RSC Advances



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PAPER



Cite this: RSC Adv., 2019, 9, 20796

Received 10th May 2019 Accepted 19th June 2019 DOI: 10.1039/c9ra03525g

rsc.li/rsc-advances

Introduction

Traditional Chinese Medicine (TCM) has been guided by its basic theories and has made significant contributions to the development of medicine and healthcare. In TCM theories, there are several different disharmony patterns (named ZHENG) such as the syndrome of blood stasis, syndrome of Qi stagnation, and Yin/Yang deficiency syndrome.¹ As one of the syndromes of blood stasis, cold coagulation is caused by coldevil. In traditional Chinese medicine theories, cold leads to

Exploring the pharmacological effects and potential targets of paeoniflorin on the endometriosis of cold coagulation and blood stasis model rats by ultra-performance liquid chromatography tandem mass spectrometry with a pattern recognition approach[†]

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This study was employed to explore the potential biomarkers of endometriosis of cold coagulation and blood stasis (ECB) model rats and the effective mechanism of action of paeoniflorin (PF). The serum metabolomics approach was carried out using the UPLC-MS technique with a pattern recognition approach to prove the possible biomarkers of the ECB model rats and the perturbed pathways. Subsequently, the mechanism of PF treatment of this disease model was elucidated. The results revealed that the serum metabolism profiles in two groups were also separated significantly. Moreover, 8 biomarkers were found in the positive mode, and 5 biomarkers were found in the negative mode. Totally, 13 biomarkers participated in the metabolism of phenylalanine, arachidonic acid, *etc.* After treatment with PF, 10 biomarkers were regulated. Among the 10 biomarkers, 4 were statistically significant: L-phenylalanine, L-tryptophan, LysoPC (18:4(6*Z*,9*Z*,12*Z*,15*Z*)), and LysoPC (16:1(9*Z*)). We initially confirmed that PF could significantly regulate the metabolic expression of multiple metabolic pathways in the ECB model rats. For the first time, this study explored the mechanism of action of PF treatment based on the metabolic pathways of the organism and demonstrated the potential of the metabolomics techniques for the study of drug action mechanisms.

blood clotting, and blood clotting leads to blood stasis; therefore, cold-evil easily leads to the syndrome of blood stasis,^{2,3} which has a serious influence on females during the menstrual period; moreover, it can cause various problems in women in the reproductive age. In addition, endometriosis is a grim challenge that may lead to female infertility. Generally, endometrium covers the luminal surface of the uterus. However, the endometrium can grow in other parts of the body due to certain factors, which can further develop into endometriosis. Thus, this disease is defined as the presence of endometrial stroma and glands;4 approximately 10-15 women of reproductive age per 100 people suffer from this disease.⁵⁻⁹ Pelvic pain, secondary dysmenorrhea and infertility¹⁰ are the main clinical manifestations of endometriosis. Regardless of any clinical manifestation, it is a severe threat to the physical and mental health of females of childbearing age. Therefore, the study of the etiology, pathogenesis, and treatment of endometriosis has become an important issue in the medical profession.

As the emerging "omic" technologies, metabolomics involves the integrated analysis of low-molecular-weight metabolites of a particular biological sample or an organism.^{11,12} The concept of metabolomics originated from the analysis of metabolic profiles proposed in the 1970s.¹³ With the

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[†] Electronic supplementary information (ESI) available. See DOI: 10.1039/c9ra03525g

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development of genomics, transcriptomics and proteomics, Fiehn¹⁴ proposed the concept of metabolomics in 1997,¹⁵ and in 1999, Nicholson¹⁶ et al. also proposed the concept of metabolomics. The emergence of metabolomics provides many valuable research methods and strategies for the development of traditional Chinese medicine. The worth of metabolomics originates from the wide diversity of low-molecular-weight endogenous metabolites, which independently or in synergies provide characteristic profiles for different human diseases such as cardiovascular, neurological and inflammatory diseases;17 moreover, studies associated with metabolomics can be beneficial to strengthen our insights into disease mechanisms and drug effects, heightening our ability to forecast individual differences in drug reaction phenotypes.18,19 On the basis of previous studies and after years of constant exploration and research, in 2001, Wang²⁰ proposed a novel strategy to address key points, for example, the property of syndromes, the evaluation of clinical efficacy of prescriptions, and the discovery of the basis of pharmacodynamics-chinmedomics (Chinese medicine metabolomics); the purpose of this study was to discover and determine the material basis for the efficacy of Chinese medicine and solve the quality problems, such as the effectiveness and safety of traditional Chinese medicines, related to the material basis of the efficacy.21,22 Therefore, highthroughput and high-resolution technologies, multivariate data analysis software and robust tools are necessary to support the large-scale data analysis. Currently, hyphenated techniques, such as the LC-MS (liquid chromatography-mass spectrometry) technology, are widely used for the determination and characterization of potential biomarkers in the metabolomics system owing to not only the high reproducibility and good linear range of the LC, but also the high definition of MS.^{23,24} Among these, ultra-performance liquid chromatography-tandem mass spectrometry (UPLC-MS) is one of the most commonly used approaches in assessing the efficacy of herbal medicine due to its high sensitivity and rapid analysis speed.^{25,26} However, the amount of raw data obtained through the UPLC-MS analysis is very large, and conventional statistical methods make it difficult to find differences among the samples or groups and the molecules that cause these differences. Thus, it is indispensable to apply special methods to analyze metabolomics data. Pattern recognition methods have been used to deal with this issue, which mainly include unsupervised methods and supervised methods. Principal components analysis (PCA) belongs to the former methods, whereas the orthogonal single correction partial least squares-discriminant analysis (OPLS-DA) and partial least squares-discriminant analysis (PLS-DA)27 belong to the latter methods, which are commonly used to filter out irrelevant information that interferes with metabolite analysis results during the metabolomics research.

The ingredients of traditional Chinese medicine are complicated, and the mechanism *via* which a substance exerts its efficacy remains unclear; *Radix Paeoniae*, the dried root of *Paeonia lactiflora* Pallas, is a Chinese herb. Paeoniflorin is an important active monomer of *Radix Paeoniae*. However, the pharmacological effect of paeoniflorin on ECB remains unclear. Therefore, in this study, ultra-performance liquid chromatography-electrospray ionization/quadrupole-time-offlight high-definition mass spectrometry with the pattern recognition approach was used to characterize the metabolites present in the serum of ECB rats after oral administration of paeoniflorin, and the therapeutic mechanisms of paeoniflorin towards ECB were elucidated.

Methods and materials

Chemicals and reagents

Paeoniflorin was obtained from Chengdu PUSH Bio. Technology Co., Ltd (batch number PS160803-06). Estradiol benzoate injection was purchased from Animal Pharmaceutical Hangzhou, China (batch number 110202511). Acetonitrile (HPLC grade) was supplied by Fisher (USA). Methanol was obtained from Fisher (USA). Formic acid (HPLC grade) was purchased from Kermel (Tianjin, China). Leucine enkephalin was supplied by Sigma-Aldrich (MO, USA). Deionized water was supplied by Watson's Food & Beverage Co., Ltd. (Guangzhou, China). All chemicals and reagents were of analytical purity.

Animals and groups

Female Wistar rats, weighing 200–240 g, were obtained from the GLP Center of Heilongjiang University of Chinese Medicine. The room temperature was set at 24–26 °C with $40 \pm 5\%$ relative humidity under 12 h light/12 h dark cycle. Before the experiment, all rats in the metabolism cages were acclimatized for a week. During this period, rats were provided with water and normal food freely. We divided the rats into 5 groups stochastically: sham groups 1 and 2 (control group), ECB groups 1 and 2 (model group), and paeoniflorin group (therapy group), with 10 rats in each group. The experimental procedures were approved by the Animal Care and Ethics Committee at Heilongjiang University of Chinese Medicine, and all experiments were performed in accordance with the declaration of Helsinki.

Animal experiments and preparation of the PF oral solution

The method endometrial auto-transplantation combined with ice water immersion was applied to prepare the ECB model.²⁸ The animal experimental procedures were executed during estrus. Moreover, the entire surgical procedure was executed under sterile environments. The rats in the five groups were anesthetized with 0.2 mL/100 g sodium pentobarbital, which placed in a supine posture and fastened them; then, routine disinfection of the skin was conducted using medical cotton dipped in an appropriate quantity of 10% povidone-iodine solution. To reach the abdominal cavity, 2-3 cm midabdominal incision was made. Then, by ligating the left uterine horn in both sides, about 1 cm tissue was cut and longitudinally opened. The sample freshly cut was put into a sterile container prefilled with 0.9% sodium chloride solution and cut into 2 pieces. The pieces were about 4 mm \times 4 mm large. Moreover, two part organizations were respectively sutured into the uterine and abdominal wall of the same rat with USP 6/0 polyglactin. After 2 mL of 0.9% sodium chloride solution was administered into the abdominal cavity, the abdominal wall was sutured. Subsequently, the midline incision was closed by a 3/0 suture line. Following this, the rats were injected with 0.5 mL gentamicin each day by intraperitoneal injection for 3 consecutive days. Subsequently, 0.1 mL estradiol was administered to rats by subcutaneous injection once every 3 days, for 3 times totally. The control group rats were merely performed an abdominal incision without tissue transplantation. Except the sham group rats, other rats were immersed in 0–1 °C ice water for 8–10 min for about 14 consecutive days. Prior to this experiment, the rats were recovered for two weeks owing to the surgery.

After ice water immersion, the treatment group was intragastrically administrated with paeoniflorin of 27.6 mg kg⁻¹ body weight. An aliquot of distilled water was given to all the rats except the paeoniflorin group. The administration process lasted for 28 days. The paeoniflorin standard powder was dissolved in distilled water and then ultrasonically mixed for 30 min to prepare an oral solution.

Calculation of the ectopic tissues volume

The rats were anesthetized with sodium pentobarbital (0.2 mL 100 g⁻¹). The ectopic tissues were separated, and the length, width and height of the grafts were measured using the vernier caliper. The graft volume was calculated using the formula V (mm³) = 0.52 × A × B × C.

Hematoxylin-eosin staining (H&E staining)

The ectopic endometrium approximately $1 \text{ mm} \times 1 \text{ mm}$ was removed and fixed in 10% formaldehyde immediately, then, it was embedded in paraffin and sectioned. The paraffin sections were fixed with the dye (hematoxylin and eosin) for histopathological analysis. The characteristics of tissue morphology in each group were observed using a microscope.

Collection of blood and preparation of serum samples

Blood was collected from the hepatic portal vein in the model group 2 and control group 2 at the last day of treatment. Subsequently, the samples were centrifuged at 4000 rpm for 15 min, and the supernatant was transferred into a clean tube for use as a serum sample. The sample was stored at -80 °C. Prior to analysis, the serum samples were thawed and vortexed. The 100 µL serum sample was carefully aspirated, and 300 µL methanol was added to the sample in order to remove proteins. After vortexing, the liquid was centrifuged at 13 000 rpm at 4 °C for 10 min. The supernatant was added to an appropriate amount of methanol in the tube and dried with nitrogen in the water bath at 40 °C. Then, 200 µL methanol was used to reconstitute the residue, followed by centrifugation 13 000 rpm at 4 $^\circ C$ for 10 min again and filtered using a 0.22 μm membrane. Finally, the contents were transferred to vials for UPLC-MS analysis.

Analysis conditions of serum samples

Chromatography conditions. An Acquity[™] UPLC (Waters, USA) and the Masslynx software were used for the process of

chromatographic analysis. The isolation of section was implemented using an ACQUITY UPLCTM phenomenex column (2.1 nm × 50 mm, 1.7 µm). The column temperature was set at 40 °C, and the optimal mobile phase contained (A) acetonitrile with 0.1% formic acid and (B) water with 0.1% formic acid in the mobile phase. The gradient elution program for UPLC analysis started from: 0–4 min, 2–40% A; 4–10 min, 40–70% A; 10–13 min, 70–100% A; 13–16 min, 100% A; 16–17 min, 100–2% A; The elution velocity was 0.4 mL min⁻¹. Then 3 µL of serum sample in the positive and negative modes was injected into the instrument, respectively.

MS conditions. The electrospray ionization source was conducted in the positive and negative ion mode in order to acquire data. The analysis parameters are as follows: ion source temperature, 110 °C; gas temperature, 350 °C; and desolvation gas flow, 750 L h⁻¹. The capillary voltage was 1.5 kV in the negative mode and 1.3 kV in the positive mode. The sample cone voltage was set at 50 V and 45 V in positive and negative ion modes, respectively. The setting of the data acquisition rate was 0.2 s per scan. Data were gathered from 50 to 1500 Da in the centroid mode. A lock-mass of 0.2 ng mL⁻¹ leucine enkephalin was used as a reference compound to monitor ion for $[M - H]^-$ = 554.2615 (negative ion mode) and $[M + H]^+$ = 556.2771 (positive ion mode) to obtain an accurate mass.

Data processing. All of the raw data were imported into the Progenesis QI software (V 2.1, Waters Corporation, MA) for achieving proper alignment, peak-picking and normalization. Then, resultant data matrices of previous step were imported to the Ezinfo software (V 3.0) for multivariate statistical analysis with pattern recognition methods. Then, the data matrix was generated to obtain information such as retention time, m/z, and peak area. The cluster profile of groups was analyzed by principal components analysis (PCA). Using Ezinfo, we acquired the variable importance in projection (VIP) values and the relative intensity of markers by orthogonal single correction partial least squares-discriminant analysis (OPLS-DA) score. We explored the characteristics across groups by the *t*-test.

Biomarkers identification. Via extracting the serum samples by the UPLC-MS system, we obtained metabolic profiles of the rats. The raw data were input into the Progenesis QI software. Then, a serial workflow containing vector editing, ion intensity detection, and base peaks ion intensity chromatogram was run to detect the small molecular species existing in ECB markers. A sequential components' statistical analysis was complied, and the .usp profile was obtained for the following data processing. Metabolic data matrices between sham group and ECB group were analyzed by OPLS-DA. A preliminary VIP value was selected for preliminary screening of potential biomarkers. The changes in the content of these potential markers were counted, and the secondary screening was carried out according to the principle that each metabolite in the control group and the model group had significant difference. The metabolites satisfying both the abovementioned conditions could be considered the possible markers. Then the candidate ions were identified by the serum biomarkers identification method, match mol file and secondary mass spectrometry information. Finally, potential biomarkers of the ECB rat model were obtained.

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Metabolic pathway analysis. Metabolomics Pathway Analysis (MetPA) was mainly used to find the metabolomic pathways associated with metabolites according to the analysis of pathway topological characteristics. The name of the compound, KEGG or HMDB number of the detected biomarkers were introduced into the website http://www.metaboanalyst.ca for metabolic pathway analysis. The metabolic pathways relevant to the ECB model were obtained and plotted, and then, schematic drawing of the respective metabolic pathways was obtained. The impact threshold was set to 0.10. Any pathway above this threshold was classified as a potential target pathway.

Statistical analysis. Data were conveyed by SPSS (V 21.0) for statistical analysis and expressed as means \pm SD. By two-sample Student's *t*-test, differences in average value were calculated to be statistically significant; p < 0.05 was considered statistically significant, and p < 0.01 indicated an extremely significant difference.

Results

Assessment of endometriosis of the cold coagulation and blood stasis rat model

Control group 1 and model group 1 were used to assess this model. The ectopic lesion volume of the control group 1 and model group 1 showed a significant difference, as shown in Fig. 1. Obviously, we observed that the ectopic tissue of the model group 2 showed a cystic vesicle with a transparent or translucent liquid inside. The results of histopathological observation (Fig. 2) in the control group 2 showed that the endometrial structure was intact; the gland was rich, and the structure was clear; the glandular epithelial cells were high columnar, arranged neatly and tightly, and the cytoplasm was rich; the interstitial cells were fusiform, arranged neatly and evenly distributed; the blood vessels were rich. In the model group 2, the epithelial cells of ectopic endometrium were short columnar and partially pseudostratified, and the cell morphology was incomplete. Compared with the case of the intimal tissue of the control group 2, the number of interstitial cells and glands increased significantly, and the blood vessels were abundant. Thus, this model was successfully established.

The therapeutic effects of PF on the endometriosis of the cold coagulation and blood stasis model rats

The control group 2, model group 2 and PF group were used to explore the therapeutic effects for this model. Fortunately, after treatment with PF, the volumes of the ectopic endometrial tissue reduced or the tissue disappeared significantly when compared with the case of the model group 2. The data of the ectopic lesion volume are shown in Table 1. The results of the H&E staining (Fig. 2) for the PF group show that the ectopic endometrial tissue has different degrees of atrophy: the cystic cavity shrinks, the capsule wall becomes thinner, and the glandular epithelial cells become short columnar or flat, partially degenerated, necrotic or shedding, and loosely arranged with irregular nuclei and uncertain position; the number of glands decreases or the glands even disappear, showing atrophy, and the inflammatory cells appear to ooze; the number of mesenchymal cells is reduced, and these cells are small, sparse, and even absent, with reduced blood vessels and fibrosis in some areas.

Metabolic profiling analysis

Metabolomics analysis was executed on the metabolic profile data of the serum samples obtained from normal rats and ECB model rats on the 28th day of modeling (Fig. 3). The serum profiles of the BPI chromatograms of the abovementioned two groups were roughly similar. However, only several peaks were adjusted. Using the unsupervised pattern recognition method-PCA, the score plots that reflected the trend between groups were obtained (Fig. 4). It can be seen from the analysis results



Con.

Mod.

GY4

Fig. 2 Endometriosis effect of shape and structure of ectopic endometrial cells in the different groups after oral administration of PF for 28 days (100× magnification). Con.: control group; Mod.: model group; GY4.: PF group.

Table 1 Effect of endometrium grafts volume in serum of GFW $(\bar{X}\pm {\rm SD})^a$

Group	Pats/n	Ectopic tissue volume/mm			
Gloup	Kats/n	Ectopic tissue volume/mm			
Control group 1	10	0.00 ± 0.00			
Model group 1	8	$87.85\pm5.09^{\Delta\Delta}$			
Control group 2	10	0.00 ± 0.00			
Model group 2	9	92.87 ± 8.54			
PF group	9	$80.07 \pm 7.12^{**}$			

 a Data are expressed as means \pm SD. Compared with normal control group 1, $^{\Delta}P$ < 0.05; compared with model group 2, *P < 0.05, **P < 0.01.

that the normal groups in the positive (Fig. 4A) and negative ion mode (Fig. 4B) cluster in the same region, and the rats in the model group are clustered, whereas the rats in the normal group are distinctly distinguished. This is due to the effect of external factors on modeling. The metabolic profile has changed.

Determination of serum biomarkers for the ECB model

To further distinguish between the control group and the model group, the supervised OPLS-DA was employed to divide samples into two clusters in the positive mode (Fig. 5A) and negative mode (Fig. 5B). In our study, the OPLS-DA score plot analysis separated the serum samples from the control group and the model group completely based on the difference in their metabolic profiles. The significant variables were founded through the VIP plot. If the VIP value of the variable is greater than 1, the variable can contribute significantly to classification. The intensity of ions with higher VIP values was selected from the *S*-plot and the VIP-plot in the positive (ESI Fig. 1A and 2A†) and negative mode (ESI Fig. 1B and 2B†). Then, we selected ions with a VIP value greater than 1 and a *t*-test *p* value less than 0.05 as biomarker candidate ions after statistical analysis of the data. Finally, the metabolites with high-resolution MS and MS/MS fragments were further analyzed using the databases HMDB, ChemSpider and KEGG.

Using the abovementioned methods, free metabolites, such as related metabolites, and mass spectra were searched, and a total of 13 blood biomarkers associated with the ECB rat model were identified, among which eight were positive ions in the positive ion mode, namely, L-benzene L-phenylalanine, cervonoyl ethanolamide, LysoPC (16:1(9Z)), LysoPC (18:2(9Z,12Z)), LysoPE (16:0/0:0), LysoPC (16:0), LysoPC (18:1(9Z)), and LysoPC (15:0), and five were negative ions in the negative ion mode: Ltryptophan, hippuric acid, 5-HETE, *etc.* The specific information is shown in Table 2 and ESI Table 1.† The statistical results



Fig. 3 UPLC-MS base peaks ion (BPI) intensity chromatograms of ECB model rats in control group (A) and model group (B) by positive and negative ion modes.



Fig. 4 Score plot of serum profile of sham group and model group scanned by positive and negative ion modes. (Data were analyzed by PCA.) Note: A: In the positive ion mode, B: in the negative ion mode; Con: the sham group, Mod: the model group.

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Fig. 5 Score plot of serum profile of sham group and model group scanned by positive and negative ion modes. (Data were analyzed by OPLS-DA.) Note: A: in the positive ion mode, B: in the negative ion mode; Con: the sham group, Mod: the model group.

of the changes in the microscopic content of latent biomarkers in the model group and sham group are shown in the ESI Fig. 3.[†]

Metabolic pathway analysis

The identification of 13 blood biomarkers by chemical name, KEGG or HMDB number in the website http:// www.metaboanalyst.ca for the Met-PA analysis indicated that six metabolic pathways were disordered: phenylalanine, tyrosine and tryptophan biosynthesis, glycerophospholipid metabolism, phenylalanine metabolism, tryptophan metabolism, arachidonic acid metabolism, and aminoacyl-tRNA biosynthesis, as shown in the ESI Fig. 4.† According to the Met-PA analysis results, the impact threshold of phenylalanine, tyrosine and tryptophan biosynthesis, phenylalanine metabolism and tryptophan metabolism was above 0.1. Therefore, they were the target pathways. The specific data results are shown in the ESI Table 2.† The results revealed that these endogenous compounds produced strong perturbations in the entire metabolic trace and were closely related to the ECB model.

Therapeutic effect of PF on the ECB model rats

The ESI-MS metabolic profile of the control group, model group, and PF group serum on the 28th day of administration was input into the Progenesis QI software for processing. Subsequently, these ions were processed by supervised PLS-DA to obtain score plots (ESI Fig. 5†) that could reflect the trend between groups. The control group and the ECB model group showed obvious clustering and separation. The results indicated that the serum metabolic profile of rats was significantly different from that of the normal group after the model was established. After administration, the model rats had

Table 2 Serum potential biomarkers related to ECB identified in positive and negative ion mode^a

No.	<i>R</i> t min	Determined mass	Calc mass	Actual mass	$\left[M-H\right]^{\!-\!}/\!\left[M+H\right]^{\!+\!}$	mDa	ppm	Proposed composition	Postulated identity	HMDB number	Trend
1	1.36	166.0869	166.0868	165.0789	$[M + H]^{+}$	0.1	0.6	C ₀ H ₁₁ NO ₂	L-Phenylalanine	HMDB00159	$\uparrow \Delta \Delta$
2	1.73	203.0827	203.0821	204.0898	[M – H]-	0.6	3	$C_{11}H_{12}N_2O_2$	L-Tryptophan	HMDB00929	ĻΔ
3	2.20	178.0504	178.0504	179.0582	$[M - H]^{-}$	0	0	C ₉ H ₉ NO ₃	Hippuric acid	HMDB00714	†ΔΔ
4	4.79	514.293	514.2934	515.3011	$[M - H]^{-}$	-0.4	-1	$C_{26}H_{46}NO_7P$	LysoPC (18:4(6Z.9Z.12Z.15Z))	HMDB10389	$\downarrow \Delta \Delta$
5	5.51	373.2745	373.2473	372.2664	$[M + H]^+$	0.2	0.5	$C_{24}H_{36}O_{3}$	Cervonoyl ethanolamide	HMDB13627	$\downarrow \Delta \Delta$
6	7.25	494.3242	494.3247	493.3168	$[M + H]^+$	-0.5	-1	C24H48NO7P	LysoPC (16:1(9Z))	HMDB10383	$\downarrow \Delta \Delta$
7	7.54	520.34	520.3403	519.3324	$\left[M + H\right]^{+}$	-0.3	-0.6	C ₂₆ H ₅₀ NO ₇ P	LysoPC (18:2(9Z,12Z))	HMDB10386	$\downarrow \Delta$
8	7.71	500.279	500.2777	501.2855	$[M - H]^{-}$	1.2	2.4	$\mathrm{C}_{25}\mathrm{H}_{44}\mathrm{NO}_{7}\mathrm{P}$	LysoPE (0:0/20:4 (8Z,11Z,14Z,17Z))	HMDB11488	$\downarrow \Delta \Delta$
9	8.13	454.2939	454.2934	453.2855	$[M + H]^{+}$	0.5	1.1	C ₂₁ H ₄₄ NO ₇ P	LysoPE (16:0/0:0)	HMDB11503	$\downarrow \Delta \Delta$
10	8.23	496.3404	496.3403	495.3324	$[M + H]^+$	0.1	0.2	C24H50NO7P	LysoPC (16:0)	HMDB10382	$\downarrow \Delta \Delta$
11	8.64	522.3563	522.356	521.3481	$[M + H]^{+}$	0.3	0.6	C ₂₆ H ₅₂ NO ₇ P	LysoPC (18:1(9Z))	HMDB02815	$\downarrow \Delta$
12	8.99	319.2278	319.2273	320.2351	$[M - H]^{-}$	0.5	1.6	$C_{20}H_{32}O_3$	5-HETE	HMDB11134	$\uparrow\Delta\Delta$
13	9.61	482.0325	482.3247	481.3168	$[M + H]^+$	0.3	0.6	$C_{23}H_{48}NO_7P$	LysoPC (15:0)	HMDB10381	$\downarrow\!\Delta\Delta$

 a $\uparrow \downarrow$ indicates that the level of the marker is increased or decreased in the urine of rats in the ECB model group; compared with the control group, $\Delta P < 0.05$, $\Delta \Delta P < 0.01$.

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a tendency similar to that of the control group; after 28 days of drug administration, by analyzing the trend of the ECB model serum biomarkers, we found that PF could affect the microbiological changes of potential biomarkers in the ECB model rats, and the content of these biomarkers tended to approach that of the blank control group. Altogether, 13 biomarkers were associated with the ECB model; among these, 10 were aminoglycoside, namely L-phenylalanine, L-tryptophan, hippuric acid, cervonoyl ethanolamide, 5-HETE, *etc*; moreover, 4 biomarkers were statistically significant. The detailed results are shown in the ESI Fig. 6 and Table 3.[†]

Discussion

Metabolism is a general term for a series of ordered chemical reactions that occur in living organisms to sustain life, which produces a series of metabolites. Metabolites are generally excreted through various body fluids. Hence, some clinically readily available biological fluids, such as urine, blood, and saliva²⁹ extracts, have become the common samples for metabolomics research. After the introduction of metabolomics, it has developed very rapidly. It has been extensively applied in disease diagnosis³⁰ and animal models, Traditional Chinese Medicine (TCM) research,³¹ toxicology,^{32,33} nutrition science34 and other research fields. Metabolomics methods and technical systems have matured and formed a complete set of research systems and strategies. A wide range of applications, its methods and technical systems are becoming more mature and form the complete research system and strategy. Metabolomics mainly analyzes the changes of small-molecular-weight metabolites of a specific biological sample or an organism^{35,36} caused by changes in the body's physiological and pathological phenomena. It uses some high-precision analytical methods and chemometrics methods to identify patterns in the diseased and normal groups to help understand the process of disease. We explored the changes of metabolites and their metabolic pathways in the body and searched for biomarkers related to diseases, which can be used for the early diagnosis and treatment of diseases, pathogenesis and prognosis. Generally, when a biological system is subjected to an exogenous stimulus or an endogenous disease, the small-molecule metabolites may change, or they may change with time, and the metabolomics techniques investigate the metabolic pathways in biological systems by this change. The specific research content is roughly divided into the following four aspects: sample processing, sample testing, data analysis, and retrieval of commonly used databases. It has been reported that paeoniflorin ameliorates acute renal injury37 and alpha-naphthylisothiocyanate (ANIT)induced cholestasis,38 suppressing inflammation.39 Moreover, paeoniflorin was identified as a quality marker (Q-makers) of Tangzhiqing tablet40,41 and Buyanghuanwu injection.42 These studies prove that it is of great significance to carry out some pharmacological studies on paeoniflorin.

In this study, a rapid, high-sensitivity and high-throughput technology, *i.e.* UPLC-Q/TOF-MS, was used to clearly discriminate rats with ECB from matched controls and confirm the potential biomarkers. Herein, 13 biomarkers associated with

the ECB disease model (5 in the negative ion mode and 8 in the positive ion mode) were identified. The biomarkers, such as LysoPC (15:0), LysoPC (16:0), LysoPC (16:1(9Z)), LysoPC (18:1(9Z)) and LysoPC (18:2(9Z,12Z)), found herein were derived from phosphatidylcholine (PC).43 LysoPE (0:0/20:4(8Z,11Z,14Z,17Z)) and LysoPE (16:0/0:0) are the metabolic products of phosphatidylethanolamine (PE).44 Both PC and PE are structural components of cell membranes,43,44 and they can be generated by phospholipase A2.44,45 Another pathway for LysoPC formation occurs by the transfer of one fatty acid of PC to cholesterol by lecithin-cholesterol acyltransferase (LCAT), which is an enzyme that converts free cholesterol into cholesteryl ester. Cholesteryl ester is a more hydrophobic form of cholesterol that is sequestered in the liver.⁴⁶ LysoPC (15:0), LysoPC (16:0), LysoPC (16:1(9Z)), LysoPC (18:1(9Z)), LysoPC (18:2(9Z,12Z)), LysoPE (0:0/20:4(8Z,11Z,14Z,17Z)) and LysoPE (16:0/0:0) were discovered in the ECB disease model involved in the glycerophospholipid metabolism, which were in accordance with previous studies.47,48 5-HETE was found in the serum of ECB rats, which participated in the arachidonic acid metabolism. In the cases of lipoxygenases and cyclooxygenases, acid49 is first converted to 5-hydroarachidonic peroxyeicosatetraenoic acid (5-HPETE).50 It can be reduced to 5hydroxyeicosatetraenoic acid (5-HETE) by ubiquitous glutathione peroxidase (GPx).50 It is one of the eicosanoids. Eicosanoids have been shown to be involved in the inflammation reaction^{51,52} in previous studies. Due to ectopic endometrial tissue, it is accompanied by an inflammatory reaction. Therefore, we speculated that PF may inhibit the inflammatory response by a certain concentration of 5-HETE in serum through the arachidonic acid metabolic pathway. In previous reports,³⁹ PF was also proved to have the effect of suppressing inflammation. The results indicated that PF probably regulated the level of 5-HETE through the arachidonic acid metabolism to exert its anti-inflammation effect for ECB.

In addition, we also discovered that amino acid metabolism (phenylalanine metabolism, tryptophan metabolism) was obviously disturbed by endometriosis of cold coagulation and blood stasis. This may be due to the fact that endometriosis results from physiological mechanism of tissue injury and repair.53 The catabolic state induced in response to injury in endometriosis leads to increased breakdown of endogenous protein and release of free amino acids in circulation. It has been reported that several amino acids in tissues of women with endometriosis are found to be dysregulated. The levels of phenylalanine in serum were higher, which further confirms our experimental results.54 After treatment, phenylalanine and tryptophan were reversed from the ECB group to the control group. Thus, we concluded that paeoniflorin may be used to treat the disease by regulating the catabolic state of the amino acid and changing the concentration of phenylalanine and tryptophan in the serum.

Biomarkers have become an integral part of the research on diseases.⁵⁴⁻⁵⁹ In this study, it was found that PF regulated 10 biomarkers by the metabolic pathways of phenylalanine, tyrosine and tryptophan biosynthesis, phenylalanine metabolism, and tryptophan metabolism, to achieve a certain therapeutic effect. Although the systematic metabolic analysis was used in our study, we can integrate the physiological indicators and pathological observations to improve this study.

Conclusions

On the basis of metabolomics technology coupled with a pattern recognition approach, we identified small-molecule metabolites in rat serum. Characterization of the profiles and pattern recognition analysis were carried out. It has been proved that PF influences the metabolic profile changes of the ECB model rats. Using the OPLS-DA pattern recognition method combined with multivariate statistical analysis, the results obtained indicated changes in 13 metabolites, which might be involved in 3 target metabolic pathways: phenylalanine, tyrosine and tryptophan biosynthesis, phenylalanine metabolism, and tryptophan metabolism. After treatment with PF, 10 biomarkers had been regulated: L-phenylalanine, L-tryptophan, hippuric acid, LysoPC (18:4(6Z,9Z,12Z,15Z)), cervonoyl ethanolamide, LysoPE (0:0/20:4(8Z,11Z,14Z,17Z)), LysoPC (16:0), LysoPC (18:1(9Z)), LysoPC (16:1(9Z)), and 5-HETE. Our study has elucidated the mechanism of action of PF treatment from the metabolic pathways of the organism and demonstrated the potential of the metabolomics techniques for the study of drug action mechanisms, which provides a better understanding of ECB pathogenesis and effective metabolic pathways for the PF treatment of ECB.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This research was supported by the grants received from the National Natural Science Foundation of China (Grant No. 81373931, 81573877), Innovative Talent Supporting Plan of Heilongjiang University of Chinese Medicine (Grant No. 2012jc01), Young Talent Lift Engineering Project of China Association of Traditional Chinese Medicine (QNRC2-B06), and Natural Science Foundation of Heilongjiang Province (YQ2019H030).

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