

Phylogenetic Analyses Reveal Monophyletic Origin of the Ergot Alkaloid Gene *dmaW* in Fungi

Miao Liu,^{1,3} Daniel G. Panaccione² and Christopher L. Schardl¹

¹201 F Plant Science Bldg. University of Kentucky, Lexington, KY 40546. ²Division of Plant and Soil Sciences, West Virginia University, PO Box 6108, Morgantown, WV 26506-6108. ³Biodiversity (Mycology and Botany), Eastern Cereal and Oilseed Research Centre, Agriculture and Agri-Food Canada, 960 Carling Avenue, Ottawa, Ontario K1A 0C6. Email: schardl@email.uky.edu.

Abstract: Ergot alkaloids are indole-derived mycotoxins that are important in agriculture and medicine. Ergot alkaloids are produced by a few representatives of two distantly related fungal lineages, the Clavicipitaceae and the Trichocomaceae. Comparison of the ergot alkaloid gene clusters from these two lineages revealed differences in the relative positions and orientations of several genes. The question arose: is ergot alkaloid biosynthetic capability from a common origin? We used a molecular phylogenetic approach to gain insights into the evolution of ergot alkaloid biosynthesis. The 4- γ , γ -dimethylallyltryptophan synthase gene, *dmaW*, encodes the first step in the pathway. Amino acid sequences deduced from *dmaW* and homologs were submitted to phylogenetic analysis, and the results indicated that *dmaW* of *Aspergillus fumigatus* (mitosporic Trichocomaceae) has the same origin as corresponding genes from clavicipitaceous fungi. Relationships of authentic *dmaW* genes suggest that they originated from multiple gene duplications with subsequent losses of original or duplicate versions in some lineages.

Keywords: mycotoxins, 4- γ , γ -dimethylallyltryptophan synthase gene, Clavicipitaceae, Trichocomaceae, amino acid sequence, GBLOCKS, gene phylogeny

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Introduction

Ergot alkaloids (EA) are a group of mycotoxins causing toxicoses in humans and animals.^{1–3} Producers of EA include plant pathogens in the genus *Claviceps*, and some grass endophytes in the genera *Epichloë*, *Neotyphodium* and *Balansia*.² These genera belong to the family Clavicipitaceae (order Hypocreales, phylum Ascomycota). EA are also produced by the common airborne fungus *Aspergillus fumigatus*, a species distantly related with clavicipitaceous fungi, belonging to the family Trichocomaceae (order Eurotiales, phylum Ascomycota).⁴ In the same family, some *Penicillium* species also produce EA.^{5,6} Relatively few species within these families produce ergot alkaloids and only one representative of the lineages in between the two orders containing these families has been reported to produce ergot alkaloids.^{7,8} Because of their significant impacts on human health and agriculture, the biochemistry and biosynthesis pathway of ergot alkaloids have attracted much attention and effort.³ Certain genes involved in the biosynthesis pathway have been characterized.^{9–15} Clustered arrangements of EA biosynthesis genes have been observed in *A. fumigatus*,¹⁰ *Claviceps fusiformis*,⁸ and *Claviceps purpurea*.¹⁴ A gene, *dmaW*, common to all these clusters, encodes 4- γ,γ -dimethylallyltryptophan (DMATrp) synthase, which has been demonstrated to be the first pathway specific step and has the key regulatory function for EA biosyntheses in *Claviceps* spp.^{15–17} Following the cloning of *dmaW* from *C. fusiformis* SD58,¹⁸ a gene cluster likely encoding enzymes of EA biosynthesis was identified by sequence analysis in the ergot fungus, *C. purpurea* P1.¹⁴ This cluster was thereafter designated *EAS* (ergot alkaloid synthesis).² Within the *EAS* cluster, four genes with known functions were named according to their functions, and the other seven genes with unknown functions were named *easA* and *easC* to *easH*. Homologs of nine *EAS*-cluster genes have been found in *C. fusiformis* SD58,^{18,19} and eight homologs have been found clustered in the *A. fumigatus* genome¹⁰ (Supplementary Fig. 1).

Based on the sequence similarity of *dmaW* genes and general clustering of other hypothetical *EAS* genes, Coyle and Panaccione¹⁰ proposed that EA biosynthesis in *A. fumigatus* has a common origin with that in clavicipitaceous fungi. However, arrangement of the cluster genes in *A. fumigatus* is drastically

different from the *EAS* clusters in *C. purpurea* and *C. fusiformis*.^{2,8} This difference in gene arrangements, and the absence of ergot alkaloids in the lineages between the two groups of EA-producing fungi, raise the question of whether the *dmaW* genes in clavicipitaceous and trichocomaceous fungi are true orthologs, homologs due to speciation. If the genes are truly orthologous, then multiple gene recombinations and inversions must have happened in either (or both) of the lineages after the divergence from their common origin, and the lineages between them have lost these genes. The alternative would be that the *A. fumigatus* *EAS* genes might have evolved independently from different gene duplication events.

Recently, multiple genes similar or related to *dmaW* have been characterized by cloning and over-expression approaches. Examples include genes for brevianamide F prenyl transferase (FtmPT1) that converts brevianamide to tryprostatin in *A. fumigatus*,²⁰ reverse prenyl transferase FGAPT1 catalyzing the final step in the biosynthesis of fumigaclavine C in the same fungus,²¹ TdiB with indole alkaloid biosynthetic ability in *A. nidulans*,²² as well as *SirD*, involved in the biosynthesis of an epipolythiodioxopiperazine (ETP) from *Leptosphaeria maculans*.²³ The available sequences of these genes allow us to look into the evolutionary relationships of *dmaW* and related genes by a phylogenetic approach.

In this study, we conduct molecular phylogenetic analysis of inferred protein sequences to test whether the known DMATrp synthases in fungi have a single origin or multiple origins. A monophyletic pattern is expected for the single origin hypotheses; whereas a polyphyletic pattern is expected for multiple origins.

Materials and Methods

Protein sequences derived from *dmaW* genes of clavicipitaceous fungi and *A. fumigatus* were obtained from previous studies.^{10,15,18} Protein sequences of FtmPT1, FGAPT1, and the *sirD* product were downloaded from GenBank (Table 1). The protein sequences from *dmaW* homologs were obtained by BLAST of the nonredundant protein sequence database in GenBank with the protein product deduced from *dmaW* from *Neotyphodium lolii* and *A. fumigatus*; sequences with relatively high similarity scores and low E-value ($<3e^{-17}$) were selected (Table 1).



Protein sequence matrix

Due to the high divergence of amino acid sequences, we used the program MAFFT ver.5.8²⁴ to align them. The alignment was conducted through the web server (<http://align.genome.jp/mafft/>). The FFT-NS-i and E-INS-i alignment strategies, iterative refinement method, were used to enhance the accuracy of the alignment. The scoring matrix (for amino acid sequences) was selected from six options by comparing fitnesses of the trees in the preliminary analysis. With the scoring matrix selected, we set gap opening penalty (OP = 1.0, 2.0, 3.0) and gap extension penalty (offset value as shown in the program, OF = 0.0, 0.5, 1.0), and again compared the overall fitness of the resulting trees in more preliminary analyses. Parameters resulting in trees of high fitness were set in the final alignment. The alignments by MAFFT were submitted to the program Gblocks 0.91b,²⁵ to eliminate the highly diverged regions and retain the conserved regions for phylogenetic analysis. Low stringency options were selected to obtain blocks.

Phylogenetic analysis

The protein matrices of ten operational taxonomic units (OTUs, six authentic *dmaW* and four related genes of known functions) resulting from MAFFT and Gblocks screening and comprised of the conserved regions of the protein alignment, were submitted separately to phylogenetic analysis in PAUP* 4.0b10.²⁶ Parsimony analyses were conducted using exhaustive search. All characters had equal weight and gaps were treated as missing data.

In order to exclude the possible bias caused by insufficient OTUs, we included multiple potential homologs of *dmaW* products, which were obtained by BLAST search, in a more extensive phylogenetic analysis. Both conserved protein regions and the whole protein sequence were used in the separate analyses. Parsimony analysis was conducted with a heuristic search with TBR (tree bisection and reconnection) branch-swapping and 100 replicates of random sequence addition. Bootstrapping analysis was based on 1000 replicates of a full heuristic search, each with 20 replicates of a random addition sequence, and tree bisection reconnection (TBR) swapping was selected and re-arrange limit was set to 5000 per replicate.

Bayesian inferences (BI) were performed using MrBayes 3.0B5²⁷ to analyze individual data sets.

The prior for amino acid model was set as mixed to allow model jumping between fixed-rate amino acid models. Maximum likelihood analysis (ML) was performed using PHYML online server²⁸ with the following settings: substitution model as JTT, transition/transversion ratio and gamma distribution as estimated by the program, bootstrap datasets 500.

Results

Protein sequence matrix

To choose the appropriate scoring matrix in the amino acid sequence alignment, we compared the overall fitness of the trees based on the alignments of six different scoring matrices in two strategies (Table 2). Setting scoring matrix as JTT200 resulted in the highest consistency index (CI), lowest homoplasy index (HI), relatively high retention index (RI), and shorter trees, thus we choose JTT200 for alignments. In the combination of various OP and OF in the alignment, OP/OF = 1.0/0.0 resulted in three shortest parsimonious trees with a relatively high CI, RI and low HI (Table 3).

For the protein sequences derived from six *dmaW* and four related genes of known functions, an analysis in which the scoring matrix was set as JTT200, OP as 1.0, and OF as 0.0 resulted in an alignment with 653 characters. The resulting alignment was put into Gblocks 0.91b to screen for conserved regions. The number of characters (amino acid positions) retained was 308. For the matrix from the extended OTU set (34 protein sequences), 933 characters resulted from MAFFT alignment, and 155 characters were retained after GBlocks screening.

Phylogenetic relationships

For the data set comprised of ten OTUs, parsimony analyses for the whole regions resulted in two most parsimonious trees (Fig. 1A). The two tree topologies differed in the order of branches ranches to *sirD* *L. maculans* and *fgaPTI* *A. fumigatus*. The analysis of the conserved regions resulted in one most parsimonious tree. The tree topology differed from those of whole gene regions in the order of divergence of *dmaW* of *A. fumigatus* and *Malbranchea aurantiaca* (Fig. 1B). The known, authentic sequences of DMATrp synthases formed a monophyletic clade with strong bootstrap support. Defining as an outgroup those prenyl transferases known or likely to catalyze production of

**Table 1.** GenBank numbers and fungal isolates of protein sequences of *dmaW* genes and homologs.

	GenBank number	Fungal species	Isolate
1	AAX56314	<i>Aspergillus fumigatus</i>	Af293
2	EAL94098	<i>Aspergillus fumigatus</i>	Af293
3	AAX08549	<i>Aspergillus fumigatus</i>	Af293
4	XP_754328	<i>Aspergillus fumigatus</i>	Af293
5	ABU51603	<i>Aspergillus nidulans</i>	unspecified
6	XP_681751	<i>Aspergillus nidulans</i>	FGSC A4
7	XP_682498	<i>Aspergillus nidulans</i>	FGSC A4
8	XP_659949	<i>Aspergillus nidulans</i>	FGSC A4
9	XP_664388	<i>Aspergillus nidulans</i>	FGSC A4
10	BAE56055	<i>Aspergillus oryzae</i>	RIB 40
11	BAE59503	<i>Aspergillus oryzae</i>	RIB 40
12	BAE62662	<i>Aspergillus oryzae</i>	RIB 40
13	BAE65189	<i>Aspergillus oryzae</i>	RIB 4
14	BAE63695	<i>Aspergillus oryzae</i>	RIB 4
15	AAP92451	<i>Balansia obtecta</i>	B249
16	AAC18893	<i>Claviceps fusiformis</i>	ATCC26245
17	CAB39314	<i>Claviceps purpurea</i>	P1
18	CAC37396	<i>Claviceps purpurea</i>	T5
19	AAZ29613	Clavicipitaceae	US2005a
20	AAZ29614	Clavicipitaceae	US2005b
21	EAS31821	<i>Coccidioides immitis</i>	RS
22	AAP81206	<i>Epichloë typhina</i> × <i>Neotyphodium lolii</i>	Lp1
23	AAV66102	<i>Fusarium heterosporum</i>	ATCC74349
24	AAS92554	<i>Leptosphaeria maculans</i>	ICBN 18
25	XP_361876	<i>Magnaporthe grisea</i>	70-15
26	XP_370025	<i>Magnaporthe grisea</i>	70-15
27	ABZ80612	<i>Malbranchea aurantiaca</i>	RRC1813
28	AAP81207	<i>Neotyphodium coenophialum</i>	ATCC90664
29	AAP81208	<i>Neotyphodium coenophialum</i>	ATCC90664
30	none	<i>Neotyphodium gansuense</i>	E818
31	XP_960156	<i>Neurospora crassa</i>	OR74A
32	AAZ29615	<i>Penicillium roqueforti</i>	IsaF09
33	AAK11526	<i>Penicillium paxilli</i>	ATCC26601
34	AAT69743	<i>Sirodesmium diversum</i>	ATCC36539

other products (4 OTUs), the most basal divergence separated the *dmaW* gene of *M. aurantiaca* and *A. fumigatus* from those of the Clavicipitaceae (Fig. 1C).

The analysis with conserved gene regions comprised of 34 OTUs resulted in six equally most parsimonious

trees, while the whole gene region (933 characters) resulted in eight equally most parsimonious trees. The variations of branch orders among these trees were mainly from the uncertain positions of the two OTUs, the putative DMATrp synthase gene from



Gene, Function	Reference
<i>ftmPT1</i> = brevianamide F prenyl transferase	(Grundmann, Li) ²¹
<i>fgaPT1</i> = easL, reverse prenyl transferase	(Unsold, Li) ²²
<i>fgaPT2</i> = <i>dmaW</i> , dimethylallyltryptophan synthase function unknown	(Unsold, Li) ⁴⁰ (Nierman et al) ⁴¹
<i>tdiB</i> = putative prenyl transferase function unknown	(Balibar) ⁴² (Galagan et al) ⁴³
function unknown	(Galagan et al) ⁴³
function unknown	(Galagan et al) ⁴³
function unknown	(Galagan et al) ⁴³
function unknown	(Machida et al) ³¹
function unknown	(Machida et al) ³¹
function unknown	(Machida et al) ³¹
function unknown	(Machida et al) ³¹
function unknown	(Machida et al) ³¹
<i>dmaW</i> = dimethylallyltryptophan synthase	(Wang et al) ¹⁵
<i>dmaW</i> = dimethylallyltryptophan synthase	(Tsai et al) ¹⁹
<i>dmaW</i> = <i>cpd1</i> , dimethylallyltryptophan synthase allele 1	(Tudzynski et al) ¹⁴
<i>dmaW</i> = <i>cpd2</i> , putative dimethylallyltryptophan synthase allele 2	(Arntz and Tudzynski, unpublished)
<i>dmaW</i> = putative dimethylallyltryptophan synthase	(Steiner et al) ³²
<i>dmaW</i> = putative dimethylallyltryptophan synthase function unknown	(Markert et al) ⁴³ (Birren et al, unpublished)
<i>dmaW</i> = dimethylallyltryptophan synthase function unknown	(Wang et al) ¹⁵ (Sims et al) ⁴⁴
<i>sirD</i> = putative tyrosine dimethylallyl transferase function unknown	(Gardiner et al) ²⁴ (Dean et al) ⁴⁵
function unknown	(Dean et al) ⁴⁶
<i>MaPT</i> = <i>dmaW</i> , dimethylallyltryptophan synthase	(Ding et al) ⁴⁷
<i>dmaW</i> = dimethylallyltryptophan synthase allele 1	(Wang et al) ¹⁵
<i>dmaW</i> = dimethylallyltryptophan synthase allele 2	(Wang et al) ¹⁵
<i>dmaW</i> = putative dimethylallyltryptophan synthase function unknown	Unpublished data (Galagan et al) ⁴⁸
<i>dmaW</i> = putative dimethylallyltryptophan synthase	(Steiner et al) ⁴⁹
<i>paxD</i> = function unknown	(Young et al) ³⁰
putative dimethylallyltyrosine synthase	Gardiner and Howlett, unpublished

Neotyphodium gansuense and *paxD* *Penicillium paxilli* AAK11526, which were partial sequences. The strict consensus trees of the six trees from conserved regions and eight trees from the whole gene regions showed the same pattern that, in the *dmaW* clade,

the two putative DMATrp synthase genes from clavicipitaceous endophyte of convolvulaceous plants (AAZ29613, AAZ29614), the clade comprised of *C. purpurea* and *C. fusiformis dmaW*, as well as the putative DMATrp synthase gene from *N. gansuense*

**Table 2.** Fitness of the trees resulting from maximum parsimony searches with different matrix scoring settings.

Strategy	Scoring matrix	Number of trees	Length	CI	RI	RC	HI	G-fit
FFT-NS-i	BLOSUM30	1	5252	0.653	0.491	0.32	0.347	-224.198
	BLOSUM45	4	5214	0.64	0.486	0.311	0.36	-219.496
	BLOSUM62	6	5230	0.637	0.491	0.313	0.363	-208.403
	BLOSUM80	8	5170	0.638	0.487	0.31	0.362	-207.579
	JTT100	5	5187	0.646	0.494	0.319	0.354	-212.253
	JTT200	2	5193	0.662	0.491	0.325	0.338	-235.767
E-INS-i	BLOSUM30	1	5200	0.636	0.486	0.309	0.364	-211.530
	BLOSUM45	6	5185	0.636	0.491	0.312	0.364	-203.114
	BLOSUM62	6	5190	0.637	0.489	0.312	0.363	-205.306
	BLOSUM80	1	5203	0.635	0.488	0.310	0.365	-204.704
	JTT100	2	5229	0.631	0.483	0.305	0.369	-206.307
	JTT200	3	5174	0.632	0.486	0.307	0.368	-204.534

collapsed as a polytomy (Fig. 2A). Outside the *dmaW* clade, *paxD* *P. paxilli* AAK11526 along with the other six OTUs appeared as unresolved branches (Fig. 2A). All trees revealed the same monophyletic group of DMATrp synthase genes (functionally tested) and putative DMATrp synthase genes.

Excluding the two partial sequences (32 OTUs included in analysis) resulted in better resolutions. Both analyses of conserved regions and the whole gene region resulted in two most parsimonious trees. The two trees from whole gene regions differed in the positions of *Magnaporthe oryzae* XP 361876, clade I

(*tdiB* *Aspergillus nidulans* ABU51603, *Neurospora crassa* XP960156 and *Magnaporthe oryzae* XP 370025) and clade II (*fgaPT1* *Aspergillus fumigatus* EAL94098, *Aspergillus oryzae* BAE65189, and *Aspergillus fumigatus* XP 754328) (Fig. 2B). Comparing the phylogenies inferred from the conserved regions and from the entire sequences, within the *dmaW* clade, these trees differed in branching orders of *B. obtecta*, *C. purpurea* and *C. fusiformis dmaW*; and in branch orders of *A. fumigatus* AAX08549, *P. roquefortii* AAZ29615 and *Malbranchea aurantiaca* ABZ80612. All clades with strong statistic support (bootstrapping value >70)

Table 3. Fitness of the trees resulting from maximum parsimony searches with different gap opening penalties (OP) and gap extension penalties (OF) (FFT-NS-i, JTT200).

OP	OF	Number of trees	Length	CI	RI	RC	HI	G-fit
1.0	0	3	5098	0.675	0.498	0.336	0.325	-245.815
1.0	0.5	1	5219	0.628	0.483	0.303	0.372	-206.925
1.0	1	1	5217	0.625	0.479	0.3	0.375	-197.856
2.0	0	11	5227	0.649	0.489	0.317	0.351	-226.988
2.0	0.5	4	5281	0.632	0.485	0.306	0.368	-206.599
2.0	1	1	5272	0.637	0.489	0.312	0.363	-204.882
3.0	0	7	5354	0.658	0.491	0.323	0.342	-239.881
3.0	0.5	4	5280	0.64	0.49	0.314	0.36	-210.476
3.0	1	12	5317	0.639	0.483	0.308	0.361	-206.114

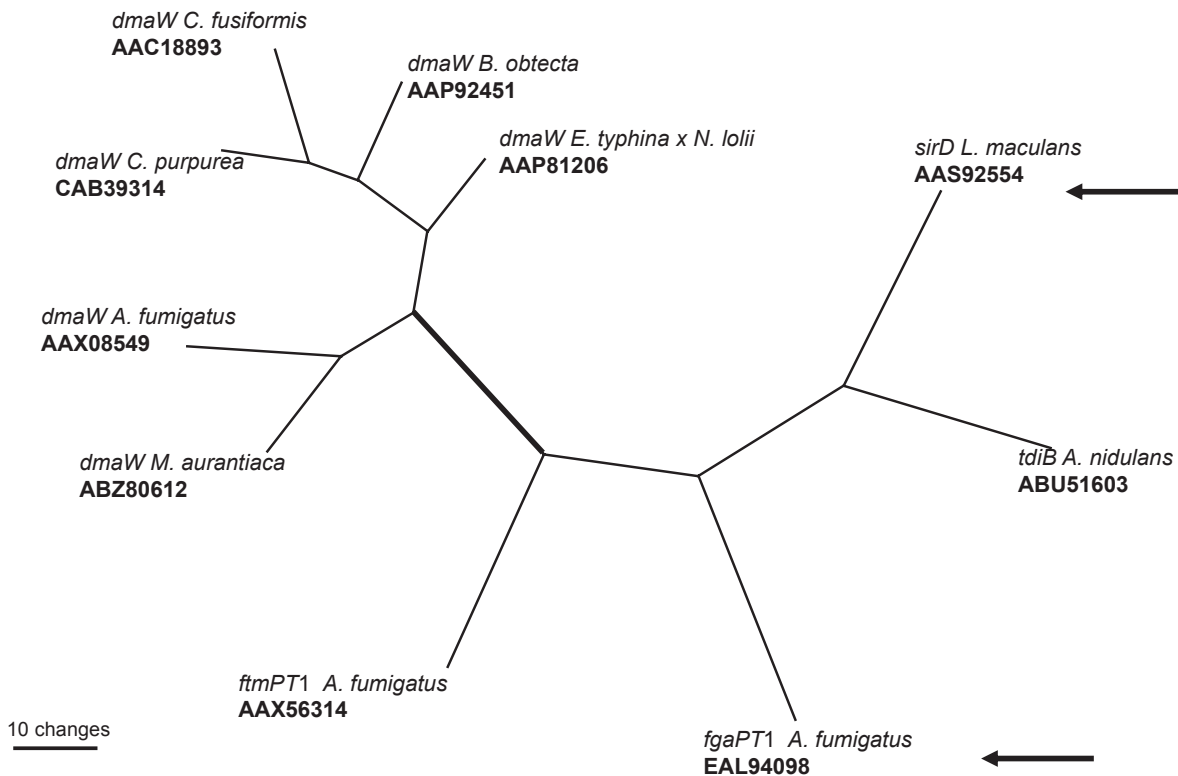
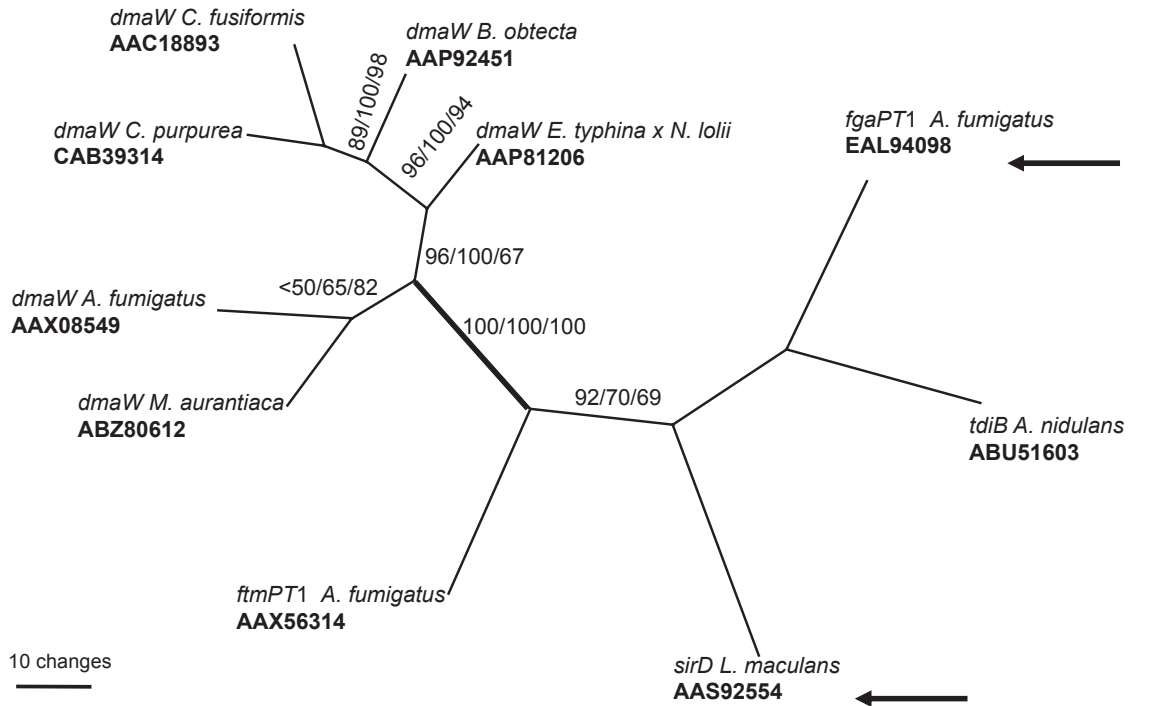


were present in all trees (Figs. 2B, C), and all trees clearly indicated monophyly of genes for authentic DMATrp synthase.

Bayesian analyses generally resulted in higher statistical support (posterior probabilities) for the

clades having high bootstrap support in parsimony analyses (Figs. 1 and 2) and internal branches. ML analyses resulted tree topologies generally congruent with MP tree, i.e. clades with strong support were congruent with MP (Figs. 1 and 2).

1A



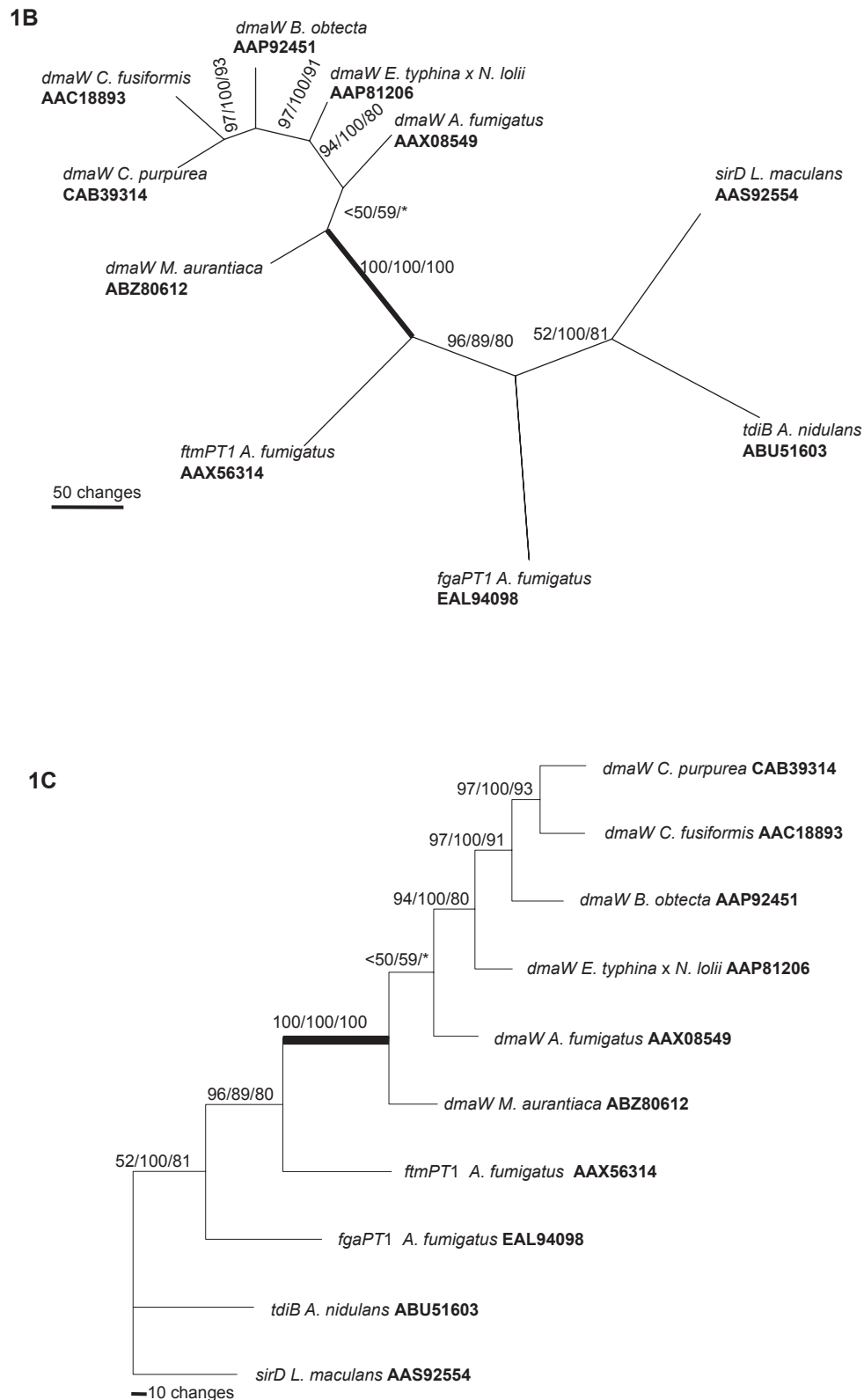


Figure 1. The most parsimonious trees relating amino acid sequences of the deduced products of six *dmaW* genes and four related genes with known functions. **A)** Tree based on whole gene region. Of 653 aligned characters, 250 characters are informative. Length = 1521, CI = 0.870, RI = 0.503. Arrows show the discrepancies of the two most parsimonious trees. **B and C)** Trees based on relatively conserved gene regions screened by GBlocks. Of 308 total aligned characters, 166 characters are informative. Length = 934, CI = 0.864, RI = 0.512. Products of *dmaW* genes formed a monophyletic group with 100% bootstrap support. **B.** Unrooted tree; **C.** Phylogram rooted by choosing the non-*dmaW* gene products as the outgroup. Numbers on branches indicate bootstrap percentage of MP/posterior probabilities of BI/bootstrap percentage of ML; *indicates that particular branch does not exist in the analysis. The thick branch separates of the outgroup from the ingroup.



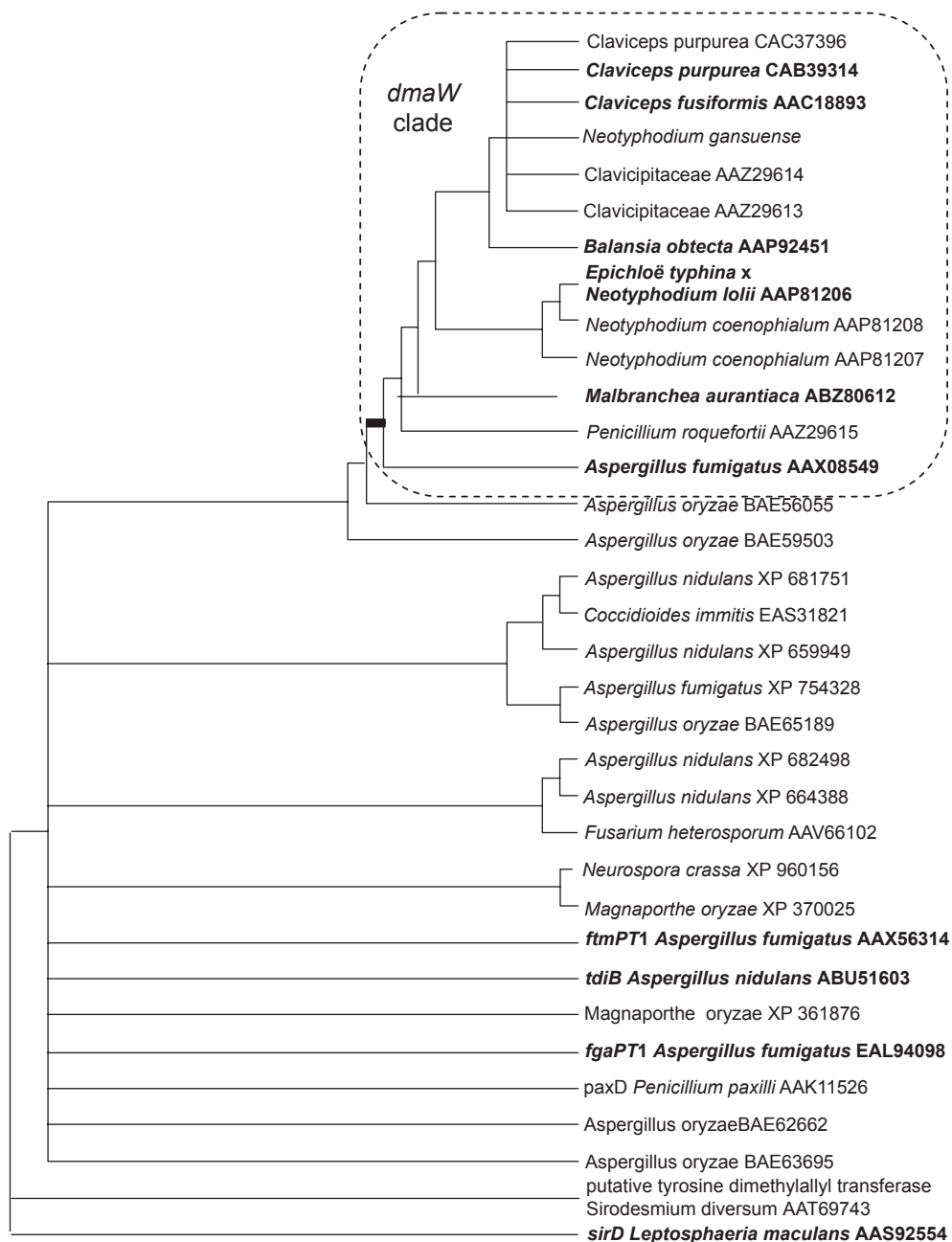
All analyses indicated that conserved gene regions selected by GBlocks did not significantly improve the phylogenetic inference for our data sets.

Genes associated with *dmaW*-related sequences

Certain *dmaW*-related genes have been identified in gene clusters that are otherwise unrelated to

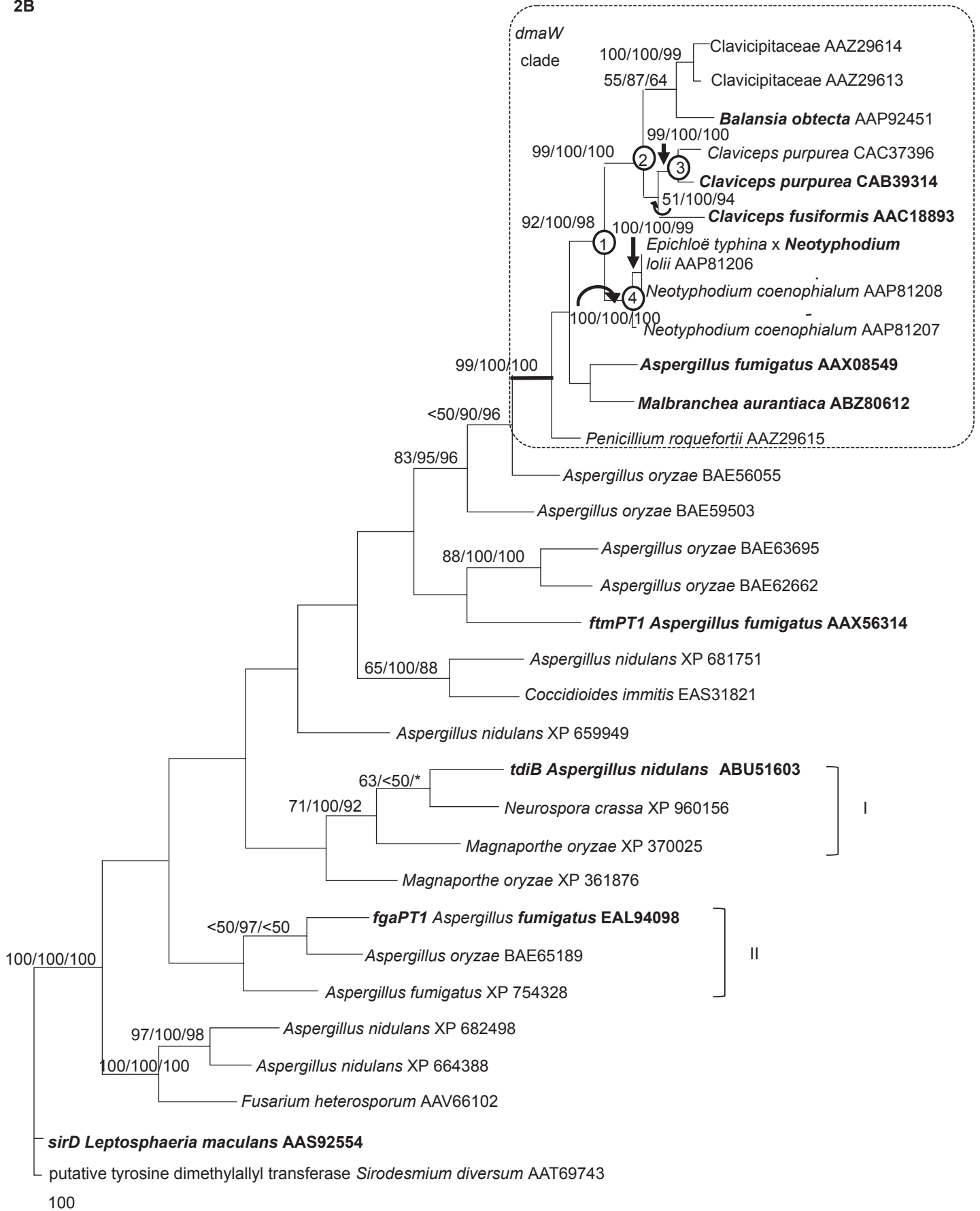
the *EAS* clusters. These include the *sirD* gene of *L. maculans*²³ and the *paxD* gene of *Penicillium paxilli*.²⁹ Because of the close relationship of two *A. oryzae* homologs, and the availability of a complete genome sequence for that fungus,³⁰ we checked the genomic contexts of these *A. oryzae* homologs. Neither of them has a comparable gene cluster. In both cases there are a few apparent secondary

2A





2B





2C

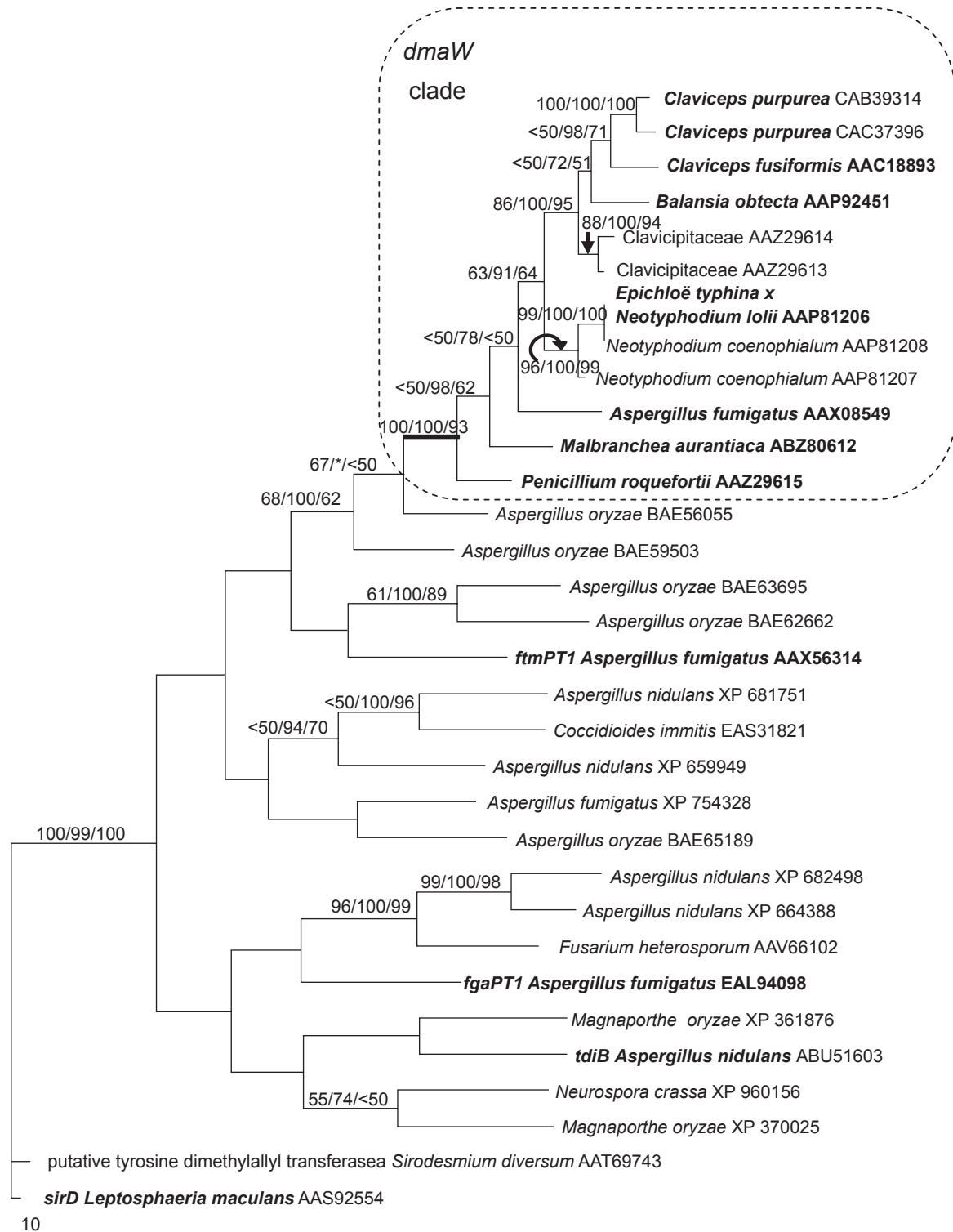


Figure 2. A) Strict consensus tree of the six most parsimonious trees for *dmaW* gene products and products of multiple related genes of 34 OTUs based on relatively conserved gene regions. Of the 155 characters, 133 were parsimony-informative; length = 1366, CI = 0.594, RI = 0.499. **B)** One of the two most parsimonious trees of 32 OTUs based on the whole gene region. Of the 933 characters, 456 were parsimony-informative characters. Length = 4974, CI = 0.682, RI = 0.499. **C)** The most parsimonious trees for *dmaW* gene products and products of multiple related genes of 32 OTUs based on relatively conserved gene regions, which were screened by Gblocks. Total aligned characters = 155, informative characters = 135; length = 1427, CI = 0.607, RI = 0.497. OTUs in bold indicate the gene products with functions that have been confirmed. Numbers on branches indicate bootstrap percentage of MP/posterior probabilities of BI/bootstrapped percentage of ML; *indicates that particular branch does not exist in the analysis; numbers in circles indicate the possible gene duplication events (also see Fig. 3).



metabolism genes nearby, but not homologs of the *EAS* cluster genes.

Discussion

Various profiles of ergot alkaloids (EA) are produced by EA-producing fungi. Evidence for diversification of EA profiles within an individual fungus, as well as among different producers was observed by Panaccione.⁷ The study reported herein is a molecular phylogenetic approach to gain additional insight into the evolution of EA synthesis among fungal producers. Due to its important role in encoding the first step in EA biosynthesis, *dmaW* was used as a marker to infer the evolutionary relationships of the pathways. Our results demonstrate that *dmaW* from *A. fumigatus* and clavicipitaceous fungi formed a monophyletic group indicating that they evolved from a common origin. Therefore we postulate that the *EAS* gene clusters of the two lineages were also from a common origin, which could be a common gene cluster encoding the shared early steps of the pathways of these two lineages. These shared steps might have been present in the most recent common ancestor of the two fungal lineages.

When 34 OTUs were included in the analysis, a hypothetical gene from *P. roqueforti* (GenBank accession number: AAZ29615.³¹ was grouped in the *dmaW* clade. *Penicillium roqueforti* produces the ergot alkaloid isofumigaclavine A,⁶ which would require *dmaW* for its biosynthesis. The product of the *P. roqueforti* gene was 63% identical with the product of *dmaW* of *A. fumigatus*, which was the top match retrieved in a BLAST search with this protein ($1e^{-112}$). These data are consistent with the *P. roqueforti* gene encoding the DMATrp synthase that catalyzes the initial prenylation in ergot alkaloid biosynthesis.

GenBank entries often are annotated completely on the basis of BLAST hits. Since the gene from *C. fusiformis* was identified as encoding DMATrp synthase,¹⁸ many sequences from other species were annotated as putative dimethylallyltryptophan synthase genes according to the similarity of their sequences to the *C. fusiformis* sequence. However many sequences annotated as dimethylallyltryptophan synthase genes are likely to encode related but nonidentical enzymes catalyzing prenylation or reverse prenylation of different co-substrates or at different positions of the indole rings. Examples of

related but functionally different prenyl transferases include the likely tyrosine prenyl transferase in sirodesmin biosynthesis from *L. maculans*,²³ and the reverse prenyl transferase from *A. fumigatus*.²¹

The rooted tree relating prenyl transferases with known functions showed the most basal separation of DMATrp synthase of *A. fumigatus* with those of the clavicipitaceous fungi. The EA profile of *A. fumigatus* includes a series of clavines, simpler tricyclic or tetracyclic alkaloids. In contrast, clavicipitaceous fungi usually produce more complex EA, ergopeptines and other amides of lysergic acid, in addition to the clavines. It is reasonable that more complex functions were gained along the evolutionary path.

Differences in the arrangement of *EAS* clusters between *A. fumigatus* and clavicipitaceous fungi were likely caused by multiple gene rearrangements through recombinations, deletions and insertions. The *EAS* gene cluster of *A. fumigatus* is in a subtelomeric region.¹⁰ Frequent recombinations associated with such regions provide a potential explanation for the differences between the *EAS* clusters of *A. fumigatus* and those of the clavicipitaceous fungi.⁷ The chromosomal locations of the clusters in clavicipitaceous species have not yet been determined. Similar rearrangements between more distant genomic locations would account for evolution of clusters, for which selection may favor their inheritance or horizontal transfer as a unit.^{32,33} Such rearrangements can be driven by repeats such as retroelements in the fungal genomes, such as observed throughout the ergot alkaloid and lolitrem biosynthesis gene clusters in *Epichloë festucae* and *Neotyphodium lolii*.^{34,35}

In fungal systematics, the molecular phylogeny indicates that genus *Claviceps* is more closely related to genus *Epichloë* (asexual stage: *Neotyphodium*) than to *Balansia*,^{36,37} (also see Fig. 3A). In our 10 OTU gene trees, the clade of *C. purpurea* and *C. fusiformis dmaW* was closer to *B. obtecta dmaW* than to *Epichloë* spp. *dmaW* (Fig. 1). The discrepancy between gene tree and species tree can be explained by polymorphic lineage sorting or incomplete sampling.³⁸ Once we included more OTUs (34 OTUs and 32 OTUs) in the analyses, the *dmaW* clade separated into two lineages. One lineage was comprised of *C. purpurea* and *C. fusiformis dmaW*, the two putative DMATrp synthase genes from clavicipitaceous endophytes of convolvulaceous plants (AAZ29613, AAZ29614),³¹



the putative DMATrp synthase gene from *N. gansuense*, and the DMATrp synthase gene from *B. obtecta*. The second lineage was comprised of putative DMATrp synthase genes from *N. coenophialum* and *dmaW* from the *E. typhina* × *N. lolii* hybrid. The divergence of these two lineages and the inconsistency of the divergence pattern with species relationships suggest that *dmaW* genes in clavicipitaceous fungi have experienced multiple gene duplications and loss of some copies. An alternative is that there may have been some instances of horizontal gene transfer, but the data are not conclusive in this respect. A scenario involving duplications and losses consistent with the

evolutionary relationships of authentic *dmaW* genes is shown in Figure 3B.

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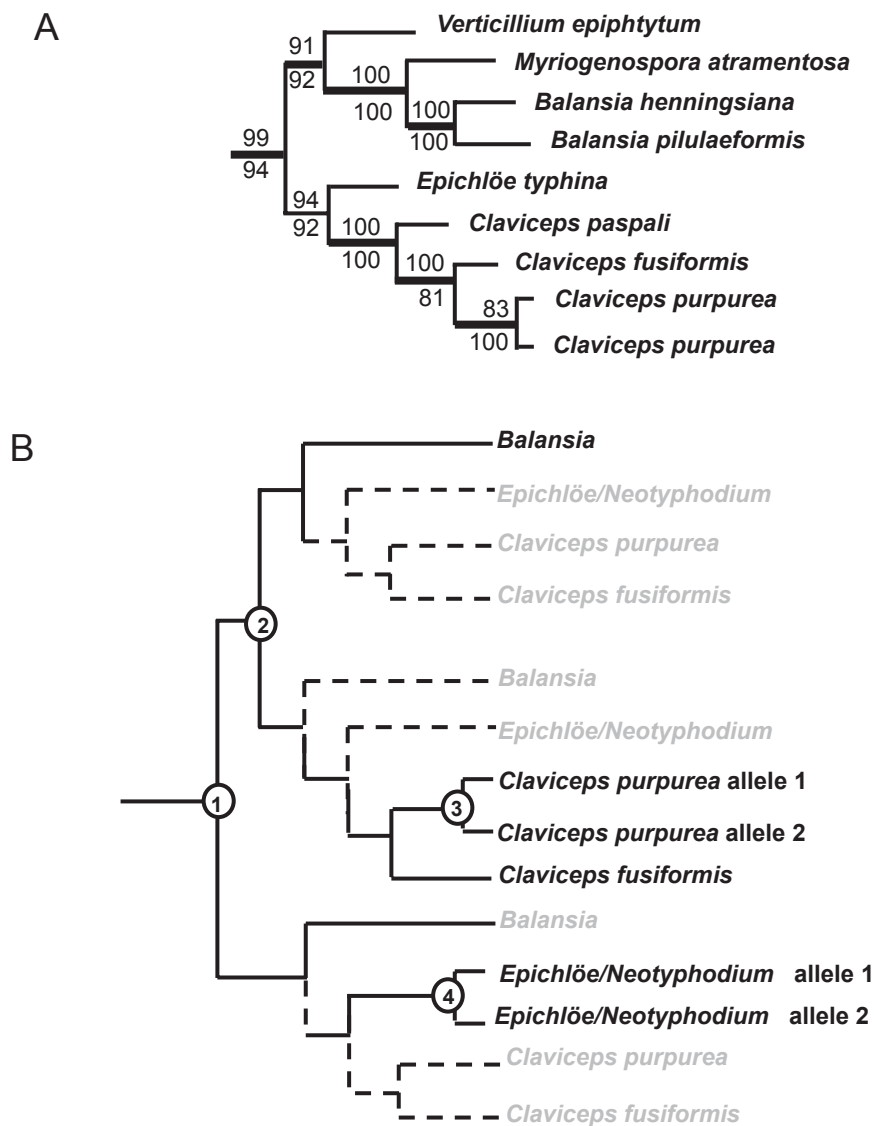


Figure 3. A) Partial phylogenetic relationships of species in Clavicipitaceae inferred by MP based on seven gene regions (Tree redrawn from the data of Sung et al).³⁷ **B)** Diagram indicating the possible gene duplication events that apparently happened in clavicipitaceous *dmaW* lineages. Numbers in circles indicate the possible gene duplication events. Dashed lines and shaded OTUs indicate the lost lineages (orthologs).



Disclosure

The authors report no conflicts of interest.

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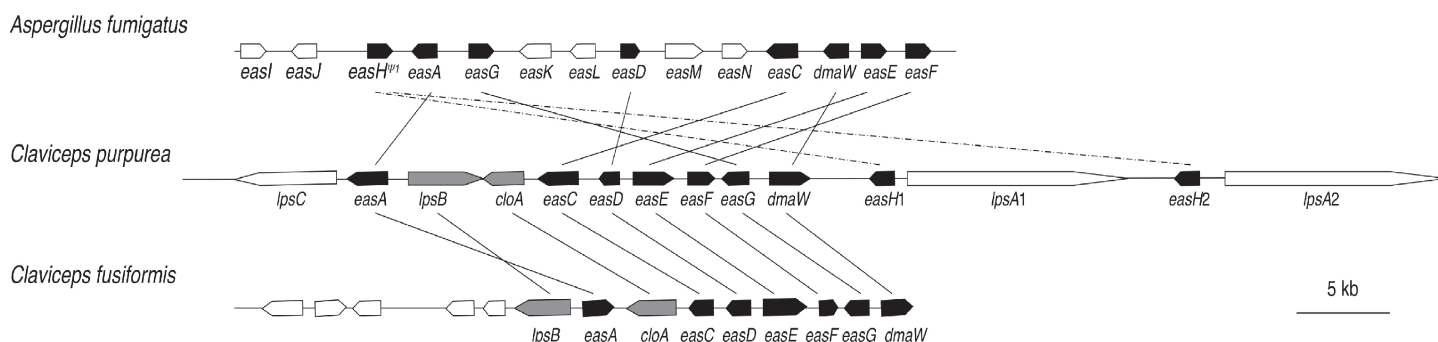


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Phylogenetic Analyses Reveal Monophyletic Origin of the Ergot Alkaloid Gene *dmaW* in Fungi

Miao Liu, Daniel G. Panaccione and Christopher L. Schardl

Supplementary Material



Supplement Figure 1. Map of EAS clusters from *Claviceps purpurea* strain P1,¹⁴ *Claviceps fusiformis* strain SD58,⁸ and *Aspergillus fumigatus*.¹⁰ Arrows indicate the directions of transcription. Black arrows indicate the genes shared in three species; shaded arrows indicate the two genes shared by *C. purpurea* and *C. fusiformis* but not *A. fumigatus*. Lines between maps connect orthologs, and dashed lines indicate the relationships of two *easH* homologues in *C. purpurea* P1 with an apparent pseudogene in *A. fumigatus*.

Supplement Figure 2. Protein sequence alignment screened by GBLOCKS 0.91b. The selected 155 positions are underlined in blue. 

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