

Exploring the Distribution of Citrinin Biosynthesis Related Genes among Monascus Species

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Citrinin, a hepato-nephrotoxic compound to humans, can be produced by the food fermentation microorganisms Monascus spp. In this study, we investigated the distribution of mycotoxin citrinin biosynthesis genes in 18 Monascus strains. The results show that the acyl-transferase and ketosynthase domains of the pksCT gene encoding citrinin polyketide synthase were found in Monascus purpureus, Monascus kaoliang, and Monascus sanguineus. Furthermore, the ctnA gene, a major activator for citrinin biosynthesis, was found in M. purpureus and M. kaoliang, but was absent in M. sanguineus. The orf3 gene encoding oxygenase, located between pksCT and ctnA, was also present in M. purpureus and M. kaoliang. The pksCT gene was highly conserved in M. purpureus, M. kaoliang, and M. sanguineus, while the ctnA and orf3 genes were shown to be highly homologous in M. purpureus and M. kaoliang. In contrast, the PCR and Southern blot analyses suggest that pksCT, ctnA, and orf3 were absent or significantly different in Monascus pilosus, Monascus ruber, Monascus barkeri, Monascus floridanus, Monascus lunisporas, and Monascus pallens. A citrinin-producing phenotype was detected only in M. purpureus and M. kaoliang using high performance liquid chromatography (HPLC). These results clearly indicate that the highly conserved citrinin gene cluster in M. purpureus and M. kaoliang carry out citrinin biosynthesis. In addition, according to the phylogenetic subgroups established with the β -tubulin gene, the citrinin gene cluster can group the species of Monascus.

KEYWORDS: Citrinin; *Monascus*; polyketide; β-tubulin

INTRODUCTION

Typically used in fermented foods, the filamentous fungi Monascus spp. are known to produce various secondary metabolites with polyketide structures such as pigments, monacolin K, and citrinin (1-3). The red pigments produced by the various species of *Monascus* have been used extensively as natural food colorants (4). Known as lovastatin and normally used for reducing serum cholesterol levels in humans, monacolin K was first isolated from the *Monascus ruber* medium (1), although it can also be found in Aspergillus terreus (5-7). Citrinin, the hepato-nephrotoxic agent and Gram-positive bacteria antibiotic, has been identified in a variety of fungi such as Aspergillus and Penicillium spp (8, 9).; the same substance is also found in Monascus as monascidin A (10, 11). Citrinin belongs to the group of polyketides synthesized by the iterative type I polyketide synthase (PKS). Its biosynthesis in Monascus originates from a tetraketide arising from the condensation of one acetyl-CoA molecule with three malonyl-CoA molecules (12). Although Monascus has been widely used in food fermentation, and shown great promise for medicinal development, its application is limited by the nephrotoxic and hepatotoxic properties of citrinin.

The citrinin biosynthetic gene cluster in Monascus purpureus BCRC33325 (IFO30873) was proposed in Shimizu's studies (3). The polyketide synthase (pksCT) and transcriptional activator (ctnA) have been proven to be involved in citrinin biosynthesis (3, 13). As such, the disruption of pksCT results in the phenotype of lost citrinin production (3). The ctnA gene that encodes the Zn(II)2Cys6 binuclear DNA binding protein is a major activator of citrinin biosynthesis. Consequently, the ctnA-disrupted strain of M. purpureus also exhibits a significant decrease in citrinin production, to a barely detectable level (13). To characterize the citrinin-producing Monascus strains, we analyzed the genotype and phenotype of the citrinin synthesized by the various species of *Monascus*. On the basis of the citrinin biosynthetic gene cluster in M. purpureus BCRC33325 (IFO30873) (3, 13), several primers were designed for polymerase chain reactions (PCR) and Southern hybridizations. Citrinin production was further determined by HPLC analysis.

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Table 1. Strains Used in This Work

strain	species ^a	polyketide synthase (pksCT) ^b	activator (ctnA)	oxygenase (orf3)	source	citrinin production
1. BCRC 31502 (ATCC 16363)	Monascus pilosus, type	_	_	_	Japan	_
2. BCRC 38072	M. pilosus	_	_	_	Taiwan	_
3. BCRC 31533 (ATCC 16246)	M. ruber, type	_	_	_	Nigeria	_
4. BCRC 31523 (ATCC 16378)	M. ruber	_	_	_	Taiwan	_
5. BCRC 31534 (ATCC 16366)	M. ruber	_	_	_	Switzerland	_
6. BCRC 31535 (ATCC 18199)	M. ruber	_	_	_	Canada	_
7. BCRC 33314 (ATCC 16371)	M. ruber	_	_	_	Taiwan	_
8. BCRC 33323 (ATCC 18199)	M. ruber	_	_	_	Canada	_
9. BCRC 31542 (ATCC 16365)	M. purpureus, type	+	+	+	Java	+
10. BCRC 31541 (ATCC 16379)	M. purpureus	+	+	+	Taiwan	+
11. BCRC 33325 (IFO 30873)	M. purpureus	+	+	+		+
12. BCRC 31615 (DSM 1379)	M. purpureus	+	+	+		+
13. BCRC 31506 (CBS 302.78)	Monascus kaoliang, type	+	+	+	Taiwan	+
14. BCRC 33446 (ATCC 200613)	Monascus sanguineus, type	+	_	_	Iraq	_
15. BCRC 33309 (ATCC 16966)	Monascus barkeri	_	_	_	Japan	_
16. BCRC 33310 (IMI 282587)	Monascus floridanus, type	_	_	_	USA	_
17. BCRC 33640 (ATCC 204397)	Monascus lunisporas, type	_	_	_	Japan	_
18. BCRC 33641 (ATCC 200612)	Monascus pallens, type	_	_	_	Iraq	_

^a "Type" indicates type strain. ^b +, positive; -, negative.

Table 2. Primers Used To Amplify Citrinin Related Genes Fragments

	<u> </u>			
primer ^a	20-mer sequence	position ^b		
F1	5'-AACGGACAGGAAGAGCGTGC	68-87 (ctnA)		
F2	5'-ACGAGTGTCAGTTCGGCTCC	483-502 (ctnA)		
F3	5'-TCGGAAGCGATCATGGACGT	1277—1296 (ctnA)		
F4	5'-CTCCTTTCCGCGCAATTCCA	760-779 (ctnA)		
F5	5'-CGTGCACCTCTACAGGGTTC	113—132 (orf3)		
F6	5'-CTACCAGGCCATGCTGAAGC	447-466 (orf3)		
F7	5'-GAGTCCCCGAGAAATGGCAT	1445-1464 (pksCT)		
F8	5'-AACTGGTCTCTTCCCCAAGC	2661-2680 (pksCT)		
F9	5'-TTAACCGTCTCCTGTCCGGC	4012-4031 (pksCT)		
F10	5'-TGCCTATCACGTCAACGGCA	5348-5367 (pksCT)		
F11	5'-ACGTGGACCATGCCGAGAAC	3313-3332 (pksCT)		
R1	5'-CGTCTGGTGCAGTTAATGCG	958-977 (ctnA)		
R2	5'-GGTATGGCATCGGTGGTGTG	1568-1587 (ctnA)		
R3	5'-GAAACGGGGGAGTGGATTGG	700-719 (orf3)		
R4	5'-GAGGATCGGATGCGGCATTT	1843-1862 (ctnA)		
R5	5'-TCTTCGATGGCAACCTGGAC	854-873 (orf3)		
R6	5'-CTGCCATCTTCCAAGCCCAA	239-258 (orf4)		
R7	5'-AACAACTACGGTGCTTCCGG	2427-2446 (pksCT)		
R8	5'-CGGGCTCTGGGTACATCAAA	3654-3673 (pksCT)		
R9	5'-CGGTCTTGAACCTGACGAGG	5090-5109 (pksCT)		
R10	5'-GAAGGTACTCGGCCAGAAGC	6486-6505 (pksCT)		
R11	5'-CAATCACATTCCAAGCGGCG	4303-4322 (pksCT)		
F12	5'-TCGTTATCTAGGCTGGGCCA	678-697 (ctnA)		
R12	5'-CGCTGTTTGCGATGCAGTAT	2977-2996 (pksCT)		
R13	5'-GCCGCCCCATTGAAGAATAC	428-447 (orf3)		

^a F, forward primer; R, reverse primer. ^b The sites corresponding to those of the *M. purpureus* BCRC33325 (IFO30873) citrinin gene cluster (GenBank accession no. AB243687).

Our results indicated that the distribution of citrinin biosynthetic genes is highly related to the phylogenetic group of *Monascus* spp.

MATERIALS AND METHODS

Strains, Media, and Growth Conditions. Eighteen strains of *Monascus* as listed in **Table 1** were used in this study. All strains were maintained on PDA (DIFCO, Detroit, MI) agar for 1 week, and spore suspensions were obtained by washing the cultured PDA agar plates with distilled water. Mycelia were harvested after incubation for 14 days at 25 °C with constant agitation in liquid medium (7% glycerol, 3% glucose, 3% monosodium glutamate, 1.2% polypetone, 0.2% NaNO₃, and 0.1% MgSO₄•7H₂O).

Genomic DNA Isolation. *Monascus* genomic DNA was extracted according to the method developed by Bingle et al. (14). Approximately 0.5 g (squeezed wet weight) of frozen mycelia was ground to a fine power under liquid nitrogen using a mortar and pestle. Proteins were

then removed by successive rounds of extraction using phenol and chloroform. Genomic DNA was recovered by precipitation using 2-propanol and then dissolved in TE buffer.

Polymerase Chain Reaction (PCR) and sequencing. The PCR amplification was carried out with a PCR system 2700 thermocycler (Applied Biosystems, Foster City, CA). The 50 μ L reaction mixture contained 100 ng of fungal DNA as template, 0.2 mM of primers, 2 units of Taq DNA polymerase, and 800 mM dNTPs. The 11 primer sets listed in **Table 2** were used for the citrinin biosynthetic gene amplification in this study. The reaction conditions included an initial denaturation for 5 min at 96 °C, which was followed by 30 cycles for 1 min at 96 °C, 1 min at 50 °C, and 2 min at 72 °C with a final extension of 10 min at 72 °C. The PCR products were resolved and recovered on 1.2% agarose gels. Cycle sequencing reactions were then carried out using the BigDye, V3.0 kit (Applied Biosystems, Foster City, CA), while DNA sequencing was performed using an ABI Prism 3730 Sequencer (Applied Biosystems, Foster City, CA).

Southern Hybridization. For Southern hybridizations, genomic DNA (10 µg per lane) was digested with the *Hin*dIII restriction enzyme and separated through 1.2% agarose gels by electrophoresis. Southern hybridization analysis was performed using the DIG system (Roche Diagnostics, Mannheim, Germany). The probes of the citrinin biosynthesis genes were labeled through PCR amplification from the genomic DNA of *M. purpureus* BCRC33325 (IFO30873), using the PCR DIG probe synthesis kit (Roche Diagnostics, Mannheim, Germany). The primer sets of citrinin biosynthesis genes were *pksCT*-F8, AACTG-GTCTCTCCCCAAGC; *pksCT*-R12, ATACTGCATCGCAAACAGCG; *ctnA*-F12, TCGTTATCTAGGCTGGGCCA; *ctnA*-R1, CGTCTGGT-GCAGTTAATGCG; *orf3*-F5, CGTGCACCTCTACAGGGTTC; and *orf3*-R13, GCCGCCCCATTGAAGAATAC.

Phylogenetic Analysis. Sequence analyses of the amplified DNA were performed using the VectorNTI 9.0 software (InforMax, Frederick, MD). The accession numbers used for the partial β -tubulin genes were as follows: Monascus species, DQ299886-DQ299896, AY498587-AY498589, AY498596, AY498598, AY498601, AY498602, and AY498604; Aspergillus flavus, M38265; Aspergillus parasiticus, L49386; and Aspergillus fumigatus, AY048754. The accession numbers used for the polyketide synthase genes were as follows: Saccharopolyspora erythraea DEBS (X56107 and X62569), Aspergillus terreus lovF (AF141925), A. terreus lovB (AF151722), Penicillium citrinum mlcA, mlcB (AB072893), Phoma sp. SQTKS (AY217789), Cochliobolus heterostrophus pks1 (U68040), Gibberella moniliformis FUM1 (AF155773), M. purpureus pksCT (AB167465), Emericella nidulans wA (X65866), E. nidulans stcA (AAC49191), Aspergillus parasiticus aflC (AY371490), Penicillium patulum 6-MSAS (X55776), A. parasiticus pksL2 (U52151), and A. terreus pksM (U31329). The phylogenetic tree was constructed using the neighbor-joining method (15) via the MEGA 3.1 software with 1000 bootstrap replicates.

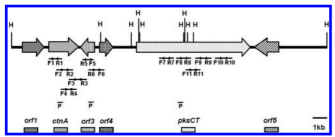


Figure 1. Citrinin gene cluster of M. purpureus BCRC33325 (IFO30873) obtained from the GenBank database using accession no. AB243687. The citrinin gene cluster includes dehydrogenase (orf1), transcriptional activator (ctnA), oxygenase (orf3), oxidoreductase (orf4), polyketide synthase (pksCT), and transporter (orf5). The abbreviations F and R indicate the forward and reverse primers, respectively, of ctnA, orf3, and pksCT for the PCR analysis; the abbreviation P indicates the probes for ctnA, orf3, and pksCT in the Southern hybridization analyses.

Measurement of Citrinin. The aliquots of the *M. pilosus* culture were cleared of cells and filtered using a 0.2 mm filter. The supernatants were analyzed using HPLC performed on a Waters system (Waters, Milford, MA) fitted with a μ Bondapak C₁₈ (10 μ m) column (Waters, Milford, MA). The HPLC parameters were as follows: solvent gradient with phosphoric acid/acetonitrile/2-propanol 60:35:5 to 25:70:5 (v:v: v) in 10 min; flow rate, 0.8 mL min⁻¹. Fluorescence detection was performed using the 2475 Multi λ Fluorescence (Waters, Milford, MA) set at 330 nm excitation wavelength and 500 nm emission wavelength. A citrinin standard compound (Sigma, St. Louis, MO) was used to confirm the HPLC analysis.

RESULTS

PCR and Southern Hybridization Analysis of Citrinin Biosynthetic Genes. Studies on fungal polyketide biosynthetic genes have indicated that metabolites are largely synthesized by the iterative multifunctional polyketide synthase systems (16). Each PKS minimally carries keto-synthase (KS), acyl-transferase (AT), and acyl carrier protein (ACP) domains to catalyze different modifications. To search for the genes related to citrinin biosynthesis, 11 primer sets were designed according to the citrinin gene cluster in M. purpureus BCRC33325 (IFO30873) (3, 13) (Table 2 and Figure 1). Eighteen strains of the Monascus species were used to amplify the genomic DNA of the citrinin biosynthetic genes (**Table 1**). The PCR results showed that the five primer sets of the pksCT gene were amplified in M. purpureus BCRC31542, BCRC31541, BCRC33325, and BCRC31615, in M. kaoliang BCRC31506, and in M. sanguineus BCRC33446. In contrast, none of the five primer sets amplified the pksCT gene from M. pilosus, M. ruber, M. barkeri, M. floridanus, M. lunisporas, and M. pallens. The PCR analyses of ctnA (encoding a transcriptional activator) and orf3 (encoding oxygenase) with six primer sets were conducted to further verify the citrinin biosynthetic gene cluster. It was found that all strains of M. purpureus and M. kaoliang possessed both ctnA and orf3 genes, but that these two genes were absent in the M. sanguineus strain and other Monascus species. Although the PCR condition adopted a low annealing temperature (50 °C), only M. purpureus and M. kaoliang could yield the PCR products typical of the pksCT, ctnA, and orf3 genes, while M. sanguineus lacked the PCR products for the ctnA and orf3 genes.

To confirm the citrinin biosynthetic gene distribution in the Monascus species, the genomic DNAs were digested by HindIII for Southern hybridization. The results revealed the presence of pksCT, ctnA, and orf3, which was consistent with the PCR results. Nevertheless, instead of a 3-kb fragment corresponding to the pksCT gene in M. purpureus and M. kaoliang, a 2-kb

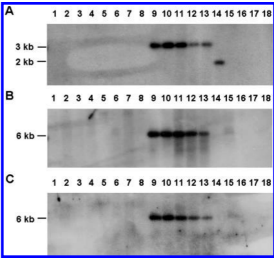


Figure 2. Southern hybridization analyses of the citrinin-related genes. Chromosomal DNA extracted from the Monascus species were digested by HindIII, separated by gel electrophoresis, and hybridized with the (A) pksCT, (B) ctnA, and (C) orf3 probes. Monascus species: lane 1, M. pilosus BCRC31502; lane 2, M. pilosus BCRC38072; lane 3, M. ruber BCRC31533; lane 4, *M. ruber* BCRC31523; lane 5, *M. ruber* BCRC31534; lane 6, M. ruber BCRC31535; lane 7, M. ruber BCRC33314; lane 8, M. ruber BCRC33323; lane 9, M. purpureus BCRC31542; lane 10, M. purpureus BCRC31541; lane 11, M. purpureus BCRC33325; lane 12, M. purpureus BCRC31615; lane 13, M. kaoliang BCRC31506; lane 14, M. sanguineus BCRC33446; lane 15, M. barkeri BCRC33309; lane 16, M. floridanus BCRC33310; lane 17, M. lunisporas BCRC33640; and lane 18, M. pallens BCRC33641.

HindIII fragment was present in M. sanguineus. Meanwhile, the 6.1-kb fragments corresponding to the ctnA and orf3 genes were present in the Southern blot results of M. purpureus and M. kaoliang, which generally agreed with the size described in the citrinin biosynthetic gene cluster from M. purpureus BCRC33325 (IFO30873) (Figure 2) (3). In addition, the ctnA and orf3 genes were consistently absent in M. sanguineus and other *Monascus* species.

Sequence and Phylogenetic Analysis of pksCT, ctnA, orf3, and β -Tubulin Genes. The candidate PCR products of pksCT, ctnA, and orf3 were further sequenced and analyzed. The result of the 1-kb amplified DNA showed that the AT domain of pksCT shared a high similarity among M. purpureus BCRC31542, BCRC31541, BCRC33325, and BCRC31615; M. kaoliang BCRC31506; and M. sanguineus BCRC33446. Both M. purpureus and M. kaoliang shared 100% identity, while M. sanguineus shared 97% identity with the other species. The sequences from M. purpureus, M. kaoliang, and M. sanguineus all contained the conserved amino acid sequence GHSXG, which was the typical active site of the AT domain. The phylogeny was constructed according to the conserved AT domain (**Figure 3**). The *pksCT* from *M. purpureus*, *M. kaoliang*, and M. sanguineus belonged to the nonreduced polyketide structural type. M. sanguineus was placed in a branch that was separate from M. purpureus and M. kaoliang. The KS domain of pksCT was further sequenced. The sequences from M. purpureus and M. kaoliang contained the conserved amino acid sequence DXACXS, which was the typical active site of the KS domain. However, the sequence revealed a gap in the upstream of the KS domain of *M. sanguineus* (Figure 4); the out-of-frame *pksCT* may result in a nonfunctional polyketide synthase. The active site of the KS domain, DXPCXS, was also different from the typical conserved sequence. Obviously, this pksCT was divergent with M. purpureus and M. kaoliang.

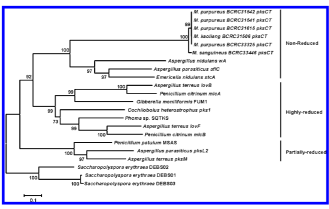


Figure 3. Phylogenetic tree of PKSs from *Monascus* and various organisms. The phylogeny of PKSs based on the conserved acyltransferase domain with 300 amino acids was constructed and rooted using the acyl-transferase domains of *S. erythraea* DEBS (X56107 and X62569). The accession numbers for the polyketide synthase genes were as follows: *A. terreus lovF* (AF141925), *A. terreus lovB* (AF151722), *P. citrinum mlcA, mlcB* (AB072893), *Phoma* sp. SQTKS (AY217789), *C. heterostrophus pks1* (U68040), *G. moniliformis* FUM1 (AF155773), *M. purpureus pksCT* (AB167465), *E. nidulans wA* (X65866), *E. nidulans stcA* (AAC49191), *A. parasiticus aflC* (AY371490), *P. patulum* 6-MSAS (X55776), *A. parasiticus pksL2* (U52151), and *A. terreus pksM* (U31329). Bootstrap values are shown in the nodes according to the 1000 replications. Only bootstrap values >50% are shown.

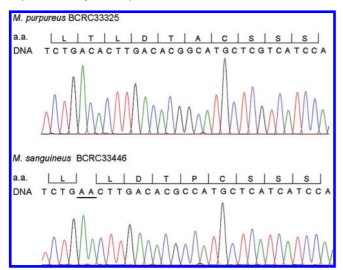


Figure 4. Sequence analyses of the *pksCT* KS domain. The deduced conserved amino acid sequence of the typical active site of the KS domain, DXACXS, was found in *M. purpureus* BCRC33325. A gap, CTGA-ACTT, was observed in the upstream of the *pksCT* KS domain of *M. sanguineus* BCRC33446. The sequencing of *M. sanguineus* BCRC33446 was carried out in triplicate.

The amplified DNA results also showed that the *ctnA* gene shared 100% similarity among *M. purpureus* BCRC31542, BCRC31541, BCRC33325, and BCRC31615, as well as *M. kaoliang* BCRC31506. The *ctnA* gene-encoded transcription factor has been suggested to be a positive regulatory protein involved in citrinin biosynthesis in *M. purpureus* BCRC33325 (IFO30873) (*13*). The arrangement of the cysteine-rich nucleotide-binding domain indicated that the consensus sequence CX₂CX₆CX₆CX₂CX₆C represented a Zn(II)2Cys6-type zinc finger (*13*).

A phylogenetic characterization using the partial β -tubulin gene as a molecular differentiation marker was likewise conducted to further explore the evolutionary history of the

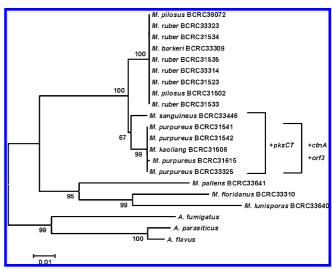


Figure 5. Phylogeny of the *Monascus* species based on the partial β -tubulin gene. The partial β -tubulin genes of the *Monascus* species with the following accession numbers were used: DQ299886—DQ299896, AY498587—AY498589, AY498596, AY498598, AY498601, AY498602, and AY498604. The accession numbers of the β -tubulin genes were used as outgroups of the following: *A. flavus* (M38265), *A. parasiticus* (L49386), and *A. fumigatus* (AY048754). Bootstrap values are shown in the nodes according to 1000 replications. Only bootstrap values >50% are shown.

various *Monascus* species. The result of the phylogenetic analysis of the partial β -tubulin gene placed both M. purpureus and M. kaoliang in the same clade (**Figure 5**). On the other hand, M. ruber, M. pilosus, and M. barkeri were placed in the same clade. In addition, M. sanguineus was placed in a branch separate from M. purpureus and M. kaoliang.

Analysis of Citrinin Production. In previous studies, the citrinin production in *M. purpureus* BCRC33325 (IFO30873) was shown to progress with the duration of cultivation (*3, 13*). In this study, the amounts of citrinin produced from *Monascus* spp. were determined by HPLC after 14 days cultivation. The results indicated that citrinin was only produced in *M. purpureus* BCRC31542, BCRC31541, BCRC33325, and BCRC31615 and *M. kaoliang* BCRC31506 (**Table 3**). In contrast, the other *Monascus* species did not produce any citrinin.

DISCUSSION

Citrinin is a known hepato-nephrotoxin found in the Aspergillus, Penicillium, and Monascus species, and has been identified as a contaminant in several foods (8, 9). Citrinin accumulation in the mitochondria induces apoptosis at the cellular level (17, 18). Monascus-related products have been used in herbal medicine because their monacolin K content shows the capability to inhibit cholesterol synthesis (1). To avoid the negative effects of citrinin, it is important to identify the non-citrinin-producing *Monascus* strains or to eliminate the production of citrinin in *Monascus*. Recently, the citrinin biosynthesis gene cluster has revealed that citrinin is synthesized by polyketide synthase in *M. purpureus* BCRC33325 (IFO30873) (3). Moreover, the transcription factor Zn(II)2Cys6 binuclear DNA binding protein is also involved in the regulation of citrinin biosynthesis (13). In this study, the distribution of the citrinin biosynthesis genes (Figure 1) and the production of citrinin were examined in various Monascus species.

Interestingly, the PCR and Southern hybridization results demonstrated that only *M. purpureus*, *M. kaoliang*, and *M. sanguineus* contained the *pksCT* gene encoding polyketide

Table 3. Concentration of Citrinin Produced by *Monascus* Species^a

		M. kaoliang:			
	BCRC31542	BCRC31541	BCRC33325	BCRC31615	BCRC31506
citrinin (mg/g mycelia) ^b	0.33 ± 0.14	15.35 ± 6.17	6.23 ± 1.74	6.56 ± 2.82	2.71 ± 0.28

a Monascus species were harvested after the cultivation of 14 days. b Citrinin was detected by HPLC with the fluorescence detector under the culture condition described in Materials and Methods.

synthase of citrinin (**Figure 2**). The sequences of the AT domain of these amplified pksCT genes verified that they were highly homologous with the known citrinin gene cluster in M. purpureus BCRC33325 (IFO30873) (Figure 3). However, a gap was found in the upstream region of the pksCT KS domain of M. sanguineus BCRC33446. This missing base could cause a frame shift that results in a nonfunctional polyketide synthase (Figure 4). In addition, M. sanguineus BCRC33446 did not contain the ctnA gene encoding Zn(II)2Cys6 binuclear DNA binding protein or the orf3 gene encoding oxygenase (Figure 2). The strain may have had a large deletion of the citrinin biosynthesis gene cluster initiated possibly by a chromosomal breakage, similar to Aspergillus oryzae RIB, which has a deletion in the aflatoxin biosynthesis gene cluster (19). The sequences of the 5' ctnA gene from M. purpureus and M. kaoliang were also verified, including the consensus sequence, CX₂CX₆CX₆CX₂CX₆C, which represents a Zn(II)2Cys6-type zinc finger. The citrinin-producing phenotype was detected in M. purpureus and M. kaoliang, whereas neither M. sanguineus BCRC33446 nor the other *Monascus* species produced citrinin. These results suggest that the citrinin gene cluster is highly conserved within M. purpureus and M. kaoliang, while the pksCT gene shows high homology in M. purpureus, M. kaoliang, and M. sanguineus. The citrinin production results were consistent with the distribution of the functional citrinin gene cluster, which was only detected in the strains of M. purpureus and M. kaoliang (**Table 1**).

Blanc et al. had initially proposed that the structure of monascidin A is identical to the citrinin from both the M. ruber ATCC96218 and M. purpureus CBS109.07 strains (10). The citrinin production of M. ruber ATCC96218 was further investigated for its biosynthetic pathway, effects of mediumchain fatty acids, and improvement of the red pigment/citrinin production ratio (12, 20, 21). Although previous studies have shown that M. ruber ATCC96218 can produce citrinin, this strain has unfortunately been misidentified and was corrected to M. purpureus (22, 23). Park et al. (22, 23) have indicated that the D1/D2 regions of the large subunit (LSU) rRNA gene and the β -tubulin gene could be used to examine the phylogenetic relationship between the *Monascus* species. Their findings are in agreement with our result that M. ruber and M. purpureus can be distinguished into two series (**Figure 5**). Moreover, the M. ruber series (M. ruber, M. pilosus, and M. barkeri) and the M. purpureus series (M. purpureus and M. kaoliang) can be grouped by determining the presence or absence of MRT non-LTR retrotransposon in the hybridization pattern, according to the phylogenetic study established with the β -tubulin gene (24). Hence, M. ruber ATCC96218 is identical to the M. purpureus series based on the D1/D2 region of the LSU rRNA gene (22). In addition, the strains of M. pilosus ATCC62949 and M. purpureus BCRC31523 (ATCC16378) were renamed as M. purpureus ATCC62949 and M. ruber BCRC31523 (ATCC16378), respectively (22). Citrinin is only produced in P. citrinum, from the examination of P. citrinum, Penicillium corylophilum, Penicillium steckii, Penicillium sizovae, and Penicillium sumatrense with 79 isolates (9). Citrinin has also been reported in Penicillium expansum and Penicillium verrucosum (8). These results show that citrinin is produced in certain species of Penicillium, which is quite similar to our findings that only the species of *M. purpureus* and *M. kaoliang* can produce citrinin.

Wang et al. have proposed that citrinin is detectable in each Monascus species under a low HPLC flow rate condition (0.15 $mL min^{-1}$) and when incubated in a YES medium (40 g L^{-1} of yeast extract and 160 g L^{-1} of sucrose) (25). However, these results may need to be reexamined, because we did not detect citrinin in some of the species listed in this paper. Moreover, the incubation and analysis conditions in the Wang et al. study may have led to false positive results. It is possible, but unlikely, that another divergent citrinin biosynthesis gene is present in the M. ruber series.

In conclusion, the citrinin production in Monascus is consistent with the presence of the functional citrinin biosynthesis genes found only in the M. purpureus series. Although M. sanguineus BCRC 33446 carries the pksCT gene, citrinin production was not detected due to a frame shift of pksCT and the absence of some other citrinin biosynthesis genes (Figures 2 and 4). The M. ruber, M. pilosus, and M. barkeri strains were grouped together on the basis of the classification of the β -tubulin gene. It was found that they did not possess the *pksCT*, ctnA, and orf3 genes and did not produce citrinin. Therefore, citrinin biosynthesis genes and citrinin production can be used for phylogenetic studies and in the exploration of gene evolution for secondary metabolism.

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