Nontargeted and targeted metabolomics analysis provides novel insight into nonvolatile metabolites in Jianghua Kucha tea germplasm (Camellia sinensis var. Assamica cv. Jianghua)

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Abstract

Jianghua Kucha (JHKC) is a special tea germplasm with high bitterness growing in China; however, the chemical characteristics of JHKC are not completely understood. In this study, 61 differential metabolites were identified between 11 wild JHKC individuals and 3 control cultivars of Fudingdabai, Yunkang 10, and Zhuyeqi using comprehensive nontargeted and targeted metabolomics approach. The JHKC accessions mainly possessed significantly higher levels of purine alkaloids of theacrine (12.06 ± 5.23 mg/g) and 1,3,7-trimethyluric acid, non-epi-form flavanols (catechin, gallocatechin, catechin gallate, and gallocatechin gallate), and methylated flavanols of epigallocatechin-3-O-(3′-O-methyl)-gallate (4.79 ± 1.45 mg/g) and epicatechin-3-O-(3′-O-methyl)-gallate (1.02 ± 0.34 mg/g), as well as significantly lower levels of flavonol glycosides, which indicated that caffeine metabolism, flavonoid biosynthesis, and flavonol and flavone biosynthesis are mostly differential metabolic pathways. Our study demonstrated that JHKC germplasm is a promising resource for breeding novel tea cultivars with high contents of theacrine, non-epi-form flavanols, and methylated flavanols.

1. Introduction

Tea (Camellia sinensis) is one of the most popular and oldest non-alcoholic beverage crop species worldwide, and its health benefits have attracted widespread attention. China is likely the place of origin of tea plants and there are >3000 tea accessions and 900 bred cultivars (Ma, Yao, & Chen, 2015).

Kucha (Camellia sinensis) is a unique tea plant variety that tastes much more bitter than common tea cultivars widely planted in China and is primarily distributed in Yunnan, Hunan, Guangdong, Jiangxi and Guizhou provinces. Although Kucha has a strong bitter taste, locals enjoy drinking it to promote health. Previous studies have demonstrated that Kucha tea extract has relatively high antioxidative activity (Li, et al., 2012) and significantly suppressed the accumulation of lipid droplets (Li, et al., 2016). Jianghua Kucha [JHKC, Camellia sinensis var. assamica (CSA) cv. Jianghua] is an ancient and specific tea germplasm resource (Li, & Lao, 2011) that grows in the Nanling Mountain region in Hunan Province, Southeastern China. JHKC plants exist as semi-arbor or small-arbor trees with an upright shape, and their height varies from 2.5 to 6 m. The morphological characteristics of JHKC differ among population accessions; for example, white-leaf Kucha, golden-leaf Kucha, cyan-leaf Kucha, bamboo-leaf Kucha, and salix-leaf Kucha, each of whose leaves are morphologically distinct (Liu, 2006). Unique germplasms constitute the foundation for the breeding of new varieties, and an increasing number of tea tree breeders are focusing on the amounts of functional ingredients of germplasms, e.g., low amount of caffeine, no caffeine, high amount of theanine, high amount of flavonols, high amount of methylated catechin, and high amount of theacrine (Chen, Apostolides & Chen, 2013; Yu, et al., 2020; Qin, et al., 2020). At present, three cultivars have been bred from JHKC by the use of traditional breeding method (individual selection), namely, Jianghua Kucha 21–1 (JHKC-21–1), Jianghua Kucha 21–3 (JHKC-21–3) and Jianghua Kucha 37–2 (JHKC-37–2), all of which are suitable for the production of black tea (Li, Huang, Zhong, & Su, 2018).

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JHKC is highly diverse, and there is an abundance of different populations of JHKC accessions. However, most researchers have analyzed the contents of only common nonvolatile compounds (for instance, water extracts, flavanols, amino acids, caffeine, and theobromine, etc.) in fresh leaves of JHKC and bred cultivars from JHKC (JHKC-B) (Li, 2012; Yang, 2013). Both a comprehensive investigation of nonvolatile metabolites and identification of new components in JHKC are needed. Nontargeted metabolomics combined with chemometric analysis enables the comprehensive detection of nonvolatile metabolites and has been widely used in tea plant studies (Dai, et al., 2018; Li, et al., 2019).

To comprehensively understand the chemical constitutions of JHKC germplasms and to utilize JHKC germplasms effectively for breeding, in this work, we investigated the differential accumulation of nonvolatile metabolites of tea varieties among representative JHKC plants, common tea cultivars and JHKC-B cultivars, based on nontargeted and targeted metabolomics analysis with ultra-high-performance liquid chromatography quadrupole time-of-flight mass spectrometry (UHPLC-QTOF/MS) and ultra-performance liquid chromatography triple quadrupole mass spectrometry (UPLC-QqQ). This study provides novel insight into the metabolite profile and intrinsic metabolic mechanisms of JHKC germplasms.

2. Materials and methods

2.1. Chemicals

Formic acid and methanol (LC-MS grade) were purchased from TIC (Tokyo, Japan) and Merck (Darmstadt, Germany), respectively. Deionized ultrapure water was obtained from Millipore (Bedford, MA, USA). Flavanols of epigallocatechin gallate (EGCG), epicatechin gallate (ECG), epicatechin (EC), epicatechin-3-O-(3'-O-methyl)-gallate (EGCG3′Me) and epicatechin-3-O-(3'-O-methyl)-gallate (EGC3′Me) were purchased from Nagara Science Co., Ltd. (Gifu, Japan). Phenylethyl primeveroside and benzyl primeveroside were synthesized by staff at the National Glycoengineering Research Center, Shandong University, Jinan, China (Li et al., 2019).

2.2. Materials

Three experimental tea groups were included in this study (Fig. 1). The JHKC group comprised 11 JHKC [Camellia sinensis var. assamica (CSA) cv. Jianghua] individuals of local accessions whose morphology was representative. The control (CK) group comprised three common tea cultivars: Fudingdabai [FDDB, Camellia sinensis var. sinensis (CSS)], Yunkang 10 (YK10, CSA), and Zhuyeqi (ZYQ, CSS). FDDB originates from Fujian Province of Southeastern China, YK10 originates from Yunnan Province of Southwestern China, and ZYQ originates from Hunan Province of Southcentral China. The JHKC-B group comprised three cultivars bred from local JHKC varieties, namely, JHKC-21–1 (CSA), JHKC-21–3 (CSA) and JHKC-37–2 (CSA). These three JHKC-bred

![Fig. 1. Workflow of samples collection and metabolomics investigation of JHKC accessions in this study: (1) the metabolomes of 11 wild JHKC individuals (JHKC group) and of three CK cultivars (FDDB, YK10, and ZYQ) were compared to investigate the most differential metabolites and differential metabolic pathways in JHKC; (2) three JHKC-bred cultivars were surveyed to further study the changes in differential metabolites after artificial breeding and cultivation.](image-url)
cultivars were registered in 2019 under the following registration numbers: GPD tea plant (2019)430028, GPD tea plant (2019)430031, and GPD tea plant (2019)430029.

The individuals in the JHKC group were collected from a mountainous region at elevations ranging from 400 to 600 m in Jianghua County, Hunan Province, while the CK and JHKC-B groups were collected from tea plantations of the Tea Research Institute, Hunan Academy of Agricultural Sciences, Changsha, Hunan, China. One bud with two leaves was removed from the tea plants on April 11–16, 2020, in triplicate, frozen in liquid nitrogen, and then freeze-dried. The freeze-dried tea leaves were subsequently ground into a powder using a mill (IKA; Staufen, Germany) and then stored at –20 °C until use.

2.3. Tea sample preparation

Tea compounds were extracted according to the methods of our previous study, with minor adjustments (Dai, et al., 2020). Fifteen ml of a prewarmed 70% (v/v) methanol solution was added to 100 mg of tea powder, after which the mixture was placed in a water bath at 70 °C for 0.5 h. The solution was subsequently centrifuged at 8000 × g for 15 min (Centrifuge 5810R, Eppendorf, Hamburg, Germany), after which the supernatant was filtered through a 0.22 μm membrane for metabolomics analysis and absolute quantification of the compounds. Each tea sample was prepared in triplicate. Quality control (QC) samples were prepared by mixing 100 μl aliquots of each tea infusion.

2.4. Nontargeted metabolomics analysis

Nontargeted metabolomics analysis was conducted according to our previous methods (Li, et al., 2018; Dai, et al., 2020). Briefly, an UHPLC system (Infinity 1290, Agilent Technologies, Santa Clara, CA, USA) connected to a Q-TOF/MS instrument (6545, Agilent Technologies, Santa Clara, CA, USA) was used for the metabolomics analysis in ESI+ mode. The compounds in the tea samples were separated with a Zorbas Eclipse Plus C18 column (150 × 3.0 mm, 1.8 μm, Agilent Technologies, Little Falls, DE, USA), and the column temperature was maintained at 40 °C; phase A consisted of 0.1% (v/v) formic acid-deionized water, phase B consisted of methanol at a flow rate of 0.4 mL/min, and the injection volume was 3 μL. The linear gradient elution procedure was as follows: 10% B (0–3 min), 10–15% B (3–4 min), 15–30% B (4–7 min), 30–50% B (7–16 min), 50–95% B (16–20 min), 95% B (20–22 min), 95–5% B (22–22.05 min), and 5% B (22.05–26 min). The MS parameters were set as follows: the temperature and the flow rate of the drying gas were 325 °C and 10 L/min, respectively; the temperature and the flow rate of the sheath gas were maintained at 325 °C and 11 L/min, respectively; the capillary voltage was 4 kV; the nebulizer pressure was 40 psi; and the mass scan range was m/z 100–1000. QC samples were injected after every nine samples to monitor the reproducibility of the metabolomic analysis results.

The raw metabolomics data acquired from UHPLC – QTOF/MS were first treated for peak picking using MassHunter Qualitative Analysis software (B.07.00 SP1, Agilent Technologies, Santa Clara, CA, USA) and for peak alignment using Mass Profiler Professional (version 13.0, Agilent Technologies, Santa Clara, CA, USA). The signal-to-noise (S/N) threshold for peak detection was set to 3. The retention time tolerance and mass tolerance for the peak alignment was set to 0.2 min and 0.01 Da, respectively. The obtained compound ion features with relative standard deviations (RSD) of mass intensities <30% in QC samples were used for multivariate and univariate analysis. Compounds were structurally identified based on authentic standards, MS² spectra, mass accuracy, metabolomics databases (Metlin and HMDB), and the previous works (Dai, et al., 2015; Dai, et al., 2017; Yang, et al., 2018; Chen, et al., 2020).

2.5. Absolute quantifications of flavanols, methylated catechins, and alkaloids

The absolute amounts of eight flavanols (EGCG, EGC, ECG, EC, GCG, GC, C, and C), two methylated flavanols (EGCG3′′Me and EGCG3′′Me), and three alkaloids (theobromine, caffeine and theacrine) were quantified by an UPLC-PDA-QTS system (Waters, Manchester, UK) in accordance with an external standard method according to our previous work (Dai, Hu, Xie, Tan, & Lin, 2020).

2.6. Statistical and data analysis

Principal component analysis (PCA) and partial least squares discriminant analysis (PLS-DA) were performed using SIMCA 14.1 software (Umetrics, Umeå, Sweden). Statistical analyses were performed using SPSS 21 software (IBM, Chicago, IL, USA) via one-way ANOVA and Student’s t-tests. A heat map was generated by MultiExperiment Viewer 4.9.0 (Oracle, Redwood, CA, USA), and pathway analysis was performed by MetaboAnalyst 4.0 (https://www.metaboanalyst.ca/).

3. Results and discussion

3.1. Differential metabolites between JHKC accessions and CK cultivars

To comprehensively investigate the differential metabolites between wild JHKC (11 JHKC individuals) and CK (FDDB, YK10 and ZYQ) cultivars, the compounds in tea samples were measured via nontargeted metabolomics analysis by the use of UHPLC-QTOF/MS. A total of 3226 metabolite ion features were acquired after peak extraction and alignment. Among them, the RSD of 2258 metabolite ion features were <30% in QC samples; these were used for further statistical analysis. The QC samples clustered together at the center of the PCA score plot (Fig. 2A), indicating good reproducibility of the metabolomic analysis. In addition, the JHKC and CK samples were clearly divided into two groups via PCA and PLS-DA (Fig. 2A-B). The JHKC accessions clustered on the left of the PCA and PLS-DA score plots, while the CK group accessions (FDDB, YK10 and ZYQ) clustered on the right. These results indicated that the metabolite patterns of the JHKC accessions and CK cultivars are significantly different. The R² and Q² intercepts were 0.296 and –0.327, respectively, indicating that the PLS-DA model was not overfitted (Fig. 2C). A total of 146 compounds (Table S1) were ultimately structurally identified or tentatively identified, among them, 61 compounds, including five alkaloids, 11 flavanols, nine dimeric flavanols, 15 flavonol/flavone glycosides, four amino acids, four phenolic acids, three aroma precursors, six lipids and four other compounds, were screened out as the main differential metabolites between the JHKC and CK groups (shown as red dots in the S-plot, Fig. 2D). A PCA biplot (in which tea samples are shown as orange diamonds and in which metabolites are shown as green dots) was constructed to display the correlations between the tea samples and the 61 main differential compounds (Fig. 2E). Metabolites with relatively high content in the tea samples are located in the same quadrant. In addition, a heatmap was created to visualize the variations of the 61 differential metabolites in the JHKC and CK groups (Fig. 3). The tea samples clustered into two groups, a JHKC group and a CK group, which was consistent with the PCA and PLS-DA results (Fig. 2A-B). In the heatmap, a red color indicates that the content of a metabolite is higher than the average level among the samples, while a green color indicates that the content of a metabolite is at a lower level. The results clearly showed that, compared with CK group (FDDB, YK10, and ZYQ), the JHKC group mainly contains higher contents of non-epiflor flavanols of C, GC, CG, and GCG, purine alkaloids of theaeneine and 1,3,7-trimethyluric acid, methylated flavanols of EGCG3′′Me and EGCG3′′Me, myricetin-3′-glucoside, ECG, procyanidin B3, and vaine. In contrast, the levels of most flavonol glycosides (kaempferol, quercetin and myricetin glycosides), EC, EGC, procyanidin B1, procyanidin B2, theobromine, theogallin, arginine, proline, and thepophan were lower.
Fig. 2. Multivariate statistical analysis of JHKC accessions and CK cultivars (FDD8, YK10, and ZYQ): (A) PCA score plot; (B) PLS-DA score plot, $R^2_X = 79.1\%$, $R^2_Y = 99.6\%$, $Q^2 = 98.7\%$; (C) cross-validation plot of PLS-DA model with 200 permutation tests; (D) S-plot of PLS-DA (the red dots represent the most differential compounds); (E) PCA biplot of the levels of differential metabolites in JHKC accessions and CK cultivars (FDD8, YK10, and ZYQ). The orange diamonds represent tea samples, and the green dots represent metabolites. Positive correlations are present between any two variables located in the same quadrant.
in the JHKC group than in the CK group (Fig. 2E and Fig. 3). Notably, purine alkaloids of theacrine and 1,3,7-trimethyluric acid, methylated flavanols of EGCG\(^3\)Me and ECG\(^3\)Me, non-epi-form flavanols of CG and GCG were 10-fold higher contents in the JHKC group than in the CK group; these compounds could therefore be considered characteristic compounds of JHKC germplasms. On the other hand, the contents of most flavonol glycosides decreased, even the flavonol glycosides with trisaccharide moieties, such as kaempferol 3-glucosylrutinoside, kaempferol 3-galactosylrutinoside, quercetin triglucoside, and quercetin 3-glucosylrutinoside, were absent in the JHKC group.

### 3.2. Differences in metabolic pathways between JHKC accessions and CK cultivars

To understand the differences in the metabolic pathways between the JHKC accessions and CK cultivars, metabolic pathway analysis of the 61 differential metabolites was performed by MetaboAnalyst 4.0 (Fig. S1). The differential metabolites were involved mainly in the metabolic pathways associated with flavonoid biosynthesis, caffeine metabolism, flavone and flavonol biosynthesis, arginine biosynthesis, arginine and proline metabolism, valine-leucine and isoleucine biosynthesis, aminoacyl-tRNA biosynthesis, glycerophospholipid metabolism, and pantothenate and CoA biosynthesis. These results suggested that the flavonoid biosynthesis, caffeine metabolism, flavone and flavonol biosynthesis pathways are the most critical metabolic pathways that
differ between JHKC accessions and CK tea cultivars. Hence, the changes of critical metabolites involved in these pathways were further mapped, the results of which are shown in Fig. 4.

Flavonoids are metabolic products of flavonoid biosynthesis, flavone and flavanol biosynthesis pathways, mainly including flavanols, dimeric flavanols and flavonol/flavone glycosides. As shown in Fig. 4 and Table 1, as JHKC belonging to CSA accessions, the composition and content of flavanols in JHKC was partly consistent with the findings of previous studies (Jin, Ma, Ma, Yao, & Chen, 2014; Yu, et al., 2020). For instance, Jin, et al. (2014) reported that CSA accessions have relatively high contents of C, EC and ECG and that CSS accessions have a relatively high content of EGC; Yu, et al. (2020) demonstrated that C, EC, ECG and GC accumulated to high levels in CSA accessions or hybrid accessions with a CSA genetic background. Interestingly, our results showed that EC accumulated at lower levels in the JHKC accessions than in the CK cultivars [YK10 (CSA accession), FDDB (CSS accession) and ZYQ (CSS accession)]. Furthermore, GCG was present at a high level (17.73 ± 1.29 mg/g, ranging from 15.75 to 19.65 mg/g) in the JHKC, while was absent in the CK cultivars (FDDB, YK10, and ZYQ) (Table 1). In general, EGCG and EGC were present at the highest levels in the tea plants, followed by ECG, EC, and C (Yu, et al., 2020). However, EGCG (64.09 ± 5.17 mg/g) and GCG (17.73 ± 1.29 mg/g) were the most abundant flavanols in the JHKC accessions. Taken together, these results suggested that JHKC is a unique tea germplasm resource.
It is well known that leucoanthocyanidin reductase (LAR), anthocyanidin synthase (ANS), anthocyanidin reductase (ANR), echinatin: 1-O-galloyl-β-D-glucose O-galloyltransf erase (EGOT) and UDP-glucose: galloyl-1-O-β-D-glucosyltransferase (UGGT) are critical enzymes involved in the flavonoid biosynthesis pathway and contribute to the differential accumulations of flavanol components in different teas. LAR catalyzes the conversion of leucoanthocyanidin/leucodelphinidin to C/GC, ANS catalyzes the conversion of leucocyanidin/leucodelphinidin to cyanidin/delphinidin, and ANR catalyzes the conversion of cyanidin/ delphinidin to EC/EGC (Yu et al., 2020). In addition, EGOT and UGGT are involved in the galloylated flavans (GG, GCG, EGCG, and ECGG) biosynthesis pathway (Liu et al., 2012). As described above, the high accumulations of C and GC in the JHKC accessions may be due to high LAR activity; in contrast, the low accumulations of EC and EGC in the JHKC accessions may be due to low ANR activity. The concentrations change of galloylated flavans exhibited higher galloylation levels in the JHKC accessions than in the CK cultivars. For example, the content of EC and EGC was 0.5-fold lower in the JHKC group than in the CK group, respectively, while the contents of EGC and ECGG were slightly higher in the JHKC group (Table 1), indicating that the UGGT and EGOT enzyme activity may be higher in the group JHKC than in the CK group.

Numerous studies have demonstrated that, compared with EGC and ECGG, EGC3’Me and EGC3’Me methylated flavans exhibit stronger antiallergic and metabolic syndrome-treating effects, respectively (Maeda-Yamamoto, Ema, Monobe, Tokuda, & Tachibana, 2012; Yang, Qiao, Zhang, Wu, & Weng, 2015; Márquez Campos, Jakobs, & Simon, 2020). Moreover, it was strongly suggested that the EGC3’Me content in tea leaves was positively correlated with the expression of the genes encoding leucoanthocyanidin reductase (LAR), dihydroflavonol-4-reductase (DFR) and caffeoyl-CoA 3-O-methyltransferase (CCoAOMT); however, some tea varieties seemed to lack these related enzymes (Luo et al., 2018). In the present study, compared with the CK cultivars, the JHKC accessions had significantly higher contents of EGC3’Me and EGC3’Me (Fig. 4, Fig. 5F, H and Table 1). The EGC3’Me level in the JHKC accessions ranged from 2.05 to 7.00 mg/g (4.79 ± 1.45 mg/g), and the EGC3’Me level ranged from 0.44 to 1.50 mg/g (1.02 ± 0.34 mg/g). Surprisingly, among the cultivars composing CK group, only the FDDB cultivar presented a small amount of EGC3’3Me (0.04 ± 0.01 mg/g) and EGC3’3Me (0.01 ± 0.00 mg/g). A previous study (Maeda-Yamamoto, et al., 2012) showed that EGC3’3’Me and EGC3’3’Me coexisted in different tea varieties and that their contents were strongly positively correlated, which was consistent with our data (Table 1). The content of EGG3’3Me did not obviously differ between the JHKC and CK groups; however, the methylated flavanol (EGCG3’3Me) content significantly differed in this study, indicating that methylation levels of flavans were higher in the JHKC accessions than in CK cultivars. It demonstrated that the JHKC accessions are good materials for breeding plants with high amounts of bioactive components of methylated flavans (ECCG3’3Me and EGC3’3Me), such as the Benifuuki cultivar in Japan (Maeda-Yamamoto, et al., 2012).

The metabolic pathway of proanthocyanidins (PAs) was also mapped, as shown in Fig. 4. Procyanidin B1 (EC[4,8]-C) and procyanidin B2 (EC[4,8]-EC) are EC-based PAs; procyanidin B3 [C(4,8)-C] and procyanidin B4 [C(4,8)-EC] are C-based PAs (Wang, et al., 2020). Procyanidin B3 abundantly accumulated in the JHKC accessions, while procyanidin B1 and procyanidin B2 highly accumulated in the CK cultivars, as C was highly enriched in the JHKC accessions, while E highly accumulated in the CK cultivars.

The glycoside forms of quercetin, myricetin and kaempferol are downstream metabolic products of naringenin (Fig. 4). In this study, with the exception of myricetin 3-glucoside, glycosides of quercetin, myricetin and kaempferol were present at significantly lower levels in the JHKC accessions than in the CK cultivars. These results indicated more vigorous biosynthesis of quercetin, myricetin and kaempferol glycosides in the CK cultivars than in the JHKC accessions. Considering that the JHKC accessions contained higher levels of flavans, we deduced that the flavonoid biosynthesis pathway tended to synthesize flavanols rather than flavonoids in the JHKC accessions.

Caffeine metabolism was one of the most differential pathways between the JHKC and CK groups (Fig. 4). This study showed that caffeine, theacrine and theobromine were the main purine alkaloid components in the JHKC accessions (Table 1). Caffeine (41.83 ± 4.49 mg/g) was the

| Table 1 | Quantitative analysis of alkaloids, flavanols and methylated flavanols in tea samples by UPLC-PDA-TQ5 (x ± SD, n = 3, mg/g). |
|---------|-----------------------------------------------------------------------------------------------------------------|
| mg/g    | JHKC                                                                                                           | CK                                                                                      | JHKC-B                                                                                     |
|         | 11 JHKC individuals | Average | FDDB | YK10 | ZYQ | Average | JHKC-21 | JHKC-21-3 | JHKC-37-2 | Average |
| TB      | 6.38-13.11           |         | 9.52 ± 2.18* |     |     |         | 10.20 ± | 20.18 ± | 10.12 ±   |        | 13.50 ± 5.78   |
| CAF     | 35.28–49.06          |         | 41.83 ± 4.49 |     |     |         | 37.13 ± | 40.72 ± | 41.48 ±   |        | 39.78 ± 3.78   |
| TC      | ND-26.33             |         | 12.96 ± 5.23** |     |     |         | –       | –       | –         |        | 6.40 ± 11.09** |
| EC      | 0.65**               |         | 2.12 ± 4.53  |     |     |         | 4.87 ± 0.55 | 2.50 ± | 4.06 ± | 3.22 ±   | 3.26 ± 0.78* |
| EGG     | 11.62–15.41          |         | 13.68 ± 1.29** |     |     |         | 9.76 ± 0.36 | 9.35 ± | 10.80 ± | 12.13 ± | 10.76 ± 1.39 |
| EGGG’3Me| 0.44–1.50            |         | 1.02 ± 0.34  |     |     |         | 0.00 ± 0.01 | 0.65 ± | 0.71 ± | 0.53 ±   | 0.07 ± 0.08 |
| EGC     | 2.46–8.35            |         | 5.33 ± 1.9**  |     |     |         | 10.86 ± | 9.26 ± | 10.66 ± | 11.21 ± | 10.37 ± 1.00 |
| EGGG     | 54.16–72.03          |         | 64.09 ± 5.17 |     |     |         | 59.46 ± | 68.46 ± | 66.01 ± | 57.7 ± | 64.05 ± 5.63 |
| CGC     | 8.43–13.17           |         | 10.88 ± 1.68** |     |     |         | 1.11 ± 0.59 | 3.16 ± | 0.48 ± | 9.19 ±   | 4.27 ± 4.46 |
| C       | 0.09–1.54            |         | 0.58 ± 0.4**  |     |     |         | 0.56 ± 0.34 | 0.56 ± | 0.02 ± | 1.10 ±   | 0.03 ± 0.03 |
| GC      | 5.87–11.01           |         | 8.20 ± 1.39** |     |     |         | 2.10 ± 0.79 | 9.08 ± | 1.57 ± | 2.85 ±   | 4.50 ± 4.01 |
| GGC     | 15.75–19.65          |         | 17.73 ± 1.42** |     |     |         | 9.68 ± 0.95 | 10.25 ± | 6.64 ± | 5.76**   | 0.49 ± 0.26 |

Note: The data represent the mean from three replications. TB, theobromine; CAF, caffeine; TC, theacrine; ND, not detected. Compared to CK, *P < 0.05, **P < 0.01.
Fig. 5. Variations in differential metabolites in the JHKC (JHKC-1, JHKC-2, JHKC-3, JHKC-4, JHKC-5, JHKC-6, JHKC-7, JHKC-8, JHKC-9, JHKC-10 and JHKC-11) and CK (FDDB, YK10, and ZYQ) groups. JHKC-M represents the average level of metabolites across 11 JHKC individuals; CK-M represents the average level of metabolites across FDDB, YK10, and ZYQ. The Y axis represents the MS intensity of the compound detected by LC-MS. The data represent the mean ± SD. The green, blue, red and yellow columns represent JHKC, CK, JHKC-M and CK-M values, respectively. ND, not detected; *, P < 0.05; **, P < 0.01.
most abundant purine alkaloid, followed by theacrine (12.06 ± 5.23 mg/g) and theobromine (9.52 ± 2.18 mg/g). A recent study indicated that theacrine had beneficial effects on sedation, anti-inflammatory, anti-depression, sleep improvement, lipid metabolism improvement, anti-influenza, and so on (Sheng, et al., 2020). Among different tea plant species, theacrine was first detected and isolated from Yunnan Kucha tea [Camellia kucha (Chang et Wang) Chang] (Ye, et al., 1999) and has subsequently been reported to exist in Puan tea (Camellia sinensis var. puanensis Kurihara) (Li, et al., 2017), Dayaoshan tea (Camellia gymnogyna Chang) (Teng, et al., 2020), Buyatcha (Camellia gymnogyna Chang) (Jin, Jiang, Yao, & Chen, 2020), and Niedu Kucha (Camellia sinensis) (Wang, et al., 2020). In terms of its synthesis, theacrine is biosynthesized first through the conversion of caffeine to a 1,3,7-trimethyluric acid intermediate by a C8-oxidase, after which the intermediate is converted to theacrine by the N9-methyltransferase (CkTcS) (Zeng, Ye, Kato, & Crozier, 2002; Wang, et al., 2020; Zhang, et al., 2020). Among the 11 JHKC individuals evaluated in this study, 7 individuals contained theacrine, and the theacrine contents of the JHKC individuals ranged from 12.27 to 26.33 mg/g (Fig. 5C-D and Table 1), and 6 JHKC individuals contained theacrine levels > 15.0 mg/g. In contrast, 1,3,7-trimethyluric acid and theacrine were not detected in the CK cultivars, which may be due to the lack of C8-oxidase. Caffeine was the dominant purine alkaloid in the JHKC individuals, which was consistent with findings concerning Niedu Kucha (Camellia sinensis) (Wang, et al., 2020) but in contrast with findings concerning Yunnan Kucha [Camellia kucha (Chang et Wang) Chang], whose predominant purine alkaloid was theacrine (Yang, Ye, Xu, & Jiang, 2007; Qin, et al., 2020).

In addition, our study showed that the levels of caffeine and EGCG of the different tea groups were not significantly different (P > 0.05, Fig. 4B, E and Table 1). These results were consistent with those of a previous study (Yu, et al., 2020) that the concentration of a few metabolites (such as caffeine and EGCG) seemed to be stable in different tea cultivars. It needs to note that picking time and plant region have an influence on compound contents in fresh tea leaves (Ahmed, et al., 2019).

For an example, methylated flavanols were found to have > 2-fold contents in tea leaves collected in summer and autumn than in spring in our previous study (Dai, et al., 2015). In this study, 11 JHKC individuals were collected from a mountainous region at elevations ranging from 400 to 600 m in Jianghu County of Hunan Province in April. The aforementioned contents of characteristic compounds (e.g., theacrine, 1,3,7-trimethyluric acid, non-epi-form flavanols, and methylated flavanols) may vary in JHKC individuals collected from different seasons and plant regions.

3.3. Changes of differential metabolites in JHKC-B cultivars

To date, three cultivars have been bred from JHKC via agronomic selection, namely, JHKC-21–1 (CSA), JHKC-21–3 (CSA) and JHKC-37–2 (CSA). To study the metabolite changes in JHKC accessions after artificial breeding and cultivation, we analyzed the metabolites in the three JHKC-bred cultivars. As shown in Fig. S2, the PCA results suggested that the metabolite profiles of the JHKC-B cultivars were more similar to those of the CK cultivars than to those of the JHKC accessions. The changes in the accumulation of metabolites in the JHKC, CK (FDDB, YK10, and ZYQ) and JHKC-B (JHKC-21–1, JHKC-21–3 and JHKC-37–2) groups are shown in Fig. S3 and Table 1. As described above, compared with the CK cultivars, the JHKC accessions had significantly higher levels of mainly of 1,3,7-trimethyluric acid and theacrine purine alkaloids, non-epi-form flavanols (C, CG, GC, and GCG), and methylated flavanols (EGCG3′Me and EGCG3′Me) and lower levels of flavonol glycosides (kaempferol, quercetin and myricetin glycosides).

After artificial breeding and cultivation, the levels of caffeine and EGCG in the JHKC-B cultivars did not significantly differ from those in the JHKC cultivars (P > 0.05, Fig. S3-B, E and Table 1). Among the three JHKC-bred cultivars, 1,3,7-trimethyluric acid and theacrine metabolites were detected in the JHKC-37–2 cultivar, and the theacrine level was 19.21 ± 1.12 mg/g (Fig. S3-C, D and Table 1). It is possible that the mother plants of the JHKC-21–1 and JHKC-21–3 varieties may lack C8-oxidase or that C8-oxidase was no longer expressed after artificial breeding and cultivation, ultimately leading to naturally free of 1,3,7-trimethyluric acid and theacrine. EGCG3′Me and EGCG3′Me methylated flavanols were detected in JHKC-21–1, JHKC-21–3 and JHKC-37–2; however, their abundance was significantly lower than that in the JHKC group (P < 0.05, Fig. S3-F, H and Table 1). Similarly, non-epi-form flavanols of C (4.27 ± 4.46 mg/g), GC (0.03 ± 0.03 mg/g), CG (4.50 ± 4.01 mg/g), and GCG (6.64 ± 5.76 mg/g) also accumulated at low levels in the JHKC-B cultivars, as their contents decreased after artificial breeding and cultivation. With respect to flavonol glycosides, JHKC-21–1, JHKC-21–3 and JHKC-37–2 contained kaempferol 3-glucosylrutinoside, kaempferol 3-galactosylrutinoside and myricetin 3-galactoside, whereas these compounds were not detected in the JHKC group (Fig. S3-K, L and N). Moreover, JHKC-21–1 and JHKC-21–3 contained quercetin triglucoside (Fig. S3-P), and HKC-21–1 and JHKC-37–2 contained quercetin 3-glucosylrutinoside (Fig. S3-Q); however, these compounds were absent in the JHKC group. In addition, three dimeric flavanols (procyanidin B1, procyanidin B2 and procyanidin B3) accumulated to relatively high levels in JHKC-37–2 but low levels in JHKC-21–1 and JHKC-21–3 (Fig. S3-B, S and T). These results may have occurred because environmental factors can strongly influence the accumulation of metabolites in tea plant (Yu, et al., 2020).

4. Conclusions

This study qualitatively and quantitatively compared the metabolite profiles of wild JHKC germplasm with those of CK and JHKC-B cultivars via metabolomic analysis combined with chemometric analysis. The JHKC germplasm has a unique metabolite profile, and a total of 61 metabolites were demonstrated to differentially accumulated between JHKC and CK cultivars; these metabolites were mainly enriched in three critical differential metabolic pathways: caffeine metabolism, flavonoid biosynthesis, and flavone and flavonol biosynthesis. High accumulations of theacrine and 1,3,7-trimethyluric acid purine alkaloids, non-epi-form flavanols (C, CG, GC, and GCG), and EGCG3′Me and EGCG3′Me methylated flavanols and low expression of flavonol glycosides (kaempferol, quercetin and myricetin glycosides) were the most significant metabolite characteristics of JHKC germplasm. Taken together, these results indicated that JHKC germplasm is a promising resource for the breeding of novel tea cultivars that have high contents of theacrine, non-epi-form flavanols, and methylated flavanols.

CRediT authorship contribution statement

Wenliang Wu: Conceptualization, Methodology, Formal analysis, Investigation, Writing – original draft, Writing – review & editing, Funding acquisition. Mелиng Lu: Formal analysis. Jіkun Peng: Investigation. Hajіpeng Lv: Formal analysis. Шіang Shi: Formal analysis. Shuguang Zhang: Investigation. Zhen Liu: Investigation. Jіhua Duan: Investigation. Dan Chen: Resources. Weidong Dai: Conceptualization, Formal analysis, Resources, Writing – review & editing, Funding acquisition. Zі Lin: Conceptualization, Resources, Writing – review & editing, Project administration, Funding acquisition.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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