

Exploration of Activated Pathways for Improving Antifungal Agent FR901469 Productivity in Fungal Species No.11243 Using Comprehensive Pathway Model

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Abstract

Secondary metabolites are important for various industrial applications. The production of secondary metabolites is often improved by the activation of substrate supply pathways for biosynthesis. However, many important pathways have remained unclear. In this study, we explored possible pathways related to substrate supply for the biosynthesis of the antifungal agent FR901469 which is a nonribosomal peptide and a fungal secondary metabolite. To clarify the unknown activated pathways, we utilized the Comprehensive Pathway Model (CPM) which was developed in our previous study. We verified that the over-expression of the hypothetical beta-alanine-aminotransferase (BAL-AT), which was included in the explored pathways, improved the FR901469 productivity. The genes encoding the BAL metabolic enzymes are considered to be important for improving the FR901469 productivity.

Keywords

Bioinformatics, Metabolic Pathway Model, Fungi, Secondary Metabolites

1. Introduction

The improvement of secondary metabolite productivity is a crucial task for var-*These authors contributed equally to this work. ious industries. Many secondary metabolites have bioactivities and are important resources as lead compounds for new drugs, active constituents of cosmetics, etc. [1] [2] [3]. Many fungal and bacterial species produce secondary metabolites [4]. In order to improve the productivity of secondary metabolites, some traditional methods, such as culture optimization and high-producing strain development, have been used [5] [6] [7]. Other approaches, in which the secondary metabolite productivity was improved by genetic technology, have been reported recently. For example, the productivity of a secondary metabolite has been enhanced by the overexpression of a transcription factor in a gene cluster [8]. In addition, *laeA*, which is known as a global regulator of secondary metabolism, has been overexpressed [9]. However, many important metabolic pathways for the substrate supply have remained unclear. Secondary metabolites are typically synthesized from primary metabolites, such as amino acids and acetyl-CoA [10]. The biosynthetic pathways of these primary metabolites are considered to be important for improving the productivities of secondary metabolites. Thus, the pathways that are activated according to the production of a secondary metabolite should be explored.

In order to explore all pathways related to the biosynthesis of useful compounds, we developed an algorithm for the construction of a Comprehensive Pathway Model (CPM) in our previous study [11]. Since a cell is regarded as a closed space, the by-products of each enzymatic reaction are used in other enzymatic reactions. The fluctuation of the by-product is considered to affect other enzymatic reactions as well as the main product. Consequently, we constructed the CPM as a metabolic pathway model in which the chemical transformations of the main products and the by-products were considered. The CPM can reveal the pathways hidden behind the main pathways. In our previous study, we explored the by-pass of galactose metabolism in *Saccharomyces cerevisiae* [11], and found some possible pathways related to galactose metabolism. Some *GAL* genes are induced and repressed by the addition of galactose and glucose, respectively [12] [13] [14]. Our explored pathways were composed not only of *GAL* genes but also other genes, which are induced and repressed in the same manner as the known *GAL* genes.

In this study, we applied the CPM to an industrial organism. Recently, the whole genome of fungal species No.11243 (No.11243 hereafter) was determined [15]. The wild type strain No.11243 was isolated from a decayed leaf sample, which was picked up at Ayabe, Kyoto, Japan [15]. This organism produces the antifungal agent FR901469 [16] [17] [18]. FR901469 is water-soluble antifungal lipopeptidolactone [16]. FR901469 is an inhibitor of 1,3-beta-glucan synthase [17]. Antifungal activity of this compound was varified against some funagl species [17]. This compound is cyclic peptide [18]. In the industrial process development of FR901469, some mutant strains with high productivity were obtained from the wild-type strain by UV irradiation [6]. Part of the genealogy is shown in Figure 1(a). The productivities of three mutant strains, #10-10, #1769, and #2889, were measured in the previous study [19]. Their relative FR901469

productivities are shown in **Figure 1(b)**. However, the mechanisms underlying the improved productivity in the mutant strains remained unclear. In this study, we explored the activated pathways that are assumed to be responsible for substrate supply. In order to find an important key gene for improving FR901469 productivity, we selected an enzymatic gene from the explored pathways. We verified that the expression of this enzymatic gene affected the productivity of FR901469.

2. Materials and Methods

2.1. Cultivation

We used special grade chemicals from Wako, unless otherwise specified. Seed medium (20 mL), containing 4% soluble starch, 2% cottonseed flour, 1% soybean flour, 1% KH_2PO_4 , and 0.2% $CaCO_3$, was poured into 100-mL Erlenmeyer flasks and sterilized at 121°C for 20 min. A quadrangular (5 mm × 5 mm) plate culture was inoculated into each flask and cultured under shaking conditions (220 rpm) at 25°C for 5 days.

Production medium (20 mL), consisting of 7% soluble starch, 1% glucose, 4% cottonseed flour, 6% corn steep liquor (Sigma-Aldrich), 1% $(NH_4)_2SO_4$, 0.5% BactoTM Peptone (Becton, Dickinson and Company), 0.44% K₂HPO₄, 1.0% KH₂PO₄, 0.1% antiform PE-L (Wako), and 0.1% antiform SI (Wako), was adjusted to pH 6.0 by adding 1 M NaOH, poured into each 100-mL Erlenmeyer flask, and sterilized at 121°C for 20 min. The resultant seed culture (2 mL) was transferred to the production medium and cultured under shaking conditions (220 rpm) at 25°C for 6 days.

In this work, we cultured one mutant No.11243 strain [6], which was obtained from the wild type strain (FERM BP-3373), and one transformant obtained in this study: #10-10 (Figure 1(a)) and $#10-10+ANO11243_080900$.



Figure 1. Mutant strains of fungal sp. No.11243 and relative FR901469 productivities. (a) The genealogy of the mutant strains of No.11243. (b) The relative productivities of the mutant strains. The bar graph represents mean values of three replicate experiments. The error bars are Mean value \pm Standard error of three replicate experiments.



2.2. Analysis of FR901469 Productivity

To extract FR901469, 200 μ L of liquid medium from 4 and 6 day cultures was placed into 1.5-mL Eppendorf tubes, mixed with 800 μ L ethanol, and incubated for 30 min at room temperature. The debris was removed by centrifugation at 15,000 rpm at room temperature for 10 min, and the supernatant was filtered using 0.22- μ m pore size filter units (NacalaiTesque). A 2- μ L aliquot of the extract was separated on an Acquity UPLC system (Waters, MA, USA) with an Acquity UPLC BEH C18 1.7 μ m 2.1 × 50 mm reversed-phase column (Waters), and eluted with a 36% - 46% aqueous acetonitrile (Wako, for LC/MS) gradient containing 0.1% formic acid at a flow rate of 0.6 mL/min. The production of FR901469 was quantified by the peak area of absorbance at 210 nm.

2.3. RNA Sequencing (RNA-Seq)

To isolate RNA, we used the 4 and 6 day production cultures. Mycelia in 2 mL of the production culture were collected by filtration, using Miracloth. The mycelia were washed with Milli-Q water, frozen by liquid nitrogen, and pulverized in a mortar. The total RNA was extracted from the frozen pulverized mycelia using 1.5 mL ISOGEN (Nippon Gene, Tokyo, Japan), and purified with an RNeasy mini kit (Qiagen, Venlo, Netherlands). The Truseq Stranded mRNA and Total RNA Sample Preparation Kit (Illumina, CA, USA) were used for library preparation from the purified RNA, and the libraries were sequenced by MiSeq (Illumina). We registered the RNA-seq data of #10-10 and #10-10+ANO11243_080900 in the DDBJ Sequence Read Archive (Accession numbers #10-10: DRX076195-DRX076200, #10-10+ANO11243_080900: DRX076201-DRX076206).

2.4. Sequence Data Processing

In this work, we used five RNA-Seq data sets. For constructing the activated pathway models, the three data sets of mutant strains #10-10, #1769, and #2889 registered in the DDBJ Sequence Read Archive were used (Accession numbers #10-10: DRX040488-DRX040490, #1769:DRX040524-DRX040526, #2889: DRX040566-DRX040568). For evaluating the activated pathways, the two data sets of #10-10 and #10-10+ANO11243_080900 from the 6 day culture periods were used (Accession numbers #10-10: DRX076198-DRX076200, #10-10+ ANO11243_080900: DRX076204-DRX076206). The reference genome was indexed by Bowtie 2 [20], and high-quality reads were mapped to the No.11243 genome by TopHat 2 [21]. By using Genome Traveler (Insilico Biology), the Reads Per Kilobase of exon model per Million mapped reads (RPKM) values were calculated, as the expression level of each gene. For the #10-10 data set, the mean RPKM value of three replicate experiments was calculated for each gene. In the same way, the mean RPKM values were also calculated for the #1769 and #2889 data sets. Using the data sets of the mean RPKM values, the Global Normalization was performed between #10-10 and #1769. The fold-change (FC) values of #1769 against #10-10 were calculated from the normalized RPKM values in each gene. The FC values were converted to the base-2 logarithmic values.

The fold-change values $\log_2(FC)$ of #2889 against #10-10 were also calculated in the same way.

2.5. Construction of Fungal sp. No.11243-Specific CPM

The No.11243-specific CPM (11243CPM) was constructed (Figure 2(a)). In the KEGG database, the K number was assigned to each ortholog group [22]. Each node that was contained in the CPM corresponded to a K number. By using KAAS [23], we assigned K numbers to the genes encoded in the No.11243 genome. The amino acid sequences of the deduced genes were retrieved from the DDBJ/EMBL/GenBank database (accession nos. DF938580~938599). Following the assignment of the K numbers, the whole CPM was constructed from KEGG (ver. 2014-04-07 release). In the CPM, the nodes and arcs corresponded to enzymes and compounds, respectively. In order to construct a 11243CPM, two simplification processes were performed as follows: 1st: Elimination of excess compounds. Some arcs indicating excess compounds were eliminated from the whole CPM [11]. Some specific compounds, for example H₂O and oxygen, connected many pairs of enzymes in the CPM. We think that large fluctuations of these compounds rarely occur in a cell. Since these compounds are usually essential for growth, sufficient amounts are assumed to exist in a cell. Thus, the arcs corresponding to these compounds were eliminated from the CPM. 2nd: Selection of organism-specific enzymes. Some nodes were selected from the CPM. The selected nodes corresponded to K numbers that were assigned to the genes encoded in the genome of No.11243. In this way, 11243CPM was constructed. The 11243CPM was defined as graph D_{11243} ,

$$D_{11243} = \left(V_{11243}, A_{11243}\right) \tag{1}$$

where V_{11243} and A_{11243} denote nodes and arcs, respectively. In D_{11243} , the nodes correspond to the enzymatic genes encoded in the No.11243 genome.

2.6. Construction of Activated Pathway Model and Exploration of Pathways

The activated pathway model was constructed (Figure 2(b), Figure 2(c)). This model was defined as the graph D_{act} . In order to construct D_{acp} we used the FC values of gene expression. The FC values of a strain with high productivity against a strain with low productivity were used. The nodes corresponding to the gene with an FC value higher than the threshold were selected from D_{11243} . The nodes that became alone by this process were eliminated. If some enzymatic genes were assigned to a node, then the gene that had the biggest fold-change value was adopted. The D_{act} is a subgraph of D_{11243} .

$$D_{act} = \left(V_{act}, A_{act}\right) \tag{2}$$

where $V_{act} \subseteq V_{11243}$ and $A_{act} \subseteq A_{11243}$ are satisfied. In this study, the gene expression changes were used from #10-10 to #1769 and from #10-10 to #2889. The constructed graphs were defined as D_{act}^{1769} and D_{act}^{2889} , respectively (Figure 2(b), Figure 2(c)).





Figure 2. Flowchart for the construction of the fungal sp. No.11243-specific model and other activated pathway models. (a) Fungal sp. No.11243-specific model. From the whole CPM, this model was constructed by two simplification processes. (b) The activated pathway model of #1769 against #10-10. This model was constructed from D_{11243} , by using gene expression profiles. (c) The activated pathway model of #2889 against #10-10. (d) The conserved activated pathway model between #1769 and #2889. The arcs that were included in both D_{act}^{1769} and D_{act}^{2889} were kept.

In order to explore the pathways that affected the productivity with high probability, the conserved activated pathways were constructed as a model (Figure 2(d)). Since random mutations occurred in the genome of the mutant strain, various genes were assumed to be induced. In order to select the genes that affect the productivity with high probability, the conserved activated pathway model was constructed. This model was defined as the graph $D_{act}^{1769,2889}$. The graph $D_{act}^{1769,2889}$ contains the common arcs between D_{act}^{1769} and D_{act}^{2889} :

$$D_{act}^{1769,2889} = \left(V_{act}^{1769,2889}, A_{act}^{1769,2889} \right), \text{ where, } A_{act}^{1769,2889} = A_{act}^{1769} \cap A_{act}^{2889}$$
(3)

The $V_{act}^{1769,2889}$ is the set of nodes that are connected by the arcs contained in the $A_{act}^{1769,2889}$.

Following the construction of $D_{act}^{1769,2889}$, we explored pathways that could be responsible for the substrate supply of FR901469 biosynthesis. FR901469 is considered to be biosynthesized from various types of amino acids [18]. Thus, the biosynthetic pathways of the substrate amino acids should be explored. The nodes corresponding to the enzymes that catalyzed the biosynthetic reactions of the substrate amino acids were defined as the beginning nodes of exploration. From the beginning nodes, the interactions between nodes were traced in the reverse direction. For the exploration of the pathways, we modified our developed Path Finding Algorithm [11]. Using this algorithm, the possible interactions between the nodes located within the constrained paths were explored comprehensively. Only the beginning node was fixed in this study, while both the beginning and end nodes were fixed in our previous study.

2.7. Overexpression of Target Gene

Overexpression plasmids were constructed by fusing four DNA fragments (ANO11243 065790 alcohol dehvdrogenase promoter, ANO11243 080900 coding sequence and 300 bp downstream region (termination), Aureobasidin A resistance gene from pAUR316 (Takara), and the pUC19/BamH I-cleaved vector (Figure 3(a)), using the In-Fusion HD Cloning Kit (Clontech, CA, USA). PCR was performed using KOD-Plus DNA polymerase (Toyobo, Osaka, Japan). The primer sets are listed in Table 1.

The preparation of protoplasts and the polyethylene glycol (PEG) transformation methods were described previously [8]. In the selection of transformants by drug resistance, Aureobasidin A (Clontech, CA, USA) was added to the regeneration agar medium, at a final concentration of 0.5 µg/mL. To confirm DNA fragment insertion, the genomic DNA of the transformant was isolated with a NucleoSpin Plant II kit (Macherey-Nagel, Düren, Germany), according to the manufacturer's protocol. The isolated genomic DNA (20 ng) was amplified by PCR, using KOD-Plus DNA polymerase and the primer pair pUC19_pADH_F and AUR_080900_R (Table 1). The PCR reaction solution (1 µL) was electrophoresed in a 0.7% (wt/vol) agarose gel containing SYBR safe DNA gel stain (Invitrogen), at a constant voltage (100 V) for 30 min in Tris-acetate-EDTA buffer (pH 8.3). The 500 bp DNA ladder dye plus (TaKaRa) was used as a marker.



Figure 3. Construction of overexpression plasmid and PCR analysis. (a) Three DNA fragments, the endogenous alcohol dehydrogenase promoter (ANO11243_065790), the ANO11243_080900 coding/termination sequence, and the Aureobasidin resistance gene, were inserted into the pUC19 BamH I-cleaved vector by in-fusion cloning. (b) Gel electrophoresis analysis of PCR check for transformants. The primer and the amplified chain length are demonstrated in Figure 3(a). Lanes M: 500 bp DNA ladder marker (Takara), 1: #10-10, 2: #10-10+ANO11243_080900.



3. Results

3.1. Fungal sp. No.11243-Specific CPM and Activated Pathway Models

In order to explore the pathways related to FR901469 productivity, we constructed 11243CPM. In total, 1,999 K numbers were assigned to 2,172 genes by using KAAS. Using the information from the assigned K numbers, the graph D_{11243} was constructed. The D_{11243} was constructed from 549 nodes and 6,933 arcs (**Table 2**). The whole image of D_{11243} is shown in **Figure 4(a)**. In this image, nodes and arcs represent enzymes and compounds, respectively. The multiplexing arcs between nodes are shown as single arcs.

In order to represent the dynamics of the metabolism, we constructed the activated pathway models D_{act}^{1769} and D_{act}^{2889} . The gene expression profiles were measured by RNA-Seq. The expression changes were calculated from #10-10 to #1769, and from #10-10 to #2889. The nodes with a base-2 logarithmic foldchange value of 1.0 and over were kept in the D_{act}^{1769} and D_{act}^{2889} . In this way, D_{act}^{1769} and D_{act}^{2889} were constructed. The D_{act}^{1769} and D_{act}^{2889} contained 44 and 80 nodes, respectively (**Table 2**). The whole images of D_{act}^{1769} and D_{act}^{2889} are shown in **Figure 4(b)** and **Figure 4(c)**, respectively.

We constructed the conserved activated pathway model $D_{act}^{1769,2889}$ (Figure 4(d)). The #1769 and #2889 were developed independently from #10-10 (Figure 1(a)). Even though #1769 and #2889 independently contained the factor for improving the productivity, a common subgraph was found between D_{act}^{1769} and D_{act}^{2889} . Thus, it is considered that the activated pathways in $D_{act}^{1769,2889}$ probably affect the FR901469 productivity. The graph $D_{act}^{1769,2889}$ comprised 30 nodes and 82 arcs, which were conserved between D_{act}^{1769} and D_{act}^{2889} (Table 2).

Primer name	Primer sequence
pUC19_pADH_F	5'-CGGTACCCGGGGATCCCCTCAACCCAGCCCTGTC
pADH_R	5'-GATCGAATCAAGTATGAATGTCGTTCAG
AUR_F	5'-CTGATGGTCAGATGGATCTGCC
AUR_pUC19_R	5'-CGACTCTAGAGGATCTTGGCTTGTTATGACTATGT ATACATATGCG
pADH_080900_F	5'-ATACTTGATTCGATCATGCCTGCCATGCCATCTGTTTC
AUR_080900_R	5'-CCATCTGACCATCAGAACAACACTCGCTACGGACTCGG

Table 2. Overview of each model.

	# of Nodes ^a	# of Arcs ^b	Cpd. ^c
$D_{_{11243}}$	549	6,933	588
$D_{\scriptscriptstyle act}^{\scriptscriptstyle 1769}$	44	124	53
$D_{_{act}}^{_{2889}}$	80	282	114
$D_{_{act}}^{_{1769,2889}}$	30	82	39

^aNumber of nodes in each graph; ^bNumber of arcs in each graph; ^cKinds of compounds in each graph.



Figure 4. Whole Image of 11243CPM. (a) The whole image of D_{11243} is shown. In this image, the green nodes are common nodes between D_{act}^{1769} and D_{act}^{2889} . The yellow nodes are unique nodes in the D_{act}^{1769} . The blue nodes are unique nodes in the D_{act}^{2889} . (b) The whole image of D_{act}^{1769} . (c) The whole image of D_{act}^{2889} . (d) The whole image of $D_{act}^{1769,2889}$.

3.2. Exploration of Substrate Supply Pathways for FR901469 Biosynthesis

In order to predict the important substrate supply pathways for FR901469 biosynthesis, we explored the activated pathways from $D_{act}^{1769,2889}$, while assuming that the FR901469 productivity is affected by the amounts of substrates available for biosynthesis. Thus, we explored the pathways related to the biosynthesis and decomposition of the amino acids that are substrates of FR901469. In the CPM, it is assumed that all enzymatic reactions have two-directions. Consequently, the pathways were explored without any distinction between biosynthesis and decomposition. The beginning nodes of exploration were determined in $D_{act}^{1769,2889}$. The cyclic peptide, FR901469, is thought to be synthesized from nine kinds of amino acids, L-Alanine, L-Glutamine, Glycine, L-Ornithine, L-Proline, L-Threonine, L-allo-Threonine, L-Tyrosine, and L-Valine [18]. We searched for the enzymes catalyzing the reaction with formulae containing the substrate amino acids. From $D_{act}^{1769,2889}$, 3 ortholog groups were found. The nodes corresponding to these 3 ortholog groups were defined as beginning nodes. From these beginning nodes, the interactions between the nodes were traced in the reverse direction. In this way, the activated pathways that reached these nodes were explored comprehensively. As a result, the maximum length of the found pathways was 3 steps. The pathways that reached the biosynthesis of L-Alanine and L-Tyrosine are shown in Figure 5. The same structures are shown in Figure 5(a) and Figure 5(b), for the common activated pathways in #1769 and #2889, respectively. In Figure 5, the activated pathways reached the nodes corresponding to K00814







Figure 5. Conserved activated pathways. All pathways explored from $D_{act}^{1769,2889}$ are shown. The labels of the nodes correspond to the K numbers in the KEGG database. These pathways reach the nodes corresponding to K00814 and K00838. (a) The fold-change values $\log_2(\#1769/\#10-10)$ are reflected to the explored pathways. The degrees of induction are represented by the colors of the nodes. Bright green represents large induction. White is used for genes with unchanged expression. Red represents repression. (b) The fold-change values $\log_2(\#2889/\#10-10)$ are reflected to the explored pathways. The color code is the same as in **Figure 5(a)**.

Ortholog Group	Gene	log ₂ (1769/10-10) ^a	log ₂ (2889/10-10) ^b	log ₂ (10-10+ANO11243_080900/10-10) ^c
K00814	ANO11243_005270	1.46	2.61	2.27
K00838	ANO11243_007360	2.25	2.60	0.22
K00128	ANO11243_066380	3.23	4.17	0.57
K13524	ANO11243_080900	6.60	4.47	10.00
K01469	ANO11243_009840	1.41	1.63	-0.34
K01580	ANO11243_077720	1.90	1.79	1.29
K01652	ANO11243_092160	1.68	2.39	0.38
K00129	ANO11243_080890	7.27	7.07	8.64
K00276	ANO11243_080880	3.06	3.27	1.16
K14163	ANO11243_034720	1.93	1.99	1.73

Table 3. Enzymatic genes in the common activated pathways.

^aLogarithmic induction ratio of 1769 to 10-10; ^bLogarithmic induction ratio of 2889 to 10-10; ^cLogarithmic induction ratio of 2889 to 10-10+*ANO*11243_080900.

3.3. Verification of the Effect of a Selected Gene from the Explored Pathways

In order to find the important key gene for improving productivity, we selected the gene, which was assumed to affect the FR901469 productivity, from the explored pathways. In order to select the target gene, we reflected the fold-change value of #1769 against #10-10, and #2889 against #10-10 to the explored pathways (Figure 5(a), Figure 5(b)). Each node was colored according to the foldchange value. The fold-change values corresponding to each node are described in **Table 3**. The 10 enzymatic genes that were included in these conserved activated pathways were assumed to affect the FR901469 productivity. Then, we tried to identify the key gene that affects the productivity the most among these 10 genes, by considering the following points. First, the expression of the genes was induced largely in the mutant strains. It is assumed that the inductions of the expression of these genes were important for improving the FR901469 productivity. Second, the nodes were connected with many nodes. The enzymatic gene corresponding to such a node is assumed to affect many other enzymatic reactions. Among the 10 genes, the ANO11243_080890 and ANO11243_ 080900 genes were induced ten-fold in both #1769 and #2889 (Table 3). Furthermore, the node corresponding to ANO11243_080900 had more connections than the node corresponding to ANO11243_080890. Consequently, we considered that ANO11243_080900 was the most likely candidate for improving the FR901469 productivity, among the 10 genes.

We empirically evaluated the effect of this selected gene by overexpression. First, ANO11243_080900 was overexpressed in the #10-10 strain (Figure 3(b)). The strain in which ANO11243_080900 was overexpressed was defined as #10-10+ANO11243_080900. Second, the FR901469 productivities of #10-10 and #10-10+ANO11243_080900 were evaluated in the same way as the other strains. The results confirmed that #10-10+ANO11243 080900 produced about 1.5-fold more FR901469 than #10-10 (Figure 6(a)). Finally, we performed RNA-Seq for #10-10 and #10-10+ANO11243 080900, under the same conditions used for #1769 and #2889. We determined the RPKM values and calculated the foldchange values of #10-10+ANO11243 080900 against #10-10. The fold-change values log₂(FC) were reflected to the explored pathways (Figure 6(b)). The ANO11243_080900 expression was induced by 1,022-fold. In Figure 6(b), the node corresponding to ANO11243 080900 is represented as a square node. Therefore, it was confirmed that the ANO11243_080900 gene affected the FR901469 productivity. In this study, only ANO11243 080900 was overexpressed. However, some other genes, for example ANO11243_080890 and ANO11243 005270, were also induced.

4. Discussion

In the explored pathways, we compared the degrees of gene expression induction between #1769 and #2889, and found that the degrees of induction were similar. The pathways for substrate supply were explored from $D_{act}^{1769,2889}$. The



Figure 6. Relative FR901469 productivity and activated pathways in the #10-10+ANO11243_080900 transformant. (a) The relative productivities of #10-10 and 10-10+ANO11243_080900 after a 6 day culture. The bar graph represents mean values of three replicate experiments. The error bars are Mean value \pm Standard error of three replicate experiments. *P = 0.043. (b) The fold-change values log₂(#10-10+ANO11243_080900/#10-10) are reflected to the explored pathways. The color code is the same as in **Figure 5**. The node corresponding to ANO11243_080900 is represented as a square node. The nodes framed in blue are BAL metabolism genes.

explored pathways were constructed from 10 genes (Figure 5, Table 3). The fold-change values corresponding to each node are described in Table 3. Between #1769 and #2889, the Pearson's correlation coefficient, which was calculated from these 10 data points, was approximately 0.92. Even though the parent of #1769 and #2889 branched at #10-10, the induction patterns of the configuration genes were similar. Thus, the alteration of these induction patterns is important for improving the FR901469 productivity.

We considered that the beta-alanine (BAL) metabolic genes affected the FR901469 productivity. In this study, we assigned K13524 to ANO11243_ 080900. Since the gamma-aminobutyric acid aminotransferase (GABA-AT) and BAL aminotransferase (BAL-AT) reactions are catalyzed by the same enzyme in human [24], K13524 corresponded to GABA-AT and BAL-AT in the KEGG database. Thus, K13524 has connections with both the GABA and BAL metabolic genes in our CPM. GABA-AT is known as a transaminase catabolizing GABA to succinate semialdehyde (SSA) in the GABA-shunt, an alternative route for the conversion of alpha-ketoglutarate to succinate [25]. The amino-acid sequence of ANO11243 080900 shows strong overall similarity to the sequences of the GABA-AT enzymes from Aspergillus nidulans GatA and Saccharomyces cerevisiae Uga1p, with identities of 67% and 53%, respectively. GatA is conserved in Aspergillus species. When grown with GABA as the sole nitrogen source, the expression of GatA in A. oryzae was strongly induced [26]. Upon the overexpression of ANO11243 080900 in this study, the GABA-shunt related genes, ANO11243_054840 (K01580) and ANO11243_010070 (K00135), which catabolize L-glutamate to GABA and SSA to succinate, respectively, remained uninduced $(\log_2(FC) = -0.37 \text{ and } -1.19, \text{ respectively})$. In contrast, the beta-alanine metabolic genes, ANO11243 080890 (K00129) and ANO11243 080880 (K00276), were co-upregulated by the ANO11243_080900 overexpression ($\log_2(FC) = 8.64$ and 1.16, respectively) (Table 3). Quite interestingly, these three genes including ANO11243 080900 are clustered on the genome, which strongly suggests that ANO11243 080900 is a member of the beta-alanine metabolism genes. Considering the co-upregulation and the co-localization of these three genes, the ANO11243_080900 product may be involved in the BAL metabolism (Figure 7). The overexpression of ANO11243 080900 could have co-upregulated the expression of the two genes encoding K00129 and K00276, which might stimulate the conversion between 1,3-diamino-propane and malonic semialdehyde through BAL metabolism. Considering the finding that the three genes are commonly induced in the two mutants, #1769 and #2889, activating the BAL transamination reaction could be at least one of the effective means to enhance the production of this compound, possibly by increasing particular or overall amino acid biosynthesis.

Furthermore, fungal sp. No.11243 has a duplicated gene encoding K13524, which is $ANO11243_016490$. The expression of this gene remained uninduced in #1769, #2889, and $ANO11243_080900$ overexpression ($\log_2(FC) = 0.80, 0.49$, and -0.13, respectively). In fungi, some species reportedly have gene duplications of K13524, but the substrates for only a few enzymes have been verified in detail. In *Saccharomyces kluyveri*, the functions reportedly differ between the duplicated genes. One of them catalyzed the reaction as GABA-AT and the other catalyzed the reaction as BAL-AT [27]. Therefore, the two duplicated genes assigned to K13524 in this study may also have different functions.

ANO11243 080880 ANO11243 080890 ANO11243 080900 Fungal sp. No. 11243 genome H₂N NH₂ H_oN \cap Malonate K00129 K13524 K00276 Beta-Aminopropion **Beta-Alanine** 1,3-Diaminopropane semialdehyde aldehyde HO ∩н $\overline{N}H_2$ 2-Oxo-glutarate L-Glutamate

Besides the above-mentioned BAL-related genes, the FR901469 productivity might be improved by the overexpression of other target genes. The fold-change

Figure 7. Location of BAL metabolite genes on the fungal sp. No.11243 genome and the BAL metabolite reaction. *ANO*11243_080880, *ANO*11243_080890, and *ANO*11243_080900 form a cluster on the genome. The encoded proteins catalyze the BAL metabolic pathway centered on Beta-alanine. *ANO*11243_080880 and *ANO*11243_080890 were co-upregulated with *ANO*11243_080900 overexpression.

values of #10-10+ANO11243_080900 against #10-10 were reflected to the explored pathways (**Figure 6(b**)). The fold-change values corresponding to each node are described in **Table 3**. Upon the overexpression of ANO11243_080900, the ANO11243_005270, ANO11243_077720, and ANO11243_034720 genes were also induced by more than two-fold. The expressions of the ANO11243_005270 and ANO11243_077720 genes were induced equally to those in the other two mutant strains. In contrast, the expression of other genes, such as ANO11243_07360 and ANO11243_092160, were scarcely induced. It is assumed that the productivity in the #10-10+ANO11243_080900 strain increases to the productivity levels in the two mutant strains by the overexpression of these genes.

In this study, we applied the CPM to the genome of fungal sp. No.11243, which produces FR901469. Some mutant strains of No.11243 have been developed independently. The conserved activated pathway model $D_{act}^{1769,2889}$ was constructed by the combination of the gene expression profiles measured in these mutant strains. Consequently, the important pathways that affect the FR901469 productivity were predicted, by using the characteristics of the independent development. We can improve the productivity of secondary metabolites effectively from many candidate pathways, which may affect the productivity with high probability. Thus, the development of many independent mutant strains is one of the important methods for exploring the key genes in bioproduction. The various target genes for improving the secondary metabolite productivity will be contained in the explored pathways. Using a genetic technology approach to identify these genes will facilitate the development of highly productive strains.

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