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1. Introduction

Traditional Chinese medicines (TCMs) play an important role in Chinese medicine due to their wide application and efficacy, particularly in the management of chronic diseases.¹ TCMs are composed of complex chemical constituents for multi-target treatment, and produce synergistic effects and reduce side effects. Although there has been a lot of research on TCMs, the active ingredients of most traditional Chinese medicine prescriptions remain unknown. Therefore, developing a fast and efficient analysis method to separate and identify complex constituents in TCMs is necessary for illustrating the pharmacological material basis of TCMs.

Sanhua decoction (SHD), a famous Chinese herbal formula, is recorded in the classic traditional Chinese medicine (TCM) book Suwen Bingji Qiyi Baomingji (Plain Questions: Discourse on Mechanism for Preserving Life). It consists of four crude herbs: Zhishi (*Aurantii fructus immaturus*), Dahuang (*Rheum palmatum* L.), Houpu (*Magnoliae Officmalis Cortex*) and Qianghuo (*Notopterygii Rhizoma Et Radix*). SHD is used widely for the

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Rapid characterization of the chemical constituents of Sanhua decoction by UHPLC coupled with Fourier transform ion cyclotron resonance mass spectrometry[†]

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Sanhua decoction, a famous Chinese herbal formula has been widely used for the treatment of stroke. In our study, a rapid, swift and straightforward analytical method with the help of UHPLC-FT-ICR-MS/MS was successfully developed for the first time to separate and identify the chemical constituents of Sanhua decoction. Chromatography was performed on a Universal XB C_{18} column (150 mm \times 2.1 mm, 1.8 μ m) using a mobile phase containing 0.1% formic acid–water (A) and acetonitrile (B). A total of 137 compounds in Sanhua decoction were identified or tentatively characterized. The findings revealed the fact that Sanhua decoction mainly contains flavonoids (in *Aurantii fructus immaturus* and *Rheum palmatum* L.), anthraquinones (in *Rheum palmatum* L.), coumarins (in *Notopterygii Rhizoma Et Radix*), phenylpropanoid glycosides, alkaloids and lignans (in *Magnoliae Officmalis Cortex*), which made up the key ingredients existing in Sanhua decoction. This study is hoped to be meaningful for the characterization of components in other traditional Chinese medicines, and lay the foundation for research on the pharmacology of Sanhua decoction.

treatment of stroke. It can improve ischemic cerebral edema and brain blood barrier (BBB) permeability.² It has been documented that Zhishi has neuroprotective, antioxidant, anti inflammatory and anti-apoptotic effects.³ Dahuang can protect neurons from hypoxic-ischemic brain damage.⁴ Houpu can protect neural damage from cerebral ischemia and reperfusion by suppressing cerebral inflammation and improving BBB function.⁵ Qianghuo has a function of anti-thrombosis, increasing cerebral blood flow and improving cerebral blood circulation.⁶ However, the chemical material basis of SHD has not been studied in detail until now.

As the complexity of traditional Chinese medicine, it is very important to establish a sensitive and reliable detection method. Chromatography-mass spectrometry (LC-MS) is widely used for its high sensitivity and specificity. Ultra high performance liquid chromatography (UHPLC) coupled with Fourier transform ion cyclotron resonance mass spectrometry (FT-ICR-MS) as the most powerful instrument with high resolution, accuracy and sensitivity play a role in analyzing different kinds of complex samples.⁷⁻¹⁰ It provides an unambiguous elemental composition and information about the isotopic abundance of ions to help determine the compositions and structures of chemical constituents. By usage of this technology, Li *et al.* characterized 179 constituents in *Rhodiola crenulata* as well as 37 prototypes and 142 metabolites in rats.¹¹ Guan *et al.* characterized 120 constituents in Sijunzi decoction¹² and Liu *et al.*

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characterized 174 constituents in Gegenqinlian decoction as well as 107 prototypes and 67 metabolites in rats.¹³ The results show that this method is useful in detecting and indenting chemical constituents and metabolites by information about accurate molecular weight and MS².

In this study, a rapid and simple approach by UHPLC-FT-ICR-MS was established for the systematical characterization of the constituents of SHD. This constituent characterization and structural elucidation provided significant information for a pharmacological study of SHD.

2. Material and methods

2.1 Chemicals and materials

The purchase of key ingredients including, Aurantii fructus immaturus (batch number: 170 901; source: Sichuan China), Rheum palmatum L. (batch number: 20 181 101; source: Gansu China), Magnoliae Officmalis Cortex (batch number: 181 001; source: Sichuan China) and Notopterygii Rhizoma Et Radix (batch number: 13 112 501; source: Sichuan China) was made from Guoda pharmacy (Shenyang, China), which followed identification by Professor Jingming Jia (Department of TCM, Shenyang Pharmaceutical University, Shenyang, China). The key source of the reference compounds (purity > 98%), including sennoside B, nodakenin, narirutin, sennoside A, neohesperidin, naringenin, hesperetin, nobiletin was Shanghai Yuanye Bio-Technology Co., Ltd. (Shanghai, China); naringin, hesperidin and honokiol was attained from the National Institute for the Control of Pharmaceutical and Biological Products (Beijing, China). In addition, acetonitrile of HPLC grade, together with formic acid of LC-MS grade was attained from Fisher Scientific (Fair Lawn, NJ, USA), followed by attaining the purified water from Wahaha (Hangzhou, China).

2.2 Preparation of SHD for analysis

Following the documentary records of SHD, four constituting herbs that included *Aurantii fructus immaturus* (40 g), *Rheum palmatum* L. (40 g), *Magnoliae Officmalis Cortex* (40 g) and *Notopterygii Rhizoma Et Radix* (40 g) were soaked in water for 60 min, then decocted two times in 80 mL water for a period of 1 h each time in a glass flask and the ensuing solution was mixed and dried with the help of lyophilization. Prior to the analysis, dissolution the 0.5 g of dried powder was dissoluted by 5 mL methanol-aqueous (1 : 1) solution, soaking for 15 min with 80 °C water.

2.3 Instrument and analytical conditions

The sample was analyzed by the UHPLC-DAD-FT-ICR-MS system contains an Agilent 1260 UHPLC system combined with a Bruker Solarix7.0T FT-ICR-MS system (Germany) and Bruker Compass-Hystar workstation (Bruker, Germany). Chromatography was carried out on a Universal XB C_{18} column (150 mm \times 2.1 mm, 1.8 µm; Kromat, USA) at the temperature of 35 °C. The mobile phase contained 0.1% formic acid–water (A) and acetonitrile (B). The gradient conditions were as follows: 8–12% (B) in 0 to 10 min, 12–13% (B) in 10 to 12 min, 13–18% (B) in 13

The mass spectra were performed with positive as well as negative electrospray ionization (ESI) modes, followed by setting the optimized conditions are as follows: capillary voltage, 4.5 kV; plate offset, 500 v; nebulizer gas pressure, 4.0 bar; dry gas flow rate, 8 L min⁻¹; dry gas temperature, 200 °C; ion accumulation time, 0.15 s, and flight time, 0.6 ms. Full scan mass spectrometry data were recorded between m/z 100 and 1200 amu and the collision energy was ranged from 10 eV to 30 eV for MS/MS experiments.

3. Results and discussion

The Fig. 1 reveals the base peak ion chromatograms (BPC) of SHD in positive as well as negative ion modes, together with the respective compounds. The extract ion chromatograms (EIC) of each molecule weight were correspondingly attained for detecting the associated compound followed by presenting in support information Fig. S1.† Comparing with reference compounds, eleven peaks from SHD were accurately identified by the retention times and MS fragments in positive ion mode. The peak of 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 in Fig. 1C were sennoside B, nodakenin, narirutin, naringin, sennoside A, hesperidin, neohesperidin, naringenin, hesperetin, honokiol, nobiletin, respectively. The others were predicted by Bruker workstation, whether or not the mass error values between the identified molecular weights with the measured molecular weights were below 3.0 ppm. If yes, it can infer the chemical structure of each component by the retention time and MS/MS data with published literature.14-28 In the end, a total of 137 compounds were identified and characterized, whereas their layouts were presented in support information Fig. S2.†

The MS/MS spectra of representative compounds are shown in Fig. 2, and displaying deduced fragmentation pathways are shown in Fig. 3. Except that, the formula, retention time, MS data, calculated m/z, error, ion mode and MS/MS data of other detected components were listed in Table 1.

3.1 Characterization of flavonoids in SHD

Flavonoids have a typical cyclohexene structure in the C ring, which is unstable and easy to break it for Diels–Alder reaction (RDA), and get the characteristic fragment ions 1,3A and 1,3B. Flavonoid glycosides have methoxy group, firstly losing the glycosyl groups, followed by losing the methyl groups, at last cleavage of the C ring. The mass spectral fragmentation of flavonoids by hesperidin as an example was described. In positive ion mode, adduct ion at m/z 611.19650 ([M + H]⁺) indicating an elemental composition of C₂₈H₃₄O₁₅. The MS/MS spectrum (Fig. 3A) is presented that the main fragment ions are m/z 449.14312, 303.08581, 288.09732, 153.02175 as well as 136.04821. Among them, m/z 449.14312 and 303.08581 are inferred by cleaving of the glycosidic linkage and losing C₆H₁₀O₅ (162 Da) or C₁₂H₂₀O₉ (308 Da) from the ion of [M - H]⁺,

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Fig. 1 The base peak ion chromatograms (BPC) of SHT in both positive (A) and negative (B) ion modes and the corresponding compounds (C). (1) Sennoside B, (2) nodakenin, (3) narirutin, (4) naringin, (5) sennoside A, (6) hesperidin, (7) neohesperidin, (8) naringenin, (9) hesperetin, (10) honokiol, (11) nobiletin.



Fig. 2 The MS/MS spectra of the typical compounds. (A) Hesperidin, (B) nomilin, (C) emodin-O-malonyl-glucose, (D) galloyl-glucoside, (E) epicatechin, (F) honokiol, (G) magnolol, (H) acteoside, (I) magnocurarine, (J) bergaptol.



Fig. 3 The possible fragmentation pathways of the typical compounds. (A) Hesperidin, (B) nomilin, (C) emodin-O-malonyl-glucose, (D) galloyl-glucoside, (E) epicatechin, (F) honokiol, (G) magnolol, (H) acteoside, (I) magnocurarine, (J) bergaptol.

m/z 288.09732 is the result of losing CH₃ (15 Da) from m/z 303.08581, and m/z 153.02175 and m/z 136.04821 was due to the Diels–Alder reaction from m/z 288.09732. The characterization of other flavonoids in SHD was performed based on the fragmentation patterns and related literatures.¹⁴⁻¹⁸

3.2 Characterization of triterpenoids in SHD

Triterpenoids are prone to lose H₂O (18 Da), CO (28 Da) and CO₂ (44 Da), due to the presence of lactone rings. Nomilin was selected to illustrate the fragmentation pathways. In positive ion mode, adduct ion at m/z 515.22753 ($[M + H]^+$) indicating an elemental composition of C₂₈H₃₄O₉. The MS/MS spectrum (Fig. 3B) is presented that the main fragment ions are m/z 469.21806, 455.20675, 411.21462, 161.05854. Among them, m/z 469.21806 is inferred by losing CO and H₂O (46 Da) from [M + H]⁺, m/z 455.20675 is presumed by losing C₂H₄O₂ (60 Da) from [M + H]⁺, m/z 455.20675, and m/z 161.05854 is estimated by losing C₁₈H₂₆O₇ (354 Da) from [M + H]⁺. The characterization of other

triterpenoids in SHD was performed based on the fragmentation patterns and related literatures.^{15,19}

3.3 Characterization of anthraquinones in SHD

The common characteristic of MS^2 in anthraquinones is the continuous loss of CO (28 Da). Anthraquinone glycosides can lose glycosyl and CO, continually. Aglycones can be judged by [aglycone – H]⁻. Anthraquinone glycosides can lose methyl-glucuronide (190 Da), cinnamoyl (148 Da), malonyl (86 Da), acetyl-glucose (204 Da), and glucose (162 Da). Emodin-*O*-malonylglucose was selected to illustrate the fragmentation pathways. In negative ion mode, adduct ion at m/z 517.09935 ([M – H]⁻) indicating an elemental composition of C₂₄H₂₂O₁₃. The mass spectrum (Fig. 3C) shows that the main fragment ions are m/z 473.10884, 431.09157, 269.04531 and 241.05131. Among them, m/z 473.10884 is presumed by losing of CO₂ (44 Da) from [M – H]⁻, m/z 431.09157 is inferred by losing of the C₃H₂O₃ (86 Da) from [M – H]⁻, and m/z 269.04531 is speculated by

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Table 1 UHPLC-FT-ICR-MS analysis of SHD^a

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0 1 7 8 4 10 9 7 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	$t_{ m R}$ (min) 2.07 2.64 2.91 2.91 3.21 4.85 5.59 6.04 6.04	Identification Quinic acid Galloyl-glucoside 1-Galloyl-di-glucoside	Formula	mass(m/z)	(m/z)	Ion mode	mqq	MS/MS (m/z)	source
10 6 4100786	2.07 2.64 3.21 5.59 5.59	Quinic acid Galloyl-glucoside 1-Galloyl-di-glucoside							
0 m 4 m 0 n 80	2.64 2.91 3.21 5.59 6.04	Ğalloyl-glucoside 1-Galloyl-di-glucoside	$C_7H_{1,2}O_6$	193.07066	193.07024	$[M + H]^+$	2.21	147.04051, 129.05446	U
с 4 L Q L 8 6	2.91 3.21 5.59 6.04	1-Galloyl-di-glucoside	$C_{13}H_{16}O_{10}$	331.06707	331.06759	$[M - H]^{-}$	-1.57	313.05289, 271.04772, 211.02436,	R
4 ら 0 7 8 6	3.21 4.85 5.59 6.04		$C_{19}H_{26}O_{15}$	493.11989	493.12035	$[M - H]^{-}$	-0.93	103-01230, 123-00323 373.07705, 331.06771, 313.05667,	R
4 ら 6 7 8 6	3.21 4.85 5.59 6.04	:						271.04642	
5 9 N 9 0	4.85 5.59 6.04 7.00	Gallic acid	$C_7H_6O_5$	169.01425	169.01447		-1.30	151.00284, 125.02324	R
0 1 8 6	5.59 6.04 7.00	Catechin-3- <i>O</i> -glucoside	$C_{21}H_{24}O_{11}$	451.12459	451.12518		-1.31	289.07014, 245.08287	Я
N 8 6	6.04 7.00	Rebouoside C	$C_{20}H_{30}O_{12}$	461.16645	461.16674		-0.64	299.11268, 137.05927	М
86	100	Chlorogenic acid	$C_{16}H_{18}O_9$	353.08781	353.08824	$[M - H]^{-}$	-1.23	191.09726, 179.76826, 173.10064	Z
6	/.02	Catechin-5-0-glucoside	$C_{21}H_{24}O_{11}$	451.12459	451.12504	$[M - H]^{-}$	-1.01	289.07156, 245.08176, 203.07109	R
	7.79	Magnocurarine	$\mathrm{C_{19}H_{24}NO_3}^+$	314.17507	314.17473	$[M + H]^+$	1.07	269.11618, 237.08974, 175.07514,	Μ
								107.04887	
10	10.40	Catechin	$\mathrm{C}_{15}\mathrm{H}_{14}\mathrm{O}_{6}$	289.07176	289.07211	$[M - H]^{-}$	-1.22	245.08146, 205.04962, 203.06974	R
11	10.63	Cryptochlorogenic acid	$C_{16}H_{18}O_9$	353.08781	353.08825	$[M - H]^{-}$	-1.25	191.10029, 179.14054, 173.10065	z
12	10.64	Syringin	$\mathrm{C_{17}H_{24}O_9}$	395.13125	395.13094	$[M + Na]^+$	0.80	372.08647, 343.94804, 208.09042,	Μ
								137.03937	
13	13.55	Magnoflorine	$\mathrm{C}_{20}\mathrm{H}_{24}\mathrm{NO}_4^{ op}$	342.16998	342.16967	_[H + M]	0.91	297.10956, 282.08696, 265.08507,	M
			+			+		23/.089/2	
14	16.29	(R)-Oblongine	$C_{19}H_{24}NO_3$	314.17507	314.17475	[H + M]	1.02	269.11625, 237.09024, 175.07485,	Z
ļ			;					10/.04801	ŗ
15	16.62	Epicatechin	$C_{15}H_{14}O_6$	289.0/1/6	289.0/213		-1.26	245.08235, 203.0/153	× ;
16	18.49	Teupolioside	$C_{35}H_{46}O_{20}$	785.25097	785.24931	[H - M]	2.11	649.28262, 623.38565, 477.34832	Μ
17	19.22	Aloe-emodin- <i>O</i> -di-glucoside	$C_{27}H_{30}O_{15}$	593.15119	593.15142	$[M - H]^{-}$	-0.38	431.09851, 269.04559, 239.03448	R
18	19.31	Apigenin-6,8-di- <i>C</i> -glucoside	$C_{27}H_{30}O_{15}$	595.16575	595.16502	$[M + H]^{+}$	1.23	577.15636, 505.13492, 475.12405	C
19	20.32	Xanthoplanine	$\mathrm{C}_{21}\mathrm{H}_{25}\mathrm{O}_4\mathrm{N}^+$	356.18563	356.18528	$[M + H]^+$	1.01	311.12628, 296.10825, 280.10864	Μ
20	20.74	Asimilobine	$C_{17}H_{17}O_2N$	268.13321	268.13294	$[M + H]^+$	0.99	251.10647, 236.08474, 219.08076,	М
								191.08617	
21	21.26	Echinacoside	$C_{35}H_{46}O_{20}$	785.25097	785.24959	$[M - H]^{-}$	1.76	649.28357, 623.38647, 477.34187	Μ
22	21.76	Chysoeriol-6,8-di- <i>C</i> -glucoside	$C_{28}H_{32}O_{16}$	625.17631	625.17579	$[M + H]^{+}$	0.84	607.16748, 535.14662, 505.13284	C
23	21.95	1-Cinnamoyl-6- <i>O</i> -(glucosyl)-	$C_{28}H_{32}O_{16}$	623.16176	623.16247	$[M - H]^{-}$	-1.15	461.10814, 331.06702, 313.05827,	R
		galloyl-glucoside				ţ		211.02305	
24	22.31	Scopoletin	$\mathrm{C_{10}H_8O_4}$	193.04954	193.04935	$[\mathbf{M} + \mathbf{H}]$	0.95	178.02482, 165.05373, 122.03784	U
25	22.45	2-O-Cinnamyl-β-ɒ-glucoside	$C_{15}H_{18}O_7$	309.09798	309.09841		-1.39	237.07692, 201.01967, 146.96590	К
26	22.55	Emodin-O-glucuronic acid methyl	$C_{22}H_{20}O_{11}$	459.09329	459.09358	$[M - H]^{-}$	-0.65	269.04543, 241.05106, 225.05490,	R
		ester						213.05525	
27	23.60	Decuroside V	$C_{20}H_{20}O_{10}$	447.12617	447.12576	$[M + Na]^+$	0.91	263.08937, 245.08472	z
28	23.73	Xanthotoxol	$C_{11}H_6O_4$	203.03389	203.03379	$[M + H]^+$	0.45	173.07564, 145.04536, 117.05273	U
29	23.73	Bergaptol	$C_{11}H_6O_4$	201.01933	201.01957	$[M - H]^{-}$	-1.17	185.02456, 157.17256, 129.10451,	N
								101.96320	
30	23.76	1-O-Cinnamyl-β-D-glucoside	$\mathrm{C}_{15}\mathrm{H}_{18}\mathrm{O}_{7}$	309.09798	309.09843	$[M - H]^{-}$	-1.47	248.95978, 161.04574, 146.96419	R
31	23.77	Bergaptol-O-β-ኴ-glucopyranoside	$\mathrm{C_{17}H_{16}O_9}$	365.08671	365.08641	$[M + H]^+$	0.81	201.03287, 159.17264, 131.10514	Z
32	24.94	Natsudaidain-3-0-glucoside	$C_{27}H_{32}O_{14}$	581.18648	581.18599	$[M + H]^+$	0.85	419.13174, 273.07487, 153.01846	C
33	24.97	Narirutin-4′-0-glucoside	$C_{33}H_{42}O_{19}$	743.23931	743.23875	$[\mathbf{M} + \mathbf{H}]^{+}$	0.75	581.18682, 435.12794, 273.07526	C
34	25.16	Eriocitrin	$C_{27}H_{32}O_{15}$	597.18140	597.18096	$[M + H]^{+}$	0.73	435.12791, 289.06947, 153.01849	U
35	25.35	Resveratrol-4′-0-glucoside	$C_{20}H_{22}O_8$	389.12419	389.12479	$[M - H]^{-}$	-0.83		Я

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TanteT									
No.	$t_{ m R}$ (min)	Identification	Formula	Monoisotopic mass (<i>m</i> / <i>z</i>)	Detected mass (m/z)	Ion mode	mqq	(z/m) SM/SM	Main source
								245.08196, 227.07138, 199.07567, 126.04204	
36	25.47	Isolindleyin	$C_{23}H_{26}O_{11}$	477.14024	477.14067	$[M - M]^{-}$	-0.90	125.04304 313.05691, 211.02328, 169.01307, 125.07387	R
37	25.79	Lindleyin	$C_{23}H_{26}O_{11}$	477.14024	477.14071	$[M - H]^{-}$	-1.12	313.05659, 211.02454, 169.01340,	R
38	26.19	Limonin	$C_{26}H_{30}O_{8}$	471.20134	471.20060	$[M + H]^+$	0.83	425.19343, 339.19426, 161.05897	U
39	26.19	Rhein-8- <i>O</i> -β- _D -glucoside	$C_{21}H_{18}O_{11}$	445.07763	445.07816	$[M - H]^{-}$	-1.21	285.07559, 267.03326, 257.07879,	R
40	26.28	Nicotiflorin	CHO	505 16575	595 16515	[M + H] ⁺	1 00	Z41.07041 419 13403 390 09214 287 07785	Ċ
41	26.28	Naringenin-7-0-glucoside	C27H30O15 C21H22O10	435.12857	435.12833	$[\mathbf{H} + \mathbf{M}]$	0.56	715.13403, 220.02414, 20.0010 272.07462, 227.18968, 177.22102, 165.13608	00
42	26.29	Eriodictyol	$\mathrm{C_{15}H_{12}O_6}$	289.07066	289.07051	$[M + H]^+$	0.53	153.01692, 135.06538, 107.04796	U
43	26.42	Limonin-17-β-ɒ-glucoside	$C_{32}H_{42}O_{14}$	649.25018	649.25108	$[H - H]^{-}$	-1.39	487.19725, 425.18954, 339.18671	C
44	26.49	Neoeriocitrin	$C_{27}H_{32}O_{15}$	597.18140	597.18134	$[M + H]^{+}$	0.52	435.12786, 289.06814, 153.01794	C
45	26.55	Isosakuranetin	$C_{16}H_{14}O_5$	287.09140	287.09138	$[M + H]^+$	0.38	153.01718, 133.065095	C
46	27.07	Alhagidin	$C_{34}H_{44}O_{20}$	771.23532	771.23656	$[M - H]^{-}$	-1.61	635.18786, 609.19823, 161.02394	М
47	27.17	Cinnamoyl-6-0-galloyl-	$C_{28}H_{32}O_{16}$	623.16176	623.16185	$[M - H]^{-}$	-1.50	475.10891, 331.06765, 313.05664	R
10	10 L C	uigucoside Modolonin	Сно	32767 207	701 10E0E	[M H]-	1 10	047 0467E 000 14E26 17E 41E7E	Ν
40 49	27.29	Marmesin	$C_{20}H_{20}O_{20}$	247.09649	407.09655 247.09655	$[\mathbf{M} + \mathbf{H}]^+$	-0.28	241.04029, 223.14930, 173.41373 153.01714	
50	27.38	Veronicastroside	$C_{n7}H_{an}O_{15}$	595.16575	595.16502	[W + H] ⁺	0.84	577.15147.457.10763.295.05903	0 0
51	27.40	Sennoside B	$C_{42}H_{38}O_{20}$	861.18837	861.18831	[M - M]	-1.71	(699.13514, 655.14385, 389.08791)	Я
52	27.44	Emodin-O-di-glucoside	$C_{27}H_{30}O_{15}$	593.15119	593.15112	$[M - H]^{-}$	-1.24	269.04549, 240.04318, 225.05493	R
53	28.01	Cassiachromone	$C_{13}H_{12}O_4$	231.06628	231.06653	$[M - H]^{-}$	-1.09	269.04549, 240.04321	R
54	28.96	Narirutin	$C_{27}H_{32}O_{14}$	581.18648	581.18594	$[\mathbf{M} + \mathbf{H}]^{+}$	0.94	435.12779, 419.13317, 273.07528	U
55	29.56	Isorhoifolin	$C_{27}H_{30}O_{14}$	579.17083	579.17056	$[M + H]^+$	0.47	449.14346, 303.08368, 153.01728	C
56	29.64	Chrysophanol-1,8-0-di-glucoside	$C_{27}H_{30}O_{14}$	577.15628	577.15658	$[M - H]^{-}$	-0.52	415.10387, 253.04962, 225.05395	R
57	30.04	Acteoside	$C_{29}H_{36}O_{15}$	623.19814	623.19902	$[M - H]^{-}$	-0.36	461.16314, 161.02265	М
58	30.15	1-Dihydro- <i>p</i> -coumaroyl-6- <i>O</i> - galloyl-glucoside	$C_{22}H_{24}O_{12}$	479.11950	479.12006	$[M - H]^-$	-1.16	461.10874, 313.05591, 169.01294, 125.02296	R
59	30.21	Prunin	$C_{21}H_{22}O_{10}$	435.12857	435.12887	$[M + H]^+$	-0.68	273.07329, 153.01674	U
60	30.22	Natsudaidai	$C_{21}H_{22}O_9$	419.13366	419.13349	$[M + H]^+$	0.39	390.09212, 389.09134, 371.08154	C
61	30.36	Naringin	$C_{27}H_{32}O_{14}$	581.18648	581.18605	$[M + H]^+$	0.75	435.12862, 419.13274, 273.07564	C
62	30.80	Sennoside A	$C_{42}H_{38}O_{20}$	885.18486	885.18538	$[M + Na]^+$	-0.59	701.13594, 657.14538, 391.08807, 770.03807	R
63	30.94	Isoacteoside	C ₂₀ H ₃₆ O ₁₅	623.19814	623.19925	$[M - H]^-$	-1.77	461.16741, 161.02568	М
64	30.98	Laccaic acid-D-O-glucoside	$C_{22}H_{20}O_{12}$	475.08740	475.08820	$[M - H]^{-}$	-0.91	431.09851, 269.03452, 253.05063,	R
65	31.45	Rhoifolin	$C_{27}H_{30}O_{14}$	579.17083	579.17057	$[M + H]^+$	0.57	240.04252 433.10764, 271.05967, 153.02389	C
<u>66</u>	31.45	Neodiosmin	$C_{28}H_{32}O_{15}$	609.18140	609.18107	$[M + M]^+$	0.77	301.07091, 286.04659	U
67	31.48	Isosakuranin	$C_{22}H_{24}O_{10}$	449.14422	449.14408	$[M + H]^+$	0.96	301.06877, 287.09012, 286.04485	C
68	31.59	Hesperidin	$C_{28}H_{34}O_{15}$	611.19705	611.19650	$[M + H]^+$	0.10	449.14312, 303.08581, 288.09732,	U
69	32.03	Nodakenetin	$\mathrm{C}_{14}\mathrm{H}_{14}\mathrm{O}_4$	245.08193	245.08221	$[M - H]^-$	-1.14	$153.02175, \ 229.10047, 211.64807, 175.42487$	Z
						,			

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Table 1 (Contd.)

ומחוב ד									
No.	$t_{ m R}$ (min)	Identification	Formula	Monoisotopic mass (<i>m</i> /z)	Detected mass (<i>m</i> / <i>z</i>)	Ion mode	udd	(z/ı/l) SW/SW	Main source
70	32.36	Physcion-1-0-glucuronic acid	$C_{22}H_{20}O_{11}$	459.09329	459.09368	$[M - H]^-$	-0.86	283.06124, 268.03815, 212.04737,	R
i					07607 600	_[11 _ 11]	010		Ç
1/	97.78	DIOSITI	C28H32U15	609.18140 609.18140	609.18081 609.18081	$[H + M]^+$	-0.42 0.96	299.10023, 284.29308 301.06863 286.04525	ر
72	32.89	Neohesperidin	$C_{28}H_{34}O_{15}$	611.19705	611.19637	$[\mathbf{H} + \mathbf{H}]^+$	0.53	449.14228, 303.08417, 153.01734	C
73	32.89	Hesperetin-7-0-glucoside	$C_{22}H_{24}O_{11}$	465.13914	465.13958	$[M + H]^+$	-0.92	303.08652, 288.03284, 153.01532	U
74	32.90	Sakuranetin	$C_{22}H_{24}O_{10}$	449.14422	449.14477	$[M + H]^+$	-1.22	301.06945, 287.08765, 286.045473	C
75	32.97	Magnoloside E	$C_{28}H_{34}O_{15}$	609.18249	609.18261	$[M - H]^-$	0.19	447.15213, 315.11019, 161.02613	Μ
76	33.05	Homoeriodictyol	$\mathrm{C}_{16}\mathrm{H}_{14}\mathrm{O}_{6}$	303.08631	303.08630	$[M + H]^{+}$	0.06	288.04856, 153.01712, 117.03375	U
77	33.18	Limocitrin-3-0-(3-hydroxy-3-	$C_{29}H_{32}O_{17}$	653.17123	653.17394	$[M + H]^+$	-0.73	539.13891, 347.07494	C
Ċ	01.00	methylglutarate)-glucoside				-[11 11]			¢
8/	33.4 0	1-Dinyurocinnanioyr-o-O-ganoyr- øhienside	U22H24U11	403.12439	403.12303	[н — м]	-0.97	331.00093, 313.03030, 211.02304, 169.01332	Ч
79	34.30	Magnolignan B	$C_{18}H_{20}O_5$	315.12380	315.12417	$[M - H]^{-}$	-0.95	267.10167, 249.09178, 221.09658, 133.06619	Μ
80	34.51	Meranzin	$\mathrm{C_{15}H_{16}O_{4}}$	261.11214	261.11203	$[M + H]^+$	0.42	243.10142, 159.04375, 103.05456	C
81	34.59	Hesperidone hydrate	$C_{15}H_{18}O_5$	296.14925	296.14925	$[M + HH_4]^+$	0.00	403.12656, 418.12351, 165.05352	C
82	35.89	Diosmetin-7-0-(6"-0-acetyl)	$C_{30}H_{34}O_{17}$	667.18688	667.18653	$[M + H]^+$	-0.26	301.07182, 286.05412	C
		neohesperidoside							i
83	36.14	Natsudaıdaın-3-0-(3-hydroxy-3- methylglutarate)-glucoside	$C_{33}H_{40}O_{18}$	725.22874	725.22824	[H + M]	0.47	581.18621, 419.13224	C
84	36.29	Magnolignan A	$\mathrm{C}_{18}\mathrm{H}_{20}\mathrm{O}_4$	299.12888	299.12924	$[M - H]^-$	-1.18	239.10865, 221.10746	Μ
85	37.01	Physcion-8-0-glucuronic acid	$C_{22}H_{20}O_{11}$	459.09329	459.09369	$[M - H]^-$	-0.89	283.06138, 268.03804, 239.03501,	R
								224.04837	
86	38.08	Cinnamoyl-6-0-galloyl-glucoside	$C_{22}H_{22}O_{11}$	461.10894	461.10948	$[M - H]^-$	-1.18	401.08794, 313.05874, 285.03987, 271.04612	R
87	39.00	Didymin	$C_{28}H_{34}O_{14}$	595.20213	595.20196	$[M + H]^+$	0.70	433.14783, 287.09001, 153.01794	U
88	39.08	Beta-hydroxyacteoside	$C_{29}H_{36}O_{16}$	639.19306	639.19315	$[M - H]^-$	-0.14	503.14287, 477.16173, 161.02408, 133.03174	W
89	39.39	6-Methoxy-8-0-β-D-glucoside	$C_{20}H_{24}O_9$	407.13476	407.13521	$[M - H]^{-}$	-1.12	253.04989, 201.09011, 146.96598	R
06	39.47	Magnoloside A	$C_{29}H_{36}O_{15}$	623.19705	623.19645	$[M + H]^{+}$	0.95	461.16607, 161.02574	М
91	39.51	Chrysophanol isomer	$C_{15}H_{10}O_4$	253.05063	253.05093		-1.23	225.05517, 210.03072, 197.06054	R
92	39.52	Chrysophanol-1-0-glucoside	$C_{21}H_{20}O_9$	415.10346	415.10402	$[M - H]^{-}$	-1.35	253.04884, 225.05627, 209.05942	R
93	39.70	Emodin- <i>O</i> -glucoside	$C_{21}H_{20}O_{10}$	431.09837	431.09867	$[M - H]^{-}$	-0.7	269.04562, 241.04997, 225.05524	В
94	39.74	3,5,6,7,8,3′,4′-Heptamethoxy	$C_{22}H_{24}O_9$	433.14931	433.14910	$[M + H]^+$	0.49	418.12356, 403.12656, 165.05354	C
		flavone							
95	39.79	1-P-Coumaroyl-6-0-galloyl-di- ماہیمونامہ	$C_{29}H_{36}O_{16}$	639.19306	639.19367	[M - M]	-0.96	477.13972, 313.05688, 211.02412,	R
90	20.02	Succosuc		00100	20100 200	$[\mathbf{M} + \mathbf{U}]^+$	94.0	141:00024 150 01701 100 06100	c
06	39.8/ 20.00	Donoirin	C16П14U5	201.09142 505 20212	201.0912/ 505 20168	[M + H] ⁺	0.40	133.UL/ZL, 133.U049Z 422 14812 - 287 00110 - 152 16712	ט כ
76	00.00 00.00	FOLICITIII Mean-clocide M	C28I134U9	51202.666 502 10705	00107.060	[M + M]	67.0 00.0	433.14812, 287.09110, 133.10/12 161 16637 161 00676	נ 2
90	59.95 20.05		C29H36U15	0791.620 202 10210	023.19043	[M + M]	70.1 10.1	401.1039/, 101.023/0	M
66	39.98	6'- <i>U-trans</i> -reruloyinodakenin	$C_{30}H_{32}O_{12}$	607.17865	607.17861	[M – H] [M + Na] ⁺	-0.02	247.10278, 229.44072, 175.28676 249.11947, 177.29759	Z
100	40.24	Chrysophanol	$\mathrm{C_{15}H_{10}O_4}$	253.05063	253.05094	$[M - H]^-$	-1.21	225.05537, 210.03104, 202.88598	Я
101	40.93	Marmin	$C_{19}H_{24}O_5$	333.16965	333.16942	[H + M]	0.70	163.03837, 107.08582	U

Paper

Table 1 (Contd.)

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	2-(2'-hydroxypropyl)-5-methyl-7- hydroxychromone Magnolignan C Naringenin Procyanidin B Physcion-8-O-glucuronic acid methyl ester Emodin-O-malonyl-glucose Chrysophanol-8-O-glucoside Emodin-O-glucoside Physcion-1-O-glucoside Physc	$\begin{array}{c} C_{13}H_{14}O_4\\ C_{13}H_{24}O_4\\ C_{15}H_{12}O_5\\ C_{30}H_{26}O_{12}\\ C_{30}H_{26}O_{12}\\ C_{23}H_{22}O_{11}\\ C_{24}H_{22}O_{13}\\ C_{21}H_{20}O_{10}\\ C_{21}H_{20}O_{10}\\ C_{21}H_{20}O_{10}\\ C_{22}H_{22}O_{10}\\ C_{16}H_{14}O_5\\ C_{16}H_{14}O_5\\ C_{15}H_{14}O_5\\ C_{18}H_{18}O_4\\ \end{array}$	233.08193 299.12888 273.07575 577.13515 473.10894	233.08217	$[M - H]^-$	-1.01	189.05568.146.96597.129.97617	
	Magnolignan C Naringenin Procyanidin B Physcion-8-O-glucuronic acid methyl ester Emodin-O-malonyl-glucose Chrysophanol-8-O-glucoside Emodin Aloe-emodin-O-glucoside Physcion-1-O-glucos	$\begin{array}{c} C_{18}H_{20}O_4\\ C_{15}H_{12}O_5\\ C_{30}H_{26}O_{12}\\ C_{30}H_{26}O_{12}\\ C_{23}H_{22}O_{11}\\ C_{24}H_{22}O_{13}\\ C_{21}H_{20}O_9\\ C_{15}H_{10}O_5\\ C_{21}H_{20}O_{10}\\ C_{22}H_{22}O_{10}\\ C_{21}H_{20}O_5\\ C_{16}H_{14}O_6\\ C_{15}H_{14}O_6\\ C_{15}H_{18}O_4\\ C_{18}H_{18}O_4\\ \end{array}$	299.12888 273.07575 577.13515 473.10894					R
	Naringenin Procyanidin B Physcion-8-O-glucuronic acid methyl ester Emodin-O-malonyl-glucose Chrysophanol-8-O-glucoside Emodin Aloe-emodin-O-glucoside Physcion-1	$\begin{array}{c} C_{15}H_{12}O_5\\ C_{30}H_{26}O_{12}\\ C_{30}H_{26}O_{11}\\ C_{23}H_{22}O_{11}\\ C_{24}H_{22}O_{13}\\ C_{24}H_{20}O_5\\ C_{15}H_{10}O_5\\ C_{15}H_{10}O_5\\ C_{21}H_{20}O_{10}\\ C_{22}H_{22}O_{10}\\ C_{16}H_{14}O_6\\ C_{15}H_{14}O_6\\ C_{15}H_{18}O_4\\ C_{18}H_{18}O_4\\ \end{array}$	273.07575 577.13515 473.10894	299.12928	$[M - H]^{-}$	-1.33	258.09247, 239.10852, 197.06244	М
	Procyanidin B Physcion-8-O-glucuronic acid methyl ester Emodin-O-malonyl-glucose Chrysophanol-8-O-glucoside Emodin Aloe-emodin-O-glucoside Physcion-1-O-glucosid	$\begin{array}{c} C_{30}H_{26}O_{12}\\ C_{23}H_{22}O_{11}\\ C_{23}H_{22}O_{13}\\ C_{24}H_{22}O_{13}\\ C_{21}H_{20}O_{5}\\ C_{15}H_{10}O_{5}\\ C_{21}H_{20}O_{10}\\ C_{22}H_{22}O_{10}\\ C_{22}H_{22}O_{10}\\ C_{16}H_{14}O_{6}\\ C_{15}H_{14}O_{6}\\ C_{15}H_{18}O_{4}\\ C_{18}H_{18}O_{4}\\ \end{array}$	577.13515 473.10894	273.07562	$[M + H]^+$	0.48	153.17823, 133.06712	U
	Physcion-8-O-glucuronic acid methyl ester Emodin-O-malonyl-glucose Chrysophanol-8-O-glucoside Emodin Aloe-emodin-O-glucoside Physcion-1-O-glucoside Physcion Hesperetin Magnatriol Magnatriol Magnolignan E 5-0-Demethylnobiletin Demethylnobiletin Demethylnobiletin Obovatol	$\begin{array}{c} C_{23}H_{22}O_{11}\\ C_{24}H_{22}O_{13}\\ C_{24}H_{22}O_{13}\\ C_{21}H_{20}O_{5}\\ C_{15}H_{10}O_{5}\\ C_{21}H_{20}O_{10}\\ C_{22}H_{22}O_{10}\\ C_{16}H_{12}O_{5}\\ C_{16}H_{14}O_{6}\\ C_{15}H_{14}O_{3}\\ C_{18}H_{18}O_{4}\\ \end{array}$	473.10894	577.13535	$[M - H]^{-}$	-0.49	425.08747, 407.07637, 289.07107	R
	Emodin-O-malonyl-glucose Emodin-O-malonyl-glucoside Emodin Aloe-emodin-O-glucoside Physcion-1-O-glucoside Physcion Hesperetin Magnatriol Magnatriol Magnolignan E 5-0-Demethylnobiletin Demethylnobiletin Obovatol	$\begin{array}{c} C_{24}H_{22}O_{13}\\ C_{21}H_{20}O_9\\ C_{15}H_{10}O_5\\ C_{21}H_{20}O_{10}\\ C_{22}H_{22}O_{10}\\ C_{22}H_{22}O_{10}\\ C_{16}H_{14}O_5\\ C_{16}H_{14}O_6\\ C_{15}H_{14}O_5\\ C_{18}H_{18}O_4\\ \end{array}$		473.10946	$[M - H]^{-}$	-1.11	283.06094, 268.03811, 240.04325	R
	Chrysophanol-8-O-glucoside Emodin Aloe-emodin-O-glucoside Physcion-1-O-glucoside Physcion Hesperetin Magnatriol Magnolignan E 5-Hydroxy-6,7,3',4',5'- pentamethoxyflavone 5-0-Demethylnobiletin Demethylnobiletin Obovatol	$\begin{array}{c} C_{21}H_{20}O_{9}\\ C_{15}H_{10}O_{5}\\ C_{21}H_{20}O_{10}\\ C_{22}H_{20}O_{10}\\ C_{16}H_{12}O_{5}\\ C_{16}H_{12}O_{5}\\ C_{16}H_{14}O_{6}\\ C_{15}H_{14}O_{6}\\ C_{15}H_{18}O_{4}\\ \end{array}$	517.09876	517.09935	$[M - H]^{-}$	-1.15	473.10884, 431.09157, 269.04531, 241 05131	Я
	Emodin Aloe-emodin-O-glucoside Physcion-1-O-glucoside Physcion Hesperetin Magnatriol Magnolignan E 5-Hydroxy-6,7,3',4',5'- pentamethoxyflavone 5-o-Demethylnobiletin Demethylnobiletin Obovatol	$C_{15}^{11}H_{20}O_{5}$ $C_{15}H_{10}O_{5}$ $C_{21}H_{20}O_{10}$ $C_{22}H_{22}O_{10}$ $C_{16}H_{12}O_{5}$ $C_{16}H_{14}O_{6}$ $C_{15}H_{14}O_{6}$ $C_{18}H_{18}O_{4}$	415 10346	415 10400	[М — М]	-1 30	211:02121 253 04084 225 05618 200 05050	ď
	Aloe-emodin-O-glucoside Physcion-1-O-glucoside Physcion Hesperetin Magnatriol Magnolignan E 5-Hydroxy-6,7,3',4',5'- pentamethoxyflavone 5-o-Demethylnobiletin Demethylnobiletin Obovatol	$\begin{array}{c} C_{21}H_{20}O_{10}\\ C_{21}H_{20}O_{10}\\ C_{22}H_{22}O_{10}\\ C_{16}H_{12}O_5\\ C_{16}H_{14}O_6\\ C_{15}H_{14}O_6\\ C_{15}H_{14}O_3\\ C_{18}H_{18}O_4\\ \end{array}$	269.04555	269.04588		-1.22	241.05016. 225.05524. 197.06027	4 24
	Physcion-1-O-glucoside Physcion Hesperetin Magnatriol Magnolignan E 5-Hydroxy-6,7,3',4',5'- pentamethoxyflavone 5-o-Demethylnobiletin Demethylnobiletin Obovatol	$C_{18}^{22}H_{22}O_{10}$ $C_{16}H_{12}O_5$ $C_{16}H_{14}O_6$ $C_{15}H_{14}O_3$ $C_{18}H_{18}O_4$	431.09885	431.09886	1	-1.14	269.04462, 239.03647, 211.02524	R
	Physcion Hesperetin Magnatriol Magnolignan E 5-Hydroxy-6,7,3',4',5'- pentamethoxyflavone 5-o-Demethylnobiletin Demethylnobiletin Obovatol	$C_{16}H_{12}O_5$ $C_{16}H_{14}O_6$ $C_{15}H_{14}O_6$ $C_{15}H_{14}O_3$ $C_{18}H_{18}O_4$	445.11402	445.11460	$[M - H]^-$	-1.30	427.03257, 283.06205, 269.04607	В
	Hesperetin Magnatriol Magnolignan E 5-Hydroxy-6,7,3',4',5'- pentamethoxyflavone 5- <i>o</i> -Demethylnobiletin Demethylnobiletin Obovatol	$C_{16}H_{14}O_{6}C_{15}H_{14}O_{6}C_{15}H_{14}O_{3}C_{18}H_{18}O_{4}$	283.06120	283.06162		-1.49	283.07696, 268.04557, 240.04266	R
	Magnatriol Magnolignan E 5-Hydroxy-6,7,3',4',5'- pentamethoxyflavone 5- <i>o</i> -Demethylnobiletin Demethylnobiletin Obovatol	$C_{15}H_{14}O_3$ $C_{18}H_{18}O_4$	303.08631	303.08608	$[M + H]^{+}$	0.77	153.01741, 110.04783	C
	Magnolignan E 5-Hydroxy-6,7,3',4',5'- pentamethoxyflavone 5-0-Demethylnobiletin Demethylnobiletin Obovatol	$\mathrm{C}_{18}\mathrm{H}_{18}\mathrm{O}_4$	241.08702	241.08730	$[M - H]^{-}$	-1.17	223.07597, 197.09716, 133.06719	М
	5-Hydroxy-6,7,3',4',5'- pentamethoxyflavone 5-0-Demethylnobiletin Demethylnobiletin Obovatol		297.11323	297.11365	$[M - H]^{-}$	-1.39	225.09132, 184.05786, 183.04759	М
	pentamethoxyflavone 5-0-Demethylnobiletin Demethylnobiletin Obovatol	$\mathrm{C}_{20}\mathrm{H}_{20}\mathrm{O}_{8}$	389.12309	389.12283	$[M + H]^+$	0.67	374.09631, 359.08221, 356.08174,	C
	5-0-Demethylnobiletin Demethylnobiletin Obovatol							
	Demethylnobiletin Obovatol	$\mathrm{C}_{20}\mathrm{H}_{20}\mathrm{O}_7$	373.12818	373.12781	$[M + H]^+$	0.73	358.10001, 343.07224, 325.06554,	U
	DemetnyInobiletin Obovatol	:			+[***		297.06973	C
	UD0Vat01	$C_{20}H_{20}O_8$	389.12309	389.12282	[H + M]	0.79	359.08112, 360.07831, 341.06984	נ :
		$C_{18}H_{18}U_3$	2601.1052	20211122	[M – M]	-1.20	20/.101/4, 249.08513	N (
	Sinensetin	$C_{20}H_{20}O_7$	373.12818	373.12790	$\left[H + M \right]$	0.74	175.01549, 147.03194, 119.01593	ວ :
	Honokiol	$C_{18}H_{18}O_2$	265.12340	265.12413	[M - M]	-0.42	224.08126, 223.07514, 197.05923	M
	O-Prenyl-umbelliferone	$\mathrm{C}_{14}\mathrm{H}_{14}\mathrm{O}_3$	229.08072	229.08726	[H - M]	-1.04	160.01707, 142.17814	Z
	Aloe-emodin	$C_{15}H_{10}O_5$	269.04555	269.04598	[H - M]	-1.60	253.05121, 240.04247, 211.03968	К
	Magnaldehyde D	$C_{16}H_{14}O_3$	253.08702	253.08735	$[M - H]^{-}$	-1.32	235.07963, 207.08634, 194.04012	Μ
	Phelloptein	$C_{17}H_{16}O_5$	323.08899	323.08875	$[M + Na]^{+}$	0.76	269.07627, 231.01796	Z
	Notopterol	$C_{21}H_{22}O_5$	353.13945	353.13988	$[M - H]^{-}$	-1.21	203.17578, 159.09786, 147.63478	Z
			377.13594	377.13559	$[M + Na]^+$	0.95	205.20652, 161.11072	
	Rhein	$C_{15}H_8O_6$	283.02481	283.02504	$[M - H]^{-}$	-0.82	257.04507, 239.03461, 211.03958	R
	Danthron	$\mathrm{C}_{14}\mathrm{H}_8\mathrm{O}_4$	239.03498	239.03530	$[M - H]^{-}$	-1.33	186.73096, 141.51394, 130.71196	R
	Nobiletin	$C_{21}H_{22}O_8$	403.13874	403.13819	$[M + H]^+$	1.41	373.09359, 355.08584, 327.08476	C
	Nomilin	$C_{28}H_{34}O_9$	515.22756	515.22753	$[M + H]^+$	0.05	469.21806, 455.20675, 411.21462,	C
							161.05854	
	Notoptol	$C_{21}H_{22}O_5$	353.13945	353.13987	$[M - H]^{-}$	-1.19	225.17472, 207.13978, 189.14075	z
			377.13594	377.13563	$[\mathbf{M} + \mathbf{N}\mathbf{a}]^{\dagger}$	0.97	227.19076, 209.14289	
	Magnaldehyde E	$C_{16}H_{14}O_{3}$	253.08702	253.08735	$[M + H]^+$	-1.32	225.09424, 184.05624, 183.05074	Μ
133 46.17	Magnolol	$C_{18}H_{18}O_2$	265.12340	265.12394	$[M - H]^{-}$	-0.39	247.10096, 245.09517, 223.07491	М
	<i>p</i> -Hydroxyphenethyl anisate	$C_{16}H_{16}O_4$	295.09408	295.09386	$[M + Na]^+$	0.73	153.07368, 138.08654	z
135 46.54	Obovaaldehyde	$\mathrm{C}_{16}\mathrm{H}_{14}\mathrm{O}_4$	269.08193	269.08215	$[M - H]^{-}$	-0.80	152.01478, 124.02018	М
	Isosinensetin	$\mathrm{C_{20}H_{20}O_7}$	373.12818	373.12783	$[M + H]^+$	0.94	358.10398, 343.08158, 315.08276	C
137 46.70	Tangeretin	$\mathrm{C}_{20}\mathrm{H}_{20}\mathrm{O}_7$	395.11012	395.10998	$[M + Na]^+$	0.35	278.54947, 276.55982, 243.12708	C

^a Ps: C represented Aurantii fructus immaturus; R represented Rheum palmatum L.; M represented Magnoliae Officmalis Cortex; N represented Notopterygii Rhizoma Et Radix.

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cleaving of the glycosidic linkage and losing $C_6H_{10}O_5$ (162 that the Da) from m/z 431.09157, m/z 241.05131 is the result of losing 197.05 CO (28 Da) at m/z 269.04531. The characterization of C_3H_5 (losing anthraquinone compounds in SHD is inferred from this losing

3.4 Characterization of glucosides in SHD

cleavage rule and related literatures.^{20,21}

Glucosides are formed by the esterification of acid with glucose. According to the type of acid, it can be divided into cinnamoyldihydrocinnamoyl-glucose, p-cinnamoyl-glucose, glucose, dihydro-p-cinnamoyl glucose, galloyl-glucose, and resveratrolgalloyl-glucose in rhubarb. They are vulnerable to loss of neutral fragments 148 Da, 150 Da, 164 Da, 166 Da, 170 Da, and 228 Da, respectively. Galloyl-glucose was selected to illustrate the fragmentation pathways. In negative ion mode, adduct ion at m/z 331.06759 ([M – H]⁻) indicating an elemental composition of $C_{13}H_{16}O_{10}$. The mass spectrum (Fig. 3D) shows that the main fragment ions are m/z 313.05289, 271.04772, 211.02436, 169.01338, and 125.00325. Among them, m/z 313.05289 is presumed by losing H₂O (18 Da) from $[M - H]^{-}$, m/z 271.04772 is inferred by losing $C_2H_4O_2$ (60 Da) from $[M - H]^-$, m/z211.02436 is the result of losing $C_4H_8O_4$ (120 Da) from [M -H]⁻, m/z 169.01338 is speculated by cleaving of the glycosidic linkage and losing glucose $C_6H_{10}O_5$ (162 Da) from $[M - H]^-$, and m/z 125.00325 is deduced by losing CO₂ (44 Da) from m/z169.01338. The characterization of glucosides in SHD is concluded by referring to the cleavage rule and related literature.21

3.5 Characterization of epicatechin in SHD

Epicatechin was selected to illustrate the fragmentation pathways for catechins. The adduct ion at 289.07213 ($[M - H]^-$) indicating an elemental composition of $C_{15}H_{14}O_6$, and the MS/ MS spectrum (Fig. 3E) is presented that the main fragment ions are m/z 245.08235, 203.07153. Among them, m/z 245.08235 is presumed by losing CO₂ (44 Da) from $[M - H]^-$, and m/z 203.07153 is inferred by losing of C_2H_2O (42 Da) from m/z 245.08235. The characterization of catechins in SHD is concluded by referring to the cleavage rule and related literatures.²¹⁻²³

3.6 Characterization of lignans in SHD

Most of the lignans have the presence of allyl and hydroxyl groups, which relative positions are different, their cleavage methods will be different. Honokiol and magnolol were selected to illustrate the fragmentation pathways. Magnolol showed m/z 265.12394 ($[M - H]^-$) in the negative ion mode, which indicated the elemental composition of $C_{18}H_{18}O_2$. The mass spectrum (Fig. 3F) shows that the main fragment ions are m/z 247.10096, 245.09517, 223.07491. Among them, m/z 247.10096 is presumed by losing H_2O (18 Da) from $[M - H]^-$, m/z 245.09517 is inferred by losing H_2 (2 Da) from m/z 247.10096, and m/z 223.07491 is obtained by losing C_3H_6 (42 Da) from $[M - H]^-$. Honokiol showed m/z 265.12413 ($[M - H]^-$) in the negative ion mode, which indicated the elemental composition of $C_{18}H_{18}O_2$. The mass spectrum (Fig. 3G) shows

that the main fragment ions are m/z 224.08126, 223.07514, 197.05923. Among them, m/z 224.08126 is presumed by losing C_3H_5 (41 Da) from $[M - H]^-$, m/z 223.07514 is inferred by losing C_3H_6 (42 Da) from $[M - H]^-$, and m/z 197.05923 is obtained by losing of C_2H_2 (26 Da) by m/z 223.07514. The characterization of lignans in SHD is concluded by referring to the cleavage rule and related literatures.^{24,25}

3.7 Characterization of phenylpropanoid glycosides in SHD

The phenylpropanoid glycosides are composed of glucose and phenethyl alcohol, which mainly based on the loss of coffee groups (162 Da). Verbascoside is chosen as an example to explain the cleavage pathway. It showed m/z 623.19814 ([M – H]⁻) in the negative ion, which indicated an elemental composition of C₂₉H₃₆O₁₅. The mass spectrum (Fig. 3H) shows that its main fragment ion is m/z 461.16314, 161.02265. Among them, m/z 461.16314 is presumed by losing C₉H₆O₃ (162 Da) from [M – H]⁻, and m/z 161.02265 is inferred by losing C₂₀H₃₀O₁₂ (462 Da) from [M – H]⁻. The characterization of phenylpropanoid glycosides in SHD is concluded by referring to the cleavage rule and related literatures.^{24,25}

3.8 Characterization of alkaloids in SHD

The main rupture method of alkaloids is the loss of N. (*R*)-Oblongine is chosen as an example to explain the cleavage pathway. It showed an m/z 314.17473 ([M + H]⁺) in the positive ion mode, which indicated the elemental composition of $C_{19}H_{24}NO_3^+$. The mass spectrum (Fig. 3I) shows that the main fragment ions are m/z 269.11625, 237.09024, 175.07485, 107.04861. Among them, m/z 269.11625 is presumed by losing C_2H_6NH (45 Da) from [M + H]⁺, m/z 237.09024 is inferred by losing CH₄O (32 Da) at m/z 269.11625, m/z 175.07485 is obtained by losing C₆H₆O (94 Da) from m/z 269.11625, and m/z 107.04861 is speculated by losing $C_{12}H_{18}NO_2$ (207 Da) from [M + H]⁺. The characterization of alkaloids in SHD is concluded by referring to the cleavage rule and related literature.²⁵

3.9 Characterization of coumarins in SHD

The main cleavage mode of coumarins is loss of CO and CO₂, while coumarin glycosides lose glucose firstly, and then break by the above-mentioned cleavage law. Bergaptol is chosen as an example to explain the cleavage pathway. It showed an m/z 201.01957 ($[M - H]^-$) in the negative ion mode, which indicated an elemental composition of C₁₁H₆O₄. The mass spectrum (Fig. 3J) shows that the main fragment ions are m/z 185.02456, 157.17256, 129.10451, 101.96320. Among them, m/z 185.02456 is presumed by losing OH (17 Da) from $[M - H]^-$, m/z 157.17256, is presumed by losing CO (28 Da) from m/z 185.02456, and m/z 101.96320 is obtained by losing CO (28 Da) from m/z 187.17256, and m/z 101.96320 is obtained by losing CO (28 Da) from m/z 169.10451. The characterization of coumarins in SHD is concluded by referring to the cleavage rule and related literatures.²⁶⁻²⁸

4. Conclusions

In our study, a rapid, swift and straightforward analytical method with the help of UHPLC-FT-ICR-MS/MS was successfully developed for the first time to separate and identify the chemical constituents of SHD. A total of 137 compounds in SHD were identified or tentatively characterized, and 11 compounds were accurately identified by reference. The research indicated that SHD mainly contains 42 flavonoids (in Aurantii fructus immaturus and Rheum palmatum L.), 3 triterpenoids (in Aurantii fructus immaturus), 23 anthraquinones (in Rheum palmatum L.), 14 glucosides, 5 catechins and 2 anthranone (in Rheum palmatum L.), 11 phenylpropanoid glycosides, 5 alkaloids and 11 lignans (in Magnoliae Officmalis Cortex), 16 coumarins (in Notopterygii Rhizoma Et Radix and in Aurantii fructus immaturus), which has laid the foundation for prospective research associated with SHD. And it is hoped to be useful to identify and characterize the components in other TCMs. And it provided a basis for finding the prototype components and metabolites in the serum of containing SHD.

In this study, the structure of the compound, except for the comparison of the reference, was obtained through the combination of retention time, accurate primary and secondary mass spectrometry information, cracking rules of the same compound and literature comparison. The identification of compound structure by this method is limited to known compounds, because this method only mass spectrometry for identification.

Abbreviations

TCMs	Traditional Chinese medicines
SHD	Sanhua decoction
Zhishi	Aurantii fructus immaturus
Dahuang	Rheum palmatum L
Houpu	Magnoliae Officmalis Cortex
Qianghuo	Notopterygii Rhizoma Et Radix
BBB	Brain blood barrier
UHPLC-FT-ICR-	Ultra high performance liquid
MS/MS	chromatography coupled with Fourier
	transform ion cyclotron resonance mass
	spectrometry
BPC	Base peak ion chromatograms
EIC	Extract ion chromatograms
RDA	Diels–Alder reaction

Conflicts of interest

The authors declare that there are no conflicts of interest.

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