Chromosome-level genome map provides insights into diverse defense mechanisms in the medicinal fungus *Ganoderma sinense*

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Fungi have evolved powerful genomic and chemical defense systems to protect themselves against genetic destabilization and other organisms. However, the precise molecular basis involved in fungal defense remain largely unknown in Basidiomycetes. Here the complete genome sequence, as well as DNA methylation patterns and small RNA transcriptomes, was analyzed to provide a holistic overview of secondary metabolism and defense processes in the model medicinal fungus, *Ganoderma sinense*. We reported the 48.96 Mb genome sequence of *G. sinense*, consisting of 12 chromosomes and encoding 15,688 genes. More than thirty gene clusters involved in the biosynthesis of secondary metabolites, as well as a large array of genes responsible for their transport and regulation were highlighted. In addition, components of genome defense mechanisms, namely repeat-induced point mutation (RIP), DNA methylation and small RNA-mediated gene silencing, were revealed in *G. sinense*. Systematic bioinformatic investigation of the genome and methylome suggested that RIP and DNA methylation combinatorially maintain *G. sinense* genome stability by inactivating invasive genetic material and transposable elements. The elucidation of the *G. sinense* genome and epigenome provides an unparalleled opportunity to advance our understanding of secondary metabolism and fungal defense mechanisms.

*Ganoderma*, which is a macrofungus in Basidiomycetes, is well known as a medicinal fungus in Asia, where it has been used to treat various diseases for more than 2000 years. Their medicinal properties are considered as a result of a rich diversity of secondary metabolites, including sterols, alkaloids and...
terpenoids. Recently, the genome of *G. lucidum* and *G. sp.* have been sequenced for the study of the molecular mechanisms of secondary metabolism and lignin degradation\(^2\).

As a saprophytic fungus living in complex ecological niches, *Ganoderma* requires a series of powerful defense systems to protect itself from the threat of competitors and invaders. Genome defense is one of the major fungal defense systems developed to protect the genomes of the organisms from invasion by foreign genetic material and transposable elements (TEs), thus maintaining genome stability\(^3\). Repeat-induced point mutation (RIP) was the first reported fungal genome defense system and was discovered in *Neurospora crassa*\(^4\). This mechanism specifically introduces cytosine-to-thymine transitions in repetitive sequences to limit the movement of selfish DNA or mobile DNA during meiosis\(^4-7\).

DNA methylation is an ancient epigenetic DNA regulatory mechanism that is widely involved in gene silencing in eukaryotes, and this mechanism also plays an important role in fungal genome defense\(^8,9\). The methylated component of the *N. crassa* genome consists almost exclusively of relics of transposable elements that were subject to RIP\(^10\). Two additional genetic mechanisms, quelling and meiotic silencing by unpaired DNA (MSUD), have been identified and have been shown to be associated with small RNA-mediated genome defense\(^11-13\).

Fungi are a rich and sustainable source of secondary metabolites. Although the ecological role of most fungal secondary metabolites is still obscure, a growing body of evidence has demonstrated that chemical defense strategies are commonly used by fungi in response to biotic and abiotic environments, including defense against herbivores and fungivores\(^14\). A variety of fungal secondary metabolites are involved in chemical defense processes, including alkaloids, polyketides and terpenoids\(^15\). In filamentous fungi, genes required for the biosynthesis of a specific secondary metabolite are generally clustered together and are often located near telomeres\(^16\). A classic fungal gene cluster contains genes encoding the signature enzymes that synthesize the skeletons of the different classes of secondary metabolites, such as non-ribosomal peptide synthases (NRPSs), polyketide synthases (PKSs) and terpenoid synthases (TSs). These clusters also contain genes encoding tailoring enzymes that modify the skeletons, such as oxidoreductases, methyltransferases and glycosyltransferases\(^17\).

In this work, we assembled a high quality genome of *G. sinense* and compared them with the other two sequenced genomes of *Ganoderma* species. We then examined whole-genome DNA methylation patterns and sequenced the small RNA transcriptome to provide a holistic overview of the genome defense processes in *G. sinense*. We also identified a large array of genes involved in the biosynthesis of secondary metabolites and their regulation. These results will ultimately lead to a better understanding of the roles of specific genes and regulatory pathways in chemical defense systems, thereby presenting a novel avenue for drug discovery.

### Results

#### Genome Sequence and Annotation.

The genome of *G. sinense* (strain ZZ0214-1) was shotgun sequenced to \(~500x\) coverage using a combination of next-generation sequencing technologies (see Supplementary Table S1). The sequences were assembled into 69 scaffolds with a total length of 48.96 Mb, and the assembly represents the largest genome of the three *Ganoderma* species sequenced to date (see Table 1). Approximately 94% of the assembled genome was distributed over 42 scaffolds that were ordered and anchored onto the optical map scaffolds of the 12 *G. sinense* chromosomes constructed using optical mapping technology\(^18,19\) (see Fig. 1A), suggesting the high quality of the genome sequence.
assembly. The average GC content of the *G. sinense* genome (55.59%) was comparable to that of the *G. lucidum* genome (55.9%)\(^1\), but it was higher than that of other sequenced Basidiomycota fungi (average of 50.63%) (see Supplementary Fig. S1).

A total of 15,688 gene models were predicted, of which 87% were supported by the assembled RNA-Seq transcripts. In total, 85% of the genes were annotated by searching against public databases,
with 65% of the annotated genes containing Pfam domains. Additionally, 202 tRNAs, including nine pseudogenes, were predicted, with an average length of 82 bp.

Transposable elements (TEs), including retrotransposons and DNA transposons, accounted for approximately 12.2% of the *G. sinense* genome. RNA-mediated retrotransposons were more abundant than DNA-mediated transposons (see Supplementary Table S2), consistent with other *Ganoderma* species. Long terminal repeats (LTRs) were the most abundant class of retrotransposons, among which the *Gypsy* and *Copia* were the largest subclasses. In *G. sinense*, long interspersed nuclear elements (LINEs) were much more abundant than in the other two fungi (see Supplementary Table S2). DNA transposons accounted for 2.2% of the *G. sinense* genome, and these transposons were primarily comprised of terminal inverted repeats (TIRs), *helitrons* and miniature inverted-repeat transposable elements (MITEs).

**Genome Comparison and Phylogenetic Relationships.** Orthology analysis was carried out using the genome sequence data for the three sequenced *Ganoderma* species (*G. sinense*, *G. lucidum* and *Ganoderma sp.*), and 22 other fungi from Basidiomycota, Ascomycota and Chytridiomycota (see Supplementary Table S3). Approximately three-quarters of the genes in *G. sinense* were included in a total of 9,336 orthologous groups. The *G. sinense* genome was found to contain more paralogous genes than the other two *Ganoderma* species (see Fig. 1B). A total of 3,122 paralogous genes were identified, accounting for 19.9% of all the *G. sinense* genes. Almost all of these genes were found in orthologous groups, suggesting that the paralogs of *G. sinense* are primarily derived from ancient gene duplication events.

Fifty-nine single-copy genes identified in the orthology analysis were used to construct a phylogenetic tree using the maximum likelihood method (see Fig. 1B). The tree topology was generally consistent with prior analyses. Of the available sequenced genomes, *Dichomitus squalens* was the closest relative of the three *Ganoderma* species in the Polyrales clade. In addition to *Trametes versicolor*, these fungi formed a clade of five white-rot fungi. Some of the major medicinal fungal species used in China, including *Ganoderma*, *T. versicolor* and *Wolfiporia cocos*, belonged to the Polyrales clade, suggesting that this order may be a source of medicinal fungi for drug development. Based on fossil calibrations at the three nodes, including the ancestors of Boletales, Agaricales and Ascomycota, the divergence time of *Ganoderma* and *Dichomitus* was estimated to be approximately 38 Ma (millions of years ago), with *G. sinense* appearing at approximately 18 Ma.

A whole-genome comparison between *G. sinense* and *G. lucidum* using the MUMmer software revealed similar chromosome organization, with ~80% similarity between the alignment blocks (see Fig. 1A and Supplementary Fig. S2). Five major breakpoints were found between *G. sinense* and *G. lucidum*, suggesting species-specific chromosomal rearrangements (see Fig. 1A). Synteny analysis revealed 160 synteny blocks comprised of 7,524 genes and 7,509 genes in *G. sinense* and *G. lucidum*, respectively. Additionally, 115 synteny blocks were identified between *G. sinense* and *G. sp.*, comprised of 7,623 and 7,539 genes, respectively. A total of 5,729 homologous genes in *G. sinense* were not syntenic with *G. lucidum* or *G. sp.*, suggesting a number of gene rearrangement events during *Ganoderma* evolution.

**Genome Wide DNA Methylation.** Whole-genome bisulfite sequencing was used to detect 5-methylcytosine (5mC) in *G. sinense*. Approximately 1.8% of the cytosine residues were methylated in the *G. sinense* genome, among which 98.7% were specific to CpG dinucleotides (see Fig. 2A). To ascertain a possible contextual bias of DNA methylation, the methylation level (number of methylated cytosines
per kilobase) of the TEs versus the coding regions was determined. As shown in Fig. 2B, the methylation level in the TE regions was higher than in the genic regions, which is consistent with results observed in other fungal genomes9, and up to 33% of the CpG dinucleotides were subject to methylation.

DNA methylation is catalyzed by DNA methyltransferase (DMTase), which transfers the methyl group from a methyl donor onto a cytosine. Phylogenetic analysis of the DMTase sequences suggested that *G. sinense* has at least three DMTases (see Supplementary Fig. S3). These genes were found to be homologous to *Dnmt1a* (46% similarity with *Postia placenta*), *Dnmt1b* (36% with *P. placenta*) and *Dnmt2* (53% with *Coprinopsis cinerea*) (see Supplementary Table S4 and S5). Domain analysis showed that these DMTases have a relatively consistent domain structure within the same subfamily (see Supplementary Table S6 and Supplementary Fig. S4).

Trimethylation of histone H3 on lysine 9 (H3K9me3) by an H3-specific histone methyltransferase (HMTase) is essential for DNA methylation. An H3K9-specific HMTase homolog was found in *G. sinense*, and this protein contained three specific domains: pre-SET, SET and post-SET (see Supplementary Fig. S4). Additionally, putative accessory proteins for H3K9me3 and cytosine methylation were found, including CUL4 (Cullin 4), DDB1 (Damaged DNA Binding Protein-1) and HP1 (Heterochromatin Protein 1) (see Supplementary Table S5).

**Repeat-Induced Point Mutation.** Previous RIP investigations mainly focused on Ascomycetes; however, a recent study suggested the existence of RIP in Basidiomycetes21. To determine whether RIP exists in *G. sinense*, TEs with a minimum length of 400 bp and a sequence identity greater than 80% were analyzed for RIP hallmarks using a sequence alignment-based algorithm6,22. Within these repetitive regions, transitions from C:G to T:A accounted for more than 87% of all point mutations (see Fig. 3A), suggesting the existence of RIP in *G. sinense*. Similarly, *G. lucidum* and *G. sp.* also have RIP processes, as suggested by their high percentages of RIP-like transition mutations (90% and 91%, respectively; see Fig. 3A). RIP is not random, and CpN dinucleotides are preferentially altered in *G. sinense*. Quantification of the
Small RNA-mediated Silencing. To unravel the potential roles of RNA silencing in *G. sinense* genome defense, two different small RNA (smRNA) libraries were generated from mycelium and fruiting bodies in *G. sinense* and then subjected to Illumina high-throughput sequencing. To identify small RNAs specific to mycelium and fruiting bodies, total reads were grouped into 1,655,536 unique sequences. Only 65,362 (3.95%) unique sequences were common to both tissues. Of the remainder, most sequences (1,035,653; 62.56%) were specific for fruiting bodies, while 33.49% (554,521) were specific to mycelium. Sequences with length 18–30 nt were mapped to the haploid *G. sinense* genome, generating 1,812,452 (76%) and 5,705,067 (89%) genome-matched reads from mycelium and fruiting bodies, respectively (see Fig. 4A). A large number of these mapped reads were annotated as non-coding rRNA (34.04% and 53.78%), and as tRNA (11.13% and 1.51%). Only a small number of mapped reads were classified as milRNA (958 and 5,132) (see Fig. 4B) and disiRNA (594 and 10,027).

Precursors of milRNA can produce stable hairpin structures. On the basis of this characteristic, sixty-three novel milRNAs (18–25 nt) were identified in *G. sinense* (see Supplementary Table S7). Analysis of 5’-terminal nucleotide bias revealed that uracil (U) was the most frequent 5’-terminal nucleotide in milRNAs of *G. sinense*, while the frequency of adenine (A) significantly increased in fruiting bodies (see Fig. 4C).

Analysis of milRNA targets showed possible regulation of a variety of transport process. Two species of milRNAs were targeted to two members of the major facilitator superfamily (MFS), involved in transmembrane transport. milRNA-4 was predicted to act on the MFS gene *GS04616*. Meanwhile, milRNA-19 was predicted to target the MFS gene *GS13257*, whose expression correlated to milRNA-19 abundance. In mycelia, milRNA-19 was silenced and *GS13257* was highly expressed, whereas in fruiting bodies, milRNA-19 was upregulated and *GS13257* was significantly downregulated. Thus milRNA-19 may silence the expression of *GS13257* and inhibit its specific function in transmembrane transport.

In *G. sinense*, more than 40 regions of the genome were found where both strands of the DNA were almost equally mapped by strand-specific smRNA reads, suggesting that both genomic DNA strands are transcribed (see Supplementary Table S8). Therefore, double-stranded RNA formed from these regions may be further processed into disiRNAs. Seventeen of these regions were located at the ends of chromosomes, presumably close to the subtelomeric regions, which are particularly rich in TEs (see Supplementary Table S8).

In *G. sinense*, a series of complex components involved in quelling and meiotic silencing by unpaired DNA (MSUD) were discovered including candidate genes encoding the RNA-dependent RNA polymerases (RdRPs), Dicer and Argonaute proteins, using funRNA and functional annotation methods (see Supplementary Table S9). Some pathway-specific genes were also found, including two *qde-1*-like genes specific for quelling and a *sad-1*-like gene specific to the MSUD pathway. These findings suggest that small RNA-mediated genome defense mechanisms are most likely conserved in this fungus.

TE Inactivation by Genome Defense. Class I TEs, also known as retrotransposons, are mobilized in genomes through the action of their RNA intermediates. Therefore, the expression of class I TEs provides direct evidence for their activity. By mapping the transcripts from the mycelium to the TE sequences, we classified approximately one third of the class I TEs as active during this developmental stage (see Table 2). Among the remaining inactive class I TEs, 63% were affected by RIP mutations or DNA methylation. Silencing of the remaining 37% of the inactive class I TEs may be caused by RNAi, indel mutations during translocation or other unknown mechanisms. In total, approximately 93.2% of the class I TEs subjected to both RIP mutations and DNA methylation were silenced, suggesting that the strong silencing was induced by the dual effects of RIP and DNA methylation.

Unlike class I TEs, class II TEs, also known as DNA transposons, move throughout the genome using a “cut and paste” mechanism, without the need for RNA intermediates. Transposases are essential for this process. Although some nonautonomous DNA transposons do not contain transposase genes, transposases encoded by other autonomous transposons may help carry out their movement. Therefore, it is very difficult to estimate DNA transposon activity by analyzing their transcript levels. In total, 110 transposase genes were found in the *G. sinense* genome, and these genes were assigned to the TIR, MAVERICK and *helitron* superfamilies. Within the TIR superfamily, 93 members were classified as CACTA, *Tc1/Mariner*, *PIF/Harbinger*, *P* and *hAT* elements. Twenty-five transposases had potential nucleic acid binding activity (GO:0003676) or DNA binding activity (GO:0003677). Eighty-six transposase genes were methylated, and only 11 of these were expressed; therefore, DNA methylation may inactivate the class II TEs by silencing the transposase genes.
Gene Clusters Involved in Biosynthesis of Secondary Metabolites. As an important medicinal fungus, *G. sinense* produces a variety of secondary metabolites including steroids, alkaloids and terpenoids, many of which are most likely involved in chemical defense. In most fungi, the genes related to the biosynthesis of secondary metabolites are generally clustered together. In *G. sinense*, one non-ribosomal peptide synthetase (NRPS), six polyketide synthases (PKSs), seven NRPS-like genes and twenty terpene synthases were located in gene clusters predicted using the antiSMASH and/or SMURF software. Among these putative clusters, one NRPS cluster had a similar organization to that found in *G. lucidum*, which contains multiple genes encoding tailoring enzymes, transcription factors and transporters, as well as the signature NRPS gene (see Fig. 5A). Five genes in this putative cluster were co-expressed with NRPS during fungal development with a correlation coefficient of >0.9, which is consistent with the existence of a gene cluster organized around the NRPS gene. A 310-kb region of chromosome 7 contained four PKSs and a 150-kb TE-enriched fragment with varying DNA methylation and transcriptional gene silencing profiles. Most of the genes were collinear with *G. lucidum*, except for those in the TE-enriched region, indicating the conserved gene order in the non-TE regions. The origin of the TEs and duplicated PKSs should be further investigated.
CYP Gene Clusters and Heavily Methylated CYPs. Triterpenoids are derivatives of lanosterol that comprise a major class of bioactive compounds in *G. sinense*. All eleven enzymes in the biosynthetic pathway of lanosterol were identified in this fungus (see Supplementary Table S10). Cytochrome P450s (CYPs) greatly contribute to the diversity of triterpenoids through a series of oxidation reactions. In our study, a total of 237 CYPs, including nine pseudogenes, were identified and classified into 41 families. The CYP distribution in the genomes of *G. sinense* and *G. lucidum* was very different, with only 11 CYPs found in the identified syntenic regions between these two species. Compared with *G. lucidum*, *G. sinense* was missing the CYP5363 family, while the number of CYP512s and CYP5035s was greatly increased (see Supplementary Table S11). A total of 44 CYPs were located in the gene clusters shown to contain secondary metabolite biosynthesis genes. At least 68 tandemly-duplicated CYP genes were found to belong to 29 physical clusters, and each cluster was comprised of the same CYP gene family. This suggests that a number of gene duplication events contributed to the expansion of the CYP gene families in *G. sinense*. An example of CYP evolution by duplication is presented in Fig. 5B. Here, a tandem CYP cluster contains four CYP5139Gs and two CYP5359s, while the corresponding cluster in *G. lucidum* contains only two CYP5139Gs and two CYP5359s, indicating the species-specific CYP5139 gene duplication following the divergence of these two species (see Fig. 5B).

However, these CYPs were not always functionalized, in sometimes, they may be regulated by specific mechanisms, such as DNA methylation. In our study, at least 99 CYPs were located in the TE-rich regions, suggesting that some CYP gene duplications and rearrangements may be associated with TE translocation. Seven of the CYP genes found in the TE-rich regions were heavily methylated, and these genes belonged to three expanded CYP families: CYP5035, CYP5150 and CYP5359. Of the methylated CYP5150 genes, three (CYP5150D27a-c) genes showed more than 90% similarity and were adjacent to each other (see Supplementary Table S12). Alignment of these gene sequences suggested that a tandem duplication event (indicated as event I), covering two 10-kb regions (see Supplementary Fig. S5), may have occurred, and this event may have resulted from an unequal crossing over event during meiosis due to the close proximity and high sequence similarity. CYP5150D27c was located outside of these regions; thus an additional single gene-specific duplication event (indicated as event IV) may have occurred.

Regulation and Transport of Secondary Metabolites Regulated by DNA Methylation. Fungal secondary metabolism is controlled by a complex regulatory network involving multiple genetic, epigenetic and proteomic control points. Transcription factors (TFs) play a central role in regulating pathway flux and in controlling the expression of key pathway enzymes. In total, 449 transcription factors were identified in the *G. sinense* genome, and these genes were classified into 48 families (see Supplementary Table S13). Among these, at least 29 TF genes were located in secondary metabolism gene clusters, suggesting that they may encode pathway-specific TFs.
At least 74 TF genes were methylated in the fungal mycelia. Most genes belonged to the CCHC-type and Cys2His2-type zinc finger TF families and were located in the TE-rich regions. RNA-Seq revealed that 93% of the methylated TF genes were silenced. Of the 29 TF genes located in secondary metabolism gene clusters, seven were methylated and five of these genes were silenced during development. Consequently, it is conceivable that secondary metabolism gene clusters and their associated transcription factors are regulated by DNA methylation in *G. sinense*.

Fungal growth conditions in culture, including the carbon and nitrogen source, mineral ion complement, light intensity, pH and oxygen supply, can greatly influence the biosynthesis of secondary metabolites. Global TFs are involved in the response to environmental signals. In *G. sinense*, two Cys2His2-type zinc finger TFs, CreA and PacC, were identified, which control the response to the carbon source and pH signals, respectively. A GATA family TF, AreA, involved in the nitrogen response, as well as components of the Velvet complex, VeA and VelB, were also found. The Velvet complex generally regulates the cross-talk between the fungal developmental and secondary metabolism.

In *G. sinense*, seven classes of transporters were identified (see Supplementary Table S14), some of which play essential roles in the transport of secondary metabolites across cell membranes. Two major families are involved in the transport of small secondary metabolite molecules, including 235 major facilitator superfamily (MFS) and 53 ATP-binding Cassette (ABC). A total of 112 transporters were located in secondary metabolism gene clusters, suggesting their possible roles in the transport of specific...
secondary metabolites. Thirty-six of these transporters were predominantly methylated and transcriptionally silenced, suggesting that methylation-induced transcriptional gene silencing may be involved in the regulation of secondary metabolite transport in this fungus.

**Discussion**

For static creatures, physical protection is limited against the enemies, as such, chemical compound becomes the second line of defense, especially for those living in complex environments. Further, the last line of defense, genomic defense, play key roles in maintain the genetic stabilization of genome. The *Ganoderma* genus comprises a large group of saprobic mushroom-forming fungi that mostly live in subtropical/tropical regions, and many have economic value for applications in healthcare. Although the medicinal value of *Ganoderma* spp. has been extensively reported, their chemical and genomic defense machinery has never been elucidated.

Here, we report the genome sequence of *G. sinense*, one of the most valuable medicinal fungi. Together with the other two sequenced *Ganoderma* species, *G. lucidum* and *G. sp.*, the genome sequence of this fungus provided further insight into the molecular basis and evolution of defense mechanisms in Basidiomycetes, particularly the genetic and epigenetic factors underlying the environmental induction of the defense systems and the production of secondary metabolites.

In this study, the combination of optical mapping and NGS technology is a powerful way to construct a chromosome-level genome sequence map. The *G. sinense* genome sequence was constructed into twelve chromosomal pseudomolecules based on the optical map scaffolds, which is one less than in *G. lucidum*. Additionally, the comparison of the genome sequences indicated a number of translocations and duplications among the three *Ganoderma* fungal species and also revealed many non-homologous genes, suggesting rapid divergent evolution following speciation. A considerable number of active TEs were also discovered in *G. sinense*, suggesting the association between specialization and rapid genome evolution. Also, it suggests that the genome defense processes in this fungus may not be as strong as in the model species *N. crassa*.

Whole-genome bisulfite sequencing provided an overview of DNA methylation in *G. sinense* for the first time. Although DNA methylation was only noted on 1.8% of the cytosines in the genome, most of the methylated cytosines were distributed in repetitive regions. Accordingly, most cytosine methylation and RIP mutations were located in the same or adjacent regions (see Fig. 3C), suggesting the potential coordination of these mechanisms in TE silencing in *G. sinense*. Additionally, small double-stranded RNAs potentially generated from the transcription of both complimentary DNA strands in the subtelomeric regions were also identified, which may lead to RNA-mediated silencing of TE activity or telomeric heterochromatin.

As a fungi-specific genome defense system, RIP has been well studied in the model fungus *N. crassa*; however, this process remains largely unstudied in Basidiomycetes. We systemically investigated the RIP process in *G. sinense* and found that RIP mutations occurred most frequently on cytosine residues in CpG dinucleotides, compared to the CpA dinucleotides in *N. crassa* and most other fungi investigated to date. CpG dinucleotides are also the most common targets of DNA methylation. Additionally, the key protein required for RIP, RID (RIP defective) in *N. crassa* (see Fig. 3), exhibits methyltransferase activity. This raises the possibility that RID evolved from DNA methyltransferases and, therefore, that ancient RIDs may have preferentially recognized CpG dinucleotides. CpA is the second-most preferred dinucleotide for RIP in *G. sinense*. Conversion of CpG to TpG via RIP can introduce the stop codon TGA, while conversion of CpA to TpA can produce the stop codons, TAG and TAA. The introduction of stop codons can truncate the proteins essential for TE movement; therefore, RIP represents an efficient TE inactivation mechanism.

Small RNAs can functionally produce the post-transcriptional gene silencing, some of which were found with genome defense related in many organisms. In *G. sinense*, miRNAs and disiRNAs were first identified. Although there is no experimental evidence to be related to small RNA-mediated genome defense in *G. sinense*, the small RNAs derived from the TE enrichment regions may be involved in silencing or repressing transposable element activity for genome defense or for telomeric heterochromatin maintenance.

Genome sequencing revealed that *G. sinense* encodes a large set of enzymes involved in the biosynthesis of secondary metabolites. Many of these genes appear to have originated from gene duplication, resulting in the potential for accelerated evolution via three fates: neo-functionalization, sub-functionalization and increased gene-dosage advantage. The diversification of the cytochrome P450 (CYP) gene families in plants and higher fungi has led to the emergence of various biochemical pathways that produce secondary metabolites such as phenylpropanoids, alkaloids and terpenoids. Therefore, CYPs contribute to the chemical defense mechanisms of the organism. Abundant CYPs have been identified in the Polyaules order, which includes *G. sinense* and other traditional medicinal fungi, such as *G. lucidum*, *T. versicolor* and *W. cocos*. Thus, these species may have undergone expansion of CYPs for the enhanced biosynthesis of diverse secondary metabolites. In *G. sinense*, evidence for the tandem and fragmental duplication of CYPs provided some explanation for the gene expansion. The resulting diversification of CYPs during evolution could explain the divergence of triterpenoids between *G. sinense* and *G. lucidum*. More than thirty gene clusters were bioinformatically predicted in the *G. sinense* genome, and these gene clusters are expected to produce non-ribosomal peptide (NRPs), polyketide (PKs) or terpenoids.
However, with the exception of triterpenoids, few NRPs, PKs or other types of terpenoids have been isolated and identified from this fungus. This suggests that most of these gene clusters are tightly controlled and silenced under general culturing conditions, resulting in the substantial underestimation of versatile secondary metabolite production. One way to maintain such tight control is through reversible gene silencing; therefore, DNA methylation and small RNA-mediated gene silencing may play critical roles in genome defense and in the regulation of secondary metabolism. Considering that many secondary metabolites may play important roles in chemical defense, the co-cultivation of *G. sinense* with one or more microorganisms from the same ecosystem may be a promising way to activate specific gene clusters. This strategy has been demonstrated in *Aspergillus nidulans*.

In conclusion, the analyses of the whole-genome sequence, DNA methylation and small RNA sequences suggest that *G. sinense* possesses the genetic and epigenetic hallmarks of genome defense systems, as well as chemical defense systems. Our data also provide new insights into the fundamental principles of secondary metabolite production in this and other important medicinal species, with significant implications for the pharmaceutical industry.

**Methods**

**Strain and Culture Condition.** *Ganoderma sinense* strain CGMCC5.0069 was sourced from the China General Microbiological Culture Collection Center (Beijing, China, http://www.cgmcc.net/). The haploid strain ZZ0214-1 was derived from CGMCC5.0069 using the protoplast monokaryogenesis method. In order to confirm the strain status, calcofluor white staining, DAPI staining and single nucleotide polymorphism (SNP) analysis were used as previously described. Liquid cultures were shaken in potato dextrose medium at 50 rpm for 6 days. The primordia and fruiting bodies of strain CGMCC5.0069 were cultivated on *Quercus variabilis* Blume logs at HuTao Pharmaceutical Company (LuoTian, Hubei Province, China). All strains are available upon request.

**Construction of Optical Maps.** Protoplasts of the monokaryotic strain ZZ0214-1 were collected by centrifugation at 1,000 g for 10 min and were then diluted to a final concentration of 2 × 10^7 cells mL^-1^. Gel inserts were prepared as previously described and stored in NDSK buffer (0.5 M EDTA, 1% (v/v) N-lauroylsarcosine, 1 mg/ml proteinase K). Genomic DNA was released by melting DNA gel inserts at 70°C for 7 min, and then digesting with β-agarase (NEB, USA) at 42°C for 2 h. Mounted DNA molecules were digested by the restriction endonuclease SpeI in NEB Buffer 2 (50 mM NaCl, 10 mM Tris-HCl, 10 mM MgCl₂, 1 mM dithiothreitol, pH 7.9; New England Biolabs, USA) without BSA, and Triton X-100 added to a final concentration of 0.02%. Digested DNA molecules were then stained with 12 μL of 0.2 μM YOYO-1 solution (5% YOYO-1 in TE containing 20% β-mercaptoethanol; Eugene, USA). Fully automated imaging workstations were used to generate single-molecule optical data sets for map assembly.

**Genome Sequencing and Assembly.** Genomic DNA of *G. sinense* was sequenced using the Roche 454 GS FLX (Roche, USA) and Illumina HiSeq 2000 (Illumina, USA) NGS platforms. The sequencing process followed the manufacturer’s recommendations. Sequencing reads from Roche 454 were used to construct the primary assembly using CABOG and then scaffolded with Illumina paired-end reads using SSPACE version 1.1. The assembly was error checked and manually corrected to build the finished scaffolds. The final scaffolds were constructed to build the chromosome-wide pseudomolecules by optical mapping.

**Transcriptome Sequencing and Analysis.** Gene expression analysis was performed on material obtained from the three developmental stages of *G. sinense*, including the mycelia, primordia, and fruiting bodies. Frozen samples were ground to powder in liquid nitrogen. Total RNA of each sample was extracted using an RNaseasy Plant Mini Kit (Qiagen, Germany) according to manufacturer’s instructions but with 10% volume β-mercaptoethanol added to Buffer RLC before use. Genomic DNA was eliminated using DNase I (NEB, USA). RNA integrity and quality were checked using the RNA 6000 Nano II kit on a Bioanalyzer 2100 (Agilent, USA). RNA-Seq was performed as recommended by the manufacturer (Illumina, USA). And the samples were sequenced using an Illumina HiSeq 2000.

RNA-Seq data from the stages of mycelia, primordia and fruiting bodies were assembled to reconstruct the transcripts using Trinity. Transcript abundance was estimated as normalized fragments per kb of transcript per million mapped reads (FPKM) using RSEM. Differential expression analysis was executed using edgeR.

**Gene Annotation.** Gene prediction was executed using the PASA software package. Four software were used for *ab initio* gene finding, including Augustus, GeneMark, Fgenes and SNAP. Additionally, the evidence-based method was used to weight and calibrate gene structure. The assembled transcripts and proteins were aligned to the genome assembly using exonerate-est2genome and exonerate-protein2genome, respectively. The predicted genes were functionally annotated by blasting the protein databases, including NCBI non-redundant, KEGG and Uniprot/Swissprot databases. Genes were also annotated using InterProScan, and then classified according to Gene Ontology terms.
**Genome Comparison and Phylogenetic Analysis.** Syntenic genes between *G. sinense*, *G. lucidum* and *G. sp* were identified using MCscan (version 0.8). Orthologous groups from 41 fungal genomes were constructed using OrthoMCL version 2.0 (http://orthomcl.org). The BLASTP cutoff was set as E-value <1e-25. According to orthology analysis, 59 single copy proteins were selected and used for phylogenomic analysis. Individual alignments of the 59 proteins were generated using MUSCLE (version 3.6) from 25 species (see below) related to *G. sinense* and previously studied in genome defense, DNA methylation and RNA silencing. Using concatenated amino acid alignments, a maximum-likelihood phylogenetic tree was subsequently produced by the program RAxML (version 7.2.8) with the PROTGAMMAWAG model and 100 bootstrap partitions. The divergence time between these fungi was estimated with penalized likelihood (PL) method using r8s by calibrating against three different clades in the Dikarya: Ascomycota (600–452 Ma), Boletales (130–50 Ma) and Agaricales (140–92 Ma)29.

Abbreviations of related 25 species: AspNi, *Aspergillus niger*; BatDe, *Batrachochytrium dendrobatidis*; ConPu, *Coniophora puteana*; CopCi, *Coprinopsis cinerea*; DicSq, *Dichomitus squalens*; FomMe, *Fomitiporia mediterranea*; FonPi, *Fomitopsis pinicola*; GanLu, *Ganoderma lucidum*; GanSi, *Ganoderma sinense*; GanSp, *Ganoderma sp*.; LacBi, *Laccaria bicolor*; MelLa, *Melampsora larici-populina*; NeuCr, *Neurospora crassa*; PhaCh, *Phanerochaete chrysosporium*; PosPi, *Postia placentae*; PucGr, *Puccinia graminis f. sp. tritici*; RhGr, *Rhototorula graminis*; SacCe, *Saccharomyces cerevisiae*; SchCo, *Schizophyllum commune*; SchPo, *Schizosaccharomyces pombe*; SerLa, *Serpula lacrymans*; TraVe, *Trametes versicolor*; TreMe, *Tremella mesenterica*; UstMa, *Ustilago maydis*; WolCo, *Wolfiporia cocos*.

**Bisulfite Sequencing and Analysis of DNA Methylation.** The Illumina bisulfite sequencing workflow was used to get the whole genome methylation profiles of *G. sinense* mycelia. Briefly, DNA was sheared to fragments of 100–500 bp using a Covaris. Fragments were pooled using T4 DNA polymerase and Klenow enzyme. After end-repair, 3’ ends were adenylated. Commercial adaptors were ligated to adenylated fragments and then purified. The purified products were treated twice with sodium bisulfite using the EpiTect Bisulfite Kit to convert unmethylated Cs to Us. The treated DNA was amplified by 18 cycles of PCR using high fidelity polymerase. Size-selection was conducted after amplification, bands around 300 bp were used for sequencing with the Illumina HiSeq 2000 platform.

DNA methylation analysis was performed using Bismark40, which maps bisulfite treated sequencing reads to the genome. Methylation level for each gene was estimated by counting the number of methylation sites per kilobase pairs.

**Small RNA Sequencing.** Strand-specific small RNA sequencing was performed according to the Illumina TruSeq small RNA sequencing protocol. One microgram of total RNA from mycelia or fruiting bodies was used as input. After 3’ and 5’ adaptor ligation using T4 RNA ligase, reverse transcription and amplification were conducted using PCR as the manufacturer’s instruction. The amplification quality was detected using a high sensitivity DNA chip on a Bioanalyzer 2100. Amplified cDNA bands of around 150 bp were agarose gel-purified and concentrated by ethanol precipitation. Library validation was conducted on an Agilent 2100 Bioanalyzer using a high sensitivity DNA chip. Sequencing was performed on an Illumina HiSeq 2000 platform.

**Identification of Small RNAs.** Small RNAs between 18–30 nt were identified in our study. The loci of miRNAs were determined by the following criteria24: 1) Small RNAs were significantly derived from one DNA strand (at least 10 fold more than the other DNA strands); 2) Small RNAs were highly enriched at the loci with a density over 270 reads/kb; 3) Small RNAs were located on the stem region of a hairpin structure with flanking sequences using secondary structures predicted by RNAfold (http://rna.tbi.univie.ac.at/cgi-bin/RNAfold.cgi); 4) Sequences with a minimum folding energy less than −20 kcal/mol and a minimal folding free energy index (MFEI) greater than 0.8 was considered as miRNA precursors. disiRNA loci were identified by screening the whole genome. Two criteria were used to predict disiRNAs24: 1) Similar abundance from both DNA strands; 2) The density of small RNAs was significantly higher than the background (at least 60 reads/kb).

**Transposable Elements and RIP Analysis.** Transposable elements were identified and annotated by the REPET package (http://urgi.versailles.inra.fr/Tools/REPET), which contains TEdenovo and TEnannot pipelines. TEs with a minimum length of 400 bp and greater than 80% sequence identity were used for expression analysis. Expression of TEs was evaluated by mapping RNA-Seq sequences to the genome using TopHat version 2.0.9. Expressed TEs were defined according to following criteria: 1) RNA-Seq reads mapped to the genome reference sequence allowing two mismatches; 2) Paired reads that spanned over estimated inserts or mapped to different scaffolds were discarded; 3) Reads with multiple hits were filtered. Then, the remaining reads were used to determine TE expression.

TEs with a minimum length of 400 bp and greater than 80% sequence identity were also used for RIP analysis. To detect RIP sites, an alignment-based method was used to locate the variant sites according to the characteristics of C to T variation. Pre-RIP sites, which represent the state prior to RIP mutation, were also determined by sequence alignment. RIP indices were calculated using RIPCAL.
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Additional Information

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