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Introduction

The *Zhi-zi-chi* decoction (*ZZCD*), composed of *Gardenia jasminoides* Ellis (*ZZ*) and Semen Sojae Praeparatum (*DDC*), is a famous traditional Chinese medicine formula (TCMP) described in Treatise on Cold Damage Diseases (*Shang-Han-Lun*). Clinical applications over a long period of time have proved that *ZZCD* is effective in treating depression with low side effects.¹ *In vitro* studies have shown that *ZZCD* contains a variety of components, such as iridoid glycosides, isoflavones, and organic acids.² An increasing amount of research has shown that the pharmacological activity of this decoction is ascribed to the superimposed effects of the prototype components,³ metabolites or both in *vivo*⁴ *via* regulating different signaling pathways. Thus, sound research about the metabolic profile of *ZZCD in vivo* is of great significance.

Up to now, the metabolism of *ZZCD* in plasma, bile and urine has been studied.^{2,5} However, the metabolism of *ZZCD* in feces has never been explored yet. Due to the existence of gut

Metabolic profile analysis of *Zhi-zi-chi* decoction in feces of normal and chronic unpredictable mild stress-induced depression rats based on UHPLC-ESI-Q-TOF-MS/MS and multiple analytical strategies †

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Zhi-zi-chi decoction (*ZZCD*) has been verified by clinical application that it has definite curative effects and low side effects on depression. Because it is administered orally, the metabolites of *ZZCD* in the intestinal tract may influence the curative effects significantly. In this study, UHPLC-ESI-Q-TOF-MS/MS was used in combination with untargeted metabolomics-driven strategy, series product ion filtering and diagnostic fragment ion strategy for acquiring the comprehensive metabolic profile of *ZZCD* in feces of normal and chronic unpredictable mild stress (CUMS)-induced depression rats after oral administration, while the rat depression model was evaluated by behavior tests and plasma biochemical indices. Finally, a total of 56 compounds, including 35 prototype compounds and 21 metabolites, were identified or tentatively characterized in fecal samples. Among these, ten compounds were sieved as potential chemical markers that would reflect the antidepressant effect of *ZZCD*, which may offer important information for quality assessment, pharmacokinetic study and clinical security. In conclusion, the metabolic profile of *ZZCD* in normal and CUMS-induced depression rats would be helpful for the further study of anti-depression material basis and mechanism.

microbiota, the intestinal tract can act as an organ with a metabolic potential at least equal to that of the liver,^{6,7} particularly for TCMPs that are administered orally. The gut microbiota express various enzymes that can metabolically activate, inactivate, or reactivate drugs.7-12 Some chemical components of ZZCD, such as geniposide, caffeic acid, daidzin, and dihydrodaidzein, can be biotransformed by various enzymes from gut microbiota.7,13-16 The reactions include hydrolysis, oxidation, reduction, isomerization, rearrangement, esterification, and condensation.17,18 This indicates that microorganisms may be important for the metabolism of ZZCD. Moreover, researches claimed that the disposition of drugs in disease models is different from normal ones.19,20 If there are significant alterations in gut microbiota in a depressed living organism,²¹⁻²³ then the metabolic profile of chemical components from ZZCD in the intestinal tract would change. In view of the correlation among the chemical components of ZZCD, gut microbiota and depression, it is significant to reveal the pharmacodynamic material basis of ZZCD for studying its metabolic in feces of normal and depression rats.7,21-24

In view of the interference of a great variety of compounds and microorganisms in feces of a living organism, it is difficult to study the analytes of interest.²⁵⁻²⁷ For avoiding the metabolism of the microorganisms present in the rat feces samples from affecting the composition of the sample under study,

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multiple analytical strategies were adopted in this study. Liquid chromatography mass spectrometry (LC-MS) has become an efficient tool for acquiring the structural information of chemical substances in biological samples due to its high separation ability and high resolution.^{11,28} Nowadays, with the improvements in high-resolution multistage mass spectrometry and related software, such as UHPLC-Q-TOF-MS/MS,29 the information of traditional Chinese medicine prescription (TCMP) metabolites transformed by various metabolic pathways could be acquired in one injection cycle by independent data acquisition (IDA) criteria,30 which is an important data acquisition technique for acquiring the global structural information of samples. The data acquisition in IDA is alternant in low and high level of collision energy (CE). When the CE level is low, the MS spectrum is acquired, while the MS/MS spectrum is acquired under high level of CE. The parent ions are not prescreened; therefore, the global fragment ions of the chemical components in the samples can be obtained. Moreover, the XCMS online platform (xcmsonline.scripps.edu) is one of the effective tools for data processing.31,32 It is a cloud-based informatic platform, which allows users to easily upload and process liquid chromatography/mass spectrometry-based data with only a few steps. Then, a complete untargeted metabolomic workflow including feature detection, retention time correction, alignment, annotation, statistical analysis, and data visualization provided by XCMS online is operated.³³ As a result, threedimensional datasets including peak codes, m/z-retention time $(t_{\rm R})$ pairs and ion intensity are acquired,³⁴ which can be downloaded as zip files for offline analysis.33 Additionally, offline data mining technologies, such as series product ion filtering (sPIF),³⁵ diagnostic fragment ion strategy (DFIS),^{36,37} principal component analysis (PCA),³⁴ orthogonal partial least square discriminant analysis (OPLS-DA)35 and extracted ion chromatogram (XIC) provided by SIMCA-P (13.0, Umetrics, Umea, Sweden)³⁸ and Peakview[™] software (AB Sciex)³⁹ could be useful for the screening and identification of the complex chemical components and metabolites.

In this study, the global metabolic profile of *ZZCD* in the feces of normal rats and CUMS-induced depression rats was characterized, while multiple identification strategies based on UHPLC-Q-TOF-MS/MS and chemometrics were used. The exogenous constituents in rat feces could be screened and identified comprehensively. Synchronously, the correlation between depression and metabolic compounds of *ZZCD*, which could show its significance, has been examined in this study.

Experimental

Chemicals and materials

All plant materials were purchased from a Tongrentang chain drugstore (Bengbu, China), while *Gardenia jasminoides* Ellis (*ZZ*) (no. 170731) and Sojae Semen Praeparatum (*DDC*) (no. 170803) were collected from Jiangxi and Anhui, China respectively, which were authenticated by Professor Xian Li (Department of Chinese Materia Medica, Bengbu Medical University, Bengbu, China).

Methanol (HPLC-grade) was purchased from Merk (Darmstadt, Germany), while formic acid (HPLC-grade) was purchased from Sigma Chemical (St. Louis, USA). Fresh ultra-pure water was obtained from a Milli-Q system (Bedford, USA). The standards of chlorogenic acid, geniposide, *p*-cumaric acid and daidzein (purity > 98%) were purchased from Mansite Biotechnology Company (Chengdu, China). All other chemicals and reagents were analytical grade or higher.

Preparation of ZZCD sample

The *ZZCD* sample was prepared according to the original composition and preparation recorded in the Treatise on Cold Pathogenic Diseases (*Shang-Han-Lun*). The powder of *ZZ* (15 g) was decocted in water (300 mL) by boiling. Then, *DDC* (10 g) was added while the volume of water was reduced to about 200 mL. The decoction was boiled further until the water volume reduced to about 100 mL. The extract was filtered by six layers of gauze and concentrated to 50 mL. Then, the solution was labeled and stored at -20 °C until analysis.

For the LC-MS analysis, an aliquot of 2.5 mL of ZZCD was diluted to 10 mL with water and vortexed for 5 min. Then, 3 mL of the diluent was diluted to 10 mL with ethanol and vortexed for 5 min. After placing the mixed liquor for 24 h at 4 $^{\circ}$ C, it was centrifuged at 12 000 rpm for 10 min and then, the supernatant was filtered through a 0.22 μ m membrane.

Animal experiments

Prescribed by the guide of relevant national legislation and local guidelines, the animal experiments were executed. In brief, male Sprague-Dawley rats (5–6 weeks, 150 \pm 20 g) were acquired from the laboratory animal center of Bengbu Medical University (Bengbu, China). In total, 24 rats were adapted in the animal breeding room for one week, with regulated temperature (20–25 °C), humidity (40–70%), and a 12 h light/dark cycle. Purified water and food were provided *ad libitum*. All rats were fasted with free access to water for 12 h prior to the experiment.

Based on the result of the sucrose preference test (SPT) in week 2, rats were divided into three groups randomly with eight in each group. The experimental design can be seen in Fig. 1. Rats in control and normal + ZZCD groups received no treatment, while rats in the CUMS + ZZCD group were subjected to chronic unpredictable mild stress (CUMS) for 3 weeks. Besides, these rats were fed separately, each in a single cage. The protocol of CUMS consisted of ten stressors: forced swimming in ice-water for 10 min, food deprivation for 24 h, water deprivation for 24 h, noise stimulation for 1 h, restraint stress for 2 h, overnight illumination for 8 h, clipping tail for 5 min, horizontal oscillation for 20 min, cage tilting at 45° for 24 h, and a soiled cage environment (600 mL water added to 300 g sawdust bedding) for 24 h. All rats were subjected to various behavioural tests in week 4, including SPT, open-field test (OFT) and forced swimming test (FST).

ZZCD was orally administrated to both the normal + *ZZCD* group and CUMS + *ZZCD* group at a dose of 2.625 g kg⁻¹ d⁻¹ for three continuous days. Then, each rat was fed in its respective metabolic cage. Fecal samples were collected from 0 h to 24 h in

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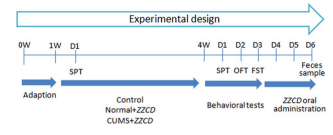


Fig. 1 Experimental schedule of the present study (4 weeks).

one bottle per rat. After drying in the darkness on a filter paper and grinding into powder at ambient temperature, the samples were labeled and stored at -20 °C until analysis.

Depression-like behaviours and biochemical analysis

Depression-like behaviours were examined using SPT, openfield test (OFT) and forced swimming test (FST). In SPT, rats were given purified water and 1% sucrose solution *ad libitum* for 1 h at the same time. Then, the preference to 1% sucrose was calculated by the consumption of purified water and 1% sucrose solution: preference (%) = consumption of 1% sucrose solution/(consumption of 1% sucrose solution + consumption of purified water). In OFT, square boxes (length: 50 cm, width: 50 cm, height: 25 cm), which were divided to 9 squares on average, were used. The score of each rat was based on the time of square-crossing and standing in 3 min. In FST, rats were allowed to swim for 6 min, while the time in immobility in last 4 min was recorded.

For biochemical analysis, the plasma of each rat was collected. In short, blood samples of about 1.0 mL were collected in heparinized tubes from the orbital venous plexus, and centrifuged at 3000 rpm for 5 min. Then, the supernates were stored at -20 °C. For analysis, 0.5 mL supernates were acquired for the enzyme-linked immunosorbent assay (Jianglaibio, China). As a result, the concentration of serotonin (5-HT) and dopamine (DA) in plasma levels were measured.

Preparation of feces samples

For the extraction and concentration of chemical components in rat feces, all fecal samples were prepared as follows. Initially, 5 mL 50% methanol-water was chosen as the solvent for dissolving 0.5 g of each sample. Then, ultrasonic treatment at 40 kHz and centrifugation at 4500 rpm were performed for 30 min and 10 min, respectively, in sequence. After transferring to another glass tube, the supernatant was dried to residues by a gentle stream of nitrogen at 30 °C. For LC-MS analysis, 200 μ L 50% methanol-water was added to the obtained residues for reconstitution. The mixtures were vortex-mixed (2 min) and centrifuged (12 000 rpm, 10 min, 4 °C), and an aliquot of 10 μ L of the supernatant was injected per sample. For the correction of rats' individual differences, samples which were pooled by the supernatants of homologous samples in equal volume were analyzed by LC-MS for typical chromatograms.

Chromatographic and mass spectrometric conditions

Samples were analyzed by UHPLC-ESI-Q-TOF-MS/MS systems, which consisted of a Shimadzu Nexera UHPLC LC-30A system coupled to a hybrid quadrupole time-of-flight tandem mass spectrometer equipped with Turbo V sources and a Turbo ion spray interface (AB Triple-TOF 5600+, AB SCIEX, Foster City, CA) to obtain the accurate mass and structure information of xenobiotic prototype components and metabolites of ZZCD in vivo. The linear elution gradient of the mobile phase consisting of A (acetonitrile) and B (CH₃COOH : $H_2O = 0.1 : 100, v/v$) was as follows: 0-10 min, 0-50% A; 10-13 min, 50-95% A; 13-15 min, 95-95% A; 15-16 min, 95-0% A; and 16-18 min, 0-0% A, at a constant flow rate of 0.3 mL min⁻¹. A Waters HSS T3 column (150 \times 3 mm, 1.8 μ m) was used to separate the analytes at the column temperature of 35 °C. The liquid chromatography effluent was split and then introduced into the inlet of the MS. Each mass detector was equipped with an electrospray ionization (ESI) source and operated in both positive and negative ionization modes.

The conditions of UHPLC-ESI-Q-TOF-MS/MS were optimized and set as follows: capillary voltage: 5000 V (+)/4500 V (-); mass range of Q-MS and TOF MS/MS scanning (m/z): 100–1500 and 50-1000; scan mode: IDA criteria; capillary temperature: 500 °C; declustering potential (DP): 60 V; collision energy (CE): 35 V. The collision energy spread (CES) was set at 15 eV, enabling us to obtain average EPI scan spectra when the CE was 20, 35 and 50 eV. A real-time multiple mass defect filter (MMDF) and dynamic background subtraction (DBS) were used to fulfill the IDA criteria. Data were acquired by the Analyst® TF 1.6 software (AB SCIEX, Foster City, CA). For calibrating the mass detector, the calibrant delivery system (CDS), under which the correction solution was injected by an independent secondary probe, was used to automatically adjust the ion mass in real time during the procedure of sample analysis. The correction solution was a polypropylene glycol (PPG) solution (2.0×10^{-7} M for positive ion mode and 3.0 \times 10⁻⁵ M for negative ion mode). The m/zvalues 175.1330 and 906.6730 were chosen as PPG preventative maintenance values for positive ion mode, while m/z values 59.0130 and 933.64 were chosen for negative mode. The mass error and mass deviation were less than 2 ppm and ± 0.1 Da, respectively.

Strategies for screening and identification of ZZCD in rat feces

In order to figure out the metabolic profile of *ZZCD* in rat feces, an identification strategy combined with on-line data acquisition and off-line data analysis techniques was used in this study. It could be divided into three steps (Fig. 2).

In step one, the IDA criterion was chosen as the mass spectrometer on-line analysis strategy. Then, the MS spectra and MS/MS spectra of the samples were acquired in the condition of low and high collision energy accordingly in both positive and negative ionization modes. The structural information of chemical ingredients in the samples, including parent ions, fragment ions and neutral loss, were obtained globally.

In step two, post data mining for exogenous components in the samples was carried out through an untargeted

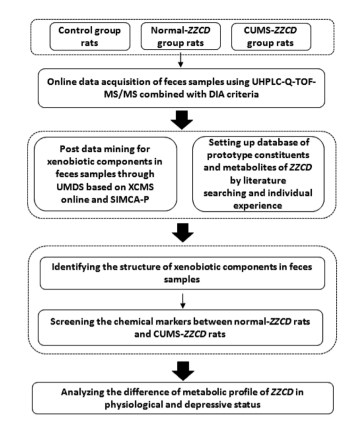


Fig. 2 Workflow of identification strategies for ZZCD in rats.

metabolomics-driven strategy. In brief, data extraction and data analysis were performed in this step. The XCMS online platform, a preprocessing tool of LC-MS chromatograms, was used for data extraction.^{35,40} It can automatically align the retention time and pick peaks of multiple compounds. As a result, threedimensional (3D) datasets including peak codes, m/z-retention time $(t_{\rm R})$ pairs and ion intensity for both blank group and drugcontaining group were acquired. For data analysis, the resulting 3D datasets were introduced to the SIMCA-P software (13.0.3 version, Umetrics, Umea, Sweden). Then, PCA and OPLS-DA were executed; the ions with VIP values greater than 1.0 in the score plots and S-plots were regarded as xenobiotics in the fecal samples. At the same time, a database of ZZCD was established in an Excel electronic form through literature search and previous research, in which the information including component names, accurate m/z of quasi-molecular ions, MS/MS fragmentation, formulae, molecular weight and relative polarity between the isomeride of prototype compounds and metabolites were listed clearly.

In step three, the xenobiotics identified above were matched based on the database hits through comparing their MS information with the database by sPIF and DIFS.^{34,35} Then, the chemical compositions were identified by sPIF through matching the database of *ZZCD*. When two or more fragment ions were matched, the chemical compounds in the database were chosen as the candidate components quickly. For the identification of the candidate components, DFIS was used to compare the characteristic ions related to the nucleus structure, possible metabolic and fragment pathways between candidates, and chemical compositions. With the view of further verification of the result, the chemical composition information was characterized by XIC in original MS spectra.

Chemical markers screening and analysis

In this step, the 3D datasets including peak codes, $m/z-t_{\rm R}$ pairs and ion intensity of exogenous compounds present in normal + ZZCD feces and CUMS + ZZCD feces were imported into the SIMCA-P software for multivariate data analysis. Then, the chemical compounds that most significantly contributed to the discrimination (VIP > 1.0 and p < 0.05) of the normal + ZZCD and CUMS + ZZCD groups were screened out as the chemical markers, which may be closely related with the anti-depression material basis of ZZCD.

Results and discussion

Depression-like behaviours and biochemical analysis

The results of behavioural tests in week 4 are listed in Fig. 3. Rats in the CUMS group showed low sucrose preference in SPT (Fig. 3(A)), low score in OFT (Fig. 3(B)), and more time in immobility in FST (Fig. 3(C)). This indicated that these rats exhibited significant depression-like behaviours. Further, the plasma levels of 5-HT and DA were measured as the indices of depression, which can be seen in Fig. 3(D) and (E). The plasma levels of 5-HT and DA of CUMS-induced depression rats were lower than those of normal rats. As a result, the depression model was attested to be successful by behaviour tests and biochemical analysis.

Optimization of sample collection and preparation methods

In consideration of the excretion kinetics of multiple constituents and their metabolites of ZZCD, the collection time was set as 0 h-24 h by preliminary experiments, during which majority of the components were totally excreted in the feces. Due to the different polarity and the different metabolic pathways of the chemical components, the metabolic profile of ZZCD in rat feces is complex. Therefore, the preparation method that can retain the most xenobiotics should be chosen. In this study, the solvents used for both the preparation assisted by ultrasonic treatment and reconstitution were compared for fecal samples, including 20% methanol-water, 50% methanol-water, 80% methanol-water and pure methanol. As a result, 50% methanol-water was chosen as the solvent in both preparation and reconstitution because the signal strength of majority of the compounds was the highest, which indicates that it has better extractability for the constituents having different partition coefficients.

Optimization of LC and MS conditions

Mobile phase compositions, which were closely linked to the peak shape and separation of the constituents in biological samples and *ZZCD* samples, were screened in this study. For aqueous phases, acetic acid and formic acid of different concentrations in water were compared, while methyl alcohol

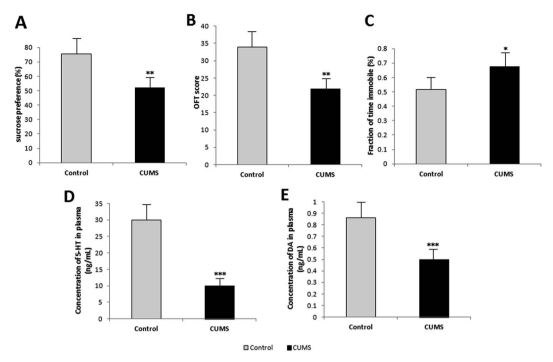


Fig. 3 Rats exhibited alterations in depression-like behaviors and biochemical criterion (n = 8). Rats in CUMS group showed lower sucrose preference in SPT (A), lower score in OFT (B), more time in immobility in FST (C), lower plasma levels of 5-HT (D) and DA (E) in week 4. These data are presented as the mean \pm s.e.m. *p < 0.05 **p < 0.01 ***p < 0.001, between two groups, as measured by independent samples *t*-test.

and acetonitrile were compared for the choice of the organic phase. As a result, acetonitrile and 0.1% acetic acid in water were selected as the mobile phases based on gradient elution.

The parameters of Q-TOF-MS/MS spectra, which were related to the resolution, greater baseline stability and higher ionization efficiency, including capillary voltage, mass range of m/z, scan mode, capillary temperature, declustering potential (DP), collision energy (CE), and collision energy spread (CES), were optimized. For fulfilling IDA criteria, real-time MMDF and DBS were used. The typical total ion chromatograms (TICs) of control rat feces, normal + *ZZCD* rat feces and CUMS + *ZZCD* rat feces in both positive and negative ion modes are listed in Fig. S1 and S2.† The TICs of standard compounds and *ZZCD* samples are listed in Fig. S3.†

Metabolic profile identification of ZZCD in rat feces

Screening the xenobiotics in rat feces. The chemometrics softwares for the integrated identification strategy in this study were used for mining the mass spectrometry information of exogenous components in biological samples. Due to the ability of the XCMS online platform for processing and visualizing mass-spectrometry-based and entire untargeted metabolomic data, an Excel file comprising the information of peak codes, *m*/*z*-retention time (t_R) pairs and ion intensity in both blank sample and drug-dose sample could be given as the result of the online analysis.^{31,41} Then, the SIMCA-P software was used for multivariate data analysis. According to the results of PCA analysis, CUMS + *ZZCD* samples, normal + *ZZCD* samples and control samples were divided into three detached groups in

score plots, which indicate that the chemical constituents of rat feces changed due to the dose of ZZCD, as shown in Fig. 4(A) and (B). In the OPLS-DA analysis, S-plots and VIP values could be helpful in visualizing and filtering the variables that influenced the model. The abscissa axis expresses covariance, indicating the visualization of the contribution, while the vertical axis expresses correlation, indicating reliability. The theoretical value ranges of the vertical axis are from -1 to +1.42As can be seen in Fig. 4(C) and (D), each triangle point in the Splot represents one of the variables of the 3D dataset, and the distance from the original point shows the confidence level of the variables contributing to the clustering observed in the PCA scatter. Farther distance means higher values, like purple ones (in red pane), which are the potential purpose points in this study. For further mining of data, the VIP variable was chosen for the recognition of constituents that only existed in the drugcontaining feces. The $m/z-t_{\rm R}$ pairs with VIP > 1.0 and p < 0.05could be regarded as chemical compounds from ZZCD. Here, m/ z-t_R pairs M411-T6.1 and M227-T6.1 in positive ion mode are chosen as examples, while the trend plots are listed in Fig. 4(E) and (F).

Identification of prototype constituents and metabolites of *ZZCD*. The fragment ion structure information of xenobiotics and metabolites of *ZZCD* mined by MUDS were listed in Excel files. According to the IDA criteria of UHPLC-Q-TOF-MS/MS, quasi-molecular ions, neutral loss and fragment ions were included in the dataset. For identifying exogenous components from *ZZCD* in fecal samples, sPIF and DIFS, which were based on comparison and filtration of m/z- t_R pair groups with the database of prototype constituents and metabolites of *ZZCD*

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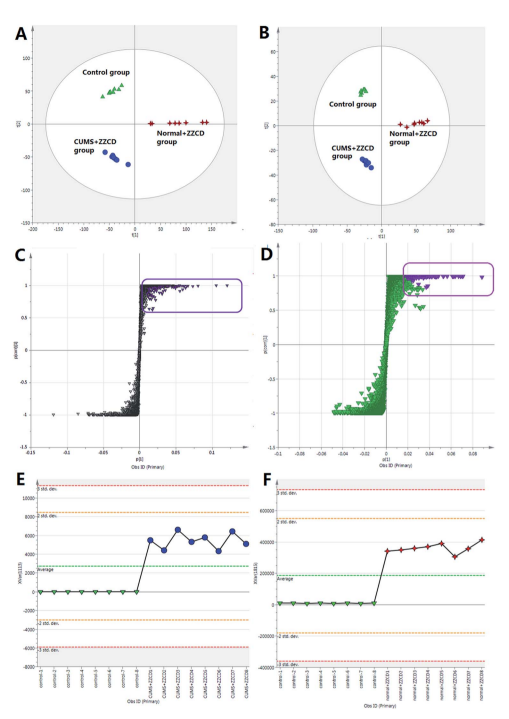
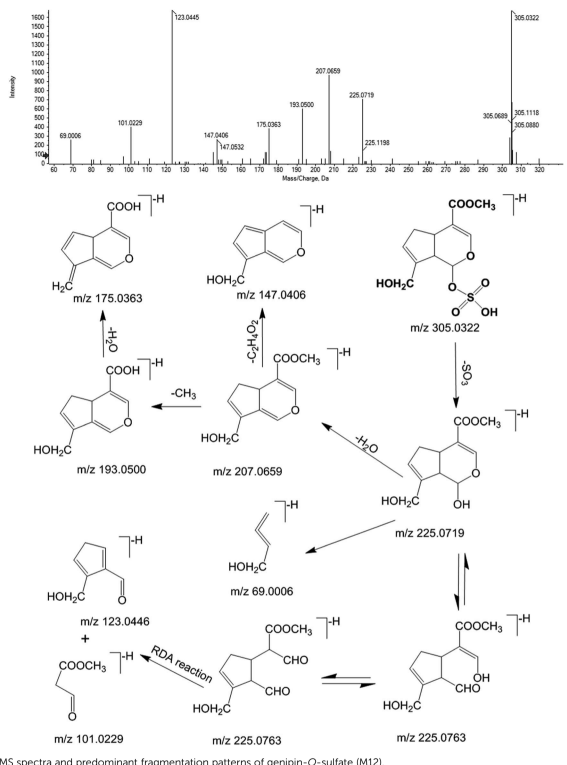


Fig. 4 The statistic studies by PCA and OPLS-DA. PCA score plot of all analyzed samples in positive-ion mode with the statistical parameters ($R^2X = 0.698$, $Q^2 = 0.572$) (A); OPLS-DA *S*-plot between CUMS + *ZZCD* group and control group in positive-ion mode (C) ($R^2X = 0.984$, $R^2Y = 0.953$, $Q^2 = 0.927$); PCA score plot of all analyzed samples in negative-ion mode with the statistical parameters ($R^2X = 0.694$, $Q^2 = 0.563$) (B); OPLS-DA *S*-plot between normal + *ZZCD* group and control group in negative-ion mode (D) ($R^2X = 0.905$, $R^2Y = 0.943$, $Q^2 = 0.912$); the trend plot of t_R -*m*/z 6.1–411 between CUMS + *ZZCD* group and control group in positive ion mode (E); the trend plot of t_R -*m*/z 6.1–227 between normal + *ZZCD* group and control group in positive ion mode (F).

(described in part "Strategies for screening and identification of *ZZCD* in rat feces", step two), were used. Finally, a total of 56 chemical components, comprising 35 prototype compounds and 21 metabolites, were identified or tentatively confirmed. All these components were verified back to the original mass spectrogram of drug-containing feces by XIC, while the

prototype compounds were further verified by comparing with the mass information in *ZZCD* samples. Four of these constituents, namely, chlorogenic acid (peak 4), geniposide (peak 16), *p*-coumaric acid (peak 19) and daidzein (peak 23), were further confirmed by reference substances by the consistency of $t_{\rm R}$ and MS/MS data. There were 9 iridoid glycosides, 8



MS/MS spectra and predominant fragmentation patterns of genipin-O-sulfate (M12). Fig. 5

monoterpenoids, 7 flavonoids, 6 soyasaponins, 4 quinic acid derivatives and 1 organic acid. All the information and the molecular structural formulae of prototype compounds are listed in Table S1[†] and shown in Fig. S4.[†] The information on 21 metabolites is listed in Table S2,† and the proposed metabolic profile of ZZCD is shown in Fig. 7.

Furthermore, M12 was chosen as an example for expounding the process of the detailed identification by an integrated identification strategy. The $m/z-t_{\rm R}$ pairs of fecal samples in the negative ion mode, including M305-T8.15, M225-T8.15, M207-T8.15, M193-T8.15, M175-T8.15, M123-T8.15, and M101-T8.15, were screened as xenobiotics from ZZCD based in their

performance in the OPLS-DA S-plot and VIP values (VIP > 1.0 and p < 0.05). Then, they were pooled as a fragment ion group due to the same retention time, which could be regarded as fragment ions coming from one chemical component. The strategy of sPIF was used to screen the possible candidate components by comparing the fragment ion group with the database of ZZCD, which was established through literature search and previous research.^{2,5,34,35,43-47} Genipin-related prototype compounds or metabolites were chosen as the candidate components of M12 due to two or more fragment ions congruously, including m/z 207, 193 and 175. In order to further confirm the structure of M12, an extracted ion chromatogram (XIC) was obtained to find out the retention time and structure information of M12, while DIFS was used to identify the possible metabolic and fragment pathway by characteristic ions. As can be seen from Fig. 5, m/z 305 and m/z 225 were chosen as diagnostic fragment ions, and M12 was defined as the sulfate conjugate due to the neutral loss fragment (NLF) of 80 Da $(-SO_3)$. The calculated molecular formula of M12, for which the quasi-molecular ions in negative ESI mode of Q-TOF-MS/MS was $305.0322 [M - H]^{-}$ (ppm = -4.6), was speculated to be C11H14O8S based on the analysis of elemental composition and fractional isotope abundance. The m/z 225, which could be found in Fig. 5, was the characteristic ion of genipin $[M - H]^{-}$. By losing H₂O, CH and CO continuously, genipin could give the series of product ions of m/z 207, 193 and 175, which accord with the information of the fragment ion group. Additionally, m/z 123 and 101 are from the Retro Diels-Alder (RDA) reaction of the iridoid skeleton. By comparing the information with the reports in related literature,2,13 M12 was tentatively identified as genipin-O-sulfate, and the molecular weight of all the fragment ions followed the nitrogen rule of mass spectrometry.35 The proposed metabolite pathway of iridoid glycosides-related metabolites in rat feces after oral administration of ZZCD is shown in Fig. 7(A).

M11 was chosen as another example for expounding the process of identification. M11 ($t_{\rm R} = 8.005 \text{ min}, m/z$ (+) 431.0963) was screened as the xenobiotic from *ZZCD* in feces based on its

performance in the OPLS-DA *S*-plot (VIP > 1.0 and p < 0.05). The calculated molecular formula was speculated to be C₂₁H₁₈O₁₀ based on the analysis of its elemental composition and fractional isotope abundance (ppm = -2.3). The fragment ions, including that of m/z 431, were chosen as the product ions of m/zz 431.0963 based on the same $t_{\rm R}$, which were verified by XIC in the original MS spectra. The NLF of 176 Da (C₆H₈O₆) indicated that M11 was a glucuronide conjugate. Then, m/z 255 and m/z199 were chosen as the diagnostic ions of M11 due to the higher relative abundance. As a result, daidzein-O-glucuronide was found to conform to the fragment pathway.48 The fragment ions with m/z 255, m/z 227, m/z 199 and m/z 137 were identified as $[M-GluA + H]^+$, $[M-GluA-CO + H]^+$, $[M + H-2CO]^+$ and [M + H- C_8H_6O ⁺, respectively. The XIC of M11 in the positive ion mode and proposed fragmentation pathway are illustrated in Fig. 6. The proposed metabolic profiles of flavonoid-related metabolites are shown in Fig. 7(C).

As a result, 36 prototype constituents and 21 metabolites of *ZZCD* were identified or tentatively confirmed by the integrated data-mining strategy, and two xenobiotic components were unidentified. M10 ($m/z-t_R$ 367.1150–7.727) was speculated as a sulfate conjugate due to the loss of a neutral fragment of 80 Da (-SO₃), while the possible structure was still unclear. In the same way, M16 ($m/z-t_R$ 465.1896–9.104) was speculated as a sulphate and glucuronide conjugate. The MS/MS spectra of M10 and M16 are illustrated in Fig. S5.†

Analysis of metabolic profile of ZZCD in rat feces. In this study, a total of 56 ZZCD-related exogenous components, namely, 35 prototype components and 21 metabolites, were identified or characterised tentatively after oral administration. As can be seen from the results, iridoid glycosides, flavanones, monoterpenoids, organic acids, and soyasaponins were the main prototype components existing in feces. Glucuronidation and sulfoconjugation were the main phase II metabolic pathways of ZZCD, while ring-opening cracking, hydroxylation, Nheterocyclization, hydrolysis reaction, loss of CH_2 and hydrogenation metabolic pathway were also detected in rat feces (list in Table 1).

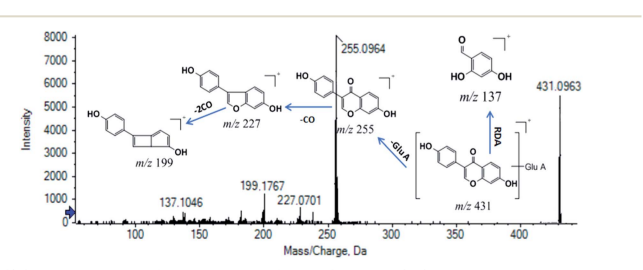


Fig. 6 MS/MS spectra and predominant fragmentation patterns of daidzein-O-glucuronide (M11).

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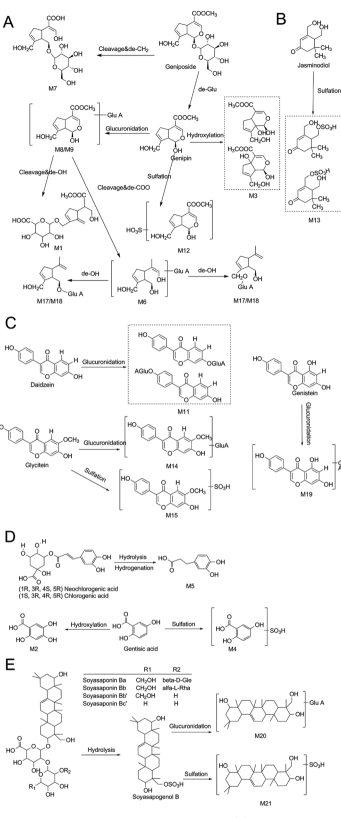


Fig. 7 The proposed metabolic profiles of iridoid glycosides-related metabolites (A), monoterpenoids-related metabolites (B), flavonoidsrelated metabolites (C), organic acids-related metabolites (D) and soyasaponins-related metabolites (E) in rat feces after ZZCD oral administration.

Iridoid glycosides, soyasaponins and isoflavones were the main constituents of ZZ and DDC, respectively. In this study,

nine iridoid glycosides-related metabolites, such as M3, M6, and M7, were identified, while the main exitence form of

	Туре	Amount of detected constituents
Prototype compounds	In total	35
	Iridoid glycosides	9
	Monoterpenoids	8
	Flavonoids	7
	Soyasaponins	6
	Quinic acid derivatives	4
	Organic acids	1
Metabolites	In total	21
	Hydroxylation	2
	Dihydroxylation	1
	Ring-opened and de-hydroxy of mono-glucuronidation	2
	Loss of CH ₂ and cleavage	1
	Ring-opened and methyl formate removal derivative of mono- glucuronidation	1
	Mono-sulfate conjugation	5
	Mono-glucuronide conjugation	4
	Sulfate and glucuronide conjugation	1
	Hydrolysis and glucuronidation	1
	Hydrolysis and sulfation	1
In total		56

soyasaponins and isoflavones in rat feces was prototype. Since studies have shown that geniposide or genipin, which is the main metabolite of geniposide, have hepatotoxicity *in vivo*,⁴⁹ the results of this study may suggest that *ZZCD* reduced the toxicity and enhanced the efficacy after oral administration. A previous research claimed that *DDC* has different degrees of regulatory effects on gut microbes,⁵⁰ and changes in intestinal flora can affect the development of depression.^{21,22} It can be speculated

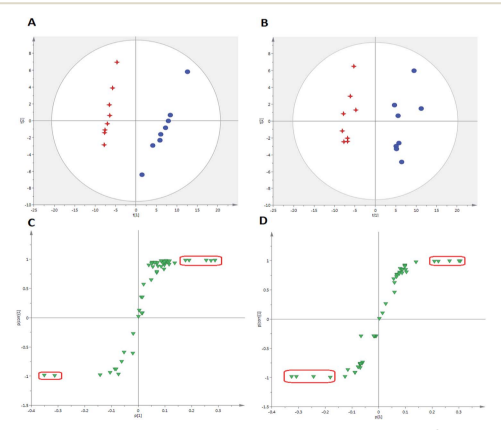


Fig. 8 PCA score plot with the statistical parameters of 56 exogenous compounds in positive-ion mode ($R^2X = 0.796$, $Q^2 = 0.815$) (A) and negative-ion ($R^2X = 0.76$, $Q^2 = 0.712$) (B); S-plot of OPLS-DA with the statistical parameters of 56 exogenous compounds in positive-ion mode ($R^2X = 0.823$, $R^2Y = 0.967$, $Q^2 = 0.95$) (C) and negative-ion ($R^2X = 0.78$, $R^2Y = 0.923$, $Q^2 = 0.95$) (D).

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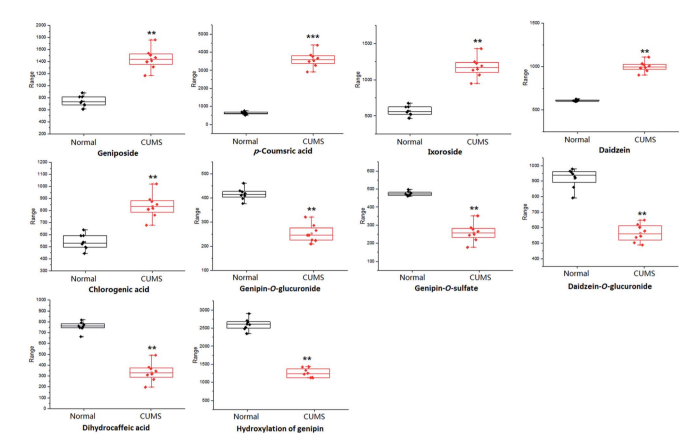


Fig. 9 Changes in the intensities of 10 potential chemical markers in the plasma samples of the normal + ZZCD group and CUMS + ZZCD group. **p < 0.01, ***p < 0.001 compared with the normal + ZZCD group (n = 8 for each group).

that gut microbes may be one of the antidepressant targets of *ZZCD*.

Chemical markers screening and identification

For the purpose of studying the chemical composition difference between the normal and CUMS-induced depression rats, the 3D datasets (peak code, $t_{\rm R}$ -m/z pairs, and ion intensity) of the 56 exogenous compounds were imported into the SIMCA-P software for multivariate data analysis, the results of which can better elucidate the material basis of ZZCD in antidepression. It can be seen in Fig. 8(A) and (B) that the two groups are clearly divided into two areas, which indicates that there are differences in the distributions of these 56 xenobiotics in the feces of normal and CUMS-induced rats. For the further study of screening the chemical markers, which may be the potential bioactive components of ZZCD in vivo, the S-plots were acquired in both positive-ion mode (Fig. 8(C)) and negative-ion mode (Fig. 8(D)). In addition, 10 chemical components with VIP > 1.0 were screened out as the potential chemical markers. Among these, the content of geniposide, p-coumaric acid, ixoroside, daidzein, chlorogenic acid in feces of CUMS-induced rats was higher than that in normal rats, while the content of genipin-Oglucuronide, genipin-O-sulfate, daidzein-O-glucuronide, dihydrocaffeic acid and hydroxylation of genipin were lower in the depression model rats (Fig. 9).

Conclusion

The metabolic profile of ZZCD in rat feces could provide the basis for studying the effective components of antidepressants due to the correlation between intestinal flora and depression. For identifying the cause of signality and complicacy, an effective and systematic identification strategy is needed urgently. In this study, multiple strategies based on UHPLC-ESI-Q-TOF-MS/ MS and data mining softwares were used to screen and identify the prototype components and metabolites of ZZCD in rat feces after oral administration. According to the results, 36 prototype components and 21 metabolites were identified or temporarily confirmed. It can be seen that the strategies mentioned in this study were easy to automate, for which the time for screening and identifying is distinctly shorter. The results indicated that mono-sulfate conjugation and mono-glucuronide conjugation were the main metabolic pathways of ZZCD in rat feces. For iridoid glycosides, hydrolysis, ring-opening, hydroxylation and loss of CH₂ metabolic pathways were observed. For organic acids, the hydrolysis metabolic pathway was identified. Even for soyasaponins, the hydrolysis metabolic pathway was identified. The study of the metabolic profile of ZZCD in rat plasma revealed that glucuronidation was the main metabolic pathway,⁵ compared with which the metabolic pathways of ZZCD in feces were more complex. Furthermore, there were differences in the metabolic pathways of geniposide and glycitein among rat bile, urine and feces.² For bile and urine,

homocysteine conjugation, methylation, *N*-acetylcysteine, ketone formation and N-heterocyclization conjugation pathways were identified. For feces, the loss of OH and CO₂ pathways was determined. The formation of the specific metabolites of *ZZCD* in rat feces may be due to the gut microbiota.³⁵

There were no differences in fecal metabolite types between the normal and CUMS-induced depression rats. However, the concentration of some constituents was significantly different. Further, a total of 10 chemical markers, which significantly contributed to the discrimination of the metabolic profile of ZZCD in rat feces, were screened as the potential bioactive components of ZZCD by statistical analysis between the normal + ZZCD group and CUMS + ZZCD group. The discrepancy in these metabolites on concentration may relate to the distinct perturbation of the composition of gut microbiota in induced depression rats, such as Bacteroides, Clostridium, Coprococcus, Pseudobutyrivibrio and Dorea genus, which could change the alteration of gastrointestinal motility.21,51 Furthermore, previous studies have claimed that some chemical components of ZZCD, such as daidzein, genistein, chlorogenic acid, p-coumaric acid, geniposide, caffeic acid, daidzin, and dihydrodaidzein, could modulate the gut microbiota composition,^{7,10,24,50,52} while the change in the composition of gut microbiota can influence the functioning of the hypothalamic-pituitary-adrenal (HPA) and immune system and thus, it may play a crucial role in depression through the brain-gut axis.^{21,22,51} The ZZCD-related chemical components, whose concentration changed significantly in feces of normal and depression rats, including excreted components and the residual components from the initial ingested decoction, may be responsible for the antidepressant effects of ZZCD in vivo.

In conclusion, this study could be helpful for the research of anti-depression material basis and mechanism of *ZZCD*, and the strategy proposed in this study would be a practical and systematic solution for screening and identifying untargeted compounds (including prototype constituents and metabolites) and bioactive components of TCMP *in vivo*.

Ethical statement

This study was performed in strict accordance with the NIH guidelines for the care and use of laboratory animals (NIH Publication No. 85–23 Rev. 1985) and was approved by the Institutional Animal Care and Use Committee of Bengbu Medical University (Bengbu, China).

Conflicts of interest

There are no conflicts to declare.

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