Research Article

Unraveling dynamic metabolomes underlying different maturation stages of berries harvested from *Panax ginseng*

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**A B S T R A C T**

Background: *Ginseng* berries (GBs) show temporal metabolic variations among different maturation stages, determining their organoleptic and functional properties.

Methods: We analyzed metabolic variations concomitant to five different maturation stages of GBs including immature green (IG), mature green (MG), partially red (PR), fully red (FR), and overmature red (OR) using mass spectrometry (MS)–based metabolomic profiling and multivariate analyses.

Results: The partial least squares discriminant analysis score plot based on gas chromatography–MS datasets highlighted metabolic disparity between preharvest (IG and MG) and harvest/postharvest (PR, FR, and OR) GB extracts along PLS1 (34.9%) with MG distinctly segregated across PLS2 (18.2%). Forty-three significantly discriminant primary metabolites were identified encompassing five developmental stages (variable importance in projection > 1.0, p < 0.05). Among them, most amino acids, organic acids, 5-C sugars, ethanolamines, purines, and palmitic acid were detected in preharvest GB extracts, whereas 6-C sugars, phenolic acid, and oleamide levels were distinctly higher during later maturation stages. Similarly, the partial least squares discriminant analysis based on liquid chromatography–MS datasets displayed preharvest and harvest/postharvest stages clustered across PLS1 (11.1 %); however, MG and PR were separated from IG, FR, and OR along PLS2 (56.8%). Overall, 24 secondary metabolites were observed significantly discriminant (variable importance in projection > 1.0, p < 0.05), with most displaying higher relative abundance during preharvest stages excluding ginsenosides Rg1 and Re. Furthermore, we observed strong positive correlations between total flavonoid and phenolic metabolite contents in GB extracts and antioxidant activity.

Conclusion: Comprehending the dynamic metabolic variations associated with GB maturation stages rationalize their optimal harvest time per se the related agroeconomic traits.

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1. Introduction

*Ginseng* is among the most popular medicinal herbs worldwide, and it is particularly valued in Asia and North America for its pharmacological activities, namely, anticancer, antiobesity, neuroprotective, and restorative activities [1,2]. The various pharmacologically active compounds in ginseng include ginsenosides, alkaloids, polysaccharides, polyacetylenes, and phenolic compounds [3]. However, the ginsenosides, which are structural glycosides with a dammarane skeleton, present in ginseng are primarily responsible for its pharmacological properties [4]. These compounds are distributed in many parts of the ginseng plant, including the root, stem, leaves, flowers, seeds, and fruits [5].

However, because of the disparate spatial distribution of ginsenosides in the ginseng plant, the pharmacological activities related to the different plant parts should vary substantially [6]. Previously, the antiaging, antiinflammatory, antioxidant, antiobesity, anticancer, hypoglycemic, and atopy-alleviating effects of the ginseng berry (GB) were reported [7–10]. Recently, we have shown that GB extracts possess considerably higher antioxidant potential than the roots [11]. These findings were congruent with the reportedly higher ginsenoside content in GB extracts than that in the other ginseng plant parts [12]. However, GBs are seldom considered superior, despite being rich in bioactive and functional compounds.

Mature GBs are typically harvested after 3–4 years of plant maturation depending on the variety [12]. Initially, a small berry...
develops which turns green and then red in the subsequent developmental stages. At the biомolecular level, the fruit maturation and ripening processes are genetically coordinated via a complex interplay of growth regulators, plant hormones, and dynamic metabolomes, influenced by numerous biological and environmental factors [13]. Metabolite alterations in plants play an instrumental role in fruit growth and maturation [14]. In addition, the overall repertoire of metabolites in fruits determines their organoleptic properties and functional components.

Essentially comprehensive and nonbiased, metabolomics has emerged as an important discipline for evaluating plant growth and development and involves the untargeted analysis of dynamically catabolized metabolic compounds [15]. In recent years, mass spectrometry (MS)—based metabolomic approaches for the demarcation of the stages of fruit growth and development have been adopted increasingly [16,17]. Enhancement of the sensitivity and precision of high-throughput techniques, such as gas chromatography (GC)—MS, liquid chromatography—mass spectrometry, and capillary electrophoresis—mass spectrometry, has enabled the rapid screening of complex metabolomes [18]. Previously, metabolomic profiling of ginseng plants has substantiated the spatial disparity in metabolite levels between the different plant parts including the roots [19,20]. However, the metabolic changes accompanying the different stages of GB growth and maturity remain largely unknown.

In this study, we used hyphenated MS-based metabolomic methods for delineating the dynamic metabolic events occurring during five different GB maturation stages. Furthermore, we established a correlation between the altered metabolomes and their effects on the functional bioactive compounds in ginseng.

2. Materials and methods

2.1. Chemicals and reagents

HPLC-grade solvents including methanol, acetonitrile, hexane, and water were purchased from Fisher Scientific (Pittsburgh, PA, USA). All standard compounds and analytical grade reagents used in this study were purchased from Sigma-Aldrich (St. Louis, MO, USA).

2.2. Plant materials

GBs in five different stages of development and maturity were harvested from 4-year-old *P. ginseng* plants cultivated in Geumsan County, Chungcheong Province, the Republic of Korea. Different staged berries were harvested at 14, 29, 45, 60, and 76 days after flowering. The five different time points at 15-day intervals were between June and August 2016 (from 03 June 2016 to 04 August 2016), representing the five different maturation stages, namely, immature green (IG): light green fruit collected on the 14th day, mature green (MG): dark green fruit collected on the 29th day, partially red (PR): light red fruit collected on the 45th day, fully red (FR): dark red fruit collected on the 60th day, and overmature (OR): deep brown fruit collected on the 76th day (Fig. 1). GBs are usually harvested when the pericarp color changes from green to red, and therefore, we further classified these five different maturation stages into three broader categories, namely, preharvest (IG and MG), harvest (PR), and postharvest (FR and OR) stages. The freshly harvested GBs were stored under deep-freezing conditions (−20°C) until further analyses. The GBs were freeze-dried for 3 days and homogenized in a blender for metabolite extraction.

2.3. Sample preparation

Each pulverized GB sample (600 mg) was extracted with 6 mL of 70% methanol using a Retsch MM400 mixer mill (Retsch GmbH, Haan, Germany), operated at 30 Hz/s for 10 min. Subsequently, the samples were subjected to sonication in an ultrasonic water bath (Power Sonic 305; Hwashin Technology Co., Seoul, Korea) for 5 min and centrifuged at 17000 rpm at 4°C for 15 min. The supernatant was filtered using a 0.2-μm polytetrafluoroethylene filter and concentrated using a speed vacuum concentrator (Modulpin 31; Biotron, Incheon, Korea). The samples collected finally were weighed and reconstituted in 70% methanol. The final concentration of the samples was 50 mg/mL for ultrahigh performance liquid chromatography—electrospray ionization—tandem mass spectrometry (UHPLC—ESI—MS/MS), ultraperformance liquid chromatography—quadrupole time-of-flight (UPLC—Q-TOF)—MS, and GC—TOF—MS analysis. The samples for GC—TOF—MS analysis were derivatized in a two-step reaction. First, 50 μL of methoxyamine hydrochloride (20 mg/mL in pyridine) was added to the dried samples and heated at 30°C for 90 min. Then, 50 μL of the derivatization agent MSTFA (N-Methyl-N-(trimethylsilyl)trifluoroacetamide) was added to the samples, and the samples were heated at 37°C for 30 min.

The pooled quality control samples were prepared from 50 μL blends of each sample. The analytical samples were analyzed in blocks of 10 runs followed by an intermittent quality control analysis procedure to ensure the data quality and robustness of the method. Three biological replicates and three analytical replicates of each GB sample extract were maintained. Similarly, the sample preparation for antioxidant activity assays involved the previously described extraction procedure.

2.4. GC—TOF—MS analysis

We used an Agilent 7890A GC system (Agilent Technologies, Palo Alto, CA, USA), equipped with an Agilent 7893 autosampler.

![Fig. 1. Different stages of ginseng berry (GB) development and maturation. The morphological changes shown here represent GBs harvested at different stages from the berry of four-year-old Panax ginseng Meyer. The five maturation stages were further subdivided into three broader categories depending on usual GB harvest time: preharvest (IG and MG), harvest (PR), and postharvest (FR and OR) stages. FR, fully red (60 days); IG, immature green (14 days); MG, mature green (29 days); OR, overmature red (76 days); PR, partially red (45 days).](image-url)
and a TOF Pegasus III mass spectrometer (LECO, St. Joseph, MI, USA) for GC–TOF–MS analysis. Metabolites were separated on an Agilent Rtx-5MS capillary column (J&W Scientific, Folsom, CA, USA) with an internal diameter, film thickness, and length of 0.25 mm, 0.25 μm, and 30 m, respectively. The derivatized samples (1 μL) were injected into the GC–TOF–MS system at a split ratio of 10:1 (v/v). Helium was used as the carrier gas at a constant flow rate of 1.5 mL/min. The injector temperature was maintained at 250°C, whereas the ion source temperature was set at 230°C. The oven temperature was held at 75°C for 2 min, increased to 300°C at a rate of 15°C/min, and then maintained at 300°C for 3 min. The mass acquisition rate was set at 20 scans/s for a mass scan range of 45–1000 m/z. The ionization energy for electron ionization was 70 eV.

2.5. UHPLC–ESI–MS/MS and UPLC–Q-TOF–MS analyses

An LTQ XL ion trap mass spectrometer (Thermo Fisher Scientific, San Jose, CA, USA) comprising an electrospray interface coupled with a Dionex UltiMate 3000 RS Pump, RS autosampler, RS column compartment, and RS diode array detector (Dionex Corporation, Sunnyvale, CA, USA) was used in this study. The sample, with an injection volume of 10 μL, at a constant flow rate of 0.3 mL/min, was separated on a Thermo Scientific SynchroNis C18 UHPLC column (100 mm × 2.1 mm i.d. × 1.7 µm [particle size]). The gradient mobile phase consisted of solvent A (water + 0.1% formic acid) and solvent B (acetonitrile + 0.1% formic acid). The LC gradient was increased from 10% solvent B to 100% solvent B in 15 min, maintained for 3 min, and then reequilibrated to the initial condition in 4 min. The photodiode array detector was tuned to a wavelength range of 200–600 nm for metabolite detection, managed by a three-dimensional field. The instrument was operated in a full-scan mode with a mass scan range of 150–1500 m/z. The operating parameters were as follows: capillary temperature was 270°C and sheath gas flow and auxiliary gas flow were 40 and 20 arbitrary units, respectively. The conditions in positive ion (and negative ion) mode for ESI were as follows: capillary voltage of 45 kV (–31 kV), source voltage of 5 V (4.5 V), and tube lens voltage of 120 V (–60 V).

UPLC was performed using a Waters Micromass Q-ToF Premier mass spectrometer with an ACQUITY UPLC system (Waters Corporation, Milford, MA, USA), equipped with a Waters ACQUITY UPLC tunable UV detector, autosampler, and binary solvent delivery system. The chromatographic operations were performed using an ACQUITY UPLC BEH C18 column (100 mm × 2.1 mm [i.d.] × 1.7 µm [particle size]) (Waters Corporation, Milford, MA, USA) at a flow rate of 0.3 mL/min. The mobile phase comprised water (A) and acetonitrile (B) with 0.1% formic acid (v/v), initially maintained at 5% B for 1 min. The mobile phase gradients were increased from 5% to 100% B over 10 min, maintained at 100% B for 1 min, reduced to 5% B over 2 min, and then, held constant for 1 min. The sample (5 μL) was injected at a constant flow rate of 0.3 mL/min. Mass spectra were recorded in full-spectrum mode over a range of 100–1500 m/z. The ion source and desolvation temperatures were set at 100°C and 200°C, respectively, and the desolvation gas flow rate was fixed at 700 L/h. The cone voltage was 40 V and capillary voltage was 2.8 kV in positive ion mode. In negative ion mode, the cone voltage was 60 V, whereas capillary voltage was 2.5 kV.

2.6. Data processing and statistical analysis

The MS raw data files were converted to Network Common Data Form (NetCDF) (*.cdf) format using ChromaTOF (version 4.44, LECO) and Xcalibur software (version 2.2; Thermo Fisher Scientific). After conversion, the NetCDF files were processed using the MetAlign software package (http://www.metalign.nl) to obtain baseline correction, peak alignment, peak detection, accurate masses, and normalized peak intensities [21]. Parameters of MetAlign were set according to the specific scaling requirements and chromatographic and mass spectrometric conditions used in the experiments (Table S1). The resulting data matrix, which contained the sample name and peak area information as variables, was processed using SIMCA-P+ 12.0 (Umetrics, Umeå, Sweden) for multivariate statistical analysis. The data sets were log-transformed and UV-scaled before principal component analysis (PCA) and partial least squares discriminant analysis (PLS-DA) modeling. PCA and PLS-DA were performed to compare the different maturation stages of GB. The significantly discriminant metabolite with a variable importance in projection (VIP) value exceeding 1.0 and p value < 0.05 was selected using the PLS-DA model. The p-values for different metabolite-based clusters were determined using Statistica (version 7.0; StatSoft, Tulsa, OK, USA). Variables were detected using MetAlign with the liquid chromatography–mass spectrometry data set.

PASW Statistics 18 (SPSS Inc., Chicago, IL, USA) was used to calculate Pearson’s correlation coefficient. The significance of data on total flavonoid content (TFC), total phenolic content (TPC), and antioxidant activity (2,2-azino-bis-(3-ethylbenzothiazoline-6-sulfonic acid) diammonium salt [ABTS] and ferric reducing antioxidant power [FRAP]) was determined using one-way analysis of variance and Duncan’s multiple range test using SPSS.

2.7. Identification and visualization of metabolites

The metabolites detected by GC–TOF–MS were putatively identified using standard compounds, an in-house library, and databases of the National Institute of Standards and Technology (NIST MS Search Program, version 2.0, Gaithersburg, MD, USA) by comparing their retention time and mass spectrometry data. Similarly, the metabolites analyzed by UHPLC–ESI–MS/MS were tentatively identified by comparing their retention time, molecular weight, ultraviolet absorption, and MS² fragment patterns based on standard compounds and published references. Accurate mass and elemental compositions were determined using MassLynx (Waters Corporation) in UPLC-Q-TOF-MS. Variations in the relative abundance of significantly discriminant metabolites among the maturation stages were visualized using a heat map generated using MultiExperiment Viewer (version 4.8.1, http://www.tm4.org/). MultiExperiment Viewer was additionally used to generate correlation heat maps between metabolite levels and antioxidant activity.

2.8. Evaluation of antioxidant activity by ABTS and FRAP assays

The methods of the ABTS free radical scavenging and FRAP assays were adopted from those developed by Lee et al [11]. All experiments were conducted for the three biological replicates of the GB extracts.

2.9. Determination of TFC and TPC

The TFC and TPC were estimated according to the method described by Lee et al [11]. All experiments were conducted for the three biological replicates of the GB extracts.

3. Results and discussion

3.1. GC–TOF–MS–based primary metabolite profiling for GB extracts

Primary metabolite profiles for the GB extracts across the five different developmental and maturation stages were characterized
Multivariate analyses of the aligned datasets indicated a clustered pattern for the extracts of GB under different stages in the PCA (Fig. S1A) and PLS-DA (Fig. 2A) models. Based on the PLS-DA score plot, the patterns of the primary metabolite profiles for the different maturation stages of GB were clustered into three main groups, namely, IG, MG, and three ripened (PR, FR, and OR) stages. The observed satisfaction values of X and Y variables in the PLS-DA model were 0.789 (R²X) and 0.989 (R²Y), respectively, with a prediction accuracy of 0.972 (Q²). Similar patterns were observed in the corresponding PCA (Fig. S1A). The preharvest (IG and MG) stages were separated from the harvest/postharvest (PR, FR, and OR) stages along PLS1 (34.9%), whereas MG was separated from the other four stages along PLS2 (18.2%). Totally, 43 significantly discriminant metabolites, including 16 amino acids and amines, 12 organic acids, 11 sugars and sugar derivatives, and four fatty acids with miscellaneous metabolites, were selected at a VIP value > 1.0 and p value < 0.05 using the PLS-DA model (Table S2). Moreover, the significantly discriminant metabolite variables were indicated in the corresponding loading plots (Fig. 2B), with their relative abundance in the extracts of GB under different stages displayed using the heat map (Fig. 2C).

Under the preharvest stages, the GB extracts exhibited a higher relative abundance of amino acids, specifically alanine (1), valine (2), isoleucine (4), proline (5), glycine (6), serine (7), threonine (8), aspartic acid (9), pyroglutamic acid (10), γ-Aminobutyric acid (GABA) (11), phenylalanine (13), glutamine (14), and lysine (15). Reportedly, increased amino acid metabolism in early maturation stages affects fruit development through enhanced respiration, endocarp hardening, phenylpropanoid precursor levels, and aroma compound synthesis [22,23]. Particularly, the higher abundance of serine in the aforementioned ground tissues (leaves, stems, and fruits) is mostly linked with higher rates of photorespiration [24].
However, the relative abundance of amino acids decreased during the later stages of GB growth and ripening (Fig. 2C). Ethanolamine (3) levels showed a tendency similar to that of amino acids during the early maturation stages. Biochemically, ethanolamines are synthesized through the decarboxylation of amino acids (mostly serine) and help to promote the overall growth and development of plants [25]. Similarly, the relative abundance of purine nucleosides, such as adenosine (42) and guanosine (43), were observed to be higher during the preharvest stages of GB development. Nucleosides serve as major energy carriers and the precursors toward the synthesis of nucleotide cofactors and subunits of nucleic acids [26]. Reports, high nucleoside levels affect high metabolic fluxes, promoting plant growth and fruit ripening [27]. Therefore, the relatively higher abundance of amino acids, amines, and nucleosides potentially determine higher growth and cell expansion rates, essentially in the early stages of GB development, i.e., IG and MG.

The contents of organic acids in fruits determine their characteristic sour tang, which masks the sweet taste attributed to sugars and certain amino acids [28]. In our study, we observed a distinctive stage-specific pattern of the abundance of organic acids, specifically succinic acid (17), glyceric acid (18), fumaric acid (19), malic acid (20), threonine acid (21), 2-hydroxythreonine acid (22), and quinic acid (27), in the GB extracts. In general, the relative organic acid levels increased linearly until the preharvest stages (IG and MG) and decreased sharply in the subsequent maturation stages. Generally, malic acid occurs as the most abundant organic acid in fruits, acting in conjunction with succinic acid and fumaric acid as the key intermediates of the Krebs cycle, potentially influencing the fruit respiration and ripening [29]. Contrarily, a relatively linear tendency of the levels of certain organic acids, including isobutaric acid (22), 4-hydroxybenzoic acid (24), acetic acid (25), citric acid (26), and chlorogenic acid (28), were observed during the later stages of GB maturation (Fig. 2C). The higher deposition of chlorogenic acid (~20%) was reportedly linked with the early-stage development of the endosperm and metabolic rerouting to lignin biosynthesis in the later stages concomitant to the temporal maturation of coffee seeds [30]. Hence, we conjectured that phenolic compounds are the specific metabolite biomarkers representing the harvest and postharvest stages of GB maturation (Fig. 2). Functionally, phenolic compounds are known to have potential pharmacological activities against chronic ailments, namely, diabetes, Alzheimer’s disease, cancer, and certain bacterial infections [31].

Fruit ripening is characterized by reduced pulp firmness, acidity, and chlorophyll content, coupled with the increased contents of total sugars and aroma volatile compounds [32]. In addition, a balanced quantitative ratio of sugar to organic acid levels is crucial for characterizing the flavor and acidity of fruit pulp [33]. Concerning fruit physiology, the production of sugars via photosynthesis and chlorophyll content, coupled with the increased contents of amino acids (mostly serine) and help to promote the overall growth and development of plants [25]. Similarly, the relative abundance of purine nucleosides, such as adenosine (42) and guanosine (43), were observed to be higher during the preharvest stages of GB development. Nucleosides serve as major energy carriers and the precursors toward the synthesis of nucleotide cofactors and subunits of nucleic acids [26]. Reports, high nucleoside levels affect high metabolic fluxes, promoting plant growth and fruit ripening [27]. Therefore, the relatively higher abundance of amino acids, amines, and nucleosides potentially determine higher growth and cell expansion rates, essentially in the early stages of GB development, i.e., IG and MG.

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Although a limited number of fatty acids and related compounds were detected in the extracts of GBs harvested at different stages, we observed a reciprocal pattern of the relative levels of fatty acids and their derivatives, including palmitic acid (40) reportedly accumulating at high levels in leaves and fruits. In olive fruits, unsaturated fatty acid (oleic and linoleic acid) levels increase during ripening, whereas those of saturated fatty acids generally decrease [39]. We observed a relatively higher abundance of oleic acid (41), an oleic acid derivative, during the later stages of GB maturation. Contrastingly, palmitic acid levels were higher during the preharvest stages and decreased subsequently. Generally, saturated or polyunsaturated fatty acids are synthesized during development, whereas monounsaturated fatty acids are mainly synthesized during fruit growth [40]. Our study revealed a similar pattern during GB maturation, although only two fatty acids were observed to be significantly discriminant.

3.2. UHPLC-ESI–MS/MS–based secondary metabolite profiling for GB extracts

Secondary metabolite profiling for the GB extracts representing the different maturation stages was performed by UHPLC–ESI–MS/MS, followed by multivariate analysis of the datasets. As shown in Fig. 3A, the secondary metabolite profiles of the GB extracts exhibited a clustered distribution pattern according to the maturation stage along PLS1 and PLS2, with an overall PLS-DA score plot data variability of 16.7% (PLS1: 11.1% and PLS2: 5.6%). The quality parameters for the PLS-DA model were verified with R²X = 0.257, R²Y = 0.993, and Q²Y = 0.849. The pattern of the secondary metabolite datasets in the PLS-DA model was congruent with the corresponding PCA score plot (Fig. S1B). The datasets for the preharvest stages (IG and MG) were separated from those for the harvest/postharvest stages (PR, FR, and OR) along PLS1, whereas MG and PR were separated from IG, FR, and OR along PLS2. Significantly discriminant secondary metabolites in the GB extracts under different maturation stages were selected at a VIP value > 1.0 and p value < 0.05 using the PLS-DA model. Overall, 24 metabolites, including a phenolic acid, two flavonoids, five notoginsenosides, six ginsenosides, three malonyl (Ma)-ginsenosides, and seven non-identified (NI) metabolites, were selected as significantly discriminant metabolites (Table S3).

The corresponding loading plot for the UHPLC–ESI–MS/MS datasets represented the distribution of the significantly discriminant metabolites along PLS1, signifying these metabolites for the GB maturation stages (Fig. 3B). Including ginsenosides Rg1 and Re, most of the ginsenosides were relatively abundant in the preharvest stages. As shown in Fig. 3C, the relative levels of notoginsenoside Fe (51), notoginsenoside Fd (52), and 20(S)-ginsenoside Rg3 (53) were higher in the stage IG. However, ginsenoside Rb1 (47), Ma-ginsenoside Rb1 (48), Ma-ginsenoside Re (49), ginsenoside Rd (50), notoginsenoside R1 (54), Ma-ginsenoside Re (57),
notoginsenoside R3 (58), ginsenoside Rf (59), and notoginsenoside R2 (60) were observed to be relatively abundant in the MG stage. Furthermore, the levels of ginsenoside Rg1 (55) and ginsenoside Re (56) increased gradually until the stage PR and decreased sharply thereafter. Ginsenosides are the active constituents of P. ginseng, distributed spatially in its leaves, roots, and berries [41]. The average levels and functional efficacies of ginsenosides, namely, antioxidant, antiinflammatory, antidiabetic, and anticancer components, increase optimally with the plant maturation stage [42,43]. We observed the increased relative abundance of ginsenosides up to the stage MG of GB maturation, followed by a sharp decrease in the subsequent developmental stages. Intriguingly, higher relative abundance of ginsenoside Rg and ginsenoside Rd was observed in the stage PR, which might have contributed to the variance between stages MG and PR along PLS2 in the PLS-DA plot (Fig. 3A). The biosynthesis and accumulation of secondary metabolites in plants is inextricably linked to the transcriptional regulation of genes vital for plant growth and fruit development [44,45]. In agreement with our study, change in fruit color from green to red during ripening results in the marked decrease in saponin levels in holly berries [46]. Similarly, high saponin levels, which reduced considerably on maturation, were detected in unripe Phytolacca dodecandra berries [47].

In addition to ginsenosides, the biological and pharmacological properties of P. ginseng have been attributed to its phenolic acid contents [48]. The high polyphenol levels in peach, potato, apple, grape, tomato, and ginseng are reported to be related to antitumor, antioxidant, antidiabetic, and cytotoxic activities [49,50]. In addition, ferulic acid, which is present in the plant cell wall as a phenolic phytochemical, has been studied for its antioxidant, antiallergenic, hepatoprotective, and anticarcinogenic activities [51]. Generally, fruit ripening involves a series of complex biochemical reactions resulting in the production of secondary metabolites, such as carotenoids, phenolic compounds, and volatile metabolites [52].
Hence, we proposed that kaempferol 3,7-diglucoside, kaempferol 3-sophoroside, and ferulic acid were synthesized as discriminant metabolites, with relatively higher abundance in the late stages of GB maturation (Fig. 3C). However, the temporal impact of numerous other important factors, including plant nutrition, climatic conditions, and water stress, should be comprehensively studied to understand the metabolic plasticity of GBs under varying maturation stages.

3.3. Analysis of correlation between metabolites in GB extracts and related biochemical phenotypes

We evaluated the antioxidant activities (ABTS and FRAP), TFC, and TPC of the GB extracts representing the different GB maturation stages. As shown in Fig. 4, the antioxidant activities of the extracts of GB in the harvest/postharvest (PR, FR, and OR) stages were relatively higher than those of the preharvest (IG and MG) samples. Although the variations among the late maturation stages were not significantly different, the antioxidant activity determined via the FRAP assay increased markedly at the stage OR compared with the stages PR and FR (Fig. 4B). The TFC and TPC increased linearly as the TFC and TPC activities increased until the late maturation stages. Previously, antioxidant levels in fruits were reported to increase considerably during late maturation stages owing to the increased synthesis and accumulation of antioxidant metabolites [53,54].

A correlation analysis was performed to evaluate the statistical relationship between, altogether, 67 significantly discriminant metabolites and observed tendencies of the related phenotypes, namely, ABTS, FRAP, TFC, and TPC. A correlation map showing both positive (red, $0 < r < 1$) and/or negative (blue, $-1 < r < 0$) correlations was constructed using color-plotted values of normalized variables (Fig. 5). Pearson’s correlation coefficients and $p$-values for all metabolites (60 identified and seven NI metabolites) and corresponding bioactivities are presented in Table S4. Overall, 24 metabolites exhibited positive correlations with the corresponding phenotypes, whereas the remaining 43 showed negative correlations. Interestingly, 29 metabolites with higher relative abundance in the preharvest stage, namely, pyroglutamic acid (10), γ-Aminobutyric acid (11), succinic acid (17), 2-hydroxyglutaric acid (23), glycero (29), arabinose (31), xylitol (32), ribitol (33), adenosine (42), guanosine (43), notoginsenoside Fe (51), notoginsenoside Fd (52), 20(S)-ginsenoside Rg1 (53), and three NI metabolites, showed significantly negative correlations ($p < 0.05$) with antioxidant activity. However, the metabolites abundant in the harvest/postharvest stages, specifically tryptophan (16), isobarbituric acid (22), 4-hydroxybenzoic acid (24), aceric acid (25), citric acid (26), chlorogenic acid (28), fructose (34), mannose (35), galactinol (39), oleamide (41), kaempferol 3-sophoroside (44), kaempferol 3,7-diglucoside (45), and one NI metabolite, showed significantly positive ($p < 0.05$) correlations with antioxidant activity. The metabolites exhibiting positive correlations with

![Fig. 4. Antioxidant activity tests. (A) 2,2-azino-bis-(3-ethylbenzothiazoline-6-sulfonic acid) diammonium salt (ABTS), (B) ferric reducing antioxidant power (FRAP), and (C) total flavonoid and (D) total phenolic contents of ginseng berry (GB) extracts from different maturation stages. Here, values are expressed as the average of three biological replicates ($n = 3$). Bar graphs denoted by the same letter were not significantly different, according to Duncan’s multiple range test ($p < 0.05$). FR, fully red; IG, immature green; MG, mature green; OR, overmature red; FR, partially red.](image-url)
antioxidant activity mainly included phenolic compounds with substantial antioxidant potentials [55]. Typically, 4-hydroxybenzoic acid, chlorogenic acid, kaempferol 3-sophoroside, and kaempferol 3,7-diglucoside are well-studied phenolic compounds in ginseng [48]. However, despite being a phenolic compound, ferulic acid showed weak positive correlations in our study. Previously, chlorogenic acid, kaempferol, and ferulic acid were mainly described as phenolic antioxidants in ginseng with reportedly higher abundance in the fruits [48].

The tendency of phenolic compound levels in the GB extracts increased linearly with maturation stages, especially in OR, characterized morphologically by the deep brown coloration of GBs. Fruit browning is an important biochemical phenomenon in floral-induced by increased polyphenol oxidase activity, causing fruits to soften and darken with deterioration of organoleptic properties [56]. However, this textural transformation negatively affects the flavor, aroma, and overall organoleptic properties of fruits [57]. Hence, optimal fruit maturity is an important agronomic trait as immature fruits lack desirable organoleptic qualities and over-mature ones have short shelf lives.

3.4. Scheme of metabolic pathways representing significantly discriminant metabolites among the GBs harvested at different maturation stages

In this study, we examined relevant metabolic pathway maps adopted from the Kyoto Encyclopedia of Genes and Genomes database for more than 60 significantly discriminant primary and secondary metabolites among GBs harvested at different maturation stages. Intriguingly, these metabolites were connected across 10 different pathways, including the citric acid cycle (TCA cycle); glycolysis; pathways of pyrimidine, amino acid, fatty acid, proline, sugar, and purine; ginsenoside metabolism; and shikimate-phenylpropanoid biosynthesis (Fig. 6).
Carbon metabolism primarily affects the bioenergetics of organisms and provides C skeletons for the synthesis of complex biomolecules regulating growth, development, and metabolism. In plants, C metabolic pathways, namely, the TCA cycle, glycolysis, sugar metabolism, and shikimate-phenylpropanoid biosynthesis, are chiefly related to plant growth and senescence and regulate plant physiological processes [58]. Changes in metabolite levels in different GB maturation stages were accompanied by an alteration in carbohydrate metabolism, especially glycolysis and the TCA cycle, which play central roles in determining the final metabolite composition in berries [59]. Sucrose and its interconversion into glucose and fructose, with numerous intermediates, feed the TCA cycle and glycolysis, generating energy molecules and reducing power sources, such as Adenosine triphosphate (ATP) and Nicotinamide adenine dinucleotide reduced (NADH) [60]. Together with amino acids and organic acids, these sugars constitute the energy source combining both photosynthesis and respiration in plants [61]. Glycolysis is the first step in the degradation of glucose to ATP, simultaneously generating precursors for the synthesis of organic acids, amino acids, anthocyanins, and numerous other secondary metabolites, including aroma compounds [62]. Reportedly, accumulated organic acids promote fruit ripening by affecting amino acid catabolism, essential for fruit maturation [63]. In our study, the preharvest stages, involving the transformation of GBs from IG to MG, include size increase and pericarp color changes (Fig. 1). These maturation stages showed high abundance of metabolite precursors for glycolysis and the TCA cycle.

Furthermore, shikimate-phenylpropanoid pathways provide alternative ways for aromatic compound synthesis, resulting in the biosynthesis of aromatic amino acids, including tyrosine, phenylalanine, and tryptophan [64]. In the preharvest GB extracts, we detected relatively higher phenylalanine levels, which decreased sharply in the later stages. Functionally, this generates several antioxidants (flavonoids, lignins, and phenols) and their precursors (aromatic amino acids and shikimic acid), which inhibit the activities of reactive oxygen species in plants and protect cell proteins, membrane lipids, and nucleic acids from ensuing damage [65]. The alkaloids and phenolic compounds derived from aromatic amino acids perform similar physiological functions.

The characteristic functional metabolites in GBs, i.e., ginsenosides, are a class of tetracyclic triterpenoid saponins, synthesized via dammarenediol-II hydroxylation by the Cyt P450 enzymes, followed by a glycosylation step catalyzed by glycosyltransferases [66]. Consequently, after the addition of monosaccharides to triterpene aglycones by glycosyltransferases, diverse ginsenosides are produced [67]. In our study, we detected 15 ginsenosides,
including protopanaxadiol and protopanaxatriol types, with most of them showing higher abundance during the preharvest stages. Saponin levels in plants are reportedly in including protopanaxadiol and protopanaxatriol types, with most economic traits and market values of GBs. Moreover, progressive increase in oxidative stress during fruit development due to reduced activities of antioxidant enzymes, namely, superoxide dismutase, lipoxygenase, and catalase, potentially favors antioxidant accumulation [69]. Hence, we inferred that the observed higher relative ginsenoside levels in the preharvest stages were related to the ripening and maturation of GB.

Moreover, we observed higher abundance of phenolic compounds derived from the shikimate-phenylpropanoid pathway in the harvest/postharvest (PR, FR, and OR)-staged extracts of GBs. These metabolites were verified to be significantly discriminant in the later stages and correlated positively with synchronous antioxidant activity. Considering metabolites to be more generic markers than the corresponding species-specific transcripts, we conjectured about the conserved metabolic pathways potentially influencing berry maturation across different ginseng species.

4. Conclusion

We assert that the metabolome of the GBs varies markedly at different maturation stages, accompanying its biomolecular, physiological, and morphological condition, which in turn regulates its organoleptic and functional properties. Notably, we demonstrated that C metabolism–related metabolites (organic acids and C– sugars) and most ginsenosides were abundant during the preharvest stages, i.e., IG and MG. However, antioxidant metabolites, i.e., flavonoids and phenolic compounds, accumulated at higher levels in the later harvest/postharvest (PR, FR and OR) stages of GB maturation. Therefore, we assume that detailed understanding of maturation stage–related metabolic tendency and related phenotypes of GBs would provide a metabolomic “snapshot” of the subtle transcriptomic–metabolomic networks governing the agro-economic traits and market values of GBs.

Conflicts of interest

The authors declare no conflict of interest.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.jgr.2019.02.002.

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