Metabolomics Analysis Reveals Potential Mechanisms in Bupleurum L. (Apiaceae) Induced by Three Levels of Nitrogen Fertilization

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Abstract: Bupleurum (Apiaceae) is widely used in traditional Chinese medicine to treat inflammatory and infectious diseases. Although roots are the only used parts in China, other countries use the whole plant. The yield and quality of Bupleurum depend mainly on fertilizers, especially nitrogen. The current study aimed to assess the relationship between the nitrogen fertilization level and the quality and metabolomic response of different parts (flowers, main shoots, lateral shoots and roots) of Bupleurum to three nitrogen fertilization levels (control group: 0 kg·ha\(^{-1}\); low-nitrogen group: 55 kg·ha\(^{-1}\); high-nitrogen group: 110 kg·ha\(^{-1}\)). The results showed that a high nitrogen level increases Bupleurum yield and quality parameters only in aerial parts, especially flowers, but has no significant effect on roots. The HPLC method was exploited for simultaneous quantification of three saikosaponins (A, C and D), which are the main bioactive components in the plant. It was found that the total content of saikosaponins decreased with high nitrogen fertilization in roots but significantly increased in flowers. Moreover, nitrogen fertilizer promoted the content of saikosaponin A but inhibited saikosaponins C and saikosaponins D in most parts of the plant. To study the response of primary metabolites, we adopted gas chromatography–mass spectrometry (GC–MS) analysis; 84 metabolites were identified that were mostly up-regulated with a high nitrogen level in flowers but down-regulated in roots. Four differential metabolites—D-fructose, lactose, ether and glycerol—were recognized as key metabolites in Bupleurum under nitrogen fertilization. Meanwhile, The Kyoto Encyclopedia of Genes and Genomes (KEGG) enrichment results explained that the impact of nitrogen fertilization on Bupleurum was attributed to the C-metabolism, N-metabolism, and lipids metabolism. This research put forward new insights into potential mechanisms and the relationship between the quality and yield of Bupleurum and nitrogen fertilization.

Keywords: Bupleurum L.; metabolite profiles; nitrogen application; GC–MS

1. Introduction

Among the essential plant nutrients, nitrogen (N), phosphorus (P) and potassium (K) are considered indispensable macronutrients for plant growth, development, and yield; thus, they play vital roles in many aspects of plant metabolism [1]. Nitrogen is the most important nutrient element for plant growth, physiology and the regulation of carbohydrates. It is a constituent of proteins, chlorophyll, alkaloids and amino acids [2]. In modern agriculture, nitrogen fertilizer is largely used to improve quality, increase yield and nourish plants. With the help of cell division, nitrogen fertilizer facilitates plant growth, protein percent, sucrose content and growth rate in sweet sorghum [3]. So far, lots of
research has been done on the absorption, assimilation and uptake of nitrogen fertilizer and its influence on plant development [4,5]. Therefore, proper nitrogen supplies are imperative to enhance yield and quality, as well as to promote plant viability under biotic [6] and abiotic stresses [7]. However, excessive application of fertilizer may disturb the nutrient balance, reduce the efficiency of fertilizer utilization and result in environmental pollution [8,9]. The quality and development of *Bupleurum* are restricted by nutritional requirements—too little or too many nutrients may restrain the growth and development. Furthermore, it is imperative to investigate the effect of nitrogen application on the metabolism of *Bupleurum* to guarantee the quality and yield.

*Bupleurum* L. is one of the largest genera in the Apiaceae family, with about 200 species all over the world. Forty-two species, including 20 endemics [10], are recorded in the flora of China. The dried roots of *Bupleurum scorzonerifolium* Wild. and *Bupleurum chinense* DC. are known as Radix Bupleuri (Chaihu), which is formally included in the *Chinese Pharmacopoeia* [11]. As traditional Chinese medicine, roots have great therapeutic effects on the treatment of common coughs, fevers and influenza hepatitis, malaria, menoxenia and so on [12–14]. *Bupleurum* L. are annual or perennial herbs, but excessive growth of *Bupleurum* extremely reduces the root biomass accumulation and secondary metabolite biosynthesis [15]. The biomasses of the aboveground part of *Bupleurum* account for about 38.1% of the whole plant, while the roots could not account for 20% [16]. Therefore, the aboveground parts of *Bupleurum* are used as anti-inflammatory drugs or topical antisepsics [17]; however, in the southeast area of China, all plant parts are used as medicines. To determine whether the aerial parts of *Bupleurum* could be used instead of roots or not, it is necessary to reveal the discrepancy in the types of metabolites among the different organs of *Bupleurum*.

Saikosaponins are the main active components of *Bupleurum* belonging to the pentacyclic triterpenoid oleanane-type compound. The biosynthesis of saikosaponins is mainly through the mevalonic acid (MVA) pathway in the cytoplasm and methylerythritol phosphate (MEP) pathway in the plastid [18]. The MVA pathway uses acetyl-CoA as the starting substrate and undergoes a six-step condensation reaction to generate IPP, and the MEP pathway uses pyruvate and glyceraldehyde 3-phosphate as the starting substrate to synthesize IPP through a seven-step reaction [19]. Among them, the MVA pathway plays a leading role in the biosynthesis of triterpene saponins [20–22]. These pathways in the early stages of biosynthesis provide a variety of precursors for secondary metabolites, for example, the aromatic amino acids tyrosine, tryptophan and phenylalanine. The exact link between primary metabolites and saikosaponins accumulation remains largely unknown, but it is closely related to pyruvate, acetyl-CoA and glyceraldehyde 3-phosphate. Due to the important role of *Bupleurum* as a medicinal compound, it is necessary to solve this problem. The regulation of saikosaponins and main compounds significantly affects the metabolic composition of the whole plant of *Bupleurum*.

Metabolomics have been proverbially used as a formidable method to analyze a lot of compounds from plant species, supplying a wide vision for the shoots to adjust metabolic processes [23,24]. Metabolomics include the quantitative, qualitative and dynamic research of the whole endogenous small molecules within organisms, organs, tissues and even cells under specific environmental conditions and at a specific time. Actually, many metabolomics studies have been conducted to comprehend the mechanism of a plant’s response to abiotic stress, for example, drought [25,26], salinity [27,28], heat [29], flooding [30], radiation [31–33], chilling [34], heavy metal toxicity [35,36] and combined multiple stresses [37–39]. Plant metabolomics have unique advantages in the identification and quality evaluation of Chinese medicinal materials. It can distinguish between different genes, different origins and even different growth cycles of medicinal plants [40,41]. Meanwhile, metabolomics research was conducted on biosynthetic pathways [42], signal transduction [43,44] and the ecological environment [45] of secondary metabolites of medicinal plants. This technique is gaining interest as a diagnosis tool for crop improvement and breeding [46]. Moreover, plant metabolomics integrate with genomes [47,48], transcri-
omes, proteomes and other omics in order to carry out quality control and germplasm improvement of medicinal plants effectively.

In order to analyze and compare the metabolite profiles of *Bupleurum* dependent on nitrogen fertilizer, we performed the technique of gas chromatography–mass spectrometry (GC–MS) to compare four different organs (roots, main shoots, lateral shoots and flowers) of *Bupleurum*. We aimed to: (1) evaluate the effect of different levels of nitrogen fertilization on yield and quality trait parameters of above and underground parts of *Bupleurum*; (2) reveal the effect of low and high nitrogen treatment on the total sakosaponins content and on the percentage of three saikosaponins in the different tissues of *Bupleurum*; (3) elicit changes in the metabolite profiles of *Bupleurum*’s different tissues caused by nitrogen fertilization; (4) compare metabolic changes to explain key metabolites involved in biosynthesis regulation.

2. Results

2.1. Comparative Analysis of Quality Traits Parameters among Different Nitrogen Fertilization Groups

In this experiment, the whole *Bupleurum* plants, growing under three levels of nitrogen fertilization (control nitrogen group (CN), low-nitrogen group (LN) and high-nitrogen group (HN)), were divided into four parts, including flower (F), main shoot (MS), lateral shoot (LS) and root (R) (Figure 1A). *Bupleurum* showed different biomass accumulation under nitrogen fertilization. At the LN level, shoots were thicker, the flowers were more flourishing and roots became slightly thicker. However, at the HN level, the increase in stem thickness and flowers flourishing was more significant than that at LN. Meanwhile, no significant increase in root thickness was observed at HN (Figure 1B). Figure 1C showed that the fresh weight of aboveground (flowers, main shoots and lateral shoots) and underground (roots) tissues in *Bupleurum* increased with application of the LN level. However, at HN levels, no significant increase in root fresh weight was observed, while the fresh weight of aerial parts, especially flowers, was extremely increased. Although there was no obvious discrepancy between HN and LN observed in root length, the total length was significantly higher at HN supply (Figure 1D), indicating that a high level of nitrogen fertilizer plays a greater role in promoting the growth of the aerial parts than roots in *Bupleurum*.

2.2. Total Saikosaponins Content Accumulated in Bupleurum Different Tissues under Three Levels of Nitrogen Fertilization

Initially, this study exploited an HPLC method for the simultaneous quantification of three saikosaponins: saikosaponin A, saikosaponin C and saikosaponin D. Simultaneously, we also measured the total content of saikosaponins in different *Bupleurum* tissues under the three levels of nitrogen fertilization. For the different nitrogen fertilization levels, the total saikosaponins display an analogous tendency and mostly accumulated in the flowers and the roots. In addition, the LN level decreased the total saikosaponins content in flowers and roots; however, at the HN level, flowers showed a significant increase in the total saikosaponins content, while the total content in roots remained stable (Figure 2A). When the total saikosaponins content was fixed, we compared saikosaponin A, saikosaponins C and saikosaponins D. We found that the percentage of saikosaponins A was lower than that of saikosaponins C and saikosaponins D without nitrogen. Nitrogen fertilizer promoted the content of saikosaponin A, and its increment was accompanied by a decrease in saikosaponins C in aerial parts (flowers, main shoots and lateral shoots); however, it was accompanied by a decrease in saikosaponins D in roots (Figure 2B).
Three nitrogen fertilization levels—CN: control nitrogen group, LN: low-nitrogen group, and HN: high-nitrogen group; four organs—R: root, MS: main shoots, LS: lateral shoots, F: flower, and W: whole plant. * and ** signify significant (p < 0.05) and extremely significant (p < 0.01), respectively.

Figure 1. Growth performances of Bupleurum at three nitrogen fertilization levels within four organs. (A) The whole Bupleurum was divided into four organs, including root, main shoots, lateral shoots and flower; (B) Appearances of Bupleurum growing under CN, LN and HN, respectively; (C) The four organs’ fresh weights (n = 3); (D) The length of the whole plant and the different parts (n = 3). Three nitrogen fertilization levels—CN: control nitrogen group, LN: low-nitrogen group, and HN: high-nitrogen group; four organs—R: root, MS: main shoots, LS: lateral shoots, F: flower, and W: whole plant. * and ** signify significant (p < 0.05) and extremely significant (p < 0.01), respectively.

Figure 2. Saikosaponins content at three nitrogen fertilization levels within four organs. (A) Total saikosaponins content, including saikosaponin A, saikosaponin C and saikosaponin D; (B) The percentage of saikosaponin A, saikosaponin C and saikosaponin D content. Three nitrogen fertilization levels: control nitrogen group (CN), low-nitrogen (LN) and high-nitrogen group (HN); four organs: flower (F), main shoots (MS), lateral shoots (LS) and root (R).
2.3. Overview of the Metabolites Profiles in Response to Three Levels of Nitrogen Fertilization

Under nitrogen fertilization, the metabolomic analysis of *Bupleurum* based on GC-MS resulted in the identification of 84 metabolites, which were divided into 9 classes, including 28 organic acids and derivatives, 16 sugars, 6 polyols, 8 amino acid and derivatives, 3 glycosides, 6 alkaloids, 7 lipids and derivatives, 9 alkyl and 1phenylpropanoid (Figure 3A, Table 1). To further clarify the difference between metabolites and accumulation patterns among the different plant organs under nitrogen fertilization levels, we adopt normalized numerical methods to accomplish cluster analysis and a heat map (Figure 3B). The heat map indicated that metabolites within some organs under the low-nitrogen fertilization level were up-regulated but down-regulated under the high-nitrogen fertilization level, suggesting that *Bupleurum* growing under the low-nitrogen fertilization level undergo significantly different metabolic processes compared with *Bupleurum* under the high-nitrogen fertilization level. Notably, the low-nitrogen fertilization level mostly caused an increased abundance of metabolites in flowers and roots. However, high nitrogen mostly increased the abundance of metabolites in flowers and reduced it in roots.

![Figure 3A](image)

**Figure 3.** (A) Classification of 84 metabolites based on *Bupleurum*. (B) The cluster analysis and heat map for 84 metabolites under different levels of nitrogen fertilization and in different plant parts. CN: control nitrogen; LN: low nitrogen; HN: high nitrogen; R: root; MS: main shoots; LS: lateral shoots; F: flower. The up-regulated and down-regulated metabolites are represented as yellow and blue, respectively. The green color means that abundance is 0, as shown in the upper right-hand bar.
| Classification            | Quantity | Metabolites                                                                 |
|---------------------------|----------|-----------------------------------------------------------------------------|
| Organic acids and derivatives | 28       | 1-Aminocyclopentanecarboxylic acid                                           |
|                           |          | 5-O-Feruloylquinic acid                                                    |
|                           |          | Benzoic acid                                                               |
|                           |          | Butanedioic acid                                                            |
|                           |          | Caffeic acid                                                               |
|                           |          | Chlorogenic acid                                                            |
|                           |          | Citric acid                                                                |
|                           |          | D-Glucic acid                                                              |
|                           |          | Gallic acid                                                                |
|                           |          | Glycolic acid                                                              |
|                           |          | Hydroxybenzoic acid                                                        |
|                           |          | Isophthalic acid                                                           |
|                           |          | Lactic Acid                                                                |
|                           |          | Malic acid                                                                 |
|                           |          | Oxalic acid                                                                |
|                           |          | Palmitic Acid                                                              |
|                           |          | Phenyllactic acid                                                           |
|                           |          | Phosphonic Acid                                                            |
|                           |          | Phthalic acid                                                              |
|                           |          | Pipoelic acid                                                              |
|                           |          | Propanedioic acid                                                          |
|                           |          | Propenoic acid                                                             |
|                           |          | Protocatechoic acid                                                        |
|                           |          | Quinic acid                                                                |
|                           |          | Succinic acid                                                              |
|                           |          | Sulfurous acid                                                             |
|                           |          | Triflorobenzoic acid                                                       |
| Sugars                    | 16       | Sucrose                                                                     |
|                           |          | Arabinofuranose                                                            |
|                           |          | Arabinose                                                                  |
|                           |          | D-Cellobiose                                                               |
|                           |          | D-Fructose                                                                 |
|                           |          | D-Glucose                                                                 |
|                           |          | D-Mannose                                                                  |
|                           |          | D-Xylose                                                                   |
|                           |          | Erythritol                                                                 |
|                           |          | Galactoseoxime                                                             |
|                           |          | Lactose                                                                    |
|                           |          | Levogluosan                                                                |
|                           |          | L-Rhamnosan                                                                |
|                           |          | Maltose                                                                    |
|                           |          | Sedoheptulose                                                              |
|                           |          | α-Mannobiose                                                               |
| Polyols                   | 6        | Benzylaminooctanol                                                         |
|                           |          | Benzenediol                                                                |
|                           |          | Cuminy1 alcohol                                                            |
|                           |          | Inositol                                                                   |
|                           |          | Muco-Inositol                                                              |
|                           |          | Ribitol                                                                    |
| Glycosides                | 3        | α-Lyoxofuranoside                                                          |
|                           |          | α-D-glucopyranoside                                                        |
|                           |          | L-Galactopyranoside                                                        |
Table 1. Cont.

| Classification          | Quantity | Metabolites                                      |
|-------------------------|----------|-------------------------------------------------|
| Alkyl                   | 9        | Cyclohexene                                     |
|                         |          | Decane                                          |
|                         |          | Disiloxane                                      |
|                         |          | Ether                                            |
|                         |          | Heptane                                         |
|                         |          | Nonane                                          |
|                         |          | Silane                                          |
|                         |          | Trisiloxane                                     |
|                         |          | Undecane has                                    |
| Phenylpropanoid         | 1        | Dihydroxybenzoate                                |
| Alkaloids               | 6        | Isoquinolinium                                  |
|                         |          | Ethanolamine                                    |
|                         |          | Copper phthalocyanine                           |
|                         |          | Heptabarbital                                   |
|                         |          | Benzopyran-4-one                                |
|                         |          | Carbamate                                       |
| Amino acid and derivatives | 8     | Alanine                                         |
|                         |          | Aminobutanoic acid                              |
|                         |          | Glycine                                         |
|                         |          | L-5-Oxoproline                                  |
|                         |          | L-Proline                                       |
|                         |          | Serine                                          |
|                         |          | Urea                                            |
|                         |          | L-Valine                                        |
| Lipids and derivatives  | 7        | Glyceric acid                                   |
|                         |          | Glycerol monostearate                           |
|                         |          | Stearic acid                                    |
|                         |          | Glycerol                                        |
|                         |          | Ribonic acid                                    |
|                         |          | Adenosine                                       |
|                         |          | 1-Monopalmitin                                  |

2.4. Identification of Differential Metabolites Relationships under Nitrogen Fertilization

In order to reveal the differential metabolites relationships under nitrogen fertilization, PLS-DA analysis was conducted on 84 metabolites. Our results revealed that the differences in the metabolomic profiles among the three levels of nitrogen fertilization groups were greater within roots and flowers of *Bupleurum* than that within the main shoots and lateral shoots. For flowers of *Bupleurum*, the samples of CN, LN as well as HN basically accomplished a better classification. The model prediction illustrating the gap between R²X and R²Y were not large, showing an excellent model (Figure 4A). According to the characteristic variables, differential metabolites in flowers were D-gluconic, lactose, glyceral, silane, l-rhamnose, phenyllactic acid, gallic acid, 5-o-feruloylquinic acid, phosphonic acid and isophthalic acid. PLS-DA of lateral shoots samples were distinctly segregated, and the model prediction results demonstrated a high-quality model (Figure 4B). The loading plot of lateral shoots demonstrated that lactose, 5-o-feruloylquinic acid, succinic acid, d-fructose, isoquinolinium, pipercolic acid, glycerol, phthalic acid, trisiloxane and D-xylose were differential metabolites. PLS-DA of the main shoots completely displays no separation, which indicted a bad model quality (Figure 4C). Propenoic acid, pipercolic acid, benzoic acid, isophthalic acid, caffeic acid, chlorogenic acid, copper phthalocyanine, urea, sucrose, and citric acid were the principal metabolites that contribute to dispersion for the main shoots. The score plots of roots metabolites displayed clear cluster trends (Figure 4D), with the red (HN) located between the green (CN) and blue (LN). Furthermore, the predictability of pattern goodness was noticed, demonstrating this model was evidently predictable and categorizable. D-xylose, α-mannobiose, lactose, silane, phthalic acid, ether,
propanedioic acid, aminobutanoic acid, citric acid and D-fructose contribute to different metabolites. Hence, we can say that various metabolites were potential biomarkers for different organs of *Bupleurum* growing under nitrogen fertilization. However, we need to further find out the key metabolites that affect the metabolism of *Bupleurum* under nitrogen fertilization.

Figure 4. PLS–DA score plot classifying the CN, LN and HN groups from metabolites of *Bupleurum* in flowers, lateral shoots, main shoots and roots (A–D), respectively. Different markers and colors represent different meanings: ● represents roots (R), ◆ represents flowers (F), ▲ represents lateral shoots (LS), ★ represents main shoots (MS); red, green and blue represent high nitrogen (HN) groups, low nitrogen (LN) and control nitrogen (CN), respectively.

2.5. Metabolite Profiling of Bupleurum under Nitrogen Fertilization in KEGG Enrichment Analysis and Volcanic Map

In order to seek out the key metabolites under different levels of nitrogen fertilization, metabolites with a fold change ≤ 0.5 and a *p*-value ≥ 1 were selected for further screening. The discrepancy in metabolite expression could be found clearly by volcano plots. Comparing CN with LN, there were 14 metabolites with an obvious change, including 11 up-regulated (3 F, 1 LS, 5 MS, 2 R) and 3 down-regulated metabolites in the roots (Figure 5A). Twenty-four metabolites had been changed when comparing HN to LN (Figure 5C), with 9 up-regulated (5 F, 4 R) and 15 down-regulated metabolites (4 F, 4 LS, 4 MS, 3 R). Compared with CN, 27 metabolites were significantly changed in HN group, including 24 up-regulated (14 F, 5 LS, 2 MS, 3 R) and 3 down-regulated metabolites in the root (Figure 5E).
Figure 5. (A–F) The volcanic map of differential metabolites and enrichment analysis of the Kyoto Encyclopedia of Genes and Genomes (KEGG). The green and red represent metabolism contents that were down–regulated and up–regulated significantly under nitrogen fertilization, respectively. High nitrogen (HN) groups, low nitrogen (LN) and control nitrogen (CN); ▼ represents roots; ▲ represents main shoots; ■ represents lateral shoots; ● represents flowers. The \( p \)-value means degree of enrichment; the smaller the \( p \)-value, the more significant enrichment degree. The size of the dot means the number of differential metabolites. (G) The Venn diagram of differential metabolites.
According to the KEGG annotation and enrichment results, the impact of different nitrogen fertilization levels (LN vs. CN, LN vs. HN, and HN vs. CN) on metabolites was mainly associated with the C-metabolism, N-metabolism and lipids metabolism. For LN vs. CN, metabolic pathways of differential metabolites were involved with the reductive carboxylate cycle (CO$_2$ fixation), glyoxylate and dicarboxylate metabolism, carbon fixation in photosynthetic organisms, pentose phosphate pathway, D-alanine metabolism and galactose metabolism (Figure 5B). For HN vs. LN, the difference of KEGG enrichment classification was also related on six metabolic pathways: insulin signaling pathway, reductive carboxylate cycle (CO$_2$ fixation), streptomycin biosynthesis, carbon fixation in photosynthetic organisms, glyoxylate and dicarboxylate metabolism, galactose metabolism and so on (Figure 5D). In HN vs. CN, various metabolites were mostly enriched in the glycerolipid metabolism, pentose phosphate pathway, starch and sucrose metabolism, galactose metabolism and the glyoxylate and dicarboxylate metabolism (Figure 5F).

2.6. Metabolic Network Diagram and Potential Metabolites in Bupleurum under Three Levels of Nitrogen Fertilization

The appraisal of potential metabolites involved into nitrogen stress might contribute to the growth and development of *Bupleurum*. We performed a Venn diagram (Figure 5G) to depict shared metabolites of different expressions among HN vs. CN, CN vs. LN and HN vs. CN. We found that the abundances of D-fructose, lactose, ether and glycerol had an obvious change in the response to nitrogen fertilization; therefore, these four overlapping metabolites could be regarded as key metabolites. Moreover, 1, 4 and 6 differential metabolites existed solely in (LN vs. CN) vs. (HN vs. LN), (LN vs. CN) vs. (HN vs. CN) and (HN vs. LN) vs. (HN vs. CN), respectively. Furthermore, on the basis of the KEGG annotation and enrichment data, significant metabolites were mapped to C-metabolism, N-metabolism, lipids metabolism pathways to guarantee clear changes in the metabolic regulation of nitrogen (Figure 6). The overlap of differentially expressed metabolites was observed, suggesting the partial similarity of mechanisms of *Bupleurum* in response to nitrogen fertilization. The metabolic network diagram went on to verify the above hypothesis. Responses to nitrogen fertilization of *Bupleurum* were dynamic and included intricate pathways. Therefore, nitrogen fertilization may activate some key physiological and metabolic activities that lead to the growth and development of *Bupleurum*.
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Figure 6. Changes of metabolites in Bupleurum under nitrogen stress are mapped to metabolic pathways. The yellow rectangle means that the content of metabolites is significantly up-regulated; the blue rectangle means that the content of metabolites is significantly down-regulated; the green rectangles mean that there is no significant difference in metabolites content. In each box, every row represents nitrogen treatment (upper row: CN; middle row: LN; lower row: HN), and every column means various organs (F: flower; LS: lateral shoots; MS: main shoots; R: roots).

3. Discussion

Bupleurum is an outstanding traditional medicinal plant in China and broadly used for treatment of inflammatory diseases. The yield and quality of Bupleurum mainly depend on the availability of fertilizers, such as nitrogen, phosphorus and potassium, especially on the availability of nitrogen. Thus, it is indispensable to probe the effect of different levels of nitrogen fertilization on Bupleurum to improve its cultivation level.

3.1. Influences of Nitrogen Fertilizer on Bupleurum Quality and Yield

For the common nutrient elements—nitrogen, phosphorus and potassium—nitrogen has the greatest effect under normal circumstances, followed by phosphorus and lastly potassium fertilizer [49–51]. Nitrogen is a component of proteins, nucleic acids and many cofactors, as well as of primary and secondary metabolites [44]. Nitrogen could also ensure plant growth and development, and therefore, its addition is needed to increase the plant yield.

Our results show that in the CN group, due to the lack of nitrogen, the shoots were weak, the leaves color turned yellow, and they fall off prematurely. The current study confirmed that plants could change the root system and leaf production to minimize
adverse effects under low nitrogen stress [52,53]. The HN fertilizer treatment significantly increased the biomass and the quality of *Bupleurum*; the shoots became thicker; the flowers were more flourish; the fresh weight and total length also increased. However, no significant effect happened to the roots. It is well documented that previous findings that high-nitrogen level treatment significantly increased plant height and ear length [54], as well as protein content and quality [55]. Additionally, Ottaiano [56] designed three nitrogen fertilization levels (N0, N30, N60), and it seemed that plants under the highest N dose showed a higher consumption of nitrogen, which is accumulated in leaves without increasing yield. In other words, high nitrogen stress promoted the accumulation of the aboveground parts, which had little effect on the yield of the underground parts. This was also consistent with our point. Saikosaponins represent the principle secondary metabolites within *Bupleurum* that have a wide range of pharmacological activities, such as their anti-inflammatory, antioxidant and hepatoprotective activity [12]. In *Bupleurum*, saikosaponins A and D are the most important bioactive constituents [57]. Our results showed that the total saikosaponins display an analogous tendency and mostly accumulated in the flowers and roots. In addition, the low-nitrogen level decreased the total saikosaponins content in flowers and roots; however, at the high-nitrogen level, flowers showed a significant increase in the total saikosaponins content, while the total content in roots remained stable. Moreover, high nitrogen fertilizer increased the content of saikosaponin A, and its increment was accompanied by a decrease in saikosaponins C in aerial parts (flowers, main shoots and lateral shoots); however, it was accompanied by a decrease in saikosaponins D in roots, indicating that high nitrogen promotes the quality of aboveground parts, especially flowers, but reduces the quality of roots. Our results were consistent with previous research that nitrogen fertilizer significantly increased the biomass and the content of saikosaponins A in *B. chinense* roots but had no significant effect on root saikosaponin D content [58]. In addition, we found saikosaponin D content decreased in roots, which suggests that saikosaponin D is less responsive to high nitrogen fertilization than saikosaponin A. In addition, increased root yield and total saikosaponin content in *B. falcatum* in response to nitrogen fertilizer was reported [59]. However, the application of moderate amounts of nitrogen fertilizer is best, as over fertilization resulted in decreased concentrations of medicinal compounds [60]. Either too little or too much nitrogen could be harmful to the growth and development. The fruit commodity rate of three nitrogen levels was highest in 5A and lowest in 3A [51]. Hence, appropriate nitrogen fertilizer rate can promote plant growth and maintain the content of active ingredients in specific parts.

3.2. Metabolic Profiling of *Bupleurum* Organs under Nitrogen Stress

PLS-DA is a multivariate statistical analysis method with supervised pattern recognition, which can distinguish groups and identify different metabolites to the greatest extent. Four groups were included in the PLS-DA: root, main shoots, lateral shoots and flower. The flowers and roots separated clearly between the N-treatments and control groups, suggesting that nitrogen fertilizer play a key role during *Bupleurum* growth and fruit maturation [61,62]. Previous studies have supported the above result that N greatly influences photosynthetic processes and grain growth during the maturation of wheat [63]. In addition, the differences in metabonomic characteristics between LN groups were greater than those in HN groups. Compared with the LN group, more negative correlations of metabolites were found in the HN group in roots, suggesting that the HN group caused metabolic changes that were not conducive to the development of *Bupleurum*. Qu [64] also came up with this conclusion; at higher nitrogen levels, the direct effects of stem–leaf ratio and leaf area index on dry matter yield are both negative. In these models, R2X and R2Y represented the interpretation rate to the X and Y matrices, respectively, and Q2 represented the prediction ability. According to PLS-DA analysis, Q2 > 0.2 and 0 < R2X-Q2Y < 0.2, suggesting there was an obvious difference between the N treatment group and the control group, and all results confirmed the model was meaningful. The clustering analysis findings were also consistent with PLS-DA. From this, we can guess that nitrogen
is an essential element for plant growth and development, and nitrogen deficiency can easily lead to metabolic disorders and environmental pollution \[51,65–67\].

3.3. KEGG Enrichment Analysis and Comprehensive Metabolic Pathways

The basis of the biological phenotype are metabolites, which facilitate a more intuitive and efficient understanding of biological processes and mechanisms \[68\]. This research was based on a non-targeted metabolic technique that detected 84 metabolites under nitrogen stress (CN, LN and HN). A common feature of nitrogen metabolism is that abundant nitrogen can significantly facilitate the biosynthesis and photosynthesis of proteins, amino acids and organic acids in plants \[69\]. The primary metabolism is widespread in plants, and the TCA cycle is most significant. Data analysis showed that the TCA cycle was extremely restrained under nitrogen stress. Compared with the CN group, the contents of citric acid, succinic acid and malic acid under nitrogen stress were lower (Figure 6). Moreover, the contents of citric acid and succinic acid in HN were much lower than CN and LN. Simultaneously, the reduced accumulation of malic acid was recorded under LN stress. Indeed, the HN group led to major changes in the content of carbohydrates in Bupleurum, which were mostly related to the glyoxylate and dicarboxylate metabolism and galactose metabolism. Furthermore, in the HN group, the increase in available carbon resources (for instance, glucose and sucrose) also clearly indicates a decrease in the metabolism of the TCA cycle (Figure 6). The research suggested HN prevented the utilization of the carbon skeleton by restraining the TCA cycle pathway \[70\]. In addition, high nitrogen might lead to an increase in nitrogenous compounds, organic acids and lipids \[71\], while limited nitrogen could promote the synthesis and accumulation of metabolites: flavones \[72\], phenolic \[73\] and total alkaloids \[74\].

Several metabolites were significantly up-regulated under high nitrogen stress for Bupleurum, which were mostly related to the C-metabolism and N-metabolism. In this research, some important sugars related to the C-metabolism were found, such as sucrose, mannose-6-P, fructose, glucose, ribose, galactose and sedoheptulose. The above-mentioned metabolites contributed to the metabolism and biosynthesis of Bupleurum, which further affect the yield of Bupleurum. The biosynthesis of saikosaponins \[18,75\] is mainly based on the mevalonic acid (MVA) pathway and methylerthyritol phosphate pathway (MEP). Furthermore, the starting points of the above metabolic pathway are pyruvate, acetyl-CoA and glyceraldehyde 3-phosphate \[18,19\], which are also related to fructose, fructose 6-phosphate and so on. Furthermore, it is noticeable that sucrose accumulated more in flowers than in other organs. These similar results support the validity. Sucrose is the main end-product of photosynthesis and the main form of carbohydrate transported over long distances in the phloem from photosynthetic source organs (leaves) to fruits, roots and shoot tips, which is extremely critical to the allocation of resources \[76,77\]. We predicted that sucrose was transported from the flower to the root as a signal molecule, and the essence may be due to the lack of nutrient nitrogen for root growth. In other words, more sucrose accumulation allowed better root growth. For most plants, carbohydrates mean the key energy storage \[78–80\]. Thus, high nitrogen stress can increase the utilization rate of sugar, and sugar is one of the most significant determining factors of yield and quality \[54,81\].

The TCA cycle could increase energy of amino acid synthesis \[82\] and thereby enhance the contents of protein and amino acid. Most amino acids are primarily related to nitrogen storage and utilization \[83,84\]. Amino acids are downstream products of nitrogen metabolism, and their abundance increases with elevated nitrogen levels \[85,86\]. Our results are consistent with previous studies; the levels of serine, glycine, alanine and valine shows a consistent pattern of change under high nitrogen. It is striking that proline and leucine had an extremely high content in LN instead of HN, which were inconsistent with other amino acids. In addition, in the N-metabolism, nitrogen usually accumulates larger in the main shoots and lateral shoots, and with the increase in N stress, it accumulates more. It is an active process of their absorption by the roots based on membrane
transport proteins \[87\]. We speculated that higher or lower nitrogen stress may lead to changes in amino acid biosynthesis flux to some extent \[88\]. There is a significant impact on amino acid metabolism for abiotic stress, particularly in some amino acid biosynthesis or degradation \[71\].

4. Conclusions

The findings of the current research highlighted the effect of nitrogen fertilizer on the traditional Chinese medicine *Bupleurum*. The LN level seems to be an ideal dose for the yield and quality of *Bupleurum*. The LN level significantly increased biomass accumulation and made *Bupleurum* more fruitful. However, the total saikosaponins decreased in the roots but increased significantly in the flowers under the HN level. Moreover, the non-targeted metabolomics analysis identified 84 kinds of metabolites of *Bupleurum* under nitrogen stress (CN, LN and HN). Under the HN level, most of the metabolites in flowers were up-regulated but down-regulated in roots. Therefore, the HN level promotes the quality and metabolites of flowers, while reducing those of roots. Because the root of *Bupleurum* (Apiaceae) is a Chinese herbal medicine and widely used to treat inflammatory and infectious diseases, we hope to accumulate more saikosaponins and metabolites in the root. In addition, four differentially accumulated metabolites—D-fructose, lactose, ether and glycerol—were identified as key metabolites under nitrogen stress of *Bupleurum*. The identified metabolites were mostly organic acids, amino acids, carbohydrates and lipids involved in carbon and nitrogen metabolisms, which set up a primary metabolic network diagram associated with *Bupleurum*. Nevertheless, it is necessary to further explore the relationship between saikosaponins and secondary metabolites in order to guide reasonable fertilization (nitrogen, phosphorus and potassium).

5. Materials and Methods

5.1. Plant Source and Experimental Design

The medicinal plant used in our study was cultivar *Bupleurum scorzonerifolium* Willd., which was widely cultivated in the northeast of China. The dried roots of *Bupleurum chinense* DC. And *Bupleurum scorzonerifolium* Willd. were the only two authentic sources of Chaihu \[11\]. Two-year-old *Bupleurum* seedlings were cultivated in natural environmental conditions at the research site of Lin Dian, Da Qing, Heilongjiang province, China (47°18′ N, 124°87′ E). The chemical prosperities of the soil \[89\] were pH 7.78; electrical conductivity, 134 us·cm\(^{-1}\); organic matter, 4.86 g·kg\(^{-1}\); total N 16.4 mg·kg\(^{-1}\); available N 2.1 mg·kg\(^{-1}\), respectively. In the 2018~2019 growing season, the annual average sunshine hours at the test site was 2807 h, the average annual temperature was 4 °C and the average annual precipitation was 417.2 mm \[90\].

The three nitrogen levels (N) were distributed in a randomized block design. Each treatment was replicated three times, and each plot measured 3 m × 1.2 m. Based on previous experience, urea (NH\(_2\))\(_2\)CO was used as the nitrogen fertilizer. *Bupleurum* were processed for CN, LN and HN treatments by adjusting the nitrogen concentration in the current study \[91–93\]. The three nitrogen fertilization levels were: CN (0 kg·ha\(^{-1}\)), LN (55 kg·ha\(^{-1}\)) and HN (110 kg·ha\(^{-1}\)).

On the 30th day (August 2018) after the second application of the nitrogen fertilizer, *Bupleurum* plants were harvested and divided into flowers, main shoots, lateral shoots and roots. Then, they were transferred quickly within three hours to the laboratory. Each group obtained three technical replicates, then the fresh weight and length were measured. Simultaneously, each sample is divided into two batches: one batch was dried in a forced air oven at 42 °C until constant weight and used for saikosaponins extraction (Section 5.2), and the other batch was stored at −80 °C for samples processing of GC–MS (Section 5.3).

5.2. Determination of the Saikosaponins Content

In total, 500 mg of the sample was dissolved in 25.0 mL of 8% NH\(_3\) methanol. The sample was sonicated at 30 °C for 30 min, filtered and evaporated, then diluted to a
10 mL volumetric flask. The solution was passed through a 0.45 µm filter membrane for HPLC analysis. The content of saikosaponins A, saikosaponins C and saikosaponins D were measured separately. A Hitachi HPLC system was equipped with an L-2000 high-performance liquid chromatograph and L-2200 autosampler, and a reversed phase column (Diamonsil C 18, 250 mm × 4.6 mm, 5 µm) was applied. Acetonitrile and methanol were used as solvents A and B, respectively, which was applied to the gradient elution as follows: 0–50 min, A: 25–90% and B: 75–10%; 50–55 min, A: 90% and B: 10%. The flow rate was maintained at 0.8 mL·min⁻¹, and the detection wavelength was set at 210 nm.

5.3. Sample Preparation and Extraction

In total, 60 mg of different organs of Bupleurum (roots, main shoots, lateral shoots and flowers) were moved to tubes. Each sample was extracted by the addition of 540 µL of methanol and 60 µL of the internal standard, followed by vortexing for 2 min and sonicating for 30 min. Then, 300 µL of chloroform and 600 µL of water were added, and the continued vortexing and sonicating continued. Subsequently, it was centrifuged at 14,000 rpm for 10 min. The supernatant was evaporated to dryness with a fast centrifugal concentrator. The dried residue was dissolved in 400 µL of methoxyamine pyridine solution and incubated at 37 °C for 90 min. Subsequently, 400 µL N,O–Bis (trimethylsilyl) trifluoroacetamide (BSTFA) and 60 µL Hexane were added, vortexed for 2 min and derivatized for 60 min at 70 °C. The solution was centrifuged at 12,000 rpm for 5 min to obtain the supernatant for GC–MS analysis.

5.4. GC–MS Analysis

In total, 1 µL of the derivatization solution was injected into the 7890A-5975C GC-MS of Agilent. The sample was separated using a non-polar DB-5 capillary column (length = 30 m, df = 0.25 µm, ID = 250 µm, I and W Scientific Folsom, CA, USA), and 1.0 mL·min⁻¹ high-purity helium gas was used as carrier gas. The temperature program started from 60 and then raised to 125 °C at 8 °C·min⁻¹ temperature ramps, 210 °C at 4 °C·min⁻¹ temperature ramps, 270 °C at 5 °C·min⁻¹ temperature ramps, 305 °C at 10 °C·min⁻¹ temperature ramps and a final maintenance at 305 °C. The electron impact ion source was maintained at 260 °C with a filament bias of −70 V. Full scan mode (m/z 50–600) was applied, with an acquisition rate of 20 spectrum·second⁻¹.

5.5. Statistical Analysis

GC–MS data were converted into Computable Document Format (CDF) and peak areas normalized to the internal stand. The metabolites were exhaustively contrasted by adopting a heat map and partial least squares discriminant analysis (PLS-DA) methods. In the Student’s t-test analysis, p-values of more than 1 and fold changes of less than 0.05 were statistically significant, and the volcano map was used to select the differentially expressed metabolites. The Kyoto Encyclopedia of Genome and Genome (KEGG) (http://www.genome.jp/kegg/) was used to analyze the metabolic pathways that had the greatest impact. The difference in the biomass and metabolites among CN, LN and HN were tested using a one-way analysis of variance (ANOVA). The graphs involved were drawn by GraphPad Prism 9.0.

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