatty acid uptake by primary human placental trophoblasts. J Clin Endocrinol Metab. 2005;90(7):2467–75.

133. Lager S, Jansson N, Olsson AL, Wennegren M, Jansson T, Powell TL. Effect of IL-6 and TNF-alpha on fatty acid uptake by primary human placental trophoblasts. Placenta. 2011;32(2):121–7.

134. Schaff WT, Bildirici I, Cheong M, Chen PL, Nelson DM, Sadowsky Y. Peroxisome proliferator-activated receptor-gamma and retinoid X receptor signaling regulate fatty acid uptake by primary human placental trophoblasts. J Clin Endocrinol Metab. 2005;90(7):2467–75.

135. Mueckler M, Thoren B. The SLC2 (GLUT) family of membrane transporters. Mol Asp Med. 2013;34(2-3):121–38.

136. Jansson T, Wennegren M, Illsley NP. Glucose transporter protein expression in human placenta throughout gestation and in intrauterine growth retardation. J Clin Endocrinol Metab. 1993;77(6):1554–62.
147. Halestrap AP. The SLC16 gene family - structure, role and regulation in health and disease. Mol Asp Med. 2013;34(2-3):337–49.
148. Settle P, Mynett K, Speake P, Champion E, Doughty IM, Sibley CP, D’Souza SW, Glazier J. Polarized lactate transporter activity and expression in the syncytiotrophoblast of the term human placenta. Placenta. 2004;25(6):496–504.
149. Price NT, Jackson VN, Halestrap AP. Cloning and sequencing of four new mammalian monocarboxylate transporter (MCT) homologues confirms the existence of a transporter family with an ancient past. Biochem J. 1998;329(Pt 2):321–8.
150. Settle P, Sibley CP, Doughty IM, Johnston T, Glazier JD, Powell TL, Jansson T, D’Souza SW. Placental lactate transporter activity and expression in intrauterine growth restriction. J Soc Gynecol Investig. 2006;13(5):357–63.
151. Kay HH, Zhu S, Tsoi S. Hypoxia and lactate production in trophoblast cells. Placenta. 2007;28(8-9):854–60.
152. Pramod AB, Foster J, Carvelli L, Henry JK. SLC6 transporters: structure, function, regulation, disease association and therapeutics. Mol Asp Med. 2013;34(2-3):197–219.
153. Bottalico B, Larsson I, Brodzki J, Hernandez-Andrade E, Cassilen B, Marsal K, Hanson SR. Norepinephrine transporter (NET), serotonin transporter (SERT), vesicular monoamine transporter (VMAT2) and organic cation transporters (OCT1, 2 and EMT) in human placenta from pre-eclamptic and normotensive pregnancies. Placenta. 2004;25(8):518–29.
154. Bzoskie L, Yen J, Tseng YT, Blount L, Kashiwai K, Padbury JF. Human placental norepinephrine transporter mRNA: expression and correlation with fetal condition at birth. Placenta. 1997;18(2-3):205–10.
155. Ramamurthy S, Prasad PD, Kulanthavel P, Leibach FH, Blakely RD, Ganapathy V. Expression of a cocaine-sensitive norepinephrine transporter in the human placental syncytiotrophoblast. Biochemistry. 1993;32(5):1346–53.

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