Abstract

Metabolic profiles of amniotic fluid and maternal blood are sources of valuable information about fetus development and can be potentially useful in diagnosis of pregnancy disorders. In this study, we applied $^1$H NMR-based metabolic profiling to track metabolic changes occurring in amniotic fluid (AF) and plasma (PL) of healthy mothers over the course of pregnancy. AF and PL samples were collected in the 2nd (T2) and 3rd (T3) trimester, prolonged pregnancy (PP) until time of delivery (TD). A multivariate data analysis of both biofluids reviled a metabolic switch-like transition between 2nd and 3rd trimester, which was followed by metabolic stabilization throughout the rest of pregnancy probably reflecting the stabilization of fetal maturation and development. The differences were further tested using univariate statistics at $\alpha = 0.001$. In plasma the progression from T2 to T3 was related to increasing levels of glycerol, choline and ketone bodies (3-hydroxybutyrate and acetoacetate) while pyruvate concentration was significantly decreased. In amniotic fluid, T2 to T3 transition was associated with decreasing levels of glucose, carnitine, amino acids (valine, leucine, isoleucine, alanine, methionine, tyrosine, and phenylalanine) and increasing levels of creatinine, succinate, pyruvate, choline, $N,N$-dimethylglycine and urocanate. Lactate to pyruvate ratio was decreased in AF and conversely increased in PL. The results of our study, show that metabolomics profiling can be used to better understand physiological changes of the complex interdependencies of the mother, the placenta and the fetus during pregnancy. In the future, these results might be a useful reference point for analysis of complicated pregnancies.

Introduction

Pregnancy is associated with the onset of many adaptation processes that are likely to change over the course of gestation [1]. In particular, metabolic composition of blood and amniotic fluid should reflect these biochemical dynamics. Maternal blood remains in constant exchange with fetus through the placenta providing nutrients required for growth and development.
Amniotic fluid originates from maternal, fetal and placental tissues therefore its metabolic profile is the net result of metabolite synthesis/degradation, fetal maturation (particularly of the kidneys and lungs), and biochemical exchanges [2,3,4]. Because of their biochemical nature plasma and amniotic fluid should recapitulate the physiological processes of fetal development what makes them an extremely valuable material for fetal health diagnostics [5,6].

NMR-based metabolomics is an established technique for studying complex biological samples (e.g., plasma, urine or amniotic fluid) [7–11]. Metabolomics exploits high-throughput analytical measurements to identify and quantify metabolites allowing to describe dynamic changes in phenotype and system homeostasis [12]. The metabolic profiles of amniotic fluid and the maternal plasma offer new insights to better understand the organ systems and biofunctions that contribute to fetal well-being during normal pregnancy [6,9,13,14]. The great advantage of such an approach is that all of the metabolites (being present in high enough concentration in the biological sample) are measured simultaneously, and a pattern of several metabolites (metabolic profile) can be more informative than the measurement of a single metabolite/analyte [15]. It should be however emphasized that the use of metabolomics in prenatal and perinatal medicine is still in its infancy [9].

Current understanding of the relationship between metabolite composition of amniotic fluid and maternal plasma during normal pregnancy is still not complete. Few studies have monitored healthy pregnancy using either amniotic fluid or plasma but these biofluids have not been studied in combination by metabolomics [16–19]. Other reports have mainly focused on selected stages or disorders of pregnancy [20–22], such as fetal malformations [23], gestational diabetes mellitus [24], macrosomia [25], preeclampsia [26,27], preterm delivery [13,14], spina bifida [28] and very-low birth weight [29], and did not clearly show the changes that occur between trimesters. To the best of our knowledge, the present study is the first combined metabolomics analysis of amniotic fluid and maternal plasma during the progression of normal pregnancy, starting at the 2nd trimester (T2) and continuing through the 3rd trimester (T3), prolonged pregnancy (PP) until time of delivery (TD). The aim of our investigation was to characterize the metabolic signature of human amniotic fluid and maternal plasma over the course of normal pregnancy. This was achieved by analyzing amniotic fluid and maternal plasma using 1H NMR-based metabolomics.

In the study reported in this manuscript the biggest metabolic differences in both amniotic fluid and maternal plasma were detected between 2nd and 3rd trimester suggesting the metabolic switch-like transition occurring in fetus and mother organism.

**Materials and Methods**

**Patients and methods**

This study was conducted at the Department of Obstetrics and Gynaecology, Wroclaw Medical University, Wroclaw, Poland. All of the samples were collected after written informed consent was obtained from the individual women. The study was approved by the Ethics Committee at Wroclaw Medical University, Wroclaw, Poland (protocol number KB-227/2011). Amniotic fluid (AF) was collected by transabdominal amniocentesis under ultrasonographic guidance during the 2nd (T2) and 3rd trimesters (T3), and prolonged pregnancy (PP) until time of delivery (TD). The aim of our investigation was to characterize the metabolic signature of human amniotic fluid and maternal plasma over the course of normal pregnancy. This was achieved by analyzing amniotic fluid and maternal plasma using 1H NMR-based metabolomics.

In the study reported in this manuscript the biggest metabolic differences in both amniotic fluid and maternal plasma were detected between 2nd and 3rd trimester suggesting the metabolic switch-like transition occurring in fetus and mother organism.
In all cases, an accurate gestational age was established by the last menstrual period confirmed by ultrasonographic evaluation. There were no signs of inflammation, vaginal infection or chorioamnionitis. Immediately after amniocentesis, all of the amniotic fluid samples were centrifuged at 3000xg for 20 minutes to separate the cells from the supernatant, which was then aliquoted and stored at −76°C until use. Amniotic fluid samples that were contaminated with blood or meconium were discarded. All of the participating women were healthy (age range 21–38 y.o.) and delivered healthy newborns without malformations, chromosomal abnormalities or postmaturity syndrome.

All of the blood samples (PL) were collected into anticoagulant-treated tubes (sodium citrate) in the morning 12 hours after the last meal and were gently mixed to dissolve the anticoagulant. Next, blood samples were centrifuged at 2000xg for 15 minutes to obtain the plasma, which was then aliquoted and stored at -76°C until use.

Sample preparation
Prior to the analysis, aliquots of human amniotic fluid and plasma were thawed at room temperature for 1 hour. For the NMR measurements, 200 µL of plasma was mixed with 400 µL of saline solution (NaCl 0.9% in 10% D2O). The mixture was centrifuged at 10,000xg for 10 minutes, and then 550 µL of supernatant was transferred into a 5-mm NMR tube. For the amniotic fluid analysis, 200 µL of 0.5 M sodium phosphate buffer (pH = 7 containing 33.3% of D2O, 0.5 mM 3-(trimethylsilyl)-2,2',3,3'-tetradeuteropropionate sodium salt TSP-d4 (TSP) as an internal standard and 3 mM sodium azide to prevent microbial contamination) was added to 400 µL of AF. The mixture was then centrifuged at 10,000xg for 10 minutes, and 550 µL of the supernatant was transferred into a 5-mm NMR tube.

1H NMR measurements
The NMR spectra were recorded using Bruker Avance spectrometer operating at the proton frequency of 600.58 MHz. Sample temperature was set at 300 K. A one-dimensional Carr-Purcell-Meiboom-Gill (CPMG) NMR spin echo pulse sequence with water suppression was employed to filter out broad spectral resonances arising from the macromolecules.

For each AF sample, 256 following scans were collected using 3.5 s relaxation delay and an acquisition time of 1.36 s, resulting in 32k data points and a spectral width of 20.01 ppm. The spectra were processed with a line broadening of 1.0 Hz and were manually phased and baseline-corrected using the Topspin 1.3 software (Bruker, GmBH, Germany). The spectra were referenced to the TSP signal (δ = 0.00 ppm), and alignment was performed using the icoshift algorithm [30]. Finally, after the residual water (4.60–5.00 ppm), urea and α-proton of glucose (5.00–6.00 ppm) regions were removed, the dataset was binned into 7951 integrals of equal width (0.001 ppm).

For each plasma sample, 128 following scans were collected using 3.5 s relaxation delay and acquisition time of 2.73 s, resulting in 32k data points and a spectral width of 20.01 ppm. The spectra were processed with a line broadening of 0.3 Hz and were manually phased and baseline-corrected using the Topspin 1.3 software (Bruker, GmBH, Germany). The whole spectrum was referenced to the α-glucose signal (δ = 5.225 ppm) and alignment was performed using the icoshift algorithm [30]. Finally, after the residual water region (4.40–5.00 ppm) and citrate (2.48–2.69 ppm) were selectively removed, the dataset was binned into 8716 integrals of equal width (0.001 ppm).

Multivariate data analysis and statistical analysis
All of the spectra were normalized using the probabilistic quotient normalization (PQN) method and the Matlab program [31]. The data matrices containing either the binned spectra...
or the areas of the metabolites were transferred into SIMCA-P (v 13.0, Umetrics, Sweden), where the principal component analysis (PCA) was conducted. The data were scaled using Pareto scaling (binned spectra) or unit variance scaling (quantified metabolites). In the case of quantified metabolites (signal areas) only the signals without overlap were used. Orthogonal Projections to Latent Structures-Discriminant Analysis (OPLS-DA) was used for testing the differences between metabolic profiles and considered statistically significant at $\alpha = 0.001$ (CV-ANOVA). STATISTICA software (v 10, StatSoft, Tulsa, USA) was utilized for the statistical analysis of the quantified metabolites. In a univariate analysis, statistical importance was determined using Student’s $t$-test because the data were characterized by normal distribution (S1 Matrix; S2 Matrix). The signal originating from the $\beta$-proton of glucose was removed during the multivariate analysis but considered in univariate statistics. Bonferroni’s correction for multiple testing was used in the statistical analysis thus, a variable was considered statistically significant at $\alpha = 0.001$.

**Results**

Table 1 reports the number of PL, AF and paired samples (AF and PL collected from the same mother) obtained at different epochs of pregnancy and analyzed by $^1$H NMR spectroscopy. Representative examples of the $^1$H NMR spectra of amniotic fluid and plasma are shown in Fig 1A and 1B. All of the spectra included several metabolite groups, such as amino acids, lipids, organic acids, carbohydrates and nucleotides. A total of 34 metabolites in the amniotic fluid and 30 metabolites in the plasma were identified (Table A in S1 File). Additionally, two compounds in each biofluid remained unassigned. For further analysis, the unassigned metabolites in the AF were marked as AU1, AU2, while those in the PL were marked as PU1 and PU2.

The resonances of metabolites were identified using assignments published in the literature [28,32,33] and the on-line databases (www.hmdb.ca) [34]. Generally, inspection of $^1$H NMR spectra of AF and PL divided into the chemical shift regions showed the greatest abundance of the following metabolites: aliphatic region (0–3 ppm), the amino acids (leucine, isoleucine, valine, alanine, threonine (AF), lysine, glutamate (AF), proline, glutamine and methionine (AF)) and the organic acids (2-hydroxybutyrate (AF)), lactate, acetate, pyruvate, succinate (AF) acetoacetate (PL) and citrate (AF)). In all of the analyzed samples, the carbohydrate profiles (3–5.5 ppm) were composed mainly of glucose with an additional small fraction of mannose (PL). In the aromatic region (5.5–10 ppm), the aromatic amino acids (tyrosine, phenylalanine and histidine (AF)) and organic acids (fumarate (AF), formate) were the most abundant. Moreover, all of the amniotic fluid spectra showed the presence of urocanate [6], which was confirmed using Statistical Total Correlation Spectroscopy (STOCSY). Briefly, the resonances putatively assigned to urocanate molecule exhibited high correlation coefficient values in analyzed set of spectra (Fig A in S1 File).

In order to visualize the data a chemometric approach was used. Since plasma and amniotic fluid samples were not paired the unsupervised multivariate data analysis was carried out

| Sampling stages | Mean gestational age at sampling in weeks | PL | AF | Paired samples * |
|-----------------|-----------------------------------------|----|----|----------------|
| T2              | 15.4 ±0.96                               | 7  | 6  | 1              |
| T3              | 37.7±1.68                                | 16 | 21 | 15             |
| TD              | 40.1±1.13                                | 29 | 33 | 26             |
| PP              | 41.3±0.46                                | 11 | 13 | 8              |

*PL and AF samples obtained from the same patient

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separately for amniotic fluid and plasma data sets. In both cases PCA was conducted based only on the metabolite signals that were quantified and additionally based on the whole binned spectra (Fig B in S1 File). Score and loadings plots obtained for both biofluids are shown in Fig 2.

Regardless of analyzed biofluid the observed grouping was similar, with well separated T2 group and high degree of overlap between rest of the samples (T3, TD and PP groups). This was fully achieved by first two principal components (PC). Introduction of additional PCs did not resolve any novel grouping patterns (Fig 2). The analysis of AF loadings plot showed that all of the quantified amino acids were grouped close together and thus exhibit similar pattern (Fig 2A). Conversely, such trend was not observed in PL (Fig 2B).

To further investigate whether metabolic profiles of amniotic fluid and plasma are changing between T2, T3, TD and PP periods a discriminant analysis was employed. The following set of comparisons was subjected to OPLS-DA modeling: T3 vs. T2; TD vs. T3; PP vs. T3; and PP vs. TD. Consistent with PCA results, a clear separation between T2 and all other groups was detected. All three discriminant models involving T2 group (T2 vs T3, T2 vs TD, T2 vs PP) exhibited high level of statistical significance and high values of R2X(cum) and Q2Y(cum) parameters (Table B in S1 File) proving that T2 period is characterized by unique AF and PL metabolic profile. Similar, trend was observed when discriminant analysis was carried out using whole profile (binned spectra) (Table C in S1 File). Conversely, all other comparisons were not-statistically significant (Table B in S1 File) suggesting limited metabolic changes in composition of either AF or PL during transition from T3 to PP period. Again, analogous tendency was observed when whole profile was used for chemometric modeling (Table C in S1 File).

For determination of statistical significance of individual metabolites a univariate statistical analysis was performed with Bonferroni’s correction for multiple testing. Metabolites that were statistically significant in at least one comparison were listed in Table 2. The highest number of significantly affected metabolites was observed in the T3 vs. T2 comparison, in which sixteen metabolites in the AF (leucine, valine, isoleucine, alanine, methionine, tyrosine, phenylalanine, N,N-dimethylglycine, carnitine, creatine, succinate, pyruvate, urocanate, glucose, choline and

Fig 1. A representative example of the 1H NMR spectrum of amniotic fluid (A) and plasma (B). The numbers in the figure correspond to the numbers in Table A in S1 File. 1 –leucine, 2 –valine, 3 –isoleucine, 4 –threonine, 5 –alanine, 6 –lysine, 7 –methionine, 8 –glutamine, 9 –glutamate, 10 –glycine, 11 –tyrosine, 12 –phenylalanine, 13 –histidine, 14 –N,N–dimethylglycine, 15 –creatinine, 16 –creatinine, 17 –choline, 18 –carnitine, 19 –dimethylamine, 20 –sarcosine, 21 –dimethylsulfoxide, 22 –trimethylamine N-oxide (TMAO), 23–laetate, 24 –acetate, 25 –pyruvate, 26 –succinate, 27 –citrate, 28 –urocanate, 29 –fumarate, 30 –formate, 31 –acetocacetate, 32 –2-hydroxyisovalerate, 33–2-hydroxybutyrate, 34 –isobutyrate, 35–3-hydroxybutyrate, 36–3-hydroxyisovalerate, 37 –proline, 38 –N-acetyl groups, 39 –NAC, 40 –VLDL/LDL, 41 –acetone, 42 –methanol, 43 –glycerol, 44 –mannose, 45 –glucose, 46 –AU1, 47 –AU2, 48 –PU1, 49 –PU2.

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AU2) and five metabolites in the PL (pyruvate, 3-hydroxybutyrate, acetoacetate, glycerol, and choline) were found.

Changes in the other comparisons were much less pronounced and limited to AF only. Two metabolites namely (citrate and pyruvate) were significantly down regulated in PP vs T3 comparison, while in PP vs TD carnitine was decreased. No statistically significant changes were detected in TD vs T3 comparison.

It was not possible to perform a correlation analysis due to the fact that most of plasma and amniotic fluid samples were not paired. However, for metabolites that were detected in both biofluids (and significantly changed in at least one comparison) a relationship analysis was conducted by plotting group average and standard deviation values (Fig 3). For pyruvate and choline a strong relationship between changes in plasma and amniotic fluid was observed.

While pyruvate was increasing in AF it was decreasing in PL during T2 to T3 transition. This change was than followed by stabilization in PL pyruvate level and moderate decrease in AF (from T3 to PP). In case of choline a simultaneous significant increase was detected in both AF and PL during T2 to T3 transition. No strong intra-biofluid relationship was found neither for 3-hydroxybutyrate nor for glucose.

Due to availability of lactate and pyruvate in PL and AF, a lactate to pyruvate ratio (L/P) was calculated. The L/P ratio is expressed in arbitrary units since absolute concentrations were
not available. Metabolite quantification was conducted using only signals without overlap thus, calculated L/P ratio values can be used only for relative comparisons and do not reflect physiological level of this parameter. Driven mostly by changes in pyruvate the L/P ratio was significantly diminished in AF and noticeably elevated (although not statistically significant, p value = 0.0066 at \( \alpha = 0.001 \)) in PL in T2 vs T3 comparison. L/P ratio was stable in T3, TD and PP groups in both biofluids. The 3-hydroxybutyrate/acetoacetate (3-HB/AcAc) ratio was calculated only for PL samples, but no differences were detected (data not shown).

**Discussion**

Collection of AF by amniotomy has certain limitations that can influence the composition of amniotic fluid metabolic profile. In particular the risk of introducing substances from vaginal environment should be taken into consideration. Our analysis did not show statistically significant differences between T3 and TD samples, which were collected using transabdominal amniocentesis or amniotomy respectively. Therefore, it is rather unlikely that contaminating metabolites were a substantial constituent of AF metabolic profile measured using \(^1\)H NMR spectroscopy. However, in case of a more sensitive analytical platform e.g. LC-MS such contamination likely will be detected.

The strongest differences in the metabolic composition of both analyzed biofluids were found between the T2 and T3 stages of normal pregnancy (Table 2, Fig 2), while with the

### Table 2. Differences of metabolites (expressed as percentage) between the four gestational age periods.

| Metabolites         | Biofluid | T3 vs. T2 | TD vs. T3 | PP vs. T3 | PP vs. TD |
|---------------------|----------|-----------|-----------|-----------|-----------|
| Leucine             | AF       | -52.81    | 29.34     | 7.17      | -17.14    |
| Valine              | AF       | -62.29    | 24.84     | 2.54      | -17.86    |
| Isoleucine          | AF       | -50.68    | 25.87     | 14.17     | -9.29     |
| Alanine             | AF       | -44.60    | 1.92      | -4.21     | -6.02     |
| Methionine          | AF       | -44.41    | 11.91     | 1.02      | -9.73     |
| Tyrosine            | AF       | -46.70    | 17.81     | 6.78      | -9.36     |
| Phenylalanine       | AF       | -32.40    | 15.78     | 9.37      | -5.54     |
| N,N-dimethylglycine | AF       | 55.26     | -1.89     | -3.54     | -1.68     |
| Carnitine           | AF       | -44.37    | -2.00     | -17.65    | -15.97    |
| Creatinine          | AF       | 84.92     | -5.84     | 1.99      | 8.32      |
| Citrate             | AF       | 1.69      | 7.23      | -26.18    | -31.16    |
| Succinate           | AF       | 114.06    | -0.57     | -11.19    | -10.68    |
| Pyruvate*           | AF       | 238.48    | -22.64    | -36.35    | -17.74    |
| Pyruvate*           | PL       | -45.37    | 13.27     | 5.34      | -6.99     |
| Acetoacetate        | PL       | 61.32     | 2.88      | 16.52     | 14.04     |
| Urocanate           | AF       | 672.58    | 7.19      | 15.23     | 7.50      |
| 3-hydroxybutyrate    | PL       | 101.48    | 22.84     | -15.85    | -31.50    |
| Glycerol            | PL       | 49.83     | 1.48      | 0.95      | 0.52      |
| Glucose             | AF       | -50.24    | 24.23     | -13.42    | -30.31    |
| Choline*            | AF       | 32.71     | 0.84      | 11.23     | 10.31     |
| Choline*            | PL       | 176.58    | 7.34      | 6.11      | -1.15     |
| AU2                 | AF       | 304.66    | -8.97     | -9.95     | -1.08     |
| Lac/Pyr             | AF       | -72.34    | 20.38     | 42.56     | 18.42     |

The percent difference was calculated using the mean values of the relative signal integrals in each group. Bold type indicates a statistical significance of \( p < 0.001 \) according to Student’s \( t \)-test.

* Metabolites present in both biofluids.

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progression of pregnancy, metabolic stabilization becomes evident. This interesting finding suggests a metabolic switch-like transition occurring in mother’s system between T2 and T3 periods. Therefore, our observations are most likely related to the change in fetus growth dynamics, namely transition into fast weight-gain phase, which requires considerably higher rates of anabolic processes. In contrast to plasma, the amniotic fluid showed significant decrease in the levels of amino acids during transition mentioned above (Table 2, Fig B in S1 File). This is likely associated with fetal maturation and the increased demand for elementary building blocks, which are necessary for protein synthesis [35] and might be utilized for many other processes required to maintain fetal homeostasis during rapid growth [36]. Amino acids are crucial for proper energy balance and maintenance of TCA cycle span by providing carbon backbone exchange via various anaplerotic and cataplerotic pathways [37]. It is known that a shortage of amino acids, may strongly influence fetal protein biosynthesis [36].

It is established that during first two-thirds of gestation the mother’s organism is in anabolic condition, while during the T3 maternal metabolism is switched towards catabolic activity [38], which is than opposite from the intensive anabolic processes occurring in fetus.

An example of catabolic/anabolic switch is the decrease in the level of glucose in amniotic fluid, which overlaps with increasing energy demand during the pregnancy progression. It is known that mothers can become hypoglycemic even though their gluconeogenesis rates are increased [39]. This is due to fetus high glucose uptake through the placenta. Therefore, small decrease in plasma glucose levels in T3 might be attributed with depleting glycogen stores, though influence of diet cannot be excluded.

The significantly increased mother’s plasma glycerol in T3 vs T2 comparison is most likely a result of triglycerides breakdown caused by high lipolitic activity of adipose tissue [38]. Glycerol can be further utilized either as a substrate for gluconeogenesis and triglycerides biosynthesis [38] or to help to maintain proper redox balance being a part of glycerol phosphate shuttle [40]. Interestingly it was suggested that glycerol can be a preferential substrate for gluconeogenesis during pregnancy [41] however, since mentioned study compared glycerol with pyruvate and alanine, it was more an evaluation of pyruvate carboxylase anaplerotic flux. Triglycerides hydrolysis results in a high concentration of free fatty acids that undergo beta oxidation which is supplementing TCA cycle and ketone bodies synthesis by providing acetyl-CoA moieties. Indeed, we observed significantly elevated levels of 3-hydroxybutyrate and acetacetate in T3 vs T2 comparison [42]. In contrast, carnitine, which is required for activation of fatty acids and relegating them towards beta oxidation in mitochondria was found to be consequently decreasing throughout the course of pregnancy (Table 2). Moreover, the sufficient level of carnitine was postulated to have an important role in fetal growth maturation by increased birth weight and postnatal growth rate [43].

TCA cycle is a nexus point of central carbon metabolism elegantly balancing amino acids, carbohydrates and lipids metabolism [44]. Besides free fatty acids, pyruvate is one of the major sources of acetyl-CoA for TCA cycle. Pyruvate fluctuation in plasma moderately corresponded to the level of glucose, which was higher in T2 than in the subsequent periods of pregnancy. Surprisingly, the pyruvate level in the amniotic fluid exhibited a reverse trend against glucose (Fig 3A and 3B).

All discussed pathways: beta-oxidation, ketone bodies production, glycolysis and TCA cycle are linked via oxidative phosphorylation to cellular redox state [45]. In vivo redox readout can
be challenging, yet one of the most common approaches is utilization of redox pairs. In particular lactate/pyruvate (L/P) and 3-hydroxybutyrate/acetoacetate (3-HB/AcAc) ratios are proven to be valuable. L/P is considered to reflect cytosolic, while 3-HB/AcAc mitochondrial redox potential. There is an inverse relationship between L/P and oxygen consumption [46]. Albeit not being statistically significant after Bonferroni’s correction for multiple testing (p value = 0.0066 at α = 0.001), the mothers plasma L/P ratio was noticeably elevated between T2 and T3 and stayed unchanged in TD and PP. It has been reported that pregnant women have decreased levels of pyruvate in blood comparing to non pregnant control resulting in elevated L/P ratio [47]. Moreover, L/P is even further elevated in twin comparing to singleton pregnancies [48] due to anaerobic metabolism. Interestingly, opposite to plasma, amniotic fluid L/P ratio was significantly (p = 0.0002) decreased between T2 and T3. Decrease in L/P ratio is considered to correspond to the elevated oxygen consumption thus, our observations might reflect the increase in fetus aerobic metabolism. In conclusion, reverse L/P trends in AF and PL suggest that transition from 2nd to 3rd trimester is coupled with the change in oxygen utilization by fetus and mother body.

Changes in choline concentration were evident in both the AF and the PL. Increased choline level might be correlated with higher phospholipid demand for the fetus brain development during later periods of pregnancy [49]. The same trend, but for amniotic fluid only, was found for N,N-dimethylglycine, which is an intermediate metabolite in glycine’s biosynthesis from choline.

Creatinine, a marker of the function and maturation of fetal kidneys [50], increased significantly during T3 and then remain stable once the kidneys have fully matured. Note that the composition of AF reflects the fetal renal system, thus, the degree of fetal renal maturation should significantly influence the level of almost all of the metabolites in the amniotic fluid [50,2,51].

To the best of our knowledge, this is for the first time that urocanate has been observed in amniotic fluid using 1H NMR. We confirmed presence of urocanate by STOCSY analysis [6, 52]. This metabolite is an important photoprotectant predominantly located in the skin. It is generated from histidine by enzyme histidinase and contributes to acidic mantle [53]. Throughout most of pregnancy fetus skin is transparent thus high levels of urocanate in AF in T2 might reflect rapid growth of skin and requirement for photoprotectants.

It is worth noticing that even though amniotic fluid samples were not paired with plasma samples a high degree of agreement in metabolic trends was observed. This was especially true in case of transition from T2 to T3 period (Fig 3). Correlation between plasma and amniotic fluid metabolites opens possibilities to investigate fetus metabolic status by mother’s plasma metabolomics monitoring. It is highly probable that more metabolic trends could be identified by using paired samples. Yet, for medical and ethical reasons, it is extremely difficult to obtain amniotic fluid from healthy mothers.

Even though in our study the biggest differences were observed between T2 and T3, we cannot exclude the possibility that even more pronounced transition occurs between T1 and T2. A substantial decrease in lysophosphatidylcholines and lysophosphatidylethanolamines plasma content was found between T1 and T2 [18]. Furthermore, a number of amino acids, non-esterified fatty acids and acylcarnitines was shown to change between 1st and 2nd trimester of healthy pregnancy [19].

Conclusions

The second trimester was easily distinguished from the other investigated periods while no significant discrimination could be achieved between other groups. A metabolic switch-like
transition in both biofluids was discovered between second and third trimester, which was followed by metabolic stabilization throughout the rest of pregnancy. To the best of our knowledge, this study provides the first metabolomics analysis of amniotic fluids and plasma collected from healthy pregnant women at different stages of pregnancy and during delivery. In the future, these results might be a useful reference point for analysis of complicated pregnancy and potentially transfer to some extent to the clinical arena.

Supporting Information

S1 File. All figures and tables of Supporting Information. Legend: Fig A. STOCSY analysis of the amniotic fluid exhibited high correlations between the urocanate signals. Fig B. The PCA score plot results obtained for the ¹H NMR data corresponding to the pregnancy stages. Green circles=prolonged pregnancy (PP), blue squares=delivery (TD), red triangles=3rd trimester (T3) and yellow inverted triangles=2nd trimester (T2). Table A. Metabolites identified in amniotic fluid and plasma using ¹H NMR spectroscopy. Table B. Parameters of OPLS-DA model (two components) calculated using quantified metabolites (signal areas). Table C. Parameters of OPLS-DA model (two components) calculated using binned spectra.

S1 Matrix. The data matrix of AF, which contains all necessary data needed to replicate experiment.

S2 Matrix. The data matrix of PL, which contains all necessary data needed to replicate experiment.

Author Contributions

Conceived and designed the experiments: PM MO-P. Performed the experiments: EJ SD. Analyzed the data: SD PM EJ AZ. Contributed reagents/materials/analysis tools: PM LH. Wrote the paper: SD PM EJ MO-P.

References

1. Lain KY, Catalano PM. Metabolic changes in pregnancy. Clin Obstet Gynecol. 2007; 50: 938–948. PMID: 17982337
2. Underwood MA, Gilbert WM, Sherman MP. Amniotic fluid: not just fetal urine anymore. J Perinatol. 2005; 25: 341–348. PMID: 15861199
3. Lind T. The biochemistry of amniotic fluid. In: Sandler M, ed. Amniotic fluid and its clinical significance. 1st ed. New York: Marcel Dekker Inc; 1981
4. Briese V, Kunkel S, Plath C, Wutzke KD, Plesse R. Sialic acid, steroids and proteohormones in maternal, cord and retroplacental blood. Z Geburtshilfe Neonatol. 1999; 203: 63–68. PMID: 10420512
5. Michaels J-EA, Dasari S, Pereira L, Reddy AP, Lapidus JA, Lu X et al. Comprehensive proteomic analysis of the human amniotic fluid proteome: gestational age-dependent changes. J Proteome Res. 2007; 6: 1277–1285. PMID: 17373941
6. Menon R, Jones J, Gunst PR, Kacerovsky M, Fortunato SJ, Saade GR et al. Amniotic fluid metabolic analysis in spontaneous preterm birth. Reprod Sci. 2014; 21: 791–803. doi: 10.1177/1933719113518987 PMID: 24440998
7. Silwood CJ, Lynch E, Claxson AW, Grootveld MC. 1H and 13 C NMR spectroscopic analysis of human saliva. J Dent Res. 2002; 81: 422–427. PMID: 12097436
8. Zhang J, Wei S, Liu L, Nagana Gowda GA, Bonney P, Stewart J et al. NMR based metabolomics study of canine bladder cancer. Biochim Biophys Acta 2012; 1822: 1607–1614. doi: 10.1016/j.bbadis.2012.08.001 PMID: 22867815
9. Fanos V, Atzori L, Makarenko K, Melis GB, Ferrazzi E. Metabolomics application in maternal-fetal medicine. Biomed Res Int. 2013; 2013:720514. doi: 10.1155/2013/720514 PMID: 23841090

10. Graça G, Moreira AS, Correia AJ, Goodfellow BJ, Barros AS, Duarte IF et al. Mid-infrared (MIR) metabolic fingerprinting of amniotic fluid: A possible avenue for early diagnosis of prenatal disorders? Anal Chim Acta 2013; 764: 24–31. doi: 10.1016/j.aca.2012.12.023 PMID: 23374211

11. Duarte IF, Diaz SO, Gil AM. NMR metabolomics of human blood and urine in disease research. J Pharm Biomed Anal. 2014; 93: 17–26. doi: 10.1016/j.jpba.2013.09.025 PMID: 24854435

12. Krastanov A. Metabolomics—the art of art. Biotechnol &Biotechnol Eq. 2010; 24: 1537–1543.

13. Romero R, Mazaki Tovi S, Vaisbuch E, Kusanovic JP, Chaiworapongsa T, Gomez R et al. Metabolomics in premature labor: a novel approach to identify patients at risk for preterm delivery. J Matern Fetal Neonatal Med. 2010; 23: 1344–1359. doi: 10.3109/14767058.2010.482618 PMID: 20504069

14. Tea I, Le Gall G, Küster A, Guignard N, Alexandre-Gouabau MC, Darmaun D et al. 1H-NMR-Based Metabolic Profiling of Maternal and Umbilical Cord Blood Indicates Altered Materno-Foetal Nutrient Exchange in Preterm Infants. PLoS One 2012; 7: e29947. doi: 10.1371/journal.pone.0029947 PMID: 22291897

15. Bock JL. Metabolic profiling of amniotic fluid by proton nuclear magnetic resonance spectroscopy: correlation with fetal maturation and other clinical variables. Clin Chem. 1994; 40: 56–61. PMID: 8267545

16. Cohn BR, Fukuchi EY, Joe BN, Swanson MG, Kurhanewicz J et al. Calculation of gestational age in pregnancy by NMR metabolic profiling of amniotic fluid: A possible avenue for early diagnosis of prenatal disorders? Anal Chim Acta 2013; 764: 24–31. doi:10.1016/j.aca.2012.12.023 PMID: 23374211

17. Pinto J, Barros AS, Domingues MR, Goodfellow BJ, Galhano E et al. Following Healthy Pregnancy by Maternal Plasma. J Proteome Res. 2014; 13: 1527–1536. doi: 10.1021/pr401068k PMID: 24450375

18. Luan H, Meng N, Liu P, Feng Q, Lin S et al. Pregnancy-Induced Metabolic Phenotype Variations in Maternal Plasma. J Proteome Res. 2014; 13: 1527–1536. doi: 10.1021/pr401068k PMID: 24450375

19. Lindsay KL, Hellmuth C, Uhl O, Buss C, Wadhwa PD et al. Longitudinal Metabolic Profiling of Amino Acids and Lipids across Healthy Pregnancy. PLoS One. 2015; doi:10.1371/journal.pone.0127418.

20. Graça G, Duarte IF, Barros AS, Goodfellow BJ, Diaz SO, Pinto J et al. Impact of prenatal disorders on the metabolic profile of second trimester amniotic fluid: a nuclear magnetic resonance metabolomic study. J Proteome Res. 2010; 9: 6016–6024. doi: 10.1021/pr100151q PMID: 20949080

21. Graça G, Diaz SO, Pinto J, Barros AS, Duarte IF, Goodfellow BJ et al. Can Biofluids Metabolic Profiling Help to Improve Healthcare during Pregnancy? Spectroscopy: An International Journal 2012; 27: 515–523.

22. Diaz S, Pinto J, Graca G, Duarte I. Metabolic Biomarkers of Prenatal Disorders: An Exploratory NMR Metabonomics Study of Second Trimester Maternal Urine and Blood Plasma. J Proteome Res. 2011; 10: 3732–3742. doi: 10.1021/pr200352m PMID: 21649438

23. Graça G, Duarte IF, Barros AS, Goodfellow BJ, Diaz S, Carreira IM et al. 1H NMR-Based Metabolomics of Human Amniotic Fluid for the Metabolic Characterization of Fetal Malformations. J Proteome Res. 2009; 8: 4144–4150. doi: 10.1021/pr900386f PMID: 19453139

24. Sachse D, Sletner L, Markid K, Jenum AK, Birkeland KI, Rise F et al. Metabolic changes in urine during and after pregnancy in a large, multiethnic population-based cohort study of gestational diabetes. PLoS One 2012; 7:e52399. doi:10.1371/journal.pone.0052399 PMID: 23285025

25. Ciborowski M, Zbucka-Kretowska M, Bomba-Opon D, Wielgos M, Brawura-Biskupski-Samaha R, Pierzynski P et al. Potential first trimester metabolic biomarkers of abnormal birth weight in healthy pregnancies. Prenat Diagn. 2014; 34: 870–877. doi: 10.1002/pd.4386 PMID: 24733416

26. Bahado-Singh RO, Akolekar R, Mandal R, Dong E, Xia J, Kruger M et al. Metabolomics and first-trimester prediction of early-onset preeclampsia. J Matern Fetal Neonatal Med. 2012; 25: 1840–1847. doi: 10.3109/14767058.2012.680254 PMID:22494326

27. Austdal M, Skråstad RB, Gundersen AS, Austgulen R, Iversen AC, Bathen TF. Metabolomic biomarkers in plasma and urine in women with preeclampsia. PLoS One 2014; 9:e81923. doi:10.1371/journal.pone.0081923 PMID:23850250

28. Groenen PM, Engelke UF, Wevers RA, Hendriks JC, Eskes TK, Merkus HM et al. High-resolution 1H NMR spectroscopy of amniotic fluids from spina bifida fetuses and control. Eur J Obstet Gynecol Reprod Biol. 2004; 112: 16–23. PMID: 14687733

29. Alexandre-Gouabau MC, Courant F, Moyon T, Küster A, Le Gall G, Tea I et al. Maternal and cord blood LC-HRMS metabolomics reveal alterations in energy and polyamine metabolism, and oxidative stress in very-low birth weight infants. J Proteome Res. 2013; 12: 2764–2778. doi: 10.1021/pr400122v PMID: 23527880
30. Savorani F, Tomasi G, Engelsen SB. Icoshift: A versatile tool for the rapid alignment of 1D NMR spectra. J Magn Reson 2010; 202: 190–202. doi: 10.1016/j.jmr.2009.11.012 PMID: 20004603

31. Dieterle F, Ross A, Schlottbeck G, Senn H. Probabilistic quotient normalization as robust method to account for dilution of complex biological mixtures. Application in 1H NMR metabonomics. Anal Chem 2006; 78: 4281–4290. PMID: 16808434

32. Graça G, Duarte IF, Goodfellow BJ, Barros AS, Carreira IM, Couceiro AB et al. Potential of NMR spectroscopy for the study of human amniotic fluid. Anal Chem. 2007; 79: 8367–8375. PMID: 17918968

33. Graça G, Duarte IF, Goodfellow B, Carreira IM, Couceiro AB, Domingues Mdo R et al. Metabolite profiling of human amniotic fluid by hyphenated nuclear magnetic resonance spectroscopy. Anal Chem. 2008; 80: 6085–6092. doi: 10.1021/ac800907f PMID: 18564856

34. http://www.ncbi.nlm.nih.gov/pubmed/17202168

35. Athanasiadis AP, Michaelidou AM, Fotiou M, Menexes G, Theodoridis TD, Ganidou M et al. Correlation of 2nd trimester amniotic fluid amino acid profile with gestational age and estimated fetal weight. J Matern Fetal Neonatal Med. 2011; 24: 1033–1038. doi: 10.3109/14767058.2010.545909 PMID: 21271782

36. Levy HL, Montag PP. Free amino acids in human amniotic fluid. A quantitative study by ion-exchange chromatography. Pediat. Res. 1969; 3: 113–120 PMID: 5771792

37. Xi L, Brown K, Woodward J, Shim K, Johnson B, Odle J. Maternal Dietary l-carnitine supplementation influence fetal carnitine Status and Stimulates Carnitine Palmitoyltransferase and Pyruvate Dehydrogenase Complex Activities in Swine. J Nutr. 2008; 12: 2356–2362.

38. Sun F, Dai C, Xie J, Hu X. Biochemical Issues in Estimation of Cytosolic Free NAD/NADH Ratio. PLoS One. 2012; 7: 1–10.

39. Madeira VMC. Overview of Mitochondrial Bioenergetics. Mitochondrail Bioenergetics. 2011; 810: 1–6.

40. Zeisel SH. Choline: Critical Role During Fetal Development and Dietary Requirements in Adults. Annu Rev Nutr. 2006; 26: 229–250. PMID: 16848706

41. Oliveira FR, Barros EG, Magalhães JA. Biochemical profile of amniotic fluid for the assessment of fetal and renal development. Braz J Med Biol Res. 2002; 35: 215–222. PMID: 11847525

42. Imbard A, Blom HJ, Schlummer D, Barto R, Czerkiewicz I, Rigal O et al. Methylation metabolites in amniotic fluid depend on gestational age. Prenat Diagn. 2013; 33: 848–855. doi: 10.1002/pd.4142 PMID: 23613283

43. Fluhr JW., Elias PM, Man M-Q, Hupe M, Selden C et al. Is the Filaggrin-Histidine-Urocanic Acid Pathway Essential for Stratum Corneum Acidification? J Investig Dermatolog. 2010; 130: 2141–2144.