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Comparative transcriptomic analysis reveals the regulatory effects of inorganic nitrogen on the biosynthesis of *Monascus* pigments and citrinin†

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Monascus spp. and its secondary metabolites have been widely applied in foods and medicines for thousands of years in eastern Asia. Nitrogen sources are essential nutrients for the growth and metabolism of *Monascus* spp. Our previous study found that inorganic nitrogen sources (especially NH₄Cl and NH₄NO₃) promoted the biosynthesis of *Monascus* pigments (MPs) and inhibited the production of citrinin. The objective of the present study was to investigate the regulatory mechanism of inorganic nitrogen on the biosynthesis of MPs and citrinin by the comparative transcriptional approach (RNA sequencing combined with RT-qPCR). Results indicated that the submerged fermentation of *M. purpureus* M3103 with NH₄Cl or NH₄NO₃ as the sole nitrogen source can significantly increase the yields of MPs (especially for *Monascus* orange and red pigments) and decrease citrinin production, compared with the organic nitrogen source (peptone group). Comparative transcriptomic profiling by RNA sequencing found that the numbers of differentially expressed genes (DEGs) between different experimental groups—M group (peptone group) vs. ML group (NH₄Cl group), and M group (peptone group) vs. MX group (NH₄NO₃ group), were 722 and 1287, respectively. Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway analysis revealed that genes involved in carbon and nitrogen metabolism, biosynthesis of amino acids were up-regulated by NH₄Cl and NH₄NO₃, which would produce more biosynthetic precursors for MPs. Whereas, the inorganic nitrogen source (both of NH₄Cl and NH₄NO₃) down-regulated the expression levels of genes involved in tyrosine metabolism. In addition, NR analysis indicated that the essential genes and transcription factors involved in the biosynthesis pathway of citrinin were down-regulated by NH₄Cl and NH₄NO₃. These results indicated that NH₄Cl or NH₄NO₃ as a nitrogen source for *M. purpureus* M3103 can significantly promote the precursor synthesis of *Monascus* pigments, but reduce the transcription of polyketide synthase for citrinin, and therefore significantly increase *Monascus* pigments production and decrease citrinin formation. These findings will facilitate a comprehensive understanding of the regulatory mechanisms of inorganic nitrogen in the biosynthesis of secondary metabolites in *M. purpureus*, and would benefit the application of *M. purpureus* in the production of MPs.

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1. Introduction

Red yeast rice is a kind of rice koji made from rice as the main raw material, and fermented by the filamentous fungus *Monascus*. It has been widely applied in traditional Chinese medicine, food additives, wine fermentation and other fields with a long history in eastern Asia.^{1,2} *Monascus* spp. can synthesize abundant useful secondary metabolites, such as pigments, dimerumic acid, γ-GABA, monacolin K, citrinin, and so on.³ *Monascus* pigments (MPs), a mixture of polyketones, are widely utilized as food colorants in various foods in China.⁴ Over the past few years, MPs have been reported to exhibit antimicrobial,⁵ hypolipidaemic,⁶ anti-atherosclerotic⁷ and photosensitizer activities.⁸ However, most strains of *Monascus* spp. also produce mycotoxin citrinin with strong nephrotoxic,



hepatotoxic⁹ and teratogenic¹⁰ effects on animals and humans, which greatly limits the application of *Monascus* fermented products. Therefore, the safety of *Monascus* fermented products has attracted much attention in many countries, especially Japan and European countries.¹¹ Thus, how to promote the biosynthesis of functional MPs but reduce the content of citrinin has become an urgent problem in the industrial production and applications of *Monascus* fermented products.

The regulation of the fermentation process is important and the most commonly used approach to improve the quality and safety of *Monascus* related products. In recent years, many researchers have tried to control the fermentation process to increase the production of MPs with low-yield or no citrinin, mainly through physical and chemical methods. The physical methods used to regulate the production of MPs and citrinin mainly focus on controlling oxygen supply,¹² temperature,¹³ pH¹⁴ and light¹⁵ during the fermentation process. Apparently, regulation of fermentation process through physical methods is difficult to block citrinin biosynthesis completely in *Monascus* spp. Regulation of fermentation process by chemical methods are mainly through the further supplementation of carbon and nitrogen sources, and other exogenous chemicals to the fermentation substrate.^{16–18} Over the past several decades, a considerable amount of studies have investigated the effects of different types of carbon and nitrogen sources on the production and secretion of secondary metabolites by *Monascus* spp. During the fermentation process, carbon source can form the carbon scaffolds of the secondary metabolites. It was found that mannitol had the best effect in enhancing the MPs production.¹⁹ By contrast, nitrogen source is essential for the biosynthesis of amino acids through the ammonia pathway and other nitrogen-containing organic compounds, such as *Monascus* red pigments (rubropunctamine [$C_{21}H_{23}NO_4$] and monascorubramine [$C_{23}H_{27}NO_4$]).

There are significant differences in the ability of utilizing inorganic and organic nitrogen sources for secondary metabolism by *Monascus* spp. Previous studies found that the addition of inorganic nitrogen source significantly increased the yields of *Monascus* red and yellow pigments, compared with the same amount of organic nitrogen source during the solid-state fermentation. Ammonium chloride (NH_4Cl) has been proved to be the most favorable for the biosynthesis of MPs, followed by ammonium nitrate (NH_4NO_3).^{20,21} Our previous experimental results also found that inorganic nitrogen (especially NH_4Cl and NH_4NO_3) not only significantly promoted the biosynthesis of MPs, but also significantly inhibited the production of citrinin. In recent years, the biosynthetic pathways involved in MPs and citrinin have been gradually revealed and understood.^{22,23} However, the regulatory mechanisms of inorganic nitrogen on the biosynthesis of pigments and citrinin in *Monascus* spp. have not been systematically investigated.

With the rapid development of high-throughput sequencing technology, comparative transcriptomics analysis based on RNA sequencing (RNA-seq) has been widely applied to investigate gene expression profiles under different types of conditions.^{24,25} In this study, RNA-seq was used to study the regulatory mechanisms of inorganic nitrogen on the biosynthesis of pigments

and citrinin in high pigment producing *Monascus purpureus* M3103. Besides, RT-qPCR was also used to detect the relative expression of MPs and citrinin related genes. The transcriptional levels of the three experimental groups were compared by transcriptome analysis. Results would enhance our knowledge of the molecular mechanisms behind the biosynthesis of secondary metabolites in *Monascus* spp.

2. Materials and methods

2.1 Strain and culture conditions

The high pigment producing strain *Monascus purpureus* M3103 (isolated from *Hongqu* provided by Lianjiang *Hongqu* Factory (Lianjiang county, Fujian Province, China)) could be obtained from the collection of Food Biotechnology Laboratory, Fujian Agriculture and Forestry University (Fuzhou, China). *M. purpureus* M3103 used in this study was obtained by strain screening from red yeast rice of Fujian Province in China. The strain was inoculated on potato dextrose agar (PDA) slant and incubated at 28 °C for 7 days, and then kept at 4 °C.

2.2 Inoculum preparation and fermentation mediums

To prepare the inoculum, a 5 day-old mycelium of *M. purpureus* M3103 on a PDA slant was washed by 10 mL dH₂O and then transferred to the fermentation medium by adjusting the initial pH to 5.00 and the concentration of suspension spores at about 1×10^5 spores per mL. The inoculum was incubated in a 500 mL Erlenmeyer flask containing 100 mL of medium for 7 days in a rotary shaker at 180 rpm under 30 °C. The fermentation mediums using peptone, NH_4Cl , and NH_4NO_3 as nitrogen sources were named as M, ML, and MX, respectively. After fermentation for 7 days, the harvested *Monascus* mycelia were immediately frozen in liquid nitrogen and stored at –80 °C until the RNA extraction. Fermentation mediums: M (peptone group): 5% soluble starch, 0.4% fishmeal peptone (Sinopharm Chemical Reagent Co., Ltd, China), 0.1% 7H₂O·MgSO₄, 0.25% KH₂PO₄, 100 mL distilled water; ML (NH_4Cl group): 5% soluble starch, 0.4% NH_4Cl (moles of nitrogen was 7.5 mmol), 0.1% 7H₂O·MgSO₄, 0.25% KH₂PO₄, 100 mL distilled water; MX (NH_4NO_3 group): 5% soluble starch, 0.4% NH_4NO_3 (moles of nitrogen was 10.0 mmol), 0.1% 7H₂O·MgSO₄, 0.25% KH₂PO₄, 100 mL distilled water. These mediums were sterilized at 121 °C for 20 min.

2.3 Determination of *Monascus* pigment and citrinin

The dry mycelium of *Monascus* (0.100–0.105 g) and the fermentation broth (0.1 mL) were transferred into a tube and extracted with 1 mL 70% ethanol in 200 W ultrasonic bath for 30 min at 50 °C, and then shaken on a rotary shaker at 200 rpm for 1.5 h. The supernatant was analyzed by ultraviolet spectrophotometer. The pigment concentration was measured by the corresponding absorbance (390, 465 and 510 nm for yellow, orange and red pigments, respectively). Subsequently, the supernatant liquid was filtered with a 0.22 μm pore size filter into a 1 mL vial to be used for citrinin determination by HPLC-FLD.^{26,27} The excitation and emission wavelengths were 331 nm



and 500 nm, respectively. ZORBAX SB-C18 column (150 mm \times 4.6 mm, 5 μ m; Agilent, USA) was used at 30 °C, and isocratic elution was performed for 20 min using acetonitrile : water (1 : 1, v/v; adjusted pH to 2.5 by acetic acid) as the mobile phase at 1 mL min $^{-1}$.²⁷

2.4 UPLC-Orbitrap MS analysis of the composition of *Monascus* pigments

The main components of MPs were identified using a water UPLC system (Waters, MA, USA) and a Thermo Scientific TM Orbitrap mass spectrometer (Thermo, MA, USA). The mobile phase A: water (0.1% formic acid), and the mobile phase B: methanol (0.1% formic acid). A T3 column (2.1 mm \times 100 mm, 3 μ m) (Waters, USA) was kept at 30 °C. The gradient elution: 98% A–90% A at 0–0.5 min, 99% A–37.5% A at 0–0.5 min, 37.5% A–16.8% A at 0.5–3.5 min, 16.8% A–7% A at 3.5–10.5 min, 7% A–0.1% A at 10.5–13 min, and 0.1% A at 13–14 min; finally, equilibrated with 99% A for 3 min. The flow rate: 0.40 mL min $^{-1}$, and the injection volume: 5 μ L. The mass spectrometry detection: positive ion electrospray mode. The scan range: 100 to 1000 m/z , and the source temperature: 120 °C. The capillary voltage: 3.2 kV at ESI $^+$. The collision energy was 15 to 40 eV for the energy scan.

2.5 RNA extraction and of high-throughput sequencing

Total RNA was extracted by using Spin Column Fungal Total RNA Purification Kit (Sangon Biotech, code no. B518659). Measurement of the concentration and integrity of the final RNA samples were referenced in ref. 28. The total RNA of samples were used for cDNA library construction and high-throughput sequencing at Biomarker Technologies Corporation (Beijing, China) on the Illumina HiSeq $^{\text{TM}}$ 2500 sequencing platform (Illumina $^{\text{TM}}$, San Diego, CA, USA) with 150 bp paired-end reads.

2.6 Sequence reads assembly, mapping and annotation

Raw data (raw reads) were provided in FASTQ format. The clean reads were obtained by removing reads containing adapter, reads containing ploy-N and low quality reads from raw data. The clean reads were mapped to *M. purpureus* YY-1 genomic sequence (NCBI accession number: samn08978766) using TopHat (<http://ccb.jhu.edu/software/tophat/>). Q20, Q30, GC-content and sequence duplication level of the clean data were also calculated. The transcript or gene expression levels were estimated by fragments per kilobase of transcript per million fragments mapped.²⁹ The sequencing data were deposited to the Sequence Read Archive (SRA) database at the National Center for Biotechnology Information (NCBI) with accession number SRR8380849 (ML group), SRR8380850 (MX group) and SRR8380851 (M group).

2.7 Quantification and differential expression analysis of transcripts

Differential expression analysis of two groups was performed using the DESeq R package (1.10.1). DESeq provided statistical

routes for determining differential expression in digital gene expression data using a model based on the negative binomial distribution. The resulting *P* values were adjusted using the Benjamini and Hochberg's approach for controlling the false discovery rate. Genes with an adjusted *P*-value < 0.01 found by DESeq were assigned as differentially expressed.

2.8 GO and KEGG enrichment analysis of differentially expressed genes (DEGs)

Gene Ontology (GO) enrichment analysis of the DEGs was implemented by the GOseq R packages based Wallenius non-central hyper-geometric distribution,³⁰ which can adjust for gene length bias in DEGs. KEGG³¹ is a database resource for understanding high-level functions and utilities of the biological system, such as the cell, the organism and the ecosystem, from molecular-level information, especially large-scale molecular datasets generated by genome sequencing and other high-throughput experimental technologies (<http://www.genome.jp/kegg/>). KOBAS software was used to test the statistical enrichment of differential expression genes in KEGG pathways.³²

2.9 Expression level of pigments and citrinin biosynthesis-related genes by RT-qPCR

Quantitative real-time reverse transcription polymerase chain reaction (RT-qPCR) analysis of pigments biosynthetic gene and citrinin biosynthetic gene. cDNA synthesis was performed using the PrimeScript $^{\text{TM}}$ RT reagent kit with gDNA Eraser (Takara, Dalian, China). Real-time qPCR was conducted to monitor gene expression levels using the SYBR $^{\text{®}}$ Premix Ex Taq $^{\text{TM}}$ II (Takara, Dalian, China). All primers used in this study were listed in Table S1.† RT-qPCR was performed in duplicate using AB7300 Real-Time PCR System (Applied Biosystems, USA). The relative levels of target mRNAs were determined using the $2^{-\Delta\Delta C_t}$ method and were normalized to the β -actin mRNA signals in each sample.

2.10 Statistical analysis

All results are expressed as the means \pm standard deviation (SD) ($n = 6$). GraphPad Prism 7 was used for all analyses. The statistical significance was determined by one-way analysis of variance (ANOVA) with Duncan's multiple test.

3. Results

3.1 Effects of inorganic nitrogen on the growth and the production of MPs and citrinin

After 7 days of cultivation, there are significant difference in *Monascus* biomass among the 3 groups. The dried cell weight concentration was 1.49%, 1.20% and 1.11% in M, ML and MX group, respectively (Fig. 1A). The pH values of the final products of M, ML and MX groups were 4.09, 2.07 and 3.68, respectively (Fig. 1B). The effects of inorganic nitrogen (NH_4Cl or NH_4NO_3) on the yields of MPs and citrinin in fermentation of *Monascus* were evaluated. The relative yields of MPs and citrinin were shown in Fig. 1. The intracellular MPs production by NH_4Cl and NH_4NO_3 were nearly 87.6% and 69.2% higher than the M group

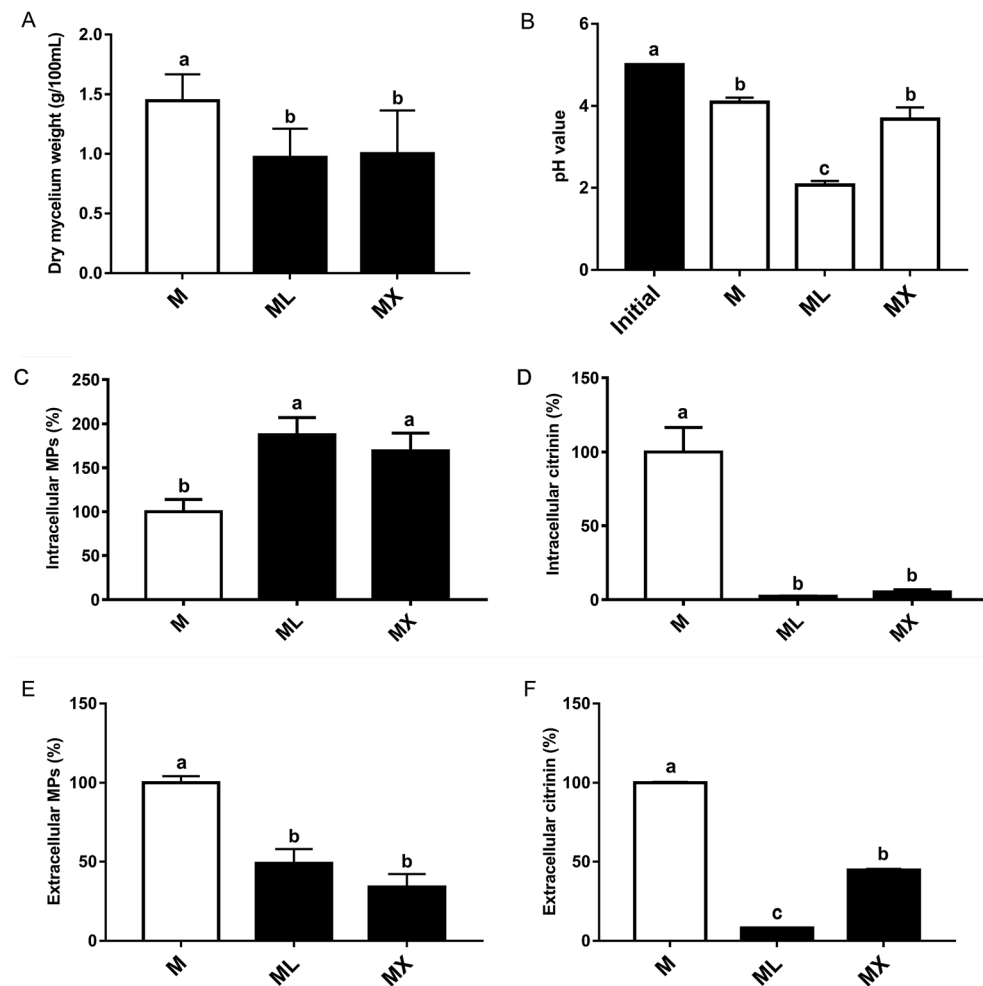


Fig. 1 Effects of inorganic nitrogen sources (NH_4Cl and NH_4NO_3) on the growth and production of *Monascus* pigments (MPs) and citrinin in *M. purpureus* M3103. (A) Dry mycelium weight; (B) initial and final pH value; (C) relative yield of intracellular MPs (%); (D) relative yield of intracellular citrinin; (E) relative yield of extracellular pigments; (F) relative yield of extracellular citrinin.

($P < 0.05$), respectively (Fig. 1C). While the extracellular pigments by NH_4Cl and NH_4NO_3 were nearly 51.1% and 66.2% less than the M group ($P < 0.01$), respectively (Fig. 1E). Surprisingly, the yields of intracellular citrinin by NH_4Cl and NH_4NO_3 were significantly lower than the peptone group ($P < 0.01$) (Fig. 1D). Which was consistent with the result of extracellular citrinin production (Fig. 1F). Thus, the above findings indicated that inorganic nitrogen source (NH_4Cl and NH_4NO_3) can significantly promote the biosynthesis of MPs and inhibited the production of citrinin in the submerged fermentation of *Monascus*, which would greatly reduce the risk of *Monascus*-related products and would provide a foundation for the study on the regulatory mechanisms of the biosynthesis of pigments and citrinin in *M. purpureus*.

3.2 Effects of inorganic nitrogen on the chemical composition of MPs

In order to better understand the effects of inorganic nitrogen sources (NH_4Cl and NH_4NO_3) on the composition of MPs, UPLC-MS/MS was used to detect the differences of the composition of MPs

among the M, ML and MX groups. As shown in Fig. 2, S1 and Table S2,† a total of 8 different MPs components were detected, including four yellow pigments (monasfluore A, monascin, monasfluore B and ankaflavin), two orange pigments (rubropunctatin and monascorubrmine) and two red pigments (rubropunctamine and monascorubrmine). Compared with other pigment components, the peak of rubropunctatin (P8) is not obvious in the total ion chromatogram due to its low relative abundance in total pigment (Fig. 2). Therefore, the specific ion of rubropunctatin (m/z 355.1572) was selectively extracted from the total ion chromatogram to eliminate the interference of other pigments with high abundance.

The pigment composition in the fermentation products of ML group and MX group was similar (Fig. 2B and C). It is worth noting that the production of rubropunctamine and monascorubrmine (two red pigments) were significantly increased by inorganic nitrogen sources (NH_4Cl and NH_4NO_3). These findings indicated that inorganic nitrogen sources (NH_4Cl and NH_4NO_3) could significantly increase the yields of *Monascus* red pigments (mainly rubropunctamine and monascorubrmine) and change

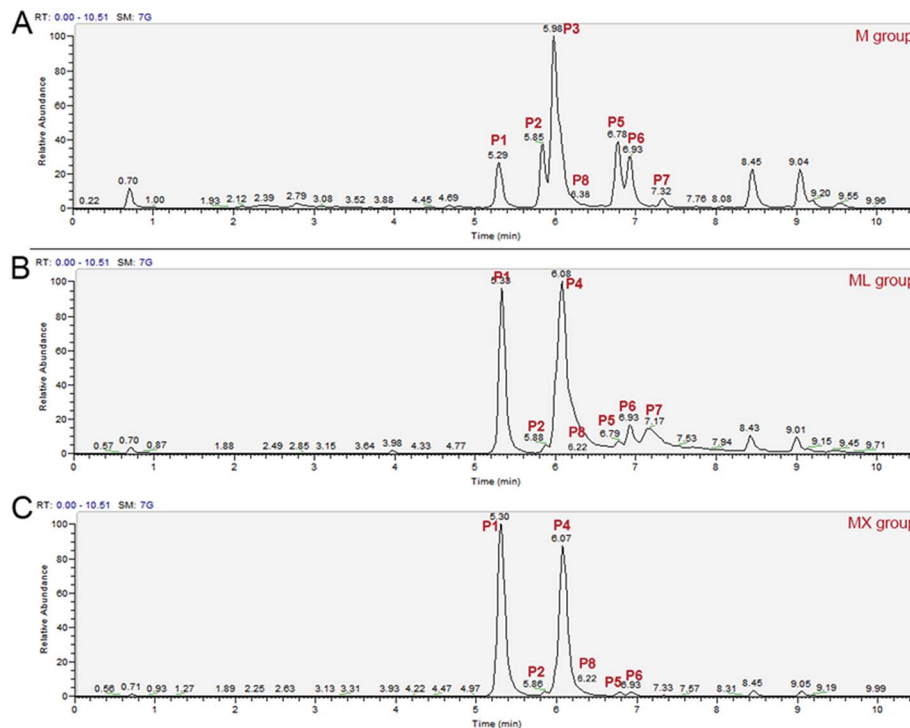


Fig. 2 Effects of inorganic nitrogens (NH_4Cl and NH_4NO_3) on the chemical composition of MPs. (A) M group; (B) ML group; (C) MX group. Note: P1 – rubropunctamine; P2 – monasfluore A; P3 – monascin; P4 – monascorubramine; P5 – monasfluore B; P6 – ankaflavin; P7 – monascorubrin; P8 – rubropunctatin.

the composition of MPs compared with organic nitrogen (peptone).

3.3 High-throughput sequencing and *de novo* assembly

To better explore the regulatory mechanisms of inorganic nitrogens (NH_4Cl and NH_4NO_3) on the biosynthetic pathways of pigments and citrinin in *M. purpureus*, high-throughput sequencing was used to investigate the transcription. A total of 222.98 million clean readings were obtained from the Illumina high-throughput sequencing platform after quality control and data filtering. The GC% of the sequencing data from 9 samples was 42.56%, and the average quality score more than 30 reading percentages was not less than 92.31%, indicating that the accuracy and quality of the sequencing data were sufficient for further analysis. General sequencing statistics were shown in ESI Table S3.† From the comparison results, the alignment efficiency of reads and reference genomes of 9 samples ranged from 86.27% to 91.17%, as shown in Table 1. The Pearson correlation coefficient was used as an indicator of biological relevance.³³ The strong correlation ($R^2 > 0.95$) between biological replicates indicated that the samples from three different groups had good repeatability (ESI Fig. S2†).

3.4 Analysis of differentially expressed genes (DEGs)

In order to reveal the regulatory mechanisms inorganic nitrogens (NH_4Cl and NH_4NO_3) on the secondary metabolism of *M. purpureus*, comparative transcriptomic analysis was carried out.

The fragments per kilobase of exon model per million mapped reads (FPKM) were calculated as the normalized expression values of each annotated gene. Venn diagram illustrated that 722, 1287 and 523 DEGs were found between the two sample groups: “M vs. ML”, “M vs. MX” and “ML vs. MX”, respectively (ESI Fig. S3†). The number of unique DEGs belonging to “M vs. MX” group (506) was much greater than the number belonging to “M vs. ML” group (145) and “ML vs. MX” group (92). The differences in gene expression levels and statistical significance between the two sample groups (“M vs. ML” and “M vs. MX”) were shown in the volcano plots, respectively (Fig. 3A and B). The number of up-regulated and down-regulated DEGs in “M vs. ML” groups were 401 and 321, respectively (Fig. 3A), while the DEGs in “M vs. MX” groups were 823 up-regulated and 464 down-regulated (Fig. 3B). Hierarchical clustering analysis of the screened differentially expressed genes was also presented in Fig. 3C.

3.5 Gene annotation and GO functional classification

Annotated numbers of the DEGs by different functional databases were shown in Table 2. A total of 1892 DEGs were annotated by eight frequently used functional databases, including Clusters of Orthologous Groups (COG), Gene Ontology (GO), Kyoto Encyclopedia of Genes and Genomes (KEGG), Eukaryotic Orthologous Groups (KOG), nonredundant (Nr), Pfam, Swiss-Prot and evolutionary genealogy of genes: Nonsupervised Orthologous Groups (eggNOG). In order to further evaluate the



Table 1 The results of sequence alignment of sample sequencing data and selected reference genomes

Samples	Total reads	Mapped reads	Uniq. mapped reads	Multiple mapped reads	Reads map to '+'	Reads map to '-'
ML1	47 452 392	42 087 728 (88.69%)	40 402 206 (85.14%)	1 685 522 (3.55%)	20 944 939 (44.14%)	21 029 533 (44.32%)
ML2	53 871 824	47 670 074 (88.49%)	45 781 220 (84.98%)	1 888 854 (3.51%)	23 726 820 (44.04%)	23 799 139 (44.18%)
ML3	43 230 106	37 294 124 (86.27%)	35 845 197 (82.92%)	1 448 927 (3.35%)	18 529 453 (42.86%)	18 649 921 (43.14%)
MX1	51 126 592	46 609 669 (91.17%)	44 905 008 (87.83%)	1 704 661 (3.33%)	23 202 380 (45.38%)	23 297 756 (45.57%)
MX2	49 137 078	44 565 319 (90.70%)	42 961 449 (87.43%)	1 603 870 (3.26%)	22 191 415 (45.16%)	22 267 643 (45.32%)
MX3	52 234 716	47 158 784 (90.28%)	45 456 437 (87.02%)	1 702 347 (3.26%)	23 451 892 (44.90%)	23 579 692 (45.14%)
M1	52 707 596	46 529 510 (88.28%)	44 936 750 (85.26%)	1 592 760 (3.02%)	23 142 723 (43.91%)	23 257 067 (44.12%)
M2	44 711 892	39 734 351 (88.87%)	38 332 129 (85.73%)	1 402 222 (3.14%)	19 770 883 (44.22%)	19 832 956 (44.36%)
M3	56 123 470	49 317 018 (87.87%)	47 531 324 (84.69%)	1 785 694 (3.18%)	24 544 156 (43.73%)	24 635 313 (43.89%)

RNA-seq data, KOG was used to classify DEGs. The DEGs annotated by KOG were divided into 25 functional categories (Fig. 4A and B). 19.78% (72 genes) and 18.91% (122 genes) DEGs in “M vs. ML” and “M vs. MX” groups were annotated as “General Functional Prediction Only”, which was the most frequently annotated functional category. “Amino acid transport and metabolism” was the second frequently annotated, accounting for 14.01% (51 genes) DEGs in “M vs. ML” and 12.56% (81 genes) DEGs in “M vs. MX”. “Lipid transport and metabolism” was the third and sixth frequently annotated functional category in “M vs. ML” (9.62%, 35 genes) and M vs. MX” (6.20%, 40 genes), respectively. “Carbohydrate transport and metabolism” was the fourth frequently annotated functional category with 9.34% (34 genes) DEGs in “M vs. ML” and 6.82% (44 genes) DEGs in “M vs. MX”. It is interesting to note that 7.42% (27 genes) DEGs in “M vs. ML” and 6.67% (genes)

DEGs in “M vs. MX” were annotated as the class Q (secondary metabolites biosynthesis, transport and catabolism), but the specific secondary metabolic pathways involved in the biosynthesis of pigments and citrinin need to be further analyzed through the KEGG metabolic pathway.

Furthermore, according to the data from the NR database (ESI Data S1†), we found that gene5930, gene8071 and gene283 were annotated as “citrinin polyketide synthase”, “citrinin biosynthetic transcription *CtnC*” and “citrinin biosynthesis transcriptional activator *CtnR*”, respectively. Compared with the M group, the transcription levels of gene5930, gene8071 and gene283 were down-regulated by 1.60/2.98, 3.20/5.79, and 1.18/1.50 log₂FC in the ML/MX group, respectively. Conversely, genes related to pigments biosynthesis (such as “non-reducing polyketide synthase” and “Zn cluster transcription factor”) were up-regulated by 1.33 and 2.22 log₂FC in the ML and MX groups. The

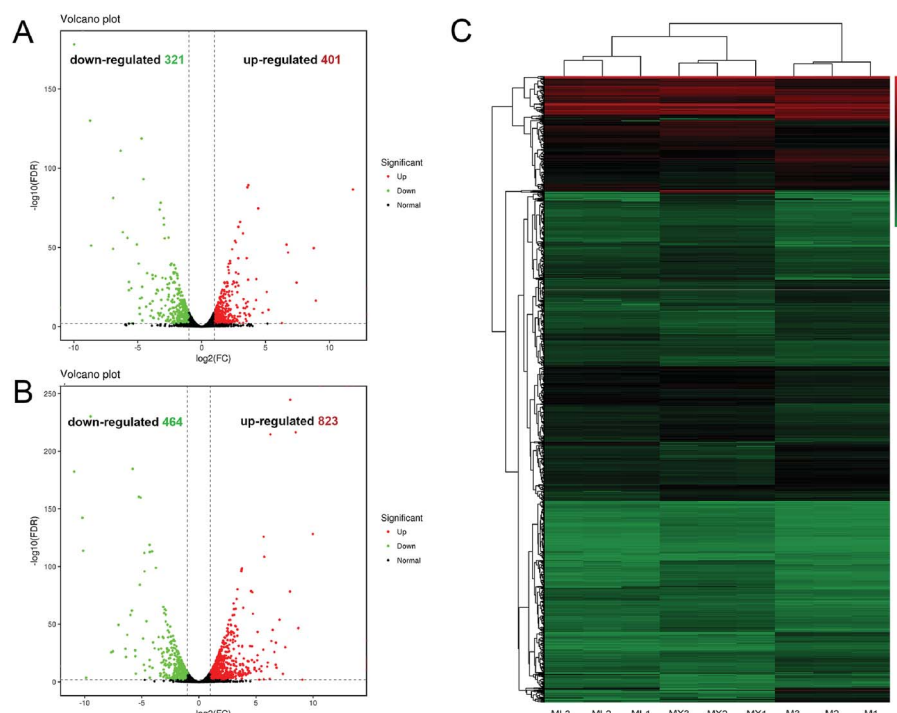


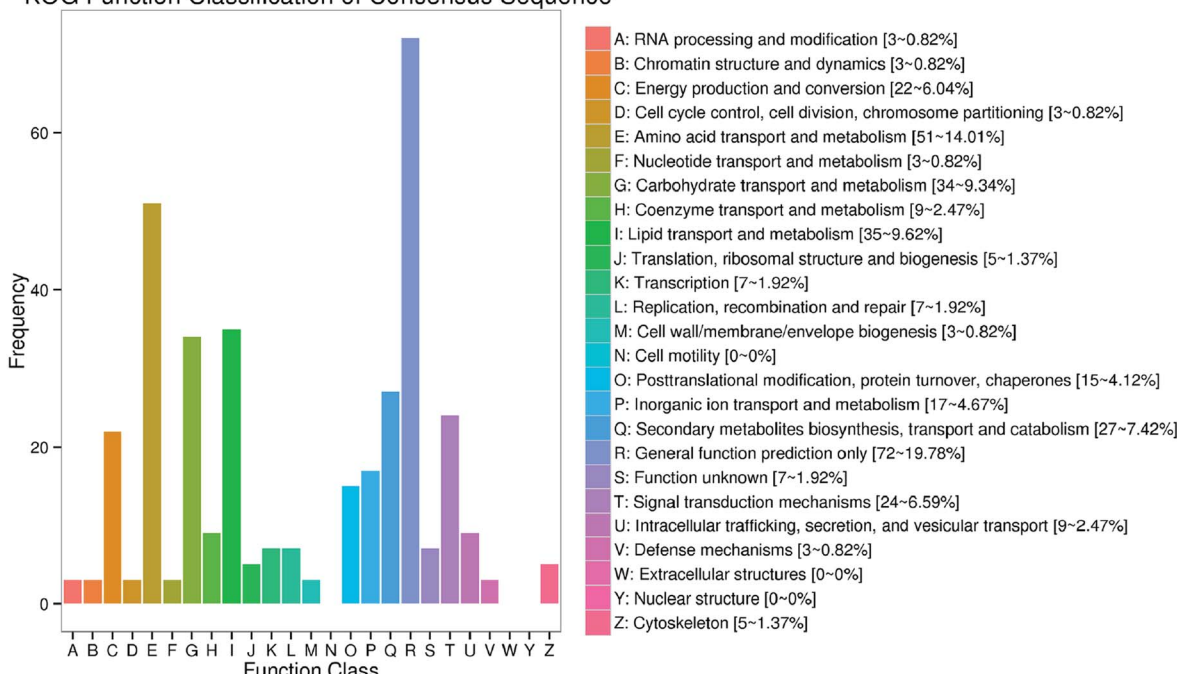
Fig. 3 The volcano plots analysis and cluster analysis of differentially expressed genes (DEGs) in M, ML and MX groups; (A and B) the volcano plots showing the comparison of DEGs in “M vs. ML” and “M and MX”, respectively. (C) The clustering map of DEGs in M, ML and MX groups.



Table 2 Annotated number of differentially expressed genes

DEG set	Total	COG	GO	KEGG	KOG	NR	Pfam	Swiss-Prot	eggNOG
M vs. ML transformants	682	306	403	187	322	681	539	402	671
M vs. MX transformants	1210	527	697	356	577	1208	918	707	1188
Total	1892	833	1100	543	899	1889	1457	1109	1859

A KOG Function Classification of Consensus Sequence



B KOG Function Classification of Consensus Sequence

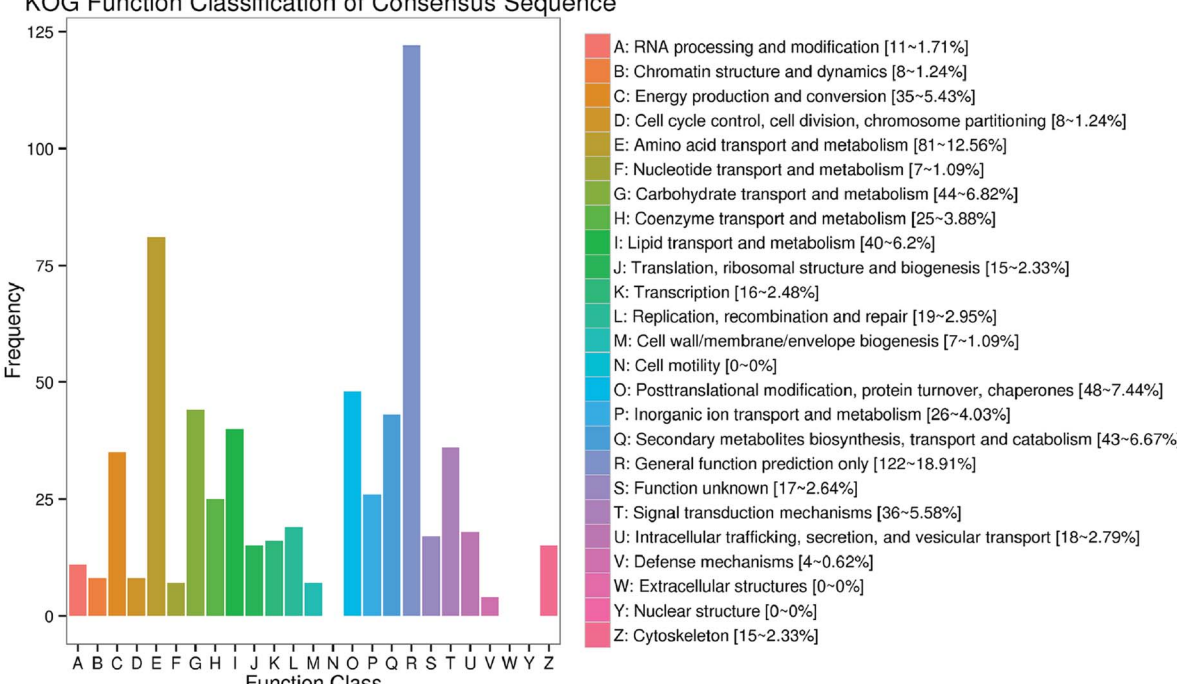


Fig. 4 KOG functional classification of DEGs in "M vs. ML" (A) and "M and MX" (B).



above findings indicated that inorganic nitrogen source (NH_4Cl and NH_4NO_3) can significantly down-regulate the genes involved in citrinin biosynthetic pathway to reduce citrinin production, and up-regulate the genes involved in pigments biosynthetic pathway to enhance pigment production.

GO is an international standardized classification system for gene functions, which can describe the properties of genes and gene products in organisms. GO enrichment analysis of the

three groups revealed that DEGs were classified into 43 functional groups, including biological processes (17 groups), cellular components (12 groups) and molecular functional classification (14 groups) (Fig. 5A and B; ESI Data S2†). “Catalytic activity” was the largest term of “molecular function”, and “metabolic process” was the largest term of “biological process” in “M vs. ML” and “M vs. MX”. “cell” and “cell part” were the

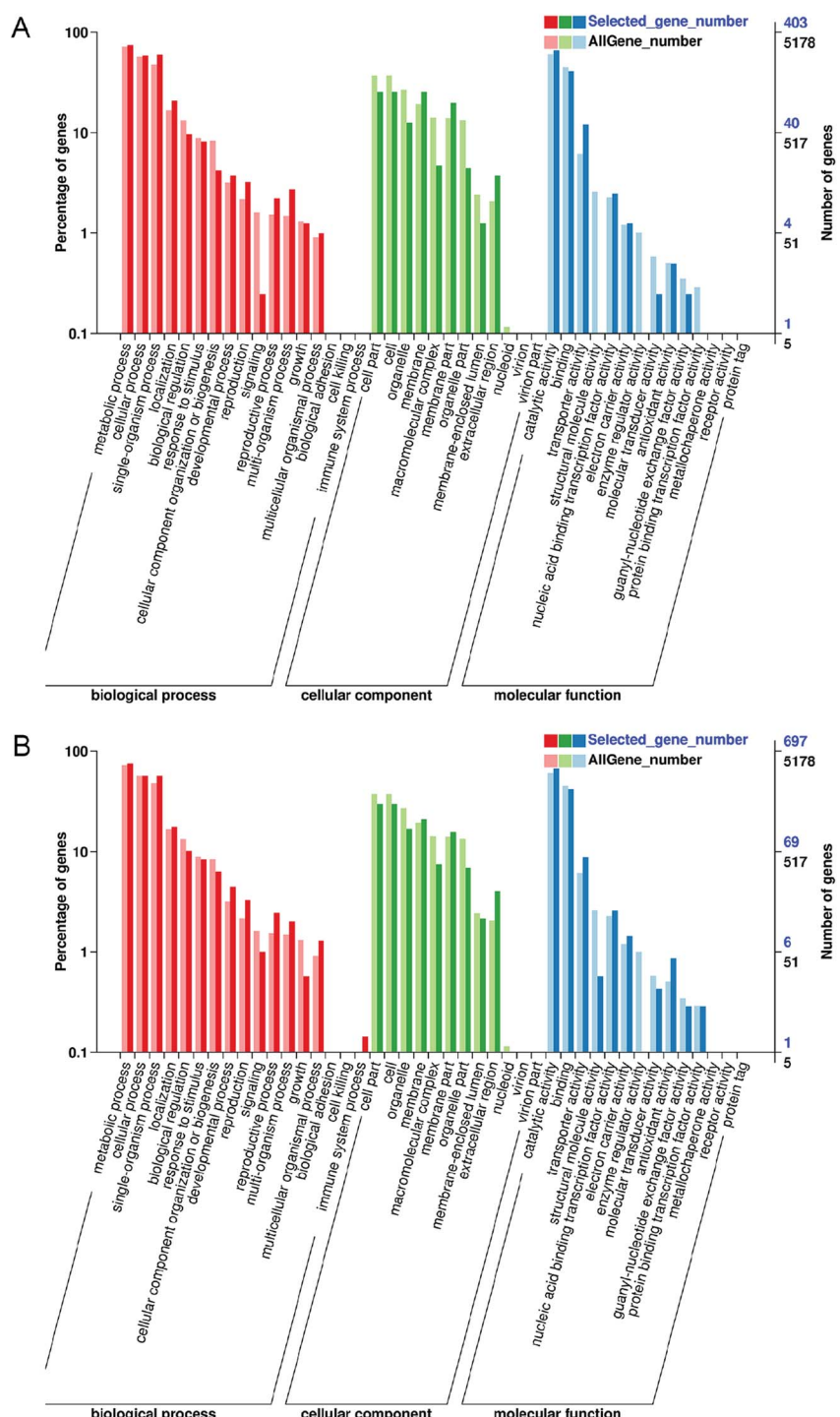


Fig. 5 GO functional classification of DEGs in “M vs. ML” (A) and “M and MX” (B).

largest terms of “cellular component” in “M vs. MX” and “M vs. ML”.

3.6 KEGG pathway enrichment of differentially expressed genes (DEGs)

Transcriptome data were further analyzed by KEGG metabolic pathway, with 75 and 99 metabolic pathways in M vs. ML and M vs. MX, respectively (ESI Data S3†). As shown in Fig. 6, the 20 most significant KEGG pathways for enrichment were revealed. “Biosynthesis of amino acids” was the pathway term with the largest number of DEGs enriched in M vs. ML and M vs. MX

(Fig. 6A and B). According to the corrected *P*-value (*q*-value), KEGG pathways in which significantly up-regulated and down-regulated enrichment in M vs. ML and M vs. MX were selected, respectively (Fig. 6A and B). “Biosynthesis of amino acids” was significantly enriched in ML and MX groups. Genes involved in aromatic amino acid metabolism pathway (which associated with “phenylalanine, tyrosine and tryptophan biosynthesis, gene7778”), and branched chain amino acid metabolism (such as “valine, leucine and isoleucine biosynthesis, gene4635”) were up-regulated in ML and MX groups, whereas genes involved in “tyrosine metabolism” were down-

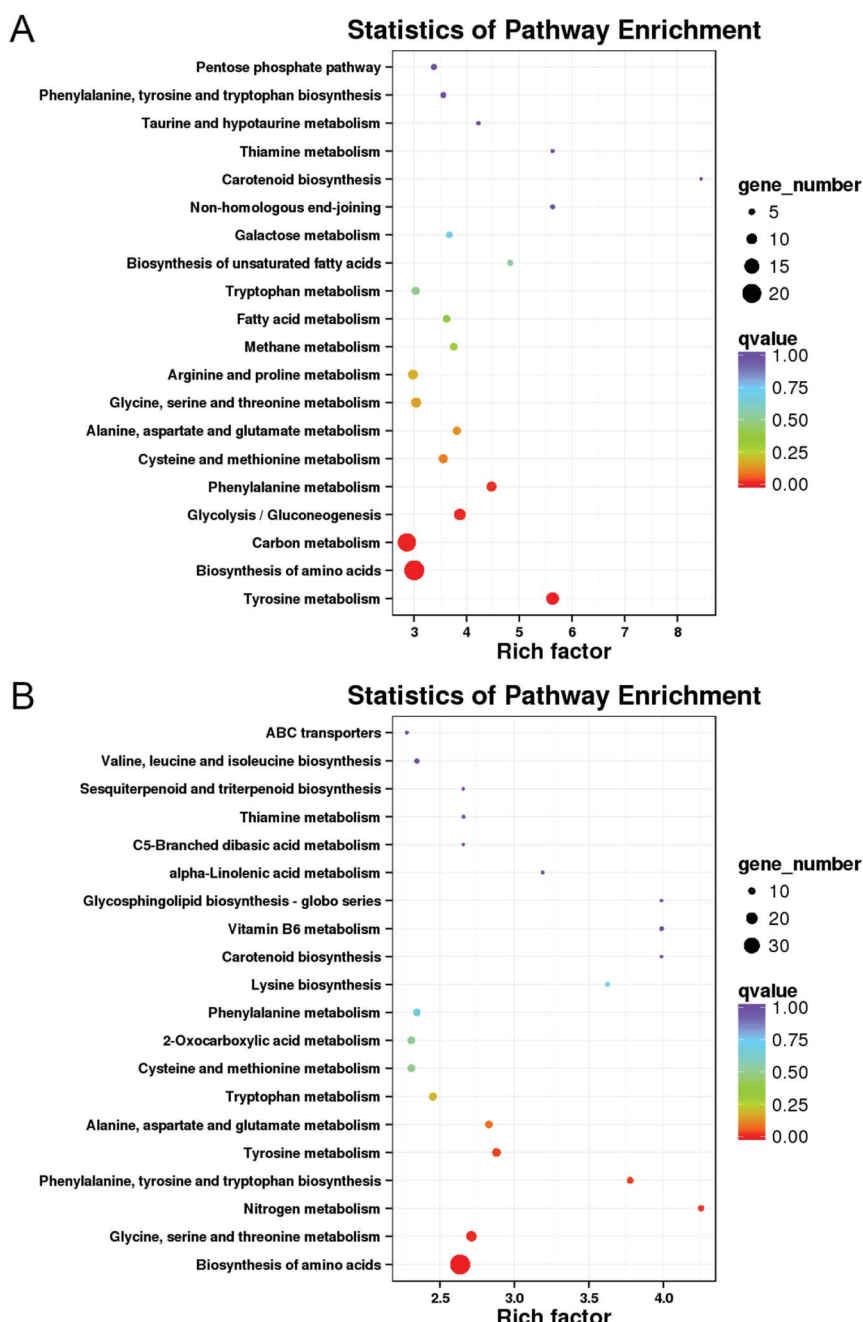


Fig. 6 KEGG pathway enrichment of “M vs. ML” (A) and “M and MX” (B). The enrichment factor indicates the number of DEGs relative to the percentage of all annotated genes involved in the pathway.



regulated in ML and MX groups, which can produce the biosynthetic precursors of pigments (ESI Data S4†). “Carbon metabolism” and “glycolysis/gluconeogenesis” were significantly up-regulated enrichment, and “phenylalanine metabolism” was significantly down-regulated enrichment in “M vs. ML”. “Phenylalanine, tyrosine and tryptophan biosynthesis”, “nitrogen metabolism” and “glycine, serine and threonine metabolism” were significantly up-regulated enrichment in “M vs. MX” (ESI Data S4†).

Expression profile of 61 DEGs in differential transcription levels involved in biosynthesis of amino acids and carbon metabolism pathway in ML and MX group compared to those in the M group ($P < 0.01$) (ESI Data S5†). Based on gene annotation, enzyme commission number (EC.number) and KEGG Orthology (KO), the transcription levels of gene1276, gene5012, gene6768, gene601, gene5806, gene1572, gene4323 involved in glycolysis/gluconeogenesis pathway in ML/MX group increased by 1.21/1.08, 1.41/0.89, 1.10/1.16, 3.46/2.97, 1.04/1.01, 1.23/0.77 and 1.09/0.78 \log_2 FC, respectively, which encoded glucose-6-phosphate isomerase (EC.5.3.1.9), 6-phosphofructokinase (EC.2.7.1.1/2.7.1.11), glyceraldehyde 3-phosphate dehydrogenase (EC.1.2.1.12), phosphoglycerate kinase (EC.2.7.2.2), 2,3-bisphosphoglycerate-independent phosphoglycerate mutase (EC.5.4.2.12), enolase (EC.4.2.1.11), pyruvate kinase (EC.2.7.1.40), respectively. Depending on these genes enriched in glycolysis/gluconeogenesis pathway, ML/MX group converts starch and sucrose to pyruvate, which can produce more biosynthetic precursors of pigments. The transcription levels of three genes (gene1276, gene3064, gene5012) involved in pentose phosphate pathway increased in ML/MX group by 1.21/1.08, 1.22/1.70, 1.41/0.89 \log_2 FC, respectively, which encoded glucose-6-phosphate isomerase (EC.5.3.1.9), gluconokinase (EC.2.7.1.12), 6-phosphofructokinase (EC.2.7.1.1/2.7.1.11), respectively. Depending on these genes enriched in pentose phosphate pathway, ML/MX group converts starch and sucrose to aromatic amino acid and NADPH, which can provide

precursors of pigments and a reducing agent for pigment synthesis. The transcription levels of five genes (gene4426, gene517, gene1078, gene5124, gene5672) involved in the biosynthesis of amino acids pathway (such as glycine, serine and threonine metabolism, cysteine and methionine metabolism and phenylalanine, tyrosine and tryptophan biosynthesis) in ML/MX group increased by 2.69/3.83, 1.57/2.76, 0.95/2.66, 3.25/4.73, and 2.54/2.59 \log_2 FC, respectively, which encoded *O*-acetylhomoserine/*O*-acetylserine sulphhydrylase (EC.2.5.1.49/2.5.1.47), aspartate aminotransferase (EC.2.6.1.1), phosphoserine aminotransferase (EC.2.6.1.52), tryptophan synthase (EC.4.2.1.20), L-serine dehydratase/L-threonine ammonia-lyase (EC.4.3.1.17/4.3.1.19), respectively. Depending on these genes enriched in amino acids pathway, ML/MX group converts serine and L-cysteine to tryptophan and pyruvate, which can produce more biosynthetic precursors of pigments.

Transcription factors play a critical role in the regulation of gene expression patterns that control the overall metabolism, including secondary metabolic pathways.^{34,35} Therefore, the expression levels of transcription factor genes in *M. purpureus* during pigment and citrinin production were analyzed. Based on Pfam annotation, 216 genes were identified as transcription factor-encoding candidates, among which 35 and 48 showed significantly different expression levels in the ML and MX group compared to those in M group (ESI Data S6†). Gene283 annotated as “citrinin biosynthesis transcriptional activator *CtnR*” in the ML/MX group were down-regulated with the \log_2 FC value 1.18/1.50, respectively. While those related to pigment biosynthesis (such as gene4184 annotated as “Zn cluster transcription factor”, gene6903 annotated as “C6 transcription factor” or “Fungal Zn(2)-Cys(6) binuclear” in the ML/MX group, gene6670 annotated as “Fungal Zn(2)-Cys(6) binuclear” in the MX group) was up-regulated by 2.22/2.66, 1.17/2.20, 1.80 \log_2 FC, respectively. This observation was inconsistent with the report that *CtnA*, which contains a typical Zn(II)₂Cys₆ DNA binding motif, functions as an activator of *pksCT* and *orf5* and that its

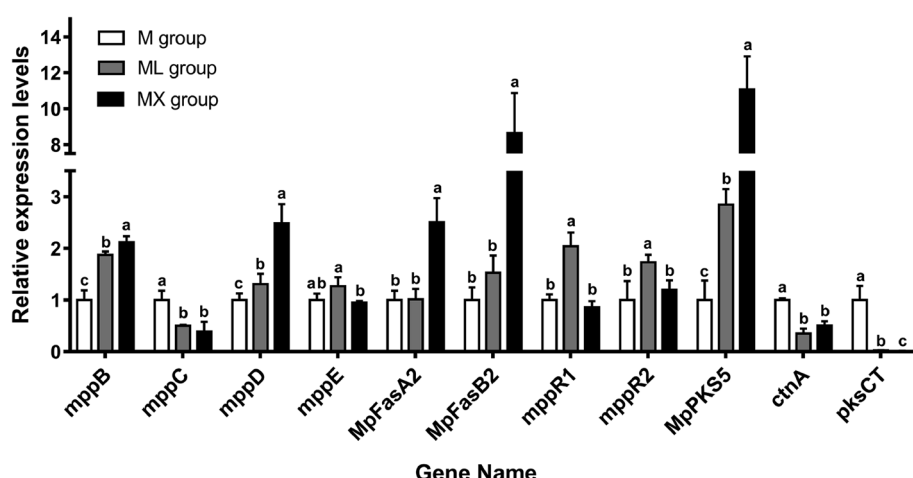


Fig. 7 Expression of pigments biosynthetic gene (*MpFasA2*, *MpFasB2*, *mppB*, *mppC*, *mppD*, *mppE*, *mppR1*, *mppR2* and *MpPKS5*) and citrinin biosynthetic gene (*ctnA* and *pksCT*) under M, ML and MX group monitored by RT-qPCR. The transcriptional levels were normalized to β -actin gene.



disruption leads to the reduction of citrinin production to barely detectable levels.³⁶ Gene5724 encoding a C₂H₂-type zinc finger in the ML/MX group, which is a key activator of sexual development in *Aspergillus nidulans*, was up-regulated by 2.07/2.95 log₂FC, respectively. Gene2534 (encoding a transcription factor *AreA*), a key transcription factor of nitrogen metabolism in filamentous fungi, was up-regulated by 0.81/1.36 log₂FC in the ML/MX group, respectively. Gene1163 (encoding nitrogen assimilation transcription factor *nit-4*), a member of the *GAL4* family of fungal transcription factors for nitrogen metabolism, was up-regulated by 1.33/1.10 log₂FC in the ML/MX group, respectively.

3.7 RT-qPCR of specific genes related to the biosynthesis of MPs and citrinin

In order to verify the effect of the medium supplemented with NH₄Cl or NH₄NO₃ on the yields of secondary metabolites of *Monascus* from the molecular level and the results of RNA-seq, the relative expression levels of the genes related to the biosynthesis of MPs and citrinin were also analyzed by RT-qPCR. The genes (*mppB*, *mppC*, *mppD*, *mppE*, *MpFasA2*, *MpFasB2*, *mppR1*, *mppR2* and *MpPKS5*) related to the biosynthesis of MPs, and the genes (*ctnA* and *pksCT*) involved in the biosynthesis of citrinin were detected (Fig. 7). It was found that NH₄NO₃ treatment resulted in significant elevations in the mRNA levels of *mppB*, *mppD*, *MpFasA2*, *MpFasB2* and *MpPKS5*, compared with the M group ($P < 0.05$). Moreover, NH₄Cl treatment made significant elevations in the mRNA levels of *mppB*, *mppD*, *mppR1*, *mppR2* and *MpPKS5*, compared with the M group ($P < 0.05$). Both NH₄NO₃ and NH₄Cl treatment resulted in reductions in the mRNA levels of *ctnA* and *pksCT*, compared with the M group ($P < 0.05$). It was noted that the mRNA levels of *mppC* (the gene responsible for the conversion of orange pigment to yellow pigment) in ML and MX were reduced ($P < 0.05$). The mRNA levels of *mppD* (the gene responsible for the conversion of yellow pigment to orange pigment) in ML and MX were increased simultaneously.

4. Discussion

Great efforts have been made to improve the fermentation conditions and medium ingredients to enhance the production of MPs in submerged fermentation.^{13,14} Although a large number of studies have been reported on the effects of different carbon and nitrogen sources on the production of MPs,^{19,20,37} the dual effects of inorganic nitrogen sources on the production of MPs and citrinin and their regulatory mechanisms are rarely reported. Previous studies have reported the use of RNA-seq technology to study the effects of blue light on *Monascus* and its related light-regulated gene regulatory networks.¹⁵ Some scholars have also explored the biosynthesis pathway of lovastatin in *M. purpureus* by RNA-seq technique.³⁸ However, there are few studies on the effect of nitrogen sources on the production of MPs and citrinin with submerged fermentation by *Monascus* through RNA-seq analysis.

Nitrogen source is a general term for nitrogen-containing compounds such as proteins and nucleic acids, which can not only affect the growth of mould, but also induce or inhibit the production of secondary metabolites in fungi. Previous studies stated that the best nitrogen sources used for good pigment production are NH₄Cl, NH₄NO₃ and peptone.^{39,40} Therefore, in this study, we use NH₄Cl, NH₄NO₃ and peptone as nitrogen sources to study their effects on the production of MPs and citrinin by liquid fermentation of *Monascus*. Compared with peptone, NH₄Cl and NH₄NO₃ can significantly increase the yields of MPs and inhibit the production of citrinin. Previous studies have shown that organic nitrogen sources are more likely to promote the growth of *Monascus* than inorganic nitrogen source but unfavourable for pigment production, probably because when *Monascus* grows too fast, it affects the transmission of nutrients and affects the synthesis of secondary metabolites.^{41,42} When *Monascus* is grown with different nitrogen sources, the *Monascus* pigment components have different characteristics, which is consistent with previous studies when nitrogen source was ammonium nitrate, which significantly increased the production of red pigments, and when peptone was used as nitrogen source, the intracellular pigments was mainly yellow pigments, and a small amount of red pigments.^{14,41} Nitrogen sources have the same ammonium ion but different negative ions (such as NH₄Cl and NH₄NO₃). Different nitrogen sources have different effects on growth and secondary metabolism.⁴³ These results indicated that different ions may affect the biosynthesis of MPs or citrinin, and also reveal the complexity and specificity of the nitrogen source for the regulation of the biosynthesis of secondary metabolites from *Monascus*.

According to the previous research reported by Chen *et al.*,^{2,44} different *Monascus* pigments are produced by the same pigment intermediate through three synthetic pathways. Two red pigments (rubropunctamine and monascorubramine) are produced in a non-catalyzed affinity ammonia reaction by orange pigments (rubropunctatin and monascorubrin). The yellow pigments (monasfluore A, monasfluore B) and (monascin, ankaflavin) are competitively synthesized by two enzyme-catalyzed reaction pathways.^{2,14,44} In this study, the red pigment synthetic pathways were preferred in the ML and MX groups, which is consistent with the production of red pigments in *Penicillium* sp. AZ.⁴⁵ Interestingly, the accumulation of yellow and orange pigments in *Penicillium* sp. AZ can be regulated by the restriction of inorganic ammonium salt in fermentation medium.⁴⁵

To better understand the effects of NH₄Cl and NH₄NO₃ on the production of MPs and citrinin by the liquid fermentation of *Monascus*, we used transcriptome sequencing to explore the differential gene expression of *Monascus purpureus* M3103. In filamentous fungi, the transcription factors *AreA*, *AreB*, *MeaB*, *TamA* and *NmrA* regulate the nitrogen source catabolism in combination.^{46,47} The *nit-4* family transcription factor (gene1163) and the GATA family transcription factor *AreA* (gene2543) were detected in our transcription data. *Nit-4*, a member of the *GAL4* family of fungal transcription factors, encodes a protein with an amino-terminal Cys₆/Zn₂ domain



that provides sequence-specific DNA binding.^{46,48} *AreA* is considered as the major regulator of a large number of genes involved in uptake and metabolism of non-favored N sources in fungi,⁴⁹ and acts as a positive regulator promoting the use of secondary nitrogen sources and participating in certain secondary metabolites of many clusters of many culture under nitrogen-limiting conditions.^{46,49} Compared with M group, the production of secondary metabolites of pigments and citrinin in ML and MX were significantly changed. And the utilization of secondary nitrogen sources were promote by upregulating *AreA* after the large consumption of ammonium nitrogen in the MX group. These results were consistent with other literature reports.^{49,50}

The sequencing data of *Monascus* mycelium in NH₄Cl group, NH₄NO₃ group and peptone group were compared. According to the KEGG pathway, nitrogen sources played an important role in the primary metabolism and secondary metabolism of *Monascus*. When NH₄Cl was used as nitrogen source, amino acid metabolism (including glycine, serine and threonine metabolism), carbon metabolism (glycolysis/glyconeogenesis, pentose phosphate pathway), and lipid metabolism were increased, and phenylalanine metabolism and tyrosine metabolism were attenuated. However, when NH₄NO₃ was used as nitrogen source, nitrogen metabolism and amino acid metabolism (such as phenylalanine, tyrosine and tryptophan biosynthesis, glycine, serine and threonine metabolism, valine, leucine and isoleucine metabolism) were enhanced, and tyrosine metabolism was attenuated. Glycine plays an important role in cell division.⁵¹ Recent research has shown that serine is an important glycolysis fuel and a product of oxidative decarboxylation of glycine, providing a reducing power (NADH) and a carbon precursor for anabolism,⁵² and affects intracellular redox metabolic activity. Serine, glycine and threonine metabolism provided energy for the TCA cycle by pyruvate and acetyl-CoA.⁵³ This indicated that NH₄Cl as nitrogen source can enhance partial amino acid metabolism to provide energy for the growth of *Monascus*. It is worth noting that phenylalanine, tyrosine tryptophan are aromatic amino acids, which can be converted into several biological compounds (including 5-hydroxytryptamine, heteroauxin, dopamine, melanin, etc.) through a series of oxidation reactions and participate in the biological physiology and growth process,² and which can produce the biosynthetic precursors of pigments. Valine, leucine and isoleucine are branched chain amino acids can be degrade to produce acetyl-CoA and methyl-CoA,⁵⁴ which can be further catalyzed by polyketide synthase to form type II polyketides, the precursors of synthetic MPs.⁵⁵ This indicated that both NH₄NO₃ and NH₄Cl attenuated the growth ability of *Monascus*, but promoted the secondary metabolic process of *Monascus*.

Further, according to the data of the NR database, it was found that when NH₄Cl was used as nitrogen source, the number of genes related to the polyketide synthase pathway and citrinin biosynthesis transcription factor were down-regulated. Similarly, such a phenomenon was found in *Monascus* with NH₄NO₃ as nitrogen source. This further confirmed that when using NH₄NO₃ or NH₄Cl as nitrogen source, the yields of MPs

were increased and the yield of citrinin was decreased. It is noteworthy that the relative expression levels of *MpPKS5*, *mppR1*, *mppR2*, *mppB* and *mppD*, were significantly up-regulated in the ML group compared with the M group. *MpPKS5* and *mppD* were the key genes in the polyketide synthase (PKS) pathway of *Monascus* pigment synthesis. And knocking out *MpPKS5* resulted in very low production of MPs.⁵⁶ Amine oxidoreductase encoded by *mppD* was responsible for the conversion of yellow pigment to orange pigment. *MppR1* was a specific transcription factor of *Monascus* with fungal Zn(2)-Cys(6) binuclear cluster domain, and ankyrin repeat protein encoded by *mppR2* was an important regulator of MPs biosynthesis.^{57,58} *MppB* encoding fatty acyltransferase in the polyketide pathway produced pigments condensed by esterification: a short-chain fatty acid (C6 or C8) and a hexaketide synthesized.⁵⁹ The up-regulation of these gene expression levels was consistent with the up-regulation of the yield of MPs. The expression level of pigment-related genes (*mppC* and *mppD*) were regulated by inorganic nitrogen sources to produce red pigment oxidized from yellow pigment and orange pigment. Moreover, there was no significant difference in the relative expression levels of genes *MpFasA2* and *MpFasB2* between ML group and M group, which indicated that the addition of NH₄Cl did not change the expression of these genes in liquid fermentation. *MpFasA2* and *MpFasB2* encoded the synthesis of short-chain fatty acyl coenzyme A, which was the key gene in the fatty acid pathway of MPs.² This study demonstrated that the addition of ammonium chloride in liquid fermentation stimulated the expression of genes related to the polyketide synthase (PKS) pathway in the synthesis of MPs, but did not significantly affect the expression of genes related to the fatty acid pathway. The relative expression levels of *pksCT* and *ctnA* were also significantly down-regulated. Both *ctnA* and *pksCT* were very important genes related to citrinin biosynthesis.⁶⁰ The inhibitory effects of *ctnA* and *pksCT* on transcriptional expression of MX group were better than ML group, resulting in a dramatic decrease in citrinin production. These conclusions can be verified in previous NR database data and KEGG metabolic pathways.

Several novel genes involved in pigment and citrinin synthesis were found in our study (ESI Data S7†). Such as (newGene369, newGene370), (newGene371, newGene372, newGene_112, newGene281, newGene1307) and newGene_1228, annotated as non-reducing polyketide synthase, conidial yellow pigment biosynthesis polyketide synthase and polyketide synthase dehydratase, respectively, all of them may be constituted a gene cluster of polyketides. NewGene_227, newGene_979, newGene_1137, newGene_802, newGene_218 and newGene_219, annotated as pyruvate decarboxylase, pyruvate dehydrogenase, putative cytochrome p450 protein, cytochrome c oxidase subunit 7A, and FAD dependent oxidoreductase, may be involved in the biosynthesis, transport and catabolism of secondary metabolites. NewGene_185, newGene_339, newGene_510 and newGene_1227, annotated as fatty acid synthase alpha subunit, 3-oxoacyl-[acyl-carrier-protein] synthase, 3-ketoacyl-CoA ketothiolase and zinc-binding dehydrogenase, may be involved in lipid transport and metabolism. NewGene_623 and



newGene1144 annotated as citrinin polyketide synthase and Zn(2)-Cys(6) binuclear cluster domain, were closely related to the citrinin biosynthesis.

5. Conclusion

In this study, the effects of inorganic nitrogen on the secondary metabolism and gene transcription of *Monascus* spp. in liquid fermentation were studied, which provides a better understanding of the links between primary and secondary metabolism in *Monascus* spp., and is expected to lay a foundation for future studies on the molecular mechanisms underlying the effects of different substances on secondary metabolism in *Monascus* spp. Transcriptome analysis highlighted differences not only in gene expression of pigments and citrinin biosynthesis related genes, but also shed light on regulatory effects on genes involved in other energetic metabolisms and biological processes, such as carbon and nitrogen metabolism, biosynthesis of amino acids and so on.

Conflicts of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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