

## Comprehensive Transcriptomic Analysis of *Cordyceps militaris* Cultivated on Germinated Soybeans

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### ABSTRACT

The ascomycete fungus *Cordyceps militaris* infects lepidopteran larvae and pupae and forms characteristic fruiting bodies. Owing to its immune-enhancing effects, the fungus has been used as a medicine. For industrial application, this fungus can be grown on germinated soybeans as an alternative protein source. In our study, we performed a comprehensive transcriptomic analysis to identify core gene sets during *C. militaris* cultivation on germinated soybeans. RNA-Seq technology was applied to the fungal cultures at seven-time points (2, 4, and 7-day and 2, 3, 5, 7-week old cultures) to investigate the global transcriptomic change. We conducted a time-series analysis using a two-step regression strategy and chose 1460 significant genes and assigned them into five clusters. Characterization of each cluster based on Gene Ontology and Kyoto Encyclopedia of Genes and Genomes databases revealed that transcription profiles changed after two weeks of incubation. Gene mapping of cordycepin biosynthesis and isoflavone modification pathways also confirmed that gene expression in the early stage of GSC cultivation is important for these metabolic pathways. Our transcriptomic analysis and selected genes provided a comprehensive molecular basis for the cultivation of *C. militaris* on germinated soybeans.

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## 1. Introduction

The ascomycete fungus *Cordyceps militaris* can infect lepidopteran pupae, which includes over 500 species capable of infecting many other insects, spiders, and mites [1]. When this fungus colonizes dead or living Lepidoptera caterpillars, the mycelium infiltrates into the caterpillar body and finally replaces most host tissues. Later, fruiting bodies are formed on the surface of the insects' body. They are called "Dong Choong Ha Cho" in Korea, meaning that worms in winter turn into plants in summer. The fruiting bodies have been used as medicine in Asia for a long time [2]. Anti-tumor, anti-inflammatory, and anti-oxidative activities have been reported in the fruiting bodies of *C. militaris* [3–8]. One of the main bioactive ingredients of *C. militaris*, 3'-deoxyadenosine (also called cordycepin), displays anti-tumor activity via an apoptosis mechanism [9–11].

The genome of *C. militaris* has been estimated as 32.27 Mb and has 9810 genes [12]. After the publication of its whole genome sequence, possible

metabolic pathways for cordycepin biosynthesis have been suggested based on bioinformatic data [12,13]. Recently, a reverse genetics approach with liquid chromatography was applied to identify a cordycepin/pentostatin biosynthesis cluster, which consists of four genes (named *CNS1-4*) [14]. In that study, *CNS1* and *CNS2* genes were indispensable for cordycepin biosynthesis, and *CNS3* was essential for pentostatin biosynthesis. The function of *CNS4* was predicted to be the transport of pentostatin to balance the intracellular concentration of cordycepin [14]. In addition, many transcriptome studies have been performed using *C. militaris*, focusing on fruiting body formation [12,15] and optimization of cordycepin production [16–18]. Recently, many comparative transcriptomic studies have suggested metabolic pathways and key enzymes involved in cordycepin production. These studies demonstrated that it is practicable to improve the productivity of cordycepin through analysis and regulation of its biosynthetic pathways [15,19–21]. Despite many studies having been conducted on the biosynthetic

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process of cordycepin in *C. militaris*, there is no report on the biosynthetic process of other useful bioactive ingredients, such as isoflavones at the level of gene expression in *C. militaris*.

Although fruiting bodies of *C. militaris* have medicinal effects on human health, it is difficult to produce them on a large scale because of the difficulty in culturing this fungus on insect pupae. To overcome this problem, a method was developed in which germinated soybeans were used as a protein source instead of pupae for artificial mass cultivation [22]. In this method, the mycelia of *C. militaris* were placed on sterile germinated soybeans and incubated at 25 °C for 8 weeks. This solid culture is hereinafter referred to as GSC. Metabolomics and multivariate statistical analyses were conducted to compare compositional changes between germinated soybean (GS) and GSC [23]. One-week-old cultures exhibited strong GSC biological activity. Interestingly, new forms of isoflavones, a modified version of genistein and daidzein, were detected in GSCs. The isoflavones significantly inhibited degranulation in antigen-stimulated RBL-2H3 cells, suggesting that GSC is an anti-allergic agent for treating IgE-mediated allergic disease [24]. Ethanol extracts of GSC showed anti-inflammatory effects on acute colitis symptoms [25]. These results reveal the efficiency of GSC, which can be used as a promising drug or health supplement.

In this study, we conducted time-series gene expression profiling during GSC cultivation to

identify marker genes at different culture stages and to map those genes to the biosynthesis pathways of cordycepin or modified isoflavones.

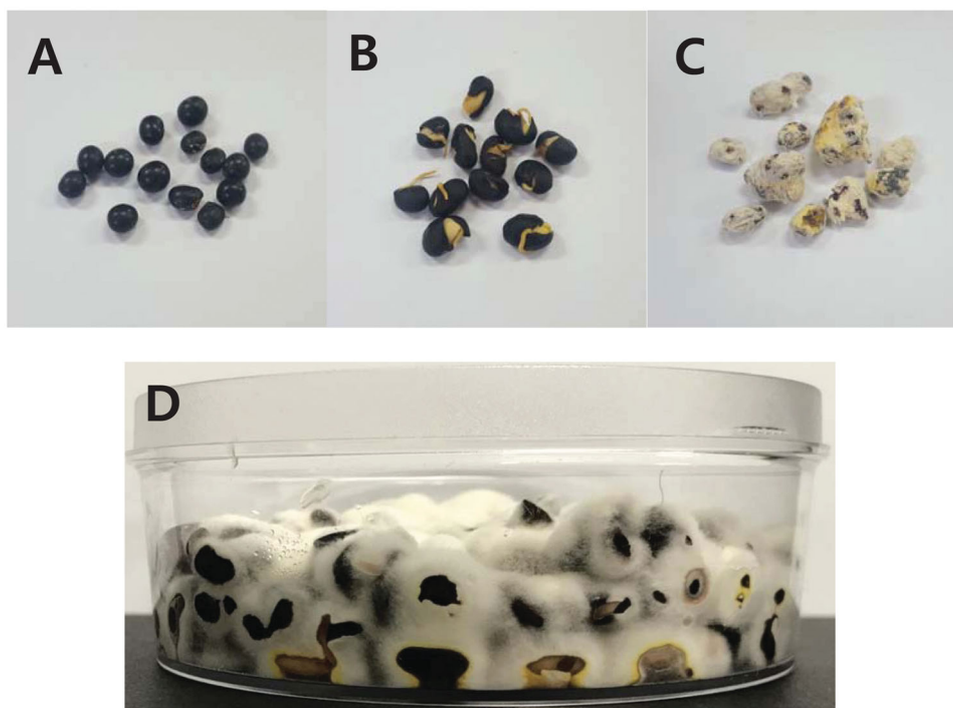
## 2. Materials and methods

### 2.1. Fungal isolates

The *C. militaris* isolate KUCARI\_0903 was obtained from the Cell Activation Research Institute (Seoul, Korea). The culture was maintained on a potato dextrose agar plate at 25 °C in the dark.

### 2.2. Fungal culture on soybeans and its sampling

Three-day-old *C. militaris* culture was filtered through a three-layered cheese cloth to remove potato dextrose broth. The fungal mycelia were ground using a sterile spoon and cloth, and 25 ml of sterile distilled water was added. Germinated soybeans (*Glycine max*, “Jui Nun Yi Cong”) were autoclaved and placed in a 30-ml plate (Fisher Scientific International, Inc., Hampton, USA) (Figure 1). Four milliliters of mycelia ground with water was used as the inoculum for each plate. The mycelia on the germinated soybeans were cultured at 25 °C for 8 weeks (Figure 1(D)) [22,23]. For RNA-Seq, fungal mycelia were sampled at 2, 4, and 7 days post-inoculation (dpi) and 2, 3, 5, and 7 weeks post-inoculation



**Figure 1.** Soybeans with *Cordyceps militaris*. (A) Soybeans (*Glycine max*, “Jui Nun Yi Cong” in Korean); (B) Germinated soybeans; (C) *C. militaris* mycelia grown on the germinated soybeans for eight weeks; (D) *C. militaris* growing on the germinated soybeans (GSC).

(wpi). Each sample harvested from three plates was pooled for cost-saving before RNA extraction [26].

### 2.3. Extraction of RNA and RNA-sequencing

Total RNA was extracted using the TRIzol<sup>®</sup> reagent (Life Technologies, MD, USA). RNA quality was confirmed using an Agilent 2100 bioanalyzer using the RNA 6000 Nano Chip (Agilent Technologies, Amstelveen, The Netherlands), and RNA quantification was performed using an NA-2000 Spectrophotometer (Thermo Fisher Scientific Inc., DE, USA). The TruSeq<sup>™</sup> RNA sample preparation kit (Illumina, CA, USA) was used for mRNA-seq library construction, following the manufacturer's instructions. High-throughput sequencing was performed as paired-end 100 sequencing using the Illumina HiSeq 2500 platform (Illumina, Inc., USA). The high-quality sequences were deposited in the NCBI Sequence Read Archive database with the accession no SAMN22563802-8 and PRJNA774439.

### 2.4. Read filtering, mapping, and expression quantification

Paired-end reads were filtered using a Flexbar (ver. 2.5). The program discarded uncalled bases, adaptor sequences, and short reads. HISAT (ver. 2.0.4) [27] was used to map the filtered reads against the reference genome of *C. militaris* [12]. The genome was 32.2 Mb in size and contained 9651 coding and 159 non-coding genes (*C. militaris* strain CM01 ver. 1.38; <http://fungi.ensembl.org>).

Differentially expressed genes were determined based on counts from unique and multiple alignments using coverage in Bedtools [28]. The read count data were processed based on the quantile normalization method using EdgeR using Bioconductor [29]. The alignment files were also used for assembling transcripts, estimating their abundances, and detecting differential expression of genes or isoforms using cufflinks. We used the fragments per kilobase of exon per million fragments (FPKM) to determine the expression level of the gene regions.

The count data were imported using “DESeqDataSetFromHTSeq” implemented in the tximport package (ver. 1.11.7). Raw counts were normalized using the DESeq2 package (ver. 1.22.2). Time-series analysis was performed using the maSigPro package (ver. 1.54.0). The *p*-values were adjusted for multiple comparisons using false discovery rate (FDR). We used “degree = 2” for three time points or “degree = 3” for four time points in the function “make.design.matrix.” The cutoff value

for the *R*-squared value of the regression model was set as 0.8.

### 2.5. Gene ontology

A weight algorithm with Fisher's exact tests implemented in the R package “topGO” version 2.12.0 was employed to calculate the GO term significance for enrichment analysis of gene sets and visualization of a GO hierarchical structure. GO terms for genes of *C. militaris* were retrieved from the Gene Ontology Annotation database (<http://www.ebi.ac.uk/GOA/>).

### 2.6. Real-time quantitative reverse transcription PCR (qRT-PCR)

RNA-Seq data were confirmed by qRT-PCR. Complementary DNA was prepared from total RNA using ImProm-II<sup>™</sup> Reverse Transcription System (Promega, Madison, WI, USA) according to the manufacturer's recommendation. PCR reactions were carried out in the mixture of cDNA template (25 ng/2 µl), reverse and forward primers (100 nM each in total 3 µl) and Power SYBR<sup>®</sup> Green PCR Master Mix (5 µl, Applied Biosystems, Foster City, CA, USA). The AB7500 Real-Time PCR system was used for amplification with the following condition: 40 cycles of 15 s at 95 °C, 30 s at 60 °C, and 30 s at 72 °C after initial denaturation (Applied Biosystems). The beta-tubulin gene was used as an endogenous control. Primers are listed in Table S1.

## 3. Results and discussion

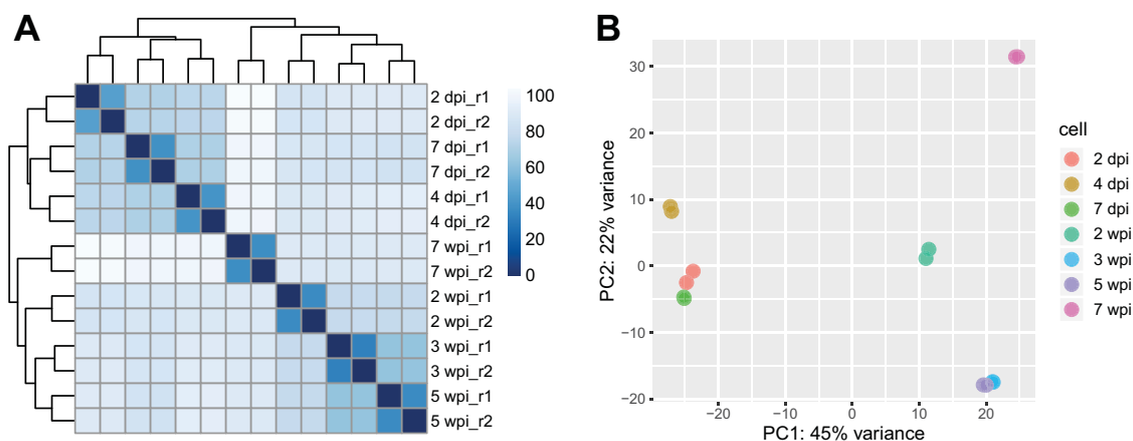
### 3.1. Overall similarity between samples

*Cordyceps militaris* was inoculated on GSCs and cultivated for eight weeks (Figure 1). We chose seven time points to investigate the global transcriptome of GSCs: 2, 4, 7 days, 2, 3, 5, and 7 weeks post-inoculation. Total RNA from seven GSC samples was isolated and sequenced using the Illumina HiSeq platform. After filtration of the reads containing adaptor or low-quality sequences, a total of 205 million paired-end reads were obtained. Approximately 75% of the filtered reads were mapped to the genome of the *C. militaris* strain CM01 (ver. 1.38; <http://fungi.ensembl.org>). As shown in Table 1, 60.8% of the mapped reads were aligned uniquely to the genomic location on average.

To assess the overall similarity between samples, a regularized logarithm transformation of the count data was applied [30]. The 6727 genes with at least ten counts across all samples were used for further analyses. A heatmap was generated based on the

**Table 1.** Summary of RNA-seq data.

	2 day	4 day	7 day	2 week	3 week	5 week	7 week	Total
Total reads (paired)	48,226,772	55,770,080	50,103,936	68,621,040	72,153,088	73,078,594	51,670,582	419,624,092
Filtered reads (paired)	46,715,997	53,698,350	48,826,453	67,354,269	70,937,302	72,092,990	50,845,749	410,471,110
Filtered reads	23,357,995	26,849,173	24,413,224	33,677,130	35,468,635	36,046,482	25,422,864	205,235,503
Mapped reads	16,445,757	19,679,987	20,108,623	26,103,309	28,373,370	23,494,164	19,694,048	153,899,257
% Mapped	70.4	73.3	82.4	77.5	80.0	65.2	77.5	75.0
Uniquely mapped	9,889,762	11,702,291	11,121,764	16,196,262	17,635,783	14,942,931	12,043,059	93,531,852
% Uniquely mapped	60.1	59.5	55.3	62.0	62.2	63.6	61.2	60.8

**Figure 2.** Overall similarity of each sample using a regularized-logarithm transformation. (A) Sample distance matrix using heatmap with Euclidean distance of 6727-gene transcriptome of each sample; (B) Principal component analysis of 6727-gene transcriptome of each sample.

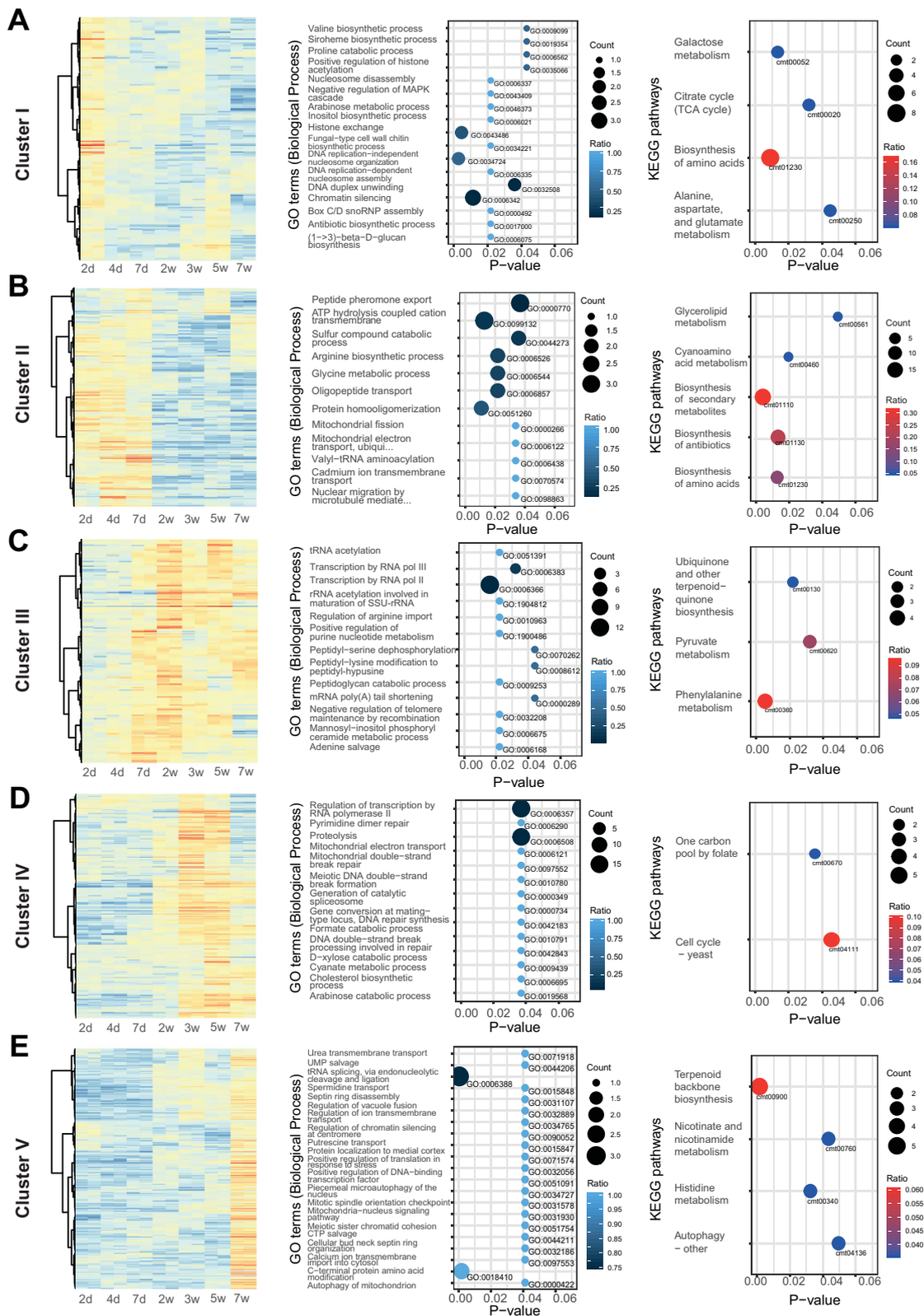
Euclidean distance between samples (Figure 2(A)). On the heatmap, the average distance between samples was 86.2 while the distance between replicates was 38.2. The day samples (2, 4, and 7 dpi) were clustered together with an average distance of 72.3, and apart from the week samples (2, 3, 5, and 7 wpi) with an average distance of 92.2. Additionally, a principal component analysis (PCA) showed similarity as a heatmap (Figure 2(B)). The day samples were far from the week samples. The “2 and 7 dpi” and “3 and 5 wpi” samples were located very close to each other. However, the “7 wpi” sample was far from the other samples based on the PC2 axis (Figure 2(B)).

### 3.2. Clustering of differentially expressed genes

Time-series differential expression analysis was performed to examine the global transcriptomic patterns of the seven GSC samples. The two-step regression strategy in the maSigPro R package was used to find a gene set with differential expression [31]. A total of 1460 significant genes were selected using the following options (degree = 4,  $Q=0.05$ ,  $\theta=5$ , and  $rsq=0.7$ ) and classified into five co-expressed gene clusters ( $k=5$ ). Of the significant genes, 181, 311, 196, 332, and 440 genes were assigned to clusters I–V, respectively (Figure 3). The expression pattern of 10 genes in the clusters was confirmed using qRT-PCR (Figure S1). Two genes

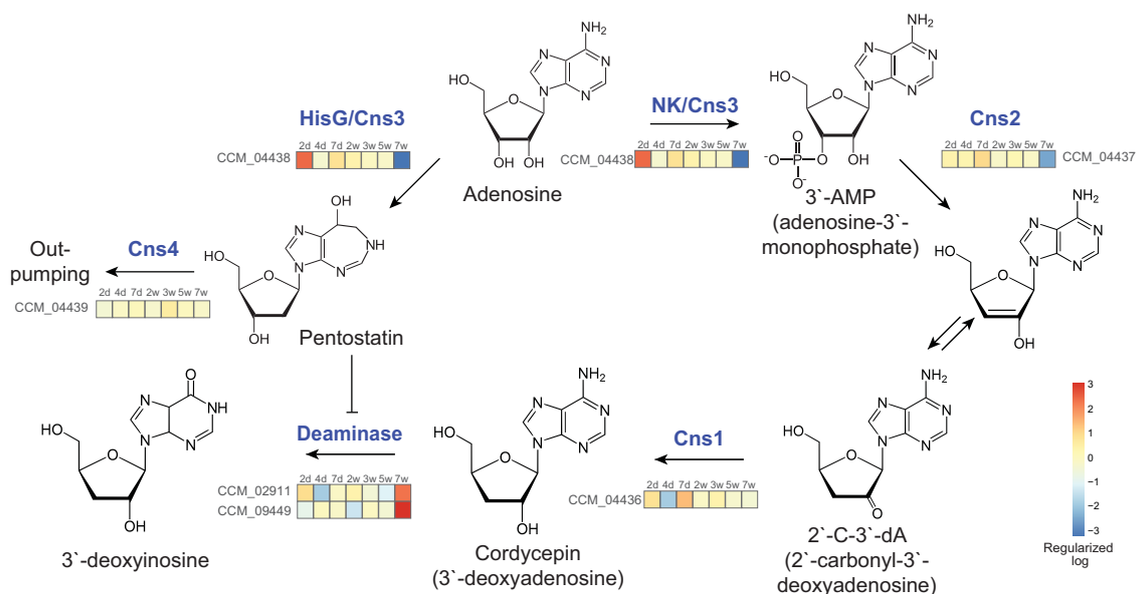
in each cluster Clusters I and II contained genes induced within 1 week, while genes in clusters III, IV, and V showed higher expression at later incubation (2–7 wpi). Interestingly, cluster V was the largest group in size and had highly expressed genes only at 7 wpi (Figure 3(E)). This explains why the “7 wpi” sample is different from the others in the PCA result (Figure 2(B)).

GO and KEGG enrichment analyses were performed for each gene cluster. In cluster I, 17 GOs were identified as significant from the 181 identified DEGs (Figure 3(A), Table S2). Among them, eight GOs were related to chromatin remodeling or reorganization of nucleosomes (GO:0000492, GO:0006335, GO:0006337, GO:0006342, GO:0032508, GO:0034724, GO:0035066, and GO:0043486). Given that the gene expression in cluster I peaked at 2 dpi, the fungal cells might make the transition from a nutrient-rich environment (i.e., PDA) to a new and poor environment (i.e., soybean surface) by remodeling chromatin structure. In particular, the histone chaperone (CCM\_05963, ASF1), a component of the FACT complex (CCM\_05775, POB3), and RuvB-like DNA helicase genes participated in chromatin remodeling. Additionally, biosynthesis of amino acids (GO: 0006021, GO:0006562, GO: 0009099, and GO: 0046373) and cell wall components (GO:0006075 and GO:0034221) were activated, indicating that this fungus was producing basic cell metabolites to grow on the germinated



soybeans. All the KEGG pathways enriched in cluster I were involved in carbohydrate and amino acid metabolism (galactose metabolism, citrate cycle, biosynthesis of amino acids, alanine aspartate, and glutamate metabolism; Table S3).

Cluster II consisted of 311 DEGs that showed strong expression within one week. Five out of 12 GOs in this cluster were involved in both energy and amino acid metabolism (GO:0099132, GO:0006122, GO:0006526, GO:0006544, and



**Cns1 (CCM\_04436): Oxidoreductase**  
**Cns2 (CCM\_04437): Phosphohydrolases**  
**Cns3 (CCM\_04438): ATP phosphoribosyltransferases (HisG and NK are domain names)**  
**Cns4 (CCM\_04439): ATP-binding cassette transporter**  
**CCM\_02911: adenosine deaminase**  
**CCM\_09449: adenosine deaminase**

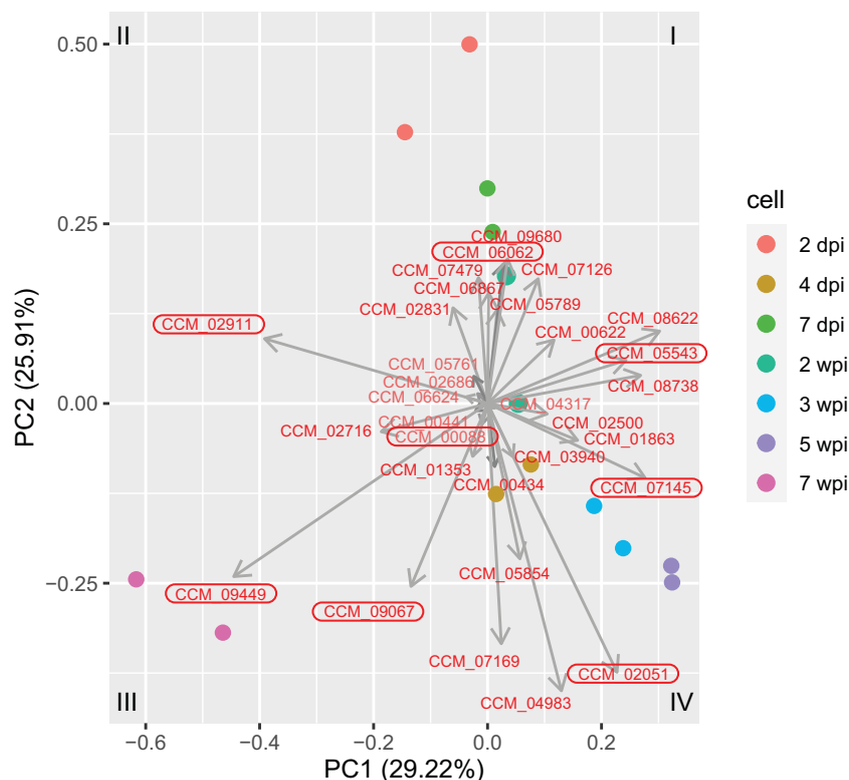
**Figure 4.** Cordycepin biosynthetic pathways. The putative pathway is based on the study by Xia et al. [14]. The heatmap was generated from the regularized-logarithm data after subtraction of the mean expression for each gene. Cns1 (CCM\_04436): oxidoreductase, Cns2 (CCM\_04437): phosphohydrolases, Cns3 (CCM\_04438): ATP phosphoribosyltransferases (HisG and NK are domain names), Cns4 (CCM\_04439): ATP-binding cassette transporter, CCM\_02911: adenosine deaminase, CCM\_09449: adenosine deaminase.

GO:0044273, Table S2). Two out of five KEGG pathways were involved in amino acid metabolism (cmt01230 and cmt00460). Thus, genes involved in the primary metabolism of energy, amino acids, and carbohydrates were included mainly in clusters II and I. This indicates that fungal growth on the germinated soybeans was active within one week of incubation on germinated soybean. Interestingly, the antibiotic (or secondary metabolite) biosynthetic pathway (cmt01130 and cmt01110) was significantly highlighted in cluster II (Figure 3(B)). The ABC multidrug transporter CCM\_07735 is one of the genes involved in this pathway. This gene was highly expressed in the aerial mycelia of *C. militaris* compared to the submerged incubation condition [32].

Cluster III contained 196 DEGs that were continuously expressed throughout the incubation after one week (Figure 3(C)). Thus, these genes may be good genetic markers for GSC cultivation. For example, the CCM\_04506 gene displayed 100-fold higher expression even in all selection points, compared to the “0 day” control. In cluster III, six of the 13 GO terms were related to transcription and translation (GO:0006366, GO:0006383, GO:0006168, GO:0051391, GO:1904812, and GO:0000289) (Figure

3(C)). Three KEGG pathways, phenylalanine metabolism, ubiquinone, and another terpenoid-quinone biosynthesis, and pyruvate metabolism (cmt00369, cmt00130, and cmt00620, respectively), were highlighted. Two genes, CCM\_00270 and CCM\_06000, which were involved in 4-hydroxyphenylpyruvate dioxygenase activities, were found in two KEGG pathways (cmt00369 and cmt00130) in common. The GO and KEGG analyses indicated that the gene expressions for catabolic and anabolic metabolism were active in this cluster.

The genes in cluster IV were up-regulated after two weeks. The biggest GO term in this cluster was related to transcription (GO:0006357, regulation of transcription by RNA polymerase II) (Figure 3(D)). This is consistent with the enriched GO in cluster III. Among the 14 GO terms, five were involved in DNA repair (GO:0000734, GO:0006290, GO:0010780, GO:0010791, and GO:0097552). In particular, three of these GO terms were enriched due to DNA repair protein rad32 (CCM\_2130), which belongs to the MRE11/RAD32 family. Likewise, “cell cycle” (cmt04111) pathway was enriched from the KEGG pathway analysis although a different set of genes contributed to the enrichment of this pathway (CCM\_00489, CCM\_01153,



**Figure 5.** Principal component analysis for the 37 genes involved in cordycepin biosynthesis. The plot was generated from the regularized logarithm values for the count data. The 37 genes were chosen based on the metabolic pathways constructed by Vongsangnak et al. [13]. The eight genes matched to the 1460 significant genes are indicated with rounded rectangles.

CCM\_05144, CCM\_07049, and CCM\_07539 in Table S3).

Cluster V was the largest among the five clusters in terms of the number of DEGs (440 DEGs), where 21 GO terms were highlighted (Figure 3(E)). Contrary to the cluster size in this cluster, GO-annotated genes were rare in this cluster. The most enriched GO term was “tRNA splicing, *via* endonucleolytic cleavage and ligation” (GO: 0006388). This term consisted of four genes, three of which (CCM\_00392, CCM\_00753, and CCM\_09522) belonged to cluster V. The second biggest GO term is “amino acid modification” (GO:0018410). This term contains two DEGs, protein-S-isoprenylcysteine O-methyltransferase (CCM\_05030) and autophagy-related protein 3 (CCM\_06512). CCM\_6512 contributed to two autophagy-related GO terms (GO:0000422 and GO:0034727). Consistently, one of the KEGG pathways was the autophagy pathway (cmt04136), where three genes in cluster V were involved (CCM\_00278, CCM\_00551, and CCM\_06512). The WD-repeat protein (CCM\_00551) also contributed to the mitochondria-nucleus signaling pathway (GO:0031930, Table S2). The last two clusters (IV and V) have many genes involved in DNA repair, proteolysis, and autophagy, suggesting that the culture turns into the death phase where dead cells are more than living cells.

### 3.3. Gene expression of cordycepin biosynthesis pathway

Cordycepin, a structural derivative of adenosine, is produced in *C. militaris*. During GSC cultivation, we found that  $180.58 \pm 1.54$  mg/g cordycepin was produced. Recently, one of the biosynthetic pathways was confirmed using mutagenesis and metabolomics approaches [14]. The characterized pathways and genes are shown in Figure 4. In summary, the precursor adenosine is converted into adenosine-3'-monophosphate (3'-AMP) by the activity of the phosphohydrolase (Cns3, CCM\_04438) NK domain. Phosphohydrolase (Cns2, CCM\_04437) catalyzes 3'-AMP into 2'-carbonyl-3'-deoxyadenosine (2'-C-3'-dA), which is further converted to cordycepin by oxidoreductase (Cns1, CCM\_04436) through a reduction reaction. When cordycepin accumulated within the cell at the toxic level, pentostatin is pumped out through the ABC transporter (Cns4, CCM\_04439) and pentostatin-suppressed deaminases convert cordycepin into the less toxic molecule, 3'-deoxyinosine. We mapped our transcriptome data to this pathway (Figure 4). Among the four genes, *CNS1*, *CNS2*, and *CNS3* were strongly expressed at most cultivation times (average regularized-logarithm values were 3.9, 4.0, and 4.9, respectively). While the expression of *CNS3* was highest at 2 dpi, both *CNS1* and *CNS2* were the

highest at 7 dpi. This time lag makes sense because adenosine is a precursor of cordycepin. The expression of *CNS2* and *CNS3* decreased at 7 wpi. Interestingly, two candidate genes for cordycepin deamination, *CCM\_02911* and *CCM\_09449*, were upregulated at 7 wpi (both belong to cluster V). The patterns of their expressions were opposite to those of *CNS2* and *CNS3*, providing two insights. First, reduction of gene expression reduces the production of both pentostatin and cordycepin directly, lowering intracellular levels. Second, a common transcriptional regulator might contribute to a sudden decrease in the transcripts of *CNS2/CNS3* and, at the same time, a sudden increase in the expression of *CCM\_02911/CCM\_09449*. In agreement with Xia et al. [14], the *CNS4* gene might also function as a suppressor of inactivation through the transport of pentostatin. It showed weak expression during most cultivation times, compared to the other genes (the average regularized-logarithm value was 0.9). However, its expression increased at 3 and 7 wpi, which might increase the out-pumping of pentostatin and activate the deaminases.

In addition, Vongsangnak et al. predicted diverse metabolic pathways in *C. militaris* using a comparative genomic approach [13]. Among the 1267 metabolic reactions, 29 metabolic reactions consisting of 37 genes were related to cordycepin biosynthesis [13]. Eight of the thirty seven genes matched with the 1460 significant genes (*CCM\_00088*, *CCM\_02051*, *CCM\_02911*, *CCM\_05543*, *CCM\_06062*, *CCM\_07145*, *CCM\_09067*, and *CCM\_09449* in Table S4). Seven of the genes were included in clusters III to V, suggesting that these genes could participate in cordycepin biosynthesis late during GSC cultivation. PCA with the 37 genes exhibited separation of “7 wpi” from the other samples (Figure 5). Interestingly, the expression of two deaminases (*CCM\_02911* and *CCM\_09449*) uniquely contributed to the separation of “7 wpi.”

### 3.4. Comparison with metabolomics of GSC cultivation

Previously, metabolomic changes in GSC cultivation were examined using both liquid chromatography and gas chromatography-mass spectrometry [23]. The total flavonoid and phenolic contents (TFC and TPC) in GSC extracts increased after co-cultivation with *C. militaris*, suggesting that GSC showed higher antioxidant activities than germinated soybean only. Isoflavones, such as genistein, daidzein, and glycitein, are major secondary metabolites of soybean and are known to have strong antioxidant effects [33,34]. In nature, isoflavones are coupled with  $\beta$ -D-glucosides, which are called genistin,

daidzin, and glycitin, respectively. They are synthesized through a phenylpropanoid pathway where cinnamic acid, p-coumaric acid, and liquiritigenin are produced in the order of phenyl alanine [35]. Among the genes related to the phenylpropanoid pathway, only two genes, cytochrome P450 oxidoreductase (similar to trans-cinnamate 4-oxygenase, *CCM\_01530*) and glycerol dehydrogenase (similar to calcane reductase, *CCM\_01682*), were found in the *C. militaris* genome (Table 2), indicating that this fungus alone cannot complete isoflavone biosynthesis. In addition, several microbial or plant enzymes involved in the metabolic modification of isoflavones (hydrolysis, hydroxylation, reduction, and methylation) have been reviewed [36]. In Table 2, we compared the key enzymes in the review of the *C. militaris* genome and transcriptome data. For example,  $\beta$ -glucosidase from *Bifidobacterium pseudocatenulatum* (Uniprot id: A0A072MX70) hydrolyzes glycosyl isoflavones into aglycones and isoflavones without attached glycosides [37]. *C. militaris* has several orthologous genes (*CCM\_06730*, *CCM\_05167*, *CCM\_02750*, and *CCM\_07910*) in its genome, three of which were found in the clusters (Table 2). Methylation is also important in the modification of isoflavones, changing the solubility and biological activity [38]. Flavonoid 4'-O-methyltransferase (Uniprot id: C6TAY1) had orthologous genes in *C. militaris* (*CCM\_01923*, *CCM\_07823*, *CCM\_05265*, and *CCM\_04907*). One of them (*CCM\_04907*, O-methyltransferase) was selected in cluster II (Table 2). During GSC cultivation, new forms of isoflavones were reported (daidzein 7-O- $\beta$ -D-glucoside 4''-O-methylate, glycitein 7-O- $\beta$ -D-glucoside 4''-O-methylate, genistein 7-O- $\beta$ -D-glucoside 4''-O-methylate, and genistein 4'-O- $\beta$ -D-glucoside 4''-O-methylate) [23]. For the reason that these new isoflavones have modifications at the isoflavone glycosides instead of those at the main isoflavones, it is still possible that other enzymes might be involved in the modification during GSC cultivation.

## 4. Conclusion

In this study, global transcriptional profiling was conducted during 8-week cultivation of *C. militaris* on germinated soybeans. Time-series transcriptome data revealed that this fungus showed active growth up to 7 dpi and autophagic status at 7 wpi. The genes related to cordycepin biosynthesis were highly expressed, supporting cordycepin production at the transcript level. Our data also support a putative controlling pathway. Isoflavone modification is unique to GSC cultivation. Although genes involved in the modification have not yet been characterized,

**Table 2.** The genes involved in phenylpropanoid pathway and isoflavone modification.

Groups	Uniprot ID	Species	Annotation	CCM ID	Annotation for <i>Corydiceps militaris</i>	Cluster
Phenylpropanoid pathway	P27991	<i>Glycine max</i>	Phenylalanine ammonia-lyase 1	NA	NA	NA
	Q42797	<i>Glycine max</i>	Trans-cinnamate 4-monoxygenase	CCM_01530	Cytochrome P450 oxidoreductase OrdA-like protein	IV
	C6TCY9	<i>Glycine max</i>	Chalcone synthase	NA	NA	NA
	W8JIN8	<i>Glycine max</i>	Chalcone reductase	CCM_01682	Glycerol dehydrogenase	-
	Q53B73	<i>Glycine max</i>	Chalcone-flavonone isomerase family protein	NA	NA	NA
	Q2XQG9	<i>Glycine max</i>	Isoflavone synthase (Fragment)	NA	NA	NA
Isoflavone modification	A0A072MX70	<i>Bifidobacterium pseudocatenulatum</i>	Beta-glucosidase	CCM_06730	Beta-glucosidase	I
				CCM_05167	Beta-glucosidase	-
				CCM_07910	Beta-glucosidase 2 precursor	II
				CCM_08029	Rieske	III
	Q46372	<i>Rhodococcus jostii</i>	Biphenyl-2,3-dioxygenase	CCM_01923	Hydroxyindole O-methyltransferase	-
	C6TAY1	<i>Glycine max</i>	Flavonoid 4'-O-methyltransferase	CCM_07823	Sterigmatocystin 8-O-methyltransferase	-
				CCM_05265	O-methyltransferase	-
				CCM_04907	O-methyltransferase	II
	Q51990	<i>Pseudomonas putida</i>	Morphinone reductase	CCM_08806	NADPH dehydrogenase	-
				CCM_04115	NADH:flavin Oxidoreductase/NADH oxidase family protein	V
Isoflavone modification (continued)				CCM_09024	NADH:flavin Oxidoreductase/NADH oxidase family protein	-
				CCM_06953	NADPH dehydrogenase	V
				CCM_06460	FMN binding oxidoreductase	-
				CCM_05885	FMN binding oxidoreductase	-
				CCM_03850	2,5-diketo-D-gluconic acid reductase A	-
	Q02198	<i>Pseudomonas putida</i>	Morphine 6-dehydrogenase	CCM_06560	Glycerol dehydrogenase	-
				CCM_06364	Aldehyde reductase	II
				CCM_08044	Aldehyde reductase I	-
	AJ011781.1	<i>Neisseria polysaccharea</i>	Amylosucrase gene	CCM_08695	Maltase	I

some candidate genes were predicted bioinformatically. Their expression was confirmed. Our transcriptomic analysis offers a comprehensive understanding of fungal responses to germinated soybeans and provides an idea to obtain bioactive ingredients effectively.

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No potential conflict of interest was reported by the authors.

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