

US008871494B2

(12) United States Patent

Keller et al.

(54) OVER-PRODUCTION OF SECONDARY METABOLITES BY OVER-EXPRESSION OF THE VEA GENE

(75) Inventors: Nancy P. Keller, Madison, WI (US);

Saori Amaike, Madison, WI (US)

(73) Assignee: Wisconsin Alumni Research

Foundation, Madison, WI (US)

(*) Notice: Subject to any disclaimer, the term of this

patent is extended or adjusted under 35

U.S.C. 154(b) by 654 days.

(21) Appl. No.: 12/799,505

(22) Filed: **Apr. 26, 2010**

(65) **Prior Publication Data**

US 2011/0076682 A1 Mar. 31, 2011

Related U.S. Application Data

(60) Provisional application No. 61/172,514, filed on Apr. 24, 2009.

(51) Int. Cl. C12N 1/00 (2006.01)

C12Q 1/68 (2006.01)

(52) **U.S. CI.** CPC *C12Q 1/6895* (2013.01); *C12Q 2600/158*

USPC 435/254.3; 435/243; 435/254.11

(2013.01)

(58) Field of Classification Search

None

See application file for complete search history.

(56) References Cited

U.S. PATENT DOCUMENTS

4,469,863 A	9/1984	Ts'o et al.
4,683,195 A	7/1987	Mullis et al.
5,034,506 A	7/1991	Summerton et al.
5,216,141 A	6/1993	Benner
5,235,033 A	8/1993	Summerton et al.
5,386,023 A		Sanghvi et al.
5,602,240 A	2/1997	De Mesmaeker et al.
5,637,684 A	6/1997	Cook et al.
5,644,048 A	7/1997	Yau
5,952,174 A	9/1999	Nikiforov et al.

OTHER PUBLICATIONS

Dreyer et al., Applied and Environmental Microbiology, 2007; 73(10): 3412-22.*

Bowie et al (Science, 1990, 257:1306-1310).*

Adams, et al., Microbiol. Mol. Biol. Rev. 62:35-54 (1998).

Bayram, et al., Fungal Genet. Biol. 45:127-138 (2008).

Bayram, et al., Science 320:1504-1506 (2008).

Brakhage, FEMS Microbiol. Lett. 148:1-10 (1997).

Bok, et al., Eukaryotic Cell 3:527-535 (2004).

Bok, et al., Eukaryotic Cell 4:1574-1582 (2005).

Bok, et al., Mol. Microbiol. 61:1636-1645 (2006).

Brodhagen, et al., Mol. Microbiol. 67:378-391 (2008).

Busch, et al., Mol. Microbiol. 49:717-730 (2003).

Busch, et al., Proc. Natl. Acad. Sci USA 104:8089-8094 (2007).

Calvo, et al., Microbiol. Mol. Biol. Rev. 66:447-459 (2002).

(10) Patent No.: US 8,871,494 B2 (45) Date of Patent: Oct. 28, 2014

Calvo, et al., Appl. Environ. Microbiol. 65:3668-3673 (1999). Calvo, et al., Appl. Environ. Microbiol. 70:4733-4739 (2004).

Chaveroche, et al., Nucleic Acids Res. 28:e97 (2000).

Cho, et al., The Journal of Microbiology 41:46-51 (2003).

Champe, et al., J. Gen. Microbiol. 133:1383-1387 (1987).

Chen, et al., Proc. Natl. Acad. Sci. USA 101:5048-5052 (2004).

Chen, et al., Genes Dev. 20:1150-1161 (2006).

Cseke, et al., Handbook of Molecular and Cellular Methods in Biology and Medicine, 2nd ed., CRC Press, Boca Raton, FL (2004), Index

Dagenais, et al., Infect. Immun. 76:3214-3220 (2008).

Diener, et al., Ann. Rev. Phytopathol. 25:249-270 (1987).

Dreyer, et al., Appl. Environ. Microbiol. 73:3412-3422 (2007).

Duran, et al., Appl. Microbiol. Biotechnol. 73:1158-1168 (2007).

Duran, et al., Open Mycol. J. 3:27-36 (2009).

Eng, et al., J. Am. Soc. Mass. Spectrom 5:976-989 (1994).

Etxebeste, et al., Eukaryotic Cell 7(1):38-48 (2008).

Feinberg, et al., Anal. Biochem. 132:6-13 (1983).

Fernandez-Abalos, et al., Mol. Microbiol. 27(1):121-130 (1998).

Fray, Ann. Bot. 89:245-253 (2002).

Georgianna, et al., Fungal Genet. Biol. 46:113-125 (2009).

Ausubel, et al. (ed.), Current Protocols in Molecular Biology, Contents, vols. 1-5, John Wiley & Sons, Inc. (2007), Index only.

Gyuris, et al., Cell 75:791-803 (1993).

Greenspan, et al., J. Cell Biol. 100:965-973 (1985).

Hanahan, et al., Methods Enzymol. 204:63-113 (1991).

Hicks, et al., The Mycota XI, Kempken (Ed.), Springer-Verlag Berlin Heidelberg, pp. 55-69 (2002).

Hu, et al., Mol. Cell 9:789-798 (2002).

Hornby, et al., Appl. Environ. Microbiol. 67:2982-2992 (2001).

Brown, et al., Appl. Environ. Microbiol. 74:5674-5685 (2008).

Jensen, et al., Appl. Environ. Microbiol. 58:2505-2508 (1992).

Kale, et al., Fungal Genet. Biol. 45:1422-1429 (2008).

Kato, et al., Eukaryotic Cell 2:1178-1186 (2003).

Kim, et al., Fungal Genet. Biol. 37:72-80 (2002). Klich, Mol. Plant Pathol. 8:713-722 (2007).

Keller, et al., Nat. Rev. Microbiol. 3:937-947 (2005).

Keller, et al., Appl. Environ. Microbiol. 60:1444-1450 (1994).

Kolar, et al., Gene 62:127-134 (1998).

Krappmann, et al., Eukaryotic Cell 4:1298-1307 (2005).

Krappmann, et al., Mol. Microbiol. 61(1):76-88 (2006).

(Continued)

Primary Examiner — Gary Nickol Assistant Examiner — Lakia Tongue

(74) Attorney, Agent, or Firm — Quarles & Brady, LLP

(57) ABSTRACT

The invention provides a general and facile method to obtain secondary metabolites from fungal sources. The invention is based on the discovery that the fungal gene veA and protein encoded thereby regulates the activity of multiple secondary metabolite gene clusters in fungi. Over expression of the gene veA provides increased production of secondary metabolites in engineered cells. In particular, such a method of increasing secondary metabolite production allows the production of improved yields of valuable secondary metabolite products.

2 Claims, 23 Drawing Sheets (13 of 23 Drawing Sheet(s) Filed in Color)

(56)References Cited

OTHER PUBLICATIONS

Kunkel, Proc. Natl. Acad. Sci. USA 82:488-492 (1985). Li, et al., Mol. Microbiol. 62(5):1418-1432 (2006).

Link, et al., Nat. Biotechnol. 17:676-682 (1999).

Lillie, et al., Histopathologic Technic and Practical Histochemistry, 4th ed., McGraw-Hill Book Company, New York, NY (1976), contents/index only.

Maggio-Hall, et al., Mol. Microbiol. 54:1173-1185 (2004).

Maggio-Hall, et al., Mol. Plant-Microbe Interact. 18:783-793 (2005).

Michailides, et al., Plant Pathol. 56:352 (2007).

Miller, et al., Mol. Cell. Biol. 5:1714-1721 (1985).

Mooney, et al., Genes Dev. 4:1473-1482 (1990).

Muyrers, et al., Genet. Eng. 22:77-98 (2000).

Muture, et al., East Afr. Med. J. 82:275-279 (2005).

Nayak, et al., Genetics 172:1557-1566 (2006).

Ni, et al., PLoS One 2(10):e970 (2007)

Perrin, et al., PLoS Pathog. 3(4):e50 (2007).

Puig, et al., Methods 24:218-229 (2001).

Punt, et al., Methods Enzymol 216:447-457 (1992).

Purschwitz, et al., Curr. Biol. 18:255-259 (2008).

Pettit, Yellow Mold and Aflatoxin, p. 35-36, in D. M. Porter, D. H. Smith, and R. Rodriguez-Kabana (ed.) Compendium of Peanut Diseases. The American Phytopathological Society, St. Paul, MN

Robens, et al., The Costs of Mycotoxin Management in the United States, p. 1-13. In H. K. Abbas (ed.) Aflatoxin and Food Safety. CRC Press, Boca Raton, FL (2005).

Rohila, et al., Plant J. 38:172-181 (2004).

Saiki, et al., Nature 324:163-166 (1986).

Sambrook, et al., Molecular Cloning: A Laboratory Manual, Second Edition, Cold Spring Harbor Laboratory Press, NY (1989), Index only.

Seiler, et al., Mol. Biol. Cell 17:4080-4092 (2006).

Shevchenko, et al., Anal. Chem 68, 850-858 (1996).

Shwab, et al., Eukaryotic Cell 6:1656-1664 (2007).

Southern, J. Mol. Biol. 98:503-517 (1975).

Sprote, et al., Arch. Microbiol. 188:69-79 (2007).

Stinnett, et al., Mol. Microbiol. 63(1):242-255 (2007).

Shchepin, et al., Chem. Biol. 10:743-750 (2003).

Shimizu, K., et al., Genetics 157:591-600 (2001).

Smart, et al., Phytopathology 80:1287-1294 (1990).

Szewczyk, et al., Nat. Protoc. 1:3111-3120 (2006).

Thompson, et al., Nucleic Acids Res 22:4673-4680 (1994).

Tsitsigiannis, et al., Mol. Microbiol. 59:882-892 (2006).

Tsitsigiannis, et al., Microbiology 151:1809-1821 (2005).

Tsitsigiannis, et al., J. Biol. Chem. 279:11344-11353 (2004).

Williams, Microbiology 153:3923-3938 (2007).

Wilson, et al., Microbiology 150:2881-2888 (2004).

Woloshuk, et al., Appl. Environ. Microbiol. 60:2408-2414 (1994).

Yu, et al., Fungal Genet. Biol. 41:973-981 (2004).

Yu, et al., Rev. Iberoam. Micol. 22:194-202 (2005).

Yu, et al., Mycotoxin Production and Prevention of Aflatoxin Contamination in Food and Feed. In G. H. Goldman and S. A. Osmani (ed.), The Aspergilli. CRC Press, Boca Raton, FL (2008), pp. 457-472.

^{*} cited by examiner

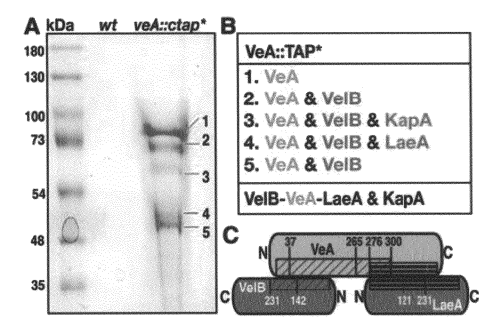


FIG. 1

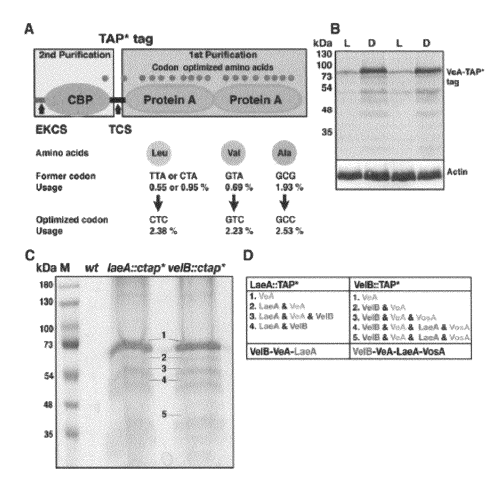


FIG. 2

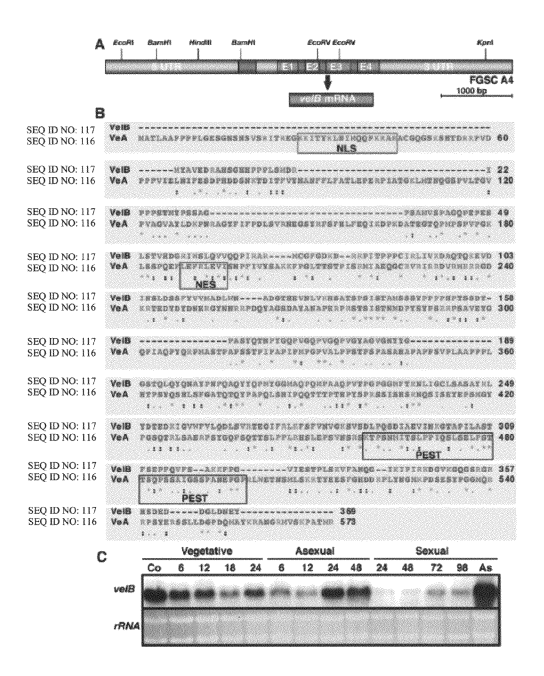


FIG. 3

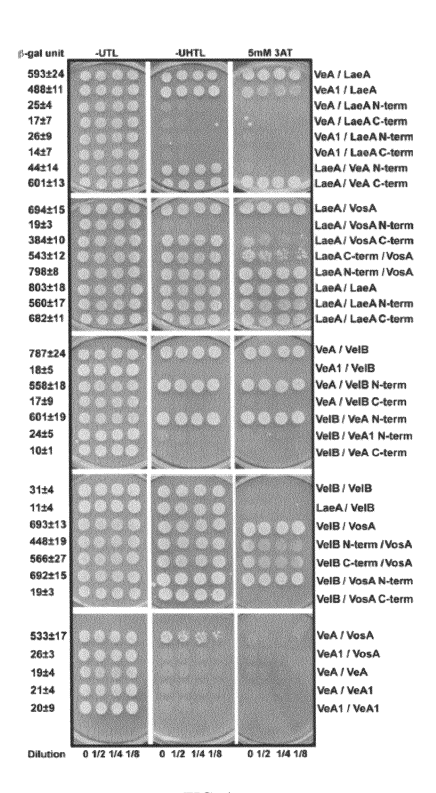


FIG. 4

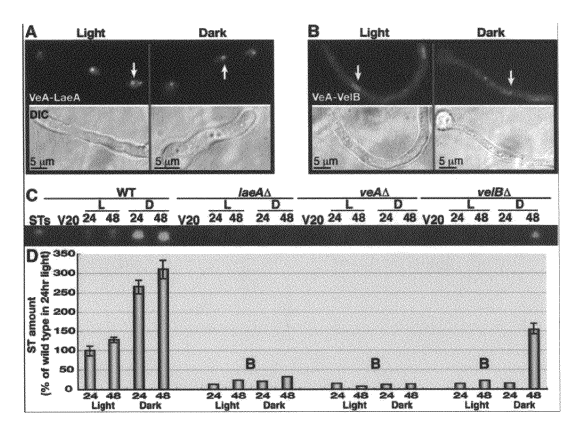


FIG. 5

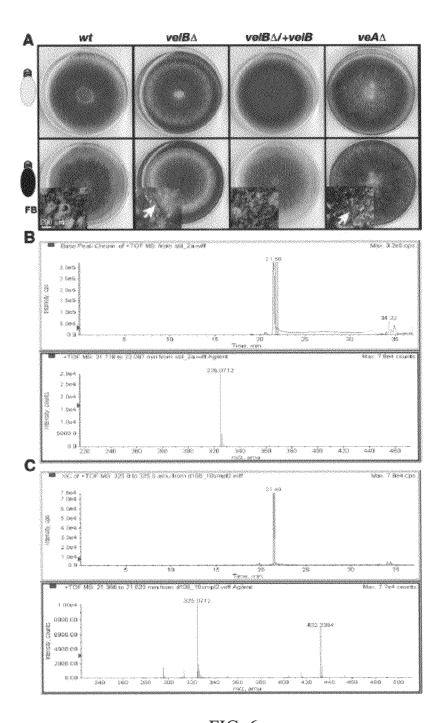


FIG. 6

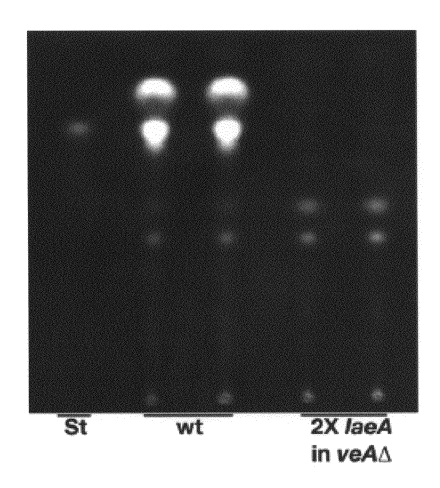


FIG. 7

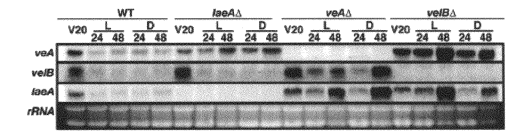


FIG. 8

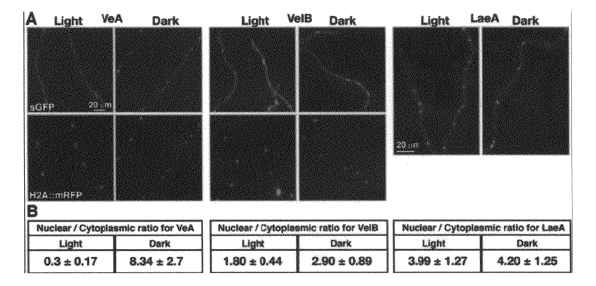


FIG. 9

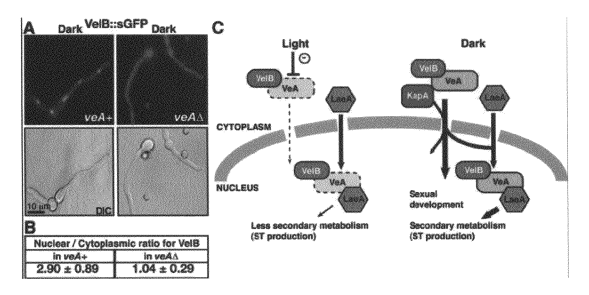


FIG. 10

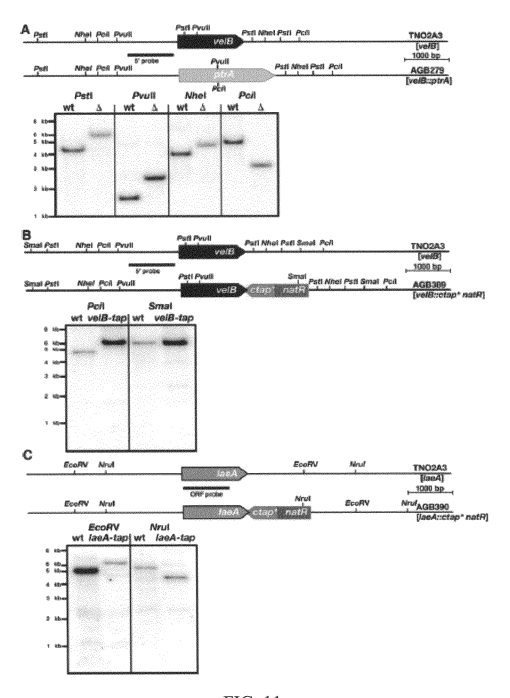


FIG. 11

SEQ ID NO: 116 - VeA protein [Aspergillus nidulans] GENBANK REF: AAD42946.

Oct. 28, 2014

matlaapppp lgesgnsnsv sritregkki tyklnimqqp kraracgqgs kshtdrrpvd 61 pppvielnif esdphddsnk tditfvynan fflfatlepe rpiatgklmt nggspvltgv 121 pvagvayldk pnragyfifp dlsvrnegsy rfsfhlfeqi kdpkdategt qpmpspvpgk 181 lsspqeflef rlevisnpfi vysakkfpgl ttstpisrmi aeqgcrvrir rdvrmrrrgd 241 krtedydydn ergynnrrpd qyagsdayan aperprstsi stnmdpysyp srrpsaveyg 301 qpiaqpyqrp mastpapsst pipapipmpg pvalppstps pasahapapp svplaapppl 361 htpsyqshls fgatqtqypa pqlshipqqt ttpthpyspr ssishsrnqs iseyepsmgy 421 pgsqtrlsae rpsygqpsqt tslpplrhsl epsvnsrskt psnmitslpp iqslselpst 481 tsqpssaigs spanepgprl wetnsmlskr tyeesfghdd rplyngmrpd sesypggmqr 541 rpsyerssll dgpdqmaykr angrmvskpa tmr

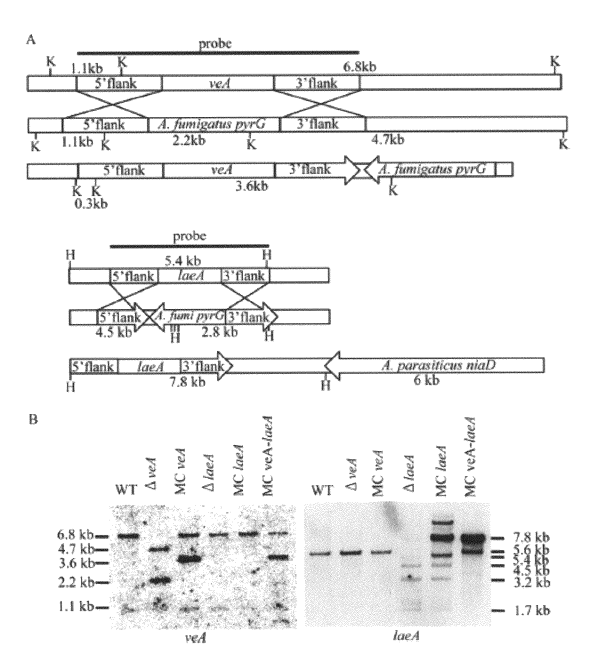


Figure 13

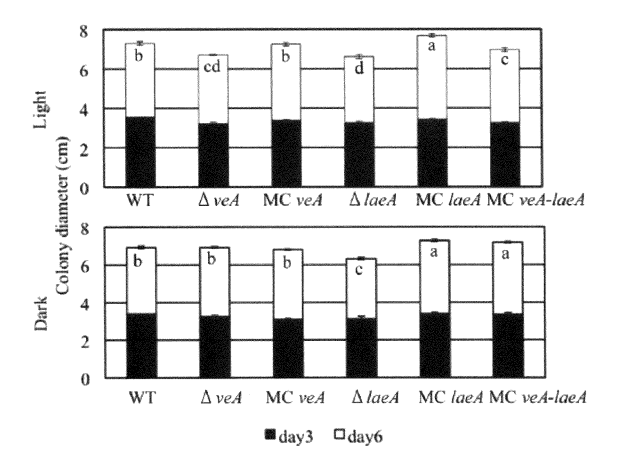


Figure 14

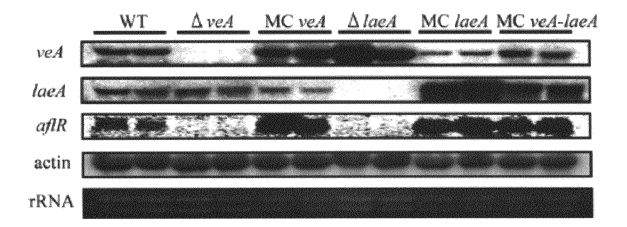


Figure 15

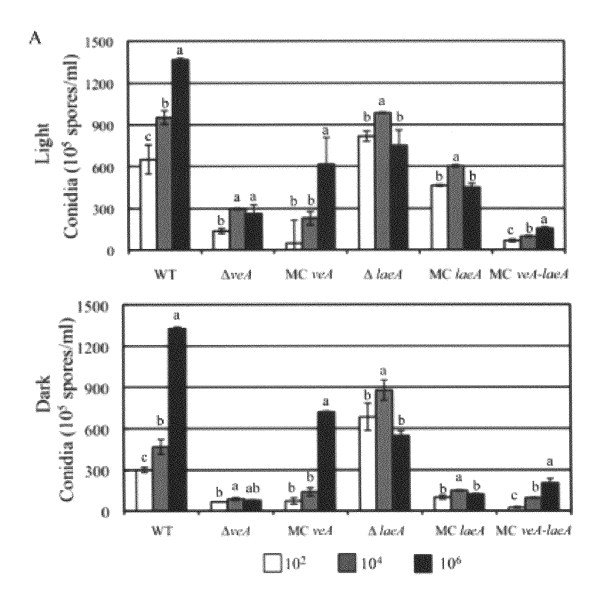


Figure 16

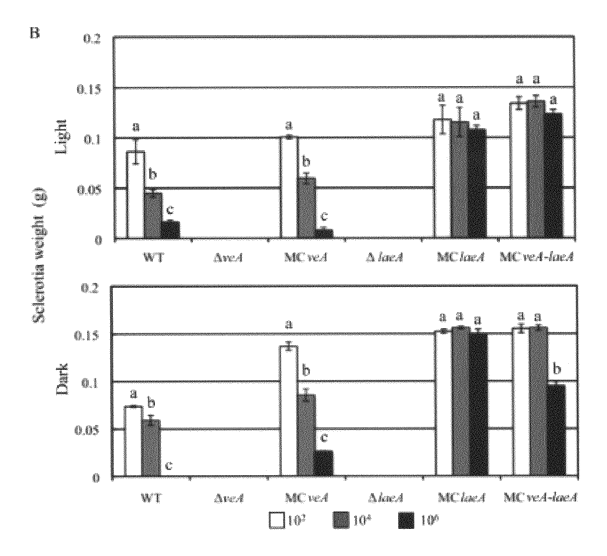


Figure 16

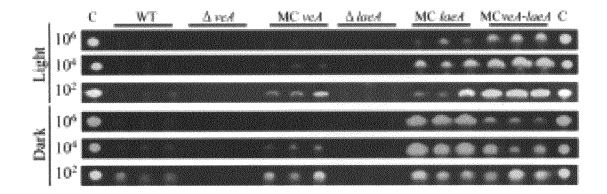


Figure 17

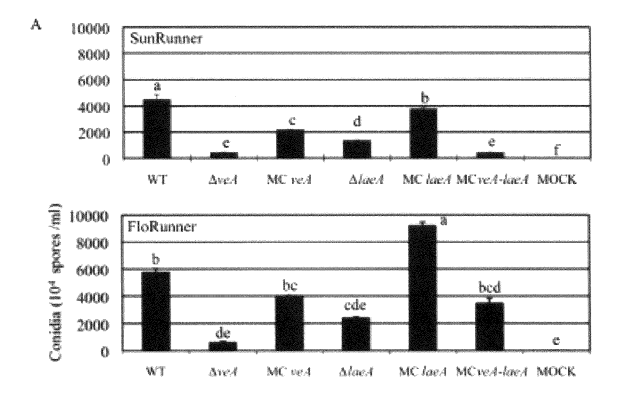


Figure 18

A - Continued

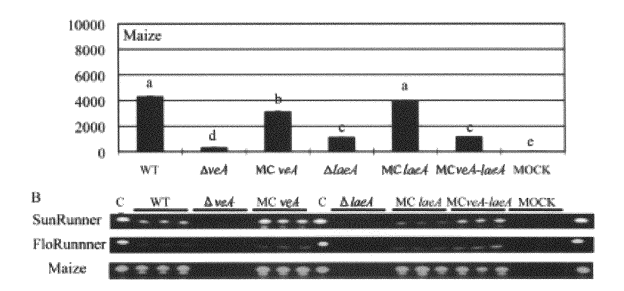


Figure 18

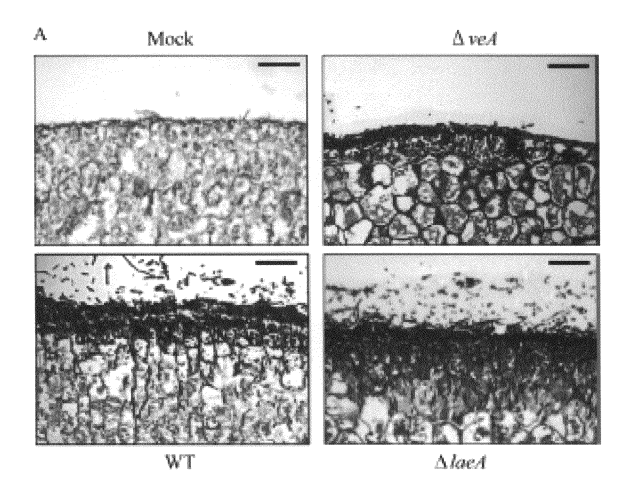


Figure 19

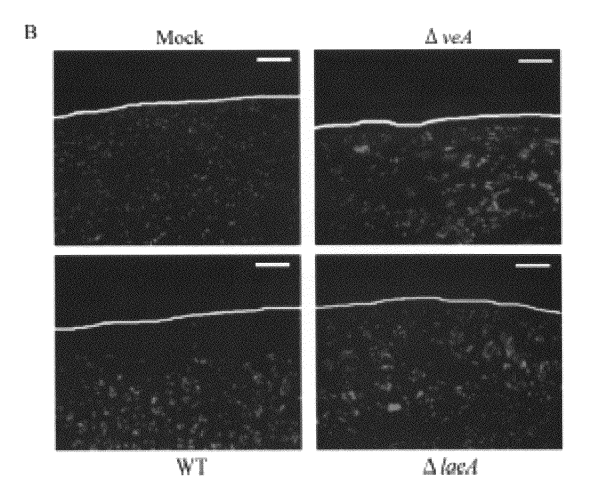


Figure 19

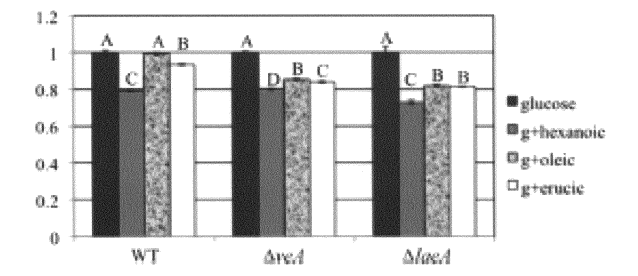


Figure 20

OVER-PRODUCTION OF SECONDARY METABOLITES BY OVER-EXPRESSION OF THE VEA GENE

CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims priority to U.S. Provisional Application No. 61/172,514 filed Apr. 24, 2009, the entirety of which is incorporated by reference herein for all purposes.

STATEMENT REGARDING FEDERALLY SPONSORED RESEARCH OR DEVELOPMENT

This invention was made with United States government 15 support awarded by the following agencies:

NSF 0236393

USDA/CSREES 09-CRHF-0-6055.

The United States government has certain rights in this invention.

FIELD OF THE INVENTION

The invention relates generally to methods of over-producing secondary metabolites. More particularly, the present 25 invention is directed to methods of over-producing secondary metabolites by manipulating fungal regulatory genes involved in the control of secondary metabolite gene clusters.

BACKGROUND OF THE INVENTION

Secondary metabolites are organic compounds that are not directly involved in the normal growth, development or reproduction of organisms. They are often used as defenses against predators, parasites and diseases, for interspecies competition, and to facilitate the reproductive processes (coloring agents, attractive smells, etc).

Secondary metabolites of fungi include both "friends and foes" of human health. For example, penicillin and derivatives produced by *Aspergillus*, *Cephalosporium* and *Penicil*-40 *lium* species are widely used antibiotics, lovastatin is a potent cholesterol-lowering drug produced by *Aspergillus terreus* and aflatoxins, produced by several *Aspergillus* species, are highly toxic carcinogens contaminating many crops.

Secondary metabolic pathways are often tightly correlated 45 with the fungal developmental program and response to external cues including light. Since secondary metabolites are usually restricted to a much more limited group of organisms, they have long been of prime importance in taxonomic research. Secondary metabolites are especially useful for 50 drug or other technological development, or as an inspiration for unnatural products. Biosynthetic genes for fungal secondary metabolites are often clustered and regulated by pathway-specific transcription factors. Secondary metabolism is also regulated at an upper hierarchic level by a global epigenetic 55 control mechanism.

However, methods of producing large amounts of secondary metabolites are difficult and provide unpredictable results. Therefore a need exists for methods of producing large amounts of secondary metabolites that address these 60 problems.

The distribution of natural products is characteristically restricted to certain fungal taxa, particularly the Ascomycetes. Perhaps the greatest number of known secondary metabolites has been ascribed to the Ascomycete genus *Emericella* (asexual stage=*Aspergillus*). Much of the current understanding of fungal secondary metabolite regulation

2

arises from studies of the genetic model *Aspergillus nidulans*. This organism produces many natural products including sterigmatocystin ST (ST; the penultimate precursor to aflatoxin) and penicillin and has been used as a heterologous host to study the biosynthesis of other natural products including lovastatin. Critical advances in understanding fungal secondary metabolism have been largely based on primary studies from *A. nidulans* and/or secondary studies in other fungi where researchers were able to exploit the knowledge gained from *A. nidulans* to their fungus of choice.

A. nidulans, a mold, produces many compounds relevant to biotechnology and human health and is a well-suited model for the analysis of the interplay between secondary metabolism, light and differentiation. A. nidulans grows vegetatively in the soil by hyphal tip extension until competent for development and secondary metabolism. In reproduction, A. nidulans forms airborne asexual spores in light but preferentially undergoes sexual reproduction in the dark. Sexual reproduction in the dark results in an increase in secondary metabolism 20 and in the formation of sexual fruit bodies called cleistothecia, which consist of different cell types. Mutations resulting in defects in fungal development often impair secondary metabolism. There is genetic evidence for a connection between fruitbody formation, secondary metabolism, and light in A. nidulans reproduction, but the molecular mechanism is not known.

Aspergillus flavus, an opportunistic pathogen of oil seeds, occurs as a saprophyte in soils worldwide and colonizes several important agricultural crops, such as maize, peanut, and cottonseed, before and after harvest. The pathogen generates asexual spores, conidia, as the source of inoculum and overwinters as sclerotia which germinate to produce conidia in the subsequent season. A. flavus and other aspergilli, such as Aspergillus parasiticus, can produce the polyketide-derived carcinogenic secondary metabolite aflatoxin. In the United States, annual yield losses in the million-dollar range from aflatoxin contamination on peanut and maize crops are frequently reported. Aflatoxin-contaminated food and feed is also a major problem in developing countries, especially in Asia and Africa. Recently, an outbreak of aflatoxin poisoning from maize was reported to have killed a hundred people in Kenya. Therefore, measures to control Aspergillus infections and aflatoxin production are urgently needed to protect human and animal health. The identification and characterization of molecules necessary for A. flavus conidial, sclerotial, and aflatoxin production are critical to develop rational control strategies.

VeA, a conserved velvet protein encoded by the veA gene, increases expression during sexual development. However, VeA transport into the nucleus is inhibited by light. It acts as a negative regulator of asexual development. VeA is required for cleistothecial production in *A. nidulans* and sclerotial production in both *A. parasiticus* and *A. flavus*. In addition, the VeA gene regulates the expression of sterigmatocystin (a precursor of aflatoxin) and penicillin genes in *A. nidulans* and aflatoxin genes in *A. parasiticus* and *A. flavus*. VeA interacts with LaeA in an as-yet-unclear mechanism, although analysis shows that VeA and LaeA negatively regulate each other at the transcript level in *A. nidulans* (1) and LaeA negatively regulates veA in *A. flavus* (21).

LaeA, another protein located in the cell nucleus, is present in numerous fungi and is a master regulator of secondary metabolism in *Aspergilli* and other fungal genera. LaeA is also necessary for sclerotial formation in *A. flavus* and affects cleistothecial development in *A. nidulans*.

The deletion of LaeA silences numerous secondary metabolite gene clusters, including those responsible for the

syntheses of the antibiotic penicillin as well as for toxins such as ST or gliotoxin. It has been suggested that LaeA might control the accessibility of binding factors to chromatin regions of secondary metabolite clusters because LaeA prevents heterochromatin maintenance of some clusters.

Other factors have been reported which link morphological development with secondary metabolism. Of particular interest are a family of oxylipin-producing oxygenases (encoded by ppo and lox genes) which have been shown to balance ascospore and conidial production in A. nidulans (40, 41) and 10 sclerotial and conidial production in A. flavus, as well as secondary metabolite production in both species. Most recently, a density-dependent switch from sclerotial-toconidial development in A. flavus was found to be affected by oxylipin production. Both oxylipin production and the 15 response to oxylipin signaling are dependent on an intact VeA protein. VeA is also required for ppoA expression, and VeAPpoA interactions affect both sexual and asexual development in A. nidulans. The impact of the loss of these proteins on pathogenesis has been explored to some degree for LaeA and 20 Ppo mutants but not yet reported for VeA.

LaeA is a key determinant in aspergillosis caused by A. fumigatus and seed rot by A. flavus and Ppo loss impacts virulence attributes of A. fumigatus, A. nidulans, and A. fla-

Despite present methodologies, a need exists for improved methods of controlling production of secondary metabolites to obtain improved production of important natural products and/or novel natural products with medicinal value.

BRIEF DESCRIPTION OF THE DRAWINGS

The file of this patent contains at least one drawing executed in color. Copies of this patent with color drawing(s) will be provided by the Patent and Trademark Office upon 35 request and payment of the necessary fee.

FIG. 1. Identification of VeA-associated proteins in A. nidulans. (A) Brilliant blue G-stained 10% SDS polyacrylamidegel electrophoresis of TAP procedure for VeA. kD, kiloaffinity purification belong to corresponding proteins (details in table 4). (C) Domain mapping of the interactions based on Y2H data (FIG. 4). N, N terminus; C, C terminus.

FIG. 2. Modified TAP tag* and VeA expression. (A) Depiction of the TAP tag. The codons for 18 amino acids were 45 changed by site-directed mutagenesis and are designated as differently colored spheres. (B) Immunoblotting with antibody against calmodulin binding peptide: 85 kDa VeA::TAP* tag. In the light (L), expression is relatively low in comparison to the expression in the dark (D); as a control antiactin anti- 50 body was used. (C) Brilliant Blue G-stained 10% SDS-PAGE gel of the TAP for VelB and LaeA. (D) The polypeptides identified from the bands of two affinity purifications belong to corresponding proteins (see Table 4).

FIG. 3. velB gene structure and alignment of VeA (SEQ ID 55 NO: 116) and VelB (SEQ ID NO: 117; Genbank Accession No. CBF89638). (A) Architecture of the velB locus of A. nidulans. Exons are indicated as E1, E2, E3, E4 (confirmed in cDNA) and recognition sites of common restriction endonucleases are shown. (B) Local alignment of the VelB and VeA 60 proteins. Identical residues are indicated by an asterisk (*), conserved amino acid substitutions (similar amino acids) as two dots (:), and semi-conserved amino acid substitutions as one dot (.). The red rectangle indicates a putative nuclear localization signal (NLS) of VeA, the blue rectangle indicates 65 a putative nuclear export signal (NES) and the black rectangle marks a conserved PEST (Pro, Glu (or Asp), Ser, Thr) motif.

Red: small and hydrophobic (including aromatic amino acid), Blue: acidic, Magenta: basic, Green: hydroxyl, amine and basic amino acids. (C) Northern hybridization of VelB during different life stages of A. nidulans. It is highly expressed in 5 asexual conidia and sexual ascospores. VelB expression stays at basal levels during vegetative growth and increases during late asexual (24, 48 h) or sexual (98 h and ascospores) development.

FIG. 4. Interaction domain mapping among VeA, VeA1, VosA, LaeA and VelB by yeast two-hybrid assay. Derivatives of yeast strain L40 expressing the different bait and prey fusion proteins were spotted in serial dilutions for growth on -UHTL (uracil, histidine, tryptophan and leucine), -UHTL with 5 mM 3-AT and -UTL media, and then incubated at 30° C. for 5 days. Their β -galactosidase activities were analyzed using ONPG.

FIG. 5. BiFC studies of velvet complex components and their effect on ST production. (A) Enhanced yellow fluorescent protein fused to the N terminus of veA gene (N-EYFP:: VeA) interacts with C-EYFP::LaeA in vivo, which is indicated as yellowish green specks in the nucleus. Histone 2A red fluorescent protein (H2A::mRFP) fusion visualizes the entire nucleus. Interaction does not take place in the whole nucleus but in certain points (gene clusters) that LaeA prob-25 ably acts on (indicated by arrows). Differential interference contrast (DIC) shows hyphal cells. (B) N-EYFP::VeA fusion protein interacts with C-EYFP::VelB in the cytoplasm and nucleus. (C) ST production in respective mutant backgrounds and WT at different time points. STs, ST standard; V20, 20 30 hours vegetative growth; L, light; D, dark. 24 and 48 hour time points are shown. (D) Quantification of ST production using thin layer chromatography: In the dark, more ST is produced in the WT. Deletion of either laeA or veA results in no ST above background (denoted by B) fluctuations. Loss of velB results in basal ST production in dark.

FIG. 6. Deletion of velB and impairment of sexual fruit body formation. (A) Phenotypic characterization of the velBΔ deletion strain AGB279. Defects are restored in AGB280 (+velB). Fruit body formation (FB) in TNO2A3 and daltons. (B) The polypeptides identified from the bands of 40 AGB280 appeared as normal (red arrows), whereas aerial hyphae (white arrows) and red pigment accumulation accompanied by a lack of fruit bodies were evident for velB Δ and $veA\Delta$ strains. Pictures of cleistothecia and hyphae were taken at 10⁸-fold magnification. (B) ST standard HPLC (retention time (RT): 21.58) and the corresponding mass spectrum. (C) Confirmation of sterigmatocystin (ST) production by LC-MS in the velB Δ mutant. HPLC condition: A=0.1% formic acid in water, B=acetonitrile+0.1% formic acid, gradient=2% B to 100% B in 30 min, re-equil=18 min/flow rate=0.200 ml/min, column=ZORBAX C-18 SB, 2.1×50 mm (100 Å, 1.8 U Agilent), Temperature=40° C. Mass spectrum condition: Agilent ESITOF, source Temp=325° C., electrospray=3500 V, drying gas=91/min, nebulizer gas=30 PSI. Tolerance=less than 3 ppm. Actual was 1.6 ppm at Mass 325.0712.

> FIG. 7. Expression of extra copy of laeA in the $veA\Delta$ background. RDIT9.32 (wild-type) and RJW108.1 (veAΔ:: argB; trpC::laeA) were grown on sexual induction condition and metabolites extracted and run on a thin layer chromatography plate (chloroform:acetone=4:1). An extra copy of laeA does not restore sterigmatocystin in the $veA\Delta$ background. ST=sterigmatocystin standard.

> FIG. 8. Northern blot analyses. Levels of veA, velB, laeA mRNA in WT. (RDIT9.32), laeAΔ (RJW41.A), veAΔ (RJW112.2) and velB Δ (RNI18.2). All strains were grown in liquid Aspergillus rich medium at 37° C., 250 rpm for 20 h (shown as V20 in the figure) and then transferred onto solid MM plus supplements with or without 0.1% casamino acids

for the concomitant induction. The strains grown on MM without casamino acids were incubated at 37° C. under white fluorescent light (shown as L in the figure), while the strains grown on MM with casamino acids were sealed with parafilm, wrapped with foil and incubated at 37° C. in the darkness (shown as D in the figure). Samples for RNA extraction were collected at 24 h and 48 h after induction. Twenty microgram of total RNA were loaded in each lane. EtBr-stained rRNA evaluated equal loading of total RNA.

FIG. 9. Subcellular localization of the subunits of the velvet complex. (A) VeA-, LaeA-, and VelB-sGFP localizations in the presence or absence of light. VeA-sGFP shows light-dependent nuclear enrichment (counterstained with H2A:: mRFP for visualization of the entire nucleus). (B) Nuclear/cytoplasmic GFP signal ratio of 100 hyphal cells each 15 (Openlab software 5.0.1). Growth in the dark results in increased nuclear and decreased cytoplasmic fluorescence for VeA. VelB and LaeA distribution is hardly affected by illumination

FIG. 10. VeA supports nuclear localization of VelB and 20 formation of the velvet complex. (A) Fluorescence patterns in strains expressing velB::sgfp in the dark in veA+ and veAD backgrounds. (B) Nuclear/cytoplasmic GFP signal ratio of 100 hyphal cells each. Nuclear signal intensity is higher in the veA+ strain background than in veAD. (C) Model: (Light) 25 VeA is mostly retained in the cytoplasm, VelB supports asexual spore formation, and LaeA shows low activity. (Dark) An increased amount of VeA is imported into the nucleus by KapA and, in addition, supports the nuclear transport of VelB. Dotted lines indicate the decreased amount of VeA that is 30 present in the cell in the light and the impairment of VeA nuclear transport in the light. VelB/VeA control development and LaeA activity by formation of the velvet complex that affects secondary metabolite clusters expression.

FIG. 11. Deletion of the velB locus and TAP tagging fusion 35 genes at the velB and laeA loci. (A) Comparative depiction of the wild-type velB locus (TNO2A3) and the velB::ptrA locus (AGB279). The black bar indicates the probe for Southern hybridization. (B) The result of TAP tagging of velB locus is depicted. Autoradiography of Southern hybridization confirms the gene replacement (C) The TAP tagged laeA locus is shown. Autoradiography of Southern hybridization confirms the homologous gene replacements for the velB and laeA loci. For the deletion of velB, the ptrA (pyrithiamin resistance gene) marker was used and for the TAP tagging of velB and laeA, the (nourseothricin resistance gene) nat marker was utilized.

FIG. 12. Sequence Listing for VeA (A. nidulans) (SEQ ID NO: 116).

FIG. 13. Deletion, MCveA, and MClaeA mutants of A. 50 flavus. (A) Diagram of the strategy of replacement of A. flavus NRRL 3357.5 veA with A. fumigatus AF293 wild-type pyrG gene shows the restriction enzyme digestion sites of KpnI for Southern analysis with veA probe. To confirm gene replacement or MC transformants using Southern analysis, at least 55 two restriction enzymes for each probe were utilized, KpnI (K) and SapI (data not shown) for veA and HindIII (H) and BamHI (data not shown) for laeA. A. fumi, A. fumigatus. (B) Southern analysis. The KpnI digest shows 6.8-kb and 1.1-kb veA fragments in the wild type and 4.7-kb, 2.2-kb, and 1.1-kb fragments in the ΔveA strain. The MCveA strain shows both wild-type 6.8-kb and 1.1-kb fragments, as well as 3.6-kb and 0.3-kb (not shown) fragments. The laeA probe presented a 5.6-kb fragment in the wild type; 4.5-kb, 3.2-kb, and 1.7-kb fragments in the ΔlaeA strain; and several extra bands in the MClaeA strain. The laeA mutants have been described before, in reference 21. WT, wild type.

6

FIG. 14. Colony diameters of veA and laeA mutants of *A. flavus*. A 5-μl amount of a suspension of 10⁶ spores/ml of each strain was point inoculated on 30 ml of 1.6% GMM. Cultures were grown at 29° C. under continuous dark or light conditions, and growth diameters measured at 3 and 6 days after inoculation. Letters indicate differences between strains that were statistically significant (P<0.05) according to the Tukey-Kramer multiple comparison test. Error bars show the standard deviations of the results of four replications. Strains were grown in both light and dark conditions. WT, wild type.

FIG. 15. Gene expression levels of veA and laeA in A. flavus mutants. Each strain was grown in liquid GMM culture with shaking (250 rpm at 29° C.) under dark conditions. Total RNA was extracted from two replicates at 48 hrs after inoculation. Northern blots were probed with internal or ORF fragments of each gene (Table 2). rRNA and actin were the loading and expression controls. WT, wild type.

FIG. 16. Effects of veA and laeA allele numbers on density-dependent conidial and sclerotial production in *A. flavus*. Each strain was grown from 10², 10⁴, and 10⁶ spores/plate as described in Materials and Methods. (A) Conidial counts. (B) Sclerotial weight. Letters indicate statistically significant differences (P<0.05) for each strain at different population levels according to the Tukey-Kramer multiple comparison test. Error bars show standard deviations of the results of four replications. WT, wild type.

FIG. 17. Aflatoxin production of veA and laeA mutants. Aflatoxin from each strain was assessed at three different spore inoculation levels. The experiment was replicated three times, as shown. C, aflatoxin B1 control; WT, wild type.

FIG. 18. Conidium production and aflatoxin production on peanut and maize seeds. Seeds of two peanut cultivars and one maize line were inoculated with 10⁵ spores/ml of the wild type and the veA and laeA mutants and incubated for either 3 days (peanut cultivar SunRunnner and maize kernels) or 5 days (peanut cultivar FloRunner) after inoculation at 29° C. under dark conditions. (A) For conidium counting, 1-ml amounts of homogenized suspensions of five peanut cotyledons or maize kernels of inoculated seeds were diluted to 1× and conidia counted. Letters indicate statistically significant differences (P<0.05) of different strains, according to Tukey-Kramer multiple comparison test. Error bars show the standard deviations of the results of three replications. (B) Aflatoxin was extracted from inoculated peanut cotyledons and maize kernels and resuspended in 500 µl of chloroform, and 10 μl of each extract was spotted on a TLC plate and separated with chloroform/acetone (95:5, vol/vol). C, aflatoxin B1 control; WT, wild type; MOCK, control inoculated with water.

FIG. 19. Histological examination reveals differences in seed ingress and lipid utilization of ΔveA and $\Delta laeA$ strains compared to these functions in the wild type. (A) Tissues were stained with Gomori methenamine-silver for detection of fungal hyphae. (B) Tissues were stained with Nile red for lipid body detection in seeds. To observe tissues, a bright-field microscope was used for Gomori stain and a tetramethyl rhodamine 5-isothiocyanate filter in a fluorescent microscope was used for Nile red. Seeds infected with the wild-type fungus show diminishment of lipid bodies near the surface (white line) of the seed. Scale bars=100 μm . WT, wild type; Mock, control inoculated with water.

FIG. 20. Loss of veA and laeA sensitizes the fungus to oleic acid. Inhibition of colony diameters of Δ veA and Δ laeA mutants but not the wild type is observed when GMM is supplemented with 6 mM oleic acid at 3 (data not shown) and 6 days after inoculation. Letters indicate statistically significant differences (P<0.05) at 6 days after inoculation with different strains, according to Tukey-Kramer multiple com-

parison test. Error bars show the standard deviations of the results of four replications. WT, wild type; g, glucose; hexanoic, hexanoic acid; oleic, oleic acid; erucic, erucic acid.

SUMMARY OF THE INVENTION

The present invention provides a novel method of increasing the amount of a secondary metabolite produced by a cell or organism. The method comprises the steps of obtaining a cell or an organism capable of biosynthesizing a secondary metabolite; transforming the cell or organism with a nucleic acid which encodes a veA polypeptide, a polypeptide having substantial sequence identity thereto, or a fragment thereof having secondary metabolite gene cluster regulating activity; and culturing the transformed cell or organism so that an 15 increase in production of the secondary metabolite occurs in the transformed cell or organism as compared to a non-transformed cell or organism. In one embodiment, the cell or organism is an Aspergillus species such as A. nidulans or A. flavus.

In another embodiment, the present invention provides a novel method of decreasing the production of a secondary metabolite by a transformed cell or organism. The method comprises the steps of: obtaining a transformed cell or organism capable of biosynthesizing a secondary metabolite, the transformed cell or organism having a defective veA gene wherein the defective veA gene is no longer biologically active and expression of secondary metabolite gene clusters is reduced; and culturing the transformed cell or organism so that a decrease in production of the secondary metabolite 30 occurs in the transformed cell or organism as compared to a non-transformed cell or organism. In one embodiment, the cell or organism is an Aspergillus species such as A. nidulans or A. flavus.

In another embodiment, the present invention provides a 35 novel method of producing an isolated secondary metabolite. The method comprises the steps of: obtaining a cell or an organism capable of biosynthesizing a secondary metabolite; transforming the cell or organism with a nucleic acid which sequence identity thereto, or a fragment thereof having secondary metabolite gene cluster regulating activity; culturing the transformed cell or organism under conditions conducive to increasing production of the secondary metabolite in the transformed cell or organism as compared to a non-trans- 45 formed cell or organism; and recovering the secondary metabolite from the transformed cell or organism or from the culture in which the transformed cell or organism was grown in an isolated form. In one embodiment, the cell or organism is an Aspergillus species such as A. nidulans or A. flavus.

In another embodiment, the present invention provides a novel method for identifying a novel secondary metabolite biosynthesis gene cluster in a fungus. The method comprises the steps of: obtaining a transformed fungus having a disrupted veA gene; isolating a sample of nucleic acids from the 55 transformed fungus, wherein the sample of nucleic acids is representative of the expressed genes of the transformed fungus; hybridizing the sample of nucleic acids isolated above or nucleic acid equivalents of same with an array comprising a plurality of nucleic acids representative of the expressed 60 genes of a non-transformed fungus under conditions conducive to forming one or more hybridization complexes; detecting the hybridization complexes; comparing the detected levels of the hybridization complexes with the level of hybridization complexes detected in a sample of nucleic acids 65 isolated from a veA-expressing fungus, wherein the nucleic acids isolated from a veA-expressing fungus are representa8

tive of the expressed genes of the veA-expressing fungus, and wherein an altered level of hybridization complexes detected above compared with a level of hybridization complexes of the sample of nucleic acids from the veA-expressing fungus correlates with and identifies at least one gene under regulatory control of a veA gene product; and examining genomic nucleotide sequence surrounding the at least one gene identified above to determine if the at least one gene is clustered with other secondary metabolite biosynthesis genes, thereby identifying a novel secondary metabolite biosynthesis gene cluster. In one embodiment, the cell or organism is an Aspergillus species such as A. nidulans or A. flavus.

DETAILED DESCRIPTION OF THE INVENTION

I. In General

Before the present materials and methods are described, it is understood that this invention is not limited to the particular 20 methodology, protocols, materials, and reagents described, as these may vary. It is also to be understood that the terminology used herein is for the purpose of describing particular embodiments only, and is not intended to limit the scope of the present invention which will be limited only by any laterfiled nonprovisional applications.

It must be noted that as used herein and in the appended claims, the singular forms "a", "an", and "the" include plural reference unless the context clearly dictates otherwise. As well, the terms "a" (or "an"), "one or more" and "at least one" can be used interchangeably herein. It is also to be noted that the terms "comprising", "including", and "having" can be used interchangeably.

The practice of the present invention will employ, unless otherwise indicated, conventional techniques of molecular biology, microbiology, recombinant DNA, and immunology, which are within the skill of the art. Such techniques are explained fully in the literature. See, for example, Molecular Cloning A Laboratory Manual, 2nd Ed., ed. by Sambrook, Fritsch and Maniatis (Cold Spring Harbor Laboratory Press: encodes a veA polypeptide, a polypeptide having substantial 40 1989); DNA Cloning, Volumes I and II (D. N. Glover ed., 1985); Oligonucleotide Synthesis (M. J. Gait ed., 1984); Mullis et al. U.S. Pat. No. 4,683,195; Nucleic Acid Hybridization (B. D. Hames & S. J. Higgins eds. 1984); Transcription And Translation (B. D. Hames & S. J. Higgins eds. 1984); Culture Of Animal Cells (R. I. Freshney, Alan R. Liss, Inc., 1987); Immobilized Cells And Enzymes (IRL Press, 1986); B. Perbal, A Practical Guide To Molecular Cloning (1984); the treatise, Methods In Enzymology (Academic Press, Inc., N.Y.); Gene Transfer Vectors For Mammalian Cells (J. H. Miller and M. P. Calos eds., 1987, Cold Spring Harbor Laboratory); Methods In Enzymology, Vols. 154 and 155 (Wu et al. eds.), Immunochemical Methods In Cell And Molecular Biology (Mayer and Walker, eds., Academic Press, London, 1987); Handbook Of Experimental Immunology, Volumes I-IV (D. M. Weir and C. C. Blackwell, eds., 1986); Cell Culture and Somatic Cell Genetics of Plants, Vol. 1 (I. K. Vasil, ed. 1984); R. V. Stanier, J. L. Ingraham, M. L. Wheelis, and P. R. Painter, The Microbial World, (1986) 5th Ed. Prentice-Hall.

> Unless defined otherwise, all technical and scientific terms used herein have the same meanings as commonly understood by one of ordinary skill in the art to which this invention belongs. Although any methods and materials similar or equivalent to those described herein can be used in the practice or testing of the present invention, the preferred methods and materials are now described. All publications and patents specifically mentioned herein are incorporated by reference

for all purposes including describing and disclosing the chemicals, instruments, statistical analysis and methodologies which are reported in the publications which might be used in connection with the invention. All references cited in this specification are to be taken as indicative of the level of skill in the art. Nothing herein is to be construed as an admission that the invention is not entitled to antedate such disclosure by virtue of prior invention.

II. Definitions

"VeA", as used herein, refers to the amino acid sequences of the VeA protein obtained from *Aspergillus nidulans*. In addition, VeA shall also refer to the amino acid sequences of VeA obtained from any species (i.e., orthologs), particularly 15 fungi (e.g. other strains and/or species of *Aspergillus*, and other genera), from any source whether natural, synthetic, semi-synthetic, or recombinant. The term encompasses proteins encoded by nucleotide sequences representing allelic variants as well as those containing single nucleotide poly- 20 morphisms (SNPs).

"veA", as used herein, refers to the nucleotide sequences of the veA gene obtained from *Aspergillus nidulans*. In addition, veA shall also refer to the nucleotide sequences of the veA gene obtained from any species, particularly fungi (e.g. other 25 strains and/or species of *Aspergillus*, and other genera), from any source whether natural, synthetic, semi-synthetic, or recombinant. The term encompasses allelic variants and single nucleotide polymorphisms (SNPs).

An "allele" or "allelic sequence", as used herein, is an 30 alternative form of the gene encoding VeA. Alleles may result from at least one mutation in the nucleic acid sequence and may result in altered mRNAs or polypeptides whose structure or function may or may not be altered. Any given natural or recombinant gene may have none, one, or many allelic forms. 35 Common mutational changes which give rise to alleles are generally ascribed to natural deletions, additions, or substitutions of nucleotides. Each of these types of changes may occur alone, or in combination with the others, one or more times in a given sequence.

"Altered" nucleic acid sequences encoding VeA, as used herein, include those with deletions, insertions, or substitutions of different nucleotides resulting in a polynucleotide that encodes the same or a functionally equivalent protein to VeA. Included within this definition are polymorphisms 45 which may or may not be readily detectable using a particular oligonucleotide probe of the polynucleotide encoding VeA, and improper or unexpected hybridization to alleles, with a locus other than the normal chromosomal locus for the polynucleotide sequence encoding VeA. The encoded protein may 50 also be "altered" and contain deletions, insertions, or substitutions of amino acid residues which produce a silent change and result in a functionally equivalent VeA. Deliberate amino acid substitutions may be made on the basis of similarity in polarity, charge, solubility, hydrophobicity, hydrophilicity, 55 and/or the amphipathic nature of the residues as long as the biological or immunological activity of VeA is retained. For example, negatively charged amino acids may include aspartic acid and glutamic acid; positively charged amino acids may include lysine and arginine; and amino acids with 60 uncharged polar head groups having similar hydrophilicity values may include leucine, isoleucine, and valine, glycine and alanine, asparagine and glutamine, serine and threonine, and phenylalanine and tyrosine.

"Amino acid sequence", as used herein, refers to an oligopeptide, peptide, polypeptide, or protein sequence, and fragment thereof. Where "amino acid sequence" is recited 10

herein to refer to a particular amino acid sequence "amino acid sequence" and like terms are not meant to limit the amino acid sequence to the complete amino acid sequence referenced but shall be understood to include fragments of the complete amino acid sequence. The term shall further encompass synthetic molecules as well as those occurring naturally. The term "portion" or "fragment", as used herein, with regard to an amino acid sequence (as in "a fragment of SEQ ID NO:1"), specifically refers to segments of that amino acid 10 sequence which are not naturally occurring as fragments and would not be found in the natural state. The segments may range in size from five amino acid residues to the entire amino acid sequence minus one amino acid. Thus, a polypeptide "comprising at least a portion of the amino acid sequence of SEQ ID NO:1" or "including an amino acid sequence as set forth in SEQ ID NO:1 or fragments thereof" encompasses the full-length VeA amino acid sequences and segments thereof.

"Amplification", as used herein, refers to the production of additional copies of a nucleic acid sequence and is generally carried out using polymerase chain reaction (PCR) technologies well known in the art (Dieffenbach, C. W. and G. S. Dveksler (1995) PCR Primer, a Laboratory Manual, Cold Spring Harbor Press, Plainview, N.Y.).

"Antisense", as used herein, refers to any composition containing nucleotide sequences which are complementary to a specific DNA or RNA sequence. The term "antisense strand" is used in reference to a nucleic acid strand that is complementary to the "sense" strand. Antisense molecules include peptide nucleic acids and may be produced by any method including synthesis or transcription. Once introduced into a cell, the complementary nucleotides combine with natural sequences produced by the cell to form duplexes and block either transcription or translation. The designation "negative" is sometimes used in reference to the antisense strand, and "positive" is sometimes used in reference to the sense strand.

"Biologically active", as used herein, refers to a protein, polypeptide, amino acid sequence, or nucleotide sequence encoding a product having structural, regulatory, or biochemical functions of a naturally occurring molecule. Preferably, a biologically active fragment of VeA will have the secondary metabolite gene cluster regulatory capabilities of a naturally occurring VeA molecule disclosed herein.

"Complementary" or "complementarity", as used herein, refer to the natural binding of polynucleotides under permissive salt and temperature conditions by base-pairing. For example, the sequence "A-G-T" binds to the complementary sequence "T-C-A". Complementary between two single-stranded molecules may be "partial", in which only some of the nucleic acids bind, or it may be complete when total complementarity exists between the single stranded molecules. The degree of complementarity between nucleic acid strands has significant effects on the efficiency and strength of hybridization between nucleic acid strands. This is of particular importance in amplification reactions, which depend upon binding between nucleic acids strands and in the design and use of PNA molecules.

A "composition comprising a given polynucleotide sequence", as used herein, refers broadly to any composition containing the given polynucleotide sequence. Compositions comprising polynucleotide sequences encoding VeA or fragments thereof, may be employed as hybridization probes. The probes may be stored in freeze-dried form and may be associated with a stabilizing agent such as a carbohydrate. In hybridizations, the probe may be deployed in an aqueous solution containing salts (e.g., NaCl), detergents (e.g., SDS) and other components (e.g., Denhardt's solution, dry milk, salmon sperm DNA, etc.).

The phrase "correlates with expression of a polynucleotide", as used herein, indicates that the detection of the presence of ribonucleic acid that is similar to SEQ ID NO:1 by northern analysis or equivalent analysis is indicative of the presence of mRNA encoding VeA in a sample and thereby correlates with expression of the transcript from the polynucleotide encoding the protein.

"Deletion", as used herein, refers to a change in the amino acid or nucleotide sequence and results in the absence of one or more amino acid residues or nucleotides.

"Derivative", as used herein, refers to the chemical modification of a nucleic acid encoding or complementary to veA or the encoded VeA protein itself. Such modifications include, for example, replacement of hydrogen by an alkyl, acyl, or amino group. A nucleic acid derivative encodes a polypeptide which retains the biological or immunological function of the natural molecule. A derivative polypeptide is one which is modified by glycosylation, or any similar process which retains the biological function of the polypeptide from which it was derived.

"Homology", as used herein, refers to sequence similarity between two peptides or between two nucleic acid molecules. Homology may be determined by comparing a position in each sequence which may be aligned for purposes of comparison. When a position in the compared sequence is occu- 25 pied by the same base or amino acid, then the molecules are homologous at that position. A degree of homology between sequences is a function of the number of matching or homologous positions shared by the sequences. A partially complementary sequence that at least partially inhibits an identical 30 sequence from hybridizing to a target nucleic acid is referred to using the functional term "substantially homologous." The inhibition of hybridization of the completely complementary sequence to the target sequence may be examined using a hybridization assay (i.e., Southern or northern blot, solution 35 hybridization and the like) under conditions of low stringency. A substantially homologous sequence or hybridization probe will compete for and inhibit the binding of a completely homologous sequence to the target sequence under conditions of low stringency. This is not to say that conditions of 40 low stringency are such that non-specific binding is permitted; low stringency conditions require that the binding of two sequences to one another be a specific (i.e., selective) interaction. The absence of non-specific binding may be tested by the use of a second target sequence which lacks even a partial 45 degree of complementary (e.g., less than about 30% identity). In the absence of non-specific binding, the probe will not hybridize to the second non-complementary target sequence.

"Identity", as used herein, means the degree of sequence relatedness between polypeptide or polynucleotide 50 sequences, as the case may be, as determined by the match between strings of such sequences. "Substantial sequence identity" as used herein means at least 80% identical, more preferably 95%, 96%, 97%, 98% or 99% identical. "Identity" and "homology" can be readily calculated by known meth- 55 ods, including but not limited to those described in (Computational Molecular Biology, Lesk, A. M., ed., Oxford University Press, New York, 1988; Biocomputing: Informatics and Genome Projects, Smith, D. W., ed., Academic Press, New York, 1993; Computer Analysis of Sequence Data, Part I, 60 Griffin, A. M., and Griffin, H. G., eds., Humana Press, New Jersey, 1994; Sequence Analysis in Molecular Biology, von Heinje, G., Academic Press, 1987; and Sequence Analysis Primer, Gribskov, M. and Devereux, J., eds., M Stockton Press, New York, 1991; and Carillo, H., and Lipman, D., 65 SIAM J. Applied Math., 48: 1073 (1988). Preferred methods to determine identity are designed to give the largest match

12

between the sequences tested. Methods to determine identity and homology are codified in publicly available computer programs. Preferred computer program methods to determine identity and homology between two sequences include, but are not limited to, the GCG program package (Devereux, J., et al., Nucleic Acids Research 12(1): 387 (1984)), BLASTP, BLASTN, and FASTA (Atschul, S. F. et al., J. Molec. Biol. 215: 403-410 (1990). The BLAST X program is publicly available from NCBI and other sources (BLAST Manual, Altschul, S., et al., J. Mol. Biol. 215: 403-410 (1990). The well known Smith Waterman algorithm may also be used to determine identity.

"Hybridization", as used herein, refers to any process by which a strand of nucleic acid binds with a complementary strand through base pairing.

"Hybridization complex", as used herein, refers to a complex formed between two nucleic acid sequences by virtue of the formation of hydrogen bonds between complementary G and C bases and between complementary A and T bases; these hydrogen bonds may be further stabilized by base stacking interactions. The two complementary nucleic acid sequences hydrogen bond in an antiparallel configuration. A hybridization complex may be formed in solution (e.g., Co t or Ro t analysis) or between one nucleic acid sequence present in solution and another nucleic acid sequence immobilized on a solid support (e.g., paper, membranes, filters, chips, pins or glass slides, or any other appropriate substrate to which cells or their nucleic acids have been fixed).

An "insertion" or "addition", as used herein, refers to a change in an amino acid or nucleotide sequence resulting in the addition of one or more amino acid residues or nucleotides, respectively, as compared to the naturally occurring molecule.

"Isolated" or "purified" or "isolated and purified" means altered "by the hand of man" from its natural state, i.e., if it occurs in nature, it has been changed or removed from its original environment, or both. For example, a polynucleotide or a polypeptide naturally present in a living organism is not "isolated," but the same polynucleotide or polypeptide separated from the coexisting materials of its natural state is "isolated", as the term is employed herein. Moreover, a polynucleotide or polypeptide that is introduced into an organism by transformation, genetic manipulation or by any other recombinant method is "isolated" even if it is still present in said organism, which organism may be living or non-living. As so defined, "isolated nucleic acid" or "isolated polynucleotide" includes nucleic acids integrated into a host cell chromosome at a heterologous site, recombinant fusions of a native fragment to a heterologous sequence, recombinant vectors present as episomes or as integrated into a host cell chromosome. As used herein, the term "substantially purified", refers to nucleic or amino acid sequences that are removed from their natural environment, isolated or separated, and are at least 60% free, preferably 75% free, and most preferably 90% free from other components with which they are naturally associated. As used herein, an isolated nucleic acid "encodes" a reference polypeptide when at least a portion of the nucleic acid, or its complement, can be directly translated to provide the amino acid sequence of the reference polypeptide, or when the isolated nucleic acid can be used, alone or as part of an expression vector, to express the reference polypeptide in vitro, in a prokaryotic host cell, or in a eukaryotic host cell.

"Exon", as used herein, refers to a nucleic acid sequence found in genomic DNA that is bioinformatically predicted

and/or experimentally confirmed to contribute contiguous sequence to a mature mRNA transcript.

"Open reading frame" and the equivalent acronym "ORF", as used herein, refer to that portion of a transcript-derived nucleic acid that can be translated in its entirety into a sequence of contiguous amino acids. As so defined, an ORF has length, measured in nucleotides, exactly divisible by 3. As so defined, an ORF need not encode the entirety of a natural

"Microarray" refers to an ordered arrangement of hybridizable array elements. The array elements are arranged so that there are preferably at least one or more different array elements, more preferably at least 100 array elements, and most preferably at least 1,000 array elements, on a 1 cm² substrate 15 surface. The maximum number of array elements is unlimited, but is at least 100,000 array elements. Furthermore, the hybridization signal from each of the array elements is individually distinguishable. In a preferred embodiment, the fungal-derived polynucleotide sequences.

"Modulate", as used herein, refers to a change in the activity of VeA. For example, modulation may cause an increase or a decrease in protein activity, binding characteristics, or any other biological, functional or immunological properties of 25

"Nucleic acid sequence" or "nucleotide sequence" or "polynucleotide sequence", as used herein, refers to an oligonucleotide, nucleotide, or polynucleotide, and fragments thereof, and to DNA or RNA of genomic or synthetic origin 30 which may be single- or double-stranded, and represent the sense or antisense strand. Where "nucleic acid sequence" or "nucleotide sequence" or polynucleotide sequence" is recited herein to refer to a particular nucleotide sequence (e.g., the nucleotide sequence set forth in SEQ ID NO:2), "nucleotide 35 sequence", and like terms, are not meant to limit the nucleotide sequence to the complete nucleotide sequence referenced but shall be understood to include fragments of the complete nucleotide sequence.

In this context, the term "fragment" may be used to spe-40 cifically refer to those nucleic acid sequences which are not naturally occurring as fragments and would not be found in the natural state. Generally, such fragments are equal to or greater than 15 nucleotides in length, and most preferably includes fragments that are at least 60 nucleotides in length. 45 Such fragments find utility as, for example, probes useful in the detection of nucleotide sequences encoding VeA.

"Sample", as used herein, is used in its broadest sense. A biological sample suspected of containing nucleic acid encoding VeA, or fragments thereof, or VeA itself may com- 50 prise a bodily fluid, extract from a cell, chromosome, organelle, or membrane isolated from a cell, a cell, genomic DNA, RNA, or cDNA (in solution or bound to a solid support, a tissue, a tissue print, and the like).

A "substitution", as used herein, refers to the replacement 55 of one or more amino acids or nucleotides by different amino acids or nucleotides, respectively. The term "conservative substitution" is used in reference to proteins or peptides to reflect amino acid substitutions that do not substantially alter the activity (specificity or binding affinity) of the molecule. 60 Typically conservative amino acid substitutions involve substitution one amino acid for another amino acid with similar chemical properties (e.g. charge or hydrophobicity). The following six groups each contain amino acids that are typical conservative substitutions for one another: 1) Alanine (A), 65 Serine (S), Threonine (T); 2) Aspartic acid (D), Glutamic acid (E); 3) Asparagine (N), Glutamine (Q); 4) Arginine (R),

14

Lysine (K); 5) Isoleucine (I), Leucine (L), Methionine (M), Valine (V); and 6) Phenylalanine (F), Tyrosine (Y), Tryp-

"Transformation", as defined herein, describes a process by which exogenous DNA enters and changes a recipient cell. It may occur under natural or artificial conditions using various methods well known in the art. Transformation may rely on any known method for the insertion of foreign nucleic acid sequences into a prokaryotic or eukaryotic host cell. The method is selected based on the type of host cell being transformed and may include, but is not limited to, viral infection, electroporation, heat shock, lipofection, and particle bombardment. Such "transformed" cells include stably transformed cells in which the inserted DNA is capable of replication either as an autonomously replicating plasmid or as part of the host chromosome. They also include cells which transiently express the inserted DNA or RNA for limited periods of time.

A "variant" of VeA, as used herein, refers to an amino acid array elements comprise polynucleotide representative of 20 sequence that is altered by one or more amino acids. The variant may have "conservative" changes, wherein a substituted amino acid has similar structural or chemical properties, e.g., replacement of leucine with isoleucine. More rarely, a variant may have "nonconservative" changes, e.g., replacement of a glycine with a tryptophan. Analogous minor variations may also include amino acid deletions or insertions, or both. Guidance in determining which amino acid residues may be substituted, inserted, or deleted without abolishing biological or immunological activity may be found using computer programs well known in the art, for example, DNASTAR software.

> The terms "polypeptide", "peptide" and "protein" are used interchangeably herein to refer to a polymer of amino acid residues. The terms apply to amino acid polymers in which one or more amino acid residue is an artificial chemical analogue of a corresponding naturally occurring amino acid, as well as to naturally occurring amino acid polymers. The term also includes variations on the traditional peptide linkage joining the amino acids making up the polypeptide. Where the terms are recited herein to refer to a polypeptide, peptide or protein of a naturally occurring protein molecule, the terms are not meant to limit the polypeptide, peptide or protein to the complete, native amino acid sequence associated with the recited protein molecule but shall be understood to include fragments of the complete polypeptide. The term "portion" or "fragment", as used herein, with regard to a protein or polypeptide (as in "a fragment of the VeA polypeptide") refers to segments of that polypeptide which are not naturally occurring as fragments in nature. The segments may range in size from five amino acid residues to the entire amino acid sequence minus one amino acid. Thus, a polypeptide "as set forth in SEQ ID NO:1 or a fragment thereof" encompasses the full-length amino acid sequence set forth in SEQ ID NO:1 as well as segments thereof. Fragments of VeA preferably are biologically active as defined herein.

> The terms "nucleic acid" or "oligonucleotide" or "polynucleotide" or grammatical equivalents herein refer to at least two nucleotides covalently linked together. A nucleic acid of the present invention is preferably single-stranded or double stranded and will generally contain phosphodiester bonds, although in some cases, as outlined below, nucleic acid analogs are included that may have alternate backbones, comprising, for example, phosphoramide (Beaucage et al. (1993) Tetrahedron 49:1925) and references therein; Letsinger (1970) J. Org. Chem. 35:3800; Sprinzl et al. (1977) Eur. J. Biochem. 81: 579; Letsinger et al. (1986) Nucl. Acids Res. 14: 3487; Sawai et al. (1984) Chem. Lett. 805, Letsinger et al.

(1988) J. Am. Chem. Soc. 110: 4470; and Pauwels et al. (1986) Chemica Scripta 26: 1419), phosphorothioate (Mag et al. (1991) Nucleic Acids Res. 19:1437; and U.S. Pat. No. 5,644,048), phosphorodithioate (Briu et al. (1989) J. Am. Chem. Soc. 111: 2321, O-methylphophoroamidite linkages (see Eckstein, Oligonucleotides and Analogues: A Practical Approach, Oxford University Press), and peptide nucleic acid backbones and linkages (see Egholm (1992) J. Am. Chem. Soc. 114:1895; Meier et al. (1992) Chem. Int. Ed. Engl. 31: 1008; Nielsen (1993) Nature, 365: 566; Carlsson et al. (1996) Nature 380: 207). Other analog nucleic acids include those with positive backbones (Denpcy et al. (1995) Proc. Natl. Acad. Sci. USA 92: 6097; non-ionic backbones (U.S. Pat. Nos. 5,386,023, 5,637,684, 5,602,240, 5,216,141 and 4,469, 15 863; Angew. (1991) Chem. Intl. Ed. English 30: 423; Letsinger et al. (1988) J. Am. Chem. Soc. 110:4470; Letsinger et al. (1994) Nucleoside & Nucleotide 13:1597; Chapters 2 and 3, ASC Symposium Series 580, "Carbohydrate Modifications in Antisense Research", Ed. Y. S. Sanghui 20 DNA or RNA sequences between two DNA or RNA moland P. Dan Cook; Mesmaeker et al. (1994), Bioorganic & Medicinal Chem. Lett. 4: 395; Jeffs et al. (1994) J. Biomolecular NMR 34:17; Tetrahedron Lett. 37:743 (1996) and non-ribose backbones, including those described in U.S. Pat. Nos. 5,235,033 and 5,034,506, and Chapters 6 and 7, ASC 25 Symposium Series 580, Carbohydrate Modifications in Antisense Research, Ed. Y. S. Sanghui and P. Dan Cook. Nucleic acids containing one or more carbocyclic sugars are also included within the definition of nucleic acids (see Jenkins et al. (1995), Chem. Soc. Rev. pp 169-176). Several nucleic acid 30 analogs are described in Rawls, C & E News Jun. 2, 1997 page 35. These modifications of the ribose-phosphate backbone may be done to facilitate the addition of additional moieties such as labels, or to increase the stability and halflife of such molecules in physiological environments. As used 35 herein, oligonucleotide is substantially equivalent to the terms "amplimers", "primers", "oligomers", and "probes", as commonly defined in the art.

The term "heterologous" as it relates to nucleic acid sequences such as coding sequences and control sequences, 40 denotes sequences that are not normally associated with a region of a recombinant construct, and/or are not normally associated with a particular cell. Thus, a "heterologous" region of a nucleic acid construct is an identifiable segment of nucleic acid within or attached to another nucleic acid mol- 45 ecule that is not found in association with the other molecule in nature. For example, a heterologous region of a construct could include a coding sequence flanked by sequences not found in association with the coding sequence in nature. Another example of a heterologous coding sequence is a 50 construct where the coding sequence itself is not found in nature (e.g., synthetic sequences having codons different from the native gene). Similarly, a host cell transformed with a construct which is not normally present in the host cell would be considered heterologous for purposes of this inven-

A "coding sequence" or a sequence which "encodes" a particular polypeptide (e.g. a methyltransferase, etc.), is a nucleic acid sequence which is ultimately transcribed and/or translated into that polypeptide in vitro and/or in vivo when 60 placed under the control of appropriate regulatory sequences. In certain embodiments, the boundaries of the coding sequence are determined by a start codon at the 5' (amino) terminus and a translation stop codon at the 3' (carboxy) terminus. A coding sequence can include, but is not limited to, 65 cDNA from prokaryotic or eukaryotic mRNA, genomic DNA sequences from prokaryotic or eukaryotic DNA, and even

16

synthetic DNA sequences. In preferred embodiments, a transcription termination sequence will usually be located 3' to the coding sequence.

The term "ortholog" refers to genes or proteins which are homologs via speciation, e.g., closely related and assumed to have common descent based on structural and functional considerations. Orthologous proteins function as recognizably the same activity in different species.

Expression "control sequences" or "regulatory elements" refers collectively to promoter sequences, ribosome binding sites, polyadenylation signals, transcription termination sequences, upstream regulatory domains, enhancers, and the like, which collectively provide for the transcription and translation of a coding sequence in a host cell. Not all of these control sequences need always be present in a recombinant vector so long as the desired gene is capable of being transcribed and translated.

"Recombination" refers to the reassortment of sections of ecules. "Homologous recombination" occurs between two DNA molecules which hybridize by virtue of homologous or complementary nucleotide sequences present in each DNA molecule.

The terms "stringent conditions" or "hybridization under stringent conditions" refers to conditions under which a probe will hybridize preferentially to its target subsequence, and to a lesser extent to, or not at all to, other sequences. "Stringent hybridization" and "stringent hybridization wash conditions" in the context of nucleic acid hybridization experiments such as Southern and northern hybridizations are sequence dependent, and are different under different environmental parameters. An extensive guide to the hybridization of nucleic acids is found in Tijssen (1993) Laboratory Techniques in Biochemistry and Molecular Biology-Hybridization with Nucleic Acid Probes. Generally, highly stringent hybridization and wash conditions are selected to be about 5° C. lower than the thermal melting point (Tm) for the specific sequence at a defined ionic strength and pH. The Tm is the temperature (under defined ionic strength and pH) at which 50% of the target sequence hybridizes to a perfectly matched probe. Very stringent conditions are selected to be equal to the Tm for a particular probe.

An example of stringent hybridization conditions for hybridization of complementary nucleic acids which have more than 100 complementary residues on a filter in a Southern or northern blot is 50% formamide with 1 mg of heparin at 42° C., with the hybridization being carried out overnight. An example of highly stringent wash conditions is 0.15 M NaCl at 72° C. for about 15 minutes. An example of stringent wash conditions is a 0.2×SSC wash at 65° C. for 15 minutes (see, Sambrook et al. (1989) Molecular Cloning—A Laboratory Manual (2nd ed.) Vol. 1-3, Cold Spring Harbor Laboratory, Cold Spring Harbor Press, NY, for a description of SSC buffer). Often, a high stringency wash is preceded by a low stringency wash to remove background probe signal. An example medium stringency wash for a duplex of, e.g., more than 100 nucleotides, is 1×.SSC at 45° C. for 15 minutes. An example low stringency wash for a duplex of, e.g., more than 100 nucleotides, is 4-6xSSC at 40° C. for 15 minutes. In general, a signal to noise ratio of 2x (or higher) than that observed for an unrelated probe in the particular hybridization assay indicates detection of a specific hybridization. Nucleic acids which do not hybridize to each other under stringent conditions are still substantially identical if the polypeptides which they encode are substantially identical.

This occurs, e.g., when a copy of a nucleic acid is created using the maximum codon degeneracy permitted by the genetic code.

"Expression vectors" are defined herein as nucleic acid sequences that are direct the transcription of cloned copies of genes/cDNAs and/or the translation of their mRNAs in an appropriate host. Such vectors can be used to express genes or cDNAs in a variety of hosts such as bacteria, bluegreen algae, plant cells, insect cells and animal cells. Expression vectors include, but are not limited to, cloning vectors, modified cloning vectors, specifically designed plasmids or viruses. Specifically designed vectors allow the shuttling of DNA between hosts, such as bacteria-yeast or bacteria-animal cells. An appropriately constructed expression vector preferably contains: an origin of replication for autonomous replication in a host cell, a selectable marker, optionally one or more restriction enzyme sites, optionally one or more constitutive or inducible promoters. In preferred embodiments, an expression vector is a replicable DNA construct in which a 20 DNA sequence encoding VeA or a fragment thereof is operably linked to suitable control sequences capable of effecting the expression of the products in a suitable host. Control sequences include a transcriptional promoter, an optional operator sequence to control transcription and sequences 25 which control the termination of transcription and translation, and so forth

A "polymorphism" is a variation in the DNA sequence of some members of a species. A polymorphism is thus said to be "allelic," in that, due to the existence of the polymorphism, some members of a species may have the unmutated sequence (i.e. the original "allele") whereas other members may have a mutated sequence (i.e. the variant or mutant "allele"). In the simplest case, only one mutated sequence may exist, and the polymorphism is said to be diallelic. In the case of diallelic diploid organisms, three genotypes are possible. They can be homozygous for one allele, homozygous for the other allele or heterozygous. In the case of diallelic haploid organisms, they can have one allele or the other, thus only two genotypes are possible. The occurrence of alternative mutations can give rise to trialleleic, etc. polymorphisms. An allele may be referred to by the nucleotide(s) that comprise the mutation.

"Single nucleotide polymorphism" or "SNPs are defined by their characteristic attributes. A central attribute of such a polymorphism is that it contains a polymorphic site, "X," most preferably occupied by a single nucleotide, which is the site of the polymorphism's variation. Methods of identifying SNPs are well known to those of skill in the art (see, e.g., U.S. Pat. No. 5,952,174).

Abbreviations used herein include "aa", amino acid; 50 "MMG", minimal media glucose; "MMT", minimal media threonine; "OE", over expression; "LB", Luria-Bertani; "nt", nucleotide; "ORF", open reading frame; "PCR", polymerase chain reaction; "PEG", polyethyleneglycol; "R", resistant; "WT", wild-type; and "TS", temperature sensitive.

III. The Invention

The present invention provides a novel method for producing secondary metabolites by inducing the over-expression of 60 the fungal gene veA (SEQ ID NO: 116—see FIG. 12). Such methods include steps of: (a) obtaining a cell or an organism capable of biosynthesizing a secondary metabolite; (b) transforming the cell or organism with an nucleic acid encoding a VeA polypeptide capable of regulating biosynthesis of the 65 secondary metabolite; and (c) culturing the transformed cell or organism so that an increase in production of the secondary

18

metabolite occurs in the transformed cell or organism as compared to a non-transformed cell or organism.

In one embodiment of the present invention, methods of increasing the amount of a secondary metabolite as described and claimed herein are practiced in an *Aspergillus* species such as *A. nidulans. A. flavus* or *A. terreus*. Secondary metabolites increased by the methods include but are not limited to lovastatin or penicillin.

The invention also provides methods of decreasing the production of a secondary metabolite in a transformed cell or organism. Such methods include the steps of: (a) obtaining a transformed cell or organism capable of biosynthesizing a secondary metabolite, the transformed cell or organism having a defective veA gene wherein the defective veA gene is no longer biologically active and expression of secondary metabolite gene clusters is reduced; and (b) culturing the transformed cell or organism so that a decrease in production of the secondary metabolite occurs in the transformed cell or organism as compared to a non-transformed cell or organism. Such a gene replacement exercise could be carried out by one of skill in the art using techniques presently known in the field. Such a method would be useful in reducing or eliminating production of toxic secondary metabolites in certain organisms. For example, a non-functional variant of veA would be useful in reducing or eliminating aflatoxin production in an A. parasiticus or A. flavus strain transformed thereby. In addition, veA may be targeted by a therapeutic such that veA's ability to regulate secondary metabolite gene cluster activity is inhibited. This approach would provide a therapeutic compound able to reduce the virulence of cells or organisms, thereby providing a treatment for medical maladies involving fungal infections. Methods of identifying inhibitors of target molecules are well known in the art.

In yet another embodiment, the present invention encompasses methods of producing an isolated secondary metabolite. These methods include steps of: (a) obtaining a cell or an organism capable of biosynthesizing a secondary metabolite; (b) transforming the cell or organism with a nucleic acid encoding a VeA polypeptide capable of regulating biosynthesis of the secondary metabolite; (c) culturing the transformed cell or organism under conditions conducive to increasing production of the secondary metabolite in the transformed cell or organism; and (d) recovering the secondary metabolite from the transformed cell or organism in an isolated form.

The invention also provides methods for identifying yet undiscovered secondary metabolite biosynthesis gene clusters in a variety of fungi based on the nucleic acids and transformed cells disclosed herein. Such methods are preferably carried out in a microarray format. For example, using standard microarray technology now commonly employed in the field, one of skill in the art may construct a microarray containing, for example, nucleic acids representative of the expressed genes of wild-type A. nidulans (see, for example, 55 D. Bowtell and J. Sambrook, DNA Microarrays: A Molecular Cloning Manual (2000) Cold Spring Harbor Laboratory Press and P. Baldi and G. W. Hatfield, DNA Microarrays and Gene Expression: From Experiments to Data Analysis and Modeling (2002) Cambridge University Press describing standard microarray techniques data analyses applicable in the present invention). The entire genome for A. nidulans has been sequenced and the sequence is available in annotated form for public use (see the Whitehead Institute/MIT Center for Genome Research website). Construction of the specific nucleic acids affixed to the array substrate may be based on, for example, an expressed sequence tag database provided by the University of Oklahoma.

Using the microarray and standard hybridization techniques known in the field, the expression levels of genes in wild-type A. nidulans, A. flavus or other wild-type fungus versus a veA deletion mutant may then be compared to identify genes whose expression is reduced or absent in the veA deletion mutant compared to the wild-type line. The artisan may subsequently examine the genomic sequence available of, for example, A. nidulans or A. flavus to identify putative secondary metabolite biosynthesis cluster genes in the immediate vicinity of the relevant gene whose expression is initially identified as affected by the absence of veA expression. As secondary metabolite biosynthesis genes are well known to occur in clustered fashion, as described in a plurality of references cited herein, new putative secondary metabolite gene clusters may be identified by this approach.

Further, genes within a putative gene cluster may subsequently be disrupted and the mutant line's production of secondary metabolite products may then be compared with wild-type production in plus/minus fashion to identify the specific natural product produced by the newly-identified 20 gene cluster. The natural product may then be isolated and characterized using standard techniques described and referenced herein.

The above-described screening strategies may be carried out not only between wild-type and veA deletion mutants but 25 also, and more preferably, between veA overexpression mutants and veA deletion mutants to obtain the greatest contrast in veA-influenced secondary metabolite biosynthesis gene expression. As well, the screening methodology described herein is not limited to any one particular fungus 30 but may be applied to any fungus having a veA ortholog (e.g., Aspergillus other than A. nidulans and A. flavus). For example, the genome for Fusarium graminearum is now available and screens utilizing veA overexpression or disruption strains to identify new F. graminearum secondary 35 metabolite gene clusters may certainly be carried out based on the novel materials and teachings provided herein (also see Whitehead Institute/MIT Center for Genomic Research website).

sequences or fragments thereof which encode VeA may be used in recombinant DNA molecules to direct expression of VeA, fragments or functional equivalents thereof, in appropriate host cells. Due to the inherent degeneracy of the genetic code, other DNA sequences which encode substantially the 45 same or a functionally equivalent amino acid sequence may be produced, and these sequences may be used to clone and express VeA.

As will be understood by those of skill in the art, it may be advantageous to produce VeA-encoding nucleotide 50 sequences possessing non-naturally occurring codons. For example, codons preferred by a particular prokaryotic or eukaryotic host can be selected to increase the rate of protein expression or to produce an RNA transcript having desirable properties, such as a half-life which is longer than that of a 55 transcript generated from the naturally occurring sequence.

The nucleotide sequences of the present invention can be engineered using methods generally known in the art in order to alter VeA-encoding sequences for a variety of reasons, including but not limited to, alterations which modify the 60 moters such as the hybrid lacZ promoter of the BLUEcloning, processing, and/or expression of the gene product. DNA shuffling by random fragmentation and PCR reassembly of gene fragments and synthetic oligonucleotides may be used to engineer the nucleotide sequences. For example, sitedirected mutagenesis may be used to insert new restriction 65 sites, alter glycosylation patterns, change codon preference, produce splice variants, introduce mutations, and so forth.

20

In another embodiment of the invention, natural, modified, or recombinant nucleic acid sequences encoding VeA may be ligated to a heterologous sequence to encode a fusion protein. For example, to screen peptide libraries for inhibitors of VeA activity, it may be useful to encode a chimeric VeA protein that can be recognized by a commercially available antibody. A fusion protein may also be engineered to contain a cleavage site located between the VeA encoding sequence and the heterologous protein sequence, so that VeA may be cleaved and purified away from the heterologous moiety.

In another embodiment, sequences encoding VeA may be synthesized, in whole or in part, using chemical methods well known in the art. Alternatively, the protein itself may be produced using chemical methods to synthesize the amino acid sequence of VeA, or a fragment thereof. For example, peptide synthesis can be performed using various solid-phase techniques and automated synthesis may be achieved, for example, using the ABI 431A Peptide Synthesizer (Perkin Elmer)

The newly synthesized peptide may be substantially purified by preparative high performance liquid chromatography. The composition of the synthetic peptides may be confirmed by amino acid analysis or sequencing. Additionally, the amino acid sequence of VeA, or any part thereof, may be altered during direct synthesis and/or combined using chemical methods with sequences from other proteins, or any part thereof, to produce a variant polypeptide.

In order to express a biologically active VeA, the nucleotide sequences encoding VeA or functional equivalents may be inserted into an appropriate expression vector, i.e., a vector which contains the necessary elements for the transcription and translation of the inserted coding sequence. Methods which are well known to those skilled in the art may be used to construct expression vectors containing sequences encoding VeA and appropriate transcriptional and translational control elements. These methods include in vitro recombinant DNA techniques, synthetic techniques, and in vivo genetic recombination.

A variety of expression vector/host systems may be uti-In another embodiment of the invention, nucleotide 40 lized to contain and express sequences encoding VeA. These include, but are not limited to, microorganisms such as bacteria transformed with recombinant bacteriophage, plasmid, or cosmid DNA expression vectors; yeast transformed with yeast expression vectors; insect cell systems infected with virus expression vectors (e.g., baculovirus); plant cell systems transformed with virus expression vectors (e.g., cauliflower mosaic virus, CaMV; tobacco mosaic virus, TMV) or with bacterial expression vectors (e.g., Ti or pBR322 plasmids); or animal cell systems. The invention is not limited by the host cell employed.

The "control elements" or "regulatory sequences" are those non-translated regions of the vector—enhancers, promoters, 5' and 3' untranslated regions—which interact with host cellular proteins to carry out transcription and translation. Such elements may vary in their strength and specificity. Depending on the vector system and host utilized, any number of suitable transcription and translation elements, including constitutive and inducible promoters, may be used. For example, when cloning in bacterial systems, inducible pro-SCRIPT phagemid (Stratagene, La Jolla, Calif.) or PSPORT1 plasmid (Gibco BRL) and the like may be used. The baculovirus polyhedrin promoter may be used in insect cells. Promoters or enhancers derived from the genomes of plant cells (e.g., heat shock, RUBISCO; and storage protein genes) or from plant viruses (e.g., viral promoters or leader sequences) may be cloned into the vector. In mammalian cell

systems, promoters from mammalian genes or from mammalian viruses are preferable. If it is necessary to generate a cell line that contains multiple copies of the sequence encoding VeA, vectors based on SV40 or EBV may be used with an appropriate selectable marker.

The following examples describing materials and methodology are offered for illustrative purposes only, and are not intended to limit the scope of the present invention.

III. Examples

Example 1

In the present invention, tandem affinity purification (TAP) was used to identify VeA-interacting proteins (FIG. 1A and 15 FIG. 2A). Final eluates of dark- and light-grown *A. nidulans* carrying the functional veA gene tagged at its C terminus by TAP tag (veA::ctap*) were analyzed by mass spectrometry. The velvet-like protein B (VelB) (FIG. 3A, 3B), the regulator LaeA, and the α importin KapA were identified as proteins 20 that interact with VeA in the dark (FIG. 1B and Table 4). (Importin is a type of protein that moves other protein molecules into the nucleus by binding to a specific recognition sequence, called the nuclear localization signal (NLS)).

In the light, tagged VeA protein is hardly expressed (FIG. 25 2B) and only copurifies with VelB. Reciprocal affinity purifications of tagged VelB and LaeA in the dark confirmed the interaction partners, except for the α importin KapA (FIGS. 2C and D). Only tagged VelB can additionally recruit the regulator of sporogenesis VosA in the dark, which seems to be 30 an alternative binding partner for this protein.

Yeast two-hybrid (Y2H) analysis confirmed the VeA-VelB and VeA-LaeA interactions, where VelB and LaeA do not interact in this assay, suggesting that VeA acts as a bridge between VelB and LaeA (FIG. 1C).

The Y2H VosA-LaeA interaction supports a role of LaeA in development (FIG. 4). The C-terminal part of VeA interacts with LaeA, whereas the N-terminal part of VeA, which includes the nuclear localization signal (NLS), is required for interaction with VelB (FIG. 1C and FIG. 4).

VelB, which is conserved in the fungal kingdom, shares 18% amino acid identity with VeA but has no typical NLS (FIG. 3B). Transcript analysis reveals that VelB expression increases like that of VeA at late developmental stages (FIG. 3C). The VeA-LaeA and VeA-VelB interactions were visualized by bimolecular fluorescence complementation (BiFC) in living cells. Distinct fluorescent specks show that the VeA-LaeA interaction occurs in the nucleus, whereas VeA and VelB interact in the cytoplasm and within the nucleus (FIGS. 5A and B).

The physical interaction of VeA with VelB, as well as with LaeA, leads to the novel understanding of the present invention that VeA and VelB are functionally interdependent. Similar to veAD, the velBD mutant (FIG. 6A) no longer displays a light-dependent developmental pattern and is unable to 55 form sexual fruit bodies, even in the dark. Asexual sporulation in velBD is impaired but not as strongly as in a veA deletion strain.

Reintroduction of the velB locus fully rescued all of the defects (FIG. 6A). The veAD/velBD double mutant exhibited a near-identical phenotype to that of the veAD single mutant. Neither VelB overexpression in a veAD background nor VeA overexpression in a velBD background rescued the defects of the individual mutants; likewise, LaeA overexpression could not rescue secondary metabolite defects of veAD (FIG. 7).

Unlike overproduction of VeA, overexpression of VelB in a veA+ background does not cause excessive production of

22

cleistothecia, but it induces a twofold increase in asexual sporulation in comparison to the wild type (WT). This suggests that VeA controls the number of sexual structures, whereas VelB has additional developmental functions. Secondary metabolism is impaired in veAD, resulting in a similar brownish pigment as is produced by the velBD strain.

Changes in gene expression and in LaeA activity were monitored in the veAD and velBD strains (FIGS. **5**C and D, FIGS. **6**B and C, and FIG. **8**). ST production is abolished in veAD and laeAD strains. In contrast, reduced and delayed but significant ST production in VelBD suggests residual activity of a VeA/LaeA complex in the dark. VeA is enriched in the nucleus in the dark, whereas VelB was found in both the nucleus and the cytoplasm and is hardly affected by illumination (FIGS. **9**A and B).

Because LaeA is constitutively nuclear (FIGS. 9A and B) and the interaction of VeA and LaeA occurs in the nucleus (FIG. 5A), VelB has to enter the nucleus, despite the lack of an obvious NLS to fully control LaeA. Localization of the VelB-sGFP fusion protein (where GFP is green fluorescent protein) in a veAD background is shifted toward the cytoplasm, whereas the presence of VeA increases the nuclear localization of VelB (FIGS. 10A and B).

This suggests that VeA can assist VelB to allow an enhanced transport into the nucleus. The data provided herein suggest that the mechanism underlying the coordinated regulation of sexual development and secondary metabolism in *A. nidulans* is the interaction between the key developmental regulatory complex VelB/VeA and LaeA.

Accordingly, in the dark the VelB/VeA/LaeA velvet complex interaction controls and presumably supports the epigenetic activity of LaeA, which subsequently controls the expression of secondary metabolite gene clusters. In the light, this interaction is diminished because less VeA protein is present, and the entrance of the bridging factor VeA to the nucleus is decreased.

Because the absence of LaeA has a minor impact on development, VeA and VelB have presumably additional functions in fungal differentiation. This is also supported by the identification of VosA, a recently identified regulator of fungal sporogenesis, as an additional binding partner of VelB (FIGS. 2C and D, and Table 4).

Light triggers asexual development, corresponding to the release of high numbers of asexual spores (conidia) into the environment. These phenotypes correlate with the light-dependent cytoplasmic localization of VeA, the constitutive nuclear function of LaeA, and the partial nuclear localization of VelB, respectively. Under light conditions, when low amounts of VeA and VelB are present in the nucleus, the secondary metabolism regulator LaeA seems to be primarily active in those hyphae that are not exposed to light.

Accordingly, the deletion of laeA results in a loss of mycelial pigmentation at the bottom of the colony. The newly described fungal protein VelB, in conjunction with VeA, connects light-dependent development to LaeA-controlled secondary metabolism in *A. nidulans*. The inventors herein present evidence that the formation of this complex is the molecular basis that synchronizes developmental and metabolic changes to the disappearance of light.

This trimeric complex is designated the "velvet complex". The VelB/VeA is part of the epigenetic control of chromatin remodeling by modulating LaeA methyltransferase activity (FIG. 10C), in which VeA is functionally active in the dark, forms a complex with increased amounts of VelB, and enhances the transport of VelB to the nucleus.

Because VeA and VelB are both partially nuclear, even in the light, we presume a certain threshold is probably necessary to initiate sexual development and control LaeA. Fungal morphogenesis and secondary metabolism have traditionally been viewed as separate fields. The VelB/VeA/LaeA velvet

complex elucidates the molecular mechanisms underlying the intimate relation between fungal development and secondary metabolism.

Strains, media, and growth conditions. Fungal strains used in this study are listed in Table 1.

TABLE 1

Strain Genotype		Fungal Strains.
FGSC4 Glasgow wild-type FGSC36 biAl, veA1 DVAR1 pabaA1, yA2; argBA:ttrpC; trpC801; veAA::argB pabaA1 pveA::weA, ptrA; pabaA1, yA2; argBA::ttrpC; trpC801, veAA::argB preA::weA.:ctap* tag, ptrA; pabaA1, yA2; argBA::ttrpC; trpC801, veAA::argB preA::weA::safp, ptrA; pabaA1, yA2; argBA::ttrpC; trpC801, veAA::argB preA::weA::safp, ptrA; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB preA::weA::safp, ptrA; pgpdA::mafr; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB prind::weB::miAT, pgpdA::mafr; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB prind::weB::miAT, pgpdA::mafr; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB proAd, pyrG89, veA pini::weB::miAT, A.f. pyrG; pgpdA::mafr; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB proAd, pyrG89, veA pini::weB::miAT, A.f. pyrG; pyroA4, pyrG89 prind::weB::miAT, A.f. pyrG; pyroA4, pyrG89 prind::weB::miAT, A.f. pyrG; pyroA4, pyrG89 proBi::weB::miAT, A.f. pyrG; pyroA4, argB2; nkuAA::argB prind::weB::miAT, A.f. pyrG; veBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB prind::weB::safp::miiAT, A.f. pyrG; veBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB prind::weA::argB, pgpdA::mafr; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB prind::weA::argB, pgpdA::mafr; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB prind::weA::miiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB prind::weA::argB, pgpdA::mafr; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB prind::deA::sgfp::miiAT, pgpdA::mafr; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB prind::deA::sgfp::miiAT, ppgdA::mafr; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB prind::deA::sgfp::miiAT, ppdA::mafr; velBA::ptrA; pyrG89, p	Strain	Genotype
FGSC26 biA1, yeA1 pdSA11 pabaA1, yA2; argBA::ttpC; ttpC801; veAA::argB pabaA1 pabaA1 AGB154 pabaA1 pabaA1 pveA::veA, ptrA; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB pveA::veA::argB pveA::veA::srgfp, ptrA; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB AGB274 pveA::veA::srgfp, ptrA; papdA::marfp::h2A, pgpdA::matR; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB pmi1a::veB::srgfp::mi1AT, pgpdA::matR; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB pmi1a::veB::mi1AT, pptGA::matR; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB pmi1a::veB::srgfp::mi1AT, A.f. pyrG; ppgdA::marfp::h2A, pgpdA::matR; pyroA4, pyrG89, veA pmi1a::veB::srgfp::mi1AT, A.f. pyrG; pyroA4, pyrG89 pmi1a::veB::srgfp::mi1AT, A.f. pyrG; pyroA4, pyrG89 pprG89, pyroA4 veBB::ptrA: pyrG89, pyroA4, argB2; nkuAa::argB prGB280 pveB::velB, pgpdA::matR; velBa::ptrA; pyrG89, pyroA4, argB2; nkuAa::argB prGB281 pni1A::velB::srgfp::mi1AT, A.f. pyrG; velBa::ptrA; pyrG89, pyroA4, argB2; nkuAa::argB prGB282 pveA::veA::srgfp::mi1AT, A.f. pyrG; velBa::ptrA; pyrG89, pyroA4, argB2; nkuAa::argB pri1A::velB::miiAT, pgpdA::matR; pabaA1, yA2; argBA::ttpC; ttpC801; veAA::argB pri1A::vleA::argB; mi1AT, pgpdA::matR; pyroA4, pyrG89		_
biA1; pyroA4, veA1 pabaA1, yA2; argBA:ttpC; ttpC801; veAA::argB pabaA1 AGB154 AGB272 pveA::weA, ptrA; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB pveA::weA,:ctap* tag, ptrA; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB pveA::weA::gfp, ptrA; pgpdA::marfp::h2A, pgpdA::matR; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB pniiA::welB::sgfp::miiAT, pgpdA::matR; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB pniiA::welB::miiTC; ttpC801, veAA::argB pniiA::welB::miiAT, pgpdA::matR; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB pyroA4, pyrG89, veA pgpdA::matR; pyroA4, pyrG89 pniiA::welB::miiAT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::matR; pyroA4, pyrG89 pniiA::welB::miiAT, A.f. pyrG; pyroA4, pyrG89 priiA::welB::miiAT, A.f. pyrG; pyroA4, pyrG89 priiA::welB::sgfp::miiAT, A.f. pyrG; pyroA4, argB2; nkuAA::argB aGB280 pvelB::welB, pgpdA::matR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB pniiA::welB::sgfp::miiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB pniiA::weA::miiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB priiA::weA::miiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB pniiA::weA::miiAT, ppdA::matR; pabaA1, yA2; argBA::ttpC; ttpC801; veAA::argB pniiA::weB::miiAT, ppdA::matR; pyroA4, pyrG89 pyroA4, argB2; nkuAA::argB pniiA::weB::miiAT, ppdA::matR; pyroA4, pyrG89 pyroA4, argB2; nkuAA::argB pniiA::weB::miiAT, ppdA::matR; pyroA4, pyrG89 pyroA4, argB2; nkuAA::argB pniiA::laeA::sgfp::miiAT, ppdA::matR; pyroA4, pyrG89 pyroA4, argB2; nk	FGSC4	Glasgow wild-type
DVAR1 pabaA1 pabaA1 AGB154 pabaA1 pobaA1 AGB272 pveA:weA, ptrA; pabaA1, yA2; argBA:ttpC; trpC801, veAA::argB AGB273 pveA:weA::ctap* tag, ptrA; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB AGB274 pveA::weA::sefgb, ptrA; pgpdA::matFp:btAA, pgpdA::matR; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB AGB275 pniiA::weB::sefgb::miiAT, pgpdA::matR; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB AGB276 pniiA::weB::miiAT, pgpdA::matR; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB AGB276 pniiA::weB::miiAT, pgpdA::matR; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB AGB152 pyroA4, pyrG89, veA ppiA::weB::seffp::miiAT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::matR; pyroA4, pyrG89 AGB278 pniiA::weB::miiAT, A.f. pyrG; pyroA4, pyrG89 AGB279 velBa::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB pveB::weB, pgpdA::matR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuA2::argB priiA::weB::seffp::miiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuA2::argB AGB281 pniiA::weB::seffp::miiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuA2::argB AGB283 pniiA::weB::miiAT, pgpdA::matR; pabaA1, yA2; argBA::ttpC; trpC801; veAA::argB AGB304 pniiA::weB::miiAT, pgpdA::matR; pabaA1, yA2; argBA::ttpC; trpC801; veAA::argB	FGSC26	biA1, veA1
AGB154 pabaA1 AGB272 pveA:veA, ptrA; pabaA1, yA2; argBA:trpC; trpC801, veAA::argB AGB273 pveA:veA:eargB pread:mrf; pabaA1, yA2; argBA:trpC; trpC801, veAA::argB AGB274 pveA:veA:sgfp, ptrA; pgpdA::mrfp::h2A, pgpdA::matR; pabaA1, yA2; argBA:trpC; trpC801, veAA::argB AGB275 pniiA:veB:sgfp::miiAT, pgpdA::matR; pabaA1, yA2; argBA:trpC; trpC801, veAA::argB AGB276 pniiA:veBB::miiAT, pgpdA::matR; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB AGB276 pniiA:veBB::miiAT, pgpdA::matR; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB AGB277 pniiA:veBB::miiAT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::matR; pyrG4, pyrG89 AGB278 pniiA:veBB::miiAT, A.f. pyrG; pyroA4, pyrG89 PyrG89, pyroA4 veBBa::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB PoelB::velB, pgpdA::matR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB pread::veA::sgfp, pgpdA::matR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB pread::veA::miiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB priiA::veB::miiAT, pgpdA::matR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB priiA::veB::miiAT, pgpdA::matR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB priiA::veB::miiAT, pgpdA::matR; pabaA1, yA2; argBA::trpC; trpC801; veAx::argB; pyroA4, pyrG89 priiA::laeA::sgfp::miiAT, pgpdA::matR; pabaA1, yA2; argBA::trpC; trpC801; veA	FGSC33	
AGB272 pveA::weA, ptrA; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB AGB273 pveA::weA::etap* tag, ptrA; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB AGB274 pveA::weA::sgfp, ptrA; pgpdA::matR; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB AGB275 pniiA::weB::sgfp::miiAT, pgpdA::natR; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB AGB276 pniiA::weB::sgfp::miiAT, pgpdA::natR; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB AGB276 pniiA::weB::sgfp::miiAT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB277 pniiA::weB::sgfp::miiAT, A.f. pyrG; pgpoAA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB278 pniiA::weB::sgfp::miiAT, A.f. pyrG; pyroA4, pyrG89 AGB279 velBa::ptrA; pyrG89, pyroA4, argB2; nkuAa::argB AGB280 pvelB::weB, pgpdA::natR; velBa::ptrA; pyrG89, pyroA4, argB2; nkuAa::argB AGB281 pniiA::weB::sgfp::miiAT, a.f. pyrG; velBa::ptrA; pyrG89, pyroA4, argB2; nkuAa::argB AGB283 pniiA::weB::miiAT, pgpdA::natR; velBa::ptrA; pyrG89, pyroA4, argB2; nkuAa::argB AGB284 pniiA::weB::miiAT, pgpdA::natR; pyrG89, pyroA4, argB2; nkuAa::argB AGB308 pniiA::weB::miiAT, pgpdA::natR; pabaA1, yA2; argBA::ttpC; ttpC801; veAA::argB AGB310 pniiA::laeA::sgfp::miiAT, pgpdA::natR; pabaA1, yA2; argBA::ttpC; ttpC801; veAA::argB AGB38		
AGB273 yeeAxiveAi:ctap* tag, ptrA; pabaA1, yA2; argBA::trpC; trpC801, yeAA::argB AGB274 pveA::veAi:saffp, ptrA; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB AGB275 proiA::veAi::saffp, ptrA; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB AGB276 pniiA::veBi::saffp::miiAT, pgdA::natR; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB AGB276 pniiA::veBi::saffp::miiAT, paf. pyrG; pgpdA::mrfp::h2A, pgdA::matR; pyroA4, pyrG89 AGB277 pniiA::veBi::saffp::miiAT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::matR; pyroA4, pyrG89 AGB278 pniiA::veBi::miiAT, A.f. pyrG; pyroA4, pyrG89 PyrG89, pyroA4 veBa::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB280 pveBi::veB, pgpdA::matR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB281 pniiA::veA::miiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB282 pveA::veA::miiAT, af, pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB283 pniiA::veA::miiAT, pgpdA::matR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB284 pniiA::veA::miiAT, pgpdA::matR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB307 pniiA::mpd::miAT, pgpdA::matR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB310 pniiA::laeA::sgfp::miiAT, pgpdA::matR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB389<		•
AGB274 veA::argB pveA::veA::sgfp, ptrA; pgpdA::mrfp::h2A, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB AGB275 pniiA::veB::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB AGB276 pniiA::velB::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801, veAA::argB AGB152 pyroA4, pyrG89, veA AGB277 pniiA::velB::sgfp::niiAT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB278 pniiA::velB::miiAT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyrG89 AGB279 velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB280 pvelB::velB, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB281 pniiA::velB::sgfp::niiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB282 pveA::veA::sgfp, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB283 pniiA::veA::niiAT, pgrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB284 pniiA::veA::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB307 pniiA::nyfp::veA::iiAT, ppgdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB310 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB388 pniiA::nyfrp::veA::niiAT-pniaD::cyfp::velB::niiaDT, A.f. pyrG; pgpdA	AGB272	
AGB274 pveA::veA::sgfp, ptrA; pgpdA::mrfp::h2A, pgpdA::matR; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB AGB275 pniiA::velB::sgfp::miiAT, pgpdA::matR; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB AGB276 pniiA::velB::miiAT, pgpdA::matR; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB AGB277 pniiA::velB::sgfp::miiAT, pgpdA::matR; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB AGB277 pniiA::velB::sgfp::miiAT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::matR; pyroA4, pyrG89 AGB278 pniiA::velB::sgfp::miiAT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::matR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB280 pvelB::velB. pgpdA::matR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB281 pniiA::velB::sgfp::miiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB282 pveA::veA::sgfp, pgpdA::matR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB283 pniiA::veA::miiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB284 pniiA::veB::miiAT, pgpdA::natR; pabaA1, yA2; argBA::ttpC; ttpC801; veAA::argB AGB307 pniiA::neA::ssgfp::miiAT, pgpdA::natR; pabaA1, yA2; argBA::ttpC; ttpC801; veAA::argB AGB308 pniiA::laeA::ssgfp::miiAT, pgpdA::natR; pabaA1, yA2; argBA::ttpC; ttpC801; veAA::argB AGB311 pniiA::laeA::ssgfp::miiAT, pgpdA::natR; pabaA1, yA2; argBA::ttpC; ttpC801; veAA::argB AGB388 pniiA::laeA::ssgfp::miiAT, pgpdA::natR; pabaA1, yA2; argBA::ttpC; ttpC801; veAA::argB AGB389 piiA::laeA::ssgfp::miiAT, pgpdA::natR; pabaA1, yA2; argBA::ttpC; ttpC801; veAA::argB AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR AGB390 laeA::pyrG; veA AIIIA: velBa:::pyrG; laeAA::metG, veA AIIIA: velBa:::pyrG; laeAA::metG, veA AIIIA: veAA::argB; laeAA::metG, veA AIIIA: veAA::argB; laeAA::metG	AGB273	
AGB275 pniiA::velB::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB AGB276 pniiA::velB::niiAT, pppdA::natR; pabaA1, yA2; argBA::ttpC; ttpC801, veAA::argB AGB152 pyroA4, pyrG89, veA AGB277 pniiA::velB::siniiAT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB278 pniiA::velB::niiAT, A.f. pyrG; pyroA4, pyrG89 AGB279 pvlBa::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB280 pvelB::velB, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB281 pviiA::veA::sigp. pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB282 pveA::veA::sigp. pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB283 pniiA::veA::miiAT, pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB284 pniiA::veA::miiAT, pgpdA::natR; pabaA1, yA2; argBA::ttpC; ttpC801; veAA::argB AGB307 pniiA::nyfp::veA::niiAT-pniaD::cyfp::laeA::niaDT, A.f. pyrG; pgpdA::matR; pyroA4, pyrG89 AGB308 pniiA::laeA::sgfp::miiAT, pgpdA::natR; pabaA1, yA2; argBA::ttpC; ttpC801; veAA::argB AGB311 pniiA::laeA::sgfp::miiAT, pgpdA::natR; velBa::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB388 pniiA::laeA::sgfp::miiAT, pgpdA::natR; velBa::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB; pyrA4; nkuAA::argB; veA1 TNI7.2 <td>AGB274</td> <td>pveA::veA::sgfp, ptrA; pgpdA::mrfp::h2A, pgpdA::natR; pabaA1,</td>	AGB274	pveA::veA::sgfp, ptrA; pgpdA::mrfp::h2A, pgpdA::natR; pabaA1,
AGB152 AGB152 AGB152 AGB152 AGB153 AGB154 AGB277 AGB277 AGB278 AGB278 AGB278 AGB278 AGB279 AGB279 AGB279 AGB279 AGB279 AGB280 AGB280 AGB280 AGB280 AGB281 AGB281 AGB281 AGB281 AGB282 AGB282 AGB282 AGB282 AGB283 AGB283 AGB283 AGB283 AGB284 AGB284 AGB284 AGB284 AGB284 AGB285 AGB285 AGB285 AGB286 AGB286 AGB287 AGB287 AGB287 AGB288 AGB288 AGB288 AGB288 AGB288 AGB288 AGB288 AGB288 AGB289 AGB289 AGB280 AGB280 AGB280 AGB280 AGB281 AGB281 AGB281 AGB281 AGB282 AGB282 AGB283 AGB283 AGB283 AGB284 AGB307 AGB308 AGB308 AGB308 AGB309 AGB308 AGB309 AGB308 AGB309 AGB308 AGB310 AGB308 AGB310 AGB308 AGB311 AGB31 AGB31 AGB31 AGB31 AGB31 AGB31 A	AGB275	pniiA::velB::sgfp::niiAT, pgpdA::natR; pabaA1, yA2;
trpC801, veAA::argB pyroA4, pyrG89, veA ppiiA::welB::sgfp::niiAT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::matR; pyroA4, pyrG89 pniiA::welB::sifp::niiAT, A.f. pyrG; pyroA4, pyrG89 priiA::welB::miiAT, A.f. pyrG; pyroA4, pyrG89 priiA::welB::miiAT, A.f. pyrG; pyroA4, pyrG89 proA4, pyrG89, pyroA4 pyrG89, pyroA4 pyrG89, pyroA4, argB2; nkuAA::argB priiA::welB::sgfp::niiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB pniiA::welB::sgfp::niiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB proA1::welB::sgfp::niiAT, a.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB pniiA::welB::niiAT, pgpdA::matR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB pniiA::welB::niiAT, pgpdA::matR; pabaA1, yA2; argBA::trpCstrpC801; veAA::argB pniiA::myfp::weA::miiAT-pniaD::cyfp::laeA::miaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::matR; pabaA1, yA2; argBa::trpC; trpC801; veAA::argB pniiA::laeA::sgfp::miiAT, pgpdA::matR; pabaA1, yA2; argBa::trpC; trpC801; veAA::argB pniiA::laeA::sgfp::miiAT, pgpdA::matR; pabaA1, yA2; argBa::trpC; trpC801; veAA::argB pniiA::laeA::sgfp::miiAT, pgpdA::matR; pabaA1, yA2; argBa::trpC; trpC801; veAA::argB priiA::laeA::sgfp::miiAT, pgpdA::matR; velBa::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB priiA::laeA::sgfp::miiAT, pgpdA::matR; velBa::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB priiA::laeA::sgfp::miiAT, pgpdA::matR; velBa::ptrA; pyrG89, pyroA4, pyrG89 pyroA4, pyrG89 pyroA4, pyrG89 pyroA4, pyrG89 pyroA4, pyrG89 pyroA4, pyrG89 pyroA4; nkuA2::argB; veAl pyrG89, pyroA4; nkuA2::argB; veAl pyrG89, pyrO44; nkuA2::argB; laeA2::metG, veA pyrG81 pyrG84::argB; laeA2::metG, veA pyrG12: pyrG12::argB::ar	ACD276	
AGB152 AGB277 pniiA::welB::sgfp::miiAT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 pniiA::welB::miiAT, A.f. pyrG; pyroA4, pyrG89 TNO2A3 pyrG89, pyroA4 AGB279 velB::welB, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB pvelB::welB, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB281 pniiA::welB::sgfp::miiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB282 pveA::weA::sgfp, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB pniiA::welB::miiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB283 pniiA::weA::miiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB284 pniiA::weB::miiAT, pgpdA::natR; pyrG89, pyroA4, argB2; nkuAA::argB AGB307 pniiA::weA::miiAT-piaD::cyfp::laeA::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 pniiA::laeA::sgfp::miiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB311 pniiA::laeA::sgfp::miiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB311 pniiA::laeA::sgfp::miiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB388 pniiA::nyfp::weA::niiAT-pniaD::cyfp::velB::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR AGB390 laeA::ctap*::p	AUB2/0	
AGB277 pniiA::velB::sgfp::niiAT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB278 pniiA::velB::niiAT, A.f. pyrG; pyroA4, pyrG89 TNO2A3 pyrG89, pyroA4 AGB279 velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB280 pvelB::velB, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB281 pniiA::velB::sgfp::niiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB282 pveA::veA::sgfp, pgpdA::matR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB283 pniiA::veA::niiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB284 pniiA::velB::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; ttpC801; veAA::argB AGB307 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pyroA4, pyrG89 pmiiA::laeA::sgfp::niiAT, pgpdA::natR; pyroA4, pyrG89 pmiiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::ttpC; ttpC801; veAA::argB AGB310 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::ttpC; ttpC801; veAA::argB AGB311 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::mipd::niiAT, pgpdA::natR; pyroA4, pyrG89 AGB388 pniiA::nyfp::veA::niiAT-pniaD::cyfp::velB::niaDT, A.f. pyrG; pgpdA::matR; pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::cta	AGB152	
pgpdA::natR; pyroA4, pyrG89 pniiA::velB::niiAT, A.f. pyrG; pyroA4, pyrG89 pyrG89, pyroA4 AGB279		
AGB278nniiA::welB::miiAT, A.f. pyrG; pyroA4, pyrG89TNO2A3pyrG89, pyroA4AGB279velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argBAGB280pvelB::welB, pgpdA::matR; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argBAGB281pniiA::welB::sgfp::miiAT, A.f. pyrG; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argBAGB282pveA::weA::sgfp, pgpdA::matR; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argBAGB283pniiA::welB::miiAT, A.f. pyrG; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argBAGB284pniiA::welB::miiAT, pgpdA::matR; pabaA1, yA2; argBΔ::trpC; trpC801; veAΔ::argBAGB307pniiA::myfp::veA1::niiAT-pniaD::cyfp::laeA::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89AGB308pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBΔ::trpC; trpC801; veAA::argBAGB310pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBΔ::trpC; trpC801; veAA::argBAGB311pniiA::laeA::sgfp::niiAT, pgpdA::natR; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argBAGB388pniiA::laeA::sgfp::niiAT-pniaD::cyfp::velB::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyrG89, pyroA4, pyrG89AGB389velB::ctap*::pgpdA::natRAGB390laeA::ctap*::pgpdA::natRAGB390laeA::ctap*::pgpdA::natRAGB390laeA::ctap*::pgpdA::natRAGB390laeA::ctap*::pgpdA::natRAGB390laeA::ctap*::pgpdA::natRAGB390laeA::ctap*::pgrG9, pyrO4, nkuAA::argB; veA1RNW108.1velBΔ::pyrG; teARNW108.1velBΔ::pyrG; laeAΔ::metG, veARNW108.1veAA::argB; taeAA::metG, veAVeNU10.2veAA::argB; la	1GB277	
TNO2A3 pyrG89, pyroA4 AGB279 velBA::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argB AGB280 pvelB::velB, pgpdA::natR; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB281 pniiA::velB::sgfp::niiAT, A.f. pyrG; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB282 pveA::veA::sgfp, pgpdA::natR; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB283 pniiA::veA::niiAT, A.f. pyrG; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB284 pniiA::veA::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB307 pniiA::nyfp::veA::niiAT-pniaD::cyfp::laeA::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB310 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB311 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB388 pniiA::laeA::sgfp::niiAT, ppdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB389 pniiA::natR; ppidA::natR; ppidA::natR; pyrG89, pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgdA::natR AGB390 laeA::ctap*::pgdA::natR <t< td=""><td>AGB278</td><td></td></t<>	AGB278	
AGB279 velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB280 pvelB::velB, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB281 pniiA::velB::sgfp::niiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB282 pveA::veA::sgfp, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB283 pniiA::veAB::niiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB284 pniiA::velB::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB307 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pyroA4, pyrG89 AGB308 pniiA::laeA::sgfp::niiAT, pgpdA::natR AGB310 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB311 pniiA::laeA::sgfp::niiAT, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB388 pniiA::nyfp::veA::niiAT-pniaD::cyfp::velB::niaDT, A.f. pyrG; pgpdA::mrlfp::h2A, pgpdA::natR; pyrG89, pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR AGB390 laeA::ctap*::pyrG; pyrG89, pyroA4; nkuAA::argB; veA1 TNT7.2 velBA::pyrG; pyrG89, pyroA4; nkuAA::argB; veA1 RW416.1 velBA::pyrG; teA		
nkuAA::argB pniiA::velB::sgfp::niiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB pveA::veA::sgfp, ppgdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB pniiA::veA::niiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB pniiA::veA::niiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB pniiA::veA::niiAT, ppgdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB pniiA::nyfp::veA::niiAT-pniaD::cyfp::laeA::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 pniiA::laeA::sgfp::niiAT, pgpdA::natR pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB pniiA::laeA::sgfp::niiAT, pgpdA::natR; pyroA4, argB2; nkuAA::argB pniiA::nyfp::veA::niiAT-pniaD::cyfp::velBa::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 pyroA4, pyrG89 ppiiA::nyfp::veA::niiAT-pniaD::cyfp::velB::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 ppiiA::tpyrG89 ppiiA::tpyrG89 ppiiA::pyrG89, pyroA4; nkuAA::argB; veA1 pyrG89, pyroA4; nkuAA::argB; veA1 pyrG89, yA2; veA welBa::pyrG; pyrG89, pyroA4; nkuAA::argB; veA1 pyrG89, yA2; veA wild type RNW10.1 velBa::pyrG; teA RNW10.1 velBa::pyrG; teA RNW10.1 veA::argB; laeAA::metG, veA RNW10.2 veAa::argB RNW11.1 vosAA::argB; laeAA::metG, veA surW11.1 vosAA::argB; laeAA::metG, veA surW11.1 vosAA::argB; laeAA::metG Saccharomyces cerevisiae	AGB279	
AGB281 pniiA::velB::sgfp::niiAT, A.f. pyrG; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argB pveA::veA::sgfp, pgpdA::natR; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argB AGB283 pniiA::veA::niiAT, A.f. pyrG; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argB AGB284 pniiA::velB::niiAT, pgpdA::natR; pabaA1, yA2; argBΔ::trpC; trpC801; veAΔ::argB AGB307 pniiA::nyfp::veA::niiAT-pniaD::cyfp::laeA::niaDT, A.f. pyrG; pgpdA::matR; pyroA4, pyrG89 AGB308 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBΔ::trpC; trpC801; veAΔ::argB AGB310 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBΔ::trpC; trpC801; veAΔ::argB AGB311 pniiA::laeA::sgfp::niiAT, pgpdA::natR; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argB AGB388 pniiA::nyfp::veA::niiAT-pniaD::cyfp::velB::niaDT, A.f. pyrG; pgpdA::mtpfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR		pvelB::velB, pgpdA::natR; velB\Delta::ptrA; pyrG89, pyroA4, argB2;
argB2; nkuAΔ::argB pveA::weA::sgfp, pgpdA::natR; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argB pniiA::veA::niiAT, A.f. pyrG; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argB AGB284 pniiA::velB::niiAT, pgpdA::natR; pabaA1, yA2; argBΔ::trpC; trpC801; veAΔ::argB AGB307 pniiA::nyfp::veA::niiAT-pniaD::cyfp::laeA::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB308 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAΔ::argB AGB310 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAΔ::argB AGB311 pniiA::laeA::sgfp::niiAT, pgpdA::natR; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argB AGB388 pniiA::nyfp::veA::niiAT-pniaD::cyfp::velB::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR RNI16.1 pyrG89, pyroA4; nkuAΔ::argB; veA1 TNI7.2 velBΔ::pyrG; pyrG89, pyroA4; nkuAΔ::argB; veA1 TNI7.2 velBΔ::pyrG; pyrG89, pyroA4; nkuAΔ::argB; veA1 RRAW16 pyrG89, yA2; veA RNI18.2 velBΔ::pyrG; veA RNI18.2 velBΔ::pyrG; laeAΔ::metG, veA RJW110.1 velBΔ::pyrG; laeAΔ::metG, veA RJW110.1 veAΔ::argB; laeAΔ::metG, veA RJW110.2 veAΔ::argB RJW114.11 vosAΔ::argB; laeAΔ::metG Saccharomyces cerevisiae		
AGB282 pveA::veA::sgfp, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB283 pniiA::veA::niiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB284 pniiA::velB::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB307 pniiA::wlpp::veA::niiAT-pniaD::cyfp::laeA::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB308 pniiA::laeA::sgfp::niiAT, pgpdA::natR AGB310 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB311 pniiA::laeA::sgfp::niiAT, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB388 pniiA::nyfp::veA::niiAT-pniaD::cyfp::velB::niiaDT, A.f. pyrG; pgpdA::mtfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR RNI16.1 pyrG89, pyroA4; nkuAA::argB; veA1 TNI7.2 velBA::pyrG; pyrG89, pyroA4; nkuAA::argB; veA1 RRAW16 pyrG89, yA2; veA RDIT9.32 wild type RJW41.A laeAA::metG; veA RJW116.1 velBA::pyrG; laeAA::metG, veA RJW106.1 velBA::pyrG; laeAA::metG, veA RJW108.1 veAA::argB; trpC::laeA RJW112.2 veAA::argB; laeAA::metG, veA RJW116.2 vosAA::argB; laeAA::metG, veA RJW116.2 vosAA::argB; laeAA::metG	AGB281	
argB2; nkuAA::argB pniiA::veA::niiAT, A.f. pyrG; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB284 pniiA::velB::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB307 pniiA::nyfp::veA::niiAT-pniaD::cyfp::laeA::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB308 pniiA::laeA::sgfp::niiAT, pgpdA::natR AGB310 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB311 pniiA::laeA::sgfp::niiAT, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB388 pniiA::nyfp::veA::niiAT-pniaD::cyfp::velB::niaDT, A.f. pyrG; pgpdA::matR; pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR RNI16.1 pyrG89, pyroA4; nkuAA::argB; veA1 velBA::pyrG; pyrG89, pyroA4; nkuAA::argB; veA1 RRAW16 pyrG89, yA2; veA RNI18.2 velBA::pyrG; veA RDIT9.32 wild type RJW41.A laeAA::metG; veA RJW1106.1 velBA::pyrG; laeAA::metG, veA RJW108.1 veAA::argB; trpC::laeA RJW112.2 veAA::argB RJW11.11 vosAA::argB; laeAA::metG, veA RJW116.2 vosAA::argB; laeAA::metG, veA RJW117.18 Saccharomyces cerevisiae	. GDaca	<i>c</i> , <i>c</i>
AGB283 pniiA::veA::niiAT, A.f. pyrG; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argB AGB284 pniiA::velB::niiAT, pgpdA::natR; pabaA1, yA2; argBΔ::trpC; trpC801; veAΔ::argB AGB307 pniiA::nyfp::veA::niiAT-pniaD::cyfp::laeA::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB308 pniiA::laeA::sgfp::niiAT, pgpdA::natR AGB310 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBΔ::trpC; trpC801; veAΔ::argB AGB311 pniiA::laeA::sgfp::niiAT, pgpdA::natR; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argB AGB388 pniiA::nyfp::veA::niiAT-pniaD::cyfp::velB::niiaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR RNI16.1 pyrG89, pyroA4; nkuAΔ::argB; veA1 TNI7.2 velBΔ::pyrG; pyrG89, pyroA4; nkuAΔ::argB; veA1 RRAW16 pyrG89, yA2; veA RNI18.2 velBΔ::pyrG; veA RDIT9.32 wild type RJW41.A laeAΔ::metG; veA RJW106.1 velBΔ::pyrG; laeAΔ::metG, veA RJW108.1 veAΔ::argB; trpC::laeA RJW112.2 veAΔ::argB RJW114.11 vosAΔ::argB; laeAΔ::metG, veA RJW116.2 vosAΔ::argB; laeAΔ::metG Saccharomyces cerevisiae	AGB282	
AGB284 pniiA::velB::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB307 pniiA::nyfp::veA::niiAT-pniaD::cyfp::laeA::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB308 pniiA::laeA::sgfp::niiAT, pgpdA::natR AGB310 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBΔ::trpC; trpC801; veAΔ::argB AGB311 pniiA::laeA::sgfp::niiAT, pgpdA::natR; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argB AGB388 pniiA::nyfp::veA::niiAT-pniaD::cyfp::velB::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR RNI16.1 pyrG89, pyroA4; nkuAA::argB; veA1 TNI7.2 velBΔ::pyrG; pyrG89, pyroA4; nkuAΔ::argB; veA1 RRAW16 pyrG89, yya2; veA RNI18.2 velBΔ::pyrG; veA RDIT9.32 wild type RJW41.A laeA::metG; veA RJW106.1 velBΔ::pyrG; laeAΔ::metG, veA RJW112.2 veAA::argB; laeAA::metG, veA RJW116.1 vosAA::argB; laeAA::metG Saccharomyces cerevisiae	AGB283	
trpC801; veAA::argB pniiA::nyfp::veA::niiAT-pniaD::cyfp::laeA::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 pniiA::laeA::sgfp::niiAT, pgpdA::natR AGB308 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB pniiA::laeA::sgfp::niiAT, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB388 pniiA::nyfp::veA::niiAT-pniaD::cyfp::velB::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 aGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR RNI16.1 pyrG89, pyroA4; nkuAA::argB; veA1 TNI7.2 velBA::pyrG; pyrG89, pyroA4; nkuAA::argB; veA1 RRAW16 pyrG89, ya2; veA RNI18.2 velBA::pyrG; veA RDIT9.32 wild type RJW41.A laeAA::metG; veA RJW106.1 velBA::pyrG; laeAA::metG, veA RJW108.1 veAA::argB; trpC::laeA RJW112.2 veAA::argB RJW114.11 vosAA::argB; laeAA::metG, veA RJW116.2 vosAA::argB; laeAA::metG, veA RJW116.2 vosAA::argB; laeAA::metG		argB2; nkuAΔ::argB
AGB307 pniiA::nyfp::veA::niiAT-pniaD::cyfp::laeA::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB308 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB310 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB311 pniiA::laeA::sgfp::niiAT, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB388 pniiA::nyfp::veA::niiAT-pniaD::cyfp::velB::niaDT, A.f. pyrG; pgpdA::mtfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR RNI16.1 pyrG89, pyroA4; nkuAA::argB; veA1 TNI7.2 velBA::pyrG; pyrG89, pyroA4; nkuAA::argB; veA1 RRAW16 pyrG89, yA2; veA RNI18.2 velBA::pyrG; veA RDIT9.32 wild type RJW41.A laeAA::metG; veA RJW106.1 velBA::pyrG; laeAA::metG, veA RJW108.1 veAA::argB; trpC::laeA RJW112.2 veAA::argB RJW114.11 vosAA::argB; laeAA::metG, veA RJW116.2 vosAA::argB; laeAA::metG, veA RJW116.1 vosAA::argB; laeAA::metG, veA RJW117.18 veAA::argB; laeAA::metG	AGB284	
pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB308 pniiA::laeA::sgfp::niiAT, pgpdA::natR AGB310 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB311 pniiA::laeA::sgfp::niiAT, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB388 pniiA::nyfp::veA::niiAT-pniaD::cyfp::velB::niaDT, A.f. pyrG; pgpdA::matR; pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR RNI16.1 pyrG89, pyroA4; nkuAA::argB; veA1 TNI7.2 velBA::pyrG; pyrG89, pyroA4; nkuAA::argB; veA1 RRAW16 pyrG89, yA2; veA RNI18.2 velBA::pyrG; veA RDIT9.32 wild type RJW41.A laeAA::metG; veA RJW106.1 velBA::pyrG; laeAA::metG, veA RJW108.1 veAA::argB; trpC::laeA RJW112.2 veAA::argB; laeAA::metG, veA RJW116.2 vosAA::argB; laeAA::metG, veA RJW116.1 vosAA::argB; laeAA::metG, veA RJW117.18 vosAA::argB; laeAA::metG Saccharomyces cerevisiae	. GD207	
AGB308 pniiA::laeA::sgfp::niiAT, pgpdA::natR AGB310 pniiA::laeA::sgfp::niiAT, pgpdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB311 pniiA::laeA::sgfp::niiAT, pgpdA::natR; velBA::ptrA; pyrG89, pyroA4, argB2; nkuAA::argB AGB388 pniiA::nyfp::veA::niiAT-pniaD::cyfp::velB::niaDT, A.f. pyrG; pgpdA::matR; pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR RNI16.1 pyrG89, pyroA4; nkuAA::argB; veA1 TNI7.2 velBA::pyrG; pyrG89, pyroA4; nkuAA::argB; veA1 RRAW16 pyrG89, yA2; veA RNI18.2 velBA::pyrG; veA RDIT9.32 wild type RJW41.A laeAA::metG; veA RJW106.1 velBA::pyrG; laeAA::metG, veA RJW108.1 veAA::argB; trpC::laeA RJW112.2 veAA::argB RJW11.11 vosAA::argB RJW11.12 vosAA::argB; laeAA::metG, veA RJW116.2 vosAA::argB; laeAA::metG RJW117.18 veAA::argB; laeAA::metG Saccharomyces cerevisiae	AGB307	
AGB310 pniiA::laeA::sgfp::niiAT, pspdA::natR; pabaA1, yA2; argBA::trpC; trpC801; veAA::argB AGB311 pniiA::laeA::sgfp::niiAT, pspdA::natR; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAa::argB AGB388 pniiA::nyfp::veA::niiAT-pniaD::cyfp::velB::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR RNI16.1 pyrG89, pyroA4; nkuAΔ::argB; veA1 TNT7.2 velBΔ::pyrG; pyrG89, pyroA4; nkuAΔ::argB; veA1 RRAW16 pyrG89, yA2; veA RNI18.2 velBΔ::pyrG; veA RNI18.2 velBΔ::pyrG; veA RDIT9.32 wild type RJW41.A laeAA::metG; veA RJW106.1 velBΔ::pyrG; laeAΔ::metG, veA RJW108.1 veAΔ::argB; trpC::laeA RJW112.2 veAΔ::argB RJW114.11 vosAΔ::argB; laeAΔ::metG, veA RJW116.2 vosAΔ::argB; laeAΔ::metG Saccharomyces cerevisiae	ACD209	
argBA::trpC; trpC801; veAA::argB pniiA::laeA::sgfp::niiAT, pgpdA::natR; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argB AGB388 pniiA::nyfp::veA::niiAT-pniaD::cyfp::velB::niaDT, A.f. pyrG; pgpdA::natR; pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR RNI16.1 pyrG89, pyroA4; nkuAΔ::argB; veA1 TNI7.2 velBA::pyrG; pyrG89, pyroA4; nkuAΔ::argB; veA1 RRAW16 pyrG89, yA2; veA RNI18.2 velBΔ::pyrG; veA RNI18.2 velBΔ::pyrG; veA RDIT9.32 wild type RJW41.A laeAΔ::metG; veA RJW106.1 veAA::argB; tpC::laeA RJW112.2 veAA::argB RJW114.11 vosAA::argB; laeAA::metG, veA RJW116.2 vosAA::argB; laeAA::metG Saccharomyces cerevisiae		
AGB311 pniiA::laeA::sgfp::niiAT, pgpdA::natR; velBΔ::ptrA; pyrG89, pyroA4, argB2; nkuAΔ::argB AGB388 pniiA::nyfp::veA::niiAT-pniaD::cyfp::velB::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR RNI16.1 pyrG89, pyroA4; nkuAΔ::argB; veA1 TNT7.2 velBΔ::pyrG; pyrG89, pyroA4; nkuAΔ::argB; veA1 RRAW16 pyrG89, yA2; veA RNI18.2 velBΔ::pyrG; veA RDIT9.32 wild type RJW41.A laeAΔ::metG; veA RJW106.1 velAΔ::argB; taeAΔ::metG, veA RJW112.2 veAA::argB; laeAΔ::metG, veA RJW114.11 vosAA::argB; laeAA::metG, veA RJW117.18 veAA::argB; laeAA::metG Saccharomyces cerevisiae	AGB310	
pyroA4, argB2; nkuAΔ::argB	ACD211	
AGB388 pniiA::nyfp::veA::niiAT-pniaD::cyfp::velB::niaDT, A.f. pyrG; pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR RNI16.1 pyrG89, pyroA4; nkuAΔ::argB; veA1 TNI7.2 velBΔ::pyrG; pyrG89, pyroA4; nkuAΔ::argB; veA1 RRAW16 pyrG89, yA2; veA RNI18.2 velBΔ::pyrG; veA RDIT9.32 wild type RJW41.A laeAΔ::metG; veA RJW106.1 velBΔ::pyrG; laeAΔ::metG, veA RJW108.1 veAΔ::argB; trpC::laeA RJW112.2 veAΔ::argB RJW114.11 vosAΔ::argB; laeAΔ::metG, veA RJW116.2 vosAΔ::argB; laeAΔ::metG Saccharomyces cerevisiae	AGBIT	
pgpdA::mrfp::h2A, pgpdA::natR; pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR RNI16.1 pyrG89, pyroA4; nkuAA::argB; veA1 TNI7.2 velBA::pyrG; pyrG89, pyroA4; nkuAA::argB; veA1 RRAW16 pyrG89, yA2; veA RNI18.2 velBA::pyrG; veA RDIT9.32 wild type RJW41.A laeAA::metG; veA RJW106.1 velBA::pyrG; laeAA::metG, veA RJW108.1 veAA::argB; trpC::laeA RJW112.2 veAA::argB RJW114.11 vosAA::argB; laeAA::metG, veA RJW116.2 vosAA::argB; laeAA::metG, veA RJW117.18 veAA::argB; laeAA::metG Saccharomyces cerevisiae	ACD300	
pyroA4, pyrG89 AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR RNI16.1 pyrG89, pyroA4; nkuAΔ::argB; veA1 TNI7.2 velBΔ::pyrG; pyrG89, pyroA4; nkuAΔ::argB; veA1 RRAW16 pyrG89, yA2; veA RNI18.2 velBA::pyrG; veA RDIT9.32 wild type RJW41.A laeAΔ::metG; veA RJW106.1 velBΔ::pyrG; laeAΔ::metG, veA RJW108.1 veAΔ::argB; tpC::laeA RJW112.2 veAΔ::argB RJW114.11 vosAΔ::argB RJW116.2 vosAΔ::argB; laeAΔ::metG, veA RJW116.2 vosAΔ::argB; laeAΔ::metG Saccharomyces cerevisiae	AGDJ66	
AGB389 velB::ctap*::pgpdA::natR AGB390 laeA::ctap*::pgpdA::natR RNI16.1 pyrG89, pyroA4; nkuAΔ::argB; veA1 TNI7.2 velBΔ::pyrG; pyrG89, pyroA4; nkuAΔ::argB; veA1 RRAW16 pyrG89, yA2; veA RNI18.2 velBΔ::pyrG; veA RDIT9.32 wild type RJW41.A laeAΔ::metG; veA RJW106.1 velBΔ::pyrG; laeAΔ::metG, veA RJW108.1 veAΔ::argB; trpC::laeA RJW112.2 veAΔ::argB; laeAΔ::metG, veA RJW114.11 vosAΔ::argB; veA RJW116.2 vosAΔ::argB; laeAΔ::metG Saccharomyces cerevisiae		
AGB390 laeA::ctap*::pgpdA::natR RNI16.1 pyrG89, pyroA4; nkuAΔ::argB; veA1 TNI7.2 velBΔ::pyrG; pyrG89, pyroA4; nkuAΔ::argB; veA1 RRAW16 pyrG89, yA2; veA RNI18.2 velBΔ::pyrG; veA RDIT9.32 wild type RJW41.A laeAΔ::metG; veA RJW106.1 velBΔ::pyrG; laeAΔ::metG, veA RJW108.1 veAΔ::argB; trpC::laeA RJW112.2 veAΔ::argB RJW114.11 vosAΔ::argB; laeAΔ::metG, veA RJW116.2 vosAΔ::argB; veA RJW117.18 veAΔ::argB; laeAΔ::metG Saccharomyces cerevisiae	AGR389	
RNI16.1 pyrG89, pyroA4; nkuAΔ::argB; veA1 TNI7.2 velBΔ::pyrG; pyrG89, pyroA4; nkuAΔ::argB; veA1 RRAW16 pyrG89, yA2; veA RNI18.2 velBΔ::pyrG; veA RDIT9.32 wild type RJW41.A laeAΔ::metG; veA RJW106.1 velBΔ::pyrG; laeAΔ::metG, veA RJW108.1 veAΔ::argB; trpC::laeA RJW112.2 veAΔ::argB RJW114.11 vosAΔ::argB; laeAΔ::metG, veA RJW116.2 vosAΔ::argB; laeAΔ::metG RJW117.18 veAΔ::argB; laeAΔ::metG		1 101
TNI7.2 velBΔ::pyrG; pyrG89, pyroA4; nkuAΔ::argB; veA1 RRAW16 pyrG89, yA2; veA RNI18.2 velBΔ::pyrG; veA RDIT9.32 wild type RJW41.A laeAΔ::metG; veA RJW106.1 velBΔ::pyrG; laeAΔ::metG, veA RJW108.1 veAΔ::argB; trpC::laeA RJW112.2 veAΔ::argB RJW114.11 vosAΔ::argB; laeAΔ::metG, veA RJW116.2 vosAΔ::argB; laeAΔ::metG RJW117.18 veAΔ::argB; laeAΔ::metG		
RRAW16 pyrG89, yA2; veA RNI18.2 velBΔ::pyrG; veA RDIT9.32 wild type RJW41.A laeAΔ::metG; veA RJW106.1 velBΔ::pyrG; laeAΔ::metG, veA RJW108.1 veAΔ::argB; tpC::laeA RJW112.2 veAΔ::argB; laeAΔ::metG, veA RJW114.11 vosAΔ::argB; veA veAJ::argB; veA veAΔ::argB; laeAΔ::metG		
RNI18.2 velBA::pyrG; veA RDIT9.32 wild type RJW41.A laeAA::metG; veA RJW106.1 velBA::pyrG; laeAA::metG, veA RJW108.1 veAA::argB; trpC::laeA RJW112.2 veAA::argB RJW114.11 vosAA::argB; laeAA::metG, veA RJW116.2 vosAA::argB; veA RJW117.18 veAA::argB; laeAA::metG Saccharomyces cerevisiae		
RDIT9.32 wild type RJW41.A laeAA::metG; veA RJW106.1 velBA::pyrG; laeAA::metG, veA RJW108.1 veAA::argB; trpC::laeA RJW112.2 veAA::argB RJW114.11 vosAA::argB; laeAA::metG, veA RJW116.2 vosAA::argB; veA RJW117.18 veAA::argB; laeAA::metG Saccharomyces cerevisiae		
RJW41.A laeAΔ::metG; veA RJW106.1 velBΔ::pyrG; laeAΔ::metG, veA RJW108.1 veAΔ::argB; trpC::laeA RJW112.2 veAΔ::argB RJW114.11 vosAΔ::argB; laeAΔ::metG, veA RJW116.2 vosAΔ::argB; veA RJW117.18 veAΔ::argB; laeAΔ::metG Saccharomyces cerevisiae		
RJW106.1 velBΔ::pyrG; laeAΔ::metG, veA RJW108.1 veAΔ::argB; trpC::laeA RJW112.2 veAΔ::argB RJW114.11 vosAΔ::argB; laeAΔ::metG, veA RJW116.2 vosAΔ::argB; veA RJW117.18 veAΔ::argB; laeAΔ::metG Saccharomyces cerevisiae		**
RJW108.1 veAΔ::argB; trpC::laeA RJW112.2 veAΔ::argB RJW114.11 vosAΔ::argB; laeAΔ::metG, veA RJW116.2 vosAΔ::argB; veA RJW117.18 veAΔ::argB; laeAΔ::metG Saccharomyces cerevisiae		
RJW112.2 veAA::argB RJW114.11 vosAA::argB; laeAA::metG, veA RJW116.2 vosAA::argB; veA RJW117.18 veAA::argB; laeAA::metG Saccharomyces cerevisiae		
RJW114.11 vosAA::argB; laeAA::metG, veA RJW116.2 vosAA::argB; veA RJW117.18 veAA::argB; laeAA::metG Saccharomyces cerevisiae		
RJW116.2 vosAA::argB; veA RJW117.18 veAA::argB; laeAA::metG Saccharomyces cerevisiae		-
RJW117.18 veAA::argB; laeAA::metG Saccharomyces cerevisiae		
Saccharomyces cerevisiae		5 /
cerevisiae		· · · · · · · · · · · · · · · · · · ·
		_

A. nidulans TNO2A3 which displays a veA+phenotype served as wild-type for the velB deletion, AGB152 and DVAR1 were used for overexpression experiments. A velB gene replacement cassette comprising 2 kb of velB upstream and downstream flanking regions and the pyrithiamine resistance gene ptrA as selection marker was created (FIG. 11A) and introduced into the nkuAΔ background strain TNO2A3. The velB deletion mutant TNI7.2 was generated by transforming RNI16.1 with the velB deletion construct with pyrG+. RNI18.2 (ΔvelB;veA+) was isolated from a meiotic cross between RRAW16 and TNI7.2. velB and laeA loci were TAP tagged in nkuAΔ background strain TNO2A3 by using clonNat resistance.

Correct gene replacement was confirmed by Southern analyses (FIG. 11A-C). AGB389 (veA+, velB::ctap*) and AGB390 (veA+, laeA::ctap*) strains were obtained from a meiotic cross between TNO2A3 and AGB154. *E. coli* DH5 α and MACH-1 (INVITROGEN) were applied for plasmid DNA and were propagated in LB medium (1% tryptone, 0.5% yeast extract, 1% NaCl) supplemented with 100-150 μ g·ml⁻¹

ampicillin. The bacterial strain KS272 for recombinogenic engineering was propagated in low-salt (0.5% NaCl) LB medium with 25 $\mu g \cdot ml^{-1}$ chloramphenicol. Minimal medium (0.52 g·l^-l KCl, 0.52 g·l^-l MgSO4, 1.52 g·l^-l KH2PO4, 0.1% trace element solution, pH6.5) was used for growth of fungal strains, supplemented with appropriate amounts of 4-aminobenzoic acid (PABA, 1 $\mu g - ml^{-1}$), Biotin (0.02 $\mu g \cdot ml^{-1}$), Uracil (50 $\mu g \cdot ml^{-1}$), Pyridoxine (0.05 $\mu g \cdot ml^{-1}$), nourseothricin-dihydrogen sulfate (100-120 $\mu g \cdot ml^{-1}$) (clonNAT, WERNER BIOAGENTS), pyrithiamine (TAKARA Bio Inc) (0.1 $\mu g \cdot ml^{-1}$); 1% D-glucose was used as the source of carbon together with 10 mM ammonium or nitrate as nitrogen source. For TAP experiments, fungal strains were grown in complete medium (0.5% yeast extract, 1% bacto-peptone, 1% glucose). Sterigmatocystin (ST) production of strains was assayed as described.

Transformation procedures. E. coli and A. nidulans cells were transformed as described.

Plasmid constructions details. The plasmids utilized in this work are listed in Table 2, oligonucleotide sequences are given in Table 3.

TABLE 2

	Plasmid Constructs.
Plasmid	Description & Characteristics
pPTRII	autonomously replicating Aspergillus plasmid [ptrA, AMA1, bla]
pPTRII	Cloning vector for the construction of LexA DNA binding domain
pGAD424	Cloning vector for the construction of GAL4 activation domain
pNJ04	veA ORF in pTLexA
pNJ05	veA ORF in pGAD424
pNJ06	veA N-terminal (1-300a.a.) in pGAD424
pNJ07	veA C-terminal (276-stop.) in pGAD424
pNJ08	veA1 ORF in pTLexA
pNJ09	veA1 ORF in pGAD424
pNJ10	veA1 N-terminal (1-265a.a.) in pGAD424
pNJ11	vos A ORF in pGAD424
pNJ12	vosA-F239 (1-239a,a) in pGAD424
pNJ13	vosA C-terminal (211a.a-stop) in pGAD424 laeA ORF in pTLexA
pNJ14 pNJ15	laeA-F231 (1-231a.a.) in pTLexA
pNJ15 pNJ16	laeA-121R (121a.astop) in pTLexA
pNJ17	laeA ORF in pGAD424
pNJ18	laeA-F231 (1-231a.a.) in pGAD424
pNJ19	laeA-121R (121a.astop) in pGAD424
pNJ20	velB ORF in pTLexA
pNJ21	velB-F231 (1-231 a.a) in pTLexA
pNJ22	velB-142R (142a.astop) in pTLexA
pNJ23	velB ORF in pGAD424
pNJ24	velB-F231 (1-231 a.a) in pGAD424
pNJ25	velB-142R (142a.astop) in pGAD424
pNV1	Dominant resistance cloning plasmid
pME3024	ptrA cassette with SfiI sites in EcoRV site of pBluescript II KS
pME3154	veA C-Terminus::ctap* tag::veA 3' UTR in pGEM5
pME3155	veA 4.6 kb HindIII genomic fragment in pUC19
pME3156	pveA::veA::ctap* tag in pUC19
pME3157	pveA::veA::ctap* tag, ptrA, in pUC19
pME3158	velB deletion cassette [velB::ptrA]
pME3159	5 kb velB genomic locus amplicon in ApaI site of pNV1
pME3160	Expression module niiAt-pniiA/pniaD-niaDt-Af pyrG, bla
pME3161	pniiA::veA cDNA in PmeI site of pME3160
pME3162	pniiA::velB cDNA in PmeI site of pME3160
pME3163	pniiA::velB::sgfp in PmeI site of pME3160
pME3164	pniiA::velB cDNA in PmeI site of pME3166 expression module
pME3165	pniiA::velB::sgfp in PmeI site of pME3166 expression module
pME3166	Expression module 2.6 kb amplicon from pME3160 with primers Sv315/318 in ApaI site of pNV1
pME3167	
	pveA::veA::sgfp, pgpdA::natR in pUC19
pME3168	pveA::veA::sgfp in pUC19
pME3169	pveA::veA::sgfp, ptrA in pUC19
pME3173	pgpdA::intron::mrfp::h2A cDNA in EcoRV and pgpdA::natR in SmaI of pBluescript II KS
pME3178	veA 4.6 kb HindIII genomic fragment in HindIII and ptrA in NotI site of pBluescript II KS
pME3188	pniiA::n-eyfp::veA cDNA in PmeI, and pniaD::c-eyfp::laeA cDNA in SwaI site of pME3160 expression module

TABLE 2-continued

	Plasmid Constructs.
Plasmid	Description & Characteristics
pME3189	pniiA::n-eyfp::veA cDNA in PmeI, and pniaD::c-eyfp::velB cDNA in SwaI site of pME3160 expression module
pME3190	pniiA::laeA::sgfp in PmeI site of pME3166 expression module

pBluescript II KS (STRATAGENE) and pUC19 (FER-MENTAS) were used as cloning plasmids. The plasmid pME3156 containing veA::ctap* tag fusion was constructed by recombineering an 800 by EarI fragment comprising a C-terminal fusion of the TAP* tag (FIG. 2A) to the veA 15 coding sequence derived from pME3154 with NaeI-linearised pME3155 in E. coli. Recombineering is genetic engineering based on homologous recombination in an E. coli host strain expressing phage-derived proteins. In order to amplified with oligonucleotides Sv129/130 from pPTRII (TAKARA) and inserted into the SmaI site of pME3156, and the final construct was used in tandem affinity purification experiments.

The veA::sgfp fusion in pME3168 was created by replac- 25 ing the C-TAP* tag module in pME3154 by an OZG28/29amplified sgfp fragment digested with NcoI/HindIII. pME3168 was digested with SmaI and a blunt ptrA (Sv129/ 130) was inserted resulting in pME3169. The Gpd1/Nat2amplified 1.4 kb pgpdA::natR cassette from pNV1 was 30 cloned into SmaI of pME3168 creating pME3167. To create a velB deletion construct, a 2 kb upstream flanking region was amplified (OZG57/58) and inserted into the EcoRV site of pBluescript KS II (STRATAGENE). The resulting plasmid was then used for insertion of a 2 kb velB downstream flanking region (OZG59/60) into SmaI site, which was then digested with SfiI to insert the SfiI-released ptrA marker from pME3024 generating pME3158, from which a 5.9 kb replacement cassette was used for deletion of velB locus.

For complementation, pME3159 was created by cloning a 40 5 kb velB genomic fragment (OZG99/100) in the ApaI site of pNV1. For overexpression and localization experiments, the nitrogen source-dependent expression module of pME3160 was exploited, which contains the A. nidulans niiA/niaD intergenic region flanked by the corresponding termination 45 regions to allow expression of two genes in a bidirectional orientation at the same time. The veA and velB cDNAs were amplified and cloned into the PmeI site of pME3160 yielding pME3161 veA and pME3162 velB overexpression constructs, respectively.

The velB cDNA::sgfp fusion construct was created by fusion PCR with OZG63/116 for velB and OZG115/29 for sgfp. To create a dominant expression module, the expression

module (niiAT::pniiA/pniaD::niaDT) of pME3160 was amplified with Sv315/318 and cloned into ApaI-digested flushed pNV1 to yield pME3166. The velB cDNA and velB cDNA::sgfp recombinant DNA fragments were cloned into the PmeI site of pME3166. To obtain the pgpdA::mrfp:h2A construct, the gpdA promoter and intron (Sv337/338), mrfp (Sv339/340), h2A cDNA containing terminator (Sv339/340) were amplified. Final products were fused using the double joint PCR procedure. The pgpdA::mrfp:h2A recombinant create pME3157, a ptrA pyrithiamine resistance cassette was $_{20}$ fragment was cloned into the EcoRV site of pBluescript KS II followed by pgpdA::natR cassette insertion into the SmaI site yielding pME3173. The n-eyfp::veA and c-eyfp::laeA fusion constructs were cloned into the PmeI and SwaI sites of the pME3160, respectively.

> For in vivo interaction analyses, n-eyfp (OZG73/74) and veA cDNA (Sv142/143, same as OZG69/70 without restriction sites) were amplified and fused and combined with c-eyfp (OZG75/76) and laeA cDNA (OZG61/62) in plasmid pME3188 and ceyfp (OZG75/77) with velB cDNA (OZG63/ 64) in pME3189, respectively. The appropriate neyfp:: veA, c-eyfp::laeA or c-eyfp::velB fusion constructs were cloned into the PmeI and SwaI sites of pME3160.

> For the construction of the laeA:: sgfp fusion plasmid (pME3190), laeA cDNA (OZG61/162) and sgfp (OZG29/ 161) were amplified, fused and inserted into the PmeI site of the pME3166 expression module under the niiA promoter.

> For construction of the velB and laeA TAP* tag fragments, velB including 400 by of the 5' UTR (OZG210/211) and laeA including 400 by of the 5' UTR (OZG201/202), velB 1.6 kbp 3'UTR (OZG211/100) and laeA 1.6 kbp 3'UTR (OZG204/ 205) were amplified from genomic DNA.

> These fragments were fused to the ctap*::natR module by fusion PCR, which creates the 5'UTR::velB::ctap*::natR:: 3'UTR (OZG223/224) and 5'UTR::laeA::ctap*::natR:: 3'UTR (OZG221/222) gene replacement fragments, respectively.

To confirm protein-protein interaction by a yeast two-hybrid assay, the ORF, N-terminal and C-terminal regions of each gene product were amplified by PCR (Table 3) from an A. nidulans cDNA library provided by Kwang-Yeop Jahng (Chonbuk University, Jeonju, Korea). The PCR product of each gene was digested with EcoRI and SalI or XhoI and cloned into the pTLexA or pGAD424 vector, respectively.

TABLE 3

	Oligonucleotides utilized for plasmid generations.
Designat:	ion Sequence Feature
OZG28	5'-TTT GGC CAT GGG TGG TAG CGG TGG TAT GGT GAG CAA sgfp-GGSGG Spacer GGG CGA GGA GCT G-3' (NcoI) (SEQ ID NO: 1)
OZG29	5'-AAA ATT TAA GCT TCT ACT TGT ACA GTT CGT CCA TGC CGT sgfp 3'end (HindIII) G-3' (SEQ ID NO: 2)

TABLE 3-continued

	Oliqonucleotides utilized for plasmid generation	ıs.
Designation	Sequence	Feature
OZG57	5'-ACT CAC GAA TCC ACG GGA TAC AT-3' (SEQ ID NO: 3)	velB 5'UTR-A
OZG58	5'-GGC CTG AGT GGC CGG GTG GGA TAC GGT CCA TCG AAA-3' (SEQ ID NO: 4)	velB 5'UTR-B (sfiI)
OZG59	5'-GGC CAT CTA GGC CGA CCG TAT ATT GTT TCA TAA ATC CTT-3' (SEQ ID NO: 5)	velB 3'UTR-A (sfiI)
OZG60	5'-TAT GAC CGC GTG AGC AAA TAG GAC-3' (SEQ ID NO: 6)	velB 3'UTR-B
OZG61	5'-ATG TTT GAG ATG GGC CCG GTG GG-3' (SEQ ID NO: 7)	laeA start
OZG62	5'-TTA TCT TAA TGG TTT CCT AGC CTG GT-3' (SEQ ID NO: 8)	laeA stop
OZG63	5'-ATG TAC GCT GTT GAG GAT AGG GC-3' (SEQ ID NO: 9)	velB start
OZG64	5'-TTA GTA TTC GTT ATC CAG ACC ATC G-3' (SEQ ID NO: 10)	velB stop
OZG68	5'-CTC GAG TTA GTA TTC GTT ATC CAG ACC ATC G-3' (SEQ ID NO: 11)	velB start (XhoI)
OZG69	5'-CCA TGG ATG GCT ACA CTT GCA GCA CCA-3' (SEQ ID NO: 12)	veA start (NcoI)
OZG70	5'-CTC GAG TTA ACG CAT GGT GGC AGG CTT TGA GA-3' (SEQ ID NO: 13)	veA stop(XhoI)
OZG73	5'-ATG GTG AGC AAG GGC GAG GAG-3' (SEQ ID NO: 14)	n-eyfp start
OZG74	5'-GGT GGT GCT GCA AGT GTA GCC ATC GTG GCG ATG GAG CGC ATG ATA TAG-3' (SEQ ID NO: 15)	n-eyfp::veA fusion maker
OZG75	5'-ATG GCC GAC AAG CAG AAG AAC-3 (SEQ ID NO: 16)	c-eyfp start
OZG76	5'-ACG AGT TCC CAC CGG GCC CAT CTC AAA CAT GTG GTT CAT GAC CTT CTG TTT CAG-3' (SEQ ID NO: 17)	c-eyfp:laeA fusion maker
OZG77	5'-GGA ATG CGC CCT ATC CTC AAC AGC GTA CAT GTG GTT CAT GAC CTT CTG TTT CAG-3' (SEQ ID NO: 18)	c-eyfp velB
OZG98	5'-TTT GAA TTC ATG CAG CAG CCC AAG CGC GCG AGA G-3' (SEQ ID NO: 19)	veA1 start
OZG99	5'-AAA GGG CCC CGA GAA TGT CCG CCT GAC CCG TGC-3' (SEQ ID NO: 20)	velB complement-A (ApaI)
OZG100	5'-CCA AGT CTG CCC GAC AAG CTC ACT G-3' (SEQ ID NO: 21)	velB complement-B
OZG115	5'-CGC CAC AGC GAC GAG GAC GAT GGT CTG GAT AAC GAA TAC GGT GGT AGC GGT GGT ATG GTG AGC AAG-3' (SEQ ID NO: 22)	velB::sgfp fusion maker
OZG116	5'-GTA TTC GTT ATC CAG ACC ATC GTC-3' (SEQ ID NO: 23)	velB nostop codon
OZG161	5'-CTG CAC ATA TAC CAG GCT AGG AAA CCA TTA AGA GGT GGT AGC GGT GGT ATG GTG AGC-3' (SEQ ID NO: 24)	laeA::sgfp fusion maker
OZG162	5'-TCT TAA TGG TTT CCT AGC CTG GTA-3' (SEQ ID NO: 25)	laeA nostop codon

TABLE 3-continued

	Oliqonucleotides utilized for plasmid qeneration	ns.
Designation	on Sequence	Feature
OZG201	5'-CCT CGC CCT CCT GCA TCA ATA TTC GG-3' (SEQ ID NO: 26)	laeA 5'UTR
OZG202	5'-GAG ACG GCT ATG AAA TTC TTT TTC CAT CTT CTC TTA CCA CCG CTA CCA CCT CTT AAT GGT TTC CTA GCC TGG TAT ATG-3' (SEQ ID NO: 27)	
OZG204	5'-GAG CAG GCG CTC TAC ATG AGC ATG CCC TGC CCC TGA GAG CAA AAG GCG ACC ACA TCC AGG-3' (SEQ ID NO: 28)	laeA 3'UTR-A (fusion maker)
OZG205	5'-TCG TCA ACC GCC TCA GCT GGA ACC-3' (SEQ ID NO: 29)	laeA 3'UTR-B
OZG210	5'-CCT CCT CGC CGC CTC TAG TAC CGT C-3' (SEQ ID NO: 30)	velB 5'UTR
OZG211	5'-GAA ATT CTT TTT CCA TCT TCT CTT ACC ACC GCT ACC ACC GTA TTC GTT ATC CAG ACC ATC GTC C-3' (SEQ ID NO: 31)	velB ctap* fusion maker
OZG212	5'-CGA GCA GGC GCT CTA CAT GAG CAT GCC CTG CCC CTG AAG ACC GTA TAT TGT TTC ATA AAT CC-3' (SEQ ID NO: 32)	velB 3'UTR-A (fusion maker)
OZG221	5'-CGG CTG TTT ACA TTG TGT TTT CTG G-3' (SEQ ID NO: 33)	laeA-NEST-A for fusion
OZG222	5'-CCG TGA AGA ACT TGG CGT TGT AG-3' (SEQ ID NO: 34)	laeA-NEST-B for fusion
OZG223	5'-GGA CCG TCT AAT TCA ACT CAC AG-3' (SEQ ID NO: 35)	velB-NEST-A for fusion
OZG224	5'-CTT CCA GCG GTT ATC CTC CGT TG-3' (SEQ ID NO: 36)	velB-NEST-A for fusion
Sv129	5'-ATC TGA CAG AGC GGC CGC AAT TGA TTA CG-3' (SEQ ID NO: 37)	ptrA-A
Sv130	5'-ATA TAT GCG GCC GCT CTT GCA TCT TTG TTT-3' (SEQ ID NO: 38)	ptrA-B
Sv315	5'-GAT ACC AAA CGG AAC TGG CTG TTA TGG-3' (SEQ ID NO: 39)	expression module A
Sv318	5'-ATC GAC GCA ACC ATC GAA GCA GC-3' (SEQ ID NO: 40)	expression module B
Sv337	5'-GAT CTT TGC CCG GTG TAT GAA ACC-3' (SEQ ID NO: 41)	gpdA promoter A (-432)
Sv338	5'-TCG GAG GAG GCC ATG GTG ATG TCT GCT CAA GC-3' (SEQ ID NO: 42)	gpdA promoter B
Sv339	5'-GAC ATC ACC ATG GCC TCC TCC GAG GAC GTC ATC-3' (SEQ ID NO: 43)	mrfp start
Sv340	5'-GGC TCC AGC GCC TGC ACC AGC TCC GGC GCC GGT GGA GTG GCG GC-3' (SEQ ID NO: 44)	mrfp stop
Sv341	5'-GGA GCT GGT GCA GGC GCT GGA GCC ACT GGC GGC AAA TCT GGT GG-3' (SEQ ID NO: 45)	h2A start
Sv342	5'-ATC TGG AGG GGA CAG GCA GTT TAT-3' (SEQ ID NO: 46)	terminator for h2A
GpdA	5'-GGG TTT CGA ACT ACA TCA AGG GTC CAA GAC CGA CAT CGA GGC TCT GTA CAG TGA CCG GTG-3' (SEQ ID NO: 47)	gpdA promoter
Nat2	5'-AGG GAA TTC TCA GGG GCA GGG CAT GC-3' (SEQ ID NO: 48)	natR stop

TABLE 3-continued

	Oligonucleotides utilized for plasmid generat	ions.
Designati	on Sequence	Feature
OMN131	5'-GAA GGT CGA TGA TGG TGT GAT G-3' (SEQ ID NO: 49)	velB 5' amplify
OMN132	5'-CTA GAG GTA AAG ATC AAG GTA G-3' (SEQ ID NO: 50)	velB 3' amplify
OMN133	5'-CTG ATG GCT GAA TGA AGC ACA G-3' (SEQ ID NO: 51)	velB 5' nested
OMN134	5'-TGC TTT ACG ACG ATA GCC ATG C-3' (SEQ ID NO: 52)	velB 3' nested
OMN135	5'-ggtg aag agc att gtt tga ggca GCG GCC AGT CTT TAG ACA AAT G-3' (SEQ ID NO: 53)	velB 5' rev with pyrG tail (bold)
OMN136	5'-agt gcc tcc tct cag aca gaa ta GGA TAA CGA ATA CTA AAG ACC G-3' (SEQ ID NO: 54)	velB 3' for with pyrG tail (bold)
OMN125	5'-TAT GCA CTG GCA CTC AAG CAA CCG-3' (SEQ ID NO: 55)	velB forward primer for probe
OMN126	5'-GTG CAT GAC GGT CGT ATC TGG TCC-3' (SEQ ID NO: 56)	velB reverse primer for probe
OKH181	5'-GGC TGT AGT CGC TTT GTT-3' (SEQ ID NO: 57)	veA forward primer for probe
OKH182	5'-GCC CAG TGT AAG AAA GGA-3' (SEQ ID NO: 58)	veA reverse primer for probe
OJA242	5'-GCT GTC GAT CTT TGT ACC CTG-3' (SEQ ID NO: 59)	laeA forward primer for probe
OJA243	5'-CGT TCC TGG ATG TGG TCG CCT-3' (SEQ ID NO: 60)	laeA reverse primer for probe
oNK11	5'-ATATAAGCTTAATGGCTACACTTGCAGCACCAC-3' (SEQ ID NO: 61)	veA forward for Y2H
oNK12	5'-ATATGTCGACTTAACGCATGGTGGCAGGCTTTG-3' (SEQ ID NO: 62)	veA reverse for Y2H
oNK13	5'-ATATAAGCTTAATGCAGCAGCCCAAGCGCGCGAG-3' (SEQ ID NO: 63)	veA1 forward for Y2H
oNK14	5'-ATATGAATTCATGAGTGCGGCGAACTATCCAG-3' (SEQ ID NO: 64)	vosA forward for Y2H
oNK15	5'-ATATGTCGACTCACCGAGGAGTTCCGTTCGCTG-3' (SEQ ID NO: 65)	vosA reverse for Y2H
oNK32	5'-ATATGAATTCATGTTTGAGATGGGCCCGGTGGGAAC-3' (SEQ ID NO: 66)	laeA forward for Y2H
oNK33	5'-ATATGTCGACTTATCTTAATGGTTTCCTAGCCTG-3' (SEQ ID NO: 67)	laeA reverse for Y2H
oNK74	5'-ATAT AAGCTT ATCAACGAGCATCAGCACAAAC-3' (SEQ ID NO: 68)	veA C-terminal forward for Y2H
oNK75	5'-ATATGTCGACTCCATATTCCACTGCCGACGGAC-3' (SEQ ID NO: 69)	veA N-terminal reverse for Y2H
oNK76	5'-ATAT GAATTC TCTGATAGGACAGCCATGCAAATC-3' (SEQ ID NO: 70)	vosA C-terminal forward for Y2H
oNK78	5'-ATAT GAATTC ATGTACGCTGTTGAGGATAG-3' (SEQ ID NO: 71)	velB forward for Y2H
oNK79	5'-ATAT GTCGAC TTAGTATTCGTTATCCAGACCA-3' (SEQ ID NO: 72)	velB reverse for Y2H
oNK130	5'-ATATGAATTCACGGTAGCGCGGGTATCGGAG-3' (SEQ ID NO: 73)	laeA-121R forward for Y2H

	Oligonucleotides utilized for plasmid generations.				
Designatio	on Sequence	Feature			
oNK132	5'-ATATGAATTCATGTCTTCATCGTATCCACCAC-3' (SEQ ID NO: 74)	velB-142R forward for Y2H			
oNK138	5'-ATAT CTCGAG ACCAGGCACCGGGACGGAGATG-3' (SEQ ID NO: 75)	laeA-F231 reverse for Y2H			
oNK140	5'-ATAT CTCGAG AGTAGGAATAGTCCCTACTCGTG-3' (SEQ ID NO: 76)	vosA-F239 reverse for Y2H			
oNK141	5'-ATAT CTCGAG TCCAGGCCCTGGAGTAACTGGCTG-3' (SEQ ID NO: 77)	velB-F231 reverse for Y2H			
jwbvelBF	5'-TTCGCTAGACAGCTCATTCTACG-3' (SEQ ID NO: 78)	velB forward primer for probe			
jwbvelBR	5'-TAGTATTCGTTATCCAGACCATCG-3' (SEQ ID NO: 79)	velB reverse primer for probe			
jwbvelAF	5'-ATACCTGGATAAACCAAATCGAGC-3' (SEQ ID NO: 80)	veA forward primer for probe			
jwbvelAR	5'-AGGTTCATTCGCAGGGCTAGAC-3' (SEQ ID NO: 81)	veA reverse primer for probe			
jwblaeAF	5'-ACCACTACAGCTACCACTCTCC-3' (SEQ ID NO: 82)	laeA forward primer for probe			
jwblaeAR	5'-TTTCGATGCTCTCTGAGACGGC-3' (SEQ ID NO: 83)	laeA reverse primer for probe			

Yeast two-hybrid analysis. pTLex (Cho et al., 2003; kindly provided by Suhn-Kee Chae at Paichai University, Daejeon, Korea) derived bait and pGAD424 (CLONTECH) derived prey constructs were cotransformed into the Saccharomyces cerevisiae reporter strain MO and transformants were selected on -UTL -trp, -leu) containing 2% glucose media. To further confirm the interactions of proteins, several transformants of each combination were tested for their coloration on the medium -UTL containing X-Gal, and the transformants were tested for β -galactosidase activity using the yeast β -galactosidase assay kit (PIERCE).

Recombinant DNA procedures, hybridization techniques and analysis of nucleic acids. For recombinant DNA technology, standard protocols were performed. Taq, Pfu (MBI FERMENTAS) and Platinum Taq DNA polymerase (INVITROGEN) were used in PCR reactions, and cloning steps were confirmed by sequencing. Fungal genomic DNA was prepared from ground mycelia, and Southern blot analyses were conducted as described. Total RNA samples were analyzed by Northern hybridization as described. The STRATAGENE Prime-It II kit was used to radioactively label hybridization probes in the presence of $[\alpha\text{-}32P]\text{dATP}.$

To produce autoradiographs, washed membranes were 55 exposed to KODAK X-Omat films. Sequence data were analyzed using the LASERGENE software package from DNASTAR, and alignments were created by the Clustal W method. PEST motifs were analyzed on the web tool and NES patterns were identified on the web tool.

TAP purification. The fungal strains AGB272, AGB273 (veA::ctap*), AGB389 (velB::ctap*) and AGB390 (laeA::ctap*) were grown in liquid culture and transferred onto CMM (minimal medium +0.1% casein hydrolysate) plates, wrapped with parafilm and covered with aluminium foil to 65 induce sexual development or were transferred onto MM and incubated under white fluorescent light without wrapping.

At 48 h post induction of sexual and 24 h post induction of asexual development, the differentiating mycelia were ground in liquid nitrogen to prepare crude extracts in B* buffer (100 mM Tris-HCl pH7.6, 250 mM NaCl, 10% glycerol, 0.05% NP-40, 1 mM EDTA, 2 mM DTT) supplemented with an EDTA-free protease inhibitors mix (ROCHE), phosphatase inhibitors (MERCK) and specified protease inhibitors as recommended in the procedure at the NCRR.

Crude extracts were centrifuged for 20 min at 15000 g and transferred into 50 ml falcon tubes. Protein extracts were incubated for 3 h on a rotator with 300 µl of IgG sepharose 6 Fast Flow (AMERSHAM) at 4° C. After that point, the standard protocol (Step 14) as outlined at the NCRR web site was followed with minor modifications. TEV cleavage was executed under rotation using 350U of AcTEV (INVITROGEN) in the presence of 1 µM E-64 (CALBIOCHEM) protease inhibitor at 4° C. for 5 h; 1 mM PMSF (phenylmethanesulfonylfluoride) was included in the calmoduline binding step on affinity resin (STRATAGENE). The TCA (trichloroacetic acid)-precipitated eluate was loaded onto a 10% polyacrylamide gel and stained with Coomassie Brilliant Blue G (Sigma). Protein bands were cut out and submitted for mass spectrometry.

Immunoblotting. For detection of the VeA::TAP* fusion protein and actin, anti-calmodulin binding peptide antibody (UPSTATE, catalog 07-482) and anti-actin antibody (MP Biomedicals, catalog 69100) were used.

LC-MS/MS Protein Identification. Excised polyacrylamide gel pieces of stained protein bands were digested with trypsin according to Shevchenko et al. Tryptic peptides extracted from each gel slice were injected onto a reversed-phase liquid chromatographic column (Dionex-NAN75-15-03-C18 PM) by using the ultimate HPLC system (Dionex, Amsterdam, Netherlands) to further reduce sample complexity prior to mass analyses with an LCQ DecaXP mass spec-

trometer (ThermoElectron Corp, San Jose, Calif.) equipped with a nanoelectrospray ion source. Cycles of MS spectra with m/z ratios of peptides and four data-dependent MS2 spectra were recorded by mass spectrometry.

The "peak list" was created with extracts provided by the Xcalibur software package (BioworksBrowser 3.1). The MS2 spectra with a total ion current higher than 10,000 were used to search for matches against a public *A. nidulans* genomewide protein sequence database of the BROAD INSTITUTE (9542 sequences, December 2005, plus 180 sequences of the most commonly appearing contaminants, e.g., keratins and proteases, provided with the BioworksBrowser package) using the TurboSEQUEST algorithm of the Bioworks software (Version 3.1, Thermo Electron Corp).

The search parameters included based on the TurboSE-QUEST algorithm were: (i) precursor ion mass tolerance less than 1.4 amu, (ii) fragment ion mass tolerance less than 1.0 amu, (iii) up to three missed tryptic cleavages allowed, and (iv) fixed cysteine modification by carboxyamidomethylation

(plus 57.05 amu) and variable modification by methionine oxidation (plus 15.99 amu) and phosphorylation of serine, threonine, or thyrosine (plus 79.97 amu).

In accordance with the criteria described by Link et al., matched peptide sequences of identified proteins had to pass the following: (i) the cross-correlation scores (Xcorr) of matches must be greater than 2.0, 2.5, and 3.0 for peptide ions of charge state 1, 2, and 3, respectively, (ii) Δ Cn values of the best peptide matches must be at least 0.4, and (iii) the primary scores (Sp) must be at least 600.

Protein identification required at least two different peptides matching these criteria. The degree of completeness of the b- and y-ion series for each SEQUEST result was manually checked for every protein identified. Peptides of identified proteins were individually blasted against the NCBI database to ensure their unambiguous assignment to the TurboSEQUEST-specified protein. See also the *Multiple Consensus Reports* for the detailed TurboSEQUEST identifications in the Table 4. The three top scoring peptides are listed for all identifications.

TABLE 4

Mass Spectrometi	Mass Spectrometry Data of Protein Identifications.						
				Delta	ι		
Peptide Sequence	MH+	Charge	:XCorr	Cn	Sp	RS	pIons
AN1052.2 (hypothetical protein					1210.	3	121(121-0-0-0-0)
similar to velvet A)1 ¹ R.LEVISNPFIVYSAK.K (SEQ ID NO: 84)	1580.85	2	5.27	0.59	1188.	8 1	21/26
R.LEVISNPFIVYSAK.K	1425.59	2	4.52	0.32	1075.	5 1	18/24
(SEQ ID NO: 85) R.LEVISNPFIVYSAK.K (SEQ ID NO: 86)	2135.24	3	6.84	0.66	2085.	3 1	37/72
AN0363.2 (hypothetical protein) ² K.IGVWFVLQDLSVR.T (SEQ ID NO: 87)	1532.81	2	5.18	0.37	1508. 1490.		151(150-1-0-0-0) 18/24
K.SVSDLPQSDIAEVINK.G (SEQ ID NO: 88)	1715.88	2	5.35	0.55	803.	4 1	24/30
R.IWSLQVVQQPIR.A (SEQ ID NO: 89)	1467.74	2	4.62	0.35	1568.	4 1	17/22
AN0807.2 (hypothetical protein) ³ K.EIHAYNILHIYQAR.K (SEQ ID NO: 90)	1741.98	2	4.85	0.61	186. 1789.		19 (18-0-2-0-0) 20/26
R.YAVAGGPAPWNR.N	1259.40	2	4.70	0.52	1632.	0 1	19/22
(SEQ ID NO: 91) R.VSESLIYAPHPINGR.F (SEQ ID NO: 92)	1641.81	2	4.17	0.57	930.	5 1	20/28
AN2142.2 (hypothetical protein similar to AF465210_1 karyopherin alpha) ⁴					660.	3	66 (66-0-0-0-0)
K.IIQVALDGLENILK.V	1539.84	2	5.09	0.57	2402.	5 1	22/26
(SEQ ID NO: 93) K.IQAVIEAGIPR.R	1167.38	2	3.87	0.44	1618.	2 1	18/20
(SEQ ID NO: 94) K.TPQPDWNTIAPALPVLAK.L (SEQ ID NO: 95)	1933.24	2	4.40	0.64	942.	9 1	19/34
AN0363.2 (hypothetical protein) ⁵ K.GTAPILASTFSEPFQVFSAK.K (SEO ID NO: 96)	2099.37	2	5.28	0.47	220. 1045.		22 (22-0-0-0-0) 22/38
K. IGVWFVLQDLSVR.T	1532.81	2	4.30	0.43	1499.	0 1	18/24
(SEQ ID NO: 97) K.SVSDLPQSDIAEVINK.G (SEQ ID NO: 98)	1715.88	2	5.34	0.48	846.	1 1	24/30
AN1052.2 (hypothetical protein similar to velvet A) ⁶					778.	4	78 (77-1-0-0-0)
K.DATEGTQPMPSPVPGK.L (SEO ID NO: 99)	1612.79	2	3.70	0.45	516.	1 1	18/30
K.KFPGLTTSTPISR.M (SEQ ID NO: 100)	1405.62	2	2.61	0.53	519.	5 1	15/24

TABLE 4-continued

Mass Spectrometi	Mass Spectrometry Data of Protein Identifications.						
				Delta	ı		
Peptide Sequence	MH+	Charg	e XCorr	Cn	Sp	RS:	pIons
K.LMTNQGSPVLTGVPVAGVAYLDKPNR.A (SEQ ID NO: 101)	2699.12	2	6.16	0.61	737.2	1	35/100
AN0807.2 (hypothetical protein) ⁷ K.EIHAYNILHIYQAR.K	1741.97	2	3.01	0.59	454.3 960.3		46(45-0-0-1-0) 17/26
(SEQ ID NO: 102) R.IQQLAADVK.S (SEQ ID NO: 103)	986.15	2	3.12	0.33	1015.4	1	15/16
R.YAVAGGPAPWNR.N (SEQ ID NO: 104)	1259.40	2	3.39	0.58	1119.0	1	17/22
AN1959.2 (hypothetical protein) ⁸ K.DVDNTDGGFFVWGDLSIK.V (SEQ ID NO: 105)	1986.13	2	4.21	0.56	660.3 931.0		66(66-0-0-0-0) 19/34
RLKDVDNTDGGFFVWGDLSIK.V (SEQ ID NO: 106)	2227.46	2	4.14	0.56	1200.5	1	21/38
AN0807.2 (hypothetical protein) ⁹ K.EIHAYNILHIYQAR.K (SEQ ID NO: 107)	1741.97	2	4.15	0.58	220.2 1819.4		22(22-0-0-0) 20/26
R.WYNLAVSESIENLSLAPFSR.V (SEQ ID NO: 108)	2297.55	2	4.30	0.54	1495.0	1	22/38
R.YAVAGGPAPWNR.N (SEQ ID NO: 109)	1259.40	2	3.90	0.54	1681.7	1	20/22
AN0363.2 (hypothetical protein) 10 K.GTAPILASTFSEPFQVFSAK.K (SEQ ID NO: 110)	2099.37	3	5.29	0.56	778.4 1021.1		78(77-1-0-0-0) 30/76
K.SVSDLPQSDIAEVINK.G (SEQ ID NO: 111)	1715.88	2	5.59	0.51	799.5	1	21/30
R.IWSLQVVQQPIR.A (SEQ ID NO: 112)	1467.74	2	4.29	0.45	2227.8	1	18/22
AN1052.2 (hypothetical protein similar to velvet A) ¹¹					454.3		46 (45-0-0-1-0)
R.LEVISNPFIVYSAK.K (SEQ ID NO: 113)	1580.85	2	4.44	0.55	1269.4	1	21/26
R.RPDQYAGSDAYANAPERPR.S (SEQ ID NO: 114)	2135.24	3	5.25	0.65	1825.0	1	36/72
R.RPSAVEYGOPIAOPYOR.P (SEQ ID NO: 115)	1961.17	3	3.99	0.56	1044.4	1	29/64
¹ Avg. Mass: 63831.1; pI: 9.43; Coverage ² Avg. Mass: 37062.4; pI: 5.97; Coverage ³ Avg. Mass: 41578.2; pI: 5.93; Coverage ⁴ Avg. Mass: 60627.4; pI: 5.00; Coverage ⁵ Avg. Mass: 37062.4; pI: 5.97; Coverage ⁶ Avg. Mass: 41578.2; pI: 5.93; Coverage ⁸ Avg. Mass: 41578.2; pI: 5.93; Coverage ⁹ Avg. Mass: 41578.2; pI: 5.93; Coverage ⁹ Avg. Mass: 41578.2; pI: 5.93; Coverage ⁹ Avg. Mass: 37062.4; pI: 5.97; Coverage ¹⁰ Avg. Mass: 37062.4; pI: 5.97; Coverage	(amino ac (amino ac (amino ac (amino ac (amino ac (amino ac (amino ac (amino ac	cids): :	30.4% 28.6% 11% 33.3% 40.6% 18.0% 31.5% 36.6%				

Fluorescence microscopy. A. nidulans spores (5.5×10⁵) were inoculated either on 18 mm×18 mm cover slips submerged in appropriately supplemented liquid medium or on large glass slides covered with a thin layer of medium and incubated at 30° C. overnight. The effect of illumination on localization of VeA and VelB was investigated by growing selected strains in darkness and light on the agar surface or in the submerged culture. Cover slips were mounted on microscope slides using spore storage solution (0.002% Tween, 0.5% NaCl) and fixed with wax.

¹¹Avg. Mass: 63831.1; pI: 9.43; Coverage (amino acids): 38.9%

Fluorescence photographs were taken with a ZEISS Axiovert S100 microscope supported with a HAMAMATSU OCRA-ER digital camera, using the Openlab TM V5.0.1 software package (IMPROVISION, Coventry, UK). For the quantification of the GFP signals, nuclei were defined as ROIs (Area of interest). Pixel intensity within the defined ROIs were analysed by using Openlab tmV5.0.1 software package

(IMPROVISION, Coventry, UK). Nuclei were verified by overlaying the GFP and Ds Red signals. Subcellular distribution was observed with a 100× objective using 495 and 558 nm extinction and emission filters. No autofluorescence was observed. All images were taken using the same exposure and microscope settings.

Sterigmatocystin extraction and thin layer chromatography (TLC) analysis. Samples (1.6 cm diameter disc with fungal samples and agar together) were collected after asexual developmental induction. The fungal samples were ground in 3 ml ddH₂O in a homogenizer, and then 3 ml chloroform was added to extract ST from the aqueous phase. About 1.8 ml chloroform containing ST was collected after centrifugation, and air-dried. The dried extracts were resuspended in 50 μl of chloroform, and 10 μl were separated in hexane:ethyl acetate (4:1) or chloroform:acetone (4:1) on TLC plates. ImageQuant TL (Amersham Biosciences Co.)

was used for ST densitometry. Data are presented as graphs with bars which stand for mean+/-standard error (FIG. 2D). For statistical analysis, data were analyzed using the JMP software package (version 3.2.6, SAS Institute, Inc, Cary, N.C.).

According to the Tukey-Kramer multiple comparison test at P \leq 0.05, the three mean values for WT in the dark are significantly different from WT in the light and velB Δ in the dark after 48 hours (FIG. 5D). The graphs without bars do not produce ST above background noise (indicated by "B").

Example 2

In this example, the inventors created several A. flavus isogenic mutants differing only in copy number of veA and

42

are restriction sites (EcoRI or SpeI). ΔlaeA (SEQ ID NO: 144) was generated by transformation with PLRM5, and contained *A. fumigatus* pyrG (nucleotides 1535-3547 in SEQ ID NO: 144), *A. flavus* 5' (nucleotides 46-1531 in SEQ ID NO: 144) and 3'flank (nucleotides 3554-4933 in SEQ ID NO: 144); nucleotides 43-45, 1532-1534, 3548-3553, and 4934 in SEQ ID NO: 144 are restriction sites. MClaeA (SEQ ID NO: 145) was generated by transformation with pLRM11, and contained *A. parasiticus* niaD (nucleotides 2-5128 in SEQ ID NO: 145):: *A. flavus* laeA (nucleotides 5134-9452 in SEQ ID NO: 145) in Invitrogen pCR bluntII TOPO plasmid (pLRM9); nucleotides 1, and 5129-5133 in SEQ ID NO: 145 are restriction sites. MCveA-laeA was prepared by co-transformation with pSA2.8 and PLRM11 plasmids using *A. flavus* laeA (SEQ ID NO: 146).

TABLE 5

Aspergillus flavus strains.						
Strain	Genotype*	Source				
NRRL 3357	Wild type	Horowitz Brownet al. 2008. Appl. Environ. Microbiol. 74: 5674-5685.				
NRRL 3357.5	pyrG ⁻	Horowitz Brownet al. 2008. Appl. Environ. Microbiol. 74: 5674-5685.				
TSA 1.54 (ΔveA)	pyrG [−] ∆veA::AfpyrG	This study				
TSA 2.46 (MCveA)	pyrG ⁻ AfpyrG veA	This study				
TJW 71.1(ΔlaeA)	pyrG ⁻ ΔlaeA::AfpyrG	Kale, et al 2008. Fungal Genet. Biol. 45: 1422-1429.				
TJW 79.13	pyrG⁻∆laeA::AfpyrG	Kale, et al 2008. Fungal Genet.				
(MClaeA)	niaD ⁻ niaD laeA	Biol. 45: 1422-1429.				
TSA 2.8 (MCveA-laeA)	pyrG ⁻ AfpyrG veA niaD laeA	Horowitz Brownet al. 2008. Appl. Environ. Microbiol. 74: 5674-5685.				

^{*}Af, A. fumigatus

laeA genes, including ΔveA , $\Delta laeA$, multicopy laeA (MClaeA), and MCveA strains and a double MC strain (MCveA-laeA). The respective VeA and LaeA mutants exhibited critical differences in cell density responses and invasion of host tissues, despite gross similarities between sclerotial and aflatoxin production.

Considering the interdependence of oxylipin function with VeA coupled with the VeA-LaeA interaction, we postulated that VeA mutants would also be impaired in seed pathogenesis in a manner similar to that of LaeA mutants and, furthermore, that both mutants could be affected in density-dependent development. To explore these hypotheses, we created several *A. flavus* isogenic mutants differing only in copy number of veA and laeA genes, including ΔveA, ΔlaeA, multicopy laeA (MClaeA), and MCveA strains and a double MC strain (MCveA-laeA). The respective VeA and LaeA 50 mutants exhibited critical differences in cell density responses and invasion of host tissues, despite gross similarities between sclerotial and aflatoxin production.

Fungal strains and growth conditions. The *Aspergillus flavus* strains used and generated in this example are listed in 55 Table 5. ΔveA (SEQ ID NO: 142); containing 5' veA flanking region (nucleotides 1-1314 in SEQ ID NO: 142), *A. fumigatus* pyrG (nucleotides 1315-3264 in SEQ ID NO: 142), and 3' flanking region of the veA open reading frame (nucleotides 3265-4556 in SEQ ID NO: 142). MCveA (SEQ ID NO: 143) 60 was generated by transformation with pSA3.X, and contained *A. flavus* veA (nucleotides 1368-3156 in SEQ ID NO: 143):: *A. fumigatus* pyrG (nucleotides 3700-5681 in SEQ ID NO: 143) in TOPO-TA cloning plasmid (TOPO-TA pCR2.1); *A.flavus* 5' (nucleotides 259-1367 in SEQ ID NO: 143) and 3'flank (nucleotides 3157-3674 in SEQ ID NO: 143); nucleotides 258, 3675-3679, 3699, 5682-5686 in SEQ ID NO: 143

All strains were maintained as stocks in glycerol and grown at 29° C. on glucose minimal medium (GMM) (36) amended with appropriate supplements for spore production.

Fusion PCR and vector construction. All primers used in this example are listed in Table 6.

TABLE 6

	Primer sequences.
Primer	Sequence (5'-3')
5'F veA For	ACAACCCTGGACTCTGGAAT (SEQ ID NO: 118)
5'F veA Rev	CGAAGAGGGTGAAGAGCATTGTTTGAGGCA GAGGACGCGTTGACTGTGATG (SEQ ID NO: 119)
3'F veA For	TGACGACAATACCTCCCGACGATACC TGGGTTGATTCCTGCTTTTCCTCC (SEQ ID NO: 120)
3'F veA Rev	TCTCGTTCTCCCATTTACCT (SEQ ID NO: 121)
A. fumigatus pyrG For	TGCCTCAAACAATGCTCTTC (SEQ ID NO: 122)
A. fumigatus pyrG Rev	CAAGGTATCGTCGGGAGGT (SEQ ID NO: 123)
Nested For	AATCACGGACCTCGAAGCAG (SEQ ID NO: 124)
Nested Rev	GGGGTCTTGATATGGCGAAT (SEQ ID NO: 125)

TABLE 0-Concluded							
Primer sequences.							
Primer	Sequence (5'-3')						
Int veA For	CAACAAGACCGACATCACCTTC (SEQ ID NO: 126)						
Int veA Rev	CCATTCTTGGGATAGCTGCAAC (SEQ ID NO: 127)						
MC veA For	CAACGA ACTAGT CCGCCTGCCCTTAACCT CCA						
	(SEQ ID NO: 128)						
MC veA Rev	GCATAC ACTAGT CTCGCATGCCAGTGGAT GGG						
	(SEQ ID NO: 129)						
veA-pyrG Rev	CATCGGTTGACTACGCTCGCA						
11	(SEQ ID NO: 130)						
laeA-niaD For	GACCTGTGGTGAAACCTGAGG						
	(SEQ ID NO: 131)						
veA Northern For	CTAGCTGGTCATTATTTGATCTCG						
	(SEQ ID NO: 132)						
veA Northern Rev	GTTGTAGAGTGGACGATCATCATG						
	(SEQ ID NO: 133)						
laeA Northern For	CCTTGTATGATGTATGTATGATGAGC						
	(SEQ ID NO: 134)						
laeA Northern Rev	GACAGCGAAAGTGAAGAGGACATC						
	(SEQ ID NO: 135)						
actin Northern For	GAAGCGGTCTGAATCTCCTG						
	(SEQ ID NO: 136)						
actin Nothern Rev	ACAGTCCAAGCGTGGTATCC (SEQ ID NO: 137)						
aflR Northern For	AGAGTCTTCCTTCAGCCAGGTC (SEQ ID NO: 138)						
aflR Northern Rev.	GTGGGGCTTTTCTTCATTCTCG (SEQ ID NO: 139)						

 $[\]boldsymbol{\star}$ Bold characters flag restriction enzyme (SpeI) site.

The veA replacement PCR products were constructed using fusion PCR following Szewczyk et al. Starting with wild type A. flavus veA (SEQ ID NO: 141, containing 1314 bp 45 of the 5' flanking region and 1292 by of the 3' flanking region of the veA open reading frame), the 1.3-kb fragments upstream and downstream of the veA coding region were amplified by PCR with primers 5'F veA For and Rev for the upstream fragment and primers 3'F veA For and Rev for the 50 downstream fragment, using NRRL 3357 (prototroph) genomic DNA as a template. Next, a 1.9-kb fragment of the pyrG auxotrophy marker gene was amplified from A. fumigatus AF293 genomic DNA using primers A. fumigatus pyrG For and Rev. These three amplified PCR products were 55 cleaned with a QIAquick gel extraction kit (Qiagen), quantified, and fused using published procedures. The PCR product was amplified with primers Nested For and Rev. All PCR steps were performed using an Expand long template PCR system (Roche Diagnostics GmbH, Mannheim, Germany) 60 according to the manufacturer's instructions.

The final construct was confirmed with endonuclease digestion and PCR using primers Int veA For and Rev for internal veA and primers *A. fumigatus* pyrG For and Rev for pyrG. The veA complementation vector was constructed in 65 two steps. First, the 1.9-kb *A. fumigatus* pyrG PCR fragment was amplified and ligated into the pCR2.1-TOPO vector (In-

44

vitrogen) to create pSA2.4. Next, a 4.4-kb SpeI fragment containing the *A. flavus* veA gene was amplified from *A. flavus* NRRL 3357 genomic DNA with primers MC veA For and Rev and ligated into the SpeI site of pSA2.4 to create the veA complementation vector, pSA3.13. The vector was confirmed by PCR with primers MC veA For and veA-pyrG Rev and endonuclease digestion.

Fungal transformation procedure and mutant confirmation.

For fungal transformation, protoplasts were produced from freshly germinated conidia of NRRL 3357.5 (pyrG auxotroph) and transformed using a polyethylene glycol method. The final fusion PCR product (5 µg) was used for replacement of veA with pyrG after gel purification using a QIAquick gel extraction kit (Qiagen) to create strain TSA 1.54 (SEQ ID NO: 142; containing 1314 bp of the 5' veA flanking region, 1950 bp of *A. fumigatus* pyrG, and 1292, by of the 3' flanking region of the veA open reading frame). The veA::pyrG vector, pSA3.13, was used alone or else cotransformed with pLRM11.1, a vector containing both laeA and niaD, to create MC strains with multiple copies of veA alone and MC strains with multiple copies of both veA and laeA (TSA 2.46 and TSA 2.8, respectively, were used for these studies).

Correct transformants were identified by analyzing genomic DNA using PCR screens followed by Southern analyses. Primers Int veA For and Rev, Nested For and Rev (4.3 kb for the wild type and 4.6 kb for transformants), and *A. fumigatus* pyrG For and Rev were used to identify pyrG replacement of veA. MC transformants were identified by PCR with primers MC veA For and veA-pyrG Rev and primers laeA-niaD For and laeA Northern Rev. Southern analysis was performed for each PCR-identified transformant to confirm single gene replacement of veA in TSA 1.54, at least 2 copies of veA in TSA 2.46, and at least 2 copies of veA and laeA in TSA 2.8. Probes were created with primers Nested For and Rev for the veA open reading frame (ORF) and primers laeA Northern For and Rev for the laeA ORF.

Northern analysis. To examine the expression of veA and laeA transcripts, Northern analysis was performed. Fiftymilliliter amounts of liquid GMM were inoculated with 10⁶ spores/ml of appropriate strains and incubated with shaking at 250 rpm at 29° C. under dark conditions. After 48 h, the mycelium was collected and total RNA was extracted by using the Trizol method (Invitrogen). Blots were hybridized with a veA fragment amplified using the primers Northern For and Rev, an laeA fragment amplified using the primers Northern For and Rev, an actin fragment amplified using the primers actin Northern For and Rev, and an aflR fragment amplified using the primers aflR Northern For and Rev from NRRL3357 genomic DNA. Detection of signals was carried out with a Phosphorimager-SI (Molecular Dynamics).

Physiological experiments. Conidial production, sclerotial formation, and colony diameter were measured for fungal strains following the methods of Horowitz Brown et al. Briefly, 8-ml amounts of 1.6% GMM plus 2% sorbitol agar were overlaid with 3-ml amounts of 0.7% agar GMM plus 2% sorbitol agar containing 10^2 , 10^4 , and 10^6 spores/plate of each A. flavus strain for culture. For conidial counts, three 1.5-cm plugs from each plate were homogenized in 5 ml of 0.01% Tween 80 (vol/vol) water, diluted to 1 x, and counted with a hematocytometer. To visualize sclerotium formation, plates were sprayed with 70% ethanol to kill and wash away conidia. The exposed sclerotia were then collected, lyophilized, and weighed (dry weight per plate). Growth diameter was measured following a point inoculation of 5 µl of 10⁶ spores/ml for each strain on 30 ml of 1.6% GMM. Cultures were grown at 29° C. under continuous dark or light conditions for 3 days

(conidia production), 7 days (sclerotia formation), and 3 and 6 days (colony diameter). Each treatment was replicated four times.

To assay for growth on different fatty acids, the wild-type, $\Delta laeA,$ and ΔveA strains were examined for growth on (i) 20 mM hexanoic acid (6 C), 6 mM oleic acid (18 C), and 4.9 mM erucic acid (22 C) as the sole carbon source, with the fatty acids substituting for the glucose in GMM, or (ii) GMM supplemented with these same molarities of fatty acids, following the method of Maggio-Hall and Keller. Growth diameter was measured following a point inoculation of 5 μl of 10^6 spores/ml for each strain on 30 ml of medium. Each treatment was replicated four times. The experiment was repeated twice

Seed infections. For seed/fungal studies, two cultivars (SunRunner and Flo-Runner) of peanut (Arachis hypogaea) and one (Northup King N33-P3) of non-fungicide treatment maize (Zea mays L.) were used. All the steps were aseptically performed as described by Kale et al. Briefly, mature peanuts 20 (20 peanut cotyledons) and maize (10 seeds) were surface sterilized and inoculated with suspensions of 10⁵ spores/ml of each respective strain, as well as with a water control (mock inoculation). Seeds were placed in 50-ml Falcon tubes containing either sterile water or the spore suspensions and 25 shaken for 30 min in a rotary shaker at 50 rpm, after which they were placed in a high-humidity chamber. Peanut cotyledons were incubated for 3 days for peanut cultivar SunRunner or 5 days for cultivar FloRunnner at 29° C. under dark conditions, and maize kernels for 3 days. All seed experiments 30 were repeated three times.

Histological study. Infected and control peanut cotyledons of cultivar Sun-Runner were collected after 3 days of inoculation and sliced with a razor blade into 2-cm pieces which were immersed in ice-cold fixative FAA (3.7% formalde- 35 hyde, 5% acetic acid, 47.5% ethanol in water) in vials with vacuum pressure for 30 min. Tissues were then removed, incubated with fresh FAA overnight, dehydrated through a tert-butanol series following the method of Cseke et al., and embedded in paraffin (Paraplast Plus). Paraffin blocks were 40 sectioned in 10-µm slices, and serial sections were placed on glass slides and incubated at 37° C. at least overnight, until tissues adhered to the slides. Dewaxing of tissues and staining with Gomori methenamine-silver were performed in the University of Wisconsin-Madison School of Veterinary Medi- 45 cine histology services laboratory. For lipid staining in peanut tissues, Nile red was applied to tissues following the method of Tsitsigiannis et al. A tetramethyl rhodamine 5-isothiocyanate filter in a fluorescent microscope (Olympus BX-60 with 546-nm excitation and 585-nm emission filters) was used to 50 observe Nile red-stained tissues.

Aflatoxin extraction from medium. Eight-milliliter amounts of 1.6% GMM-2% sorbitol agar were overlaid with 3-ml amounts of 0.7% GMM agar plus 2% sorbitol agar containing 10², 10⁴, and 10⁶ spores/plate of each fungal 55 strain. Cultures were grown for 3 days at 29° C. under dark or light conditions. Three 1.5-cm plugs from each plate were homogenized in 3 ml of 0.01% Tween 80 (vol/vol) water and vortexed vigorously for 1 min. One milliliter of chloroform was added, and the sample vortexed and incubated at room 60 temperature for 30 min. The mixture was vortexed again and then centrifuged for 15 min. The lower layer was collected, allowed to dry for 3 days, and then resuspended in 100 µl of chloroform, and 40 µl of the suspension was spotted onto TLC plates (Whatman, Maidstone, England) using a chloroform/ acetone (95:5, vol/vol) solvent system. Each treatment was repeated three times.

46

Aflatoxin extraction from seed. Peanut cotyledons and maize kernels inoculated as described above were collected in 50-ml Falcon tubes with the addition of 5 ml of 0.01% Tween 80 and vortexed vigorously for 1 min. One milliliter was removed from each sample for conidium counting prior to aflatoxin extraction. Five milliliters of acetone was then added to the samples, followed by shaking for 10 min in a rotary shaker at 150 rpm. Samples were allowed to stand for 5 min at room temperature, and then 5 ml of chloroform was added to each sample, followed by shaking for 10 min at 150 rpm. Samples were allowed to stand for an additional 10 min at room temperature, vortexed briefly, and centrifuged for 15 min at 2,000 rpm to collect the organic lower phase. This phase was placed in a new tube and then dried completely for 15 3 days. Five milliliters of 0.1 M NaCl methanol/water (55:45) and 2.5 ml of hexane were added to each tube, and the mixture vortexed vigorously at high speed for 1 min. Samples were centrifuged at 2,000 rpm for 5 min. The hexane layer was collected, the remaining aqueous phase was washed with 2.5 ml of hexane, and then the collection process repeated as described above. The hexane extracts were combined, allowed to dry, and then resuspended in 500 µl of chloroform, and 10 µl of each extract was separated on a silica gel TLC plate using the chloroform/acetone (95:5 vol/vol) solvent system. Each treatment was repeated three times.

Statistical analysis. Statistical differences were analyzed using the JMP software package, version 3.2.6 (SAS Institute, Inc., Cary, N.C.). Multiple comparisons of results for all strains were calculated for growth diameter, lipase activity, and sporulation on seed. To assess the density-dependent development of each strain, sclerotial and conidial numbers were compared at three population levels. Statistically significant mean values, indicated with different letters in the figures, are significant at P<0.05.

Results. Creation of veA and laeA mutant strains in A. flavus. This study required creating near-isogenic strains varying in the number of laeA and veA alleles in the same A. flavus isolate. As Δ laeA and MC strains of the genome-sequenced strain A. flavus 3357 already existed, the first goal was to obtain near-isogenic strains of A. flavus 3357 with loss of or overexpression of veA.

The sequence of the A. flavus 3357 veA ortholog was obtained by designing primers from the A. flavus ATCC MYA384 veA gene (GenBank DQ296645, SEQ ID NO: 140). The sequences of the two genes were found to be 99% identical. All primers and probes in this study were designed from this sequence (Table 6). FIG. 13A shows the strategy of replacement of veA with A. fumigatus pyrG. Transformants were first screened for loss of production of sclerotia on GMM plus 2% sorbitol medium, a phenotype associated with the A. flavus ATCC MYA384 AveA mutant. Several asclerotial A. flavus 3357 transformants were identified and their DNA extracted and analyzed by PCR and Southern analysis. Seventeen out of 100 transformants were found to contain the 4.6-kb and 4.3-kb fragments expected of KpnI (FIG. 13B) and SapI (data not shown) digests, respectively, as expected for a veA replacement with A. fumigatus pyrG. One of these strains, TSA 1.54, was chosen for further studies (FIG. 13B). A strain with at least two copies of veA was obtained by transforming NRRL 3357.5 with plasmid pSA3.13. Several strains were obtained, as determined by Southern analysis, and one, the MCveA strain TSA 2.46, was chosen for further studies (FIG. 13B). Next, a strain with at least two copies of both veA and laeA was obtained by transforming NRRL 3357.5 with plasmids pSA3.13 and pLRM11.1. One of these transformants, the MCveA-laeA strain TSA2.8, was chosen for further studies (FIG. 13B).

The strains with the six genotypes (the wild type and five mutants) exhibited clear differences in development and morphology, as described below, and additionally, the AlaeA strain showed a statistically significant inhibition in growth diameter compared to the growth of most other strains under 5 both light and dark conditions. Conversely, the MClaeA strain's growth diameter was greater than the growth diameters of most other strains in both light and dark regimes (FIG. 14)

veA and laeA affect each other's transcription. Kale et al. 10 recently found that laeA expression negatively affects transcription of veA in A. flavus; this result was replicated in our work (FIG. 15). We also found evidence for veA regulation of laeA expression. Although Northern analysis revealed that the Δ veA strain did not show an increase of laeA expression, 15 the MCveA strain had decreased laeA expression compared to that of the wild type. The MCveA-laeA strain showed relatively high levels of expression of both veA and laeA but not as high as the individual MC strains. We also examined the expression of the aflatoxin-specific transcription factor 20 afIR in all strains. As expected and as previously described, there was no affR expression in ΔveA and $\Delta laeA$ strains. Similarly to the MClaeA strain, both the MCveA and MCveA-laeA strain showed higher levels of aflR expression than the wild type with this treatment.

Conidial and sclerotial density-dependent production is affected by VeA and LaeA. A recent study has shown that conidial and sclerotial production is density dependent in *A. flavus*, for which low cell densities resulted in high sclerotial formation and high cell densities in low sclerotial formation, 30 with an inverse effect on conidial production. This quorum-like signaling system regulating the sclerotial-to-conidial shift was impaired in oxylipin-generating oxygenase mutants. Because VeA has been shown to be important in oxylipin signaling responses and forms a complex with LaeA 35 in the nucleus, we now show that changes in veA and laeA expression could affect the density-dependent sclerotial-to-conidial shift.

The relative abilities of the wild type and the veA and laeA mutants to form sclerotia and conidia were determined by 40 inoculating 10², 10⁴, and 10⁶ conidia onto GMM plus 2% sorbitol plates which were placed in constant dark at 29° C. for 3 (conidia) and 7 (sclerotia) days. Similar to prior results, sclerotial production diminished and conidial production increased in the wild type with increasing cell population 45 levels (FIGS. 16A and B). The veA and laeA null mutants were incapable of producing sclerotia at any population level and yielded relatively constant levels of conidial production regardless of population levels (FIGS. 16A and B).

However, clear differences between effects of loss of or 50 overexpression (MC) of veA compared to the results for cognate laeA mutants emerged in both conidial and sclerotial development. Previous studies have suggested a "balance" in sclerotial and conidial production, i.e., when sclerotial production is low, conidial is high and vice versa. This appeared 55 to hold true for the ΔlaeA strain (no sclerotial production at any cell density and high conidial counts at all densities) but not the Δ veA strain, for which conidial counts were very low at all population levels (FIG. 16A) despite the lack of sclerotial production (FIG. 16B). The MC mutants also showed clear differences in their density-dependent responses. The MCveA strain still exhibited a density-dependent response in sclerotial production with declining numbers in both light and dark regimes at high population levels (FIG. 16B). This was in contrast to the MClaeA strain, which maintained constant 65 sclerotial numbers at all population levels (FIG. 16B). The MCveA-laeA double mutant exhibited an intermediate

48

response. The trend to increased conidial numbers at high population levels was maintained in the MCveA and MCveA-laeA strains but not in the MClaeA strain (FIG. **16**A). These results are summarized in Table 7.

TABLE 7

Summary of density-dependent phenomena in A. flavus	
mutants, morphological differentiations under indicated conditions.	

Mutation	Light-Conidia	Light- Sclerotia	Dark-Conidia	Dark- Sclerotia
None (WT)	+	+	+	+
ΔveA	±	_	±	-
Δ lae A	-	-	-	-
MCveA	±	+	±	+
MClaeA	-	-	-	-
MCveA-laeA	+	±	+	±

- + indicates the presence of density-dependent development.
- ± indicates an intermediate response
- indicates the absence of density-dependent development

Density-dependent production of aflatoxin is controlled by LaeA. We also examined the strains for possible effects of laeA and veA expression on aflatoxin production at all cell densities, as aflatoxin production in the wild type is highest at low population levels. Regardless of cell densities, the Δ veA and Δ laeA strains never produced observable aflatoxin under the growth conditions used here, whereas all the MC strains produced aflatoxin in all treatments (FIG. 17). The MCveA strain also showed a density-dependent decrease of aflatoxin with increasing cell population, similar to the wild type, whereas the MClaeA strain did not, and the double mutant showed an intermediate result. Aflatoxin production correlated with sclerotial production.

VeA and LaeA are important factors for seed colonization. Recently, Kale et al. reported that laeA mutants were aberrant in host colonization and aflatoxin production on both peanut and maize seed, but there are no reports for the role of VeA in *A. flavus* pathogenicity. Here, we examined and contrasted colonization attributes of the different veA and laeA mutants on two peanut cultivars and one maize hybrid.

Each fungal strain maintained similar growth patterns regardless of the host seed. FIG. **18**A shows that both null mutants produced fewer conidia than the wild type during growth on seeds, with the ΔveA strain developing significantly fewer conidia than the $\Delta laeA$ strain. Visually, the ΔveA strain was most crippled in its ability to grow on any seed (data not shown). The MCveA and MCveA-laeA strains also produced fewer conidia than the wild type; however, the MClaeA strain was similar to the wild type in conidial production, depending on the host seed, as reported earlier. The MC strains also formed sclerotia on the seeds (data not shown).

The colonized seeds were next examined for aflatoxin contamination. All MC strains and the wild type produced aflatoxin in all hosts, in contrast to the lack of aflatoxin production by both the ΔveA and $\Delta laeA$ strain (FIG. 18B). The considerably higher aflatoxin production by some MC mutants in vitro (FIG. 17), however, was not replicated in growth on seed under the conditions in this study.

To further investigate the ability of the strains to colonize seed, histological studies were performed. We were specifically interested in assaying for maceration effects and reasoned that this could be partially measured by host cell lipid utilization. The staining techniques did not show any obvious difference in host penetration by MC strains compared to that of the wild type (data not shown). However, the two null

mutants exhibited different host invasion patterns. The results in FIGS. **19**A and B show that wild-type hyphae penetrated several layers of host epidermal and mesophyll cells, with accompanying dissolution of host lipid reserves. Although the Δ laeA strain also penetrated the host cells intracellularly, host lipid reserves were largely intact and the cell integrity appeared less damaged (FIGS. **19**A and B). In contrast, hyphae of the Δ veA strain grew intercellularly in epidermal cells and did not appear to penetrate peanut cells as well as hyphae of other strains (FIG. **19**A). This mutant, like the Δ laeA mutant, was also less able to degrade host cell lipid reserves than the wild type (FIG. **19**B). However, an in vitro assay for general lipase activity revealed no significant difference between these strains (data not shown).

The wild type and the Δ laeA and Δ veA strains were then 15 grown on media amended with different fatty acids either as sole carbon source or supplemented with glucose to determine if there might be any gross difference in the ability to utilize or be inhibited by short-, medium-, or long-chain fatty acids. The results did not support any critical difference 20 between the wild type and the two mutants when grown on a fatty acid as the sole carbon source, but the two mutants showed significant inhibition of growth compared to that of the wild type when cultured on GMM amended with oleic acid (FIG. 20). This experiment was repeated twice with 25 similar results (data not shown).

Discussion. In this study, we characterized the function and crossregulation of VeA and LaeA in *A. flavus* development and pathogenesis. The results, while confirming that VeA and LaeA share functions in regulating aflatoxin and sclerotial 30 production, also demonstrate distinct roles of VeA and LaeA in terms of vegetative growth, conidiation, density-dependent responses, and pattern of colonization of host tissues.

A requirement for LaeA in density-dependent sensing. Quorum-sensing systems in bacteria contribute to the produc- 35 tion of virulence factors and biofilm formation in interactions between bacteria and host. In fungi, a quorum-sensing system governing morphological shifts and virulence has been uncovered in the human pathogen Candida albicans. Recently, oxylipin-deficient lipoxygenase and dioxygenase 40 mutants have been found to affect a newly discovered quorum-sensing-like, density-dependent sclerotial-to-conidial morphology shift in A. flavus. Because oxylipin signaling is dependent on VeA function and VeA is part of a nuclear complex with LaeA, we asked if VeA or LaeA mutants could 45 be affected in this quorum-like morphology shift in A. flavus. Both null mutants were blocked in sclerotial formation regardless of cell population, and perhaps due to an inability to produce sclerotia, conidial production was relatively stable for each mutant at all three population levels, although it was 50 much higher in the ΔlaeA strain.

The MC strains showed clear differences in density-dependent development in that an extra copy of LaeA but not VeA abolished this quorum-like phenomenon (FIGS. 16A and B). To date, there are no chemical data identifying molecules 55 regulating the sclerotial-to-conidial switch in *A. flavus*, although oxylipins are hypothesized to fulfill this function at least in part. Quorum-sensing molecules for *Candida albicans* (farnesol and tyrosol) and *Saccharomyces cerevisiae* (phenylethanol and tryptophol) are aromatic alcohols and 60 control the morphological switch from the yeast to filamentous growth in these fungi.

Interestingly, the yeast-to-filamentous growth switch in the fungus *Ceratocystis ulmi* is attenuated by lipoxygenase inhibitors and may implicate oxylipins in quorum sensing in 65 this tree pathogen. We speculate that *A. flavus* MClaeA mutants are aberrant in oxylipin production and/or sensing

but that this can be remediated to some degree when VeA levels also increase, as demonstrated by the intermediate density-dependent phenotype of the MCveA-laeA strain. The effects of gene loss and gain on density-dependent development are summarized in Table 7.

50

VeA and LaeA feedback regulation. Both veA and laeA have been reported to be global regulators of secondary metabolites in A. flavus, as well as in other aspergilli. Here, the results indicate that the MCveA and MClaeA strainsparticularly the MClaeA strain—produce more aflatoxin and sclerotia than the wild type. The MCveA-laeA double mutant did not show increased toxin production compared to that of the single mutants or an additive effect on sclerotial production. Prior work indicated that LaeA negatively regulated veA expression, and here, we show evidence for VeA regulation of LaeA (FIG. 15), as was described for A. nidulans. These results support a mechanism of mutual repression of veA and laeA expression and may explain, in part, a dampening of the expression of both genes in the MCveA-laeA strain compared to the expression of the single genes in the MCveA and MClaeA strains which, in turn, may affect aflatoxin and sclerotial output in the double mutant.

Requirement for VeA and LaeA in host cell penetration and degradation. Host lipid reserves are depleted during seed colonization by Aspergillus, with lipase and esterase activities implicated in seed pathogenesis. Both null mutants were impaired in seed colonization, where neither strain could degrade lipid reserves despite hyphal penetration of at least some layers of the host seeds (FIGS. 19A and B). The crippled ability of both null mutants to utilize lipid reserves brings to mind several lipid biosynthesis mutants also impaired in Aspergillus colonization of seed, including β-oxidation mutants, odeA mutants [(delta)12-desaturase], and the oxylipin oxygenase mutants in A. nidulans and A. flavus. The inhibition of both null mutants by oleic acid (not seen in the wild type) (FIG. 20) suggests a possible toxic effect of this fatty acid on these strains which may relate to their impairment in growth on seed. It is less likely that the inhibition is associated with defects in β-oxidation, since the mutants grew equally as well as the wild type on oleic acid as a sole carbon source (data not shown), although we cannot rule out this possibility. Regardless of mechanism, the results of all of these studies together may support lipid utilization and/or signaling as an important factor in Aspergillus seed pathogenesis.

Interestingly, the hyphal penetration patterns of the two null mutants as revealed by Gomori staining were quite diverse, whereas hyphae of the ΔveA strain remained largely intercellular (FIG. 19A). This inability to penetrate intracellularly may indicate loss of degradative enzymes in this strain and may explain its poor production of conidia on host seed. However, we note that the strain is crippled in conidial production on medium also. The relative decrease of conidial production by the $\Delta laeA$ strain on seed (compared to its vigorous conidial production in medium) might be attributable to a loss in lipid assimilation or the possible toxicity effects mentioned above.

Histology of the MC strains presented an invasion and lipid degradation pattern similar to that of the wild type. The relatively decreased conidial production on seed from these strains is possibly a function of their skewed sclerotial development rather than an inability to obtain nutrients from the seed.

In conclusion, this example provides additional evidence for distinct roles of LaeA and VeA in the development and pathogenesis of *A. flavus* despite the considerable overlapping of functions previously reported. The loss of both genes

blocks the production of sclerotia and aflatoxin, but under our conditions, only laeA overexpression abolishes density-dependent phenomena, including a sclerotial-to-conidial shift and decreased aflatoxin production with cell population increase. The null mutants, while both were reduced in host 5 lipid utilization, displayed distinct cell ingress abilities as reflected in patterns of hyphal penetration of host cells.

Other embodiments and uses of the invention will be apparent to those skilled in the art from consideration from the specification and practice of the invention disclosed herein. All references cited herein for any reason, including all journal citations and U.S./foreign patents and patent applications, are specifically and entirely incorporated herein by reference.

It is understood that the invention is not confined to the 15 specific reagents, formulations, reaction conditions, etc., herein illustrated and described, but embraces such modified forms thereof as come within the scope of the following claims.

REFERENCES

Adams et al., Microbiol. Mol. Biol. Rev. 62, 35 (1998). Bayram et al., Fungal Genet Biol 45, 127 (2008).

Bayram et al., Science 320:1504-1506 (2008).

Brakhage, FEMS Microbiol. Lett. 148, 1 (1997).

Bok et al., Eukaryot. Cell 3, 527 (2004).

Bok et al., Eukaryot. Cell 3:527-535 (2004).

Bok et al., Eukaryot. Cell 4:1574-1582 (2005).

Bok et al., Mol. Microbiol. 61, 1636 (2006).

Brodhagen et al., Mol. Microbiol. 67:378-391 (2008).

Busch et al., Mol Microbiol 49, 717 (2003).

Busch et al., Proc. Natl. Acad. Sci. U.S.A. 104, 8089 (2007).

Calvo et al., Microbiol. Mol. Biol. Rev. 66, 447 (2002).

Calvo et al., Appl. Environ. Microbiol. 65:3668-3673 (1999).

Calvo et al., Appl. Environ. Microbiol. 70:4722-4739 (2004).

Chaveroche et al., Nucleic Acids Res 28, E97 (2000).

Cho et al., The Journal of Microbiology 41, 46 (2003).

Champe et al., J. Gen. Microbiol. 133:1383-1387 (1987).

Chen et al., Proc. Natl. Acad. Sci. USA 101:5048-5052 (2004).

Chen et al., Genes Dev. 20:1150-1161 (2006).

ogy and medicine, 2nd ed.

CRC Press, Boca Raton, Fla. (2004).

Dagenais et al., Infect. Immun. 76:3214-3220 (2008).

Diener et al., Annu. Rev. Phytopathol. 25:249-270 (1987).

Dreyer et al., Appl. Environ. Microbiol. 73, 3412 (2007).

Duran et al., Appl. Microbiol. Biotechnol. 73:1158-1168 (2007).

Duran et al., Open Mycol. J. 3:27-36 (2009).

Eng et al., J. Am. Soc. Mass 5, 976 (1994).

Etxebeste et al., Eukaryot Cell (2007).

Feinberg et al., Anal Biochem 132, 6 (1983).

Fernandez-Abalos et al., Mol Microbiol 27, 121 (1998).

Fray, Ann. Bot. 89:245-253 (2002).

Georgianna et al., 2009. Fungal Genet. Biol. 46:113-125 (2009).

Golemis, R. Brent, in Current protocols in molecular biology F. M. Ausubel et al., Eds. (Harvard Medical School, Massachusetts), vol. 3, pp. 429-454 (1996).

Gyuris et al., Cell 75, 791 (1993).

Greenspan et al., 1985. J. Cell Biol. 100:965-973.

Hanahan et al., Methods Enzymol 204, 63 (1991).

52

Hicks et al., in The Mycota, vol. 11, F. Kempken, Ed. (Springer, Berlin), pp. 55-69 (2002).

Hu et al., Mol. Cell 9, 789 (2002).

Hornby et al., Appl. Environ. Microbiol. 67:2982-2992 (2001).

Horowitz et al., Appl. Environ. Microbiol. 74:5674-5685 (2008).

Horowitz et al., Mol. Plant Microbe Interact., in press.

10 Jensen et al., Appl. Environ. Microbiol. 58:2505-2508 (1992).

Kale et al., Fungal Genet. Biol. 45:1422-1429 9 (2008).

Kato et al., Eukaryot. Cell 2:1178-1186 (2003).

Kim et al., Fungal Genet. Biol. 37:72-80 (2002).

Klich, Mol. Plant Pathol. 8:713-722 (2007).

Keller et al., Nat. Rev. Microbiol. 3, 937 (2005).

Keller et al., Appl. Environ. Microbiol. 60:1444-1460 (1994).

Kolar et al., Gene 62, 127 (1988).

20 Krappmann et al, Eukaryot. Cell 4, 1298 (2005).

Krappmann et al., Mol Microbiol 61, 76 (2006).

Kunkel, Proc Natl Acad Sci USA 82, 488 (1985).

Li et al., Mol. Microbiol. 62, 1418 (2006).

Link et al., Nat Biotechnol 17, 676 (1999).

25 Lillie, Histopathologic technique and practical histochemistry, 3rd ed. McGraw-Hill Book Company, New York, N.Y.

Maggio-Hall et al., Mol. Microbiol. 54:1173-1185 (2004).

Maggio-Hall et al., Mol. Plant Microbe Interact. 18:783-793 (2005).

Michailides et al., Plant Pathol. 56:352 (2007).

Miller et al., Mol. Cell. Biol. 5:1714-1721 (1985).

Mooney et al., Genes Dev. 4, 1473 (1990).

Muyrers et al., Genet Eng (NY) 22, 77 (2000).

Muture et al., East Afr. Med. J. 82:275-279 (2005).

Nayak et al., Genetics 172, 1557 (2.006).

Ni et al., PLoS One 2, e970 (2007).

Perrin et al., PLoS Pathog. 3:e50 (2007).

Puig et al., Methods 24, 218 (2001).

Punt et al., Methods Enzymol 216, 447 (1992).

Purschwitz et al., Curr. Biol. 18, 255 (2008).

Pettit, Yellow mold and aflatoxin, p. 35-36. In D. M. Porter, D. H. Smith, and R.

Cseke et al., Handbook of molecular/cellular methods in biol- 45 Rodeiguez-Kabana (ed.), Compendium of peanut diseases. The American Phytopathological Society, St. Paul, Minn.

> (1984).Rohila et al., Plant J. 38, 172 (2004).

Rubens, J., and K. F. Cardwell. The cost of mycotoxin management in the United States, p. 1-13. In H. K. Abbas (ed.), Aflatoxin and food safety. CRC Press, Boca Raton, Fla. (2005).

Saiki et al., Nature 324, 163(1986).

Sambrook et al., Maniatis, Molecular Cloning: A Laboratory Manual

Seiler, et al., Mol Biol Cell 17, 4080 (2006).

Shevchenko et al., Anal Chem 68, 850 (1996).

Shwab et al., Eukaryot. Cell 6, 1656 (2007).

Southern, J Mol Biol 98, 503 (1975).

Spröte et al., Arch. Microbiol. 188, 69 (2007).

Stinnett et al., Mol. Microbiol. 63, 242 (2007).

Shchepin, et al., Chem. Biol. 10:743-750 92003).

Shimizu, K. et al., Genetics 157: 591-600 (2001).

Smart et al., Phytopathology 80:1287-1294 (1990).

Szewczyk et al., Nat. Protoc. 1:3111-3120 (2006).

Thompson et al., Nucleic Acids Res 22, 4673 (1994).

Tsitsigiannis et al., Mol. Microbiol. 59:882-892 (2006). Tsitsigiannis et al., Microbiology 151:1809-1821 (2005). Tsitsigiannis et al., J. Biol. Chem. 279:11344-11353 (2004). Williams, Microbiology 153:3923-3938 (2007). Wilson et al., Microbiology 150:2881-2888 (2004). Woloshuk et al., Appl. Environ. Microbiol. 60, 2408 (1994). 54

Yu et al., Fungal Genet Biol 41, 973 (2004).
Yu et al., Rev. Iberoam. Micol. 22:194-202 (2005).
Yu et al., Mycotoxin production and prevention of aflatoxin contamination in food and feed. In G. H. Goldman and S. A. Osmani (ed.), The aspergilli. CRC Press, Boca Raton, Fla. (2008).

SEQUENCE LISTING <160> NUMBER OF SEQ ID NOS: 146 <210> SEQ ID NO 1 <211> LENGTH: 49 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide <400> SEQUENCE: 1 tttggccatg ggtggtagcg gtggtatggt gagcaagggc gaggagctg 49 <210> SEQ ID NO 2 <211> LENGTH: 40 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide <400> SEQUENCE: 2 aaaatttaag cttctacttg tacagttcgt ccatgccgtg 40 <210> SEQ ID NO 3 <211> LENGTH: 23 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide <400> SEQUENCE: 3 23 actcacgaat ccacgggata cat <210> SEQ ID NO 4 <211> LENGTH: 36 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide <400> SEQUENCE: 4 ggcctgagtg gccgggtggg atacggtcca tcgaaa 36 <210> SEQ ID NO 5 <211> LENGTH: 39 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide <400> SEQUENCE: 5 ggccatctag gccgaccgta tattgtttca taaatcctt 39 <210> SEQ ID NO 6 <211> LENGTH: 24 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: Synthetic oligonucleotide

-continued

<400> SEQUENCE: 6		
tatgaccgcg tgagcaaata ggac		24
010 GEO TE NO E		
<210> SEQ ID NO 7 <211> LENGTH: 23		
<212> TYPE: DNA		
<213> ORGANISM: Artificial Sequence		
<220> FEATURE:		
<223> OTHER INFORMATION: Synthetic oligon	ucleotide	
<400> SEQUENCE: 7		
atgtttgaga tgggcccggt ggg		23
<210> SEQ ID NO 8		
<211> LENGTH: 26		
<212> TYPE: DNA <213> ORGANISM: Artificial Sequence		
<220> FEATURE:		
<223> OTHER INFORMATION: Synthetic oligon	ucleotide	
AAA GROUPIGE A		
<400> SEQUENCE: 8		
ttatcttaat ggtttcctag cctggt		26
-210, CEO ID NO 0		
<210> SEQ ID NO 9 <211> LENGTH: 23		
<212> TYPE: DNA		
<213> ORGANISM: Artificial Sequence		
<220> FEATURE:		
<223> OTHER INFORMATION: Synthetic oligon	ucleotide	
<400> SEQUENCE: 9		
atgtacgctg ttgaggatag ggc		23
<210> SEQ ID NO 10		
<211> LENGTH: 25		
<212> TYPE: DNA <213> ORGANISM: Artificial Sequence		
<220> FEATURE:		
<223> OTHER INFORMATION: Synthetic oligon	ucleotide	
-400- CEOHENCE 10		
<400> SEQUENCE: 10		
ttagtattcg ttatccagac catcg		25
<210> SEQ ID NO 11		
<211> LENGTH: 31		
<212> TYPE: DNA		
<213> ORGANISM: Artificial Sequence		
<pre><220> FEATURE: <223> OTHER INFORMATION: Synthetic oligon</pre>	ugleotide	
(223) OTHER INFORMATION. Synthetic Offgon	ucleotide	
<400> SEQUENCE: 11		
		2.1
ctcgagttag tattcgttat ccagaccatc g		31
<210> SEQ ID NO 12		
<211> LENGTH: 30		
<212 > TYPE: DNA		
<213> ORGANISM: Artificial Sequence <220> FEATURE:		
<223 > OTHER INFORMATION: Synthetic oligon	ucleotide	
<400> SEQUENCE: 12		
ccatonaton otacactton accaccacca		30
ccatggatgg ctacacttgc agcaccacca		50
<210> SEQ ID NO 13		
~211 \ T.FNGTH + 32		

<211> LENGTH: 32

```
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 13
ctcgagttaa cgcatggtgg caggctttga ga
                                                                       32
<210> SEQ ID NO 14
<211> LENGTH: 21
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 14
atggtgagca agggcgagga g
<210> SEQ ID NO 15
<211> LENGTH: 51
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEOUENCE: 15
ggtggtggtg ctgcaagtgt agccatcgtg gcgatggagc gcatgatata g
                                                                        51
<210> SEQ ID NO 16
<211> LENGTH: 21
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 16
atggccgaca agcagaagaa c
                                                                        21
<210> SEQ ID NO 17
<211> LENGTH: 54
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 17
acgagttccc accgggccca tctcaaacat gtggttcatg accttctgtt tcag
                                                                        54
<210> SEQ ID NO 18
<211> LENGTH: 54
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 18
ggaatgcgcc ctatcctcaa cagcgtacat gtggttcatg accttctgtt tcag
                                                                       54
<210> SEQ ID NO 19
<211> LENGTH: 31
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 19
```

-continued

gaattcatgc agcagcccaa gcgcgcgaga g	31
010 dD0 tD W0 00	
<210> SEQ ID NO 20 <211> LENGTH: 33	
<212> TYPE: DNA	
<213> ORGANISM: Artificial Sequence	
<220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide	
<400> SEQUENCE: 20	
aaagggcccc gagaatgtcc gcctgacccg tgc	33
<210> SEQ ID NO 21	
<211> LENGTH: 25	
<212> TYPE: DNA <213> ORGANISM: Artificial Sequence	
<220> FEATURE:	
<223> OTHER INFORMATION: Synthetic oligonucleotide	
<400> SEQUENCE: 21	
ccaagtctgc ccgacaagct cactg	25
<210> SEQ ID NO 22	
<211> LENGTH: 66	
<212> TYPE: DNA <213> ORGANISM: Artificial Sequence	
<220> FEATURE:	
<223> OTHER INFORMATION: Synthetic oligonucleotide	
<400> SEQUENCE: 22	
cgccacagcg acgaggacga tggtctggat aacgaatacg gtggtagcgg tggtatggtg	60
agcaag	66
<210> SEQ ID NO 23	
<211> LENGTH: 24	
<212> TYPE: DNA	
<213> ORGANISM: Artificial Sequence	
<220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide	
<400> SEQUENCE: 23	
gtattcgtta tccagaccat cgtc	24
<210> SEQ ID NO 24	
<211> LENGTH: 57	
<212> TYPE: DNA <213> ORGANISM: Artificial Sequence	
<220> FEATURE:	
<223> OTHER INFORMATION: Synthetic oligonucleotide	
<400> SEQUENCE: 24	
ctgcacatat accaggctag gaaaccatta agaggtggta gcggtggtat ggtgagc	57
<210> SEQ ID NO 25	
<211> LENGTH: 24	
<212> TYPE: DNA	
<213> ORGANISM: Artificial Sequence <220> FEATURE:	
<223> OTHER INFORMATION: Synthetic oligonucleotide	
<400> SEQUENCE: 25	
tcttaatggt ttcctagcct ggta	24
<210> SEQ ID NO 26	

<211> LENGTH: 26

```
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 26
cctcgccctc ctgcatcaat attcgg
                                                                        26
<210> SEQ ID NO 27
<211> LENGTH: 78
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 27
gagacggcta tgaaattctt tttccatctt ctcttaccac cgctaccacc tcttaatggt
                                                                        60
                                                                        78
ttcctagcct ggtatatg
<210> SEQ ID NO 28
<211> LENGTH: 60
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEOUENCE: 28
gagcaggcgc tctacatgag catgccctgc ccctgagagc aaaaggcgac cacatccagg
                                                                        60
<210> SEQ ID NO 29
<211> LENGTH: 24
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 29
tegteaaceg ceteagetgg aace
                                                                        2.4
<210> SEQ ID NO 30
<211> LENGTH: 25
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 30
cctcctcgcc gcctctagta ccgtc
                                                                        25
<210> SEQ ID NO 31
<211> LENGTH: 64
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 31
gaaattcttt ttccatcttc tcttaccacc gctaccaccg tattcgttat ccagaccatc
                                                                        60
gtcc
                                                                        64
<210> SEQ ID NO 32
<211> LENGTH: 62
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
```

```
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 32
cgagcaggcg ctctacatga gcatgccctg cccctgaaga ccgtatattg tttcataaat
                                                                        60
                                                                        62
<210> SEQ ID NO 33
<211> LENGTH: 25
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
<223 > OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 33
cggctgttta cattgtgttt tctgg
                                                                        25
<210> SEQ ID NO 34
<211> LENGTH: 23
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
<223 > OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 34
                                                                        23
ccgtgaagaa cttggcgttg tag
<210> SEQ ID NO 35
<211> LENGTH: 23
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 35
ggaccgtcta attcaactca cag
                                                                        23
<210> SEQ ID NO 36
<211> LENGTH: 23
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 36
cttccagcgg ttatcctccg ttg
                                                                        23
<210> SEQ ID NO 37
<211> LENGTH: 29
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 37
atctgacaga gcggccgcaa ttgattacg
                                                                        29
<210> SEQ ID NO 38
<211> LENGTH: 30
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223 > OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 38
atatatgcgg ccgctcttgc atctttgttt
                                                                        30
```

```
<210> SEQ ID NO 39
<211> LENGTH: 27
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 39
gataccaaac ggaactggct gttatgg
                                                                       27
<210> SEQ ID NO 40
<211> LENGTH: 23
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
<223 > OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 40
                                                                       23
atcgacgcaa ccatcgaagc agc
<210> SEQ ID NO 41
<211> LENGTH: 24
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 41
                                                                       24
gatctttgcc cggtgtatga aacc
<210> SEQ ID NO 42
<211> LENGTH: 32
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 42
tcggaggagg ccatggtgat gtctgctcaa gc
                                                                       32
<210> SEQ ID NO 43
<211> LENGTH: 33
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 43
                                                                       33
gacatcacca tggcctcctc cgaggacgtc atc
<210> SEQ ID NO 44
<211> LENGTH: 44
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 44
ggetecageg cetgeaceag etceggegee ggtggagtgg egge
                                                                       44
<210> SEQ ID NO 45
<211> LENGTH: 44
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
```

```
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 45
ggagctggtg caggcgctgg agccactggc ggcaaatctg gtgg
                                                                       44
<210> SEQ ID NO 46
<211> LENGTH: 24
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 46
atctggaggg gacaggcagt ttat
                                                                       24
<210> SEQ ID NO 47
<211> LENGTH: 60
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEOUENCE: 47
gggtttcgaa ctacatcaag ggtccaagac cgacatcgag gctctgtaca gtgaccggtg
                                                                       60
<210> SEQ ID NO 48
<211> LENGTH: 26
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
<223 > OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 48
agggaattct caggggcagg gcatgc
                                                                       26
<210> SEQ ID NO 49
<211> LENGTH: 22
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 49
gaaggtcgat gatggtgtga tg
                                                                       22
<210> SEQ ID NO 50
<211> LENGTH: 22
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 50
ctagaggtaa agatcaaggt ag
                                                                       22
<210> SEQ ID NO 51
<211> LENGTH: 22
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 51
ctgatggctg aatgaagcac ag
                                                                       2.2
```

```
<210> SEQ ID NO 52
<211> LENGTH: 22
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223 > OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 52
tgctttacga cgatagccat gc
                                                                        22
<210> SEQ ID NO 53
<211> LENGTH: 45
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
<223 > OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 53
ggtgaagagc attgtttgag gcagcggcca gtctttagac aaatg
                                                                        45
<210> SEQ ID NO 54
<211> LENGTH: 45
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 54
                                                                        45
agtgcctcct ctcagacaga ataggataac gaatactaaa gaccg
<210> SEQ ID NO 55
<211> LENGTH: 24
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 55
tatgcactgg cactcaagca accg
                                                                        2.4
<210> SEQ ID NO 56
<211> LENGTH: 24
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 56
                                                                        24
gtgcatgacg gtcgtatctg gtcc
<210> SEQ ID NO 57
<211> LENGTH: 18
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 57
                                                                        18
ggctgtagtc gctttgtt
<210> SEQ ID NO 58
<211> LENGTH: 18
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
```

-continued

<400> SEQUENCE: 58		
gcccagtgta agaaagga	18	
<u> </u>		
<210> SEQ ID NO 59		
<211> LENGTH: 21		
<212> TYPE: DNA		
<213> ORGANISM: Artificial Sequence		
<220> FEATURE:		
<223> OTHER INFORMATION: Synthetic oligonucleotide		
<400> SEQUENCE: 59		
getgtegate tttgtaccet g	21	
geographic congrators g	21	
<210> SEQ ID NO 60 <211> LENGTH: 21		
<212> TYPE: DNA		
<213> ORGANISM: Artificial Sequence		
<220> FEATURE:		
<223> OTHER INFORMATION: Synthetic oligonucleotide		
<400> SEQUENCE: 60		
auttantens tataataan t	21	
cgttcctgga tgtggtcgcc t	21	
<210> SEQ ID NO 61		
<211> LENGTH: 33 <212> TYPE: DNA		
<213> ORGANISM: Artificial Sequence		
<220> FEATURE:		
<223> OTHER INFORMATION: Synthetic oligonucleotide		
<400> SEQUENCE: 61		
atataagett aatggetaca ettgeageae eae	33	
<210> SEQ ID NO 62		
<211> LENGTH: 33 <212> TYPE: DNA		
<213> ORGANISM: Artificial Sequence		
<220> FEATURE:		
<223> OTHER INFORMATION: Synthetic oligonucleotide		
<400> SEQUENCE: 62		
atatgtcgac ttaacgcatg gtggcaggct ttg	33	
<210> SEQ ID NO 63		
<211> LENGTH: 34		
<212> TYPE: DNA <213> ORGANISM: Artificial Sequence		
<220> FEATURE:		
<223> OTHER INFORMATION: Synthetic oligonucleotide		
<400> SEQUENCE: 63		
atataagctt aatgcagcag cccaagcgcg cgag	34	
<210> SEQ ID NO 64		
<211> LENGTH: 32		
<212> TYPE: DNA		
<213> ORGANISM: Artificial Sequence		
<pre><220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide</pre>		
<400> SEQUENCE: 64		
atatogatto atogotogoo oggantatoo ao	32	
atatgaattc atgagtgcgg cgaactatcc ag	32	
<210> SEQ ID NO 65		

<211> LENGTH: 33

```
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 65
atatgtcgac tcaccgagga gttccgttcg ctg
                                                                        33
<210> SEQ ID NO 66
<211> LENGTH: 36
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 66
atatgaattc atgtttgaga tgggcccggt gggaac
<210> SEQ ID NO 67
<211> LENGTH: 34
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 67
                                                                         34
atatgtcgac ttatcttaat ggtttcctag cctg
<210> SEQ ID NO 68
<211> LENGTH: 32
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 68
atataagett ateaacgage ateageacaa ac
                                                                        32
<210> SEQ ID NO 69
<211> LENGTH: 33
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 69
atatgtcgac tccatattcc actgccgacg gac
                                                                         33
<210> SEQ ID NO 70
<211> LENGTH: 34
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 70
atatgaattc tctgatagga cagccatgca aatc
                                                                        34
<210> SEQ ID NO 71
<211> LENGTH: 30
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 71
```

atatgaattc atgtacgctg ttgaggatag	30	
<210> SEQ ID NO 72		
<211> LENGTH: 32 <212> TYPE: DNA		
<213> ORGANISM: Artificial Sequence		
<220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide		
<400> SEQUENCE: 72		
atatgtcgac ttagtattcg ttatccagac ca	32	
<210> SEQ ID NO 73		
<211> LENGTH: 31		
<212> TYPE: DNA <213> ORGANISM: Artificial Sequence		
<220> FEATURE:		
<223> OTHER INFORMATION: Synthetic oligonucleotide		
<400> SEQUENCE: 73		
atatgaattc acggtagcgc gggtatcgga g	31	
<210> SEQ ID NO 74		
<211> LENGTH: 32		
<212> TYPE: DNA <213> ORGANISM: Artificial Sequence		
<220> FEATURE:		
<223> OTHER INFORMATION: Synthetic oligonucleotide		
<400> SEQUENCE: 74		
atatgaattc atgtcttcat cgtatccacc ac	32	
<210> SEQ ID NO 75		
<211> LENGTH: 32		
<212> TYPE: DNA <213> ORGANISM: Artificial Sequence		
<220> FEATURE:		
<223> OTHER INFORMATION: Synthetic oligonucleotide		
<400> SEQUENCE: 75		
atatetegag accaggeace gggaeggaga tg	32	
<210> SEQ ID NO 76		
<211> LENGTH: 33		
<212> TYPE: DNA <213> ORGANISM: Artificial Sequence		
<220> FEATURE:		
<223> OTHER INFORMATION: Synthetic oligonucleotide		
<400> SEQUENCE: 76		
atatotogag agtaggaata gtooctacto gtg	33	
<210> SEQ ID NO 77		
<211> LENGTH: 34		
<212> TYPE: DNA <213> ORGANISM: Artificial Sequence		
<220> FEATURE:		
<223> OTHER INFORMATION: Synthetic oligonucleotide		
<400> SEQUENCE: 77		
atatctcgag tccaggccct ggagtaactg gctg	34	
<210> SEQ ID NO 78		
<211> LENGTH: 23 <212> TYPE: DNA		
<212> ORGANISM: Artificial Sequence		

```
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 78
ttcgctagac agctcattct acg
                                                                       23
<210> SEQ ID NO 79
<211> LENGTH: 24
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 79
tagtattcgt tatccagacc atcg
                                                                       24
<210> SEQ ID NO 80
<211> LENGTH: 24
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 80
atacctggat aaaccaaatc gagc
                                                                       24
<210> SEQ ID NO 81
<211> LENGTH: 22
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 81
                                                                       22
aggttcattc gcagggctag ac
<210> SEQ ID NO 82
<211> LENGTH: 22
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 82
accactacag ctaccactct cc
                                                                       22
<210> SEQ ID NO 83
<211> LENGTH: 22
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 83
                                                                       22
tttcgatgct ctctgagacg gc
<210> SEQ ID NO 84
<211> LENGTH: 16
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 84
Arg Leu Glu Val Ile Ser Asn Pro Phe Ile Val Tyr Ser Ala Lys Lys
                5
                                    10
```

```
<210> SEQ ID NO 85
<211> LENGTH: 16
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 85
Arg Leu Glu Val Ile Ser Asn Pro Phe Ile Val Tyr Ser Ala Lys Lys
<210> SEQ ID NO 86
<211> LENGTH: 16
<212> TYPE: PRT
<213 > ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 86
Arg Leu Glu Val Ile Ser Asn Pro Phe Ile Val Tyr Ser Ala Lys Lys
<210> SEQ ID NO 87
<211> LENGTH: 15
<212> TYPE: PRT
<213 > ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 87
Lys Ile Gly Val Trp Phe Val Leu Gln Asp Leu Ser Val Arg Thr
                                   10
<210> SEQ ID NO 88
<211> LENGTH: 18
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 88
Lys Ser Val Ser Asp Leu Pro Gln Ser Asp Ile Ala Glu Val Ile Asn
Lys Gly
<210> SEQ ID NO 89
<211> LENGTH: 14
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 89
Arg Ile Trp Ser Leu Gln Val Val Gln Gln Pro Ile Arg Ala
<210> SEQ ID NO 90
<211> LENGTH: 16
<212> TYPE: PRT
<213 > ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 90
Lys Glu Ile His Ala Tyr Asn Ile Leu His Ile Tyr Gln Ala Arg Lys
                                   10
<210> SEQ ID NO 91
<211> LENGTH: 14
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 91
Arg Tyr Ala Val Ala Gly Gly Pro Ala Pro Trp Asn Arg Asn
      5
```

```
<210> SEQ ID NO 92
<211> LENGTH: 17
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 92
Arg Val Ser Glu Ser Leu Ile Tyr Ala Pro His Pro Thr Asn Gly Arg
<210> SEQ ID NO 93
<211> LENGTH: 16
<212> TYPE: PRT
<213 > ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 93
Lys Ile Ile Gl<br/>n Val Ala Leu Asp Gly Leu Glu As<br/>n Ile Leu Lys Val 1 \phantom{\bigg|} 5
<210> SEQ ID NO 94
<211> LENGTH: 13
<212> TYPE: PRT
<213 > ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 94
Lys Ile Gln Ala Val Ile Glu Ala Gly Ile Pro Arg Arg
<210> SEQ ID NO 95
<211> LENGTH: 20
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 95
Lys Thr Pro Gln Pro Asp Trp Asn Thr Ile Ala Pro Ala Leu Pro Val
1 5
                         10
Leu Ala Lys Leu
<210> SEQ ID NO 96
<211> LENGTH: 22
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 96
Lys Gly Thr Ala Pro Ile Leu Ala Ser Thr Phe Ser Glu Pro Phe Gln
Val Phe Ser Ala Lys Lys
<210> SEQ ID NO 97
<211> LENGTH: 15
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 97
Lys Ile Gly Val Trp Phe Val Leu Gln Asp Leu Ser Val Arg Thr
1 5
                                   10
<210> SEQ ID NO 98
<211> LENGTH: 18
<212> TYPE: PRT
<213 > ORGANISM: Aspergillus nidulans
```

```
<400> SEQUENCE: 98
Lys Ser Val Ser Asp Leu Pro Gln Ser Asp Ile Ala Glu Val Ile Asn
                                    10
Lys Gly
<210> SEQ ID NO 99
<211> LENGTH: 18
<212> TYPE: PRT
<213 > ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 99
Lys Asp Ala Thr Glu Gly Thr Gln Pro Met Pro Ser Pro Val Pro Gly
Lys Leu
<210> SEQ ID NO 100
<211> LENGTH: 15
<212> TYPE: PRT
<213 > ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 100
Lys Lys Phe Pro Gly Leu Thr Thr Ser Thr Pro Ile Ser Arg Met
<210> SEQ ID NO 101
<211> LENGTH: 28
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 101
Lys Leu Met Thr Asn Gln Gly Ser Pro Val Leu Thr Gly Val Pro Val
                                   10
Ala Gly Val Ala Tyr Leu Asp Lys Pro Asn Arg Ala
         20
<210> SEQ ID NO 102
<211> LENGTH: 16
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 102
Lys Glu Ile His Ala Tyr Asn Ile Leu His Ile Tyr Gln Ala Arg Lys
<210> SEQ ID NO 103
<211> LENGTH: 11
<212> TYPE: PRT
<213 > ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 103
Arg Ile Gln Gln Leu Ala Ala Asp Val Lys Ser
<210> SEQ ID NO 104
<211> LENGTH: 14
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 104
Arg Tyr Ala Val Ala Gly Gly Pro Ala Pro Trp Asn Arg Asn
     5
```

-continued

```
<210> SEQ ID NO 105
<211> LENGTH: 20
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 105
Lys Asp Val Asp Asn Thr Asp Gly Gly Phe Phe Val Trp Gly Asp Leu
Ser Ile Lys Val
<210> SEQ ID NO 106
<211> LENGTH: 22
<212> TYPE: PRT
<213 > ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 106
Arg Leu Lys Asp Val Asp Asn Thr Asp Gly Gly Phe Phe Val Trp Gly 1 \phantom{-} 10 \phantom{-} 15
Asp Leu Ser Ile Lys Val
<210> SEQ ID NO 107
<211> LENGTH: 16
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 107
Lys Glu Ile His Ala Tyr Asn Ile Leu His Ile Tyr Gln Ala Arg Lys
<210> SEQ ID NO 108
<211> LENGTH: 22
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 108
Arg Trp Tyr Asn Leu Ala Val Ser Glu Ser Ile Glu Asn Leu Ser Leu
                                       10
Ala Pro Phe Ser Arg Val
            20
<210> SEQ ID NO 109
<211> LENGTH: 14
<212> TYPE: PRT
<213 > ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 109
Arg Tyr Ala Val Ala Gly Gly Pro Ala Pro Trp Asn Arg Asn
<210> SEQ ID NO 110
<211> LENGTH: 22
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 110
Lys Gly Thr Ala Pro Ile Leu Ala Ser Thr Phe Ser Glu Pro Phe Gln
Val Phe Ser Ala Lys Lys
```

<210> SEQ ID NO 111

```
<211> LENGTH: 18
<212> TYPE: PRT
<213 > ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 111
Lys Ser Val Ser Asp Leu Pro Gln Ser Asp Ile Ala Glu Val Ile Asn
               5
                                    10
Lys Gly
<210> SEQ ID NO 112
<211> LENGTH: 14
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 112
Arg Ile Trp Ser Leu Gln Val Val Gln Gln Pro Ile Arg Ala
<210> SEQ ID NO 113
<211> LENGTH: 16
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 113
Arg Leu Glu Val Ile Ser Asn Pro Phe Ile Val Tyr Ser Ala Lys Lys 1 \phantom{\bigg|} 10 \phantom{\bigg|} 15
<210> SEQ ID NO 114
<211> LENGTH: 21
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEOUENCE: 114
Arg Arg Pro Asp Gln Tyr Ala Gly Ser Asp Ala Tyr Ala Asn Ala Pro
                                     10
Glu Arg Pro Arg Ser
           20
<210> SEQ ID NO 115
<211> LENGTH: 19
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 115
Arg Arg Pro Ser Ala Val Glu Tyr Gly Gln Pro Ile Ala Gln Pro Tyr
                                     10
Gln Arg Pro
<210> SEQ ID NO 116
<211> LENGTH: 573
<212> TYPE: PRT
<213> ORGANISM: Aspergillus nidulans
<400> SEQUENCE: 116
Met Ala Thr Leu Ala Ala Pro Pro Pro Pro Leu Gly Glu Ser Gly Asn
                      10
Ser Asn Ser Val Ser Arg Ile Thr Arg Glu Gly Lys Lys Ile Thr Tyr
                                25
Lys Leu Asn Ile Met Gln Gln Pro Lys Arg Ala Arg Ala Cys Gly Gln
                            40
Gly Ser Lys Ser His Thr Asp Arg Arg Pro Val Asp Pro Pro Pro Val
              55
```

Ile 65	Glu	Leu	Asn	Ile	Phe 70	Glu	Ser	Asp	Pro	His 75	Asp	Asp	Ser	Asn	80 Lys
Thr	Asp	Ile	Thr	Phe 85	Val	Tyr	Asn	Ala	Asn 90	Phe	Phe	Leu	Phe	Ala 95	Thr
Leu	Glu	Pro	Glu 100	Arg	Pro	Ile	Ala	Thr 105	Gly	ГЛа	Leu	Met	Thr 110	Asn	Gln
Gly	Ser	Pro 115	Val	Leu	Thr	Gly	Val 120	Pro	Val	Ala	Gly	Val 125	Ala	Tyr	Leu
Asp	Lys 130	Pro	Asn	Arg	Ala	Gly 135	Tyr	Phe	Ile	Phe	Pro 140	Asp	Leu	Ser	Val
Arg 145	Asn	Glu	Gly	Ser	Tyr 150	Arg	Phe	Ser	Phe	His 155	Leu	Phe	Glu	Gln	Ile 160
Lys	Asp	Pro	Lys	Asp 165	Ala	Thr	Glu	Gly	Thr 170	Gln	Pro	Met	Pro	Ser 175	Pro
Val	Pro	Gly	Lys 180	Leu	Ser	Ser	Pro	Gln 185	Glu	Phe	Leu	Glu	Phe 190	Arg	Leu
Glu	Val	Ile 195	Ser	Asn	Pro	Phe	Ile 200	Val	Tyr	Ser	Ala	Lys 205	Lys	Phe	Pro
Gly	Leu 210	Thr	Thr	Ser	Thr	Pro 215	Ile	Ser	Arg	Met	Ile 220	Ala	Glu	Gln	Gly
Сув 225	Arg	Val	Arg	Ile	Arg 230	Arg	Asp	Val	Arg	Met 235	Arg	Arg	Arg	Gly	Asp 240
Lys	Arg	Thr	Glu	Asp 245	Tyr	Asp	Tyr	Asp	Asn 250	Glu	Arg	Gly	Tyr	Asn 255	Asn
Arg	Arg	Pro	Asp 260	Gln	Tyr	Ala	Gly	Ser 265	Asp	Ala	Tyr	Ala	Asn 270	Ala	Pro
Glu	Arg	Pro 275	Arg	Ser	Thr	Ser	Ile 280	Ser	Thr	Asn	Met	Asp 285	Pro	Tyr	Ser
Tyr	Pro 290	Ser	Arg	Arg	Pro	Ser 295	Ala	Val	Glu	Tyr	Gly 300	Gln	Pro	Ile	Ala
Gln 305	Pro	Tyr	Gln	Arg	Pro 310	Met	Ala	Ser	Thr	Pro 315	Ala	Pro	Ser	Ser	Thr 320
Pro	Ile	Pro	Ala	Pro 325	Ile	Pro	Met	Pro	Gly 330	Pro	Val	Ala	Leu	Pro 335	Pro
Ser	Thr	Pro	Ser 340	Pro	Ala	Ser	Ala	His 345	Ala	Pro	Ala	Pro	Pro 350	Ser	Val
Pro	Leu	Ala 355	Ala	Pro	Pro	Pro	Leu 360	His	Thr	Pro	Ser	Tyr 365	Gln	Ser	His
Leu	Ser 370	Phe	Gly	Ala	Thr	Gln 375	Thr	Gln	Tyr	Pro	Ala 380	Pro	Gln	Leu	Ser
His 385	Ile	Pro	Gln	Gln	Thr 390	Thr	Thr	Pro	Thr	His 395	Pro	Tyr	Ser	Pro	Arg 400
Ser	Ser	Ile	Ser	His 405	Ser	Arg	Asn	Gln	Ser 410	Ile	Ser	Glu	Tyr	Glu 415	Pro
Ser	Met	Gly	Tyr 420	Pro	Gly	Ser	Gln	Thr 425	Arg	Leu	Ser	Ala	Glu 430	Arg	Pro
Ser	Tyr	Gly 435	Gln	Pro	Ser	Gln	Thr 440	Thr	Ser	Leu	Pro	Pro 445	Leu	Arg	His
Ser	Leu 450	Glu	Pro	Ser	Val	Asn 455	Ser	Arg	Ser	Lys	Thr 460	Pro	Ser	Asn	Met
Ile 465	Thr	Ser	Leu	Pro	Pro 470	Ile	Gln	Ser	Leu	Ser 475	Glu	Leu	Pro	Ser	Thr 480

-continued

Thr Ser Gln Pro Ser Ser Ala Ile Gly Ser Ser Pro Ala Asn Glu Pro 490 Gly Pro Arg Leu Trp Glu Thr Asn Ser Met Leu Ser Lys Arg Thr Tyr Glu Glu Ser Phe Gly His Asp Asp Arg Pro Leu Tyr Asn Gly Met Arg 520 Pro Asp Ser Glu Ser Tyr Pro Gly Gly Met Gln Arg Arg Pro Ser Tyr Glu Arg Ser Ser Leu Leu Asp Gly Pro Asp Gln Met Ala Tyr Lys Arg Ala Asn Gly Arg Met Val Ser Lys Pro Ala Thr Met Arg <210> SEQ ID NO 117 <211> LENGTH: 369 <212> TYPE: PRT <213 > ORGANISM: Aspergillus nidulans <400> SEQUENCE: 117 Met Tyr Ala Val Glu Asp Arg Ala His Ser Gly His His Pro Pro Pro Leu Ser Met Asp Arg Ile Pro Pro Pro Ser Thr Met Tyr Pro Ser Ser Ala Gly Pro Ser Ala Met Val Ser Pro Ala Gly Gln Pro Glu Pro Glu Ser Leu Ser Thr Val His Asp Gly Arg Ile Trp Ser Leu Gln Val Val Gln Gln Pro Ile Arg Ala Arg Met Cys Gly Phe Gly Asp Lys Asp Arg 65 70 75 80Arg Pro Ile Thr Pro Pro Pro Cys Ile Arg Leu Ile Val Lys Asp Ala Gln Thr Gln Lys Glu Val Asp Ile Asn Ser Leu Asp Ser Ser Phe Tyr Val Val Met Ala Asp Leu Trp Asn Ala Asp Gly Thr His Glu Val Asn 120 Leu Val Lys His Ser Ala Thr Ser Pro Ser Ile Ser Thr Ala Met Ser Ser Ser Tyr Pro Pro Pro Pro His Pro Thr Ser Ser Asp Tyr Pro Ala Ser Tyr Gln Thr Asn Pro Tyr Gly Gln Pro Val Gly Gln Pro Val Gly Gln Pro Val Gly Tyr Ala Gly Val Gly Asn Tyr Tyr Gly Gly Ser Thr Gln Leu Gln Tyr Gln Asn Ala Tyr Pro Asn Pro Gln Ala Gln Tyr Tyr ${\tt Gln\ Pro\ Met\ Tyr\ Gly\ Gly\ Met\ Ala\ Gln\ Pro\ Gln\ Met\ Pro\ Ala\ Ala\ Gln}$ Pro Val Thr Pro Gly Pro Gly Gly Met Phe Thr Arg Asn Leu Ile Gly Cys Leu Ser Ala Ser Ala Tyr Arg Leu Tyr Asp Thr Glu Asp Lys Ile Gly Val Trp Phe Val Leu Gln Asp Leu Ser Val Arg Thr Glu Gly Ile 265 Phe Arg Leu Lys Phe Ser Phe Val Asn Val Gly Lys Ser Val Ser Asp 280

```
Leu Pro Gln Ser Asp Ile Ala Glu Val Ile Asn Lys Gly Thr Ala Pro
                        295
Ile Leu Ala Ser Thr Phe Ser Glu Pro Phe Gln Val Phe Ser Ala Lys
305
                    310
                                        315
Lys Phe Pro Gly Val Ile Glu Ser Thr Pro Leu Ser Lys Val Phe Ala
Asn Gln Gly Ile Lys Ile Pro Ile Arg Lys Asp Gly Val Lys Gly Gln
                                345
Gly Ser Arg Gly Arg His Ser Asp Glu Asp Asp Gly Leu Asp Asn Glu
Tyr
<210> SEQ ID NO 118
<211> LENGTH: 20
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 118
                                                                      20
acaaccctgg actctggaat
<210> SEQ ID NO 119
<211> LENGTH: 51
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223 > OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 119
cgaagagggt gaagagcatt gtttgaggca gaggacgcgt tgactgtgat g
                                                                      51
<210> SEQ ID NO 120
<211> LENGTH: 50
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 120
tgacgacaat acctcccgac gatacctggg ttgattcctg cttttcctcc
                                                                      50
<210> SEQ ID NO 121
<211> LENGTH: 20
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 121
tctcgttctc ccatttacct
                                                                       20
<210> SEQ ID NO 122
<211> LENGTH: 20
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 122
tgcctcaaac aatgctcttc
                                                                      20
```

```
<210> SEQ ID NO 123
<211> LENGTH: 19
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223 > OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 123
caaggtatcg tcgggaggt
                                                                        19
<210> SEQ ID NO 124
<211> LENGTH: 20
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
<223 > OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 124
                                                                        20
aatcacggac ctcgaagcag
<210> SEQ ID NO 125
<211> LENGTH: 20
<212> TYPE: DNA
<213 > ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 125
                                                                        20
ggggtcttga tatggcgaat
<210> SEQ ID NO 126
<211> LENGTH: 22
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 126
caacaagacc gacatcacct tc
                                                                        2.2
<210> SEQ ID NO 127
<211> LENGTH: 22
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 127
                                                                        22
ccattcttgg gatagctgca ac
<210> SEQ ID NO 128
<211> LENGTH: 32
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
<400> SEQUENCE: 128
                                                                        32
caacgaacta gtccgcctgc ccttaacctc ca
<210> SEQ ID NO 129
<211> LENGTH: 32
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic oligonucleotide
```

-continued

<400> SEQUENCE: 129		
gcatacacta gtctcgcatg ccagtggatg gg	32	
<210> SEQ ID NO 130 <211> LENGTH: 21 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide		
<400> SEQUENCE: 130		
categgttga ctaegetege a	21	
<210> SEQ ID NO 131 <211> LENGTH: 21 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide		
<400> SEQUENCE: 131	21	
gacctgtggt gaaacctgag g <210> SEQ ID NO 132	21	
<211> LENGTH: 24 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide		
<400> SEQUENCE: 132		
ctagctggtc attatttgat ctcg	24	
<210> SEQ ID NO 133 <211> LENGTH: 24 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide		
<400> SEQUENCE: 133		
gttgtagagt ggacgatcat catg	24	
<210> SEQ ID NO 134 <211> LENGTH: 26 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide		
<400> SEQUENCE: 134		
ccttgtatga tgtatgtatg atgagc	26	
<210> SEQ ID NO 135 <211> LENGTH: 24 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide <400> SEQUENCE: 135		
gacagcgaaa gtgaagagga catc	24	
<210> SEQ ID NO 136		

<211> LENGTH: 20

-continued	
<pre><212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE:</pre>	
<223> OTHER INFORMATION: Synthetic oligonucleotide	
<400> SEQUENCE: 136	
gaageggtet gaateteetg 20	
<210> SEQ ID NO 137 <211> LENGTH: 20 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide	
<400> SEQUENCE: 137	
acagtecaag egtggtatee 20	
<210> SEQ ID NO 138 <211> LENGTH: 22 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide	
<400> SEQUENCE: 138	
agagtettee tteageragg te 22	2
<210> SEQ ID NO 139 <211> LENGTH: 22 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic oligonucleotide	
<400> SEQUENCE: 139	
gtggggcttt tcttcattct cg 22	2
<210> SEQ ID NO 140 <211> LENGTH: 3662 <212> TYPE: DNA <213> ORGANISM: Aspergillus flavus	
<400> SEQUENCE: 140	
acctcacage tggctgagtt eccaetggat eggecattet tettttttta ecceetette 60	
tgtgacacca tccaccttcc cccattttag tgccatccgt gctgacccaa tacccaccat 120	
atoggotoag togtgoagag ttagtgoogt ogtacataco agotocatta gaaggtactg 180)
tacatacgcc atgtatctac ggggtaattc cggttgccgt cgtcagaacg gataaacatc 240)
tcctattgac cctctcgata agaaagagca agttcagagc gaataaaatg ctcgaagcgc 300)
agaaatgcga ccgtagaccc atggtaccgg gatatttaag aacgcaagtt atgtatgggg 360	
ccgtcagccc gatcaccatg ttgttggaac ttggaaccga tggaaaccgt caccgtcggt 420)
aatcttacag taccacggct ccggagacga gacggttagg gtgttatagt tggttgcgat 480)
agtgtgtgta cctgccgatt ctcaggctga ctcgtccttt gacttttccc gtcctcgctc 540)
getecacece eteteattea trattetteg tecececegt ateaceaact tgaettttgt 600)
tactectete ceatacette ttteeteate etggaegggt caategetgg attgtetgtt 660)
gtactgcctc ggcgcagcgg acccagttta aattttttt ttttttaatt tcattatcct 720)
tetttteece ttttggtega eagttggaat etetetegtt tteegttteg eagacgaett 780	

cgagaacatc	cttgaccgac	tggtcggttt	cacggcttgc	attcgtggga	gtcacgcccg	840	_
acataaccag	cacactagaa	gaaaaccaga	ttcacacggg	aaacggggat	agattacttg	900	
cttgacgcaa	gcactttccc	atagcgttga	ttttgtttgg	ttggaccgag	ggattgccaa	960	
gtaagtccca	agcttcattg	ctagctggtc	attatttgat	ctcgaccatg	aaattgtggt	1020	
caattacgcc	cgtcgaatct	tgttgtgcta	atcctggcag	tttttcgcat	cacagtcaag	1080	
gcgtctccaa	aatggcgaca	cgagctcctt	tggcgcctcc	gccgaacgag	acggaagcct	1140	
ccgtcagccg	gatcactcga	gagggcaaga	agctcaccta	taaactcaat	gtcatgcaac	1200	
agcctgagcg	tgcgcgagcc	tgcggtgcag	gtgcaaagtg	tatgcgtcca	accagcactc	1260	
cataaccgga	tagcatcaaa	ctgatggttc	tgactttata	tagcctctgc	ggaccgtcgt	1320	
ccagtcgatc	ctccaccggt	cgtcgaactt	cgagtgtacg	agtccgatcc	caacgacgac	1380	
ctcaacaaga	ccgacatcac	cttcgcatac	aacgccaatt	tetteetgta	cgccactttg	1440	
gaaaccgctc	gtcccatggc	ccaaggccgt	tttgccccga	atccgacttg	tccagtattg	1500	
accggtgtgc	ccgtggctgg	agtggcttac	ttggaccgcc	catctcaagc	cggttacttc	1560	
atcttccccg	atctttccgt	gcggcatgaa	ggtgtatatc	gattgaactt	ccacctgtac	1620	
gaggaaacca	aggagagcaa	ggatgcgaac	gagaatgctc	cgatccagtc	catgtccaac	1680	
ccaatgccat	cgaagccgat	ggcgccgaag	tcattcctgg	agtttcgtct	cgaggtcgtt	1740	
tccgttccgt	tcaccgtatt	taacgccaag	aagttcccag	gattggccac	gagtacctcc	1800	
ctgagtcggg	tcattgcgga	gcaaggttgt	cgtgtgcgga	ttcgacgtga	tgtccgcatg	1860	
agacgtcggg	gagagaageg	caccgatgac	tacgactacg	atgaggagag	agtctaccga	1920	
tcttctgacc	gaatctctac	cccagatacc	cacgggtacg	ccggcactcc	cgttgaacgt	1980	
cctcgatcaa	ccagtaccag	cacggtggat	ccctcattcc	cctacggtgt	cgatgctcag	2040	
cgccggtcat	ctggcgcgac	cgagtatggt	ttccagggtg	cacagccgta	ccaacgacca	2100	
ttgccgcctg	ctccgggtcc	cgcaccagcc	gctgtttcca	cgcccgctcc	tcccgctcct	2160	
cccgcgccac	catcccataa	tcctggatat	caatcgcatc	tttcctttgg	ctcgactcaa	2220	
actcaatatc	cageteecea	gctgcctcca	actccacaga	ccgcgtcgac	attggcagct	2280	
ccgtactcgc	cccatccatc	gtattctcat	gctcggaatc	catcgacgag	ggccgagtat	2340	
gaaacgcccg	gttactccta	teegecatea	cggatgtcaa	cggaacgttc	cagctatccc	2400	
aagaatggct	tgcctccgct	ccgcttggaa	ccgcctaagc	cactaaatat	gccatccggc	2460	
gagccacgct	cgtccgatct	gaacgcatat	cattccgtgg	ctcaatcggc	ggcaccccgg	2520	
tctcagacac	cgtcatccag	tetggtgeet	tecetteege	ccctcaaggc	tctatcgggg	2580	
gattatccca	acaacctctc	tcaatcatcc	agcagtacct	ctcagagece	cagtcacgat	2640	
ctcggcgctg	gcaagaagtt	cttctgggat	acgggcgcca	gcctgtccaa	gcggtcgtac	2700	
gaagattegt	ttggccatga	tgatcgtcca	ctctacaacg	gcatgcgccc	cgatacggaa	2760	
agttatcctc	ggaggctgtc	agatgccagt	cggaacttct	acaacgaaac	gcgcgatgaa	2820	
atggcgtaca	aacgagccaa	cgggagaatg	gccacgaaga	tatcccctgc	actccagtaa	2880	
aacaagttga	ttcctgcttt	tcctcccgct	catatagtga	cggcgtcttg	gcgtaacggt	2940	
cgtcgattga	tttctttccc	gtaatctgtt	ccttttccta	atgtactctg	gtgtgatggg	3000	
cttcagggac	tcttttaacg	acccagactt	ttgatgttta	taccaccgtt	ctttttcttc	3060	
tttcctcgat	ctttggcatt	attgtacatg	atgctctgca	tgtggttttc	aagatattcc	3120	
ccggattgtt	cttgtcttca	gtttatatac	ggccgctctc	gtgtttatta	tccgctgtgt	3180	
-							

-continued

ttccaggtcg	gctggacctg	gggcctctcc	cttcccgcga	atagaagtga	gtgagcaata	3240
caaatgtgac	attgtccaaa	agtttggtga	tetgaaegeg	caacctggat	gcattgatcc	3300
gagacaatca	cggggtctta	gacatgcgac	atgtctgatt	cactccttcg	accatttcct	3360
tgtttatcca	tgaccatgcc	ccatccactg	gcatgcgaga	atgacgtatg	cgacagataa	3420
gatcgacgat	ctgccttata	tatccgaatt	gatccgattg	tcaatactct	ctcttagtgt	3480
tgtataagta	tatatatgtg	ctgtagagta	tgtcctggct	gtctcccata	cagaagaagc	3540
catgtcgaga	aagggtatgt	ccaccagagt	aagattgtac	attccttgga	catgtcattg	3600
tcatttcaca	ggaaatatcg	aagggtcatg	gattcggagg	aacattccag	ggaagagaca	3660
cc						3662
<210> SEQ I <211> LENGT <212> TYPE: <213> ORGAN	H: 4406 DNA	gillus flavı	ls			
<400> SEQUE	NCE: 141					
aatcacggac	ctcgaagcag	tgggccttct	ccaccctcga	actgcctgtt	caagttgtta	60
ggtttttgta	tttaattat	tattttcttt	ttgtctgaat	tttctcacaa	cttatttgtt	120
atgcagccaa	tgtctaaacc	ccaccggacc	caatgaccaa	gccggccgca	gtcgccatga	180
cccaacgacc	gtaaccgcct	gcccttaacc	tccagctggc	tgagttccca	ctggatcggc	240
cattcttctt	tttttacccc	ctcttctgtg	acaccatcca	ccttccccca	ttttagtgcc	300
atccgtgctg	acccaatacc	caccatatcg	gctcagtcgt	gcagagttag	tgccgtcgta	360
cataccagct	ccattagaag	gtactgtaca	tacgccatgt	atctacgggg	taattccggt	420
tgccgtcgtc	agaacggata	aacatctcct	attgaccctc	tcgataagaa	agagcaagtt	480
cagagcgaat	aaaatgctcg	aagcgcagaa	atgcgaccgt	agacccatgg	taccgggata	540
tttaagaacg	caagttatgt	atggggccgt	cagecegate	accatgttgt	tggaacttgg	600
aaccgatgga	aaccgtcacc	gtcggtaatc	ttacagtacc	acggctccgg	agacgagacg	660
gttagggtgt	tatagttggt	tgcgatagtg	tgtgtacctg	ccgattctca	ggctgactcg	720
tcctttgact	tttcccgtcc	tegetegete	caccccctct	cattcattat	tcttcgtccc	780
ccccgtatca	ccaacttgac	ttttgttact	cctctcccat	accttctttc	ctcatcctgg	840
acgggtcaat	cgctggattg	tctgttgtac	tgcctcggcg	cagcggaccc	agtttaaatt	900
tttttttt	tttaatttca	ttatccttct	tttccccttt	tggtcgacag	ttggaatctc	960
tctcgttttc	cgtttcgcag	acgacttcga	gaacatcctt	gaccgactgg	tcggtttcac	1020
ggcttgcatt	cgtgggagtc	acgcccgaca	taaccagcac	actagaagaa	aaccagattc	1080
acacgggaaa	cggggataga	ttacttgctt	gacgcaagca	ctttcccata	gcgttgattt	1140
tgtttggttg	gaccgaggga	ttgccaagta	agtcccaagc	ttcattgcta	gctggtcatt	1200
atttgatctc	gaccatgaaa	ttgtggtcaa	ttacgcccgt	cgaatcttgt	tgtgctaatc	1260
ctggcagttt	ttcgcatcac	agtcaaggcg	tctctgcctc	aaacaatgct	cttccaaaat	1320
ggcgacacga	gctcctttgg	cgcctccgcc	gaacgagacg	gaagcctccg	tcagccggat	1380
cactcgagag	ggcaagaagc	tcacctataa	actcaaatgt	catgcaacag	cctgagcgtg	1440
cgcgagcctg	cggtgcaggt	gcaaagtgta	tgcgtccaac	cagcactcca	taaccggata	1500
agot googt	astaattata	2011112121	aaatataaaa	agget gat ag	agt agat agt	1560

gcatcaaact gatggttctg actttatata gcctctgcgg accgtcgtcc agtcgatcct 1560

ccaccggtcg	tcgaacttcg	agtgtacgag	tccgatccca	acgacgacct	caacaagacc	1620
gacatcacct	tcgcatacaa	cgccaatttc	ttcctgtacg	ccactttgga	accgctcgtc	1680
ccatggccca	aggccgtttt	gccccgaatc	cgacttgtcc	agtattgacc	ggtgtgcccg	1740
tggctggagt	ggcttacttg	gaccgcccat	ctcaagccgg	ttacttcatc	ttccccgatc	1800
tttccgtgcg	gcatgaaggt	gtatatcgat	tgaacttcca	cctgtacgag	gaaaccaagg	1860
agagcaagga	tgcgaacgag	aatgctccga	tccagtccca	tgtccaaccc	aatgccatcg	1920
aagcccgatg	gcgccgaagt	cattcctgga	gtttcgtctc	gaggtcgttt	ccgttccgtt	1980
caccgtattt	agcgccaaga	agttcccagg	attggccacg	agtacctccc	tgagtcgggt	2040
cattgcggag	caaggttgtc	gtgtgcggat	tcgacgtgat	gtccgcatga	gacgtcgggg	2100
agagaagcgc	accgatgact	acgactacga	tgaggagaga	gtctaccgat	cttctgaccg	2160
aatctctacc	ccagataccc	acgggtacgc	cggcactccc	gttgaacgtc	ctcgatcaac	2220
cagtaccagc	acggtggatc	cctcattccc	ctacggtgtc	gatgctcagc	gccggtcatc	2280
tggcgcgacc	gagtatggtt	tccagggtgc	acagccgtac	caacgaccat	tgccgcctgc	2340
teegggteee	gcaccagccg	ctgtttccac	gecegeteet	cccgctcctc	ccgcgccacc	2400
atcccataat	cctggatatc	aatcgcatct	ttcctttggc	tcgactcaaa	ctcaatatcc	2460
agctccccag	ctgcctccaa	ctccacagac	cgcgtcgaca	ttggcagctc	cgtactcgcc	2520
ccatccatcg	tattctcatg	ctcggaatcc	atcgacgagc	gccgagtatg	aaacgcccgg	2580
ttactcctat	ccgccatcac	ggatgtcaac	ggaacgttcc	agctatccca	agaatggctt	2640
geeteegete	cgcttggaac	cgcctaagcc	actaaatatg	ccatccggcg	agccacgctc	2700
gtccgatccg	aacgcatatc	attccgtggc	tcaatcggcg	gcaccccggt	ctcagacacc	2760
gtcatccagt	ctggtgcctt	cccttccgcc	cctcaaggct	ctatcggggg	attatcccaa	2820
caacctctct	caatcatcca	gcagtacctc	tcagagcccc	agtcacgatc	tcggcgctgg	2880
caagaagttc	ttctgggata	cgggcgccag	cctgtccaag	cggtcgtacg	aagattcgtt	2940
tggccatgat	gatcgtccac	tctacaacgg	catgcgcccc	gatacggaaa	gttatcctcg	3000
gaggctgtca	gatgccagtc	ggaacttcta	caacgaaacg	cgcgatgaaa	tggcgtacaa	3060
acgagccaac	gggagaatgg	ccacgaagat	atcccctgca	ctccagtaaa	acaagttgat	3120
tcctgctttt	cctcccgctc	atataggacg	gcgtcttggc	gaacggtcgt	cgattgattt	3180
ctttcccgta	atctgttcct	tttcctaatg	tactctggtg	tgatgggctt	cagggactct	3240
tttaacgacc	cagacttttg	atgtttatac	caccgttctt	tttcttcttt	cctcgatctt	3300
tggcattatt	gtacatgatg	ctctgcatgt	ggttttcaag	atattccccg	gattgttctt	3360
gtcttcagtt	tatatacggc	egetetegtg	tttattatcc	gctgtgtttc	caggtcggct	3420
ggacctgggg	cctctccctt	cccgcgaata	gaagtgagtg	agcaatacaa	atgtgacatt	3480
gtccaaaagt	ttggtgatct	gaacgcgcaa	cctggatgca	ttgatccgag	acaatcacgg	3540
ggtcttagac	atgcgacatg	tctgattcac	tccttcgacc	atttccttgt	ttatccatga	3600
ccatgcccca	tccactggca	tgcgagaatg	acgtatgcga	cagataagat	cgacgatctg	3660
ccttatatat	ccgaattgat	tcgattgtca	atactctctc	ttagtgttgt	ataagtatat	3720
atatgtgctg	tagagtatgt	cctggctgtc	tcccatacag	aagaagccat	gtcgagaaag	3780
	ccagagtaag					3840
	ggtcatggat					3900
	gtaaacatgt					3960
goucygated	gradacatyt	agacciacat	adectigite	accecatya	cycactyaca	3,700

-continued

ctagagcata	gaggcgttta	ctttacctct	tatcaacgac	taattgacat	aaccaagcag	4020
tggaatttt	atgattcaac	gtcccagact	atagtgccta	ctgttatgat	ctacctgcta	4080
ctttgatcgg	tttcactttc	tttgttggtt	gatacttgtt	gcgtatcttc	tttgattact	4140
tatagtcaat	agtccctgtt	actatattag	cgccgttcct	agccattgta	tatcctttga	4200
tccactactt	gaaagaaaat	tcgaacaagg	gttttaatgg	atgtatccat	tctattttct	4260
cttttctttg	gatgtaatga	gatacaaagg	tgttctaatc	aaaaagtatg	tattcaattc	4320
aaatacatct	teccagacaa	tetgggaaga	ggaataagca	agagagataa	ttctccgact	4380
cacactattc	gccatatcaa	gacccc				4406
<220> FEATURE CONTROL	TH: 4556 : DNA NISM: Artif: JRE:	ON: Aspergi	nce llus flavus	veA knockou	ut with	
<400> SEQUI	ENCE: 142					
aatcacggac	ctcgaagcag	tgggccttct	ccaccctcga	actgcctgtt	caagttgtta	60
ggtttttgta	ttttaattat	tattttcttt	ttgtctgaat	tttctcacaa	cttatttgtt	120
atgcagccaa	tgtctaaacc	ccaccggacc	caatgaccaa	gccggccgca	gtcgccatga	180
cccaacgacc	gtaaccgcct	gcccttaacc	tccagctggc	tgagttccca	ctggatcggc	240
cattcttctt	tttttacccc	ctcttctgtg	acaccatcca	ccttccccca	ttttagtgcc	300
atccgtgctg	acccaatacc	caccatatcg	gctcagtcgt	gcagagttag	tgccgtcgta	360
cataccagct	ccattagaag	gtactgtaca	tacgccatgt	atctacgggg	taattccggt	420
tgccgtcgtc	agaacggata	aacatctcct	attgaccctc	tcgataagaa	agagcaagtt	480
cagagcgaat	aaaatgctcg	aagcgcagaa	atgcgaccgt	agacccatgg	taccgggata	540
tttaagaacg	caagttatgt	atggggccgt	cagcccgatc	accatgttgt	tggaacttgg	600
aaccgatgga	aaccgtcacc	gtcggtaatc	ttacagtacc	acggctccgg	agacgagacg	660
gttagggtgt	tatagttggt	tgcgatagtg	tgtgtacctg	ccgattctca	ggctgactcg	720
tcctttgact	tttcccgtcc	tegetegete	caccccctct	cattcattat	tcttcgtccc	780
ccccgtatca	ccaacttgac	ttttgttact	cctctcccat	accttctttc	ctcatcctgg	840
acgggtcaat	cgctggattg	tctgttgtac	tgcctcggcg	cageggaeee	agtttaaatt	900
tttttttt	tttaatttca	ttatccttct	tttccccttt	tggtcgacag	ttggaatctc	960
tctcgttttc	cgtttcgcag	acgacttcga	gaacatcctt	gaccgactgg	tcggtttcac	1020
ggcttgcatt	cgtgggagtc	acgcccgaca	taaccagcac	actagaagaa	aaccagattc	1080
acacgggaaa	cggggataga	ttacttgctt	gacgcaagca	ctttcccata	gcgttgattt	1140
tgtttggttg	gaccgaggga	ttgccaagta	agtcccaagc	ttcattgcta	gctggtcatt	1200
atttgatctc	gaccatgaaa	ttgtggtcaa	ttacgcccgt	cgaatcttgt	tgtgctaatc	1260
ctggcagttt	ttcgcatcac	agtcaaggcg	tctctgcctc	aaacaatgct	cttcaccctc	1320
ttcgcgggtc	tgaaataccc	tcacctggca	acagcaattg	gcgcttcatg	gctgttttc	1380
cgatctctct	acttgtacgg	ctatgtgtac	tcgggtaagc	cacaaggcaa	gggcagattg	1440
ctgggaggtt	tcttctggtt	ttctcaaggc	gctctgtggg	ctctgagtgt	gtttggtgtt	1500

gccaaagaca tgatctctta ctgagagtta ttctgtgtct gacgaaatat gttgtgtata 1560

tatatatatg	tacgttaaaa	gttccgtgga	gttaccagtg	attgaccaat	gttttatctt	1620
ctacagttct	gcctgtctac	cccattctag	ctgtacctga	ctacagagta	gtttaattgt	1680
ggttgacccc	acagtcggag	gcggaggaat	acagcaccga	tgtggcctgt	ctccatccag	1740
attggcacgc	aatttttaca	cgcggaaaag	atcgagatag	agtacgactt	taaatttagt	1800
ccccggcggc	ttctatttta	gaatatttga	gatttgattc	tcaagcaatt	gatttggttg	1860
ggtcaccctc	aattggataa	tatacctcat	tgctcggcta	cttcaactca	tcaatcaccg	1920
tcataccccg	catataaccc	tccattccca	cgatgtcgtc	caagtcgcaa	ttgacttacg	1980
gtgetegage	cagcaagcac	cccaatcctc	tggcaaagag	actttttgag	attgccgaag	2040
caaagaagac	aaacgttacc	gtctctgctg	atgtgacgac	aacccgagaa	ctcctggacc	2100
tegetgaeeg	tacggaagct	gttggatcca	atacatatgc	cgtctagcaa	tggactaatc	2160
aacttttgat	gatacaggtc	tcggtcccta	catcgccgtc	atcaagacac	acatcgacat	2220
cctcaccgat	ttcagcgtcg	acactatcaa	tggcctgaat	gtgctggctc	aaaagcacaa	2280
ctttttgatc	ttcgaggacc	gcaaattcat	cgacatcggc	aataccgtcc	agaagcaata	2340
ccacggcggt	gctctgagga	tctccgaatg	ggcccacatt	atcaactgca	gcgttctccc	2400
tggcgagggc	atcgtcgagg	ctctggccca	gaccgcatct	gcgcaagact	tcccctatgg	2460
tcctgagaga	ggactgttgg	tcctggcaga	gatgacctcc	aaaggatcgc	tggctacggg	2520
cgagtatacc	aaggcatcgg	ttgactacgc	tcgcaaatac	aagaacttcg	ttatgggttt	2580
cgtgtcgacg	cgggccctga	cggaagtgca	gtcggatgtg	tcttcagcct	cggaggatga	2640
agatttcgtg	gtcttcacga	cgggtgtgaa	cctctctcc	aaaggagata	agcttggaca	2700
gcaataccag	actcctgcat	cggctattgg	acgeggtgee	gactttatca	tegeeggteg	2760
aggcatctac	getgeteeeg	acccggttga	agctgcacag	cggtaccaga	aagaaggetg	2820
ggaagcttat	atggccagag	tatgcggcaa	gtcatgattt	cctcttggag	caaaagtgta	2880
gtgccagtac	gagtgttgtg	gaggaaggct	gcatacattg	tgcctgtcat	taaacgatga	2940
gctcgtccgt	attggcccct	gtaatgccat	gttttccgcc	cccaatcgtc	aaggttttcc	3000
ctttgttaga	ttcctaccag	tcatctagca	agtgaggtaa	gctttgccag	aaacgccaag	3060
gctttatcta	tgtagtcgat	aagcaaagtg	gactgatagc	ttaatatgga	aggtccctca	3120
ggacaagtcg	acctgtgcag	aagagataac	agcttggcat	cacgcatcag	tgcctcctct	3180
cagacagaat	aagtcgccat	aagttatcga	ccgaacctag	gtagggtata	tttggtgacg	3240
acaatacctc	ccgacgatac	ctgggttgat	tcctgctttt	cctcccgctc	atataggacg	3300
gcgtcttggc	gaacggtcgt	cgattgattt	ctttcccgta	atctgttcct	tttcctaatg	3360
tactctggtg	tgatgggctt	cagggactct	tttaacgacc	cagacttttg	atgtttatac	3420
caccgttctt	tttcttcttt	cctcgatctt	tggcattatt	gtacatgatg	ctctgcatgt	3480
ggttttcaag	atattccccg	gattgttctt	gtcttcagtt	tatatacggc	cgctctcgtg	3540
tttattatcc	gctgtgtttc	caggtcggct	ggacctgggg	cctctccctt	cccgcgaata	3600
gaagtgagtg	agcaatacaa	atgtgacatt	gtccaaaagt	ttggtgatct	gaacgcgcaa	3660
cctggatgca	ttgatccgag	acaatcacgg	ggtcttagac	atgcgacatg	tctgattcac	3720
teettegace	atttccttgt	ttatccatga	ccatgcccca	tccactggca	tgcgagaatg	3780
acgtatgcga	cagataagat	cgacgatctg	ccttatatat	ccgaattgat	tcgattgtca	3840
atactctctc	ttagtgttgt	ataagtatat	atatgtgctg	tagagtatgt	cctggctgtc	3900

-continued

60

tcccatacag	aagaagccat	gtcgagaaag	ggtatgtcca	ccagagtaag	attgtacatt	3960
ccttggacat	gtcattgtca	tttcacagga	aatatcgaag	ggtcatggat	tcggaggaac	4020
attccaggaa	gagacaccag	aaatgagttg	gcacggatca	gtaaacatgt	agatctacat	4080
aaccttgttc	attttcatga	tgcactgaca	ctagagcata	gaggcgttta	ctttacctct	4140
tatcaacgac	taattgacat	aaccaagcag	tggaattttt	atgattcaac	gtcccagact	4200
atagtgccta	ctgttatgat	ctacctgcta	ctttgatcgg	tttcactttc	tttgttggtt	4260
gatacttgtt	gcgtatcttc	tttgattact	tatagtcaat	agtccctgtt	actatattag	4320
cgccgttcct	agccattgta	tatcctttga	tccactactt	gaaagaaaat	tcgaacaagg	4380
gttttaatgg	atgtatccat	tctattttct	cttttctttg	gatgtaatga	gatacaaagg	4440
tgttctaatc	aaaaagtatg	tattcaattc	aaatacatct	tcccagacaa	tctgggaaga	4500
ggaataagca	agagagataa	ttctccgact	cacactattc	gccatatcaa	gacccc	4556

<210> SEQ ID NO 143

<400> SEOUENCE: 143

agequecaat acquaaaccq cetetececq eqeqttqqcc qattcattaa tqcaqetqqc acgacaggtt tcccgactgg aaagcgggca gtgagcgcaa cgcaattaat gtgagttagc 120 tcactcatta ggcaccccag gctttacact ttatgcttcc ggctcgtatg ttgtgtggaa 180 ttgtgagcgg ataacaattt cacacaggaa acagctatga ccatgattac gccaagcttg 240 gtaccgagct cggatccact agtccgcctg cccttaacct ccagctggct gagttcccac 300 tggatcggcc attettettt ttttaccccc tettetgtga caccatecae etteccccat 360 tttagtgcca tccgtgctga cccaataccc accatatcgg ctcagtcgtg cagagttagt 420 gccgtcgtac ataccagctc cattagaagg tactgtacat acgccatgta tctacggggt 480 aattccggtt gccgtcgtca gaacggataa acatctccta ttgaccctct cgataagaaa 540 gagcaagttc agagcgaata aaatgctcga agcgcagaaa tgcgaccgta gacccatggt 600 accgggatat ttaagaacgc aagttatgta tggggccgtc agcccgatca ccatgttgtt 660 ggaacttgga accgatggaa accgtcaccg tcggtaatct tacagtacca cggctccgga 720 gacgagacgg ttagggtgtt atagttggtt gcgatagtgt gtgtacctgc cgattctcag gctgactcgt cctttgactt ttcccgtcct cgctcgctcc accccctctc attcattatt 840 cttcgtcccc cccgtatcac caacttgact tttgttactc ctctcccata ccttctttcc tcatcctgga cgggtcaatc gctggattgt ctgttgtact gcctcggcgc agcggaccca 960 qtttaaattt ttttttttt ttaatttcat tatccttctt ttcccctttt qqtcqacaqt 1020 tggaatctct ctcgttttcc gtttcgcaga cgacttcgag aacatccttg accgactggt 1080 cggtttcacg gcttgcattc gtgggagtca cgcccgacat aaccagcaca ctagaagaaa 1140 accagattca cacgggaaac ggggatagat tacttgcttg acgcaagcac tttcccatag 1200 cgttgatttt gtttggttgg accgagggat tgccaagtaa gtcccaagct tcattgctag ctggtcatta tttgatctcg accatgaaat tgtggtcaat tacgcccgtc gaatcttgtt 1320 gtgctaatcc tggcagtttt tcgcatcaca gtcaaggcgt ctccaaaatg gcgacacgag 1380

<211> LENGTH: 9329

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: Aspergillus flavus multicopy veA strain with Aspergillus fumigatus pyrG in TOPO-TA cloning plasmid

ctcctttggc	gcctccgccg	aacgagacgg	aagcctccgt	cagccggatc	actcgagagg	1440
gcaagaagct	cacctataaa	ctcaatgtca	tgcaacagcc	tgagcgtgcg	cgagcctgcg	1500
gtgcaggtgc	aaagtgtatg	cgtccaacca	gcactccata	accggatagc	atcaaactga	1560
tggttctgac	tttatatagc	ctctgcggac	cgtcgtccag	tegateetee	accggtcgtc	1620
gaacttcgag	tgtacgagtc	cgatcccaac	gacgacctca	acaagaccga	catcaccttc	1680
gcatacaacg	ccaatttctt	cctgtacgcc	actttggaaa	ccgctcgtcc	catggcccaa	1740
ggccgttttg	ccccgaatcc	gacttgccca	gtattgaccg	gtgtgcccgt	ggctggagtg	1800
gcttacttgg	accgcccttc	tcaagccggt	tacttcatct	teccegatet	ttccgtgcgg	1860
catgaaggtg	tatatcgatt	gaacttccac	ctgtacgagg	aaaccaagga	gagcaaggat	1920
gcgaacgaga	atgctccgat	ccagtccctg	tccaacccaa	tgccatcgaa	gccgatggcg	1980
ccgaagtcat	teetggagtt	tegtetegag	gtcgtttccg	ttccgttcac	cgtatttagc	2040
gccaagaagt	tcccaggatt	ggccacgagt	acctccctga	gtcgggtcat	tgcggagcaa	2100
ggttgtcgtg	tgcggattcg	acgtgatgtc	cgcatgagac	gtcggggaga	gaagegeace	2160
gatgactacg	actacgatga	ggagagagtc	taccgatctt	ctgaccgaat	ctctacccca	2220
gatacccacg	ggtacgccgg	cactcccgtt	gaacgtcctc	gatcaaccag	taccagcacg	2280
gtggatccct	cattccccta	cggtgtcgat	gctcagcgcc	ggtcatctgg	cgcgaccgag	2340
tatggtttcc	agggtgcaca	gccgtaccaa	cgaccattgc	egeetgetee	cggtcccgca	2400
ccagccgctg	tttccacgcc	egeteeteee	geteeteeeg	cgccaccatc	ccataatcct	2460
ggatatcaat	cgcatctttc	ctttggctcg	actcaaactc	aatatccagc	tccccaactg	2520
cctccaactc	cacagaccgc	gtcgacattg	gcagctccgt	actegeecea	tccatcgtat	2580
teteatgete	ggaatccatc	gacgagcgcc	gagtatgaaa	cgcccggtta	ctcctatccg	2640
ccatcacgga	tgtcaacgga	acgttccagc	tatcccaaga	atggcttgcc	teegeteege	2700
ttggaaccgc	ctaagccact	aaatatgcca	tccggcgagc	cacgctcgtc	cgatccgaac	2760
gcatatcatt	ccgtggctca	atcggcggca	ccccggtctc	agacaccgtc	atccagtctg	2820
gtgccttccc	ttccgcccct	caaggctcta	tcgggggatt	atcccaacaa	cctctctcaa	2880
tcatccagca	gtacctctca	gagccccagt	cacgateteg	gcgctggcaa	gaagttette	2940
tgggatacgg	gcgccagcct	gtccaagcgg	tcgtacgaag	attcgtttgg	ccatgatgat	3000
cgtccactct	acaacggcat	gcgccccgat	acggaaagtt	atcctcggag	gctgtcagat	3060
gccagtcgga	acttctacaa	cgaaacgcgc	gatgaaatgg	cgtacaaacg	agccaacggg	3120
agaatggcca	cgaagatatc	ccctgcactc	cagtaaaaca	agttgattcc	tgcttttcct	3180
cccgctcata	taggacggcg	tcttggcgaa	cggtcgtcga	ttgatttctt	tecegtaate	3240
tgttcctttt	cctaatgtac	tctggtgtga	tgggcttcag	ggactctttt	aacgacccag	3300
acttttgatg	tttataccac	cgttctttt	cttctttcct	cgatctttgg	cattattgta	3360
catgatgctc	tgcatgtggt	tttcaagata	ttccccggat	tgttcttgtc	ttcagtttat	3420
atacggccgc	tctcgtgttt	attatccgct	gtgtttccag	gtcggctgga	cctggggcct	3480
ctcccttccc	gcgaatagaa	gtgagtgagc	aatacaaatg	tgacattgtc	caaaagtttg	3540
gtgatctgaa	cgcgcaacct	ggatgcattg	atccgagaca	atcacggggt	cttagacatg	3600
	gattcactcc					3660
	gagactagta					3720
	gtcaccaaat					3780
aggractytt	geodecadat	acaccccacc	Laggeregge	cyacaactta	cyycyactta	5700

ttctgtctga gaggaggcac	tgatgcgtga	taccaaacta	ttatctcttc	tgcacaggtc	3840
gacttgtcct gagggacctt					3900
agataaagcc ttggcgtttc					3960
ctaacaaagg gaaaaccttg					4020
cggacgaget categtttaa					4080
tactggcact acacttttgc					4140
taagetteee ageettettt					4200
tagatgcctc gaccggcgat					4260
tggtattgct gtccaagctt					4320
acgaaatctt catcctccga					4380
gtcgacacga aacccataac					4440
gtatactcgc ccgtagccag					4500
ctctcaggac cataggggaa					4560
ccctcgccag ggagaacgct					4620
ccgccgtggt attgcttctg					4680
atcaaaaagt tgtgcttttg					4740
tcggtgagga tgtcgatgtg					4800
tcaaaagttg attagtccat					4860
ggtcagcgag gtccaggagt					4920
tettetttge tteggeaate					4980
ctcgagcacc gtaagtcaat					5040
ggggtatgac ggtgattgat					5100
agggtgaccc aaccaaatca					5160
ccgccggga ctaaatttaa					5220
cgtgccaatc tggatggaga					5280
					5340
gggtcaacca caattaaact					5400
gaactgtaga agataaaaca atatatatat atacacaaca					5460
					5520
gtctttggca acaccaaaca					5580
acctcccagc aatctgccct gagagatcgg aaaaacagcc					5640
					5700
acccgcgaag agggtgaaga					
attetgeaga tateeateae					5760
gccctatagt gagtcgtatt					5820
aaaccctggc gttacccaac	ttaatcgcct	tgcagcacat	ccccctttcg	ccagctggcg	5880
taatagcgaa gaggcccgca	ccgatcgccc	ttcccaacag	ttgcgcagcc	tgaatggcga	5940
atggacgcgc cctgtagcgg	cgcattaagc	gcggcgggtg	tggtggttac	gcgcagcgtg	6000
accgctacac ttgccagcgc	cctagcgccc	gctcctttcg	ctttcttccc	ttcctttctc	6060
gccacgttcg ccggctttcc	ccgtcaagct	ctaaatcggg	ggctcccttt	agggttccga	6120

tttagtgctt	tacggcacct	cgaccccaaa	aaacttgatt	agggtgatgg	ttcacgtagt	6180
gggccatcgc	cctgatagac	ggtttttcgc	cctttgacgt	tggagtccac	gttctttaat	6240
agtggactct	tgttccaaac	tggaacaaca	ctcaacccta	tctcggtcta	ttcttttgat	6300
ttataaggga	ttttgccgat	tteggeetat	tggttaaaaa	atgagctgat	ttaacaaaaa	6360
tttaacgcga	attttaacaa	aattcagggc	gcaagggctg	ctaaaggaag	cggaacacgt	6420
agaaagccag	teegeagaaa	cggtgctgac	cccggatgaa	tgtcagctac	tgggctatct	6480
ggacaaggga	aaacgcaagc	gcaaagagaa	agcaggtagc	ttgcagtggg	cttacatggc	6540
gatagctaga	ctgggcggtt	ttatggacag	caagcgaacc	ggaattgcca	gctggggcgc	6600
cctctggtaa	ggttgggaag	ccctgcaaag	taaactggat	ggctttcttg	ccgccaagga	6660
tctgatggcg	caggggatca	agatctgatc	aagagacagg	atgaggatcg	tttcgcatga	6720
ttgaacaaga	tggattgcac	gcaggttctc	cggccgcttg	ggtggagagg	ctattcggct	6780
atgactgggc	acaacagaca	atcggctgct	ctgatgccgc	cgtgttccgg	ctgtcagcgc	6840
aggggcgccc	ggttcttttt	gtcaagaccg	acctgtccgg	tgccctgaat	gaactgcagg	6900
acgaggcagc	gcggctatcg	tggctggcca	cgacgggcgt	tccttgcgca	gctgtgctcg	6960
acgttgtcac	tgaagcggga	agggactggc	tgctattggg	cgaagtgccg	gggcaggatc	7020
tcctgtcatc	ccaccttgct	cctgccgaga	aagtatccat	catggctgat	gcaatgcggc	7080
ggctgcatac	gcttgatccg	gctacctgcc	cattcgacca	ccaagcgaaa	catcgcatcg	7140
agcgagcacg	tactcggatg	gaageeggte	ttgtcgatca	ggatgatctg	gacgaagagc	7200
atcaggggct	cgcgccagcc	gaactgttcg	ccaggctcaa	ggcgcgcatg	cccgacggcg	7260
aggatctcgt	cgtgacccat	ggcgatgcct	gcttgccgaa	tatcatggtg	gaaaatggcc	7320
gcttttctgg	attcatcgac	tgtggccggc	tgggtgtggc	ggaccgctat	caggacatag	7380
cgttggctac	ccgtgatatt	gctgaagagc	ttggcggcga	atgggctgac	cgcttcctcg	7440
tgctttacgg	tategeeget	cccgattcgc	agegeatege	cttctatcgc	cttcttgacg	7500
agttcttctg	aattgaaaaa	ggaagagtat	gagtattcaa	catttccgtg	tcgcccttat	7560
tccctttttt	gcggcatttt	gccttcctgt	ttttgctcac	ccagaaacgc	tggtgaaagt	7620
aaaagatgct	gaagatcagt	tgggtgcacg	agtgggttac	atcgaactgg	atctcaacag	7680
cggtaagatc	cttgagagtt	ttcgccccga	agaacgtttt	ccaatgatga	gcacttttaa	7740
agttctgcta	tgtggcgcgg	tattatcccg	tattgacgcc	gggcaagagc	aactcggtcg	7800
ccgcatacac	tattctcaga	atgacttggt	tgagtactca	ccagtcacag	aaaagcatct	7860
tacggatggc	atgacagtaa	gagaattatg	cagtgctgcc	ataaccatga	gtgataacac	7920
tgcggccaac	ttacttctga	caacgatcgg	aggaccgaag	gagctaaccg	cttttttgca	7980
caacatgggg	gatcatgtaa	ctcgccttga	tcgttgggaa	ccggagctga	atgaagccat	8040
accaaacgac	gagcgtgaca	ccacgatgcc	tgtagcaatg	gcaacaacgt	tgcgcaaact	8100
attaactggc	gaactactta	ctctagcttc	ccggcaacaa	ttaatagact	ggatggaggc	8160
ggataaagtt	gcaggaccac	ttetgegete	ggcccttccg	gctggctggt	ttattgctga	8220
taaatctgga	gccggtgagc	gtgggtctcg	cggtatcatt	gcagcactgg	ggccagatgg	8280
		ttatctacac				8340
		taggtgcctc				8400
		agattgattt				8460
						8520
ggrgaagate	cittigata	atctcatgac	CaaaatCCCCT	caacytgagt	tttegtteea	0320

				-contir	nued	
ctgagcgtca	gaccccgtag	aaaagatcaa	aggatettet	tgagatcctt	tttttctgcg	8580
		_	accgctacca			8640
tcaagagcta	ccaactcttt	ttccgaaggt	aactggcttc	agcagagcgc	agataccaaa	8700
tactgttctt	ctagtgtagc	cgtagttagg	ccaccacttc	aagaactctg	tagcaccgcc	8760
tacatacctc	gctctgctaa	tcctgttacc	agtggctgct	gccagtggcg	ataagtcgtg	8820
tcttaccggg	ttggactcaa	gacgatagtt	accggataag	gcgcagcggt	cgggctgaac	8880
ggggggttcg	tgcacacagc	ccagcttgga	gcgaacgacc	tacaccgaac	tgagatacct	8940
acagcgtgag	ctatgagaaa	gcgccacgct	tcccgaaggg	agaaaggcgg	acaggtatcc	9000
ggtaagcggc	agggtcggaa	caggagagcg	cacgagggag	cttccagggg	gaaacgcctg	9060
gtatctttat	agtcctgtcg	ggtttcgcca	cctctgactt	gagcgtcgat	ttttgtgatg	9120
ctcgtcaggg	gggeggagee	tatggaaaaa	cgccagcaac	geggeetttt	tacggttcct	9180
ggccttttgc	tggccttttg	ctcacatgtt	ctttcctgcg	ttatcccctg	attctgtgga	9240
taaccgtatt	accgcctttg	agtgagctga	taccgctcgc	cgcagccgaa	cgaccgagcg	9300
cagcgagtca	gtgagcgagg	aagcggaag				9329
<210> SEQ ID NO 144 <211> LENGTH: 7814 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Aspergillus flavus laeA knockout with Aspergillus fumigatus pyrG						
<400> SEQUE		cacatactec	cggccgccat	aaccaaaacc	tactaqttaa	60
Dagogaaceg	Jacobacac	Jourgette	- Jyccyccac	Jacogagaco	Jaseageeda	- 0

tageteegat egagtgatae gtaceateet eatgeeagae eteteeeage taggtatega 120 cttagaaccc gccaacatca agcttcaggt cgaacataag ttccaggacg tagtcaacag 180 actaagttgg aaccatgtca cattttcttc aacgagtgaa ttcgtcaccg catccacctt 240 catgaaccct gacatttacg tttgggaacg gagtcacggt tccctggtga agatcctcga 300 gggtcctaga gaagaactgg gcgtcgtgga atggcaccct tctcgcccta tggttgtcgc 360 ttgcggttta gaatctggat gcatctacac atggtcgatt gtgacgcctc aaaaatggtc 420 cgcgctggca cctgattttg gtgaagtcga ggaaaacgtc gagtatgttg agcgcgaaga 480 cgaatttgac gttcaccctg ccgaagaaat tcaccaacgc cggcttgacc aggaagacga 540 agttcctgac gtattaacga tcgagcccca caaaagcggt acggatgagg agatggaatc cttccgcatg cctgtgcttc tagatatttc tgacagcgaa agtgaagagg acatcattgc cgtcggtccc ggaacaatgc ggaggcgtag ccccggcgct ggccgtgact gggccagcgg 720 agatggtgag aaagaaagta ctggaggtag aaacggtacc tcccggggac aaaagggccg 780 ccggcgttaa agtgatatca ttgtatgagt tcacattata gtattagata atttacagga 840 gcatctgtct tgggtcattg ggtgggcggg tgtcttgaag gctatcgagg cgttttggcg 900 aaatacccaa agactggtat ctcacacact tgcatttccg gttcatgtta gctggtagtg 960 agaccttaca aacgatcaca tggataaaat tctttgcatt gtattatata aatttttata 1020 ttctttttaa gtgatatcat gttatgaatc tcgaatatag caataaccat gattggatga 1080 tgtcgtcaca gatagaatga actaaaccgg gtatacagtt atcaacatta cactcaccca 1140 1200 aaacgccata aatatgctgt ctttggcaag cgggaaagca tgcaatgcca gggaatgatg

tggtcctgaa gtgtgatgaa	ggagccacag	ccaggttaag	agcaggcaga	gggcagaggg	1260
cagagggcat gccatgccgt	gtcctattaa	tgacctgcca	gtattctgca	gccccattgg	1320
ttcgtccata ggggcaagag	ctgcatcgcg	atgtatttt	tggacgaaat	gatctttgac	1380
ttgcctgttt ggcctgggta	atttgtataa	ccccttcgc	aagacgtacg	cccgcctcat	1440
gctcggacct atgataatca	aagcaaactc	tgtatatttc	catcaacctt	ctaagtgcta	1500
ctggagtgat acaggcagct	cagtgcatcg	tegggatagg	cgttaattcg	cggcatacgg	1560
tgtctaatcc aggtatcgtc	gggaggtatt	gtcgtcacca	aatataccct	acctaggttc	1620
ggtcgataac ttatggcgac	ttattctgtc	tgagaggagg	cactgatgcg	tgatgccaag	1680
ctgttatctc ttctgcacag	gtcgacttgt	cctgagggac	cttccatatt	aagctatcag	1740
tccactttgc ttatcgacta	catagataaa	gccttggcgt	ttctggcaaa	gcttacctca	1800
cttgctagat gactggtagg	aatctaacaa	agggaaaacc	ttgacgattg	ggggcggaaa	1860
acatggcatt acaggggcca	atacggacga	gctcatcgtt	taatgacagg	cacaatgtat	1920
gcagcettee tecacaacae	tcgtactggc	actacacttt	tgctccaaga	ggaaatcatg	1980
acttgccgca tactctggcc	atataagctt	cccagccttc	tttctggtac	cgctgtgcag	2040
cttcaaccgg gtcgggagca	gcgtagatgc	ctcgaccggc	gatgataaag	teggeacege	2100
gtccaatagc cgatgcagga	gtctggtatt	gctgtccaag	cttatctcct	ttggaagaga	2160
ggttcacacc cgtcgtgaag	accacgaaat	cttcatcctc	cgaggctgaa	gacacateeg	2220
actgcacttc cgtcagggcc	cgcgtcgaca	cgaaacccat	aacgaagttc	ttgtatttgc	2280
gagcgtagtc aaccgatgcc	ttggtatact	cgcccgtagc	cagcgatcct	ttggaggtca	2340
tctctgccag gaccaacagt	cctctctcag	gaccataggg	gaagtettge	gcagatgcgg	2400
tetgggeeag ageetegaeg	atgccctcgc	cagggagaac	gctgcagttg	ataatgtggg	2460
cccattcgga gatcctcaga	gcaccgccgt	ggtattgctt	ctggacggta	ttgccgatgt	2520
cgatgaattt gcggtcctcg	aagatcaaaa	agttgtgctt	ttgagccagc	acattcaggc	2580
cattgatagt gtcgacgctg	aaatcggtga	ggatgtcgat	gtgtgtcttg	atgacggcga	2640
tgtagggacc gagacctgta	tcatcaaaag	ttgattagtc	cattgctaga	cggcatatgt	2700
attggatcca acagcttccg	tacggtcagc	gaggtccagg	agttctcggg	ttgtcgtcac	2760
atcagcagag acggtaacgt	ttgtcttctt	tgcttcggca	atctcaaaaa	gtctctttgc	2820
cagaggattg gggtgcttgc	tggctcgagc	accgtaagtc	aattgcgact	tggacgacat	2880
cgtgggaatg gagggttata	tgcggggtat	gacggtgatt	gatgagttga	agtagccgag	2940
caatgaggta tattatccaa	ttgagggtga	cccaaccaaa	tcaattgctt	gagaatcaaa	3000
tctcaaatat tctaaaatag	aagccgccgg	ggactaaatt	taaagtcgta	ctctatctcg	3060
atcttttccg cgtgtaaaaa	ttgcgtgcca	atctggatgg	agacaggcca	catcggtgct	3120
gtattcctcc gcctccgact	gtggggtcaa	ccacaattaa	actactctgt	agtcaggtac	3180
agctagaatg gggtagacag	gcagaactgt	agaagataaa	acattggtca	atcactggta	3240
actccacgga acttttaacg	tacatatata	tatatacaca	acatatttcg	tcagacacag	3300
aataactctc agtaagagat	catgtctttg	gcaacaccaa	acacactcag	agcccacaga	3360
gcgccttgag aaaaccagaa	gaaacctccc	agcaatctgc	ccttgccttg	tggcttaccc	3420
gagtacacat agccgtacaa	gtagagagat	cggaaaaaca	gccatgaagc	gccaattgct	3480
gttgccaggt gagggtattt	cagacccgcg	aagagggtga	agagcattgt	ttgaggcaat	3540

				0011011		
cactagtggc	cgcgtagtac	gagtcgtgtg	gtggtgaggc	catcgccgga	agacgctgtc	3600
cagtctggcc	gtttccaaac	attccgtgtg	gtgaagaagg	gtcggtaaga	agtttggtac	3660
tttaagatgg	tgacgagggg	aagaatgccg	acgtcgatgg	cgacaagctg	atgtcagtag	3720
cgtggacgct	tgtaggttgt	gtatatggta	tatatattga	gtgacgaata	aaacatagag	3780
tagaatgggg	ctgagcctag	tgaatggacg	gcaagttcat	tcataaattt	aagggagtga	3840
aaagcgaaac	acaagatatc	attcgcagta	ttagaaagag	agacgaggat	tcagggggag	3900
aatcagaaaa	gtaaacagag	aaaaataaga	aaagaaaaga	aaaaaaaatt	aaaatcaaaa	3960
atcaaaaatc	aaaaatcaga	atctaaaaaa	ataagaaaag	ttggagatag	gaccaggcag	4020
agaatcaaag	gtcacgtcca	cttaaacaca	gaagggcaga	aacggaaaaa	ccaagacggt	4080
gaccagccaa	aagtaccgac	aggcaaagag	ataataatag	agaacgccgt	gctgtctccc	4140
cgcgatgaag	cgatgccagt	cgggaaatcc	acagaagaag	tagcggtcac	ggcgcaagca	4200
atcactgggc	gcaaatcgtc	aacaacggac	aggagttgtc	acggctaacc	atggatgggt	4260
ggtcgatcca	aggtcatcaa	ttgggacgaa	ttcacccagc	accataccat	ggaaaagaag	4320
attaatgagc	ggagccctga	gggggcgaga	attggtagtg	gtggacgagg	gggaagcgag	4380
aagaagccct	gtaatggaaa	gcctgccttt	aaggttgttg	cagccgctgt	cttgtccccg	4440
ttgggcccac	ggctgcgagg	gcggggcggt	taaggaagaa	gcctgaggct	gtccttgcac	4500
tgtccggtaa	gtatcctttc	gtaatactct	ttttctttcc	ctctttcctc	tctcctttcc	4560
agcagaagat	gggcgtaagc	aacgctggga	tgtatgagca	tgccctctat	ttgtgtgtta	4620
tttgacagtt	caacgaacta	atcgccttaa	ataaatgtat	ccgacaacag	agtgtcgatt	4680
tagagaaggt	gttgtggttg	gtatcccatc	ggatttattg	gctggagagg	ttaaaactgc	4740
ccctccataa	accatggtat	atcccctaat	tagtacctgt	ccgtcatcaa	caaccctaca	4800
tggattatct	tttagtggtt	actagtctta	aaaagactct	ggtagcgccc	attcaaccta	4860
tccatggatc	ttgaacagcc	ggttgcacaa	cttggatata	atctaggtat	gttgtattca	4920
tgtacatata	tactgtattc	tatagtgtca	cctaaatagc	ttggcgtaat	catggtcata	4980
gctgtttcct	gtgtgaaatt	gttatccgct	cacaattcca	cacaacatac	gagccggaag	5040
cataaagtgt	aaagcctggg	gtgcctaatg	agtgagctaa	ctcacattaa	ttgcgttgcg	5100
ctcactgccc	gctttccagt	cgggaaacct	gtcgtgccag	ctgcattaat	gaatcggcca	5160
acgcgcgggg	agaggeggtt	tgcgtattgg	gegetettee	getteetege	tcactgactc	5220
gctgcgctcg	gtcgttcggc	tgcggcgagc	ggtatcagct	cactcaaagg	cggtaatacg	5280
gttatccaca	gaatcagggg	ataacgcagg	aaagaacatg	tgagcaaaag	gccagcaaaa	5340
ggccaggaac	cgtaaaaagg	ccgcgttgct	ggcgttttc	cataggetee	gcccccctga	5400
cgagcatcac	aaaaatcgac	gctcaagtca	gaggtggcga	aacccgacag	gactataaag	5460
ataccaggcg	tttccccctg	gaageteeet	cgtgcgctct	cctgttccga	ccctgccgct	5520
taccggatac	ctgtccgcct	ttetecette	gggaagcgtg	gegetttete	atagctcacg	5580
ctgtaggtat	ctcagttcgg	tgtaggtcgt	tcgctccaag	ctgggctgtg	tgcacgaacc	5640
ccccgttcag	cccgaccgct	gcgccttatc	cggtaactat	cgtcttgagt	ccaacccggt	5700
aagacacgac	ttatcgccac	tggcagcagc	cactggtaac	aggattagca	gagcgaggta	5760
tgtaggcggt	gctacagagt	tcttgaagtg	gtggcctaac	tacggctaca	ctagaagaac	5820
agtatttggt	atctgcgctc	tgctgaagcc	agttaccttc	ggaaaaagag	ttggtagctc	5880
ttgatccggc	aaacaaacca	ccgctggtag	cggtggtttt	tttgtttgca	agcagcagat	5940

tacgcgcaga aaaaaaggat ctcaagaaga tcctttgatc ttttctacgg ggtctgacgc	6000
tcagtggaac gaaaactcac gttaagggat tttggtcatg agattatcaa aaaggatctt	6060
cacctagatc cttttaaatt aaaaatgaag ttttaaatca atctaaagta tatatgagta	6120
aacttggtct gacagttacc aatgcttaat cagtgaggca cctatctcag cgatctgtct	6180
atttcgttca tccatagttg cctgactccc cgtcgtgtag ataactacga tacgggaggg	6240
cttaccatct ggccccagtg ctgcaatgat accgcgagac ccacgctcac cggctccaga	6300
tttatcagca ataaaccagc cagccggaag ggccgagcgc agaagtggtc ctgcaacttt	6360
atcogcotco atcoagtota ttaattgttg cogggaagot agagtaagta gttcgccagt	6420
taatagtttg cgcaacgttg ttgccattgc tacaggcatc gtggtgtcac gctcgtcgtt	6480
tggtatggct tcattcagct ccggttccca acgatcaagg cgagttacat gatcccccat	6540
gttgtgcaaa aaagcggtta geteettegg teeteegate gttgteagaa gtaagttgge	6600
cgcagtgtta tcactcatgg ttatggcagc actgcataat tctcttactg tcatgccatc	6660
cgtaagatgc ttttctgtga ctggtgagta ctcaaccaag tcattctgag aatagtgtat	6720
gcggcgaccg agttgctctt gcccggcgtc aatacgggat aataccgcgc cacatagcag	6780
aactttaaaa gtgctcatca ttggaaaacg ttcttcgggg cgaaaactct caaggatctt	6840
accgctgttg agatecagtt cgatgtaacc cactcgtgca cccaactgat cttcagcatc	6900
ttttactttc accagcgttt ctgggtgagc aaaaacagga aggcaaaaatg ccgcaaaaaa	6960
gggaataagg gcgacacgga aatgttgaat actcatactc ttcctttttc aatattattg	7020
aagcatttat cagggttatt gtctcatgag cggatacata tttgaatgta tttagaaaaa	7080
taaacaaata ggggttccgc gcacatttcc ccgaaaagtg ccacctgatg cggtgtgaaa	7140
taccgcacag atgcgtaagg agaaaatacc gcatcaggaa attgtaagcg ttaatatttt	7200
gttaaaattc gcgttaaatt tttgttaaat cagctcattt tttaaccaat aggccgaaat	7260
cggcaaaatc ccttataaat caaaagaata gaccgagata gggttgagtg ttgttccagt	7320
ttggaacaag agtccactat taaagaacgt ggactccaac gtcaaagggc gaaaaaccgt	7380
ctatcagggc gatggcccac tacgtgaacc atcaccctaa tcaagttttt tggggtcgag	7440
gtgccgtaaa gcactaaatc ggaaccctaa agggagcccc cgatttagag cttgacgggg	7500
aaagccggcg aacgtggcga gaaaggaagg gaagaaagcg aaaggagcgg gcgctagggc	7560
getggcaagt gtageggtea egetgegegt aaceaecaea eeegeegege ttaatgegee	7620
getacaggge gegtecatte gecatteagg etgegeaaet gttgggaagg gegateggtg	7680
cgggcctctt cgctattacg ccagctggcg aaagggggat gtgctgcaag gcgattaagt	7740
tgggtaacgc cagggttttc ccagtcacga cgttgtaaaa cgacggccag tgaattgtaa	7800
tacgactcac tata	7814
<210> SEQ ID NO 145 <211> LENGTH: 9453 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Partial sequence of plasmid containing Aspergillus parasiticusniaD and Aspergillus flavus laeA <400> SEQUENCE: 145	
atggatttcc tacgtcttca atacaaacca tcagacgtcc tggaagctat cattacgtcg	60
cgatcgctta acaagtatga tcgtcttttc aagcatctgc ttcgactcct tcgaatggtt	120
	120

tcggtcgtca	aaggtctcat	tcgtgattct	acggggagag	actccctatc	tgggcaccct	180
cgaaacgtgt	accagaaatt	tcgcattgac	tgccagcatt	tcgtgctctc	attgagtgat	240
tattgctttc	atgtcggcat	tggctcgact	tggcagcggt	tccaagatag	cctggctaag	300
attgaacgct	gcctcgaccg	cggtgatatt	gatggcacga	tagaagcagc	acattctgtc	360
cctagactta	gagattatca	tgaagatatt	ctcgatcaaa	tgctttttgc	gctctttctc	420
agcaaaagac	atgctgatgc	agcgaagctg	ctggaaagta	ttttcggtac	gattttgaca	480
tttgctccat	tgtcgaggat	ggatggaacg	ageggegtge	gccacgaaac	tgaggctatt	540
gcctatcagc	tctttgctac	attccggaaa	caaacatccc	tttttgtgaa	ttatctacgc	600
aacttagatg	gcgtgaacgc	atcttcaaag	tettteggea	ggtccggcac	gacttttgca	660
tccagagaag	cgcctacatg	tgtattcgac	cacctcctag	cgcgcttgga	tatgaggaaa	720
tattactgag	agtcgaaaac	aagctccacc	gcaccagete	ttcttggagt	tttatattaa	780
agaatattcc	cagetegttg	tattattctt	tttctaccgt	gctaatgtat	caaggacttt	840
ggtacctatt	aacgttatta	ttegtgtget	attcccaaac	ataaccctgt	atatgtttcg	900
aacgccgtta	tgacccatgt	cttacatact	cattaagtca	ttcccttgga	taatcccaat	960
ttagaagaag	tgaaggtctg	attettteea	teetteegee	aacagtatcc	tccgagccga	1020
ttcttccatg	gctggcggac	cacaaatcag	gaccatactc	tcatcttctg	gagccgcgta	1080
ctcctttagg	agctcttcgg	atatgcgtcc	tcggcggcca	gtccatgagt	ccggcgcttt	1140
ggatagggtg	tgtattatat	tacaccttct	gctgtcggtt	gccatgaagc	cgtcgagctc	1200
agcccggcaa	aggatatett	cctcctgtct	gtttccattg	aggactgtac	aagaggtggg	1260
atcttgccgg	tcctgaacca	cggcgcgcaa	gacctggaag	atcggtgtga	taccggttcc	1320
tccacaaatc	atcttaaacg	accgaacatg	gegtteette	ccacttatga	caactcgtcc	1380
atttccaagg	tattcgaatc	tgcctgtcgg	accettgeat	tccaccacgg	agcccaatgg	1440
cagcctatcc	agggccatcg	tcatcttgcc	gcctgccgag	gtggctgttg	caaagtatac	1500
tttaaccagc	aagtccacgg	tecetttetg	gctggtttca	gaaattgggg	tgtatgagcg	1560
gatgatggct	tegttgttgg	atgatgtgtc	gaggactttg	atcataagat	gctggccgac	1620
tggtaaaccc	aatgtttgat	cttcgtgttc	caatttgaaa	ctaaatattc	gtgtatccca	1680
ggatatgtct	tteetttett	tcaatgttgc	ctttgtccaa	gaccgtgatt	ggaggaacac	1740
tgggcgaatt	tcatcggtgg	aggatgatgc	atcatccttg	agtgctttta	aaccttccgg	1800
atccatcgtt	ccaatatggt	actcaggcat	catcgccttt	gccgtctcgc	tatctatgga	1860
taggtgtcaa	tagatggtac	aattgcagtg	tgatattttt	gggactcacg	aatagcaagg	1920
aattcctcag	agacatccag	accagcagag	gagataatac	tetgegetee	gccagggtgg	1980
ccttcaagaa	atgcttgacc	atcatacact	tctccattca	cgatgaacca	tggcttctca	2040
tcgcaggaat	tctccttgaa	ttcttcaaaa	ccaatcactc	ggcttagccc	gtctttcttc	2100
atattaatgt	cttgcacggg	ctccggctcc	gtcggctcct	ctccttcgtg	tctttctccc	2160
cagttaccat	tegteaggte	acccccagcc	tttttgacgc	gttccatcca	tcctgtaggc	2220
atactagggt	gggtagggtg	ctcgaatctc	aagttcccgt	tttccttcgt	aattgtaacc	2280
cggaaccacg	ggttgttcat	cattccgaga	acggaccagt	acatatcgcg	aggctgcacg	2340
cccaatgctt	cgtccatggc	tcttacaagg	atggcatcac	tgttctcaag	ctctgggatg	2400
gtgatgctta	gagaccaaaa	acaccagcag	aagcaagttt	cgcgccagta	catatctact	2460

ttgcctccaa	aaagctcgcc	ttcaaaatca	cgatacttgt	cttcggcata	ttcgatttcc	2520
gccaatctcc	aagctataag	tccgttagct	ttgataagca	ttctcacaca	tcgagcgagc	2580
gagggtgcgt	acatttgcct	ttgtctaggg	atatttctac	cctggtaacc	ctgcggcccc	2640
caccggcgta	tgcatatcct	ctgacagtat	atgacggccc	tgcgaccagg	agatttaaga	2700
cctcattgtt	ttggggatat	gcaacggcgg	agttggtgtt	taggtcataa	atcgcatacc	2760
gctcatcgtg	ccaccaattt	cggttatttg	atgccatctc	aggcgagacc	attgttctgg	2820
gttagggagt	tagacaaatg	atggaaatat	aaaataagtg	ccctttagac	atacggtaag	2880
acgcggttgt	cattgatatg	gtaccagttg	tegettggtg	catcggtcaa	gatcagcctc	2940
ttcagccact	taacacttcg	tcctcctatt	tgaccgggca	cgacggccct	cagcggacga	3000
ccatgatctg	ggcgaagaga	ctccccgttc	attttatgtg	caagcatgat	ccccctgttg	3060
gggtccaggg	cccagttcaa	tttaatagat	gtgccgtagt	gaccattggg	ctgcggcgaa	3120
cttagcaatt	atcatcataa	gatagaggta	cagcatacca	gcttatccgc	tccttccata	3180
cagacgtatt	tegetttaeg	caggggtttc	gcactgcgga	gaatateege	cagcaatggg	3240
ccagtgaaga	gggcagtcga	tagtcccgcc	gatececagg	aaaaaccttt	cgttttacgt	3300
acattgtttt	gctctttgcg	tcgattgcca	gcacatacga	gggtgatagg	cgctgttatt	3360
tggtcgtact	gctgcaacac	ttgtcggaag	tttagtacca	aaggcttctc	taccagtcta	3420
tactttggtt	aacggatgtt	tggcagagaa	cctagcacta	tactaaccct	tcgatgctaa	3480
tttcccagtg	agggatatct	tcatccttga	tatgagggac	tgggccatga	tttcgaacat	3540
agaagagctc	cggcgatgtt	aaaaaccctt	tcagagtgtg	agaatgtaac	ggctcaaggg	3600
gacaagcatg	acagccggtg	caagcaacct	gataaggata	ggagtggagc	agttataact	3660
cataccttct	ttatacagat	ctgtgagagg	tggctcaaca	ttaaacggat	gaacacccgt	3720
taatctgata	agccgagggt	cacgaggaac	atggctatct	ggagttcctt	tatctacgct	3780
cagcacttct	gtcggccgtt	ttgatggcgg	tggcagaggg	atatcagaca	ggtctctcgt	3840
cgagatctct	tegetttega	ttttgatctg	acctgtctta	agaacgaggt	cagttgggac	3900
gagcgcatcc	gtccgcacct	cggtgatggt	tgccatgtta	ccggcaggga	aggccaatga	3960
aagtaaaatt	acgagggagg	gagcatgaac	aaggatgctg	agtatgaata	agtcgaatgg	4020
tcagccagtg	cattaactcc	aaataaggag	gcaatccacc	acactaaaat	actcttgcct	4080
atcgtatgat	ggcacgcagt	acgtgttacc	catgcgcggg	cagtggacat	tctattaggt	4140
cacggcagta	actccttgtt	accatataac	gcctcggaga	aaggtcacaa	taagcaatgc	4200
tcctaggaac	ccaccagcga	tttccgcgga	gtcccaaaat	cageteatte	tgggaggtgg	4260
gacgctcgaa	attagggcaa	gccttcaggc	tggacggcgt	cccaccgctt	aaccaagcgt	4320
tgaggcaaat	aaatcgcgtt	gacccacaca	acactctcga	ggctccagcc	atttgtccgc	4380
tcaaccttgc	aggatttctt	tttcgtcata	ttaattggtt	ctttgaagaa	tgatggagac	4440
aatgccgtga	agccatgtgc	aacttccaat	tagaagtgtt	gttgcttatc	gtccgaatga	4500
gcctggttcg	cgtggagaat	gggccagatg	ggageteaeg	gctgttagag	cggagctact	4560
actctgtacg	tacccttcaa	aggaattctc	ggtaagtttg	ttagagggat	attgctcacg	4620
tttaattggc	actccaggat	cctttaaatc	caggcaaaaa	tcgctcgatc	tggcttttt	4680
tgccaatctt	ggaagtctac	cgtatacttg	tagttacacc	cttgaggatt	taccacatga	4740
		cgtgatttaa				4800
cggatatatc	ggcatctaac	ttaggatttg	tettaegtae	aatatttatq	acctgtggtg	4860
	-	3	-	9		

aaacctgagg	caacaagggg	gcgcgattta	ccagactggc	gttcacatac	caatacagtg	4920
cttaattgta	ggtctcatgg	gtggaatgag	atgaccttcc	ctttcatcta	ttcttaagag	4980
gaacagggat	ggtacccaca	ccataccccg	aagagetegt	gatgtaatag	accctttcgt	5040
agtatgcggg	tttttattga	gatgccgata	tgcaaacttg	tagtaagact	aataataaca	5100
ggtgcaatta	attgaatttg	gggcctgtta	gcttatcctc	cacaaagcct	ttcgtaaaat	5160
aactaaccaa	ctcaccacta	tcagcctaaa	tgagtcgatc	gaacctccta	ccagtaccaa	5220
tetgettegt	atataataga	aggttgtcgg	ctccttctga	caagacataa	tctaatctaa	5280
gatttaactg	tggcttttgt	gcttaatcga	tttatctttt	ttatctatct	acggagtagt	5340
tattcacgga	tttttagctg	ataatctagg	ctacattttc	agactgaatt	acgcacaatt	5400
ttggggaagt	gagaaaacag	acagaaagag	gagaaaaaaa	agaaaaagaa	aaaagcatgg	5460
gctaatgtgt	gcccactgcc	cagacatcta	taccttgtat	gatgtatgta	tgatgagcaa	5520
acataactgt	acatacatat	accatacata	ccctataata	cacccatacg	gggtatattt	5580
acgtaccttt	taggtgtacc	cgtatgtata	ttgtatgcat	gcatgtatat	atgtacatga	5640
atacaacata	cctagattat	atccaagttg	tgcaaccggc	tgttcaagat	ccatggatag	5700
gttgaatggg	cgctaccaga	gtctttttaa	gactagtaac	cactaaaaga	taatccatgt	5760
agggttgttg	atgacggaca	ggtactaatt	aggggatata	ccatggttta	tggaggggca	5820
gttttaacct	ctccagccaa	taaatccgat	gggataccaa	ccacaacacc	ttctctaaat	5880
cgacactctg	ttgtcggata	catttattta	aggcgattag	ttcgttgaac	tgtcaaataa	5940
cacacaaata	gagggcatgc	tcatacatcc	cagcgttgct	tacgcccatc	ttctgctgga	6000
aaggagagag	gaaagaggga	aagaaaaaga	gtattacgaa	aggatactta	ccggacagtg	6060
caaggacagc	ctcaggcttc	ttccttaacc	gccccgccct	cgcagccgtg	ggcccaacgg	6120
ggacaagaca	gcggctgcaa	caaccttaaa	ggcaggcttt	ccattacagg	gcttcttctc	6180
gcttccccct	cgtccaccac	taccaattct	cgccccctca	gggctccgct	cattaatctt	6240
cttttccatg	gtatggtgct	gggtgaattc	gtcccaattg	atgaccttgg	atcgaccacc	6300
catccatggt	tagccgtgac	aactcctgtc	cgttgttgac	gatttgcgcc	cagtgattgc	6360
ttgcgccgtg	accgctactt	cttctgtgga	tttcccgact	ggcatcgctt	catcgcgggg	6420
agacagcacg	gcgttctcta	ttattatctc	tttgcctgtc	ggtacttttg	gctggtcacc	6480
gtcttggttt	ttccgtttct	gcccttctgt	gtttaagtgg	acgtgacctt	tgattctctg	6540
cctggtccta	tctccaactt	ttcttatttt	tttagattct	gatttttgat	ttttgatttt	6600
tgattttaat	tttttttct	tttctttct	tatttttctc	tgtttacttt	tctgattctc	6660
cccctgaatc	ctcgtctctc	tttctaatac	tgcgaatgat	atcttgtgtt	tcgcttttca	6720
ctcccttaaa	tttatgaatg	aacttgccgt	ccattcacta	ggctcagccc	cattctactc	6780
tatgttttat	tcgtcactca	atatatac	catatacaca	acctacaagc	gtccacgcta	6840
ctgacatcag	cttgtcgcca	tcgacgtcgg	cattcttccc	ctcgtcacca	tcttaaagta	6900
ccaaacttct	taccgaccct	tcttcaccac	acggaatgtt	tggaaacggc	cagactggac	6960
agcgtcttcc	ggcgatggcc	tcaccaccac	acgactcgta	ctactcacag	tcattggcat	7020
ctagtcgatc	aaggaataac	tcggatgcta	tggatatcta	cgccatcaca	gacagagatc	7080
ctccggcacg	agaaccctct	ggttatagcc	agtggtaccg	taatggttct	ccaagtgtga	7140
attccattca	tagcaagtaa	ttetteeetq	tttccttaaa	tetgeegttq	aatttctctc	7200
	_ 3	5		, , ,		

-continued

tgggctgcgt	cagttgttca	ctctgccgaa	gctgttattg	ttggacgact	aatgttcttc	7260
tctgttgtta	ctgtctctct	ctaggagttc	tgaaaaacag	cctttctatg	aagaaaacgg	7320
acgaatgtat	catgcgtacc	gcaaaggggt	atatatgcta	ccatgcgatg	agcaggagca	7380
agatcgcctt	gatatettee	acaaattatt	cacggtggca	agggtgtcgg	atggcctaat	7440
gtacgccccg	catcccagga	acggccgatt	tctagacttg	ggctgcggga	ccgggatatg	7500
ggcaattgat	gttgccaaca	aatacccaga	cgctttcgtt	gttggggttg	accttgctcc	7560
catacagccc	tcaaaccacc	caaagaattg	cgaattttac	gccccattcg	acttcgagag	7620
teettgggee	atgggcgagg	attcttggga	cctgattcac	ctgcaaatgg	ggtgtggaag	7680
tgtgatggga	tggccgaacc	tgtaccggag	aatattcgct	caccttcgac	ccggggcttg	7740
gtttgaacag	gtggagatcg	actttgaacc	gcggtgcgat	gaccgtcctc	tcgagggact	7800
agctattcga	cagtggtatc	agtatctaaa	gcaagctaca	caagatgcca	tgcgacctat	7860
aaaccacaac	tctcgtgata	caatccgaga	tctgcaggag	gctggtttta	ccgatattga	7920
tcatcaaatg	gtggggttgc	cccttaaccc	atggcatcaa	gacgagcacg	aaagaaaggt	7980
tgctcgctgg	tacaatttgg	ctgtctcgga	gagtattgag	tegeteagta	tggcgccttt	8040
cagtcgtatt	tttaattggg	acttggacag	aatcaggcgc	atttcgtcag	aagtcaagtc	8100
ggaggcgttc	aacaaagaaa	tacacgccta	caatatcctt	catatatacc	aagcacggaa	8160
acctgcgaac	tgattcttct	accaacatgc	gcacgacgga	catccaaaca	tgcgccagca	8220
gcgtcactag	tgcccaagct	ccgagttatg	gggtgggcga	attaccatcc	aggcagtcac	8280
ctttatctgc	tcttttatga	gctctccaaa	gatgcagcga	gttgatatga	gcttgtgtgg	8340
tgccacttgt	tagcttgcac	acaacggtcc	cgagcagtca	ttgcttgcgt	actgcaaagc	8400
aaaatcgaac	tcatacgggt	ctgtgatctc	tatctatctg	gaaaatccag	tgtgtcggga	8460
gagctccatt	actgggtatt	cggtccgaga	cgtcctgttt	aggetgttgg	ttatggcagc	8520
cacgaacgat	gcactgagct	gcctgtatca	ctccagtagc	acttagaagg	ttgatggaaa	8580
tatacagagt	ttgctttgat	tatcataggt	ccgagcatga	ggcgggcgta	cgtcttgcga	8640
agggggttat	acaaattacc	caggccaaac	aggcaagtca	aagatcattt	cgtccaaaaa	8700
atacatcgcg	atgcagetet	tgcccctatg	gacgaaccaa	tggggctgca	gaatactggc	8760
aggtcattaa	taggacacgg	catggcatgc	cctctgccct	ctgccctctg	cctgctctta	8820
acctggctgt	ggctccttca	tcacacttca	ggaccacatc	attecetgge	attgcatgct	8880
ttcccgcttg	ccaaagacag	catatttatg	gcgttttggg	tgagtgtaat	gttgataact	8940
gtatacccgg	tttagttcat	tctatctgtg	acgacatcat	ccaatcatgg	ttattgctat	9000
attcgagatt	cataacatga	tatcacttaa	aaagaatata	aaaatttata	taatacaatg	9060
caaagaattt	tatccatgtg	atcgtttgta	aggtctcact	accagctaac	atgaaccgga	9120
aatgcaagtg	tgtgagatac	cagtctttgg	gtatttcgcc	aaaacgcctc	gatagccttc	9180
aagacacccg	cccacccaat	gacccaagac	agatgctcct	gtaaattatc	taatactata	9240
atgtgaactc	atacaatgat	atcactttaa	cgccggcggc	ccttttgtcc	ccgggaggta	9300
ccgtttctac	ctccagtact	ttctttctca	ccatctccgc	tggcccagtc	acggccagcg	9360
ccggggctac	gcctccgcat	tgttccggga	ccgacggcaa	tgatgtcctc	ttcactttcg	9420
ctgtcagaaa	tatctagaag	cacaggcatg	cgg			9453

<210> SEQ ID NO 146 <211> LENGTH: 1455

-continued

<212> TYPE: DNA <213> ORGANISM: Aspergillus flavus	
<400> SEQUENCE: 146	
atgtttggaa acggccagac tggacagcgt cttccggcga tggcctcacc accacacgac	60
tegtaetaet cacagteatt ggeatetagt egateaagga ataaetegga tgetatggat	120
atctacgcca tcacagacag agatcctccg gcacgagaac cctctggtta tagccagtgg	180
taccgtaatg gttctccaag tgtgaattcc attcatagca agaacggccg atttctagac	240
ttgggctgcg ggaccgggat atgggcaatt gatgttgcca acaaataccc agacgctttc	300
gttgttgggg ttgaccttgc tcccatacag ccctcaaacc acccaaagaa ttgcgaattt	360
tacgccccat tcgacttcga gagtccttgg gccatgggcg aggattcttg ggacctgatt	420
cacctgcaaa tggggtgtgg aagtgtgatg ggatggccga acctgtaccg gagaatattc	480
gctcaccttc gacccggggc ttggtttgaa caggtggaga tcgactttga accgcggtgc	540
gatgaccgtc ctctcgaggg actagctatt cgacagtggt atcagtatct aaagcaagct	600
acacaagatg ccatgcgacc tataaaccac aactctcgtg atacaatccg agatctgcag	660
gaggctggtt ttaccgatat tgatcatcaa atggtggggt tgccccttaa cccatggcat	720
caagacgagc acgaaagaaa ggttgctcgc tggtacaatt tggctgtctc ggagagtatt	780
gagtcgctca gtatggcgcc tttcagtcgt atttttaatt gggacttgga cagaatcagg	840
cgcatttcgt cagaagtcaa gtcggaggcg ttcaacaaag aaatacacgc ctacaatatc	900
cttcatatat accaagcacg gaaacctgcg aactgattct tctaccaaca tgcgcacgac	960
ggacatccaa acatgcgcca gcagcgtcac tagtgcccaa gctccgagtt atggggtggg	1020
cgaattacca tccaggcagt cacctttatc tgctctttta tgagctctcc aaagatgcag	1080
cgagttgata tgagcttgtg tggtgccact tgttagcttg cacacaacgg tcccgagcag	1140
tcatttgcgt actgcaaaag caaaatcgaa ctcatacggg tctgtgatct ctatctatct	1200
ggaaaatcca gtgtgtcggg agagctccat tactgggtat tcggtccgag acgtcctgtt	1260
taggetgttg gttatggeag ceaegaaega tgeaetgage tgeetgtate acteeagtag	1320
cacttagaag gttgatggaa atatacagag tttgctttga ttatcatagg tccgagcatg	1380
aggcgggcgt acgtcttgcg aagggggtta tacaaattac ccaggccaaa caggcaagtc	1440
aaagatcatt tcgtc	1455

What is claimed is:

- 1. A method of producing an isolated *Aspergillus* secondary metabolite comprising the steps of:
 - (a) transforming an *Aspergillus* cell or organism with a nucleic acid encoding a veA polypeptide comprising SEQ ID NO:116;
 - (b) culturing the transformed *Aspergillus* cell or organism 55 under conditions conducive to production of a second-
- ary metabolite that the Aspergillus cell or organism is capable of biosynthesizing; and
- (c) recovering the secondary metabolite from the cultured, transformed *Aspergillus* cell or organism in an isolated form
- **2**. The method of claim **1** wherein the *Aspergillus* cell or organism is *A. nidulans* or *A. flavus*.

* * * * *