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Dai et al.

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(54) **ENHANCED CITRIC ACID PRODUCTION IN ASPERGILLUS WITH INACTIVATED ASPARAGINE-LINKED GLYCOSYLATION PROTEIN 3 (ALG3), AND/OR INCREASED LAEA EXPRESSION**

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(73) Assignee: **Battelle Memorial Institute**, Richland, WA (US)

(*) Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35 U.S.C. 154(b) by 0 days.

This patent is subject to a terminal disclaimer.

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Related U.S. Application Data

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(60) Provisional application No. 61/565,018, filed on Nov. 30, 2011.

(51) **Int. Cl.**
C12P 7/48 (2006.01)
C07K 14/38 (2006.01)

(52) **U.S. Cl.**
CPC .. **C12P 7/48** (2013.01); **C07K 14/38** (2013.01)

(58) **Field of Classification Search**
CPC C12P 7/48; C07K 14/38
See application file for complete search history.

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(57) **ABSTRACT**

Provided herein are fungi, such as *Aspergillus niger*, having a dolichyl-P-Man:Man(5)GlcNAc(2)-PP-dolichyl mannosyltransferase (*Alg3*) gene genetic inactivation, increased expression of a loss of *afIR* expression A (*Lae*), or both. In some examples, such mutants have several phenotypes, including an increased production of citric acid relative to the parental strain. Methods of using the disclosed fungi to make citric acid are also provided, as are compositions and kits including the disclosed fungi.

12 Claims, 22 Drawing Sheets

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FIG. 1A

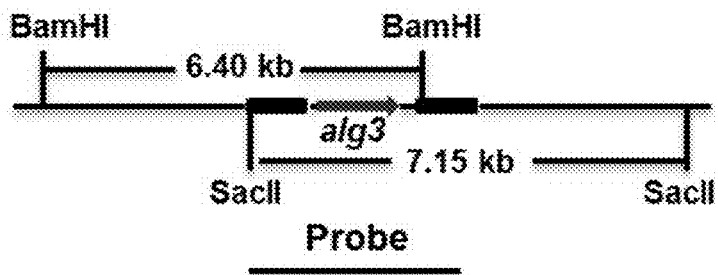


FIG. 1B

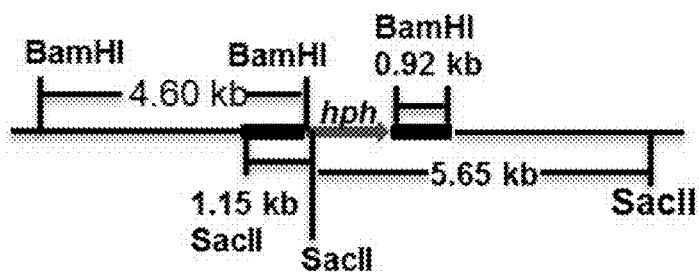
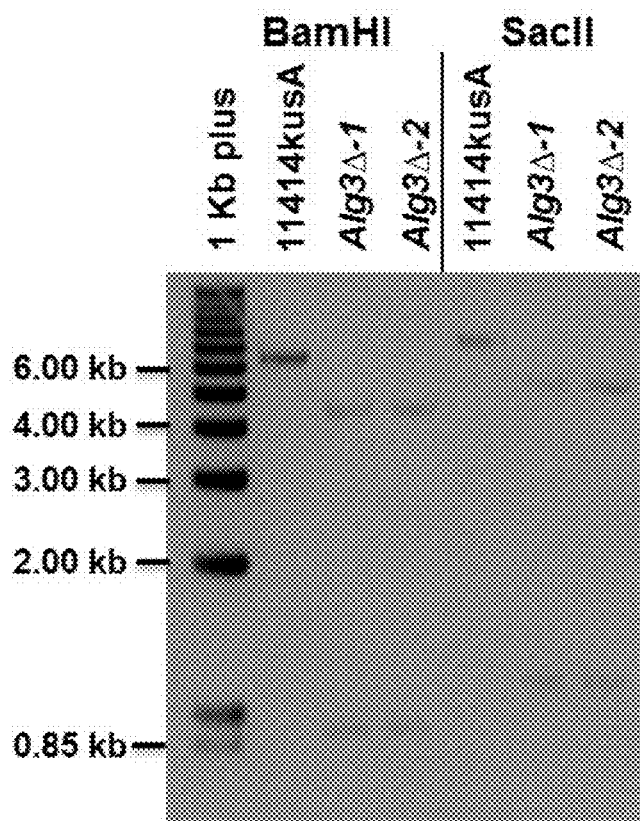


FIG. 1C



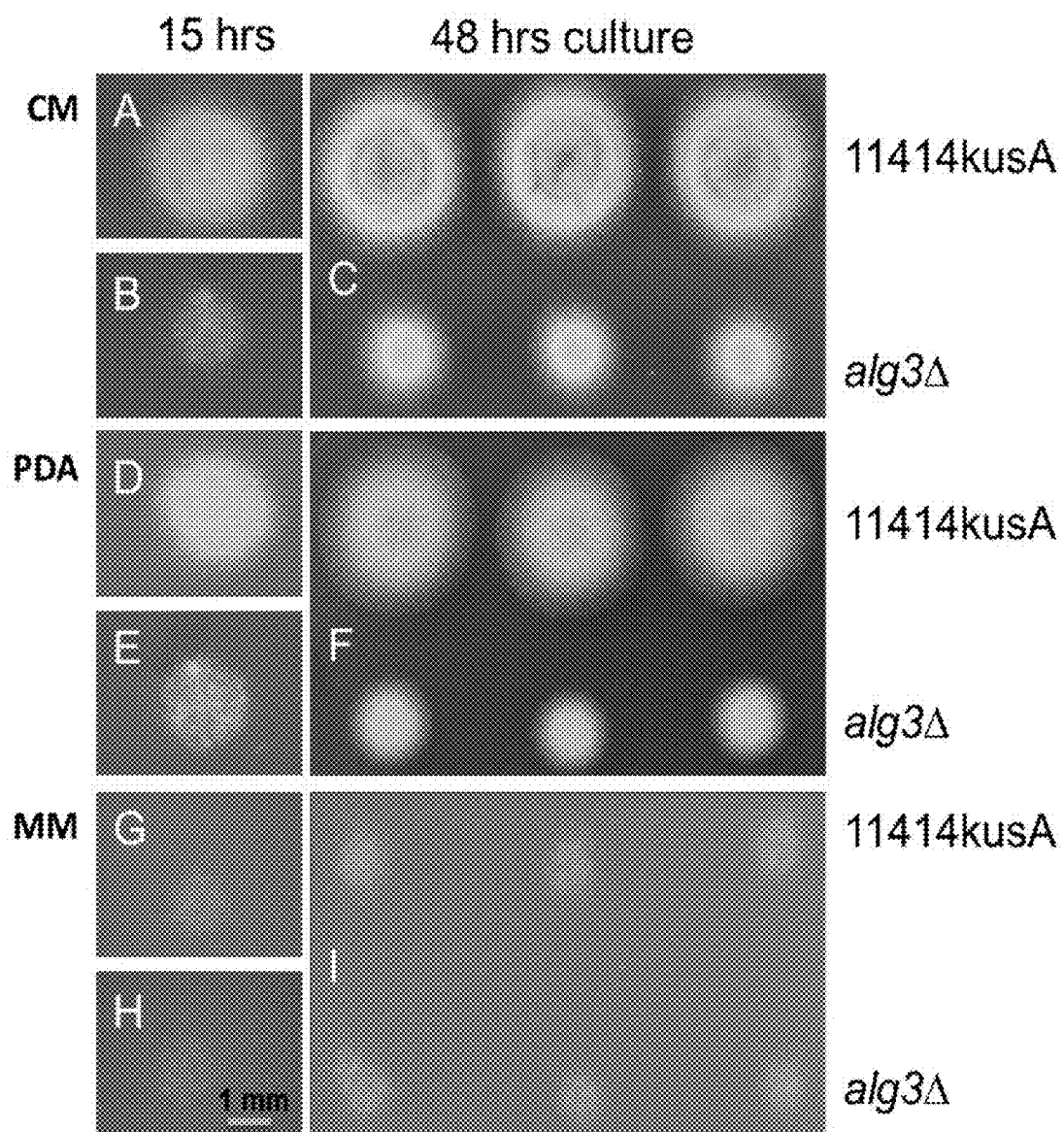


FIG. 2

FIG. 3A

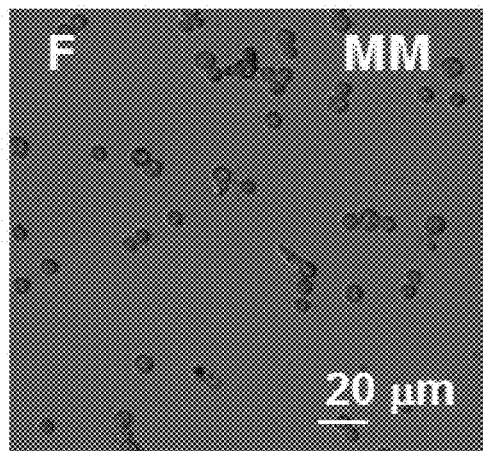
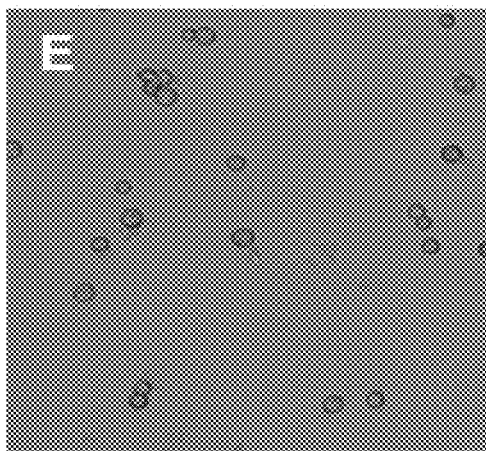
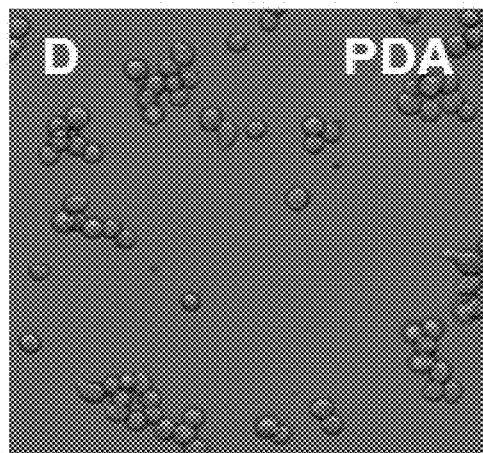
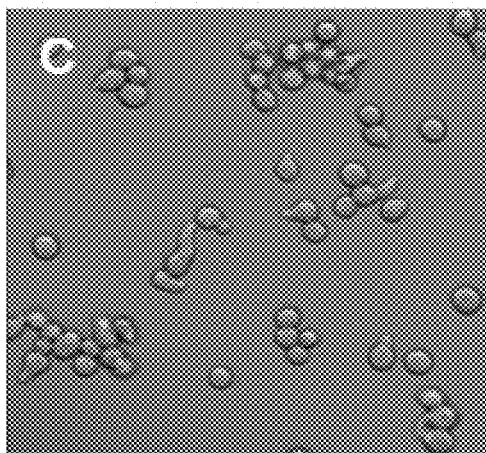
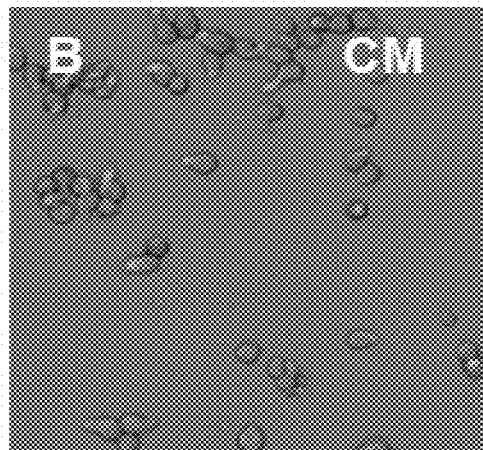
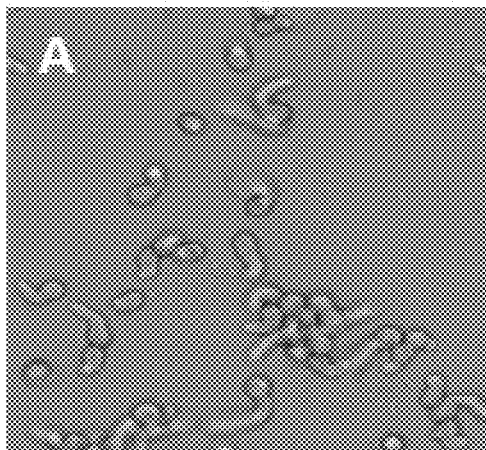
*11414kusA**alg3Δ*

FIG. 3B

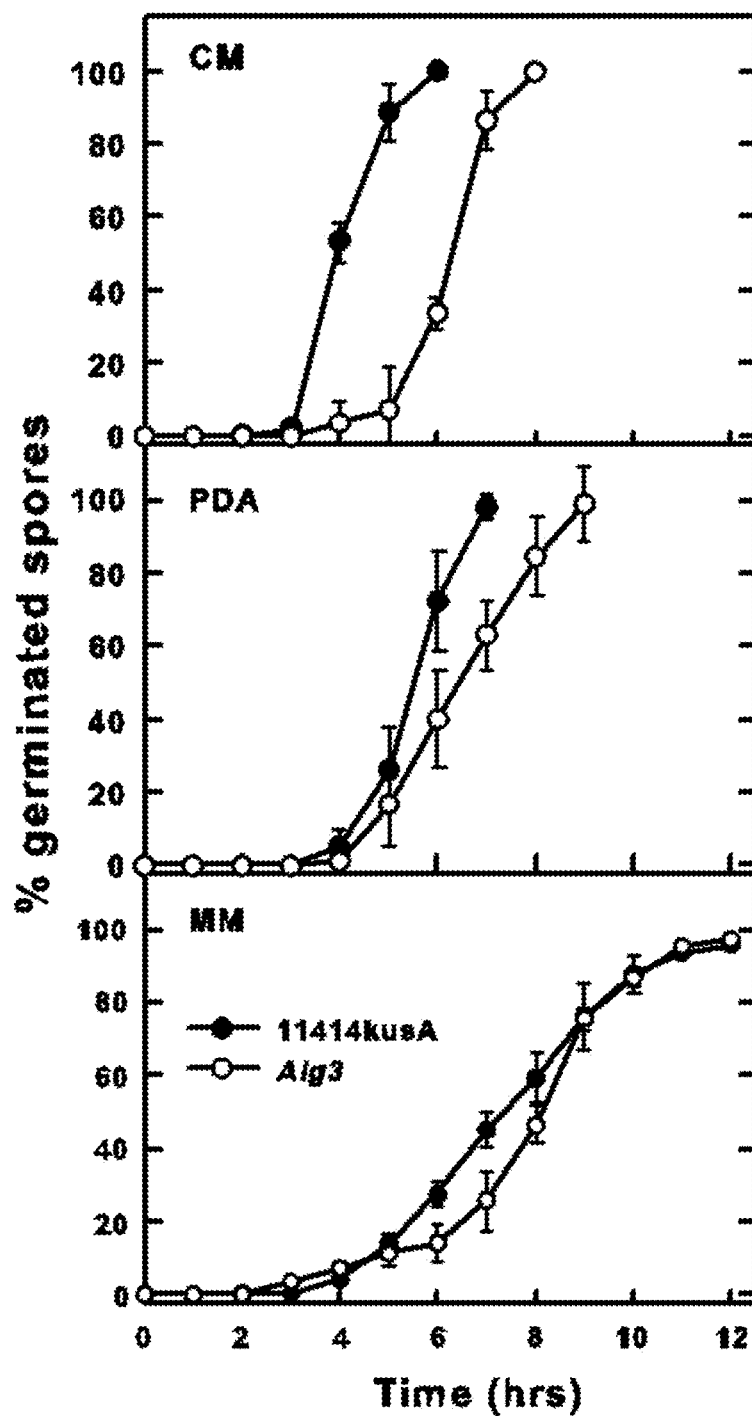
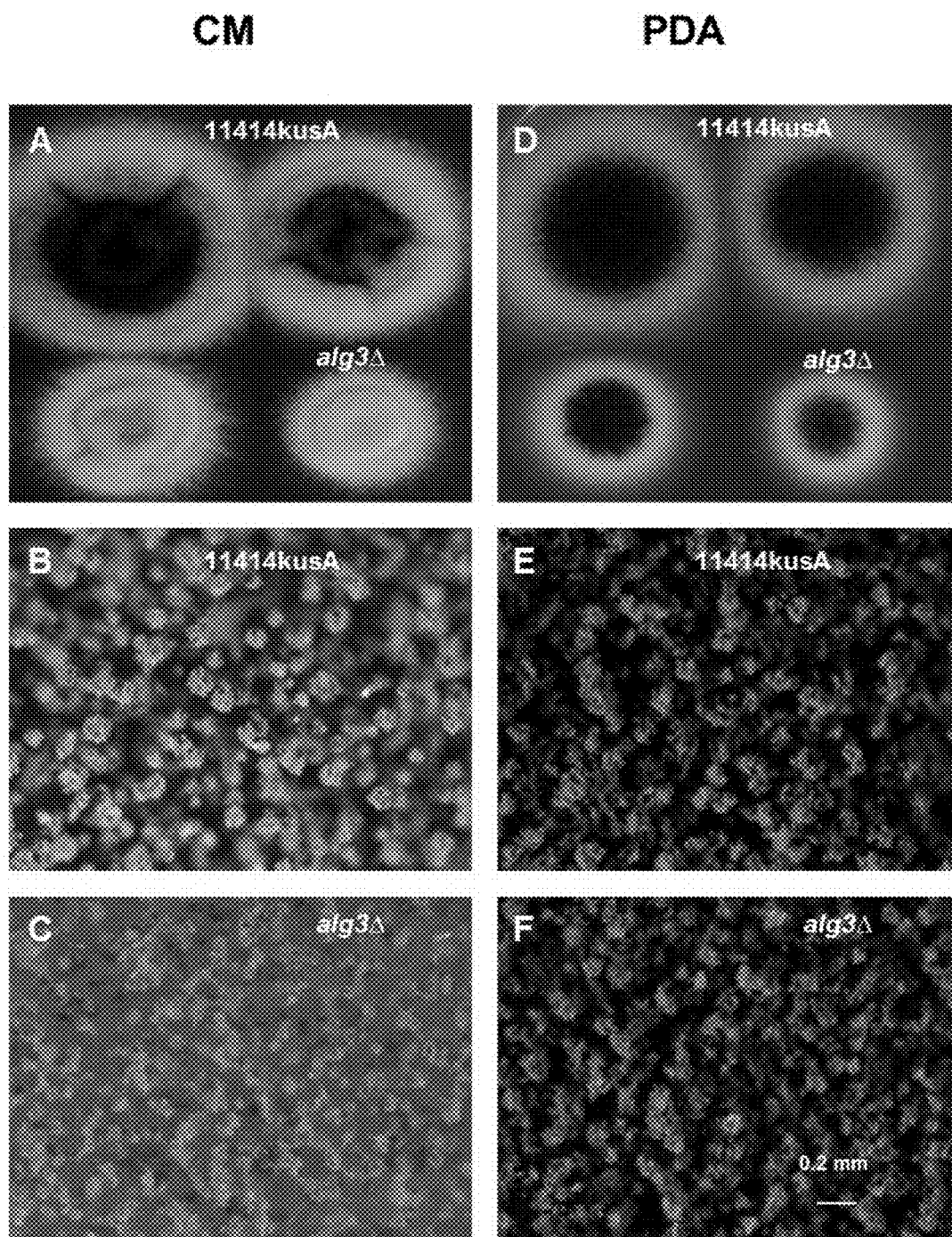


FIG. 4



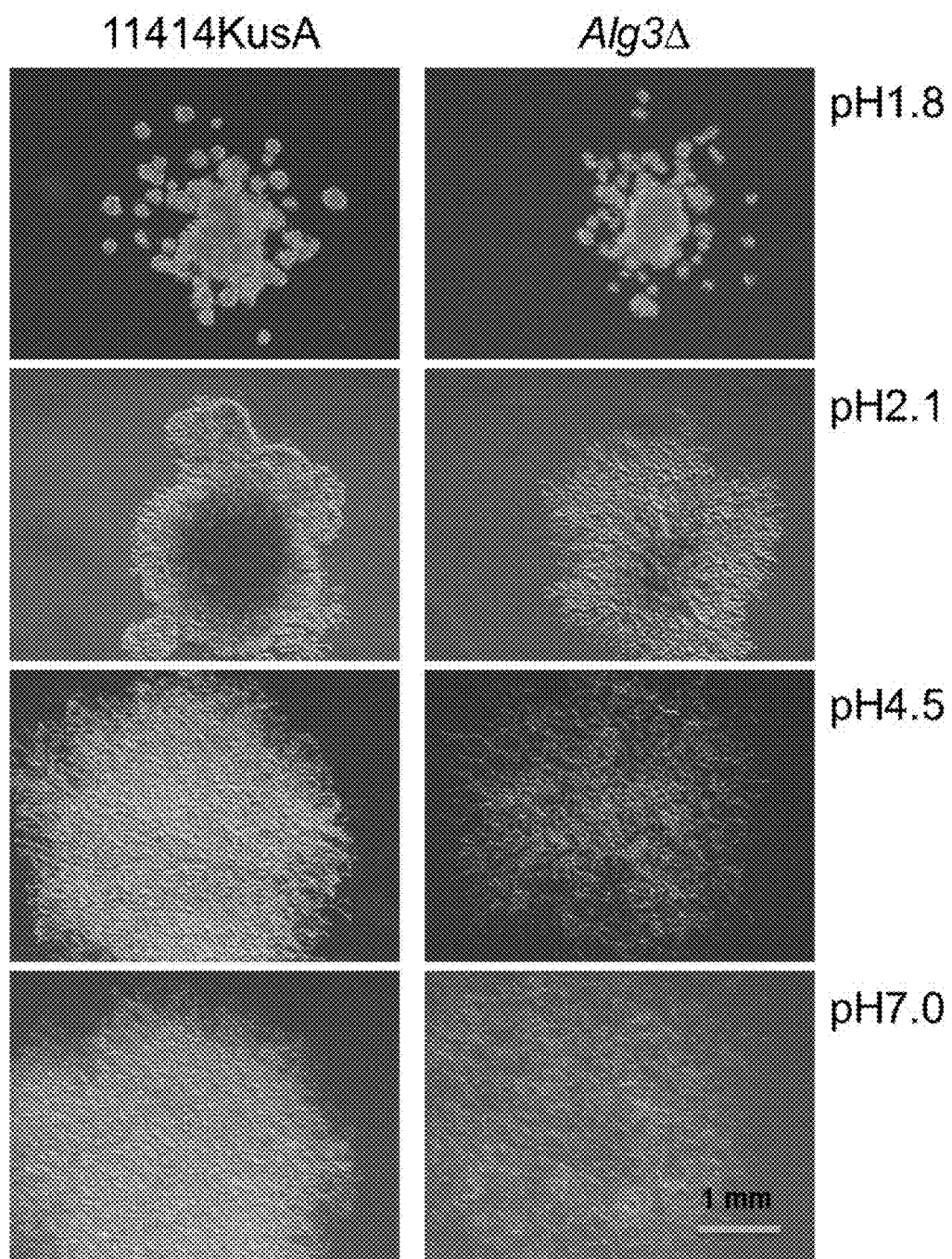
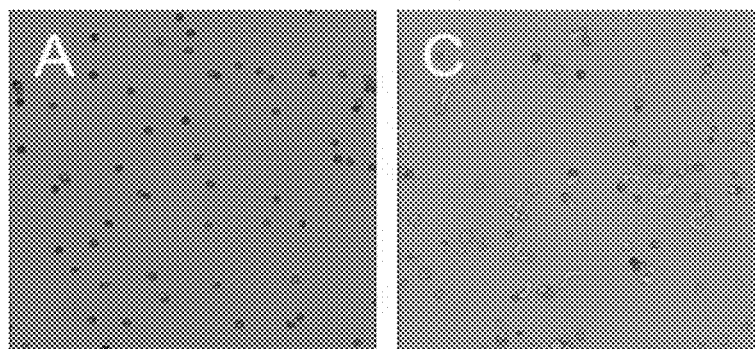


FIG. 5

FIG. 6A *11414kusaA* *Alg3Δ*
8 hrs after inoculation



15 hrs after inoculation

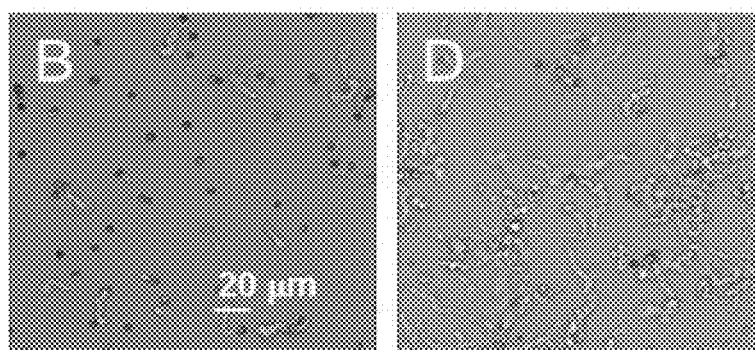
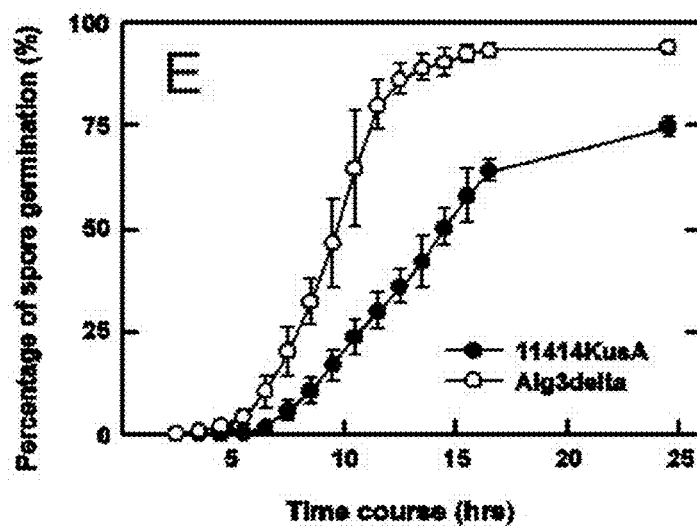


FIG. 6B



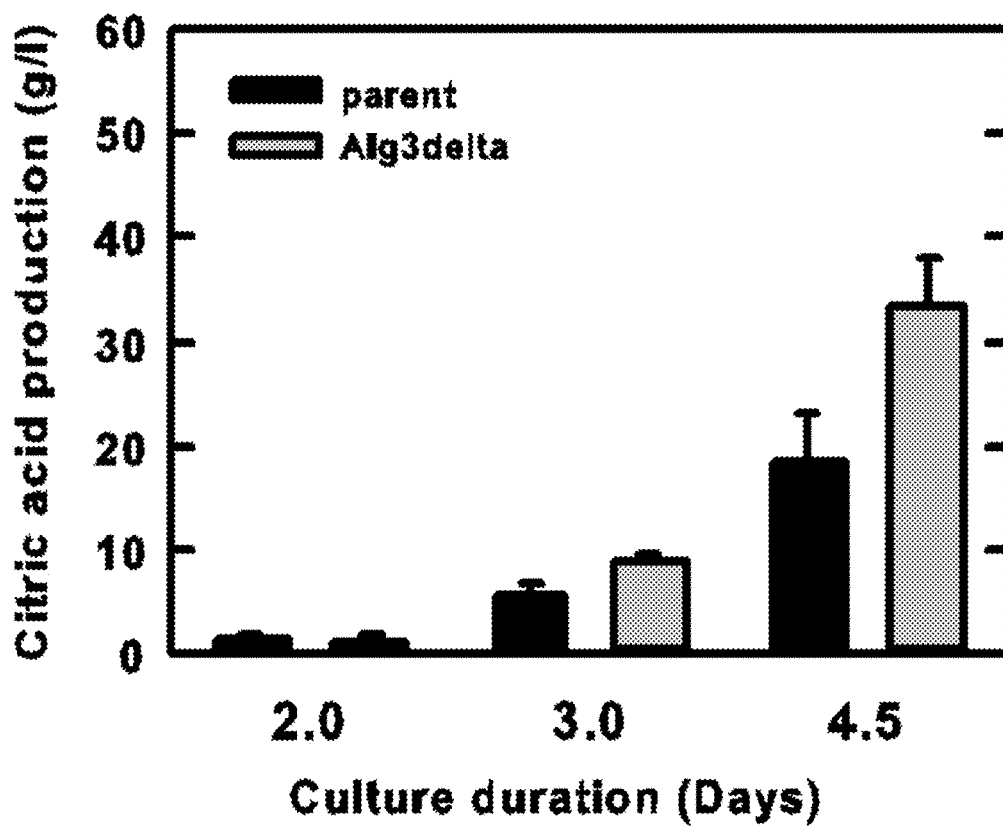


FIG. 7

Transgene fragment for complemented transformation

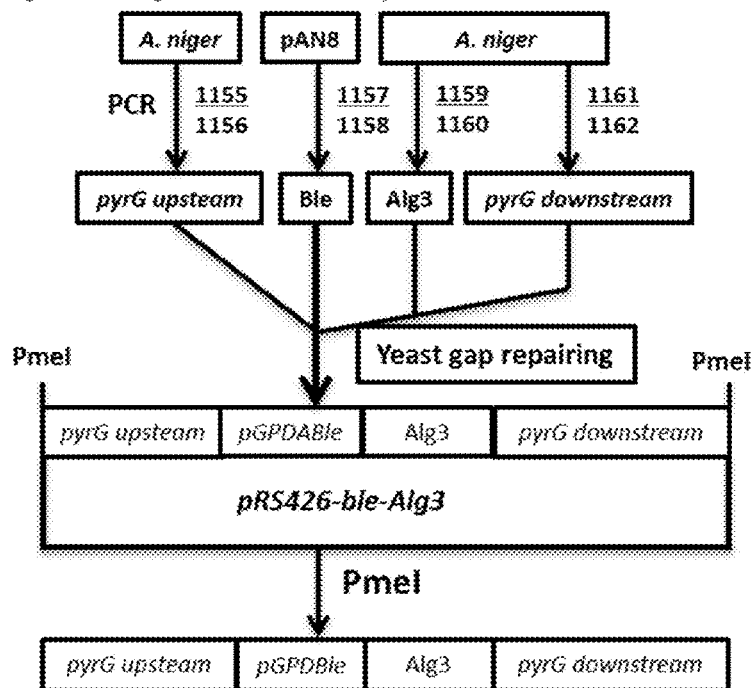


FIG. 8A

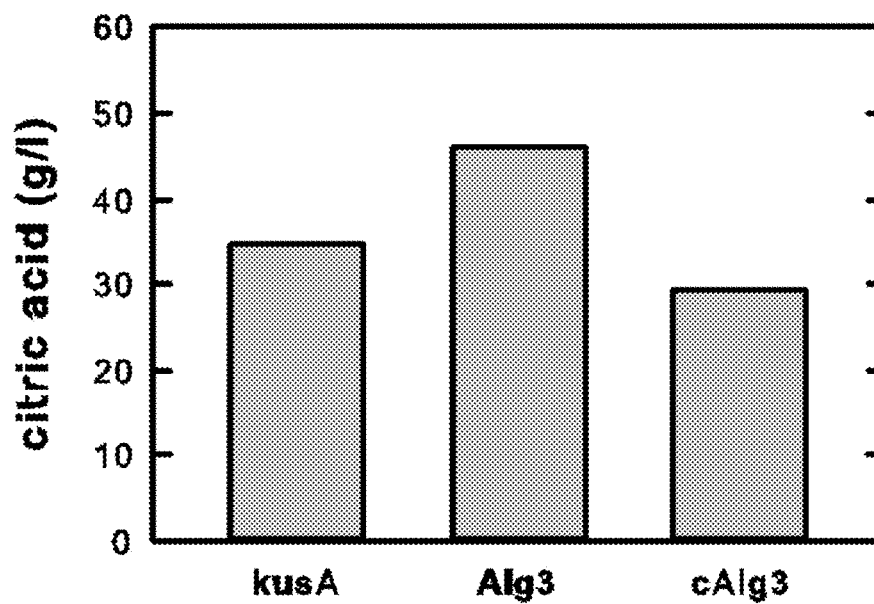


FIG. 8B

FIG. 9

GENE ID: 5996303 ADR 1 556094 | dolichyl-P-Man:Man(5)GlcNAc(2)-PP-dolichyl
mannosyltransferase [Aspergillus oryzae RIB40] (10 or fewer PubMed links)

Score = 137 bits (74), Expect = 2e-28
Identities = 400/553 (72%), Gaps = 40/553 (7%)
Strand=Plus/Plus

```
Query 801 CAGGTTTACTCGCGATACCGTTCCTACAAAACAACCCGGCGGGGTATC-TCTCGCGGGC 859
          ||||| ||| ||||| || ||||| ||||| ||||| ||||| |||||
Sbjct 643 CAGGTTCTACTAGCGATTCCCTTCCTACAGGSTAACCCCATAGGATA-CGTGCGCGGGC 701

Query 860 GTTCGAGCTAACCAGACAGTTCATGTTTAAATGGACAGTCAATGGAGATTGTGGCGA 919
          || ||| || ||||| ||||| ||||| ||||| ||||| ||||| |||||
Sbjct 702 CTTTGAGTTGACTAGACAGTTTATGTTCAAATGGACTGTCAATGGAGGTTTGTGGGTGA 761

Query 920 AGAAGTATTCTTAT-CTAAGAGCTTTTCCCTGGCATTGCT-GGCCGTCCACATTGTGCTG 977
          ||| | ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Sbjct 762 AGACTTGTTCCTATCCAAACAG-TTTCCTCTAGCCTTACTAGG-TTTCATATTTTCTG 819

Query 978 CTAGGCG-CT-TTTCGGTCACTGGTGGCTGA-GATAC-TCCAGG-TCTAGCTTSCCTG 1032
          ||| | ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Sbjct 820 CT-GG-GATTATTTGTTACCACAGGCTGG-TTACG-GCGT-CAGGATCTAACGTCCTG 874

Query 1033 -CGTTCATTGCGAATCTGCTAGC-GGGTCGACATCGCACAGT-GTCCCTCCCAACCCCT 1089
          ||| | ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Sbjct 875 AC-TTCCTCCGAGCCTACT-CCAAGGACGCCAACGCACCGTGGT-GCTTCTAAGTCT 931

Query 1090 ACATCATGAGCGTGATGCTCTCGTCTCTGACAG-TTGGCTTGTGTGCGCAAGGTCCCTT 1148
          ||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Sbjct 932 TCATAATGACCGTGATGTTGACATCGCTGGC-GATCGGGTGTGTGCGCAAGGTCCCTT 990

Query 1149 CATTACCAATTCTTCGCCCTACCTCTCCTGGGCGACACCCCT-CCTCCTCTGGCGCGCAGG 1207
          ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Sbjct 991 CATTACCAATTCTTTGCCCTATCTCTCCTGGGCTACGCC-TTGCCTTCTCTGGCGGGCTCG 1049

Query 1208 GTTTCATCCAATC-TTGCTGTAC-CTTATCTGGGCTA-TGCAAGAGTGGGCTTGAACA- 1263
          | | ||||| ||| || | || | ||||| ||||| ||||| ||||| |||||
Sbjct 1050 GCTCCATCCGATCCTTA-TATATGGG-ATCTGGGC-ACTACAGAGTGGGCTTGAATGT 1106

Query 1264 CATTCCCCAGCACCAAC-CTCAGTTCATCATT-GTTGTCTCTCCTGCTACCCAGAG 1321
          | | ||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Sbjct 1107 C-TACCCAAGCACCAATGC-CAGTTCT-TCGGTCGTTGTCTTCTCACTTGTGTTTCTAG-G 1162

Query 1322 TTT-CGGCGTCTCT 1333
          ||| ||| |||||
Sbjct 1163 TTTTCGGTGTCTCT 1175
```

[illegible]

FIG. 11

Score = 279 bits (713), Expect = 2e-87, Method: Compositional matrix adjust.
Identities = 159/405 (39%), Positives = 239/405 (59%), Gaps = 6/405 (1%)

```
Query 12  FNPRHTKWMAPLLVLGDAFLCALIIWKVPYTEIDWATYMQQISLYLSGERDYTLIRGSTG 71
          F P +T + +L + + +I KV YTEIDW YM ++ ++G DYT ++G TG
Sbjct 31  FKPEYTLTAVLWFLEIAINIWVIQKVSYTEIDWKAYMDEVEGVINGTYDYTQLKGD TG 90

Query 72  ELVYPAAHVYSYALYHMTDEGRDIFFGQILFAVLYLITLVVVLCCY-RQSGAPPYLLPL 130
          PLVYPA VY +T LY+LTD G +I GQ +FAV YLI L++V+ Y R EPPY+
Sbjct 91  ELVYPAGFVYIFTGLYYLTDHGHNIIRLGQYVFAVSYLINLLVMRIYHRTKKVPPYVFFF 150

Query 131 LVL-SKRLHSVYVLRFLFNDGLAALAMWVAILLFMNRKWTAAVAVWSTGVAIKMTLLLLAP 189
          + S R+HS+++LRLFND +A + + AI LF++ +WT A++S V++KM +LL AP
Sbjct 151 ICCASYRIHSIFILRLFNDPVAMMLCFGAINLFLDGRWTLGCALYSLAVSVKMNVLLFAP 210

Query 190 AIAVVTVLSLSLGPVSVGLGVLAVLVQVLLAIPFLQNNPAGYLSRAFELTRQFMFKWTVNW 249
          + + + L ++ L ++Q++L +PFL NP GY+SRAF+L RQF+FKWTVNW
Sbjct 211 GLLFLLLCFGLWKTFLERLALCAVIQLVLGLPFLLVNPVGYSRAFDLGRQFLFKWTVNW 270

Query 250 RFVGEEVFLSKSFSLALLAVHIVLLGAFVTCWLRYSRSSLPAFIRNLLAGRHRVTVSLPK 309
          RF+ E+VFL++ F LALL HI L FA+ W R S SS+ +++ + +
Sbjct 271 RFLPEDVFLNRYFHLALLAHITTTLLLFALKRWKR-SGSSIWTILKDPSEKETAHKVNA 329

Query 310 PYIMSVMLSSLTVGLLCARSLHYQFFAYLSWATPFLLWRAGFHP---ILLYLIWAMQEWA 366
          ++ ++ +S +G+ +RSLHYQF+ + P+LLW G +L LI + E +
Sbjct 330 DQMVLILFTSNFIGMCFSRSLHYQFYVWYFHTLPYLLWSGGVKKLARLLRVLIILGLIELS 389

Query 367 WNTFPSTNLSSTIIVVLSLATQSEFGLANSASAFYTMRSNPSGKEH 411
          WNT+PSTN SS+ + + + N A + RS K H
Sbjct 390 WNTYPSTNYSSLSLHVCHLIILLCLWLNPNPASPSHRSENKAKSH 434
```

FIG. 12

A. nidulans

Query 1 MTSPAHHNHSYHSPTSSDRGRSRQNSDAMDIQSITEREPATR-----YAVAGGPAPWN 53
M SP N+YSY S D GRSRQNSDAMDI IT +EP Y GGPA +

Sbjct 14 MASPNRNNYSYQGIESYDSGRSRQNSDAMDIHVITAQEPPREPPDNNDPYDGHGGPAGTS 73

A. niger

Query 54 RNGSPSPSPINSENRQFHEENGRTYHGFRRGMYFLPCDEQEQRDLDFHKLFTVARVSES 113
P R F+EENGRTYHG+RRG+Y LPCDEQEQRDLDFHKLFTVAR+SES

Sbjct 74 HYSKPP-----NRWLFYEENGRTYHGYRRGVYFLPCDEQEQRDLDFHKLFTVARMSSES 127

Query 114 LIYAPHPTNGRFLDLGCGTGIWAIEVANKYPDAFVAGVDLAPIQPPNHPKNCFYAPFDF 173
LIYAPHP NGRFLDLGCGTGIWAI+VA+KYP+AFVAGVDLAPIQPPNHP NCFYAPFDF

Sbjct 128 LIYAPHPNGRFLDLGCGTGIWAIDVAHKYPNAFVAGVDLAPIQPPNHPDNCFYAPFDF 187

Query 174 EAPWAMGEDSWDLIHLQMGCSSVMGWPNLYRRIEFAHLRPGAWFEQVEIDFEPRCDDRSLE 233
EAPW +GE+SWDLIHLQMGCSSV+GW NLY+RI HL+PGAWFEQVEIDFEPRCDDRSLE+

Sbjct 188 EAPWTLGENSWDLIHLQMGCSSVLGWQNLKYRILRHLQPGAWFEQVEIDFEPRCDDRSLE 247

Query 234 GTALRHWDCLKQATAETMRPIAHSSRDTIKDLQDAGFTEIDHQIVGLPLNPWHQDEHER 293
G ALR WY LKQAT +TMRPIAHSSRDTI+ L++AGFT+IDHQ+VGLPLNPWH+DEHE+

Sbjct 248 GLALREWYQYLKQATQDTMRPIAHSSRDTIRHLEEAGFTQIDHQMVGGLPLNPWHRDEHEQ 307

Query 294 KVARWYNLAVSESIENLSLAPFSRVYRWPLERIQQLAADVKSEAFNKEIHAYNILHIYQA 353
KVARWYNLA+SESIE LSLAPFSR++ W L+RI+Q+ A+VKS+AFNKEIHAYNILHIYQA

Sbjct 308 KVARWYNLAISESIETLSLAPFSRIHWDLDRIHQITAQVKSQAFNKEIHAYNILHIYQA 367

Query 354 RKP 356
RKP

Sbjct 368 RKP 370

FIG. 13

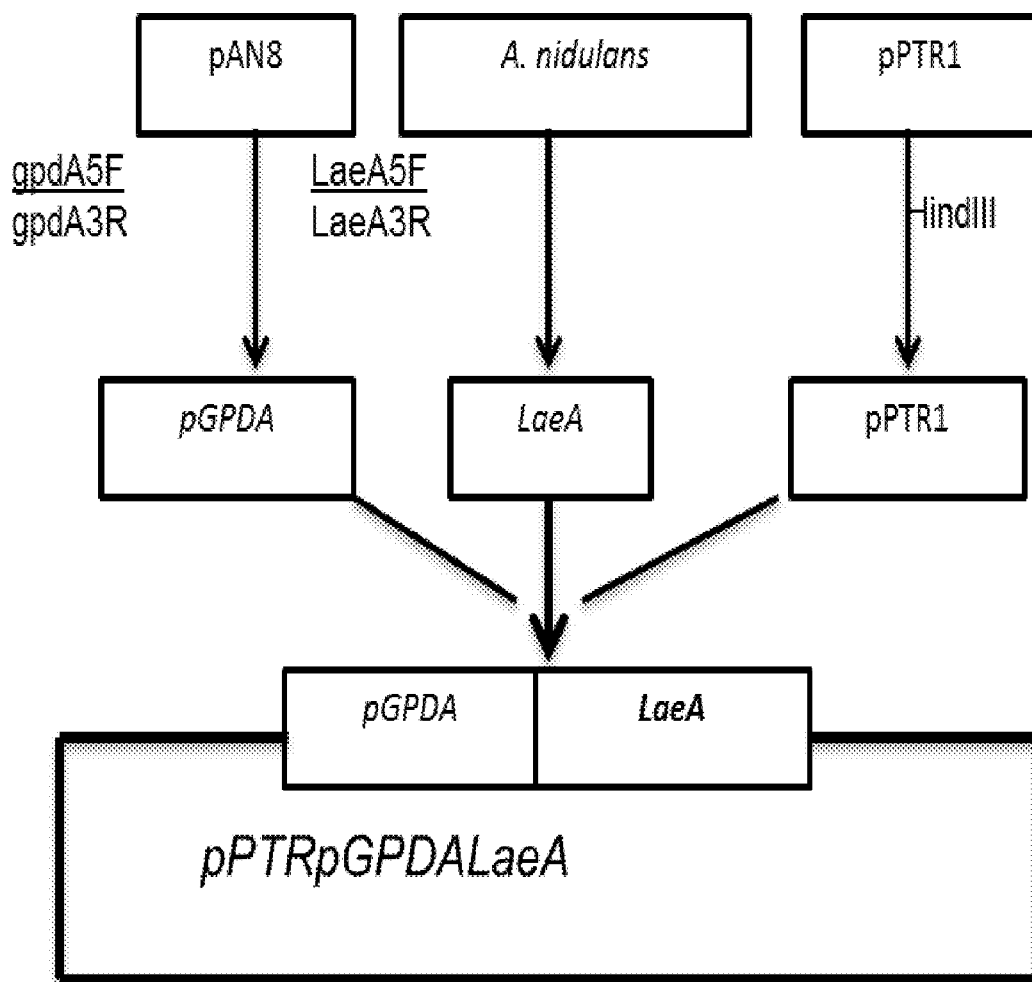


FIG. 14

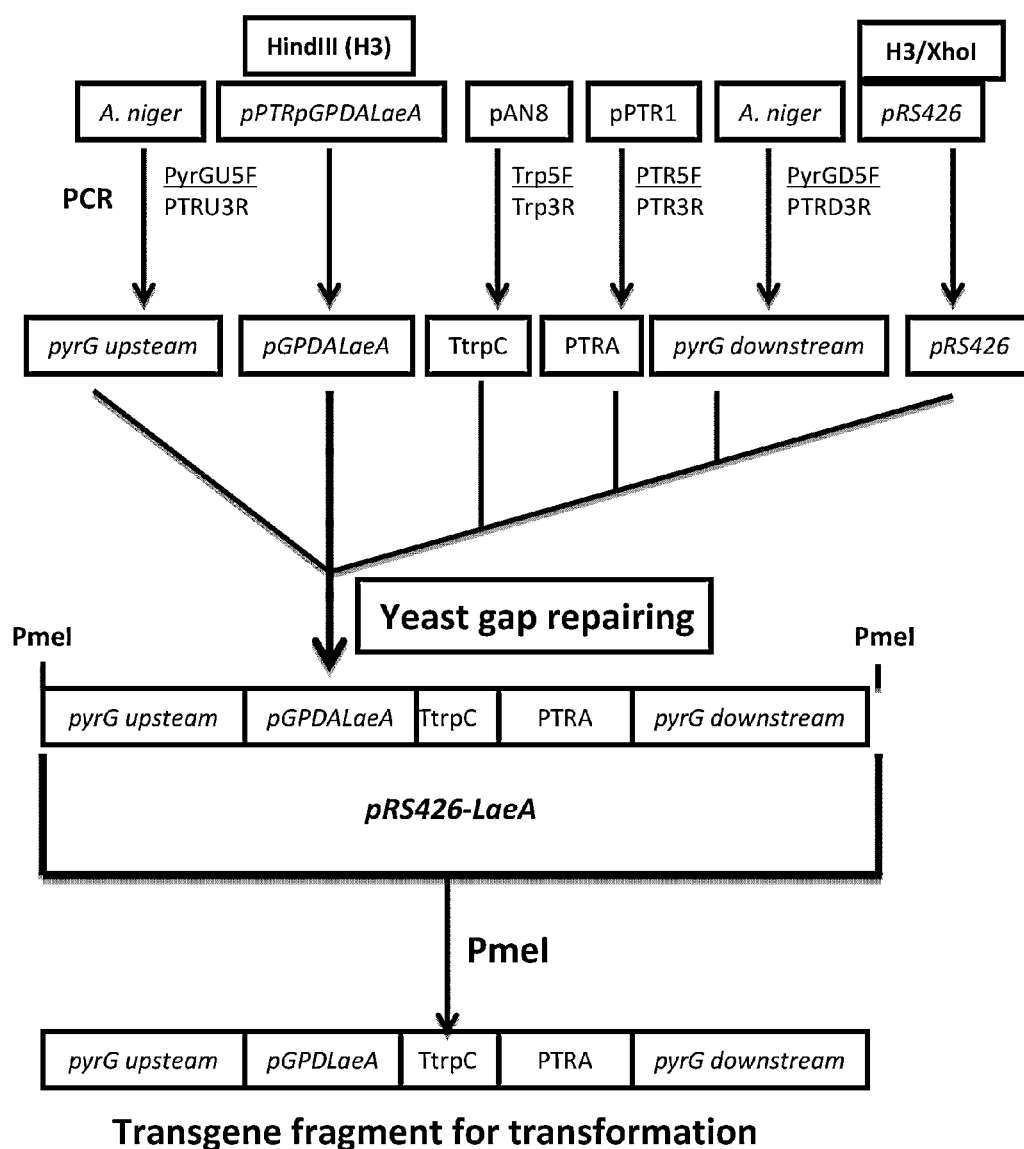


FIG. 15A. Oligo primers:PTR5F/PTR3R

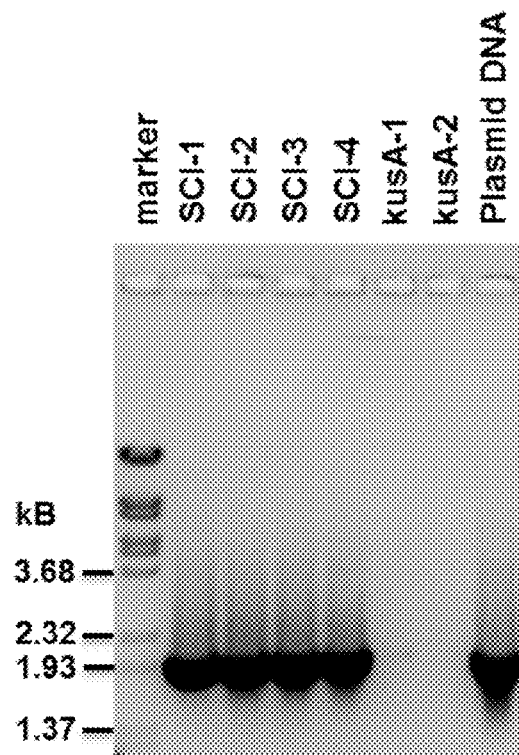
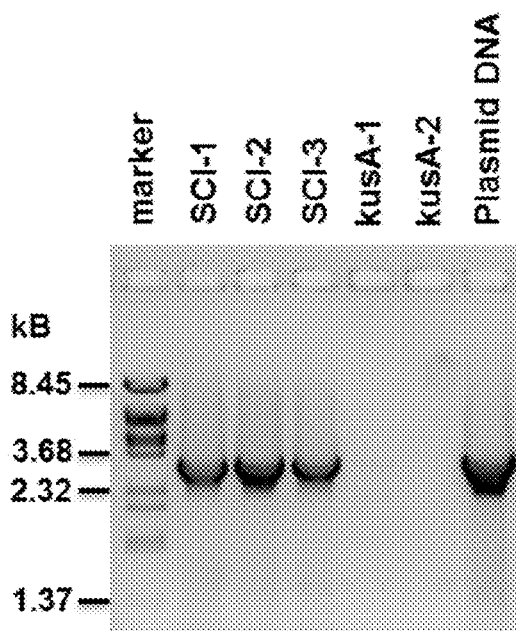


FIG. 15B. Oligo primers:LaeA5F/TRP3R



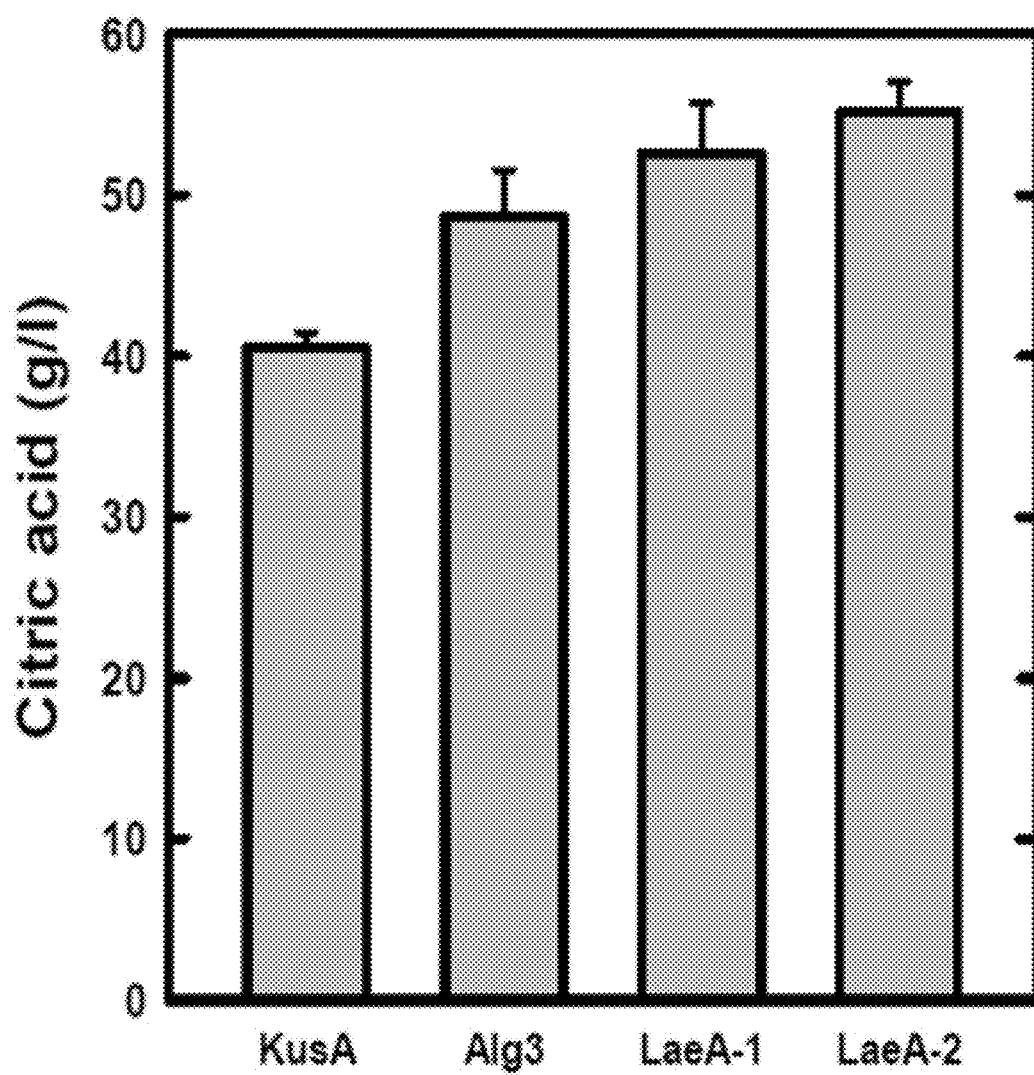


FIG. 16

FIG. 17A

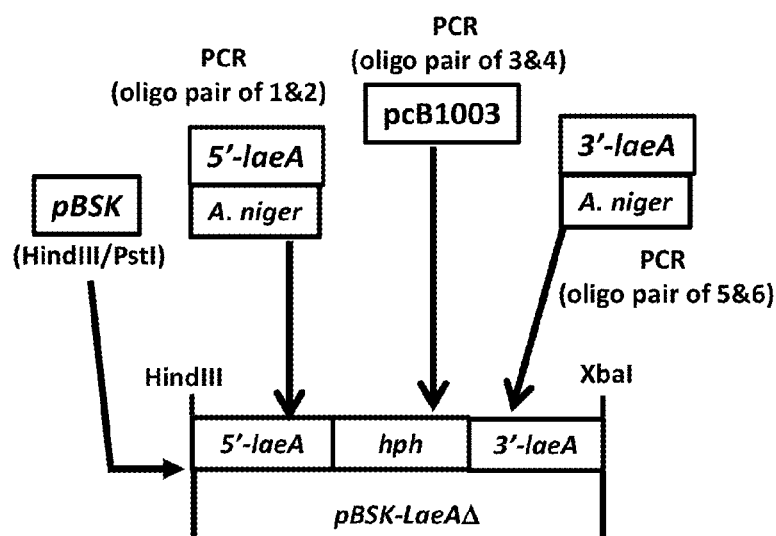
***laeA* deletion construct:**

FIG. 17B

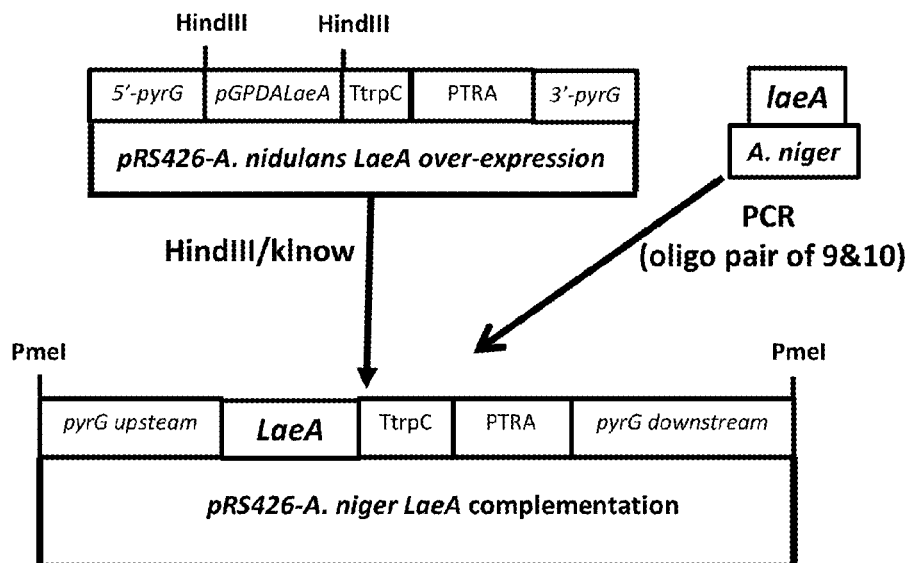
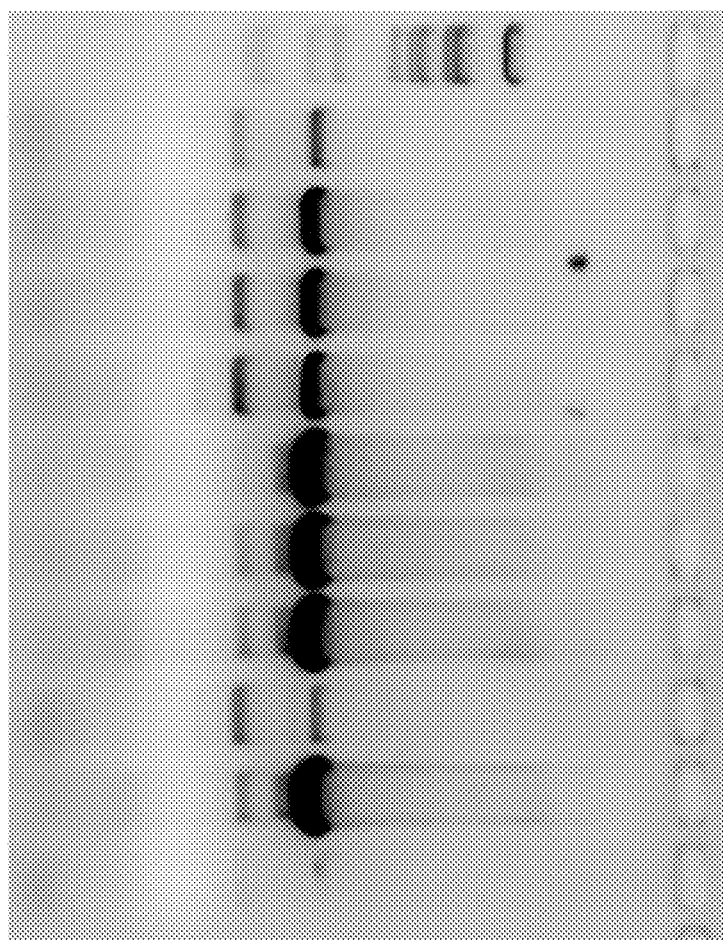
***laeA* complementation construct:**

FIG. 18

laeA gene deletion



Bλ

Clone-1

Clone-2

Clone-3

Clone-4

Clone-5

Clone-6

Clone-7

Clone-8

Clone-9

parent

FIG. 19

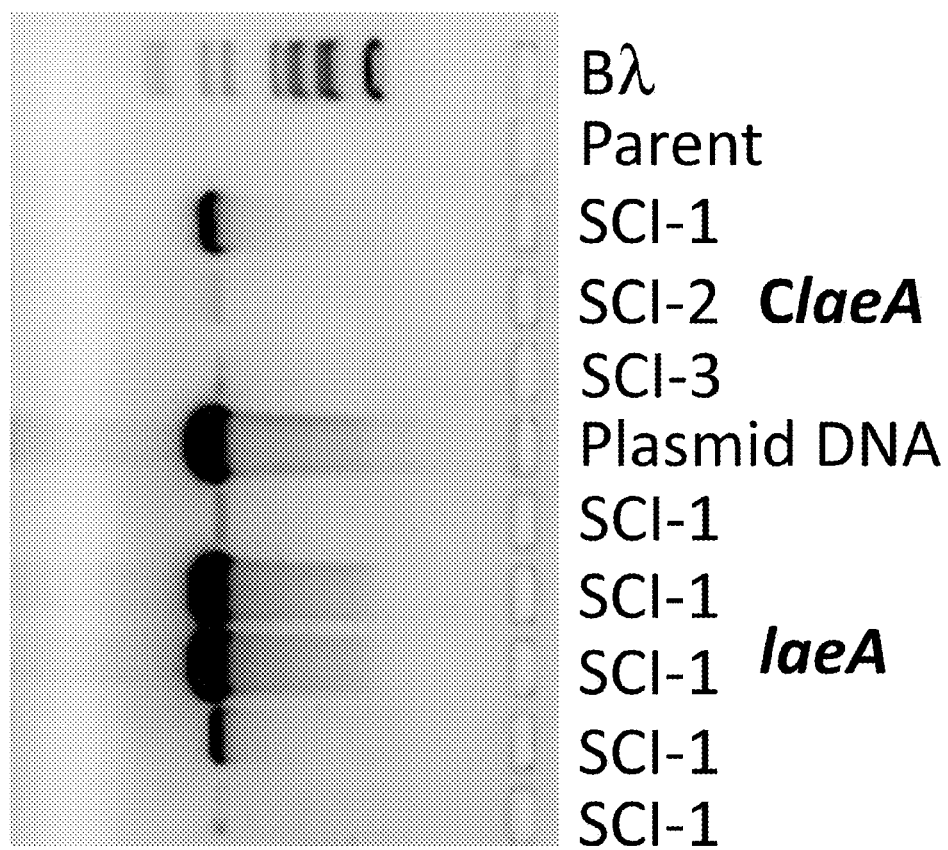
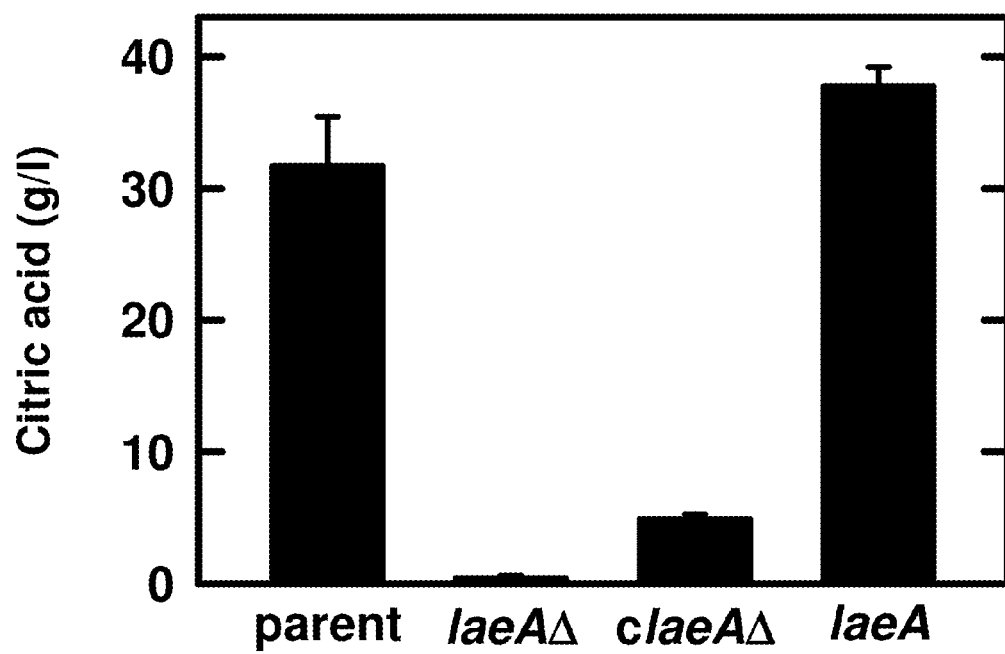


FIG. 20



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ENHANCED CITRIC ACID PRODUCTION IN ASPERGILLUS WITH INACTIVATED ASPARAGINE-LINKED GLYCOSYLATION PROTEIN 3 (ALG3), AND/OR INCREASED LAEA EXPRESSION

CROSS-REFERENCE TO RELATED APPLICATIONS

This is a continuation-in-part of U.S. application Ser. No. 13/691,396, filed Nov. 30, 2012, now U.S. Pat. No. 9,023,637, which claims the benefit of U.S. Provisional Application No. 61/565,018, filed Nov. 30, 2011. The above-referenced applications are herein incorporated by reference in their entirety.

ACKNOWLEDGMENT OF GOVERNMENT SUPPORT

This invention was made with government support under contract number DE-AC05-76RLO1830 awarded by the U.S. Department of Energy. The government has certain rights in the invention.

FIELD

This application provides recombinant *Aspergillus* fungi that are genetically inactivated for the dolichyl-P-Man:Man (5)GlcNAc(2)-PP-dolichyl mannosyltransferase (Alg3) gene, are genetically enhanced to increase the expression levels of the loss of aflR expression A (LaeA) gene, or both, which results in substantial improvement of citric acid production. Methods of using these fungi to produce citric acid are also provided.

BACKGROUND

Filamentous fungi, such as *Aspergillus niger*, are well known for their industrial applications in protein and chemical productions. They are used to produce a wide variety of products ranging from human therapeutics, glycosyl hydrolases to specialty chemicals (Punt et al., *Trends Biotechnol* 20(5):200-206, 2002; Schuster et al., *Appl Microbiol Biotechnol* 59(4-5):426-435, 2002; Gerngross, *Nat Biotechnol* 22(11):1409-1414, 2004; Nevalainen et al., *Trends Biotechnol* 23(9):468-474, 2005; Sauer et al., *Trends Biotechnol* 26(2):100-8, 2008; Magnuson and Lasure (2004). "Organic acid production by filamentous fungi." *Advances in fungal biotechnology for industry, agriculture, and medicine*, pages 307-340). Some of industrial *A. niger* strains are capable of growing on solutions of glucose or sucrose in excess of 20% (w/v) and converting approximately 90% of the supplied carbohydrate to citric acid. These remarkable properties are the reason that *A. niger* has been used to produce citric acid for more 80 years and is currently the primary source of commercial citric acid production (Magnuson and Lasure (2004). "Organic acid production by filamentous fungi." *Advances in fungal biotechnology for industry, agriculture, and medicine*, pages 307-340).

The maximum product output in fermentation processes is the result of optimal metabolic pathways and cellular formation, which are influenced by endogenous and exogenous factors. Cellular metabolisms are tightly controlled and highly interconnected, and are regulated spatially and temporally at different levels, such as transcription, post-transcription, translation, and post-translation. Therefore, different approaches have been explored to understand the regulatory

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mechanisms of metabolic processes and cellular formation for maximizing the product output in filamentous fungi. For example, comparative genomics was used to examine citric-acid-producing versus enzyme-producing *A. niger* strains (Andersen et al., *Genome Res.* 21(6): 885-97, 2011), proteomics was used to examine filamentous fungi related to enzymes or organic acid production (de Oliveira and de Graaff, *Appl. Microbiol. Biotechnol.* 89(2): 225-37, 2011), or combination of both genomics and proteomics were used to examine enzyme production (Jacobs et al., *Fungal Genetics and Biology* 46(1, Supplement):S141-S152, 2009). Although these studies examined the potential involvement of selected genes and proteins in optimizing production of organic acids or proteins in filamentous fungi, methods for altering the complex post-translation modifications (such as N-glycosylation of cellular proteins) for signal transduction, cellular formation and metabolism at different growth and development stages, which may affect product output, have not been examined.

Protein glycosylation is a ubiquitous and structurally diverse form of post translation modification, which occurs at all domains of life. More than two-thirds of eukaryotic proteins are predicted to be glycosylated (Apweiler et al., *Biochim Biophys Acta* 1473(1):4-8, 1999). N- and O-linked protein glycosylation are common types of protein glycosylation, occurring mainly on the asparagine (N) and serine/threonine (S/T) residues, respectively. N-linked glycosylation has been implicated in many biochemical and cellular processes, including protein secretion, stability and translocation, maintenance of cell structure, receptor-ligand interactions and cell signaling, cell-cell recognition, pathogen infection, and host defense in various organisms (Haltiwanger and Lowe, *Ann. Rev. Biochem.* 73(1):491-537, 2004; Dellaporta et al., *Plant Mol. Biol. Reporter* 1(4):19-21, 1983; Nam et al., *Biotech. Bioengineer.* 100(6): 1178-1192, 2008; Trombetta and Parodi, *Ann. Rev. Cell Dev. Biol.* 19(1):649-676, 2003; Tsang et al., *Fungal Genetics and Biology* 46(1): S153-S160, 2009; Pang et al., *Science*, 333(6050):1761-4, 2011).

N-glycosylation is highly complex and has been extensively studied in mammalian systems (Yan and Lennarz, *J. Biol. Chem.* 280(5):3121, 2005; Silberstein and Gilmore, *FASEB J.* 10(8): 849, 1996; Kornfeld and Kornfeld, *Annu. Rev. Biochem.* 54:631-664, 2005; Kim et al., *PLoS ONE* 4(10): e7317, 2009, 2009) and yeast (Kukuruzinska et al., *Annu. Rev. Biochem.* 56(1):915-944, 1987). The protein N-glycosylation pathways in filamentous fungi have also been identified (Deshpande et al., *Glycobiology* 18(8):626-637, 2008; Geysens et al., *Fungal Genetics and Biology* 46(1, Supplement): S121-S140, 2009) on the basis of the known genomic sequences. Several genes involved in N-glycosylation have been studied in filamentous fungi (Kotz et al., *PLoS ONE* 5(12):e15729, 2010; Kainz et al., *Appl Environ Microbiol* 74(4):1076-86, 2008; Maras et al., *J. Biotechnol.* 77(2-3):255-63, 2000; Maddi and Free, *Eukaryot Cell* 9(11):1766-75, 2010; Bowman et al., *Eukaryotic Cell* 5(3):587-600, 2006). In these studies, the effects of gene deletion on N-linked glycan patterns formation, the cell wall formation, overall protein secretion and/or the phenotypic changes were demonstrated.

Alg3 is localized in the ER and catalyzes the initial transfer of a mannose residue from dolichol pyrophosphate-mannose to lipid-linked Man5GlcNAc2-PP-Dol on the ER luminal side. It is involved in the early N-glycan synthesis in eukaryotes for the assembly of a Glc3Man9GlcNAc2 core oligosaccharide that is linked to the lipid carrier dolichol pyrophosphate. The Alg3 gene and its functions have been identified

and studied in *S. cerevisiae*, *P. pastoris*, *T. brucei*, *A. thaliana*, and human (Aebi et al., *Glycobiol.* 6(4):439-444, 1996; Korner et al., *EMBO J.* 18(23): 6816-6822, 1999; Davidson et al., *Glycobiology* 14(5):399-407, 2004; Manthri et al., *Glycobiol.* 18(5):367-83, 2008; Kajiura et al., *Glycobiol.* 20(6):736-51, 2010). In these studies, the Alg3 mutants exhibited a unique structural profile in the glycoproteins, such as Man3GlcNAc2, Man4GlcNAc2, Man5GlcNAc2, GlcMan5GlcNAc2, and Glc3Man5GlcNAc2, which affected the overall N-glycosylation by incomplete utilization of N-linked glycosites in glycoproteins. No obvious growth phenotype was observed in those Alg3Δ mutants of *S. cerevisiae*, *P. pastoris*, *T. brucei*, and plant except that the Alg3 defect in human caused severe diseases such as profound psychomotor delay, optic atrophy, acquired microcephaly, iris ophthalmos and hypsarrhythmia (Stibler et al., *Neuropediatrics* 26(5): 235-7, 1995; Sun et al., *J. Clin. Endocrinol. Metab.* 90(7):4371-5, 2005; Schollen et al., *Eur. J. Med. Genet.* 48(2):153-158, 2005; Kranz et al., *Am. J. Med. Genet.* 143A(13):1414-20, 2007; Denecke et al., *Pediatr. Res.* 58(2): 248-53, 2005).

LaeA, a global regulator gene for the secondary metabolism, was first identified in *A. nidulans* through complementing the aflR deficient mutants (Bok and Keller, *Eukaryot Cell* 3:527-535, 2004). Deletion of LaeA gene inhibits the expression of secondary metabolic gene clusters, such as sterigmatocystin, penicillin, and lovastatin, but has no effect on spore production in *A. nidulans*. The LaeA that was confirmed as a nuclear protein and a putative methyltransferase does not involve in gene clusters for nutrient utilization (Bok et al., *Mol Microbiol* 61:1636-45, 2006). Furthermore, the role of LaeA in secondary metabolism was confirmed in *Aspergillus flavus* and *Aspergillus oryzae* (Kale et al., *Fungal Genet. Biol.* 45:1422-9, 2008; Oda et al., *Biosci Biotechnol Biochem* 75:1832-4, 2011). Evidence indicates that LaeA reverses gene repression at the level of the heterochromatin state (Reyes-Dominguez et al., *Molecular Microbiology* 76:1376-86, 2010). LaeA is a component of the heterotrimeric VeA/VelB/LaeA protein complex (Bayram et al., *Science Signaling* 320:1504, 2008), which involves in the acetylation signal transduction for secondary metabolite production in *A. nidulans* (Soukup et al., *Mol. Microbiol.*, 86(2):314-30, 2012). The veA/VelB/LaeA complex may coordinately respond to environmental cues (Ramamoorthy et al., *Mol. Microbiol.*, 85(4):795-814, 2012) and has a role in fungal morphology (Calvo, *Fungal Genetics and Biology* 45:1053-61, 2008). LaeA may direct the formation of the VelB-VosA and VelB-VelA-LaeA complexes, control veA modification and protein levels, and be involved in light regulation of growth and development (Bayram et al., *PLoS genetics* 6: e1001226, 2010).

SUMMARY

Although the current commercial conversion rate of carbohydrate to citric acid in *A. niger* is more than eighty to ninety percent, further improvement in production of citric acid and other metabolites is desirable. This disclosure describes the role of the dolichyl-P-Man:Man(5)GlcNAc(2)-PP-dolichyl mannosyltransferase gene (α -1,3-mannosyltransferase, Alg3) on the spore germination, filamentous growth, sporulation, and production of citric acid in *Aspergillus niger*. In addition, the role of LaeA in citric acid production by its over-expression is shown, alone or in combination with an Alg3Δ mutant background.

Based on these observations, provided herein are isolated fungi (such as filamentous fungi) having a gene inactivation

(also referred to herein as a gene deletion) of a dolichyl-P-Man:Man(5)GlcNAc(2)-PP-dolichyl mannosyltransferase (Alg3) gene (referred to herein as Alg3Δ strains), a gene enhancement (e.g., overexpression) of a LaeA gene (referred to herein as upregulated LaeA strains), or both. Any strain of fungi can be used, such as a filamentous fungi, for example *Aspergillus niger* (*A. niger*) or particular strains thereof (for example *A. niger* strain 11414 or 11414KusA). In particular examples, an Alg3Δ strain exhibits one or more of the following characteristics: slower growth on citric acid production (CAP) medium, complete medium (CM) or potato dextrose agar (PDA) medium; earlier spore germination and a higher germination rate in CAP medium; delayed spore germination in CM or PDA medium; reduced sporulation on complete medium; or combinations thereof. In some examples, such increases or decreases are relative to *A. niger* strain 11414KusA grown under the same conditions. The combination of Alg3Δ and over-expression of LaeA resulted in some improvement of sporulation on CM.

In particular examples, such Alg3Δ strains, up-regulated LaeA strains, or Alg3Δ-upregulated LaeA strains, produce more citric acid when grown in CAP medium, such as at least 20%, at least 50%, or at least 70% more than *A. niger* strain 11414KusA under identical growing conditions after at least 4 days, at least 5 days or at least 10 days. Thus, one strategy to increase citric acid production is to reduce the carbohydrate consumption for protein glycosylation and cellular formation, as altering protein glycosylation can augment the carbohydrate flux into citric acid production in *A. niger*.

Also provided herein are compositions (such as fermentation broth) and kits that include a fungal Alg3Δ strain, up-regulated LaeA strain, or Alg3Δ-upregulated LaeA strain.

Also provided herein are methods of making citric acid using the disclosed fungal Alg3Δ strains, up-regulated LaeA strains, and Alg3Δ-upregulated LaeA strains. For example, such a method can include culturing an isolated Alg3Δ fungus, up-regulated LaeA fungus, or Alg3Δ-upregulated LaeA fungus, under conditions that permit the fungus to make citric acid, thereby making citric acid. For example, the Alg3Δ fungus, up-regulated LaeA fungus, or Alg3Δ-upregulated LaeA fungus, can be cultured in CAP medium. In some examples, the method further includes isolating the citric acid produced.

The foregoing and other objects and features of the disclosure will become more apparent from the following detailed description, which proceeds with reference to the accompanying figures.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1A is a schematic drawing showing a restriction map of the 10.9 kb fragment containing the *A. niger* Alg3 gene.

FIG. 1B is a schematic drawing showing the introduction of the hyg-selective marker, which was flanked by the upstream and downstream DNA sequences of Alg3. Integration of the linear molecules by homologous recombination replaces Alg3 with hph in the chromosome.

FIG. 1C is a digital image of a Southern blot showing the genomic DNA hybridization of parent and Alg3Δ strains. One parent and two selected Alg3Δ strains are shown, which have the correct enzyme restriction pattern.

FIG. 2 is a series of digital images showing the Alg3Δ and parent 11414KusA strains grown on agar plates of complete medium (CM), potato dextrose (PDA), and minimal medium (MM) at 30° C. for 15 hrs or 48 hrs. Parent strain (panel A), Alg3Δ strain (panel B), and both parent and Alg3Δ strains (panel C) grown on complete medium plate. Parent strain

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(panel D), Alg3Δ strain (panel E), and both parent and Alg3Δ strains (panel F) were grown on PDA agar plates. Parent strain (panel G), Alg3Δ strain (panel H), and both parent and Alg3Δ strains (panel I) were grown on MM agar plates.

FIG. 3A is a series of digital images showing the spore germination of parent 11414kusA and Alg3Δ strains in liquid cultures of complete medium (CM), potato dextrose medium (PDA), and minimal medium (MM) at 30° C. for 7 hours. Panels A and B are germinated spores in CM liquid culture. Panels C and D are germinated spores in PDA liquid culture. Panels E and F are germinated spores in MM liquid culture. The left panels for the parent 11414kusA strain and the right panels for the Alg3Δ strain.

FIG. 3B provides graphs showing the time courses of the percentage of spore germination of parent 11414kusA and Alg3Δ strains grown in the liquid cultures of complete medium (CM; top graph), potato dextrose medium (PDA; middle graph), or minimal medium (MM; bottom graph). The solid filled cycle is the parent 11414kusA strain and open cycle is the Alg3Δ strain.

FIG. 4 shows a series of digital images (panels A and D) and stereo microscopy digital images (panels B, C, E and F) of parent and Alg3Δ strains grown on agar plates of complete medium (CM) and potato dextrose (PDA) at 30° C. for 4 days.

FIG. 5 provides stereo microscopy digital images parent 11414kusA and Alg3Δ strains grown on citric acid production (CAP) medium plates at different pH levels for 27 hrs. The left panels for parent strain 11414kusA and right panels for the Alg3Δ strain.

FIGS. 6A-6B show spore germination of parent 11414kusA and Alg3Δ strains in citric acid production (CAP) liquid medium. (FIG. 6A, top left and bottom left panel) Inverted microscopic images for parent 11414kusA strain. (FIG. 6A, top right and bottom right panel) Inverted microscopic images for Alg3Δ strain. (FIG. 6A, top panels) Strains grown in CAP liquid culture at 30° C. for 8 hrs. (FIG. 6A, bottom panels) Strains grown in CAP liquid culture at 30° C. for 15 hrs. (FIG. 6B) Time course of the spore germination rate (%) of parent and Alg3Δ strains grown in CAP liquid culture. The solid cycle for parent 11414kusA strain and open cycle for the Alg3Δ strain.

FIG. 7 is a graph showing the time course of citric acid production by parent 11414kusA and Alg3Δ strains in the liquid culture of citric acid production at 30° C. and 200 rpm.

FIGS. 8A and 8B show (FIG. 8A) a schematic diagram showing the construction of the transgene used to complement the Alg3Δ mutant, and (FIG. 8B) a graph showing the citric acid production by parent strain 11414kusA (kusA), Alg3Δ mutant (Alg3) and Alg3Δ mutant complemented with Alg3 gene (cAlg3) after growth at 30° C., 200 rpm for 12 days

FIG. 9 shows an alignment of Alg3 nucleic acid sequences from *A. niger* (top strand, nucleotides 1986-2518 of SEQ ID NO: 1) and *A. oryzae* (bottom strand, nucleotides 643-1175 of SEQ ID NO: 3).

FIGS. 10A and 10B show an alignment of Alg3 protein sequences from *A. niger* (SEQ ID NO: 2), *A. nidulans* (SEQ ID NO: 31), *Fusarium oxysporum* (SEQ ID NO: 32), *Neurospora crassa* (SEQ ID NO: 33), *S. cerevisiae* (SEQ ID NO: 34), *Arabidopsis thaliana* (SEQ ID NO: 35), and *Homo sapiens* (SEQ ID NO: 36). The signs at the top of the alignment show: '-' the average weight of column pair exchanges is less than weight matrix mean value; '.' is less than mean value plus one SD; '+' is less than mean value plus two SD; and '*' is more than mean value plus two SD.

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FIG. 11 shows an alignment of Alg3 protein sequences from *A. niger* (top strand, amino acids 12-411 of SEQ ID NO: 2) and *S. cerevisiae* (bottom strand, amino acids 31-434 of SEQ ID NO: 34).

FIG. 12 shows an alignment of LaeA protein sequences from *A. nidulans* (top strand, amino acids 14-372 of SEQ ID NO: 41) and *A. niger* (bottom strand, amino acids 14-370 of SEQ ID NO: 59).

FIG. 13 is a schematic diagram showing the construction of pPTRpGPDALaeA plasmid vector. Both the glyceraldehyde 3-phosphate dehydrogenase (gpdA) promoter and LaeA (loss of aflR expression A) gene coding sequence of genomic DNA were isolated by overlap PCR from pAN7-1 plasmid vector and *A. nidulans* with additions of HindIII restriction enzyme sites at 5'-end of gpdA promoter and 3'-end of LaeA gene, confirmed by DNA sequence, and ligated into pPTR1 plasmid vector at HindIII restriction enzyme site.

FIG. 14 is a schematic illustrating a plasmid vector pRS426-LaeA, which contains the upstream region of pyrG gene of *A. niger*, the coding region of LaeA gene under the control of gpdA promoter and transcriptional terminator of trpC gene from *A. nidulans*, the pyriothiamine resistance (ptrA) gene from *A. oryzae*, and the downstream region of pyrG gene of *A. niger*. The unique restriction enzyme PmeI site was introduced at the both end of transgene expression fragment.

FIGS. 15A and 15B show digital images of the results of polymerase chain reaction (PCR) analysis of LaeA gene insertion in the transgenic *A. niger* genome of heterologous expression of *A. nidulans* LaeA gene. (FIG. 15A) PCR products of *A. oryzae* ptrA gene detected in selected single spore colony isolate (SCI) of LaeA gene transgenic mutants and parent kusA and Alg3Δ are control strains. (FIG. 15B) PCR products of transgene expression DNA fragment including the gpdA promoter, LaeA coding region and trpC gene transcriptional terminator. The SCI-1 to SCI-4 is the individual single spore colony of LaeA gene transgenic mutants and parent kusA and Alg3Δ are control strains. Lambda DNA marker is the restriction fragment of BstEII restriction enzyme.

FIG. 16 is a bar graph showing the results of citric acid production after 10 days in culture of parent strain (kusA), alg3Δ mutant (Alg3), and over-expression of LaeA gene in alg3Δ (LaeA-1 and LaeA-2) mutants. The data for each strain is the average of three replicates.

FIG. 17A is a schematic diagram showing the construction of the pBSK-LaeA deletion plasmid vector. The DNA fragments of the 5' and 3' ends of *A. niger* LaeA (lack of aflR expression) gene and the hph (hygromycin phosphotransferase) expression cassette were isolated from *A. niger* genomic DNA (LaeA) or pCB 1003 plasmid vector DNA by PCR with oligonucleotide pairs 1 and 2; 3 and 4; and 5 and 6. The PCR DNA fragments were assembled together by Gibson assembly cloning kit in pBSK backbone vector prepared by HindIII/PstI double digestion. The plasmid DNA fragment containing the LaeA gene deletion cassette was prepared by restriction enzyme digestion with restriction endonucleases of HindIII and XbaI and used for *A. niger* transformation.

FIG. 17B is a schematic diagram showing the construction of the pRS426-*A. niger* LaeA complementation plasmid vector. The LaeA gene containing both its promoter and transcriptional terminator was isolated by PCR with oligonucleotide pair 9 and 10 from *A. niger* genomic DNA. The PCR fragment was cloned into the plasmid vector pRSB426-LaeA prepared by restriction endonuclease HindIII digestion and klenow treatment with blunt-end ligation. The new plasmid DNA vector contains the upstream region of the pyrG gene of

A. niger, the entire region of *A. niger* LaeA gene, the transcriptional terminator of the trpC gene from *A. nidulans*, the pyrithiamine resistance (ptrA) gene from *A. oryzae*, and the downstream region of the pyrG gene of *A. niger*. The unique restriction endonuclease PmeI site was introduced at both ends of the transgene expression fragment.

FIG. 18 shows a digital image of the results of PCR analysis of the LaeA gene deletion in the transgenic *A. niger* genome by homologous recombination. The PCR products corresponding to the 5' end of the *A. niger* LaeA gene and part of the hph expression cassette were amplified with oligonucleotide pair 7 and 8, and were detected in selected single spore colony isolate (SCI) of LaeA gene deletion transgenic mutants.

FIG. 19 shows a digital image of the results of PCR analysis of the LaeA gene complementation (ClaeA) in the genetic background of laeA deletion mutant or *A. nidulans* laeA over-expression (laeA) in the genetic background of 11414kusA of those transgenic *A. niger* genome by homologous recombination. PCR products of *A. oryzae* ptrA gene detected in selected single spore colony isolate (SCI) of LaeA gene transgenic mutants and parent kusA (negative) and plasmid DNA of pRS426-laeA (positive) represent control DNA.

FIG. 20 is a bar graph showing the results of citric acid production after 5 days in culture of the parent strain (kusA), the LaeAΔ mutant, *A. niger* LaeA gene complementation in the LaeAΔ mutant (cLaeAΔ), and over-expression of *A. nidulans* LaeA gene in the parent (KusA) strain. The data for each strain is the average of at least three biological replicates.

SEQUENCE LISTING

The nucleic and amino acid sequences listed in the accompanying sequence listing are shown using standard letter abbreviations for nucleotide bases, and three letter code for amino acids, as defined in 37 C.F.R. 1.822. Only one strand of each nucleic acid sequence is shown, but the complementary strand is understood as included by any reference to the displayed strand. The Sequence Listing is submitted as an ASCII text file, created on Oct. 20, 2015, 125 KB, which is incorporated by reference herein. In the accompanying sequence listing:

SEQ ID NOS: 1 and 2 are exemplary Alg3 nucleic acid and protein sequences, respectively, from *A. niger*.

SEQ ID NOS: 3 and 4 are exemplary Alg3 nucleic acid and protein sequences, respectively, from *A. oryzae*.

SEQ ID NOS: 5-30 show exemplary primer sequences.

SEQ ID NOS: 31-36 are exemplary Alg3 protein sequences from *A. nidulans*, *Fusarium oxysporum*, *Arabidopsis thaliana*, *Neurospora crassa*, *S. cerevisiae*, and *Homo sapiens*, respectively.

SEQ ID NO: 37 is an exemplary *Aspergillus nidulans* glyceraldehyde 3-phosphate dehydrogenase (gpdA) promoter sequence.

SEQ ID NOS: 38 and 39 are exemplary forward and reverse primers, respectively, that can be used to isolate or amplify an *A. nidulans* gpdA promoter.

SEQ ID NOS: 40 and 41 are exemplary *Aspergillus nidulans* methyltransferase (LaeA) coding and protein sequences, respectively.

SEQ ID NOS: 42 and 43 are exemplary forward and reverse primers, respectively, that can be used to isolate or amplify an *A. nidulans* LaeA sequence.

SEQ ID NO: 44 is the nucleic acid sequence of the pGPDA-LaeA fragment described in FIG. 13 and Example 5.

SEQ ID NO: 45 is the upstream region of *A. niger* pyrG gene.

SEQ ID NO: 46 is the trpC transcriptional terminator of *A. nidulans*.

SEQ ID NO: 47 is the pyrithiamine resistance gene (ptrA) of *Aspergillus oryzae*.

SEQ ID NO: 48 is the downstream region of *A. niger* pyrG gene.

SEQ ID NOS: 49 and 50 are exemplary forward and reverse primers, respectively, that can be used to isolate or amplify an upstream region of *A. niger* pyrG.

SEQ ID NOS: 51 and 52 are exemplary forward and reverse primers, respectively, that can be used to isolate or amplify the trpC transcriptional terminator of *A. nidulans*.

SEQ ID NOS: 53 and 54 are exemplary forward and reverse primers, respectively, that can be used to isolate or amplify ptrA of *Aspergillus oryzae*.

SEQ ID NOS: 55 and 56 are exemplary forward and reverse primers, respectively, that can be used to isolate or a downstream region of *A. niger* pyrG.

SEQ ID NO: 57 is the nucleic acid sequence of the transgene fragment described in FIG. 13 and Example 6.

SEQ ID NOS: 58 and 59 are exemplary *Aspergillus niger* LaeA coding and protein sequences, respectively.

SEQ ID NO: 60 is the nucleic acid sequence of the transgene fragment used to complement the alg3Δ mutant with the original alg3 gene at pyrG locus.

SEQ ID NO: 61 is the nucleic acid sequence of the 5' end of the *A. niger* LaeA gene.

SEQ ID NO: 62 is the nucleic acid sequence of the 3' end of the *A. niger* LaeA gene.

SEQ ID NO: 63 is the nucleic acid sequence of the hph expression cassette.

SEQ ID NO: 64 is the nucleic acid sequence of the *A. niger* LaeA gene.

SEQ ID NO: 65 is the nucleic acid sequence of the LaeA deletion cassette.

SEQ ID NO: 66 is the nucleic acid sequence of the LaeA complementation construct.

SEQ ID NOS: 67-78 are oligonucleotide primers.

DETAILED DESCRIPTION

Unless otherwise explained, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this disclosure belongs. Definitions of common terms in molecular biology may be found in Benjamin Lewin, *Genes V*, published by Oxford University Press, 1994 (ISBN 0-19-854287-9); Kendrew et al. (eds.), *The Encyclopedia of Molecular Biology*, published by Blackwell Science Ltd., 1994 (ISBN 0-632-02182-9); and Robert A. Meyers (ed.), *Molecular Biology and Biotechnology: a Comprehensive Desk Reference*, published by VCH Publishers, Inc., 1995 (ISBN 1-56081-569-8).

The singular terms "a," "an," and "the" include plural referents unless context clearly indicates otherwise. Similarly, the word "or" is intended to include "and" unless the context clearly indicates otherwise. Hence "comprising A or B" means including A, or B, or A and B. It is further to be understood that all base sizes or amino acid sizes, and all molecular weight or molecular mass values, given for nucleic acids or polypeptides are approximate, and are provided for description. Although methods and materials similar or equivalent to those described herein can be used in the practice or testing of the present disclosure, suitable methods and materials are described below. All references and GENBANK™ Accession numbers mentioned herein are incorporated by reference (the sequence available on Nov. 30, 2011).

The materials, methods, and examples are illustrative only and not intended to be limiting.

In order to facilitate review of the various embodiments of the disclosure, the following explanations of specific terms are provided:

Alg3 (Dolichyl-P-Man: Man(5)GlcNAc(2)-PP-Dolichyl Mannosyltransferase):

Also known as asparagine-linked glycosylation 3 and α -1,3-mannosyltransferase. Alg3 encodes an enzyme which catalyzes the addition of the first dol-p-man derived mannose in an α -1,3 linkage to Man5GlcNAc2-PP-Dol. The term Alg3 (or Alg3) includes any Alg3 gene (such as a fungal Alg3 sequence), cDNA, mRNA, or protein, that is an Alg3 involved in catalyzing the addition of the first dol-p-man derived mannose in an α -1,3 linkage to Man5GlcNAc2-PP-Dol, and when genetically inactivated results in a fungus that has an ability to produce more citric acid than the parent strain (such as at least 20%, at least 30%, at least 50%, at least 60%, or at least 70% more than a parent strain under the same growing conditions).

Alg3 sequences are publicly available for many species of *Aspergillus*. For example, GENBANK™ Accession Nos: XM_001823992.2 and XP_001824044 disclose *Aspergillus oryzae* RIB40 Alg3 nucleic acid and protein sequences, respectively; GENBANK™ Accession Nos: XM_001398659.2 and XP_001398696.2 disclose *Aspergillus niger* CBS 513.88 Alg3 nucleic acid and protein sequences, respectively (SEQ ID NOS: 1 and 2); and GENBANK™ Accession Nos: XM_748359.1 and XP_753452 disclose *Aspergillus fumigatus* Af293 Alg3 nucleic acid and protein sequences, respectively. Additional exemplary Alg3 sequences are provided in SEQ ID NOS: 1-4 and 31-36. However, one skilled in the art will appreciate that in some examples, an Alg3 sequence can include variant sequences (such as allelic variants and homologs) that retain Alg3 activity but when genetically inactivated in *Aspergillus* results in a fungus that has an ability to produce more citric acid than the parent strain (such as at least 20%, at least 30%, at least 50%, at least 60%, or at least 70% more under the same growing conditions).

Detectable: Capable of having an existence or presence ascertained. For example, production of citric acid is detectable if the signal generated is strong enough to be measurable.

Genetic Enhancement or Up-Regulation: When used in reference to the expression of a nucleic acid molecule, such as a gene, refers to any process which results in an increase in production of a gene product. A gene product can be RNA (such as mRNA, rRNA, tRNA, and structural RNA) or protein. Examples of processes that increase transcription include those that facilitate formation of a transcription initiation complex, those that increase transcription initiation rate, those that increase transcription elongation rate, those that increase processivity of transcription and those that relieve transcriptional repression (for example by blocking the binding of a transcriptional repressor). Gene up-regulation can include inhibition of repression as well as stimulation of expression above an existing level. Examples of processes that increase translation include those that increase translational initiation, those that increase translational elongation and those that increase mRNA stability. In one example, additional copies of genes are introduced into a cell in order to increase expression of that gene in the resulting transgenic cell.

Gene up-regulation includes any detectable increase in the production of a gene product. In certain examples, production of a gene product increases by at least 1.5-fold, at least 2-fold, or at least 5-fold), such as LaeA. For example, a genetic enhancement of a LaeA gene in *Aspergillus* (e.g., *A. niger*)

results in an *Aspergillus* strain having increased levels of the LaeA protein relative to the parent strain, which can increase the ability of the fungus to produce more citric acid. Genetic enhancement is also referred to herein as “enhancing or increasing expression.”

Genetic Inactivation or Down-Regulation: When used in reference to the expression of a nucleic acid molecule, such as a gene, refers to any process which results in a decrease in production of a gene product. A gene product can be RNA (such as mRNA, rRNA, tRNA, and structural RNA) or protein. Therefore, gene down-regulation or deactivation includes processes that decrease transcription of a gene or translation of mRNA.

For example, a mutation, such as a substitution, partial or complete deletion, insertion, or other variation, can be made to a gene sequence that significantly reduces (and in some cases eliminates) production of the gene product or renders the gene product substantially or completely non-functional. For example, a genetic inactivation of an Alg3 gene in *Aspergillus* (e.g., *A. niger*) results in *Aspergillus* having a non-functional or non-existent Alg3 protein, which results in an ability of the fungus to produce more citric acid. Genetic inactivation is also referred to herein as “functional deletion”.

Isolated: To be significantly separated from other agents. An “isolated” biological component (such as a nucleic acid molecule or protein) has been substantially separated, produced apart from, or purified away from other biological components in the cell of the organism in which the component occurs, for example, other chromosomal and extra-chromosomal DNA and RNA, and proteins. Nucleic acid molecules and proteins which have been “isolated” include nucleic acid molecules and proteins purified by standard purification methods. The term also embraces nucleic acid molecules and proteins prepared by recombinant expression in a host cell as well as chemically synthesized proteins and nucleic acids. Samples of isolated biological components include samples of the biological component wherein the biological component represents greater than 90% (for example, greater than 95%, such as greater than 98%) of the sample.

An “isolated” microorganism (such as an Alg3A strain of *Aspergillus*) has been substantially separated or purified away from microorganisms of different types, strains, or species. Microorganisms can be isolated by a variety of techniques, including serial dilution and culturing and resistance to certain chemicals.

LaeA (loss of aflR Expression A): LaeA encodes a protein which regulates secondary metabolite production in *Aspergillus*. The term LaeA (or LaeA) includes any LaeA gene (such as a fungal LaeA sequence), cDNA, mRNA, or protein, that is an LaeA involved in secondary metabolite production, and when its expression is increased, for example in combination with a genetically inactivated Alg3 gene, results in a fungus that has an ability to produce more citric acid than the parent strain (such as at least 20%, at least 30%, at least 40%, 50%, at least 60%, or at least 70% more than a parent strain under the same growing conditions).

LaeA sequences are publicly available for many species of *Aspergillus*. For example, GENBANK™ Accession Nos: AB267276 and BAF74528.1 disclose *Aspergillus oryzae* LaeA nucleic acid and protein sequences, respectively; GENBANK™ Accession No. EHA27020.1 discloses an exemplary *Aspergillus niger* ATCC1015 LaeA protein sequence, a parent strain of 11414kusA (other exemplary sequences are provided in SEQ ID NOS: 58 and 59); GENBANK™ Accession No: CBF88745 discloses an *Aspergillus nidulans* LaeA protein sequence; and GENBANK™ Accession Nos:

AY422723 and AAR01218 disclose *Aspergillus fumigatus* LaeA nucleic acid and protein sequences, respectively. Additional exemplary LaeA sequences are provided in SEQ ID NOS: 40-41 and 58-59. However, one skilled in the art will appreciate that in some examples, an LaeA sequence can include variant sequences (such as allelic variants and homologs) that retain LaeA activity and when genetically up-regulated in *Aspergillus* (for example with addition of copy or in combination with Alg3Δ) results in a fungus that has an ability to produce more citric acid than the parent strain (such as at least 20%, at least 30%, at least 40%, at least 50%, at least 60%, or at least 70% more under the same growing conditions).

Mutation: A change in a nucleic acid sequence (such as a gene sequence) or amino acid sequence, for example as compared to a nucleic acid or amino acid sequence present in a wild-type or native organism. In particular examples, a mutation is introduced into an Alg3 gene in *Aspergillus*. Mutations can occur spontaneously, or can be introduced, for example using molecular biology methods. In particular examples, a mutation includes one or more nucleotide substitutions, deletions, insertions, or combinations thereof. In particular examples, the presence of one or more mutations in a gene can significantly inactivate that gene.

Recombinant: A recombinant nucleic acid molecule or protein is one that has a sequence that is not naturally occurring or has a sequence that is made by an artificial combination of two otherwise separated segments of sequence. In particular examples, this artificial combination is accomplished by chemical synthesis or by the artificial manipulation of isolated segments of nucleic acids, for example, by genetic engineering techniques such as those described in Sambrook et al. (ed.), *Molecular Cloning: A Laboratory Manual*, 3d ed., vol. 1-3, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y., 2001. The term recombinant includes nucleic acid molecules that have been altered solely by addition, substitution, or deletion of a portion of the nucleic acid molecule.

Sequence identity/similarity: The identity/similarity between two or more nucleic acid sequences, or two or more amino acid sequences, is expressed in terms of the identity or similarity between the sequences. Sequence identity can be measured in terms of percentage identity; the higher the percentage, the more identical the sequences are. Sequence similarity can be measured in terms of percentage similarity (which takes into account conservative amino acid substitutions); the higher the percentage, the more similar the sequences are.

Methods of alignment of sequences for comparison are well known in the art. Various programs and alignment algorithms are described in: Smith & Waterman, *Adv. Appl. Math.* 2:482, 1981; Needleman & Wunsch, *J. Mol. Biol.* 48:443, 1970; Pearson & Lipman, *Proc. Natl. Acad. Sci. USA* 85:2444, 1988; Higgins & Sharp, *Gene*, 73:237-44, 1988; Higgins & Sharp, *CABIOS* 5:151-3, 1989; Corpet et al., *Nuc. Acids Res.* 16:10881-90, 1988; Huang et al. *Computer Appls. in the Biosciences* 8, 155-65, 1992; and Pearson et al., *Meth. Mol. Bio.* 24:307-31, 1994. Altschul et al., *J. Mol. Biol.* 215: 403-10, 1990, presents a detailed consideration of sequence alignment methods and homology calculations.

The NCBI Basic Local Alignment Search Tool (BLAST) (Altschul et al., *J. Mol. Biol.* 215:403-10, 1990) is available from several sources, including the National Center for Biological Information (NCBI, National Library of Medicine, Building 38A, Room 8N805, Bethesda, Md. 20894) and on the Internet, for use in connection with the sequence analysis

programs blastp, blastn, blastx, tblastn and tblastx. Additional information can be found at the NCBI web site.

BLASTN is used to compare nucleic acid sequences, while BLASTP is used to compare amino acid sequences. To compare two nucleic acid sequences, the options can be set as follows: -i is set to a file containing the first nucleic acid sequence to be compared (e.g., C:\seq1.txt); -j is set to a file containing the second nucleic acid sequence to be compared (e.g., C:\seq2.txt); -p is set to blastn; -o is set to any desired file name (e.g., C:\output.txt); -q is set to -1; -r is set to 2; and all other options are left at their default setting. For example, the following command can be used to generate an output file containing a comparison between two sequences: C:\B12seq -i c:\seq1.txt -j c:\seq2.txt -p blastn -o c:\output.txt -q -1 -r 2.

To compare two amino acid sequences, the options of B12seq can be set as follows: -i is set to a file containing the first amino acid sequence to be compared (e.g., C:\seq1.txt); -j is set to a file containing the second amino acid sequence to be compared (e.g., C:\seq2.txt); -p is set to blastp; -o is set to any desired file name (e.g., C:\output.txt); and all other options are left at their default setting. For example, the following command can be used to generate an output file containing a comparison between two amino acid sequences: C:\B12seq -i c:\seq1.txt -j c:\seq2.txt -p blastp -o c:\output.txt. If the two compared sequences share homology, then the designated output file will present those regions of homology as aligned sequences. If the two compared sequences do not share homology, then the designated output file will not present aligned sequences.

Once aligned, the number of matches is determined by counting the number of positions where an identical nucleotide or amino acid residue is presented in both sequences. The percent sequence identity is determined by dividing the number of matches either by the length of the sequence set forth in the identified sequence, or by an articulated length (e.g., 100 consecutive nucleotides or amino acid residues from a sequence set forth in an identified sequence), followed by multiplying the resulting value by 100. For example, a nucleic acid sequence that has 1166 matches when aligned with a test sequence having 1554 nucleotides is 75.0 percent identical to the test sequence (i.e., $1166 \div 1554 \times 100 = 75.0$). The percent sequence identity value is rounded to the nearest tenth. For example, 75.11, 75.12, 75.13, and 75.14 are rounded down to 75.1, while 75.15, 75.16, 75.17, 75.18, and 75.19 are rounded up to 75.2. The length value will always be an integer. In another example, a target sequence containing a 20-nucleotide region that aligns with 20 consecutive nucleotides from an identified sequence as follows contains a region that shares 75 percent sequence identity to that identified sequence (i.e., $15 \div 20 \times 100 = 75$).

For comparisons of amino acid sequences of greater than about 30 amino acids, the Blast 2 sequences function is employed using the default BLOSUM62 matrix set to default parameters, (gap existence cost of 11, and a per residue gap cost of 1). Homologs are typically characterized by possession of at least 70% sequence identity counted over the full-length alignment with an amino acid sequence using the NCBI Basic Blast 2.0, gapped blastp with databases such as the nr or swissprot database. Queries searched with the blastn program are filtered with DUST (Hancock and Armstrong, 1994, *Comput. Appl. Biosci.* 10:67-70). Other programs use SEG. In addition, a manual alignment can be performed. Proteins with even greater similarity will show increasing percentage identities when assessed by this method, such as at least 75%, 80%, 85%, 90%, 95%, or 99% sequence identity.

Nucleic acid sequences that do not show a high degree of identity may nevertheless encode identical or similar (conserved) amino acid sequences, due to the degeneracy of the genetic code. Changes in a nucleic acid sequence can be made using this degeneracy to produce multiple nucleic acid molecules that all encode substantially the same protein. Such homologous nucleic acid sequences can, for example, possess at least 60%, 70%, 80%, 90%, 95%, 98%, or 99% sequence identity determined by this method.

One of skill in the art will appreciate that these sequence identity ranges are provided for guidance only; it is possible that strongly significant homologs could be obtained that fall outside the ranges provided.

Transformed: A cell, such as a fungal cell, into which a nucleic acid molecule has been introduced, for example by molecular biology methods known in the art. As used herein, the term transformation encompasses all techniques by which a nucleic acid molecule might be introduced into such a cell, including, but not limited to transfection with viral vectors, conjugation, transformation with plasmid vectors, and introduction of naked DNA by chemical-mediated, electroporation, lipofection, and biolistic particle delivery.

Overview

This disclosure provides the first demonstration that genetic inactivation of Alg3, a gene involved in protein N-linked glycosylation, can result in substantial improvement of citric acid production in *A. niger*, while the total biomass is similar to the parent strain. The core oligosaccharide Glc3Man9GlcNAc2 is synthesized by a series of membrane-bound glycosyltransferases, which begins on the cytoplasmic side of the membrane of the endoplasmic reticulum (ER) and flips into the luminal side of the ER membrane to complete its synthesis. The lipid-linked core Glc3Man9GlcNAc2 is subsequently transferred to a nascent protein in the ER, where the glycoproteins are folded and then shuttled to the Golgi for additional, but divergent processing. The Alg3 gene encodes the enzyme α -1,3-mannosyltransferase that converts Man5GlcNAc2-Dol-PP to Man6GlcNAc2-Dol-PP on the ER membrane of the luminal side. Provided herein is a homolog of *Saccharomyces cerevisiae* Alg3 identified from *Aspergillus niger* (e.g., see SEQ ID NOS: 1 and 2).

It is shown herein that genetic inactivation of Alg3 in *A. niger* resulted in a significant reduction of growth on complete medium (CM) and potato dextrose agar medium (PDA), but no effect on minimal medium (MM). The Alg3 deletion also caused the substantial reduction in spore production of *A. niger* on CM, but no significant change on the PDA. When the spores were germinated in CM or PDA liquid culture medium, the Alg3 Δ strain showed pronounced delay in spore germination. This growth phenotype is similar to the mutants with defects in signal transduction pathways observed in *A. nidulans* and *A. niger* (Fillinger et al., *Mol. Microbiol.* 44(4): 1001-16, 2002; Saudohar et al., *Microbiol.* 148(8):2635-45, 2002; Xue et al., *Eukaryot Cell* 3(2):557-60, 2004). Deletion of *pkaA*, *cycaA* or *schA/pkaA* in *A. nidulans* substantially reduces its growth on CM medium plates and spore germination rate in MM liquid culture medium (Fillinger et al., *Mol. Microbiol.* 44(4):1001-1016, 2002) and similar growth phenotypes were observed in the strains with the deletion of *pkaR*, *pkaC* or double deletion of *pkaR/pkaC* in *A. niger* (Saudohar et al., *Microbiol.* 148(8):2635-2645, 2002). However, functional deletion of the MAP kinase *SakA* in *A. fumigatus* delays the spore germination in liquid CM, but stimu-

lates spore germination in MM liquid medium (Xue et al., *Eukaryot Cell* 3(2): 557-560, 2004).

Furthermore, the Alg3 deletion reduced the overall growth on citric acid production (CAP) medium plates at different pHs. In contrast, the Alg3 deletion triggered early spore germination and substantially improved spore germination rate in CAP liquid culture medium. Citric acid production in CAP liquid culture medium was significantly improved in *A. niger*. When the alg3 Δ mutant was complemented with the original alg3 gene at *pyrG* locus (FIG. 8A; SEQ ID NO: 60), its transcription levels was similar to parent strain with the cycle threshold (Ct) values about 23, while the Ct value for alg3 Δ mutant was 30.8 in CAP liquid culture conditions, which was determined by real-time reverse-transcription PCR. Consequently, the citric acid production in the resulted complemented mutant strains was similar to the parent strain as shown in FIG. 8B. The results shown herein demonstrate the involvement of Alg3 on the growth and development and citric acid production in *A. niger*.

It is proposed that inactivation of Alg3 influences the N-glycosylation of those proteins involving in signal transduction pathways. The N-glycosylation consensus sequence (N-glycosite) for N-glycosylation in those proteins from the signal transduction pathways was observed. Most of those proteins contained 1 to 7 N-glycosites, such as, 6 N-glycosites found in *sskB* (map kinase kinase kinase), 7 in *Ste11/SteC*, 5 in *acyA*, 5 in *rgsA*, 6 in *rgsC*, 4 in *gprA*, 4 in *pkaC2*, 4 in *flbA* and 3 in *G β* . Comparison of these results with previous studies indicates that the effects of the Alg3 deletion on spore germination and growth may be regulated by altering the N-glycosylation in those proteins involved in signal transduction pathways in *A. niger*.

When the Alg3 Δ strain was grown on CM medium, spore production of Alg3 Δ mutants was dramatically reduced as compared to the parent strain, while maintaining a similar level when grown on PDA medium. This phenotype of sporulation production may be influenced by both endogenous and exogenous factors. For example, protein glycosylation was greatly influenced by culture conditions in filamentous fungi, such as fully glycosylated *Cel7A* only isolated from MM culture medium (Stals et al., *Glycobiology* 14(8):725-737, 2004). In addition, higher amounts of proteases were secreted by the Alg3 Δ strain than the parent in liquid MM culture supplemented 1 g/l yeast extract, which further influenced nutrient uptakes, cellular formation and overall N-glycosylation. This would alter the yield and N-glycosylation in G protein system in *A. niger*, where G protein signaling is crucial for detection of major environmental stimuli for food acquisition, asexual sporulation, and spore germination (Chang et al., *Genetics* 167(3):305, 2004; Li et al., *Annu. Rev. Microbiol.* 61:423-452, 2007).

The spores of parent strain germinated more slowly and had a lower germination rate than the Alg3 Δ strain in CAP liquid culture medium, which contains limited nitrogen source (3.1 g/l of NH_4NO_3), similar to MM. A similar phenotype was observed when the stress activating kinase, a MAP kinase, was deleted in *A. fumigatus* (SakAA strain) and grown in MM liquid culture medium (Xue et al., *Eukaryot Cell* 3(2): 557-560, 2004). The spore germination of SakAA strain was dramatically influenced by nitrogen sources. For example, similar rates of spore germination between parent and SakAA strains were observed on MM containing 10 mM NH_4Cl or 10 mM Pro, while the spore germination rates of SakAA strain was much higher than the parent strain in the MM culture medium containing 10 mM NaNO_3 , NaNO_2 , or Phe. In addition, the CAP medium contains high level of glucose and low pH, which contributes additional stresses to

A. niger growth. Although the Alg3Δ strain had earlier and higher germination in CAP medium, its biomass formation was less than the parent strain at early stages. The dried biomass yields for both parent and Alg3Δ strains were similar after growth in CAP medium for four and half days. However, more citric acid was produced by the Alg3Δ strain than the parent strain. This indicates more glucose was directly converted to citric acid by influence citric acid metabolism and reduction of glucose consumption for complex N-glycan formation and sequentially for other cellular metabolisms.

This disclosure also provides the first demonstration that genetic inactivation of Alg3, in combination with an increase in expression of the loss of aflR expression A (LaeA) gene, can result in substantial improvement of citric acid production in *A. niger*. It is proposed that increased expression of LaeA can also improve citric acid production in *A. niger* or other filamentous fungi. To increase expression of LaeA in fungal cells, a transgene was generated and expressed in *A. niger* as follows. The LaeA gene of *A. nidulans* was operably controlled by glyceraldehyde 3-phosphate dehydrogenase (gpdA) promoter and trpC transcriptional terminator (TtrpC) of *A. nidulans*. This chimeric gene was flanked with the upstream of *A. niger* pyrG gene, the pyrithiamine resistance (ptrA) gene of *A. oryzae* and the downstream of *A. niger* pyrG gene. The transgene expression fragment containing the chimeric gene was used to transform the protoplasts of alg3Δ mutants of *A. niger*.

The present disclosure also describes the generation of an *A. niger* strain having a deletion of the LaeA gene (LaeAΔ), as well as an *A. niger* LaeAΔ strain complemented with a transgene encoding LaeA (cLaeAΔ) at the pyrG locus. These strains were used to evaluate the role of LaeA expression on citric acid production in *Aspergillus*. The data disclosed herein demonstrates that deletion of the LaeA gene in *A. niger* (LaeAΔ) results in a loss of citric acid production in culture. When the original *A. niger* LaeA gene was used for complementation in the LaeAΔ mutant (cLaeAΔ) at the pyrG locus, citric acid production was partially recovered, which indicates the importance of the chromosomal location of LaeA gene. When *A. nidulans* LaeA was over-expressed in the *A. niger* parent strain, citric acid production was higher than the parent strain. These results indicate that LaeA enhances citric acid production in *A. niger*.

In summary, the deletion of Alg3, increasing expression of LaeA, or both, can be used to increase citric acid production in fungi (such as filamentous fungi, e.g., *A. niger*). In addition, deletion of Alg3 alters the overall N-glycosylation and further influences the spore germination, filamentous growth, sporulation and other organic acid production in *A. niger*.

Alg3Δ Fungi

The present disclosure provides isolated fungi having its Alg3 gene inactivated, wherein such inactivation results in increased citric acid production by the fungi. Such fungi are referred to herein as Alg3Δ fungi. It is disclosed herein that genetic inactivation of Alg3 results in *Aspergillus* fungi that can increase citric acid production as compared to *Aspergillus* having a native Alg3 sequence.

Contemplated herein are isolated fungi containing a genetic inactivation of a dolichyl-P-Man:Man(5)GlcNAc(2)-PP-dolichyl mannosyltransferase gene (Alg3). Any fungus can be used, such as any genus or variety of *Aspergillus*. In particular examples, the disclosed *Aspergillus* fungus is *A. niger*, such as *Aspergillus niger* strain 11414 (American Type Culture Collection (ATCC) No. 11414; NRRL 2270); 1015 (ATCC No. 1015; NRRL 328, CBS 113.46); NRRL 3 (ATCC

No. 9029, CBS 120.49, N400); NRRL 3122 (ATCC No. 22343); or 11414KusA-. In other specific examples, the *Aspergillus* is *A. aculeatus*, *A. awamori*, *A. carbonarius*, *A. wentii*, *A. foetidus*, *A. oryzae*, *A. terreus*, or *A. fumigatus*.

In addition, any method for genetic inactivation can be used, as long as the expression of the gene is significantly reduced or eliminated, or the function of the expressed protein is significantly reduced or eliminated. In particular examples, the Alg3 gene is genetically inactivated by complete or partial deletion mutation or by insertional mutation. In some examples genetic inactivation need not be 100% genetic inactivation. In some embodiments, genetic inactivation refers to at least 50%, at least 60%, at least 70%, at least 80%, at least 90%, or at least 95% gene or protein inactivation. The term "reduced" or "decreased" as used herein with respect to a cell and a particular gene or protein activity refers to a lower level of activity than that measured in a comparable cell of the same species. For example, a particular fungi lacking Alg3 activity has reduced Alg3 activity if a comparable fungi not having an Alg3 genetic inactivation has detectable Alg3 activity.

Alg3 sequences are disclosed herein and others are publicly available, for example from GENBANK™ or EMBL. In some examples, the Alg3 gene functionally deleted encodes a protein having at least 80%, at least 90%, at least 95%, at least 97%, or at least 98% sequence identity to SEQ ID NO: 2, 4, 31, 32, 33, 34, 35, or 36. In some examples, the Alg3 gene functionally deleted comprises at least 80%, at least 90%, at least 95%, at least 97%, or at least 98% sequence identity to SEQ ID NO: 1 or 3 or nucleotides 1186-2582 of SEQ ID NO: 1.

The inactivation of Alg3 results in many phenotypes in the fungi. For example, Alg3Δ mutants can have one or more of the following phenotypes: slower growth on citric acid production (CAP) medium, earlier spore germination in CAP medium (for example germination in at least 3 hours, at least 4 hours, or at least 5 hours after inoculation, such as within 3 hours of inoculation), increased spore germination rate in CAP medium, increased citric acid production in CAP medium, slower growth on complete medium (CM) or potato dextrose (PDA) medium, delay initiation of spore germination in CM or PDA medium, reduced sporulation on CM, or combinations thereof.

Such changes (such as increases or decreases) can be relative to a fungi having a wild-type Alg3 gene, such as a parental strain (e.g., *A. niger* strain 11414KusA), grown under the same conditions as the Alg3Δ mutant. In some examples, an increased germination rate is germination of at least 20%, at least 25%, or at least 30% of the spores from an Alg3Δ fungus have germinated 8 hours after inoculation in CAP medium (such as 20% to 35%, such as 32%), as compared to no more than 20%, no more than 15%, or no more than 10% (such as 5 to 15%, or 10%) for *A. niger* strain 11414KusA. In some examples, an increased germination rate is germination of at least 80%, at least 85%, or at least 90% of the spores from an Alg3Δ fungus have germinated 15 hours after inoculation in CAP medium (such as 80% to 95%, such as 90%), as compared to no more than 60%, no more than 65%, or no more than 75% (such as 55 to 65%, or 60%) for *A. niger* strain 11414KusA. In some examples, increased citric acid production in CAP medium is an increase of at least 10%, at least 20%, at least 30%, at least 40%, at least 50%, at least 60%, at least 65%, or at least 70%, by an Alg3Δ fungus as compared to *A. niger* strain 11414KusA. In some examples, reduced sporulation on complete medium is a reduction of sporulation by at least 20%, at least 30%, at least 40%, at least 50%, or at

least 60%, (such as a 40% to 60% reduction) by an Alg34 fungus as compared to *A. niger* strain 11414KusA.

One skilled in the art will appreciate that additional genes can also be inactivated, wherein the additional genes may or may not provide additional enhancement of citric acid production to the fungus. In one example KusA (e.g., GENBANK™ Accession No. EF061656) is also genetically inactivated.

Also provided by the present disclosure are compositions that include isolated Alg3Δ fungi, such as a growth medium. Also provided by the present disclosure are kits that include isolated Alg3Δ fungi, such as a kit that includes a medium for culturing, storing, or growing the fungus. Exemplary mediums include solid medium (such as those containing agar, for example CM, PDA or MM) and liquid media (such as a fermentation broth, such as CM, MM, or CAP medium).

A. Methods of Functionally Deleting Genes

As used herein, an “inactivated” or “functionally deleted” gene means that the gene has been mutated, for example by insertion, deletion, or substitution (or combinations thereof) of one or more nucleotides such that the mutation substantially reduces (and in some cases abolishes) expression or biological activity of the encoded gene product. The mutation can act through affecting transcription or translation of the gene or its mRNA, or the mutation can affect the polypeptide product itself in such a way as to render it substantially inactive.

Genetic inactivation of one or more genes (which in some examples is also referred to as functional deletion) can be performed using any conventional method known in the art. In one example, a strain of *Aspergillus* is transformed with a vector which has the effect of down-regulating or otherwise inactivating an Alg3 gene. This can be done by mutating control elements such as promoters and the like which control gene expression, by mutating the coding region of the gene so that any protein expressed is substantially inactive, or by deleting the Alg3 gene entirely. For example, an Alg3 gene can be functionally deleted by complete or partial deletion mutation (for example by deleting a portion of the coding region of the gene) or by insertional mutation (for example by inserting a sequence of nucleotides into the coding region of the gene, such as a sequence of about 1-5000 nucleotides). Thus, the disclosure in some examples provides transformed fungi that include at least one exogenous nucleic acid molecule which genetically inactivates an Alg3 gene (such as a nucleic acid sequence encoding SEQ ID NO: 2 or 4). In one example, such a transformed cell produces more citric acid, for example relative to a comparable fungus with a native Alg3 sequence.

In particular examples, an insertional mutation includes introduction of a sequence that is in multiples of three bases (e.g., a sequence of 3, 9, 12, or 15 nucleotides) to reduce the possibility that the insertion will be polar on downstream genes. For example, insertion or deletion of even a single nucleotide that causes a frame shift in the open reading frame, which in turn can cause premature termination of the encoded Alg3 polypeptide or expression of a substantially inactive polypeptide. Mutations can also be generated through insertion of foreign gene sequences, for example the insertion of a gene encoding antibiotic resistance (such as hygromycin or bleomycin).

In one example, genetic inactivation is achieved by deletion of a portion of the coding region of the Alg3 gene. For example, some, most (such as at least 50%) or virtually the entire coding region can be deleted. In particular examples, about 5% to about 100% of the gene is deleted, such as at least

20% of the gene, at least 40% of the gene, at least 75% of the gene, or at least 90% of the Alg3 gene.

Deletion mutants can be constructed using any of a number of techniques known in the art. In one example, allelic exchange is employed to genetically inactivate one or more genes in *Aspergillus*. A specific example of such a method is described in Example 2 below.

In one example, a strategy using counterselectable markers can be employed which has been utilized to delete genes. For a review, see Reyrat et al. (*Infec. Immun.* 66:4011-4017, 1998). In this technique, a double selection strategy is employed wherein a plasmid is constructed encoding both a selectable and counterselectable marker, with flanking DNA sequences derived from both sides of the desired deletion. The selectable marker is used to select for fungi in which the plasmid has integrated into the genome in the appropriate location and manner. The counterselectable marker is used to select for the very small percentage of fungi that have spontaneously eliminated the integrated plasmid. A fraction of these fungi will then contain only the desired deletion with no other foreign DNA present.

In another technique, the cre-lox system is used for site specific recombination of DNA (for example see Steiger et al., *Appl. Environ. Microbiol.* 77(1):114, 2011). The system includes 34 base pair lox sequences that are recognized by the bacterial cre recombinase gene. If the lox sites are present in the DNA in an appropriate orientation, DNA flanked by the lox sites will be excised by the cre recombinase, resulting in the deletion of all sequences except for one remaining copy of the lox sequence. Using standard recombination techniques, the targeted gene of interest (e.g., Alg3) can be deleted in the *Aspergillus* genome and to replace it with a selectable marker (for example a gene coding for kanamycin resistance) that is flanked by the lox sites. Transient expression (by electroporation of a suicide plasmid containing the cre gene under control of a promoter that functions in *Aspergillus*) of the cre recombinase should result in efficient elimination of the lox flanked marker. This process will produce a mutant containing the desired deletion mutation and one copy of the lox sequence.

In another method, an Alg3 gene sequence in the *Aspergillus* genome is replaced with a marker gene, such as green fluorescent protein, β-galactosidase, or luciferase. In this technique, DNA segments flanking a desired deletion are prepared by PCR and cloned into a suicide (non-replicating) vector for *Aspergillus*. An expression cassette, containing a promoter active in *Aspergillus* and the appropriate marker gene, is cloned between the flanking sequences. The plasmid is introduced into wild-type *Aspergillus*. Fungi that incorporate and express the marker gene are isolated and examined for the appropriate recombination event (replacement of the wild type Alg3 gene with the marker gene).

Thus, for example, a fungal cell can be engineered to have a disrupted Alg3 gene using common mutagenesis or knock-out technology. (Methods in Yeast Genetics (1997 edition), Adams, Gottschling, Kaiser, and Sterns, Cold Spring Harbor Press, 1998; Datsenko and Wanner, *Proc. Natl. Acad. Sci. USA* 97: 6640-5, 2000; and Dai et al., *Appl. Environ. Microbiol.* 70(4):2474-85, 2004). Alternatively, antisense technology can be used to reduce or eliminate the activity of Alg3. For example, a fungal cell can be engineered to contain a cDNA that encodes an antisense molecule that prevents Alg3 from being translated. The term “antisense molecule” encompasses any nucleic acid molecule or nucleic acid analog (e.g., peptide nucleic acids) that contains a sequence that corresponds to the coding strand of an endogenous Alg3 gene. An antisense molecule also can have flanking sequences (e.g.,

regulatory sequences). Thus, antisense molecules can be ribozymes or antisense oligonucleotides. A ribozyme can have any general structure including, without limitation, hairpin, hammerhead, or axehead structures, provided the molecule cleaves RNA. Further, gene silencing can be used to reduce the activity of Alg3.

B. Measuring Gene Inactivation

A fungus having an inactivated Alg3 gene can be identified using any method known in the art. For example, PCR and nucleic acid hybridization techniques, such as Northern and Southern analysis, can be used to confirm that a fungus has an inactivated Alg3 gene. Alternatively, real-time reverse transcription PCR (qRT-PCR) can be used for detection and quantification of targeted messenger RNA, such as mRNA of Alg3 gene in the parent and mutant strains as grown at the same culture conditions. Immunohisto-chemical and biochemical techniques can also be used to determine if a cell expresses Alg3 by detecting the expression of the Alg3 peptide encoded by Alg3. For example, an antibody having specificity for Alg3 can be used to determine whether or not a particular fungus contains a functional nucleic acid encoding Alg3 protein. Further, biochemical techniques can be used to determine if a cell contains a particular gene inactivation by detecting a product produced as a result of the expression of the peptide. For example, structural determination of N-glycans excised from glycoproteins can indicate that a fungal cell contains an inactivated Alg3 gene. In addition, measurements of sporulation, germination, secondary metabolite production, and citric acid production can be measured using the methods described herein.

C. Measuring Citric Acid Production

Methods of determining whether a genetic inactivation of Alg3 in *Aspergillus* increases citric acid production, for example relative to the same strain with a native Alg3 sequence (such as a parental strain), are routine in the art. Although particular examples are disclosed herein, the methods are not limiting.

For example, production of citric acid by *Aspergillus* (such as an Alg3Δ strain) can be measured using a spectrophotometric assay. In one example citric acid production can be determined with an endpoint spectrophotometric enzyme assay (for example see, Bergmeyer, H. U. 1985. *Metabolites 2: tri- and dicarboxylic acids, purines, pyrimidines and derivatives, coenzymes, inorganic compounds*, p. 5-10. In *Citric acids*. VCH Publishers, Weinheim, Germany). Citric acid can also be measured by liquid chromatography (LC) or high-performance liquid chromatography (HPLC) methods.

D. Alg3 Sequences

Alg3 protein and nucleic acid sequences are publicly available and specific examples are provided herein. In addition, Alg3 sequences can be identified using routine molecular biology methods.

Examples of Alg3 nucleic acid sequences shown in SEQ ID NOS: 1 and 3. However, the disclosure also encompasses variants of SEQ ID NOS: 1 and 3 which retain the ability to encode an Alg3 protein. One skilled in the art will understand that variant Alg3 nucleic acid sequences can be inactivated. Variant sequences may contain a single insertion, a single deletion, a single substitution, multiple insertions, multiple deletions, multiple substitutions, or any combination thereof (e.g., single deletion together with multiple insertions). In addition, the degeneracy of the code permits multiple nucleic acid sequences to encode the same protein. For example, FIG. 9 shows an alignment of Alg3 nucleic acid sequences from *A. niger* (nucleotides 1986-2518 of SEQ ID NO: 1) and *A. oryzae* (nucleotides 643-1175 of SEQ ID NO: 3), which permits one to identify nucleotides that can tolerate substitu-

tion (e.g., those that are not conserved between species) and those that may not (e.g., those that are conserved between species). Such nucleic acid molecules can share at least 80%, at least 85%, at least 90%, at least 95%, at least 97%, at least 98%, or at least 99% sequence identity to any known Alg3 nucleic acid sequence, such as SEQ ID NO: 1 or 3 or nucleotides 1186-1306, 1393-1916 and 1989-2582 of SEQ ID NO: 1.

Examples of Alg3 protein sequences shown in SEQ ID NOS: 2, 4, 31, 32, 33, 34, 35, and 36. However, the disclosure also encompasses variants SEQ ID NOS: 2, 4, 31, 32, 33, 34, 35, and 36 which retain Alg3 activity. One skilled in the art will understand that variant Alg3 enzyme sequences can be inactivated. Variant sequences may contain a single insertion, a single deletion, a single substitution, multiple insertions, multiple deletions, multiple substitutions, or any combination thereof (e.g., single deletion together with multiple insertions). Such polypeptides share at least 80%, at least 85%, at least 90%, at least 95%, at least 97%, at least 98%, or at least 99% sequence identity to an Alg3 sequence, such as SEQ ID NO: 2, 4, 31, 32, 33, 34, 35, or 36.

Variant sequences can be identified, for example by aligning known Alg3 sequences. For example, FIGS. 10A and 10B show the alignment of seven different Alg3 sequences from different organisms. In addition, FIG. 11 shows a detailed alignment of Alg3 protein sequences from *A. niger* (amino acids 12-411 of SEQ ID NO: 2) and *S. cerevisiae*, indicating amino acids that are identical, conserved (+) or not conserved (space). Based on these alignments, variants of Alg3 sequences can be identified. For example, amino acid residues that are conserved between organisms are ones that should not be substituted (such as amino acids M50, T70, Y81, Q100 and D150 based on the numbering for *A. niger*), while amino acid residues that are not conserved between organisms are ones likely to tolerate substitution (such as amino acids R8, L160, S395 and N405 based on the numbering for *A. niger*). Similarly, amino acid positions in FIGS. 10A and 10B indicated with different amino acids at the same position are ones likely to tolerate substitution, while positions with the same amino acid (*) are not.

In some examples, an Alg3 sequence that is to be genetically inactivated encodes or includes one or more conservative amino acid substitutions. A conservative amino acid substitution is a substitution of one amino acid (such as one found in a native sequence) for another amino acid having similar biochemical properties. Typically, conservative substitutions have little to no impact on the activity of a resulting peptide. In one example, an Alg3 sequence (such as any of SEQ ID NOS: 2, 4, 31, 32, 33, 34, 35, or 36) includes one or more amino acid substitutions (for example at 1, 2, 5 or 10 residues). Examples of amino acids which may be substituted for an original amino acid in a protein and which are regarded as conservative substitutions include: Ser for Ala; Lys for Arg; Gln or His for Asn; Glu for Asp; Ser for Cys; Asn for Gln; Asp for Glu; Pro for Gly; Asn or Gln for His; Leu or Val for Ile; Ile or Val for Leu; Arg or Gln for Lys; Leu or Ile for Met; Met, Leu or Tyr for Phe; Thr for Ser; Ser for Thr; Tyr for Trp; Trp or Phe for Tyr; and Ile or Leu for Val. Further information about conservative substitutions can be found in, among other locations in, Ben-Bassat et al., (*J. Bacteriol.* 169:751-7, 1987), O'Regan et al., (*Gene* 77:237-51, 1989), Sahin-Toth et al., (*Protein Sci.* 3:240-7, 1994), Hochuli et al., (*Bio/Technology* 6:1321-5, 1988), WO 00/67796 (Curd et al.) and in standard textbooks of genetics and molecular biology.

The Alg3 gene inactivated in a fungus, in particular examples, includes a sequence that encodes an Alg3 protein having at least at least 80%, at least 85%, at least 90%, at least

95%, at least 97%, at least 98%, or at least 99% sequence identity to an Alg3 sequence, such as SEQ ID NO: 2, 4, 31, 32, 33, 34, 35, or 36, wherein the protein can catalyze the addition of the first dol-p-man derived mannose in an α -1,3 linkage to Man5GlcNAc2-PP-Dol. In a specific example, the Alg3 gene inactivated in a fungus encodes an Alg3 protein shown in SEQ ID NO: 2, 4, 31, 32, 33, 34, 35, or 36.

The Alg3 gene that is to be inactivated in a fungus, in particular examples, includes a sequence having at least at least 80%, at least 85%, at least 90%, at least 95%, at least 97%, at least 98%, or at least 99% sequence identity to an Alg3 nucleic acid sequence, such as SEQ ID NO: 1 or 3 or nucleotides 1186-1306, 1393-1916 and 1989-2582 of SEQ ID NO: 1, and encode an Alg3 protein that can catalyze the addition of the first dol-p-man derived mannose in an α -1,3 linkage to Man5GlcNAc2-PP-Dol. In a specific example, the Alg3 gene inactivated in a fungus is shown in SEQ ID NO: 2 or 4.

One skilled in the art will appreciate that additional Alg3 sequences can be identified using any method such as those described herein. For example, Alg3 nucleic acid molecules that encode an Alg3 protein can be identified and obtained using common molecular cloning or chemical nucleic acid synthesis procedures and techniques, including PCR. In addition, standard nucleic acid sequencing techniques and software programs that translate nucleic acid sequences into amino acid sequences based on the genetic code can be used to determine whether or not a particular nucleic acid has any sequence homology with known Alg3 sequences. Sequence alignment software such as MEGALIGN (DNASTAR, Madison, Wis., 1997) can be used to compare various sequences.

In addition, nucleic acid hybridization techniques can be used to identify and obtain a nucleic acid molecule that encodes an Alg3 protein. Briefly, any known Alg3 nucleic acid molecule, or fragment thereof, can be used as a probe to identify similar nucleic acid molecules by hybridization under conditions of moderate to high stringency. Such similar nucleic acid molecules then can be isolated, sequenced, and analyzed to determine whether the encoded protein is an Alg3 protein.

Any method can be used to introduce an exogenous nucleic acid molecule into a fungal cell, for example to genetically inactivate Alg3. For example, chemical mediated-protoplast transformation, electroporation, *Agrobacterium*-mediated transformation, fusion of protoplasts, and biolistic delivery are common methods for introducing nucleic acid into fungal cells. (See, e.g., Ito et al., *J. Bacteriol.* 153:163-8, 1983; Durrens et al., *Curr. Genet.* 18:7-12, 1990; Sambrook et al., *Molecular cloning: A laboratory manual*, Cold Spring Harbour Laboratory Press, New York, USA, third edition, 2001; and Becker and Guarente, *Methods in Enzymology* 194:182-7, 1991. An exogenous nucleic acid molecule contained within a particular cell of the disclosure can be maintained within that cell in any form. For example, exogenous nucleic acid molecules can be integrated into the genome of the cell or maintained in an episomal state. That is, a cell can be a stable or transient transformant.

Fungi with Increased LaeA Expression or Increased LaeA Expression and Alg3 Deletion

The present disclosure provides isolated fungi having increased LaeA expression, wherein such increased expression or activity (for example in combination with an Alg3 functional inactivation, Alg3 Δ) results in increased citric acid production by the fungi. Such fungi are referred to herein as increased LaeA fungal strains. It is disclosed herein that

increased expression of LaeA (for example in combination with genetic inactivation of Alg3, Alg3 Δ) results in *Aspergillus* fungi that can increase citric acid production as compared to *Aspergillus* having native LaeA levels of expression.

Contemplated herein are isolated fungi having increased LaeA activity/expression, for example in combination with a genetic inactivation of Alg3. Any fungus can be used, such as any genus or variety of *Aspergillus*. In particular examples, the *Aspergillus* fungus is *A. niger*, such as *Aspergillus niger* strain 11414 (American Type Culture Collection (ATCC) No. 11414; NRRL 2270); 1015 (ATCC No. 1015; NRRL 328, CBS 113.46); NRRL 3 (ATCC No. 9029, CBS 120.49, N400); NRRL 3122 (ATCC No. 22343); or 11414KusA-. In other specific examples, the *Aspergillus* is *A. aculeatus*, *A. awamori*, *A. carbonarius*, *A. wentii*, *A. foetidus*, *A. fumigatus*, *A. oryzae*, or *A. terreus*.

Any method for genetic enhancement or up-regulation can be used, as long as the expression of the gene and/or gene product is significantly increased, or the function of the expressed protein is significantly increased. In particular examples, LaeA gene expression is up-regulated by transformation of the fungi with one or more copies of a LaeA coding or genomic sequence (which can be a native or non-native LaeA sequence). In some embodiments, up-regulation refers to an increase in gene or protein expression of at least 20%, at least 40%, at least 50%, at least 100%, at least 150%, at least 200%, at least 300%, or at least 500%, for example relative to the parental fungal strain without the additional copies of an LaeA gene. The term "increased" or "up-regulated" as used herein with respect to a cell and a particular gene or protein activity refers to a higher level of activity than that measured in a comparable cell of the same species. For example, a particular fungi having increased or up-regulated LaeA activity has increased LaeA activity if a comparable fungi having native LaeA activity has less detectable LaeA activity (for example as measured by gene or protein expression).

LaeA sequences are disclosed herein and others are publicly available, for example from GENBANK™ or EMBL. In some examples, the LaeA gene up-regulated encodes a protein having at least 80%, at least 90%, at least 95%, at least 97%, or at least 98% sequence identity to SEQ ID NO: 41 or 59. In some examples, the LaeA gene upregulated comprises at least 80%, at least 90%, at least 95%, at least 97%, or at least 98% sequence identity to SEQ ID NO: 40 (e.g., nt 1-236 and 367-1252 of SEQ ID NO: 41) or 58 (e.g., nt 1-230 and 373-1267 of SEQ ID NO: 58).

Increasing LaeA activity (for example in combination with genetic inactivation of Alg3) results in many phenotypes in the fungi. For example, such recombinant fungi exhibit increased citric acid production in CAP medium. Such increases can be relative to a fungi having a native or wild-type level of LaeA (or LaeA and Alg3) gene or protein expression, such as a parental strain (e.g., *A. niger* strain 11414KusA), grown under the same conditions as the fungi with increased LaeA activity (or increased LaeA activity and decreased Alg3 activity). In some examples, increased citric acid production in CAP medium is an increase of at least 20%, at least 30%, at least 50%, at least 60%, at least 65%, or at least 70%, by such a recombinant fungus as compared to *A. niger* strain 11414KusA. In some examples, recombinant fungi with increased LaeA activity (for example in combination with genetic inactivation of Alg3) have increased sporulation relative to *A. niger* Alg3 Δ on MM, accumulate red color pigments to *A. niger* strain 11414KusA on complete medium, or both.

One skilled in the art will appreciate that additional genes can also be inactivated or upregulated, wherein the additional

genes may or may not provide additional enhancement of citric acid production to the fungus. In one example KusA (e.g., GENBANK™ Accession No. EF061656) is also genetically inactivated.

Also provided by the present disclosure are compositions that include isolated LaeA up-regulated fungi, such as a growth medium. Also provided by the present disclosure are kits that include isolated LaeA up-regulated fungi, such as a kit that includes a medium for culturing, storing, or growing the fungus. Exemplary mediums include solid medium (such as those containing agar, for example CM, PDA or MM) and liquid media (such as a fermentation broth, such as CM, MM, or CAP medium).

A. Methods of Up-Regulating Gene and/or Protein Expression

As used herein, an “activated” or “up-regulated” gene means that expression of the gene or gene product (e.g., protein) has been up-regulated, for example by introduction of additional copies of the appropriate gene or coding sequence into the fungus (or other common molecular biology methods), such that the introduce nucleic acid sequence is expressed, resulting in increased expression or biological activity of the encoded gene product.

Increasing expression of one or more genes (which in some examples is also referred to as up-regulation) can be performed using any conventional method known in the art. In one example, a strain of *Aspergillus* is transformed with a vector which has the effect of up-regulating or otherwise activating a LaeA gene (such as a native or non-native LaeA gene). This can be done by introducing one or more LaeA coding sequences (such as a gene sequence), whose expression is controlled by elements such as promoters and the like which control gene expression, by introducing a nucleic acid sequence which itself (or its encoded protein) can increase LaeA protein activity in the fungus, or by introducing another molecule (such as a protein or antibody) increases LaeA protein activity in the fungus. For example, a LaeA gene can be up-regulated by introduction of a vector that includes one or more LaeA sequences (such as 1, 2, 3, 4, 5, 6, 7, 8, 9 or 10 LaeA sequences or copies of such sequences) into the desired fungus. In some examples, such LaeA sequences are from different fungal species, can be multiple copies from a single species, or combinations thereof, such as LaeA sequences from at least 2, 3, 4, 5, 6, 7, 8, 9, or 10 different fungal species. In some examples, the LaeA sequence(s) introduced into the fungus is optimized for codon usage. Thus, the disclosure in some examples provides transformed fungi that include at least one exogenous nucleic acid molecule which includes a LaeA gene or coding sequence (such as a nucleic acid sequence encoding SEQ ID NOs: 41 or 59), for example in combination with Alg3Δ. In one example, such transformed cells produce more citric acid, for example relative to a comparable fungus with a native LaeA sequence (or a native LaeA sequence combined with a native Alg3 sequence).

In another technique, the cre-lox system is used for site specific recombination of DNA (for example see Steiger et al., *Appl. Environ. Microbiol.* 77(1):114, 2011). The system includes 34 base pair lox sequences that are recognized by the bacterial cre recombinase gene. If the lox sites are present in the DNA in an appropriate orientation, DNA flanked by the lox sites will be excised by the cre recombinase, resulting in the deletion of all sequences except for one remaining copy of the lox sequence. Using standard recombination techniques, the targeted gene of interest (e.g., LaeA) can be deleted in the *Aspergillus* genome and replaced with one or more copies of a non-native LaeA sequence (for example in *A. niger*, replacing one or both *A. niger* LaeA sequences with one or more, or

combination of, LaeA sequences from *A. nidulans*, *A. flavus*, *fusarium oxysporum*, *penicillium chrysogenum*, which have high secondary metabolite production) flanked by the lox sites. Transient expression (by electroporation of a suicide plasmid containing the cre gene under control of a promoter that functions in *Aspergillus*) of the cre recombinase should result in efficient elimination of the lox flanked marker. This process will produce a fungus containing the desired insertion mutation and one copy of the lox sequence.

In one example, one or more LaeA genes are introduced into fugal cells by chemical mediated proteoplast transformation in combination of yeast-gap repairing method for transgene expression construction.

In one example, a transgene is generated and expressed in the desired fungal cell, such as an Alg3Δ cell, to increase LaeA expression. For example, such a transgene can include a LaeA genomic or cDNA sequence (such as one having at least 80%, at least 85%, at least 90%, at least 95%, at least 97%, at least 98%, or at least 99% sequence identity to any known LaeA sequence, such as SEQ ID NO: 40 or 58), for example operably linked to a promoter, such as a glyceraldehyde 3-phosphate dehydrogenase (gpdA) promoter or other promoter, such as one that has high activity in CAP culture medium, for example a polyubiquitin promoter, Arsa-7, and A-37 from *A. niger*. In one example, the promoter has at least 80%, at least 85%, at least 90%, at least 95%, at least 97%, at least 98%, or at least 99% sequence identity to SEQ ID NO: 37. In one example, the promoter comprises or consists of the sequence shown in SEQ ID NO: 37. In some examples, the transgene further includes pyrG upstream and downstream sequences (for example that are at the 5'- and 3'-end, respectively, of the transgene). The pyrG gene in *A. niger* is mutated and has lost its original functions. Thus, other non-essential gene loci can be used as long as it is not influenced by the native neighbor genes. In one example, the pyrG upstream and downstream sequences have at least 80%, at least 85%, at least 90%, at least 95%, at least 97%, at least 98%, or at least 99% sequence identity to SEQ ID NO: 45 and 48, respectively. In one example, the pyrG upstream and downstream sequences comprise or consist of the sequence shown in SEQ ID NO: 45 or 48, respectively. In some examples, the transgene further includes a trpC transcriptional terminator sequence of *A. nidulans*, for example downstream of the LaeA sequence. As an alternative to trpC, other transcriptional terminators can be used, such as promoters which include a transcriptional terminators (e.g., Arsa7, Arsa-37, polyubiquitin (ubi4)). In one example, the trpC transcriptional terminator has at least 80%, at least 85%, at least 90%, at least 95%, at least 97%, at least 98%, or at least 99% sequence identity to SEQ ID NO: 46. In one example, the trpC transcriptional terminator comprises or consists of the sequence shown in SEQ ID NO: 46. In some examples, the transgene further includes a ptrA sequence, for example downstream of the trpC transcriptional terminator sequence. As an alternative to ptrA, the bleomycin gene or bar gene can be used. In one example, the ptrA sequence has at least 80%, at least 85%, at least 90%, at least 95%, at least 97%, at least 98%, or at least 99% sequence identity to SEQ ID NO: 47. In one example, the ptrA sequence comprises or consists of the sequence shown in SEQ ID NO: 47. In one example, the transgene comprises a sequence having at least 80%, at least 85%, at least 90%, at least 95%, at least 97%, at least 98%, or at least 99% sequence identity to SEQ ID NO: 44 or 57. In one example, the transgene comprises or consists of the sequence shown in SEQ ID NO: 44 or 57.

Thus, for example, a fungal cell can be engineered to have increased copies of LaeA using common recombinant technology methods.

B. Measuring Gene Activation or Up-Regulation

A fungus having an activated or up-regulated LaeA gene can be identified using any method known in the art. For example, PCR and nucleic acid hybridization techniques, such as Northern, RT-PCR, and Southern analysis, can be used to confirm that a fungus has an up-regulated LaeA gene, such as an increase in the LaeA copy number. Immunohistochemical and biochemical techniques can also be used to determine if a cell expresses LaeA by detecting the expression of the LaeA peptide encoded by LaeA. For example, an antibody having specificity for LaeA can be used to determine whether or not a particular fungus has increased LaeA protein expression. Further, biochemical techniques can be used to determine if a cell has increased LaeA expression by detecting a product produced as a result of the expression of the peptide. For example, measurement of secondary metabolites can indicate that a fungal cell contains an up-regulated LaeA gene. In addition, measurements of citric acid production can be measured using the methods described herein.

C. Measuring Citric Acid Production

Methods of determining whether a genetic up-regulation of LaeA (alone or in combination with inactivation of Alg3) in *Aspergillus* increases citric acid production, for example relative to the same strain with a native LaeA sequence, Alg3 sequence, or both (such as a parental strain), are routine in the art. Although particular examples are disclosed herein (see above and in the examples below), the methods are not limiting.

D. LaeA Sequences

LaeA protein and nucleic acid sequences are publicly available and specific examples are provided herein. In addition, LaeA sequences can be identified using routine molecular biology methods.

Examples of LaeA nucleic acid sequences shown in SEQ ID NOS: 40 and 58. However, the disclosure also encompasses variants of SEQ ID NOS: 40 and 58 (such as the coding regions nt 1-236 and 367-1252 of SEQ ID NO: 41 and nt 1-230 and 373-1267 of SEQ ID NO: 58) which retain the ability to encode a LaeA protein. One skilled in the art will understand that variant LaeA nucleic acid sequences can be used to increase expression of LaeA. Variant sequences may contain a single insertion, a single deletion, a single substitution, multiple insertions, multiple deletions, multiple substitutions, or any combination thereof (e.g., single deletion together with multiple insertions). In addition, the degeneracy of the code permits multiple nucleic acid sequences to encode the same protein. Thus, in one example, a LaeA sequence having at least 80%, at least 85%, at least 90%, at least 95%, at least 97%, at least 98%, or at least 99% sequence identity to any known LaeA sequence, such as SEQ ID NO: 40 or 58 (such as the coding regions nt 1-236 and 367-1252 of SEQ ID NO: 41 and nt 1-230 and 373-1267 of SEQ ID NO: 58) can be expressed in a fungal cell to increase LaeA expression in the fungal cell.

For example, FIG. 12 shows an alignment of LaeA protein sequences from *A. nidulans* (aa 14-372 of SEQ ID NO: 41) and *A. niger* (aa 14-370 of SEQ ID NO: 59), which permits one to identify amino acids that can tolerate substitution (e.g., those that are not conserved between species) and those that may not (e.g., those that are conserved between species). Based on these alignments, variants of LaeA sequences can be identified. For example, amino acid residues that are conserved between organisms are ones that should not be substituted (such as amino acids S16, M42, and P52 based on the

numbering for *A. nidulans*), while amino acid residues that are not conserved between organisms are ones likely to tolerate substitution (such as amino acids T15, S44, and N325 based on the numbering for *A. nidulans*). Similarly, amino acid positions in FIG. 12 indicated with a space are ones likely to tolerate substitution, while positions with the same amino acid are not.

Such protein molecules can share at least 80%, at least 85%, at least 90%, at least 95%, at least 97%, at least 98%, or at least 99% sequence identity to any known LaeA nucleic acid sequence, such as SEQ ID NOS: 41 and 59, and such variants can be used to increase LaeA activity in a fungal cell. One skilled in the art will understand that variant LaeA enzyme sequences can be used to increase LaeA activity in a fungal cell. Variant sequences may contain a single insertion, a single deletion, a single substitution, multiple insertions, multiple deletions, multiple substitutions, or any combination thereof (e.g., single deletion together with multiple insertions).

In some examples, a LaeA sequence whose expression is to be up-regulated encodes or includes one or more conservative amino acid substitutions. A conservative amino acid substitution is a substitution of one amino acid (such as one found in a native sequence) for another amino acid having similar biochemical properties. Typically, conservative substitutions have little to no impact on the activity of a resulting peptide. In one example, a LaeA sequence (such as any of SEQ ID NOS: 41 and 59) includes one or more amino acid substitutions (for example at 1, 2, 5 or 10 residues). Examples of amino acids which may be substituted for an original amino acid in a protein and which are regarded as conservative substitutions include those discussed above for Alg3.

The LaeA gene up-regulated in a fungus, in particular examples, includes a sequence that encodes a LaeA protein having at least at least 80%, at least 85%, at least 90%, at least 95%, at least 97%, at least 98%, or at least 99% sequence identity to an LaeA sequence, such as SEQ ID NO: 41 or 59, wherein the protein can regulate secondary metabolite production in *Aspergillus*. In a specific example, the LaeA gene up-regulated in a fungus encodes a LaeA protein shown in SEQ ID NO: 41 or 59.

The LaeA gene up-regulated in a fungus, in particular examples, includes a sequence having at least at least 80%, at least 85%, at least 90%, at least 95%, at least 97%, at least 98%, or at least 99% sequence identity to a LaeA nucleic acid sequence, such as SEQ ID NO: 40 or 58 (or to the coding regions nt 1-236 and 367-1252 of SEQ ID NO: 41 or nt 1-230 and 373-1267 of SEQ ID NO: 58), and encodes a LaeA protein which can regulate secondary metabolite production in *Aspergillus*. In a specific example, the LaeA gene upregulated in a fungus is shown in SEQ ID NO: 40 or 58 (or includes the coding regions nt 1-236 and 367-1252 of SEQ ID NO: 41 or nt 1-230 and 373-1267 of SEQ ID NO: 58).

One skilled in the art will appreciate that additional LaeA sequences can be identified and obtained using any method such as those described herein. For example, LaeA nucleic acid molecules that encode a LaeA protein can be identified and obtained using common molecular cloning or chemical nucleic acid synthesis procedures and techniques, including PCR. In addition, standard nucleic acid sequencing techniques and software programs that translate nucleic acid sequences into amino acid sequences based on the genetic code can be used to determine whether or not a particular nucleic acid has any sequence homology with known LaeA sequences. Sequence alignment software such as MEGA-LIGN (DNASTAR, Madison, Wis., 1997) can be used to compare various sequences.

In addition, nucleic acid hybridization techniques can be used to identify and obtain a nucleic acid molecule that encodes a LaeA protein. Briefly, any known LaeA nucleic acid molecule, or fragment thereof, can be used as a probe to identify similar nucleic acid molecules by hybridization under conditions of moderate to high stringency. Such similar nucleic acid molecules then can be isolated, sequenced, and analyzed to determine whether the encoded protein is a LaeA protein. The gene specific oligonucleotide pair can also be designed, synthesized and used for real-time RT-PCR to quantify the LaeA gene transcription level.

Any method can be used to introduce an exogenous nucleic acid molecule into a fungal cell, for example to genetically enhance LaeA expression. For example, chemical mediated-protoplast transformation, electroporation, *Agrobacterium*-mediated transformation, fusion of protoplasts, and biolistic delivery are common methods for introducing nucleic acid into fungal cells. (See, e.g., Ito et al., *J. Bacteriol.* 153:163-8, 1983; Durrens et al., *Curr. Genet.* 18:7-12, 1990; Sambrook et al., *Molecular cloning: A laboratory manual*, Cold Spring Harbour Laboratory Press, New York, USA, second edition, 1989; and Becker and Guarente, *Methods in Enzymology* 194:182-7, 1991). An exogenous nucleic acid molecule contained within a particular cell of the disclosure can be maintained within that cell in any form. For example, exogenous nucleic acid molecules can be integrated into the genome of the cell or maintained in an episomal state. That is, a cell can be a stable or transient transformant.

E. LaeA Deletion and Complementation

Disclosed herein is a LaeA deletion cassette for generating a fungus, such as an *Aspergillus* species fungus, having a deletion in the LaeA gene. A schematic of the LaeA deletion cassette is shown in FIG. 17A. To generate the deletion cassette, DNA fragments comprising the 5' and 3' ends of the *A. niger* LaeA gene (SEQ ID NOs: 61 and 62, respectively) were isolated from *A. niger* genomic DNA and the hph expression cassette (SEQ ID NO: 63) (hph) was isolated from pCB 1003 plasmid vector DNA by PCR. The PCR DNA fragments were assembled in the pBSK backbone vector to produce the LaeA deletion cassette (SEQ ID NO: 65).

Provided herein is a nucleic acid molecule comprising a nucleotide sequence at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98% or at least 99% identical to SEQ ID NO: 65. In some embodiments, the nucleic acid molecule comprises or consists of the nucleotide sequence of SEQ ID NO: 65. Further provided is an isolated fungus transformed with a nucleic acid molecule comprising a nucleotide sequence at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98% or at least 99% identical to SEQ ID NO: 65. In some embodiments, the isolated fungus is transformed with a nucleic acid molecule comprising or consisting of the nucleotide sequence of SEQ ID NO: 65. In some embodiments, the fungus is a species of *Aspergillus*, such as *A. niger*.

Further provided herein are isolated fungi (such as filamentous fungi) having a gene inactivation (also referred to as a gene deletion) of a LaeA gene. Any strain of fungi can be used, such as a filamentous fungi, for example *A. niger* or particular strains thereof (for example *A. niger* strain 11414 or 11414KusA). In particular examples, the LaeA gene is genetically inactivated by complete or partial deletion mutation or by insertional mutation. In some examples genetic inactivation need not be 100% genetic inactivation. In some embodiments, genetic inactivation refers to at least 50%, at least 60%, at least 70%, at least 80%, at least 90%, or at least 95% gene or protein inactivation. The term "reduced" or "decreased" as used herein with respect to a cell and a particular gene or protein activity refers to a lower level of

activity than that measured in a comparable cell of the same species. For example, a particular fungi lacking LaeA activity has reduced LaeA activity if a comparable fungi not having an LaeA genetic inactivation has detectable LaeA activity. In some embodiments, the isolated fungi having a gene inactivation are generated using the LaeA deletion construct set forth herein as SEQ ID NO: 65.

Also provided herein is a LaeA complementation construct for complementing expression of LaeA in fungi having a deleted LaeA gene. A schematic of the LaeA complementation construct is shown in FIG. 17B. To generate the complementation construct, the LaeA gene containing both its promoter and transcriptional terminator was isolated by PCR from *A. niger* genomic DNA. The PCR fragment was cloned into the plasmid vector pRSB426-LaeA. The new plasmid DNA vector contained the upstream region of the pyrG gene of *A. niger*, the entire region of *A. niger* LaeA gene, the transcriptional terminator of the trpC gene from *A. nidulans*, the pyrithiamine resistance (ptrA) gene from *A. oryzae*, and the downstream region of the pyrG gene of *A. niger* (SEQ ID NO: 66).

Provided herein is a nucleic acid molecule comprising a nucleotide sequence at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98% or at least 99% identical to SEQ ID NO: 66. In some embodiments, the nucleic acid molecule comprises or consists of the nucleotide sequence of SEQ ID NO: 66. Further provided is an isolated fungus transformed with a nucleic acid molecule comprising a nucleotide sequence at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98% or at least 99% identical to SEQ ID NO: 66. In some embodiments, the isolated fungus is transformed with a nucleic acid molecule comprising or consisting of the nucleotide sequence of SEQ ID NO: 66. In some embodiments, the fungus is a species of *Aspergillus*, such as *A. niger*.

Further provided herein are isolated fungi (such as filamentous fungi) having a gene inactivation (also referred to as a gene deletion) of a LaeA gene and further transformed by a nucleic acid construct comprising a nucleotide sequence at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98% or at least 99% identical to SEQ ID NO: 66. In some embodiments, the isolated fungus is transformed with a nucleic acid molecule comprising or consisting of the nucleotide sequence of SEQ ID NO: 66. In some embodiments, the fungus is a species of *Aspergillus*, such as *A. niger*. Any strain of fungi can be used, such as a filamentous fungi, for example *A. niger* or particular strains thereof (for example *A. niger* strain 11414 or 11414KusA).

Production of Citric Acid Using Alg3Δ Mutants, Fungi with Increased LaeA Expression, or Both

The fungi provided herein, namely Alg3Δ fungi, up-regulated LaeA fungi, and fungi with both Alg3Δ and up-regulated LaeA, can be used to produce citric acid, as well as derivatives thereof such as hydroxycitric acid (for example for medical applications). Such fungi can be from any species, such as *Aspergillus* or *Rhizopus* cells. For example, the disclosure provides methods of making citric acid, which can include culturing Alg3Δ fungi, up-regulated LaeA fungi, or fungi with both Alg3Δ and up-regulated LaeA, under conditions that permit the fungus to make citric acid, for example in CAP medium.

Citric acid (2-hydroxy-propane-1,2,3-tricarboxylic acid) combines a pleasant taste with low toxicity and palatability and is a ubiquitous food additive. It is also able to complex heavy metal ions, like iron and copper, and is therefore

applied in the stabilization of oils and fats or ascorbinic acid during metal ion-catalyzed oxidation reactions. Consequently, it is today one of the bulk products produced by fermentation, most of which occurs with the fungus *Aspergillus niger*, although a small portion is also produced by fermentation with yeast, such as *Candida oleophila* and *Candida lipolytica*.

Citric acid production generally requires a unique combination of several unusual nutrient conditions (e.g., excessive concentrations of carbon source, 1-1%, and dissolved oxygen, or suboptimal concentrations of certain trace metals and phosphate), which synergistically influence the yield of citric acid. Table 1 below shows the environmental parameters that influence citric acid accumulation.

TABLE 1

Parameters that influence citric acid accumulation by <i>A. niger</i>	
Parameter	Requirement for citric acid accumulation
Carbon source concentration	Higher than 50 g/l
Carbon source type	Enable rapid catabolism
Nitrogen source	Consumption leads to some decrease in pH
Phosphate concentration	Suboptimal
Aeration	In excess
Trace metal ions	Limiting, especially Mn ²⁺
pH	Below pH 3

Methods of making citric acid, which can include culturing Alg3Δ fungi, up-regulated LaeA fungi, or fungi with both Alg3Δ and up-regulated LaeA, under conditions that permit the fungus to make citric acid, are provided. In general, the culture media and/or culture conditions can be such that the fungi grow to an adequate density and produce citric acid efficiently. In one example the Alg3Δ fungi, up-regulated LaeA fungi, or fungi with both Alg3Δ and up-regulated LaeA, are cultured or grown in a liquid medium that includes sucrose and/or glucose as the carbon source, for example at a concentration of at least 50 g/liter, such as at least 100 g/l, or at least 140 g/l. Thus, a fungus within the scope of the disclosure in some examples can utilize a variety of carbon sources. In one example the Alg3Δ fungi, up-regulated LaeA fungi, or fungi with both Alg3Δ and up-regulated LaeA, are cultured or grown in a liquid medium that includes a very small amount of manganese, such as less than 100 parts per billion (ppb), less than 50 ppb, less than 20 ppb, less than 15 ppb, for example 5 ppb to 15 ppb or 10 ppb to 15 ppb, such as 5, 10, 13, 15 or 20 ppb. In one example the Alg3Δ fungi, up-regulated LaeA fungi, or fungi with both Alg3Δ and up-regulated LaeA, are cultured or grown in a liquid medium having an initial pH of less than 3, such as less than 2.5, for example about pH 1.8 to 3, 1.8 to 2.5, 1.8 to 2.2, 1.9 to 2.1, for example pH 1.8, 1.9, 2, 2.1, 2.2, 2.3, 2.4, 2.5, 2.6, 2.7, 2.8 or 2.9. In some examples the Alg3Δ fungi, up-regulated LaeA fungi, or fungi with both Alg3Δ and up-regulated LaeA, are cultured or grown in a liquid medium at about 25 to 35° C. (such as 28 to 32° C., or 30° C.) with rotation of 180 to 300 rpm.

In a specific example, the Alg3Δ fungi, up-regulated LaeA fungi, or fungi with both Alg3Δ and up-regulated LaeA, are grown in citric acid production (CAP) medium. In a specific example, the CAP medium includes 140 g of glucose/liter, 3.1 g of NH₄NO₃/liter, 0.15 g of KH₂PO₄/liter, 0.15 g of NaCl/liter, 2.2 g of MgSO₄·7H₂O/liter, 6.6 mg of ZnSO₄·7H₂O/liter, and 0.1 mg of FeCl₃/liter adjusted to about pH 2 with 4 M H₂SO₄. Cations can be removed from the glucose solution by ion exchange on Dowex 50W-X8, 100/200-mesh, H cation exchange resin (Fisher Scientific, Pittsburgh, Pa.) prior to adding the other nutrient components. The manganese concentration in the medium can be adjusted by the addition of appropriate volumes of a stock solution of MnCl₂·4H₂O (10

mM). In one example, the manganese concentration is less than 50 ppb, such as less than 20 ppb, for example 5 to 15 ppb, such as 10 ppb.

Methods of culturing *Aspergillus* to enable citric acid production are well known in the art. In one example, the fungi are grown in culture containers (such as baffled flasks, and in some examples are silanized (5% solution of dichlorodimethylsilane in heptane (Sigma, St. Louis, Mo.)). The Alg3Δ fungi, up-regulated LaeA fungi, or fungi with both Alg3Δ and up-regulated LaeA, provided herein can be grown in CAP media containing low amounts of Mn²⁺ (e.g., 10 ppb) at 30° C. with rotation (e.g., 200 to 250 rpm) for at least 3 days (e.g., 3 to 7 days). Each culture container is inoculated with spores (such as at least 10⁶ spores/ml) and incubated for at least 12 or at least 15 hours at 30° C. and 200 to 250 rpm to obtain properly pelleted morphology.

In one example, the Alg3Δ fungi, up-regulated LaeA fungi, or fungi with both Alg3Δ and up-regulated LaeA, produce more citric acid than a corresponding fungus with wild-type Alga. In specific examples, the Alg3Δ fungi, up-regulated LaeA fungi, or fungi with both Alg3Δ and up-regulated LaeA, produce at least 25 g/l of citric acid (for example at least 30 g/l, at least 32 g/l, at least 35 g/l, at least 40 g/l, at least 42 g/l, at least 45 g/l, at least 50 g/l, at least 52 g/l or at least 55 g/l), for example after at least 4 days (such as at least 5 days, at least 6 days, at least 7 days, at least 8 days, or at least 10 days, such as after 4 to 6 days, 8 to 10 days, or 4 to 5 days) when grown in CAP medium at 30° C. with 200 rpm shaking.

In some examples, the method further includes isolating the citric acid made by the Alg3Δ fungi, up-regulated LaeA fungi, or fungi with both Alg3Δ and up-regulated LaeA. Once produced, any method can be used to isolate the citric acid. For example, common separation techniques can be used to remove the fungal biomass from the culture medium, and common isolation procedures (e.g., filtration, distillation, precipitation, electrodialysis, and ion-exchange procedures) can be used to obtain the citric acid from the broth (such as a fungi-free broth). In addition, the citric acid can be isolated from the culture medium after the citric acid production phase has been terminated.

EXAMPLE 1

Materials and Methods

This example describes methods used in the experiments described in Examples 2-5 below.

Strains and Media. The *Escherichia coli* strains Top10 and *Saccharomyces cerevisiae* strain YVH10 were used as hosts for routine cloning and gap repair experiments. *A. niger* strain ATCC 11414 (American Type Culture Collection, Rockville, Md.), was grown on potato dextrose agar plates (PDA) and complete medium (CM) agar plates at 30° C. for culture maintenance and spore preparation, respectively. The mutant strain *Aspergillus niger* 11414KusA was generated by the deletion of kusA in *A. niger* strain 11414 by the replacement with *A. fumigatus* pyrG gene, which encodes the ortholog of the ku70 protein that involves in the non-homologous end joining pathway of DNA repair for the integration of a DNA fragment into the genome in other eukaryotes, and was confirmed by Southern blotting analysis. The 11414KusA strain with high rate of homologous replacement was mainly used as a parent strain. The cultures on PDA or complete medium (CM) agar plates were incubated for four days at 30° C. and the spores were harvested by washing with sterile 0.8% Tween 80 (polyoxyethylenesorbitan monooleate). The CM medium contains 20 g of D-glucose/liter, 5 g yeast extract/liter, 2 g trypticase peptone/liter, 1 g casamino acids/liter, 6 g NaNO₃/liter, 0.52 g KCl/liter, 0.52 g MgSO₄·7H₂O/liter, 1.52

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g KH_2PO_4 /liter, 36.7 mg $\text{ZnSO}_4 \cdot 7\text{H}_2\text{O}$ /liter, 18.3 mg H_3BO_3 /liter, 8.3 mg $\text{MnCl}_2 \cdot 4\text{H}_2\text{O}$ /liter, 8.3 mg $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$, 2.8 mg $\text{CoCl}_2 \cdot 6\text{H}_2\text{O}$ /liter, 2.7 mg $\text{CuSO}_4 \cdot 5\text{H}_2\text{O}$ /liter, 2.5 mg $\text{Na}_2\text{MoO}_4 \cdot 2\text{H}_2\text{O}$ /liter, 83.3 mg Na_2EDTA /liter, 1 mg biotin/liter, 1 mg pyridoxin/liter, 1 mg thiamine/liter, 1 mg riboflavin/liter, 1 mg p-aminobenzoic acid/liter and 1 mg nicotinic acid/liter. The PDA medium contains 4 g/liter potato starch and 20 g/liter dextrose. Conidia were enumerated with a hemacytometer. Aliquots of the resulting spore suspension (1×10^9 spores/ml) were used to inoculate baffled-flask liquid cultures. The citric acid production (CAP) medium contained 140 g/l of glucose, 3.1 g/l NH_4NO_3 , 0.15 g/l KH_2PO_4 , 0.15 g/l NaCl, 2.2 g/l $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$, 6.6 mg/l $\text{ZnSO}_4 \cdot 7\text{H}_2\text{O}$, and 0.1 mg/l FeCl_3 adjusted to pH 2.1 with 4 M H_2SO_4 . Cations were removed from the glucose solution by ion-exchange on Dowex 50W-X8, 100-200 mesh, H cation exchange resin (Fisher Scientific, Pittsburgh, Pa.) prior to adding the other nutrient components.

Culture Methods. Glass baffled-flasks of 250 ml or 1000 ml were sterilized by rinsing in a 5% solution of dichlorodimethylsilane in heptane (Sigma, St. Louis, Mo.) to minimize leaching of metals. For citric acid production tests, 1×10^6 spores/ml of parent or mutant strains were grown in 80 ml CAP media containing 10 ppb Mn^{2+} in 250 ml baffled flasks or 220 ml CAP media in 1000 ml baffled flasks at 30° C. and 200 rpm. Samples for citric acid analysis were taken at intervals. The biomass of transgenic clones and parent strain were prepared from 2 ml CM station cultures with proper antibiotics and grown in 16x125 mm glass culture-tubes at 30° C. without shaking. The biomass formed on the surface of the culture medium was collected, frozen immediately in liquid nitrogen and dried in the lyophilizer.

Dried Biomass Measurement. After proper cultivation, the cell mass from citric acid production culture was collected by centrifugation at room temperature and 4500xg for 5 min in Sorvall floor centrifuge with swinging-bucket rotor. The cell mass was then transferred onto the Whatman Grade No 1 filter paper or left in centrifuge tubes for freeze-drying. The biomass was then dried in high temperature oven at 80° C. or freeze-dried in the lyophilizer. Prior to being used, the centrifuge tube or Whatman filter paper was weighted and re-weighted after the biomass was completely dried.

Total Genomic DNA Isolation for PCR and Southern Blotting Analysis. Total genomic DNA was isolated from *A. niger* according to the SDS extraction method described previously by Dellaporta et al. (*Plant Molecular Biology Reporter* 1(4): 19-21, 1983) with some modifications. Briefly, fungal biomass from 2 ml station cultures was looped and transferred into a 1.5 ml microcentrifuge tube. A needle size hole on the cap was punched with 18 gauge needle. The tube was immediately frozen in liquid N₂ for 5 minutes and biomass in the tube was dried in a VirTis benchtop manifold freeze dryer (SP Scientific, Gardiner, N.Y.) overnight. The dried biomass and two 3.5 mm diameter glass beads were transferred into the 2 ml polypropylene microvial, where biomass was pulverized into fine power with Mini-Beadbeater-8 (Bio Spec Products Inc., Bartlesville, Okla.) for one minute. Then, 500 µl of 60°

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C. extraction buffer and 80 µl of 15% SDS were added into the microcentrifuge tube and incubated at 65° C. for at least 30 minutes with occasionally swirling to mix. Two-hundred microliters of 5M potassium acetate was added, mixed and incubated on ice for 30 minutes. The supernatant was collected by centrifugation at 12,000 g for 10 minutes at 4° C. and transferred into the new microcentrifuge tube. The total nucleic acids were precipitated with 780 µl of 2-propanol for 30 minutes at -20° C. and centrifuged at 12,000 g for 10 minutes. The nucleic acids were re-suspended in 200 µl 50TE buffer containing 2 µl RNase (10 µg/µl stock solution) and incubated in Eppendorf thermomixer at 50° C. and 500 rpm for 30 minutes. The proteins and cell debris was removed by being added and well mixed 20 µl 3M sodium acetate and equal volume of phenol:chloroform and centrifuged at 15,000 g for 5 minutes. The supernatant was transferred to new DNase-free microcentrifuge tube containing 220 µl of 2-propanol, mixed well and incubated at room temperature for 5 minutes. The genomic DNA was pelleted by centrifugation at 15,000 g for 10 minutes and washed with 500 µl of 70% ethanol. The genomic DNA was re-suspended in 80 µl 10 mM TrisHCl (pH8.0) buffer and determined with Qubit fluorometer (Invitrogen, Carlsbad, Calif.). One microgram of total genomic DNA was digested with restriction endonuclease BamH and SacII. The genomic DNA fragments were separated in 1% agarose gel electrophoretically and transferred onto the zeta-probe membrane (BioRad) with alkaline capillary transfer method. A 3.8 kb genomic DNA fragment containing the Alg3 sequence was used for preparation of the biotin-labeled probe. The genomic DNA in Zeta-probe membrane was hybridized with the biotin-labeled probe overnight in 60° C. hybridization oven. The genomic DNA on hybridized membrane was visualized with North2South chemiluminescent detection kit (Pierce Protein Research Products, Rockford, Ill.).

Spore Production and Germination. The spore production on the PDA or CM agar plates described above was excised with plastic closures of culture tubes in 27 mm diameter and transferred into the 50 ml centrifuge tubes containing 25 ml 0.8% tween 80. The spores were released from the agar surface by scraping with plastic loops and vortexed with vortex mixer at top speed. The spores were diluted properly and enumerated with a hemacytometer. The spore production in a unit area (cm^2) was determined. For spore germination, 1×10^5 spores per well were added into each well of 24 well Schwarz sensoplate and incubated in the microscopic incubator with temperature control at 30° C. The spore germination was automatically imaged hourly for 24 hrs through the Olympus inverted system microscope (Olympus America Inc., Center Valley, Pa., USA). The spore germination was visualized with Adobe Photoshop CSS (San Jose, Calif.) and counted manually.

Citric acid measurements. Citric acid concentrations were determined with an end-point spectrophotometric enzyme assay as described in the instruction from the manufacturer (R-Biopharm AG/Roche, Darmstadt, Germany) with a proper dilution.

Table 2 shows oligonucleotides used in the methods.

TABLE 2

Oligonucleotides		
Name	Sequence	Product size (kb)
Alg3A construction		
Alg3-ForScr	CGGTTTCCTTCAGTTCCAGT (SEQ ID NO: 5)	

TABLE 2-continued

Oligonucleotides		
Name	Sequence	Product size (kb)
Alg3-1	GTAACGCCAGGGTTTTCCAGTCACGACGTCATAACTTCTCTCCCTCC (SEQ ID NO: 6)	1.06
Alg3-2	ATCCACTTAACGTTACTGAAATCTCCAACTTTCATGGACACACAGACC (SEQ ID NO: 7)	
Hph-F	GGTCTGTGTGTCCATGAAGTTGGAGATTTAGTAACGTTAAGTGGAT (SEQ ID NO: 8)	1.49
Hph-R	GCTACTACTGATCCCTCTGCGTCGGAGACAGAAGATGATATTGAAGGAG (SEQ ID NO: 9)	1.03
Alg3-3	CTCCTTCAATATCATCTTCTGTCTCCGACGCAGAGGGATCAGTAGTAGC (SEQ ID NO: 10)	
Alg3-4	GCGGATAACAATTTACACAGGAAACAGCCGTGAGAGGTTTGTAGTACG (SEQ ID NO: 11)	
Alg3-RevScr	AAGCTGAGAGCGACATCTTCA (SEQ ID NO: 12)	
hyg-RevScr	GTAATTCTACACAGCCATCGGTCCA (SEQ ID NO: 13)	
hyg-ForScr	GTAATTCTACACAGCCATCGGTCCA (SEQ ID NO: 14)	
Alg3A Alg3 construction		
pryGScr	TCTGCTGTCTTGCATGAGGTCCTT (SEQ ID NO: 15)	
pyrGScr	Agcgtaggacaaggctcgtctctgt (SEQ ID NO: 16)	2.34
5-pyrG5F	GTAACGCCAGGGTTTTCCAGTCACGACGttaaacATGCATCATCTCCCGCTTTGT (SEQ ID NO: 17)	1.69
5-pyrG3R	agaaagagtcaccggtcacGacatcgccaatcacctcaatcac (SEQ ID NO: 18)	1.47
ble5F	gtgattgaggtgattggcgatgtCgtgaccggtgactctttct (SEQ ID NO: 19)	1.23
Ble3R	TCCAACCTTGTAGCAACCAAAGCTTCGAGCGTCCCAAAACCT (SEQ ID NO: 20)	
Alg3-5F1	<u>AGGTTTGGGACGCTCGAAGCTTTGGTTGCTACAAGGTGGA</u> (SEQ ID NO: 21)	
Alg3-3R1	TCAAGTAGAGCACAGCAATAGTATCTGA (SEQ ID NO: 22)	
Alg3-5F2	TCAGATACTATTGCTGTGCTCTACTTGA (SEQ ID NO: 23)	
Alg3-3R2	ttgatccttgtgccacaccaTCTACGTGGTCATCGATACCA (SEQ ID NO: 24)	
3-pyrG5F	<u>TGGTATCGATGACCAGTAGGA</u> tgggtgtggcacaaggatcaa (SEQ ID NO: 25)	
3-pyrG3R	GCGGATAACAATTTACACAGGAAACAGCgtttaaactgtgccagtcattgtccgaagt (SEQ ID NO: 26)	
Alg3Seq-1	TACAGACGCGGTACGCATGT (SEQ ID NO: 27)	
Alg3seq-1	TGCTATTGTCCACAGATACCGAGA (SEQ ID NO: 28)	
Alg3seq-3	GAGCTAACCAGACAGTTCATGT (SEQ ID NO: 29)	
Alg3seq-4	Tcgtcgtaccgcattgaccc (SEQ ID NO: 30)	

EXAMPLE 2

Genetic Inactivation of Alg3 in *A. niger*

This example describes methods used to genetically clone and then inactivate Alg3 in *A. niger* strain 11414KusA. Based on these teachings, one skilled in the art will appreciate that Alg3 can be similarly inactivated in other strains of *Aspergillus*.

Alg3 has been identified and characterized in *Arabidopsis thaliana*, *Homo sapiens*, *Pichia pastoris*, *Trypanosoma brucei* and *Saccharomyces cerevisiae* (see for example Korner et al., *EMBO J.* 18(23): 6816-6822, 1999; Davidson et al., *Glycobiology* 14(5):399-407, 2004; Manthri et al., *Glycobiology* 18(5): 367-383, 2008; Kajiura et al., *Glycobiology* 20(6):736-

751, 2010). A database search based on the amino acid sequence of *S. cerevisiae* Alg3 identified a putative α -1,3-mannosyltransferase gene in JGI (DOE Joint Genome Institute)-*A. niger* genome database (jgi|Aspni5|42720). The *A. niger* Alg3 gene contains two introns and its 1400 by open reading frame (nt 1186-1306, 1393-1916 and 1989-2582 of SEQ ID NO: 1) encodes a protein consisting of 413 amino acids (SEQ ID NO: 2), which contains one potential N-glycosite at the amino acid position 374. The predicted Alg3 amino acid sequence has 39% sequence identity to the *S. cerevisiae* Alg3.

Alg3 was functionally inactivated in *A. niger* using a gene deletion vector constructed by yeast gap repairing approach. The 5'- and 3'-end of the hygromycin marker (hph) gene was flanked with about 1 kb upstream and downstream fragments

of Alg3 coding region that were isolated by PCR from *A. niger* genomic DNA. The DNA sequence of the upstream and downstream fragments was confirmed by DNA sequencing analysis. The Alg3 in *A. niger* was deleted by homologous replacement with hygromycin marker (hph) gene in the kusA deletion background of *A. niger*, where the kusA gene, encoding the ortholog of the Ku70 protein in other eukaryotes, was deleted for dramatically improved homologous integration efficiency. FIGS. 1A and 1B show the predicted restriction enzyme digestion patterns of genomic DNA of the parent and mutant strains with BamHI and SacII. FIG. 1C shows the Southern blotting analysis of the digested genomic DNA of parent and mutant strains. The results confirm that the Alg3 coding region in *A. niger* was replaced by the hygromycin selection marker gene (hph) in the Alg3Δ strains.

EXAMPLE 3

Effects of Alg3 Deletion on *A. niger* Growth and Development

This example describes methods used to determine the effect genetically inactivating Alg3 in *A. niger*.

It was previously demonstrated that the deletion of Alg3 in different organisms causes underglycosylation, but no obvious phenotype changes were observed at the selected culture condition in those studies (Aebi et al., *Glycobiology* 6(4): 439-444, 1996; Korner et al., *EMBO J.* 18(23): 6816-6822, 1999; Davidson et al., *Glycobiology* 14(5):399-407, 2004; Manthri et al., *Glycobiology* 18(5): 367-383, 2008; Kajiura et al., *Glycobiology* 20(6):736-751, 2010).

The effects of the Alg3 deletion were examined on CM, PDA and MM plates. As exhibited in FIGS. 2A-I, the Alg3Δ strain grew much slower than the parent strain when grown on either CM or PDA medium plate, but there was no significant difference between the Alg3Δ mutants and parent strain when grown on the MM medium plate. When both Alg3Δ strain and parent strain were grown in the liquid culture of CM and PDA, the initiation of spore germination of Alg3Δ strain was pronouncedly delayed (FIG. 3A), but the spore germination rate was not affected by the Alg3 deletion (FIG. 3B). These results demonstrate that deletion of Alg3 has significant effects on *A. niger* growth on nutrient rich media.

The effects of Alg3 deletion on spore production on both CM and PDA plates were also examined. The Alg3 deletion had a substantial reduction of sporulation on CM medium plate, while no obvious difference was exhibited on PDA plate (FIG. 4). The spore production at given area was enumerated with hemocytometer (Table 3). The average spore production of Alg3Δ was 2.64×10^7 spores/cm², about 40% of parent strain (6.44×10^7 spores/cm²) on CM medium plates, while average spore production per a square millimeter was similar between Alg3Δ mutant (7.72×10^7 spores/cm²) and parent strains (7.89×10^7 spores/cm²) on PDA medium plates.

TABLE 3

Average spore production in PDA and CM media plates ($\times 10^7$ sp/cm ²). This was averaged from four cuts and 3 replicate counting.		
Strain	CM pates	PDA
11414-kusA	6.44 ± 1.24	7.72 ± 1.78
Alg3Δ	2.64 ± 0.49	7.89 ± 1.18

EXAMPLE 4

Effects of Alg3 Deletion on Spore Germination, Growth and Citric Acid Production

This example describes methods used to measure spore germination, growth, and citric acid production in the Alg3Δ *A. niger* strain generated in Example 1. Based on these teachings, one skilled in the art will appreciate that spore germination, growth, and citric acid production can be similarly measured in other Alg3Δ strains of *Aspergillus*.

A. niger strain ATCC11414 is a strain developed for industrial production of citric acid. *A. niger* morphology plays a role in citric acid production. The fungal morphology affect overall molecular regulation in response to the endogenous and exogenous factors, which include the regulations of transcription, post-transcription, translation and post-translation. Therefore, the effects of Alg3 deletion on *A. niger* growth on CAP agar plates at different pHs or in CAP liquid culture conditions was determined.

FIG. 5 shows the effects of Alg3 deletion in fungi grown on CAP medium plates at different pH conditions after 28 hrs in culture. When the Alg3Δ strain was grown on the CAP medium plates at pH 1.8, its growth was relatively slower than the parent strain, where its colonies were much smaller than the parent strain. At pH 2.1, the Alg3Δ strain formed a less tight pellet than that of parent strain. Growth of the Alg3Δ strain on the CAP medium at pH 4.5 and pH 7.0 was affected more profoundly than that of parent strain, where the Alg3Δ strain formed thinner layer of biomass on the agar plates than that of parent strain. These results show that the Alg3 deletion reduces the normal growth of *A. niger* at different levels on CAP culture medium plates at different pHs.

The spore germination of the Alg3Δ and parent strains in CAP liquid culture medium was also examined by using automated microscopic imaging, enumerating the germination manually in a same visual unit area, and expressing spore germination as a percentage of total spores at the same visual unit area. FIG. 6 shows the different dynamics of spore germination between the Alg3Δ and parent strains. The spore germination of Alg3Δ strain began as early as 3 hrs after spore inoculation, while the parent strain was not initiated until 6 hrs after spore inoculation. After 8 hrs inoculation, more than 32% spores of Alg3Δ mutant germinated, while only 10% of parent strain spores did (FIG. 6A, top panels; and FIG. 6B). After 15 hrs growth in CAP liquid medium, more than 90% spores of Alg3Δ strain germinated, while only 50% spores of parent strain did (FIG. 6A, bottom panels; and FIG. 6B). After 24 hrs of growth in the liquid culture, about 75% spores of parent strain germinated, while the Alg3Δ strain achieved 94% of germination rate (FIG. 6B). The Alg3 deletion leads to earlier germination and a higher germination rate than parent strain.

The effect of Alg3 deletion on citric acid production was determined in CAP flask cultures. FIG. 7 shows the time course of citric acid production in CAP liquid medium. The yield of citric acid production was similar between the Alg3Δ mutant and parent strain in 2 days culture. After 3 days culture, the average citric acid production by Alg3Δ mutants was 8.8 g/l citric acid, while the parent strain only produced 5.8 g/l. After 4.5 days of culture, the parent strain accumulated 18.8 g/l citric acid and the Alg3Δ mutant produced 33.3 g/l citric acid (more than 70% higher than the parent strain). Thus, the Alg3 deletion substantially improves the citric acid production in *A. niger*.

The effect of Alg3 deletion on citric acid production was also examined by complementation of its original gene into

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the *alg3Δ* mutant. FIG. 8B shows the citric acid production in CAP liquid medium after 10 days of culture. The yield of citric acid production was similar between the *alg3Δ* complemented (*cAlg3Δ*) mutant and parent strain, but much lower than the *alg3Δ* mutant in 10 days culture. After 10 days culture, the average citric acid production by *Alg3Δ* mutants was 46.1 g/l citric acid, while the parent and *calg3Δ* strain only produced 34.8 and 29.4 g/l, respectively.

EXAMPLE 5

pPTRpGPDALaeA Plasmid Vector Construction

This example describes methods used to generate the pPTRpGPDALaeA plasmid vector (FIG. 13). One skilled in the art will appreciate that although *gpdA* and *LaeA* sequences were used from *Aspergillus nidulans*, one skilled in the art will appreciate that variants of these sequences can be used in the fungi and methods provided herein, such as *gpdA* and *LaeA* sequences from other *Aspergillus* species. In one example, a *gpdA* sequence having at least 80%, at least 90%, at least 95%, or at least 98% sequence identity to SEQ ID NO: 37 is used in the fungi and methods provided herein.

The *Aspergillus nidulans* glyceraldehyde 3-phosphate dehydrogenase (*gpdA*) promoter (SEQ ID NO: 37) was isolated from the pAN8-1 plasmid DNA using the primer set *gpdA5F/gpdA3R* (*gpdA5F*: CGCAGATCTC AAGCTGTAAG GATTCGGCA SEQ ID NO: 38; *gpdA3R*: CACCGGGCCC ATCTCAAACA TTGTGATGTC TGCTCAAGCG SEQ ID NO: 39) and the *LaeA* coding sequence of genomic DNA from *A. nidulans* (SEQ ID NO: 40) obtained by PCR using *LaeA5F/LaeA3R* (*LaeA5F*: CGCTTGAGCA GACATCACAA TGTTTGAGAT GGGCCCGGTG; SEQ ID NO: 42; *LaeA3R*: CGCAGATCTG AGGATTATGA GAAGGGAGC; SEQ ID NO: 43).

The DNA fragment of *pGPDALaeA* was filled together by overlap PCR and a HindIII restriction enzyme site was introduced at both 5'- and 3'-end of the DNA fragment. The DNA fragment (SEQ ID NO: 44) and pPTR1 plasmid DNA were cut with Hind III and ligated together by a quick DNA ligation kit at 25° C. for 30 min. The ligated plasmid DNA was transferred into the Top 10 *E. coli* competent cells by lithium acetate mediated transformation. The transformed bacterial colonies were screened for the DNA fragment insertion by PCR with the primers *gpdA5F* (SEQ ID NO: 38) and *LaeA3R* (SEQ ID NO: 43). The plasmid DNA for the selected transformed colonies was prepared for restriction enzyme confirmation and further expression vector construction.

EXAMPLE 6

pRS426-LaeA Vector Construction

This example describes methods used to generate a transgene containing *A. niger* *LaeA* (FIG. 14).

PCR was performed to isolate DNA fragments of *A. niger* *pyrG* upstream region (SEQ ID NO: 45), *trpC* transcriptional terminator of *A. nidulans* (SEQ ID NO: 46), pyrithiamine resistance gene (*ptrA*) of *Aspergillus oryzae* (SEQ ID NO: 47), and *A. niger* *pyrG* downstream region (SEQ ID NO: 48), using primers *pyrGU5F/PTRU3R* (*pyrGU5F*: GTAACGC-CAG GGTTCCTCC GTCACGACGT TTAAACATGC ATCATCTCC CGCTTTGT, SEQ ID NO: 49; *pyrGU3R*: TGCCGAAATC CTTACAGCTT GAAGCTTCAT CGC-CAATCAC CTCAATCAC, SEQ ID NO: 50), *Trp5F/Trp3R* (*Trp5F*: AGCTCCCTTC TCATAATCCT CAAGCTTGGA CCGATGGCTG TGTAAGAGT, SEQ ID NO: 51; *Trp3R*:

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CGTAATCAATTGCCCGTCTGTCAGAGAGCG GATTC-CTCAG TCTCGT; SEQ ID NO: 52), *PTR5F/PTR3R* (*PTR5F*: ACGAGACTGA GGAATCCGCT CTCTGACAGAC GGGCAATTG ATTACG, SEQ ID NO: 53; *PTR3R*: ACAGCAGTGC TTATCTGCGA TGACGAGCCG CTCTGTCATC TTTGT, SEQ ID NO: 54) and *PyrGD5F/PTRD3R* (*pyrGD5F*: ACAAAGATGC AAGAGCGGCT CGTCATCGCA GATAAGCACT GCTGT; SEQ ID NO: 55, *pyrGD3R*: TGAGACGCTG TTTCACCGAG TACATCGCCA ATCACCTCAA TCAC, SEQ ID NO: 56), respectively.

As shown in FIG. 14, the DNA fragment of *pGPDALaeA* (SEQ ID NO: 44) was isolated from pPTRpGPDALaeA (FIG. 13) by HindIII digestion. The yeast gap repairing vector pRS426 was double digested with restriction enzyme HindIII and XhoI. Hundred nanograms of each DNA fragment generated from PCR (i.e., SEQ ID NOS: 45-56) or restriction enzyme digestions were used for *S. cerevisiae* transformation. The gap repairing plasmid DNA in the total *S. cerevisiae* genomic DNA was isolated by transferred into the Top10 *E. coli* cells.

The transformed plasmid DNA was confirmed by PCR and digested with PmeI. The PmeI DNA fragment (SEQ ID NO: 57) was used for *A. niger* transformation.

One skilled in the art will appreciate that although the *pyrG* upstream and downstream sequences, *trpC* transcriptional terminator sequence, and *ptrA* sequence used were from particular organisms, one skilled in the art will appreciate that variants of these sequences can be used in the fungi and methods provided herein, such as those from other *Aspergillus* species. In one example, *pyrG* upstream and downstream sequences having at least 80%, at least 90%, at least 95%, or at least 98% sequence identity to SEQ ID NO: 45 and 48 are used in the fungi and methods provided herein. In one example, a *trpC* transcriptional terminator sequence having at least 80%, at least 90%, at least 95%, or at least 98% sequence identity to SEQ ID NO: 46 is used in the fungi and methods provided herein. In one example, a *ptrA* sequence having at least 80%, at least 90%, at least 95%, or at least 98% sequence identity to SEQ ID NO: 47 is used in the fungi and methods provided herein.

EXAMPLE 7

Expression of *pGPDALaeA*-Containing Transgene in *A. niger* *alg3Δ*

This example describes methods used to introduce *pGPDALaeA* (SEQ ID NO: 57) into *A. niger*.

The originally transformed *A. niger* colonies were picked from the minimal medium plates with 0.1 μg/ml pyrithiamine hydrobromide selection on minimal medium agar plates without thiamine supplementation. The single spore colonies were picked for spore production after the initial transformant spores were grown on the same selection medium plates. The biomass was harvested from the single spore colony isolates grown in minimal medium with pyrithiamine selection and dried in the VirTis bench top freeze dryer. The genomic DNA was prepared for PCR confirmation of *pGPDALaeA* insertion in transgenic *A. niger*. As shown in FIG. 15A, the primer set of *PTR5F* (SEQ ID NO: 53) and *PTR3R* (SEQ ID NO: 54) was used to confirm the presence of *A. oryzae* pyrithiamine resistance gene (*ptrA*) in transgenic *A. niger* with the expected size of 2kb PCR DNA fragment. As shown in FIG. 15B, the primer set *LaeA5F* (SEQ ID NO: 42) and *TRP3R* (SEQ ID NO: 52) was used to demonstrate that the transgene *A. nidulans* *LaeA* was under the control of *gpdA* promoter

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and trpC transcriptional terminator of *A. nidulans* with the expected 3.4 kb PCR fragment size. The genomic DNA of parent strain and the plasmid DNA of transgene vector carrying the pGDPALaeA fragment were used for negative and positive references.

EXAMPLE 8

Increased Production of Citric Acid

This example describes methods used to demonstrate that citric acid production was increased in the presence of increased expression of LaeA, alone or in combination with deletion of Alg3.

Citric acid was produced as described in Example 1.

As shown in FIG. 16, of citric acid production was increased in the Alg3Δ mutant (Alg3), and in the mutants over-expressing LaeA in Alg3Δ (LaeA-1 and LaeA-2), as compared to the parent strain (kusA).

EXAMPLE 9

pBSK-LaeAA Plasmid Vector Construction

This example describes methods used to generate the pBSK-LaeAA plasmid vector (FIG. 17A). The 5' (SEQ ID NO: 61) and 3' (SEQ ID NO: 62) ends of the *Aspergillus niger* LaeA gene were isolated from *A. niger* genomic DNA by PCR using oligonucleotide sets D1 & D2, and D5 & D6 (see Table 4 below). The hph expression cassette was isolated from plasmid vector DNA of pCB1003 (SEQ ID NO: 63) by PCR using the oligonucleotide sets D3 & D4. The DNA fragments were assembled together into the backbone plasmid vector of pBSK linearized with restriction endonucleases of both HindIII and PstI using the Gibson assembly cloning kit. The assembled plasmid DNA was transferred into the Top10 *E. coli* competent cells by lithium acetate mediated transformation. The transformed bacterial colonies were screened for the DNA fragment insertion by restriction endonuclease digestion of PvuII and XhoI. The plasmid DNA for the transformed colonies was prepared and digested with endonucleases of HindIII and XbaI for further LaeA deletion in *A. niger*.

TABLE 4

Oligonucleotide Primers for <i>A. niger</i> LaeA Deletion		
Name	Sequence	SEQ ID NO:
D1	gtcgacggtatcgataAGCTTCAAGACAGCGGCTGCAA	67
D2	gccgacgctGGTAGAGATACAGGGGTTT	68
D3	ctctaccaCGGTCGGCATCTACTCTATTC	69
D4	gcgactgaGCTGGAGCTAGTGGAGGT	70
D5	gctccagctCAGTCGCATCTTTCACTG	71
D6	atcccccgggctgcaTGGTAGGCTGTCTCAAGG	72
SC7	ACTGCAGCAGAGATGCCATCT	73
SC8	CGTTATGTTTATCGGCACCTTGTCAT	74

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EXAMPLE 10

pRS426-*A. niger* LaeA Complementation Plasmid Vector Construction

This example describes methods used to generate the pBSK-LaeAA plasmid vector (FIG. 17B). The entire LaeA gene (2.34kb; SEQ ID NO: 64) containing 740 bp of promoter region and 330 bp of transcriptional terminator region was isolated from *A. niger* genomic DNA by PCR using oligonucleotide set CP9 & CP10 (see Table 5 below). The DNA fragment was ligated into the plasmid DNA of pRS426-*A. nidulans* LaeA over-expression vector at a HindIII restriction endonuclease site after the pGDPALaeA fragment in the vector was removed by agarose gel separation. The assembled plasmid DNA was transferred into the Top10 *E. coli* competent cells by lithium acetate mediated transformation. The transformed bacterial colonies were screened for the DNA fragment insertion by restriction endonuclease digestion of BamHI and XhoI. The plasmid DNA for the transformed colonies was prepared and digested with PmeI restriction endonuclease for *A. niger* LaeA expression at pyrG locus to complement the LaeA deletion in *A. niger* LaeAA mutants.

TABLE 5

Oligonucleotide Primers for <i>A. niger</i> LaeA Complementation		
Name	Sequence	SEQ ID NO:
CP9	agggtgattggcgatgaTGCCGTTACGTGTCTGC	75
CP10	acagccatcggtccaTCTCTCTCGTAACGCCTG	76
SC11	ggatagaatcgggtgccgctgatct	77
SC12	gagaaccatggcaccgaaggt	78

EXAMPLE 11

Deletion of *A. niger* LaeA Gene in *A. niger*

This example describes methods used to introduce the HindIII/XbaI DNA fragment contain the LaeA deletion cassette (SEQ ID NO: 65) into *A. niger*.

The originally transformed *A. niger* colonies were picked from the minimal medium plates with 100 μg/ml Hygromycin B selection on minimal medium agar plates. The single spore colonies were picked for spore production after the initial transformant spores were grown on the same selection medium plates. The biomass was harvested from the single spore colony isolates grown in complete medium with Hygromycin B selection and dried in the VirTis bench top freeze dryer. The genomic DNA was prepared for PCR confirmation of the replacement of LaeA gene coding region in transgenic *A. niger*. As shown in FIG. 18, the primer set SC7 and SC8 (see Table 4) was used to confirm the hph replacement of the LaeA gene in transgenic *A. niger* with the expected 2kb PCR DNA fragment. The genomic DNA of parent strain was used for negative and positive references.

EXAMPLE 12

Expression of pGDPALaeA-Containing Transgene in *A. niger*

This example describes methods used to introduce the *A. niger* LaeA gene into the pyrG locus in the *A. niger* LaeAA

mutant to complement the loss of *LaeA* function with *PmeI* DNA fragment (SEQ ID NO: 66).

The originally transformed *A. niger* colonies were picked from the minimal medium plates with 0.1 µg/ml pyrithiamine hydrobromide selection on minimal medium agar plates without thiamine supplementation. The single spore colonies were picked for spore production after the initial transformant spores were grown on the same selection medium plates. The biomass was harvested from the single spore colony isolates grown in minimal medium with pyrithiamine selection and dried in the VirTis bench top freeze dryer. The genomic DNA was prepared for PCR confirmation of *LaeA* gene insertion in transgenic *A. niger*. As shown in FIG. 19, the primer set of PTR5F (SEQ ID NO: 53) and PTR3R (SEQ ID NO: 54) was used to confirm the presence of *A. oryzae* pyrithiamine resistance gene (*ptrA*) in transgenic *A. niger* with the expected size of 2kb PCR DNA fragment. The genomic DNA of the parent strain and the plasmid DNA of the transgene vector carrying the *A. oryzae* pyrithiamine resistance gene (*ptrA*) fragment were used for negative and positive references.

EXAMPLE 13

The Effects of *LaeA* Gene Deletion and Complementation and Over-Expression of *A. nidulans* *LaeA* Gene on Citric Acid Production

This example describes methods used to demonstrate that citric acid production in *A. niger* mutant strains was significantly influenced by perturbing the *LaeA* gene expression in *A. niger*. At least three individual clones per mutant strain were selected for spore production on complete medium

plates at 30° C. for 4 days. The spores were counted with a hemocytometer. Citric acid production culture was initiated by inoculation with a 1×10⁶ spore/ml of 75 ml citric acid production culture medium in a 250 ml baffled Erlenmeyer glass flask siliconized with a 5% solution of dichlorodimethylsilane in hexane solvent. The culture was maintained at 30° C. and 220 rpm in shaker incubator for 5 days. One microliter of culture was harvested and briefly spun down in a microcentrifuge at full speed. The supernatants of cultures were diluted 200-fold with dH₂O prior to citric acid measurement. The citric acid in the supernatant was quantified with the citric acid assay kit from R-Biopharm AG. The detailed procedure for the citric acid assay was performed essentially according to the manufacturer's description. The results in FIG. 20 show that deletion of the *LaeA* gene in *A. niger* (*LaeAΔ*) led to loss of citric acid production in citric acid production culture. When the original *A. niger* *LaeA* gene was used for complementation in the *LaeAΔ* mutant (*cLaeAΔ*) at the *pyrG* locus, the citric acid production was partially recovered, which indicates the importance of chromosome location of *LaeA* gene. When *A. nidulans* *LaeA* was over-expressed in the *A. niger* parent strain (*LaeA*), the citric acid production was higher than the parent strain. This indicates that the *LaeA* is involved in citric acid production in *A. niger*.

In view of the many possible embodiments to which the principles of the disclosed invention may be applied, it should be recognized that the illustrated embodiments are only examples of the disclosure and should not be taken as limiting the scope of the invention. Rather, the scope of the invention is defined by the following claims. We therefore claim as our invention all that comes within the scope and spirit of these claims.

SEQUENCE LISTING

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<210> SEQ ID NO 1

<211> LENGTH: 3888

<212> TYPE: DNA

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<222> LOCATION: (1307) .. (1392)

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cattgttctt ttacaccgaa actaacgcat gcccgcgccc tagtgaacgc atcgctgcta 240

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gca agg tcc ctt cat tac caa ttc ttc gcc tac ctc tcc tgg gcg aca Ala Arg Ser Leu His Tyr Gln Phe Phe Ala Tyr Leu Ser Trp Ala Thr 330 335 340	2369
ccc ttc ctc ctc tgg cgc gca ggg ttt cat cca atc ttg ctg tac ctt Pro Phe Leu Leu Trp Arg Ala Gly Phe His Pro Ile Leu Leu Tyr Leu 345 350 355	2417
atc tgg gct atg caa gag tgg gct tgg aac aca ttc ccc agc acc aac Ile Trp Ala Met Gln Glu Trp Ala Trp Asn Thr Phe Pro Ser Thr Asn 360 365 370	2465
ctc agt tcc atc att gtt gtc ctc tca ctt gct acc cag agt ttc ggc Leu Ser Ser Ile Ile Val Val Leu Ser Leu Ala Thr Gln Ser Phe Gly 375 380 385 390	2513
gtc ctt gcg aat agt gcc agc gcc ttt tat acc atg cgt tcg aac cct Val Leu Ala Asn Ser Ala Ser Ala Phe Tyr Thr Met Arg Ser Asn Pro 395 400 405	2561
agc ggt aaa gag cat aac caa tagaagtgc acccgccag tatcgagatc Ser Gly Lys Glu His Asn Gln 410	2612
gggctgtgac aggtgcatcg ataatcgcaa tcagtcttgt acccatgaga atccctgaaa	2672
aagtaagact gctctgtcag gtatgccatt gcccatgcga taggttcgga cgcctaaagg	2732
atcaatcaag atgccaatca agcatccgac tcateggaag aaggcatctt gccgacattg	2792
gactcatcct ctctgtccga gtcgtggcg acaacagcag cttgcttagc gaactccttt	2852
ggcctgcaga gggatcagta gtatccccag gcaccgcgat tgagggatcc actcaccaaa	2912
acggggcttt gcgccacgct ttctcatcac gaacgatgtt gactatttct cccttgagct	2972
tggtgtcgcg gccctcaaga gcattggtat cgatgaccac gtaggaaccc cgcttgatcc	3032
agatggtcga tcggaagcgg gcaggagct ccacccaaac cgactccttc gagggaggtt	3092

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ccaccgagta gatgttgttt ccggtcgctt tgatagcccg ggcaattaga tggccctgag 3152
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atgaacacca aagagactga agaaagagtg agcaaatgc gagaaacctg cgactgcgga 3332
gagttttact cgaatgaaga aattcgggcg gcagaaaatc cccgggtggg agtgtgggtc 3392
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<210> SEQ ID NO 2

<211> LENGTH: 413

<212> TYPE: PRT

<213> ORGANISM: *Aspergillus niger*

<400> SEQUENCE: 2

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Met Asp Trp Met Arg Leu Ile Arg Asp Leu Cys Phe Asn Pro Arg His
1          5          10          15

Thr Lys Trp Met Ala Pro Leu Leu Val Leu Gly Asp Ala Phe Leu Cys
20          25          30

Ala Leu Ile Ile Trp Lys Val Pro Tyr Thr Glu Ile Asp Trp Ala Thr
35          40          45

Tyr Met Gln Gln Ile Ser Leu Tyr Leu Ser Gly Glu Arg Asp Tyr Thr
50          55          60

Leu Ile Arg Gly Ser Thr Gly Pro Leu Val Tyr Pro Ala Ala His Val
65          70          75          80

Tyr Ser Tyr Thr Ala Leu Tyr His Leu Thr Asp Glu Gly Arg Asp Ile
85          90          95

Phe Phe Gly Gln Ile Leu Phe Ala Val Leu Tyr Leu Ile Thr Leu Val
100         105         110

Val Val Leu Cys Cys Tyr Arg Gln Ser Gly Ala Pro Pro Tyr Leu Leu
115         120         125

Pro Leu Leu Val Leu Ser Lys Arg Leu His Ser Val Tyr Val Leu Arg
130         135         140

Leu Phe Asn Asp Gly Leu Ala Ala Leu Ala Met Trp Val Ala Ile Leu
145         150         155         160

Leu Phe Met Asn Arg Lys Trp Thr Ala Ala Val Ala Val Trp Ser Thr
165         170         175

Gly Val Ala Ile Lys Met Thr Leu Leu Leu Leu Ala Pro Ala Ile Ala
180         185         190

Val Val Thr Val Leu Ser Leu Ser Leu Gly Pro Ser Val Gly Leu Gly
195         200         205

Val Leu Ala Val Leu Val Gln Val Leu Leu Ala Ile Pro Phe Leu Gln
210         215         220

Asn Asn Pro Ala Gly Tyr Leu Ser Arg Ala Phe Glu Leu Thr Arg Gln

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225	230	235	240
Phe Met Phe Lys Trp Thr Val Asn Trp Arg Phe Val Gly Glu Glu Val			
	245	250	255
Phe Leu Ser Lys Ser Phe Ser Leu Ala Leu Leu Ala Val His Ile Val			
	260	265	270
Leu Leu Gly Ala Phe Ala Val Thr Gly Trp Leu Arg Tyr Ser Arg Ser			
	275	280	285
Ser Leu Pro Ala Phe Ile Arg Asn Leu Leu Ala Gly Arg His Arg Thr			
	290	295	300
Val Ser Leu Pro Lys Pro Tyr Ile Met Ser Val Met Leu Ser Ser Leu			
	305	310	315
Thr Val Gly Leu Leu Cys Ala Arg Ser Leu His Tyr Gln Phe Phe Ala			
	325	330	335
Tyr Leu Ser Trp Ala Thr Pro Phe Leu Leu Trp Arg Ala Gly Phe His			
	340	345	350
Pro Ile Leu Leu Tyr Leu Ile Trp Ala Met Gln Glu Trp Ala Trp Asn			
	355	360	365
Thr Phe Pro Ser Thr Asn Leu Ser Ser Ile Ile Val Val Leu Ser Leu			
	370	375	380
Ala Thr Gln Ser Phe Gly Val Leu Ala Asn Ser Ala Ser Ala Phe Tyr			
	385	390	395
Thr Met Arg Ser Asn Pro Ser Gly Lys Glu His Asn Gln			
	405	410	

<210> SEQ ID NO 3

<211> LENGTH: 1923

<212> TYPE: DNA

<213> ORGANISM: Aspergillus oryzae

<220> FEATURE:

<221> NAME/KEY: CDS

<222> LOCATION: (1)..(1242)

<400> SEQUENCE: 3

atg gag ttg aag cac ttc atc cac gaa ctc tgc cta aac ccc aga cat	48
Met Glu Leu Lys His Phe Ile His Glu Leu Cys Leu Asn Pro Arg His	
1 5 10 15	
aca aaa tgg att gca cgc ctt ctt gtc ata ggc gat gcc ttc cta tgt	96
Thr Lys Trp Ile Ala Pro Leu Leu Val Ile Gly Asp Ala Phe Leu Cys	
20 25 30	
gct ctt atc atc tgg aag atc cca tat act gag atc gac tgg acg acg	144
Ala Leu Ile Ile Trp Lys Ile Pro Tyr Thr Glu Ile Asp Trp Thr Thr	
35 40 45	
tac atg cag cag ata gcg ctt tac atc tct ggc gaa cgt gac tat acc	192
Tyr Met Gln Gln Ile Ala Leu Tyr Ile Ser Gly Glu Arg Asp Tyr Thr	
50 55 60	
ctg atc aag ggg tcc act gga ccc ctt gta tac ccg gcc gcc cac gta	240
Leu Ile Lys Gly Ser Thr Gly Pro Leu Val Tyr Pro Ala Ala His Val	
65 70 75 80	
tat agc tac atg gca ctc tat cac cta aca gat gaa ggc cga gat att	288
Tyr Ser Tyr Met Ala Leu Tyr His Leu Thr Asp Glu Gly Arg Asp Ile	
85 90 95	
ctt ttc ggg cag ata tta ttt gct gtc ctt tac ctt gtc acg cta gca	336
Leu Phe Gly Gln Ile Leu Phe Ala Val Leu Tyr Leu Val Thr Leu Ala	
100 105 110	
gtt gtg atg gtt tgc tac agg cag tca ggt gcc cct ccg tac ctg ttt	384
Val Val Met Val Cys Tyr Arg Gln Ser Gly Ala Pro Pro Tyr Leu Phe	
115 120 125	
cct ctt ctt gtc ctt tcc aag cgg ctc cac agt gtc ttt gtt ttg cgc	432

Pro	Leu	Leu	Val	Leu	Ser	Lys	Arg	Leu	His	Ser	Val	Phe	Val	Leu	Arg																																																				
																130																	135																	140																	
ctt	ttc	aat	gat	ggc	ctc	gcg	gtc	tgt	gcc	atg	tgg	ata	gcg	att	ctg	480																																																			
Leu	Phe	Asn	Asp	Gly	Leu	Ala	Val	Cys	Ala	Met	Trp	Ile	Ala	Ile	Leu																																																				
																145																	150																	155																	160
ctc	ttc	cag	aat	aag	aaa	tgg	acg	gct	ggg	gtt	acg	gcc	tgg	act	gtt	528																																																			
Leu	Phe	Gln	Asn	Lys	Lys	Trp	Thr	Ala	Gly	Val	Thr	Ala	Trp	Thr	Val																																																				
																165																	170																	175																	
ggg	gtt	ggc	att	aag	atg	acg	tta	ctg	ctc	ctt	gcg	cca	gcc	att	gcg	576																																																			
Gly	Val	Gly	Ile	Lys	Met	Thr	Leu	Leu	Leu	Leu	Ala	Pro	Ala	Ile	Ala																																																				
																180																	185																	190																	
gtg	gta	act	gtg	ctc	agc	ctt	tcc	ctc	gtg	cct	agt	att	cga	ctt	gga	624																																																			
Val	Val	Thr	Val	Leu	Ser	Leu	Ser	Leu	Val	Pro	Ser	Ile	Arg	Leu	Gly																																																				
																195																	200																	205																	
att	cta	gca	ttg	ctc	att	cag	gtt	cta	cta	gcg	att	ccc	ttc	cta	cag	672																																																			
Ile	Leu	Ala	Leu	Leu	Ile	Gln	Val	Leu	Leu	Ala	Ile	Pro	Phe	Leu	Gln																																																				
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Phe	Met	Phe	Lys	Trp	Thr	Val	Asn	Trp	Arg	Phe	Val	Gly	Glu	Asp	Leu																																																				
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ttc	cta	tcc	aaa	cag	ttt	tct	cta	gcc	tta	cta	ggg	ttg	cat	att	ttt	816																																																			
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ctg	ctg	gga	tta	ttt	gtt	acc	aca	ggc	tgg	tta	cgg	ccg	tca	gga	tct	864																																																			
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aac	gtc	cct	gac	ttc	ctc	cgg	agc	cta	ctc	caa	gga	cgc	caa	cgc	acc	912																																																			
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gtg	gtg	ctt	tct	aag	tct	ttc	ata	atg	acc	gtg	atg	ttg	aca	tcg	ctg	960																																																			
Val	Val	Leu	Ser	Lys	Ser	Phe	Ile	Met	Thr	Val	Met	Leu	Thr	Ser	Leu																																																				
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gcg	atc	ggg	ttg	ttg	tgc	gca	agg	tcc	ctt	cat	tac	caa	ttc	ttt	gcc	1008																																																			
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tat	ctc	tcc	tgg	gct	acg	cct	tgc	ctt	ctc	tgg	cgg	gct	cgg	ctc	cat	1056																																																			
Tyr	Leu	Ser	Trp	Ala	Thr	Pro	Cys	Leu	Leu	Trp	Arg	Ala	Arg	Leu	His																																																				
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cgg	atc	ctt	ata	tat	gcg	atc	tgg	gca	cta	cag	gag	tgg	gct	tgg	aat	1104																																																			
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<210> SEQ ID NO 4
<211> LENGTH: 413
<212> TYPE: PRT
<213> ORGANISM: Aspergillus oryzae

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<400> SEQUENCE: 4

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Met Glu Leu Lys His Phe Ile His Glu Leu Cys Leu Asn Pro Arg His
1          5          10          15

Thr Lys Trp Ile Ala Pro Leu Leu Val Ile Gly Asp Ala Phe Leu Cys
20        25        30

Ala Leu Ile Ile Trp Lys Ile Pro Tyr Thr Glu Ile Asp Trp Thr Thr
35        40        45

Tyr Met Gln Gln Ile Ala Leu Tyr Ile Ser Gly Glu Arg Asp Tyr Thr
50        55        60

Leu Ile Lys Gly Ser Thr Gly Pro Leu Val Tyr Pro Ala Ala His Val
65        70        75        80

Tyr Ser Tyr Met Ala Leu Tyr His Leu Thr Asp Glu Gly Arg Asp Ile
85        90        95

Leu Phe Gly Gln Ile Leu Phe Ala Val Leu Tyr Leu Val Thr Leu Ala
100       105       110

Val Val Met Val Cys Tyr Arg Gln Ser Gly Ala Pro Pro Tyr Leu Phe
115       120       125

Pro Leu Leu Val Leu Ser Lys Arg Leu His Ser Val Phe Val Leu Arg
130       135       140

Leu Phe Asn Asp Gly Leu Ala Val Cys Ala Met Trp Ile Ala Ile Leu
145       150       155       160

Leu Phe Gln Asn Lys Lys Trp Thr Ala Gly Val Thr Ala Trp Thr Val
165       170       175

Gly Val Gly Ile Lys Met Thr Leu Leu Leu Leu Ala Pro Ala Ile Ala
180       185       190

Val Val Thr Val Leu Ser Leu Ser Leu Val Pro Ser Ile Arg Leu Gly
195       200       205

Ile Leu Ala Leu Leu Ile Gln Val Leu Leu Ala Ile Pro Phe Leu Gln
210       215       220

Gly Asn Pro Ile Gly Tyr Val Ala Arg Ala Phe Glu Leu Thr Arg Gln
225       230       235       240

Phe Met Phe Lys Trp Thr Val Asn Trp Arg Phe Val Gly Glu Asp Leu
245       250       255

Phe Leu Ser Lys Gln Phe Ser Leu Ala Leu Leu Gly Leu His Ile Phe
260       265       270

Leu Leu Gly Leu Phe Val Thr Thr Gly Trp Leu Arg Pro Ser Gly Ser
275       280       285

Asn Val Pro Asp Phe Leu Arg Ser Leu Leu Gln Gly Arg Gln Arg Thr

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290	295	300
Val Val Leu Ser Lys Ser Phe Ile Met Thr	Val Met Leu Thr Ser Leu	
305	310	315 320
Ala Ile Gly Leu Leu Cys Ala Arg Ser Leu His Tyr Gln Phe Phe Ala		
	325	330 335
Tyr Leu Ser Trp Ala Thr Pro Cys Leu Leu Trp Arg Ala Arg Leu His		
	340	345 350
Pro Ile Leu Ile Tyr Ala Ile Trp Ala Leu Gln Glu Trp Ala Trp Asn		
	355	360 365
Val Tyr Pro Ser Thr Asn Ala Ser Ser Ser Val Val Val Phe Ser Leu		
	370	375 380
Ala Val Gln Val Phe Gly Val Leu Leu Asn Ser Arg Asn Ala Leu Ser		
	385	390 395 400
Asp Ala Pro Pro Arg Arg Lys Gly Lys Glu His Ile Gln		
	405	410

<210> SEQ ID NO 5
 <211> LENGTH: 22
 <212> TYPE: DNA
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
 <223> OTHER INFORMATION: Alg3-ForScr primer

<400> SEQUENCE: 5

cggtttccct tcagtttcca gt 22

<210> SEQ ID NO 6
 <211> LENGTH: 49
 <212> TYPE: DNA
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
 <223> OTHER INFORMATION: Alg3-1 primer

<400> SEQUENCE: 6

gtaacgccag ggttttccca gtcacgacgt cataacttct ctcccctcc 49

<210> SEQ ID NO 7
 <211> LENGTH: 49
 <212> TYPE: DNA
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
 <223> OTHER INFORMATION: Alg3-2 primer

<400> SEQUENCE: 7

atccacttaa cggtactgaa atctccaact tcatggacac acacagacc 49

<210> SEQ ID NO 8
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 <212> TYPE: DNA
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
 <223> OTHER INFORMATION: Hph-F primer

<400> SEQUENCE: 8

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<210> SEQ ID NO 9
 <211> LENGTH: 49
 <212> TYPE: DNA
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
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<400> SEQUENCE: 9

gtactactg atccctctgc gtcggagaca gaagatgata ttgaaggag 49

<210> SEQ ID NO 10

<211> LENGTH: 49

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: Alg3-3 primer

<400> SEQUENCE: 10

ctccttcaat atcatcttct gtctccgacg cagagggatc agtagtagc 49

<210> SEQ ID NO 11

<211> LENGTH: 49

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: Alg3-4 primer

<400> SEQUENCE: 11

gcggataaca atttcacaca ggaaacagcc gtgagagggt ttagtagc 49

<210> SEQ ID NO 12

<211> LENGTH: 21

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: Alg3-RevScr primer

<400> SEQUENCE: 12

aagctgagag cgacatcttc a 21

<210> SEQ ID NO 13

<211> LENGTH: 25

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: hyg-RevScr primer

<400> SEQUENCE: 13

gtacttctac acagccatcg gtcca 25

<210> SEQ ID NO 14

<211> LENGTH: 25

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: hyg-ForScr primer

<400> SEQUENCE: 14

gtacttctac acagccatcg gtcca 25

<210> SEQ ID NO 15

<211> LENGTH: 24

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: pryGScr primer

<400> SEQUENCE: 15

tctgtgtct tgcagaggt cett 24

<210> SEQ ID NO 16

<211> LENGTH: 24

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<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: primer

<400> SEQUENCE: 16

agcgtaggac aaggtcgtct ctgt 24

<210> SEQ ID NO 17
<211> LENGTH: 58
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: 5-pyrG5F primer

<400> SEQUENCE: 17

gtaacgccag gggtttccca gtcacgacgt ttaaacaatgc atcattctcc cgetttgt 58

<210> SEQ ID NO 18
<211> LENGTH: 43
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: 5-pyrG3R primer

<400> SEQUENCE: 18

agaaagagtc accggtcacg acatcgccaa tcacctcaat cac 43

<210> SEQ ID NO 19
<211> LENGTH: 43
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: ble5F primer

<400> SEQUENCE: 19

gtgattgagg tgattggcga tgcgtgacc ggtgactctt tct 43

<210> SEQ ID NO 20
<211> LENGTH: 42
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Ble3R primer

<400> SEQUENCE: 20

tccaaccttg tagcaaccaa agcttcgagc gtcccaaaac ct 42

<210> SEQ ID NO 21
<211> LENGTH: 42
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Alg3-5F1 primer

<400> SEQUENCE: 21

agggttttggg acgctcgaag ctttggttgc tacaaggttg ga 42

<210> SEQ ID NO 22
<211> LENGTH: 29
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Alg3-3R1 primer

<400> SEQUENCE: 22

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tcaagtagag cacagcaaat agtatctga 29

<210> SEQ ID NO 23
<211> LENGTH: 29
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Alg3-5F2 primer

<400> SEQUENCE: 23

tcagatacta ttgctgtgc tctacttga 29

<210> SEQ ID NO 24
<211> LENGTH: 42
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Alg3-3R2 primer

<400> SEQUENCE: 24

ttgatccttg tgccacacca tcctacgtgg tcatcgatac ca 42

<210> SEQ ID NO 25
<211> LENGTH: 42
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: 3-pyrG5F primer

<400> SEQUENCE: 25

tggtatcgat gaccacgtag gatggtgtgg cacaaggatc aa 42

<210> SEQ ID NO 26
<211> LENGTH: 60
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: 3=pyrG3R primer

<400> SEQUENCE: 26

gcggaataca atttcacaca ggaaacagcg tttaaactgt gccagtcaat tgtccgaagt 60

<210> SEQ ID NO 27
<211> LENGTH: 21
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Alg3seq-1 primer

<400> SEQUENCE: 27

tacagacgcg tgtacgcatg t 21

<210> SEQ ID NO 28
<211> LENGTH: 24
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Alg3seq-2 primer

<400> SEQUENCE: 28

tgctattgtc cacagatacc gaga 24

<210> SEQ ID NO 29
<211> LENGTH: 22
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence

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<220> FEATURE:

<223> OTHER INFORMATION: Alg3seq-3 primer

<400> SEQUENCE: 29

gagctaacca gacagttcat gt

22

<210> SEQ ID NO 30

<211> LENGTH: 21

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: Alg3seq-4 primer

<400> SEQUENCE: 30

tcgctcgtagc gcattgatcc t

21

<210> SEQ ID NO 31

<211> LENGTH: 413

<212> TYPE: PRT

<213> ORGANISM: Aspergillus nidulans

<400> SEQUENCE: 31

Met Ala Leu Thr Asp Leu Val Ser Gly Leu Cys Ser Asn Pro Lys His
1 5 10 15Thr Lys Trp Ile Ala Pro Ile Leu Asn Ile Ala Asp Gly Leu Leu Cys
20 25 30Ala Phe Ile Ile Trp Lys Val Pro Tyr Thr Glu Ile Asp Trp Thr Thr
35 40 45Tyr Met Gln Gln Val Lys Leu Tyr Leu Ser Gly Glu Arg Asp Tyr Thr
50 55 60Leu Ile Lys Gly Ser Thr Gly Pro Leu Val Tyr Pro Ala Ala His Val
65 70 75 80Tyr Ser Tyr Ser Leu Phe His His Leu Thr Asp Glu Gly Arg Asp Ile
85 90 95Val Phe Gly Gln Ile Ile Phe Ala Phe Leu Tyr Leu Ile Cys Leu Thr
100 105 110Val Val Met Ala Cys Tyr Arg Arg Val Gly Ala Pro Pro Tyr Leu Phe
115 120 125Pro Leu Leu Val Leu Ser Lys Arg Leu His Ser Val Tyr Met Leu Arg
130 135 140Leu Phe Asn Asp Gly Leu Ala Ala Leu Ala Met Trp Gly Ser Ile Trp
145 150 155 160Leu Phe Ile Asn Arg Lys Trp Thr Pro Ala Val Val Leu Trp Ser Leu
165 170 175Gly Leu Gly Val Lys Met Thr Leu Ile Leu Leu Val Pro Ala Val Met
180 185 190Val Val Leu Ala Leu Ser Leu Asp Ile Gly Arg Cys Ile Arg Leu Ala
195 200 205Gly Leu Ala Leu Gly Ile Gln Ile Leu Leu Ala Ile Pro Phe Leu Lys
210 215 220Thr Asn Pro Ser Gly Tyr Phe Glu Arg Ala Phe Glu Phe Gly Arg Gln
225 230 235 240Phe Met Phe Lys Trp Thr Val Asn Trp Arg Phe Val Gly Glu Asp Ile
245 250 255Phe Leu Ser Lys Gly Phe Trp Ala Gly Leu Ile Val Leu His Leu Leu
260 265 270

Ile Leu Val Val Leu Gly Phe Thr Cys Phe Leu Asn Pro Ser Gly Thr

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275	280	285
Ser Leu Pro Asp Phe Ala Gly	Arg Phe Leu Thr Gly	Gln His Arg Gly
290	295	300
Ile Ala Leu His Pro Ser Phe	Ile Met Ser Ala Leu Leu Thr Ser Leu	
305	310	315
Ser Val Gly Leu Leu Cys Ala Arg Ser Leu His Tyr Gln Phe Phe Ala		335
325	330	
Tyr Leu Ser Trp Ala Thr Pro Phe Leu Leu Trp Gln Ala Gly Tyr His		350
340	345	
Pro Ile Leu Val Tyr Ala Leu Trp Leu Val Gln Glu Trp Ala Trp Asn		365
355	360	
Val Tyr Pro Ser Thr Asn Leu Ser Ser Ala Ala Val Val Leu Leu Leu		380
370	375	
Gly Ala Gln Val Leu Gly Val Leu Val Asn Arg Asp Arg Ala Phe Pro		400
385	390	395
Ser Ser Pro Pro Thr Pro Lys Ala Lys Gln His Val Gln		
405	410	
<210> SEQ ID NO 32		
<211> LENGTH: 434		
<212> TYPE: PRT		
<213> ORGANISM: Fusarium oxysporum		
<400> SEQUENCE: 32		
Met Pro Glu Ser Ala Ser Gly Thr Leu Ser Gln Gly Val Arg Phe Leu		
1	5	10
Arg Asn Val Leu Asn Gly Arg His Ala Leu Ser Lys Leu Ile Pro Ile		
20	25	30
Ala Leu Trp Leu Val Asp Ala Leu Gly Cys Gly Leu Ile Ile Trp Lys		
35	40	45
Ile Pro Tyr Thr Glu Ile Asp Trp Val Ala Tyr Met Gln Gln Ile Ser		
50	55	60
Gln Phe Val Ser Gly Glu Arg Asp Tyr Thr Lys Met Glu Gly Asp Thr		
65	70	75
Gly Pro Leu Val Tyr Pro Ala Ala His Val Tyr Thr Tyr Thr Gly Leu		
85	90	95
Tyr Tyr Ile Thr Asp Lys Gly Thr Asn Ile Leu Leu Ala Gln Gln Ile		
100	105	110
Phe Ala Val Leu Tyr Met Ala Thr Leu Ala Val Val Met Leu Cys Tyr		
115	120	125
Trp Lys Ala Lys Val Pro Pro Tyr Met Phe Ile Phe Leu Ile Ala Ser		
130	135	140
Lys Arg Leu His Ser Leu Phe Val Leu Arg Cys Phe Asn Asp Cys Phe		
145	150	155
Ala Val Phe Phe Leu Trp Leu Thr Ile Phe Leu Phe Gln Arg Arg Gln		
165	170	175
Trp Thr Val Gly Ser Leu Val Tyr Ser Trp Gly Leu Gly Ile Lys Met		
180	185	190
Ser Leu Leu Leu Val Leu Pro Ala Ile Gly Val Ile Leu Phe Leu Gly		
195	200	205
Arg Gly Leu Trp Pro Ser Leu Arg Leu Ala Trp Leu Met Ala Gln Ile		
210	215	220
Gln Phe Ala Ile Gly Leu Pro Phe Ile Thr Lys Asn Pro Arg Gly Tyr		
225	230	235
		240

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Ala	Ala	Arg	Ala	Phe	Glu	Leu	Ser	Arg	Gln	Phe	Gln	Phe	Lys	Trp	Thr
				245					250					255	
Val	Asn	Trp	Arg	Met	Leu	Gly	Glu	Glu	Val	Phe	Leu	Ser	Lys	Tyr	Phe
			260					265					270		
Ala	Leu	Ser	Leu	Leu	Ala	Cys	His	Ile	Leu	Val	Leu	Leu	Ile	Phe	Ile
		275					280					285			
Ser	Lys	Arg	Trp	Ile	Gln	Pro	Thr	Gly	Arg	Ser	Leu	Tyr	Asp	Leu	Ile
	290					295					300				
Pro	Ser	Phe	Leu	Arg	Leu	Lys	Ser	Pro	Phe	Thr	Met	Gln	Glu	Gln	Leu
305					310					315					320
Arg	Ile	Ser	His	Tyr	Val	Thr	Pro	Glu	Tyr	Ala	Met	Thr	Thr	Met	Leu
			325						330					335	
Thr	Ala	Asn	Leu	Ile	Gly	Leu	Leu	Phe	Ala	Arg	Ser	Leu	His	Tyr	Gln
		340						345					350		
Phe	Tyr	Ala	Tyr	Leu	Ala	Trp	Ala	Thr	Pro	Tyr	Leu	Leu	Trp	Arg	Ala
		355					360					365			
Thr	Glu	Asp	Pro	Val	Ile	Val	Ala	Ile	Ile	Trp	Ala	Ala	Gln	Glu	Trp
	370					375					380				
Ala	Trp	Asn	Val	Tyr	Pro	Ser	Thr	Asp	Leu	Ser	Ser	Thr	Ile	Ala	Val
385					390					395					400
Asn	Thr	Met	Leu	Ala	Thr	Val	Val	Leu	Val	Tyr	Leu	Gly	Thr	Ala	Arg
			405						410					415	
Arg	Ala	Val	Pro	Ala	Pro	Ala	Ala	Gln	Val	Gly	Asn	Val	Asp	Asp	Lys
		420						425					430		

Asn Lys

<210> SEQ ID NO 33

<211> LENGTH: 502

<212> TYPE: PRT

<213> ORGANISM: Neurospora crassa

<400> SEQUENCE: 33

Met	Ala	Ala	Pro	Ser	Ser	Arg	Pro	Glu	Ser	Asn	Pro	Pro	Leu	Tyr	Lys
1				5					10					15	
Gln	Ala	Leu	Asp	Phe	Ala	Leu	Asp	Val	Ala	Asn	Gly	Arg	His	Ala	Leu
		20						25					30		
Ser	Lys	Leu	Ile	Pro	Pro	Ala	Leu	Phe	Leu	Val	Asp	Ala	Leu	Leu	Cys
		35				40					45				
Gly	Leu	Ile	Ile	Trp	Lys	Val	Pro	Tyr	Thr	Glu	Ile	Asp	Trp	Ala	Ala
	50				55					60					
Tyr	Met	Glu	Gln	Val	Ser	Gln	Ile	Leu	Ser	Gly	Glu	Arg	Asp	Tyr	Thr
65				70					75					80	
Lys	Val	Arg	Gly	Gly	Thr	Gly	Pro	Leu	Val	Tyr	Pro	Ala	Ala	His	Val
			85					90						95	
Tyr	Ile	Tyr	Thr	Gly	Leu	Tyr	His	Leu	Thr	Asp	Glu	Gly	Arg	Asn	Ile
		100					105						110		
Leu	Leu	Ala	Gln	Gln	Leu	Phe	Ala	Gly	Leu	Tyr	Met	Val	Thr	Leu	Ala
		115				120					125				
Val	Val	Met	Gly	Cys	Tyr	Trp	Gln	Ala	Lys	Ala	Pro	Pro	Tyr	Leu	Phe
	130					135					140				
Pro	Leu	Leu	Thr	Leu	Ser	Lys	Arg	Leu	His	Ser	Ile	Phe	Val	Leu	Arg
145				150					155					160	
Cys	Phe	Asn	Asp	Cys	Phe	Ala	Val	Leu	Phe	Leu	Trp	Leu	Ala	Ile	Phe
			165					170						175	

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Phe	Phe	Gln	Arg	Arg	Asn	Trp	Gln	Ala	Gly	Ala	Leu	Leu	Tyr	Thr	Leu
			180					185					190		
Gly	Leu	Gly	Val	Lys	Met	Thr	Leu	Leu	Leu	Ser	Leu	Pro	Ala	Val	Gly
		195					200					205			
Ile	Val	Leu	Phe	Leu	Gly	Ser	Gly	Ser	Phe	Val	Thr	Thr	Leu	Gln	Leu
	210					215					220				
Val	Ala	Thr	Met	Gly	Leu	Val	Gln	Ile	Ala	Ile	Gly	Leu	Pro	Phe	Ile
225					230					235					240
Thr	Lys	Asn	Pro	Arg	Gly	Tyr	Ala	Ala	Arg	Ala	Phe	Glu	Leu	Ser	Arg
				245					250					255	
Gln	Phe	Gln	Phe	Lys	Trp	Thr	Val	Asn	Trp	Arg	Met	Leu	Gly	Glu	Glu
			260					265					270		
Val	Phe	Leu	Ser	Lys	Tyr	Phe	Ala	Leu	Ser	Leu	Leu	Ala	Cys	His	Ile
		275					280					285			
Leu	Val	Leu	Leu	Ile	Leu	Ile	Gly	Val	Pro	Phe	Leu	Ala	His	Tyr	Pro
	290					295					300				
Thr	Glu	Tyr	Leu	Ser	Arg	Ala	Phe	Glu	Leu	Ser	Arg	Gln	Phe	Phe	Phe
305					310					315					320
Lys	Trp	Thr	Val	Asn	Trp	Arg	Phe	Val	Gly	Glu	Glu	Ile	Phe	Leu	Ser
				325					330					335	
Lys	Gly	Phe	Ala	Leu	Thr	Leu	Leu	Ala	Leu	His	Val	Leu	Val	Leu	Gly
			340					345					350		
Ile	Phe	Ile	Thr	Thr	Arg	Trp	Ile	Lys	Pro	Ala	Arg	Lys	Ser	Leu	Val
		355					360					365			
Gln	Leu	Ile	Ser	Pro	Val	Leu	Leu	Ala	Gly	Lys	Pro	Pro	Leu	Thr	Val
	370					375					380				
Pro	Glu	His	Arg	Ala	Ala	Ala	Arg	Asp	Val	Thr	Pro	Arg	Tyr	Ile	Met
385					390					395					400
Thr	Thr	Ile	Leu	Ser	Ala	Asn	Ala	Val	Gly	Leu	Leu	Phe	Ala	Arg	Ser
				405					410					415	
Leu	His	Tyr	Gln	Phe	Tyr	Ala	Tyr	Val	Ala	Trp	Ser	Thr	Pro	Phe	Leu
			420					425					430		
Leu	Trp	Arg	Ala	Gly	Leu	His	Pro	Val	Leu	Val	Tyr	Leu	Leu	Trp	Ala
		435					440					445			
Val	His	Glu	Trp	Ala	Trp	Asn	Val	Phe	Pro	Ser	Thr	Pro	Ala	Ser	Ser
	450					455					460				
Ala	Val	Val	Val	Gly	Val	Leu	Gly	Val	Thr	Val	Ala	Gly	Val	Trp	Phe
465					470					475					480
Gly	Ala	Arg	Glu	Glu	Trp	Glu	Pro	Gly	Met	Lys	Ser	Ser	Ser	Lys	Lys
				485					490					495	
Glu	Glu	Ala	Ala	Met	Arg										
				500											

<210> SEQ ID NO 34

<211> LENGTH: 434

<212> TYPE: PRT

<213> ORGANISM: *Saccharomyces cerevisiae*

<400> SEQUENCE: 34

Met	Ala	Gly	Gly	Lys	Lys	Lys	Ser	Ser	Thr	Ala	Pro	Ser	Arg	Phe	Gln
1				5					10					15	

Lys	Thr	Leu	Ser	Ser	Ile	Trp	Gln	Asp	Lys	His	Thr	Val	Leu	Phe	Lys
		20						25					30		

Pro	Glu	Tyr	Thr	Leu	Leu	Val	Thr	Ala	Val	Leu	Trp	Phe	Leu	Glu	Ile
		35					40					45			

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Ala	Ile	Asn	Ile	Trp	Val	Ile	Gln	Lys	Val	Ser	Tyr	Thr	Glu	Ile	Asp
50						55					60				
Trp	Lys	Ala	Tyr	Met	Asp	Glu	Val	Glu	Gly	Val	Ile	Asn	Gly	Thr	Tyr
65					70					75					80
Asp	Tyr	Thr	Gln	Leu	Lys	Gly	Asp	Thr	Gly	Pro	Leu	Val	Tyr	Pro	Ala
			85						90					95	
Gly	Phe	Val	Tyr	Ile	Phe	Thr	Gly	Leu	Tyr	Tyr	Leu	Thr	Asp	His	Gly
		100						105					110		
His	Asn	Ile	Arg	Leu	Gly	Gln	Tyr	Val	Phe	Ala	Val	Ser	Tyr	Leu	Ile
	115						120					125			
Asn	Leu	Leu	Leu	Val	Met	Arg	Ile	Tyr	His	Arg	Thr	Lys	Lys	Val	Pro
130						135					140				
Pro	Tyr	Val	Phe	Phe	Phe	Ile	Cys	Cys	Ala	Ser	Tyr	Arg	Ile	His	Ser
145					150					155					160
Ile	Phe	Ile	Leu	Arg	Leu	Phe	Asn	Asp	Pro	Val	Ala	Met	Met	Leu	Cys
			165						170					175	
Phe	Gly	Ala	Ile	Asn	Leu	Phe	Leu	Asp	Gly	Arg	Trp	Thr	Leu	Gly	Cys
		180						185					190		
Ala	Leu	Tyr	Ser	Leu	Ala	Val	Ser	Val	Lys	Met	Asn	Val	Leu	Leu	Phe
	195					200					205				
Ala	Pro	Gly	Leu	Leu	Phe	Leu	Leu	Leu	Cys	Glu	Phe	Gly	Leu	Trp	Lys
210					215						220				
Thr	Leu	Pro	Arg	Leu	Ala	Leu	Cys	Ala	Val	Ile	Gln	Leu	Leu	Val	Gly
225				230						235					240
Leu	Pro	Phe	Leu	Ile	Thr	Tyr	Pro	Val	Ser	Tyr	Ile	Ala	Asn	Ala	Phe
			245						250				255		
Asp	Leu	Gly	Arg	Val	Phe	Ile	His	Phe	Trp	Ser	Val	Asn	Phe	Lys	Phe
		260						265					270		
Val	Pro	Glu	Arg	Val	Phe	Val	Ser	Lys	Glu	Phe	Ala	Val	Cys	Leu	Leu
		275				280					285				
Ile	Ala	His	Leu	Phe	Leu	Leu	Val	Ala	Phe	Ala	Leu	Lys	Arg	Trp	Lys
290					295						300				
Arg	Ser	Gly	Ser	Ser	Ile	Trp	Thr	Ile	Leu	Lys	Asp	Pro	Ser	Glu	Arg
305				310						315					320
Lys	Glu	Thr	Ala	His	Lys	Val	Asn	Ala	Asp	Gln	Met	Val	Leu	Ile	Leu
			325					330						335	
Phe	Thr	Ser	Asn	Phe	Ile	Gly	Met	Cys	Phe	Ser	Arg	Ser	Leu	His	Tyr
		340					345						350		
Gln	Phe	Tyr	Val	Trp	Tyr	Phe	His	Thr	Leu	Pro	Tyr	Leu	Leu	Trp	Ser
	355					360					365				
Gly	Gly	Val	Lys	Lys	Leu	Ala	Arg	Leu	Leu	Arg	Val	Leu	Ile	Leu	Gly
370					375					380					
Leu	Ile	Glu	Leu	Ser	Trp	Asn	Thr	Tyr	Pro	Ser	Thr	Asn	Tyr	Ser	Ser
385				390					395						400
Leu	Ser	Leu	His	Val	Cys	His	Leu	Ile	Ile	Leu	Leu	Cys	Leu	Trp	Leu
			405					410						415	
Asn	Pro	Asn	Pro	Ala	Ser	Pro	Ser	His	Arg	Ser	Glu	Asn	Lys	Ala	Lys
		420						425					430		
Ser	His														

<210> SEQ ID NO 35

<211> LENGTH: 437

<212> TYPE: PRT

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<213> ORGANISM: *Arabidopsis thaliana*

<400> SEQUENCE: 35

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Met Ala Gly Ala Ser Ser Pro Ala Ser Leu Arg Ala Ser Arg Ser Arg
 1           5           10          15
Arg Leu Gly Lys Glu Thr Asn Arg Ser Asp Leu Phe Lys Lys Pro Ala
 20          25          30
Val Pro Phe Ala Phe Ala Leu Ile Leu Ala Asp Ala Ile Leu Val Ala
 35          40          45
Leu Ile Ile Ala Tyr Val Pro Tyr Thr Lys Ile Asp Trp Asp Ala Tyr
 50          55          60
Met Ser Gln Val Ser Gly Phe Leu Gly Gly Glu Arg Asp Tyr Gly Asn
 65          70          75          80
Leu Lys Gly Asp Thr Gly Pro Leu Val Tyr Pro Ala Gly Phe Leu Tyr
 85          90          95
Val Tyr Ser Ala Val Gln Asn Leu Thr Gly Gly Glu Val Tyr Pro Ala
100          105          110
Gln Ile Leu Phe Gly Val Leu Tyr Ile Val Asn Leu Gly Ile Val Leu
115          120          125
Ile Ile Tyr Val Lys Thr Asp Val Pro Trp Trp Ala Leu Ser Leu Leu
130          135          140
Cys Leu Ser Lys Arg Ile His Ser Ile Phe Val Leu Arg Leu Phe Asn
145          150          155          160
Asp Cys Phe Ala Met Thr Leu Leu His Ala Ser Met Ala Leu Phe Leu
165          170          175
Tyr Arg Lys Trp His Leu Gly Met Leu Val Phe Ser Gly Ala Val Ser
180          185          190
Val Lys Met Asn Val Leu Leu Tyr Ala Pro Thr Leu Leu Leu Leu Leu
195          200          205
Leu Lys Ala Met Asn Ile Ile Gly Val Val Ser Ala Leu Ala Gly Ala
210          215          220
Ala Leu Val Gln Ile Leu Val Gly Leu Pro Phe Leu Ile Thr Tyr Pro
225          230          235          240
Val Ser Tyr Ile Ala Asn Ala Phe Asp Leu Gly Arg Val Phe Ile His
245          250          255
Phe Trp Ser Val Asn Phe Lys Phe Val Pro Glu Arg Val Phe Val Ser
260          265          270
Lys Glu Phe Ala Val Cys Leu Leu Ile Ala His Leu Phe Leu Leu Val
275          280          285
Ala Phe Ala Asn Tyr Lys Trp Cys Lys His Glu Gly Gly Ile Ile Gly
290          295          300
Phe Met Arg Ser Arg His Phe Phe Leu Thr Leu Pro Ser Ser Leu Ser
305          310          315          320
Phe Ser Asp Val Ser Ala Ser Arg Ile Ile Thr Lys Glu His Val Val
325          330          335
Thr Ala Met Phe Val Gly Asn Phe Ile Gly Ile Val Phe Ala Arg Ser
340          345          350
Leu His Tyr Gln Phe Tyr Ser Trp Tyr Phe Tyr Ser Leu Pro Tyr Leu
355          360          365
Leu Trp Arg Thr Pro Phe Pro Thr Trp Leu Arg Leu Ile Met Phe Leu
370          375          380
Gly Ile Glu Leu Cys Trp Asn Val Tyr Pro Ser Thr Pro Ser Ser Ser
385          390          395          400

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Gly Leu Leu Leu Cys Leu His Leu Ile Ile Leu Val Gly Leu Trp Leu
 405 410 415
 Ala Pro Ser Val Asp Pro Tyr Gln Leu Lys Glu His Pro Lys Ser Gln
 420 425 430
 Ile His Lys Lys Ala
 435
 <210> SEQ ID NO 36
 <211> LENGTH: 433
 <212> TYPE: PRT
 <213> ORGANISM: Homo sapiens
 <400> SEQUENCE: 36
 Arg Lys Arg Gly Arg Ser Gly Ser Ala Ala Gln Ala Glu Gly Leu Cys
 1 5 10 15
 Lys Gln Trp Leu Gln Arg Ala Trp Gln Glu Arg Arg Leu Leu Leu Arg
 20 25 30
 Glu Pro Arg Tyr Thr Leu Leu Val Ala Ala Cys Leu Cys Leu Ala Glu
 35 40 45
 Val Gly Ile Thr Phe Trp Val Ile His Arg Val Ala Tyr Thr Glu Ile
 50 55 60
 Asp Trp Lys Ala Tyr Met Ala Glu Val Glu Gly Val Ile Asn Gly Thr
 65 70 75 80
 Tyr Asp Tyr Thr Gln Leu Gln Gly Asp Thr Gly Pro Leu Val Tyr Pro
 85 90 95
 Ala Gly Phe Val Tyr Ile Phe Met Gly Leu Tyr Tyr Ala Thr Ser Arg
 100 105 110
 Gly Thr Asp Ile Arg Met Ala Gln Asn Ile Phe Ala Val Leu Tyr Leu
 115 120 125
 Ala Thr Leu Leu Leu Val Phe Leu Ile Tyr His Gln Thr Cys Lys Val
 130 135 140
 Pro Pro Phe Val Phe Phe Phe Met Cys Cys Ala Ser Tyr Arg Val His
 145 150 155 160
 Ser Ile Phe Val Leu Arg Leu Phe Asn Asp Pro Val Ala Met Val Leu
 165 170 175
 Leu Phe Leu Ser Ile Asn Leu Leu Leu Ala Gln Arg Trp Gly Trp Gly
 180 185 190
 Cys Cys Phe Phe Ser Leu Ala Val Ser Val Lys Met Asn Val Leu Leu
 195 200 205
 Phe Ala Pro Gly Leu Leu Phe Leu Leu Leu Thr Gln Phe Gly Phe Arg
 210 215 220
 Gly Ala Leu Pro Lys Leu Gly Ile Cys Ala Gly Leu Gln Val Val Leu
 225 230 235 240
 Gly Leu Pro Phe Leu Leu Glu Asn Pro Ser Gly Tyr Leu Ser Arg Ser
 245 250 255
 Phe Asp Leu Gly Arg Gln Phe Leu Phe His Trp Thr Val Asn Trp Arg
 260 265 270
 Phe Leu Pro Glu Ala Leu Phe Leu His Arg Ala Phe His Leu Ala Leu
 275 280 285
 Leu Thr Ala His Leu Thr Leu Leu Leu Leu Phe Ala Leu Cys Arg Trp
 290 295 300
 His Arg Thr Gly Glu Ser Ile Leu Ser Leu Leu Arg Asp Pro Ser Lys
 305 310 315 320
 Arg Lys Val Pro Pro Gln Pro Leu Thr Pro Asn Gln Ile Val Ser Thr
 325 330 335

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Leu Phe Thr Ser Asn Phe Ile Gly Ile Cys Phe Ser Arg Ser Leu His
 340 345 350

Tyr Gln Phe Tyr Val Trp Tyr Phe His Thr Leu Pro Tyr Leu Leu Trp
 355 360 365

Ala Met Pro Ala Arg Trp Leu Thr His Leu Leu Arg Leu Leu Val Leu
 370 375 380

Gly Leu Ile Glu Leu Ser Trp Asn Thr Tyr Pro Ser Thr Ser Cys Ser
 385 390 395 400

Ser Ala Ala Leu His Ile Cys His Ala Val Ile Leu Leu Gln Leu Trp
 405 410 415

Leu Gly Pro Gln Pro Phe Pro Lys Ser Thr Gln His Ser Lys Lys Ala
 420 425 430

His

<210> SEQ ID NO 37
 <211> LENGTH: 1291
 <212> TYPE: DNA
 <213> ORGANISM: *Aspergillus nidulans*

<400> SEQUENCE: 37

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aagcttcaag ctgtaaggat ttcggcacgg ctacggaaga cggagaagcc caccttcagt    60
ggactcgagt accatttaat tctatttggt ttgatcgag acctaataca gccctacaa    120
cgaccatcaa agtcgtatag ctaccagtga ggaagtggac tcaaatcgac ttcagcaaca    180
tctcttggt aaactttaag cctaaactat acagaataag atggtggaga gcttataaccg    240
agctcccaaa tctgtccaga tcatggttga ccggtgcctg gatcttcccta tagaatcatc    300
cttattcggt gacctagctg attctggagt gaccagagg gtcagtactt gagcctaaaa    360
tccgcccgt ccaccatttg tagaaaaatg tgacgaactc gtgagctctg tacagtgacc    420
ggtgactctt tctggcatgc ggagagacgg acggacgcag agagaagggc tgagtaataa    480
gcgccactgc gccagacagc tctggcggct ctgaggtgca gtggatgatt attaaccgg    540
gaccggccgc cctccgccc cgaagtggaa aggctgggtg gcccctcggt gaccaagaat    600
ctattgcac atcgagaat atggagcttc atcgaatcac cggcagtaag cgaaggagaa    660
tgtgaagcca ggggtgtata gccgtcggcg aaatagcatg ccattaacct aggtacagaa    720
gtccaattgc ttccgatctg gtaaaagatt cacgagatag taccttctcc gaagtaggta    780
gagcgagtac ccggcgcgta agctccctaa ttggcccatc cggcatctgt agggcggtcca    840
aatatcgtgc ctctcctgct ttgcccggtg tatgaaaccg gaaaggccgc tcaggagctg    900
gccagcggcg cagaccggga acacaagctg gcagtcgacc catccggtgc tctgcactcg    960
acctgctgag gtccctcagt cctggttagg cagctttgcc ccgtctgtcc gcccggtgtg   1020
tcggcggggt tgacaaggtc gttgcgtcag tccaacattt gttgccatat tttcctgctc   1080
tccccaccag ctgctctttt cttttctctt tcttttccca tcttcagtat attcatcttc   1140
ccatccaaga acctttatct cccctaagta agtactttgc tacatccata ctccatcctt   1200
cccatccctt attcctttga acctttcagt tcgagcttcc ccacttcac gcagcttgac   1260
taacagctac cccgcttgag cagacatcac a                                     1291

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<210> SEQ ID NO 38
 <211> LENGTH: 30
 <212> TYPE: DNA
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:

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<223> OTHER INFORMATION: gpdA5F primer

<400> SEQUENCE: 38

cgcagatctc aagctgtaag gatttcggca 30

<210> SEQ ID NO 39

<211> LENGTH: 40

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: gpdA3R primer

<400> SEQUENCE: 39

caccgggccc atctcaaaca ttgtgatgtc tgctcaagcg 40

<210> SEQ ID NO 40

<211> LENGTH: 1255

<212> TYPE: DNA

<213> ORGANISM: Aspergillus nidulans

<220> FEATURE:

<221> NAME/KEY: CDS

<222> LOCATION: (1)..(236)

<220> FEATURE:

<221> NAME/KEY: Intron

<222> LOCATION: (237)..(366)

<220> FEATURE:

<221> NAME/KEY: CDS

<222> LOCATION: (367)..(1252)

<400> SEQUENCE: 40

atg ttt gag atg ggc cgg gtg gga act cgt ctc ccc gcc atg acc tct 48
Met Phe Glu Met Gly Pro Val Gly Thr Arg Leu Pro Ala Met Thr Ser
1 5 10 15

cca gcg cac aac cac tac agc tac cac tct ccc acc tcc agc gac aga 96
Pro Ala His Asn His Tyr Ser Tyr His Ser Pro Thr Ser Ser Asp Arg
20 25 30

ggc cgg tca agg cag aac tcg gat gcc atg gac atc cag tcc atc act 144
Gly Arg Ser Arg Gln Asn Ser Asp Ala Met Asp Ile Gln Ser Ile Thr
35 40 45

gaa cga gag ccg gcg acc aga tac gcg gtt gcg ggc ggc cct gcg ccc 192
Glu Arg Glu Pro Ala Thr Arg Tyr Ala Val Ala Gly Gly Pro Ala Pro
50 55 60

tgg aat cgc aac ggg tct ccg agc atg agc cct atg tat agc aa 236
Trp Asn Arg Asn Gly Ser Pro Ser Met Ser Pro Met Tyr Ser Asn
65 70 75

gtacatctct cttacccctc cgttttcttc tgcttttcta ccaccccatc cctctttcca 296

gtctgagtcc aggcttggtc cgcttgaagt ggctaagtgt atcctcgtct tctctctttc 356

tgtgttttag c aat tcc gag cga aac cag ttt cat gaa gag aac gga cgc 406
Asn Ser Glu Arg Asn Gln Phe His Glu Glu Asn Gly Arg
80 85 90

acc tac cat ggc ttt cgc agg gga atg tat ttt ctt ccg tgc gat gag 454
Thr Tyr His Gly Phe Arg Arg Gly Met Tyr Phe Leu Pro Cys Asp Glu
95 100 105

caa gaa cag gat cgc ctc gac atc ttc cat aag cta ttc acg gta gcg 502
Gln Glu Gln Asp Arg Leu Asp Ile Phe His Lys Leu Phe Thr Val Ala
110 115 120

cgg gta tcg gag agt ctg atc tac gcg ccc cat cca acc aac ggc cgg 550
Arg Val Ser Glu Ser Leu Ile Tyr Ala Pro His Pro Thr Asn Gly Arg
125 130 135 140

ttt ctg gac cta gga tgt gga act ggt atc tgg gcg atc gag gta gcg 598
Phe Leu Asp Leu Gly Cys Gly Thr Gly Ile Trp Ala Ile Glu Val Ala
145 150 155

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aac aag tac cct gat gcg ttt gtc gct ggt gtg gat ttg gct cct att	646
Asn Lys Tyr Pro Asp Ala Phe Val Ala Gly Val Asp Leu Ala Pro Ile	
160 165 170	
cag cct ccg aac cac ccg aag aac tgc gag ttc tac gcg ccc ttc gac	694
Gln Pro Pro Asn His Pro Lys Asn Cys Glu Phe Tyr Ala Pro Phe Asp	
175 180 185	
ttc gaa gcg cca tgg gcc atg ggg gag gat tcc tgg gat cta atc cat	742
Phe Glu Ala Pro Trp Ala Met Gly Glu Asp Ser Trp Asp Leu Ile His	
190 195 200	
ctg cag atg ggt tgc ggt agt gtc atg ggc tgg cca aac ttg tat cga	790
Leu Gln Met Gly Cys Gly Ser Val Met Gly Trp Pro Asn Leu Tyr Arg	
205 210 215 220	
agg ata ttc gca cat ctc cgt ccc ggt gcc tgg ttt gag cag gtt gag	838
Arg Ile Phe Ala His Leu Arg Pro Gly Ala Trp Phe Glu Gln Val Glu	
225 230 235	
atc gat ttc gag cct cga tgt gat gat ccg tca cta gat gga acg gca	886
Ile Asp Phe Glu Pro Arg Cys Asp Arg Ser Leu Asp Gly Thr Ala	
240 245 250	
ttg cgg cat tgg tac gat tgt ctt aaa cag gcg aca gca gag acc atg	934
Leu Arg His Trp Tyr Asp Cys Leu Lys Gln Ala Thr Ala Glu Thr Met	
255 260 265	
ccg cca atc gcc cat agc tcc cgc gat aca ata aaa gac ctg cag gac	982
Arg Pro Ile Ala His Ser Ser Arg Asp Thr Ile Lys Asp Leu Gln Asp	
270 275 280	
gct ggg ttc acg gag atc gac cat caa ata gtg gga ctc ccg ctc aac	1030
Ala Gly Phe Thr Glu Ile Asp His Gln Ile Val Gly Leu Pro Leu Asn	
285 290 295 300	
ccg tgg cat cag gac gaa cac gag ccg aag gtg gca cgt tgg tat aac	1078
Pro Trp His Gln Asp Glu His Glu Arg Lys Val Ala Arg Trp Tyr Asn	
305 310 315	
ctg gcc gtc tca gag agc atc gaa aac ctc agt ctg gct ccc ttc agt	1126
Leu Ala Val Ser Glu Ser Ile Glu Asn Leu Ser Leu Ala Pro Phe Ser	
320 325 330	
cgt gtc tat cgc tgg ccc ctg gag aga atc cag caa ctc gcc gca gat	1174
Arg Val Tyr Arg Trp Pro Leu Glu Arg Ile Gln Gln Leu Ala Ala Asp	
335 340 345	
gtg aag tcc gaa gca ttc aac aaa gag atc cat gcc tac aat ata ctg	1222
Val Lys Ser Glu Ala Phe Asn Lys Glu Ile His Ala Tyr Asn Ile Leu	
350 355 360	
cac ata tac cag gct agg aaa cca tta aga taa	1255
His Ile Tyr Gln Ala Arg Lys Pro Leu Arg	
365 370	

<210> SEQ ID NO 41

<211> LENGTH: 374

<212> TYPE: PRT

<213> ORGANISM: Aspergillus nidulans

<400> SEQUENCE: 41

Met Phe Glu Met Gly Pro Val Gly Thr Arg Leu Pro Ala Met Thr Ser
1 5 10 15

Pro Ala His Asn His Tyr Ser Tyr His Ser Pro Thr Ser Ser Asp Arg
20 25 30

Gly Arg Ser Arg Gln Asn Ser Asp Ala Met Asp Ile Gln Ser Ile Thr
35 40 45

Glu Arg Glu Pro Ala Thr Arg Tyr Ala Val Ala Gly Gly Pro Ala Pro
50 55 60

Trp Asn Arg Asn Gly Ser Pro Ser Met Ser Pro Met Tyr Ser Asn Asn
65 70 75 80

Ser	Glu	Arg	Asn	Gln	Phe	His	Glu	Asn	Gly	Arg	Thr	Tyr	His	Gly	
				85					90					95	
Phe	Arg	Arg	Gly	Met	Tyr	Phe	Leu	Pro	Cys	Asp	Glu	Gln	Glu	Asp	
				100					105					110	
Arg	Leu	Asp	Ile	Phe	His	Lys	Leu	Phe	Thr	Val	Ala	Arg	Val	Glu	
				115					120					125	
Ser	Leu	Ile	Tyr	Ala	Pro	His	Pro	Thr	Asn	Gly	Arg	Phe	Leu	Asp	
				130					135					140	Leu
Gly	Cys	Gly	Thr	Gly	Ile	Trp	Ala	Ile	Glu	Val	Ala	Asn	Lys	Tyr	
				145					150					155	Pro
Asp	Ala	Phe	Val	Ala	Gly	Val	Asp	Leu	Ala	Pro	Ile	Gln	Pro	Asn	
				165					170					175	
His	Pro	Lys	Asn	Cys	Glu	Phe	Tyr	Ala	Pro	Phe	Asp	Phe	Glu	Ala	
				180					185					190	Pro
Trp	Ala	Met	Gly	Glu	Asp	Ser	Trp	Asp	Leu	Ile	His	Leu	Gln	Met	
				195					200					205	Gly
Cys	Gly	Ser	Val	Met	Gly	Trp	Pro	Asn	Leu	Tyr	Arg	Arg	Ile	Phe	
				210					215					220	Ala
His	Leu	Arg	Pro	Gly	Ala	Trp	Phe	Glu	Gln	Val	Glu	Ile	Asp	Phe	
				225					230					235	Glu
Pro	Arg	Cys	Asp	Asp	Arg	Ser	Leu	Asp	Gly	Thr	Ala	Leu	Arg	His	
				245					250					255	Trp
Tyr	Asp	Cys	Leu	Lys	Gln	Ala	Thr	Ala	Glu	Thr	Met	Arg	Pro	Ile	
				260					265					270	Ala
His	Ser	Ser	Arg	Asp	Thr	Ile	Lys	Asp	Leu	Gln	Asp	Ala	Gly	Phe	
				275					280					285	Thr
Glu	Ile	Asp	His	Gln	Ile	Val	Gly	Leu	Pro	Leu	Asn	Pro	Trp	His	
				290					295					300	Gln
Asp	Glu	His	Glu	Arg	Lys	Val	Ala	Arg	Trp	Tyr	Asn	Leu	Ala	Val	
				305					310					315	Ser
Glu	Ser	Ile	Glu	Asn	Leu	Ser	Leu	Ala	Pro	Phe	Ser	Arg	Val	Tyr	
				325					330					335	Arg
Trp	Pro	Leu	Glu	Arg	Ile	Gln	Gln	Leu	Ala	Ala	Asp	Val	Lys	Ser	
				340					345					350	Glu
Ala	Phe	Asn	Lys	Glu	Ile	His	Ala	Tyr	Asn	Ile	Leu	His	Ile	Tyr	
				355					360					365	Gln
Ala	Arg	Lys	Pro	Leu	Arg										
				370											

<400> SEQUENCE: 42

cqcttgaqca qacatcacaa tqtttgaqat qqqcccqqtg

40

<400> SEQUENCE: 43

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cgcagatctg aggattatga gaaggagc

29

<210> SEQ ID NO 44

<211> LENGTH: 2866

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: pGPDA promoter LeaA fragment

<400> SEQUENCE: 44

aagcttcaag ctgtaaggat ttccggcacgg ctacggaaga cggagaagcc caccttcagt 60
ggactcgagt accatttaaat tctattttgtg ttgatcgag acctaataca gccctacaa 120
cgaccatcaa agtcgtatag ctaccagtga ggaagtggac tcaaatcgac ttcagcaaca 180
tctcctggat aaactttaag cctaaactat acagaataag atggtggaga gcttataccg 240
agctcccaaa tctgtccaga tcatggttga ccggtgcctg gatcttcta tagaatcatc 300
cttattcggt gacctagctg attctggagt gaccagagg gtcatgactt gagcctaaaa 360
tccgcccgtt ccaccatttg tagaaaaatg tgacgaactc gtgagctctg tacagtgacc 420
ggtgactctt tctggcatgc ggagagacgg acggacgcag agagaagggc tgagtaataa 480
gcgccactgc gccagacagc tctggcggct ctgaggtgca gtggatgatt attaaccgg 540
gaccggccgc cctccgcgc cgaagtggaa aggctggtgt gccctcgtt gaccaagaat 600
ctattgcac atcggagaat atggagcttc atcgaatcac cggcagtaag cgaaggagaa 660
tgtgaagcca ggggtgtata gccgtcggcg aaatagcatg ccattaacct aggtacagaa 720
gtccaattgc ttccgatctg gtaaaagatt cacgagatag taccttctcc gaagtaggta 780
gagcgagtac ccggcgcgta agctccctaa ttggcccatc cggcatctgt agggcgtcca 840
aatatcgtgc ctctcctgct ttgcccggtg tatgaaacgg gaaaggccgc tcaggagctg 900
gccagcggcg cagaccggga acacaagctg gcagtcgacc catccggtgc tctgcaactc 960
acctgctgag gtcctcagct cctcggtagg cagctttgcc ccgtctgtcc gcccggtgtg 1020
tcggcggggt tgacaaggtc gttgcgtcag tccaacattt gttgccatat ttctctgctc 1080
tccccaccag ctgctctttt cttttctctt tcttttccca tcttcagtat attcatcttc 1140
ccatccaaga acctttatth ccctaagta agtactttgc tacatccata ctccatcctt 1200
cccatccctt attcctttga acctttcagt tcgagcttcc ccacttcac gcagcttgac 1260
taacagctac ccgcttgag cagacatcac aatgtttgag atgggcccgg tgggaactcg 1320
tctccccgcc atgacctctc cagcgacaaa ccactacagc taccactctc ccacctccag 1380
cgacagaggg cggtaaggc agaactcgga tgccatggac atccagcca tcaactgaac 1440
agagccggcg accagatagc cggttgcggg cggccctgcg cctggaatc gcaacgggtc 1500
tccgagcatg agccctatgt atagcaagta catctctctt acccctccgt ttctttctgc 1560
ttttctacca ccccatccct ctttccagtc tgagtcagg cttgttccgc ttgaagtggc 1620
taatgtgatc ctgctcttct ctctttctgt gttttagcaa ttccgagcga aaccagtttc 1680
atgaagagaa cggacgcacc taccatggct ttccgagggg aatgtattht cttccgtgcg 1740
atgagcaaga acaggatgc ctgcacatct tccataagct attcacgga gcgcggttat 1800
cggagagtct gatctacgc ccccatccaa ccaacggcgg gtttctggac ctaggatgtg 1860
gaactggtat ctggcgatc gaggtagcga acaagtacc tgatgcgttt gtcgctggtg 1920
tggaatttgc tcctattcag cctccgaacc acccgaagaa ctgcgagttc tacgcgccct 1980
tcgacttcga agcgccatgg gccatggggg aggattcctg ggatctaact catctgcaga 2040

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tgggttcg	tagtgtcatg	ggctggccaa	acttgtatcg	aaggatatcc	gcacatctcc	2100
gtcccgggtg	ctggtttgag	cagggtgaga	tcgatttcga	gcctcgatgt	gatgatcggt	2160
cactagatgg	aacggcattg	cggcattggg	acgattgtct	taaacaggcg	acagcagaga	2220
ccatgcggcc	aatcgcccat	agctcccgcg	atacaataaa	agacctgcag	gacgtgggt	2280
tcacggagat	cgaccatcaa	atagtgggac	tcccgcctca	cccgtggcat	caggacgaac	2340
acgagcggaa	gggtggcact	tggtataacc	tgcccgctct	agagagcatc	gaaaacctca	2400
gtctggctcc	cttcagtcgt	gtctatcgct	ggccctcgga	gagaatccag	caactcgccg	2460
cagatgtgaa	gtccgaagca	ttcaacaaag	agatccatgc	ctacaatata	ctgcacatat	2520
accaggctag	gaaaccatta	agataagagc	aaaaggcgac	cacatccagg	aacgcaaaac	2580
gaaaaggagg	aaaactgcta	gcgcaagttt	atgtcacgct	ggcacacgcc	cagccatcag	2640
aaatctcaac	agcgaaagtt	atgaaccgca	tcaaccgagt	atgaacgaca	attcgctccat	2700
cacacacctc	tcggttctct	tcgcaggccc	agcatggcgc	cctatcaacc	tgctttacga	2760
cgctgtatat	actggcgaag	tatctctctc	atctactctg	gcgctctaga	taccgtgaag	2820
atgcagacaa	aattggccga	gctcccttct	cataatcctc	aagctt		2866

<210> SEQ ID NO 45

<211> LENGTH: 1279

<212> TYPE: DNA

<213> ORGANISM: *Aspergillus niger*

<400> SEQUENCE: 45

atgcatcatt	ctcccgtttt	gtttttgggc	ccaaactaac	cgagtaggtg	tggcatttgc	60
gggcatgatg	tttcaactac	cgctgatcat	tatcacccgc	ccattagaga	agatccaaga	120
ccctactggg	aagggtgatg	gcaattccat	tttctgggtt	agtttttgct	ttgtcggcca	180
gcctttggga	gctttgtctg	acttctttgc	ctggcaagcg	aagtatggca	gtgtgagccg	240
aatgtgaatg	taaaagcacg	cacgtgtccg	ctgtttgtca	tagatgtaaa	taaatgccaa	300
caacttcagc	cattttttga	aaagcaaaag	aaccgaagta	aacgatcctg	taccatcagc	360
gctcttcaca	atggaatctt	ttagatgttt	ctgttccatt	catcttgctt	actgcaatgt	420
tcttttcgcg	tttgactaat	tctccggatg	ttgaatggca	acgctgtcgg	cgctgggtct	480
tcagggatcc	gccaaaggatg	ctctggatcc	gcacccggcc	gctcttgccg	cccatcaatc	540
gcccactat	aaatcgaaact	actttcggca	tcttctagac	ttcctaatac	cgcttagtca	600
tagcagattc	aagctgagaa	caccacaagt	aaatatcacc	catcatgctt	accctgaccg	660
tccctgaaaa	ctacgggtat	gtgccaatc	tacaattcct	tgcagacaa	gccattctcc	720
ccatgaagtc	tgatgctaac	tatctgcag	ctctgtcatt	gccgtcgctc	tgggtgccat	780
ccccgtcctg	agcttcgtcc	atggcgccgt	cgtgtctcgt	ctccgcaagg	aagctgattg	840
cccctaccct	cactgctatg	cgaccgtaga	gcagtgcag	accaacgtaa	gccaacctca	900
cacaaacagg	attctctgag	ctaacataca	ttccgaaccg	tgcagcccaa	ggcgagcag	960
ttcaactcgg	ctcagcgcg	tcatgccaac	ttccttgaga	actccagcca	aactatgctc	1020
ttcctcctgg	tagctggact	gaagtacccc	cagttggcga	ctggcctcgg	aagcatctgg	1080
gtcctcggtc	gctcactgtt	cctttacgga	tatgtgtact	cggcaagcc	gcggggtcgc	1140
ggctggtttg	acggcagctt	ctacttgctt	gcacagggag	ctctctgggg	cttgacgtct	1200
tttgagttg	cgaggaggtt	gatttcctac	ttctaagttt	ggactgaatc	cgtgggtgtga	1260

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 ttgaggtgat tggcgatgt 1279

<210> SEQ ID NO 46
 <211> LENGTH: 472
 <212> TYPE: DNA
 <213> ORGANISM: *Aspergillus nidulans*
 <220> FEATURE:
 <221> NAME/KEY: misc.feature
 <222> LOCATION: (361)..(361)
 <223> OTHER INFORMATION: n is a, c, g, or t

<400> SEQUENCE: 46

ggaccgatgg ctgtgtagaa gtactcgccg atagtggaaa ccgacgcccc agcactcgtc	60
cgagggcaaa ggaatagagt agatgccgac cgcgggatcc acttaacgtt actgaaatca	120
tcaaacagct tgacgaatct ggatataaga tcgttggtgt cgatgtcagc tccggagttg	180
agacaaatgg tgttcaggat ctcgataaga tacgttcatt tgtccaagca gcaaagagt	240
ccttctagtgt atttaatagc tccatgtcaa caagaataaa acgcgttttc gggtttacct	300
cttcagata cagctcatct gcaatgcatt aatgcattga ctgcaaccta gtaacgectt	360
ncaggctccg gcgaagagaa gaatagctta gcagagctat tttcattttc gggagacgag	420
atcaagcaga tcaacgggtcg tcaagagacc tacgagactg aggaatccgc tc	472

<210> SEQ ID NO 47
 <211> LENGTH: 2005
 <212> TYPE: DNA
 <213> ORGANISM: *Aspergillus oryzae*

<400> SEQUENCE: 47

tctgacagac gggcaattga ttacgggac ccatggtaa cgaaatgtaa aagctaggag	60
atcgctccgc gatgtcagga tgatttcaact tgtttcttgt ccggctcacc ggtcaaagct	120
aaagaggagc aaaaggaacg gatagaatcg ggtgccgctg atctatacgg tatagtgcc	180
ttatcacgtt gactcaaccc atgctattta actcaacccc tccttctgaa ccccaccatc	240
ttcttctctt tcctctcatc ccacacaatt ctctatctca gatttgaatt ccaaaagtcc	300
tcggacgaaa ctgaacaagt ctctctccct tcgataaacc tttggtgatt ggaataactg	360
accatcttct atagtccca aaccaaccga caatgtaaat acactcctcg attagccctc	420
tagagggcat acgatggaag tcattggaata cttttggctg gactctcaca atgatcaagg	480
tatcttaggt aacgtctttg gcgtgggccc gtgttcgttc ccagtcacg atgcattcac	540
atgcctctcc taagctgggc cctagactct aggatcctag tctagaagga catggcatcg	600
atggactggg ttcgttctga gattatacgg ctaaaacttg atctggataa taccagcgaa	660
aagggtcatg ccttctctcg ttcttctgt tgatggaatg gctaacagat gatagtcatt	720
gcaacttgaa acatgtctcc tccagctgcc atctacgaac ccactgtggc cgctaccggc	780
ctcaagggtg aggtcgtggt ttctgagacc gtccccgttg agggagcttc tcagaccaag	840
ctgttggaac atttcggtgg caagtgggac gaggttcaagt tcgcccctat ccgcgaaagc	900
caggtctctc gtgccatgac cagacgttac tttgaggacc tggacaagta cgctgaaagt	960
gacgttgtea ttgttggtgc tggttcctgc ggtctgagca ctgcgtacgt cttggccaag	1020
gctcgtccgg acctgaagat tgctatcgtc gaggccacgg tctctcctgg tcagtagtcc	1080
atgatggatt gccttgcaact cagctttccg gaactaacgt gcaatagggt gcggtgctg	1140
gttggtggc caactctttt ctgctatggt catgcgccgt cccgcggaag tcttctgaa	1200
cgagctgggt gttccttacg aagaggacgc aaacccaac tacgttgtcg tcaagcacgc	1260

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ctccctgttt acctcgacac tcattgtgaa ggttctctcc tcccccaatg tcaagctctt 1320
caatgctacc gctgttgagg acttgatcac ccgtccgacc gagaacggca acccccagat 1380
tgctgggtgt gtcgtcaact ggacgctggt cacccttcac cagcatgac actcctgcat 1440
ggacccaac actatcaacg ctctgtcat catcagtacc actggtcacg atgggccatt 1500
cggcgccctc tgtgcgaagc gcttggtgtc catgggcagc gtcgacaagc taggtggcat 1560
gcgtgggtct gacatgaact cggccgagga tgccatcgtc aagaacaccc gcgaggttac 1620
taagggttg ataatcgagg gtatggagct gtctgaaatt gatggcttta accgcatggg 1680
ccctaccttc ggtgccatgg ttctcagtgg tgtcaaggct gccgaggagg cattgaagg 1740
gttcgacgag cgtcagcgcg agtgtgctga gttaaactgact cactaccga atgggttcag 1800
tgcatgaacc ggatttgtct tacggtcttt gacgataggg gaatgatgat tatgtgatag 1860
ttctgagatt tgaatgaact cgtagctcg taatccacat gcatatgtaa atggctgtgt 1920
cccgtatgta acggtggggc attctagaat aattatgtgt aacaagaaag acagtataat 1980
acaaacaaag atgcaagagc ggctc 2005

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<210> SEQ ID NO 48
<211> LENGTH: 1052
<212> TYPE: DNA
<213> ORGANISM: Aspergillus niger

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<400> SEQUENCE: 48

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gtcatcgag ataagcactg ctgtcttgca tccaagtcag cgtcagcaga aatacgggac 60
ttccgaaagt atatggcaaa attaaagaac ttgactctcc agcaatgttt tgccctgacc 120
gtcgctaaaa cgttactacc cctatacccg tctgtttgtc ccagcccgag gcattaggtc 180
tgactgacag cacggcgcca tgcgggcttg ggacgccatg tccgtcgcgt gataagggtt 240
gatccatgca gctactatcc ttccatcgtt ccattcccat ccttgctcta tctccatcct 300
tgaaacttta ctagttagt tgtagctcg agcttgctct cggctactcc gtccaatgga 360
taagaccccg atgcgggtcc tcattggtct ccagctggtg tgcaccaac cttcgtgtga 420
tcgctctctc gcttccctc atcatcatta ctaactagta catccaaaag ccatcccagt 480
gcttccctc acccttgccc aagacattcc aagtgggcct tcggtggaa aacatggacc 540
cattggttcc atcgataagc tagctcctcg tccgttacc cagattgata ccagataaca 600
ttgaccagcg gcttatcacc gaggtctgcg ggtgagaccc cccctgcgac aagttagata 660
aaagaaactc gcctcattgt gcttccgatg gggtcggatg acgagccttc ggaaagagct 720
ggcgccctct taaaggggac agctgtcgcc aagttgtgaa attctccgat aactactaac 780
aatctctccc ttcttcccg ctactgttgt caccaaatca actctctttt ctggccaag 840
atctaactg gcggatgaga agactgaaaa gtctcccca ccatgacgg tggtatgagga 900
gactggcaca acagagggaa ttgaccgac aatggcaaag catacgaagg atgcagacga 960
ggcactggcg gtcttcgaag acctccatgg tgaagtcac acacttgatg aggagacaaa 1020
caaaaggata cttcgacaaa ttgactggca ca 1052

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<210> SEQ ID NO 49
<211> LENGTH: 58
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: pyrGU5F primer

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<400> SEQUENCE: 49

gtaacgccag gggttttccca gtcacgacgt ttaaacaatgc atcattctcc cgctttgt 58

<210> SEQ ID NO 50

<211> LENGTH: 49

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: pyrGU3R primer

<400> SEQUENCE: 50

tgccgaaatc cttacagctt gaagcttcat cgccaatcac ctcaatcac 49

<210> SEQ ID NO 51

<211> LENGTH: 49

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: Trp5F primer

<400> SEQUENCE: 51

agctcccttc tcataatcct caagcttgga ccgatggctg tgtagaagt 49

<210> SEQ ID NO 52

<211> LENGTH: 46

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: Trp3R primer

<400> SEQUENCE: 52

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<210> SEQ ID NO 53

<211> LENGTH: 46

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: PTR5F primer

<400> SEQUENCE: 53

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<210> SEQ ID NO 54

<211> LENGTH: 45

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: PTR3R primer

<400> SEQUENCE: 54

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<210> SEQ ID NO 55

<211> LENGTH: 45

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: PyrGD5F primer

<400> SEQUENCE: 55

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<210> SEQ ID NO 56

<211> LENGTH: 44

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<212> TYPE: DNA
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
 <223> OTHER INFORMATION: PyrGD3R primer

<400> SEQUENCE: 56

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<210> SEQ ID NO 57
 <211> LENGTH: 7689
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 <213> ORGANISM: Artificial Sequence
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 <223> OTHER INFORMATION: transgene
 <220> FEATURE:
 <221> NAME/KEY: misc_feature
 <222> LOCATION: (4513)..(4513)
 <223> OTHER INFORMATION: n is a, c, g, or t

<400> SEQUENCE: 57

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<210> SEQ ID NO 58
<211> LENGTH: 1270
<212> TYPE: DNA
<213> ORGANISM: Aspergillus niger
<220> FEATURE:
<221> NAME/KEY: CDS
<222> LOCATION: (1)..(230)
<220> FEATURE:
<221> NAME/KEY: Intron
<222> LOCATION: (231)..(372)
<220> FEATURE:
<221> NAME/KEY: CDS
<222> LOCATION: (373)..(1267)

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<400> SEQUENCE: 58

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cgc aat cgc aat aac tac agc tac caa ggg ata gaa tcc tat gat tcc 96
Pro Asn Arg Asn Asn Tyr Ser Tyr Gln Gly Ile Glu Ser Tyr Asp Ser
20 25 30

ggc cgt tcc agg caa aac tcg gat gct atg gac att cac gtc att acg 144
Gly Arg Ser Arg Gln Asn Ser Asp Ala Met Asp Ile His Val Ile Thr
35 40 45

gcc caa gaa cct cct cga gaa ccc ccg gac aac aac gat cct tat gat 192
Ala Gln Glu Pro Pro Arg Glu Pro Pro Asp Asn Asn Asp Pro Tyr Asp
50 55 60

ggc cat ggg ggt cca gct ggg act agc cat tat agc aa gtacttctcc 240
Gly His Gly Gly Pro Ala Gly Thr Ser His Tyr Ser Lys

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cgtttaatct ag g cct cca aac aga tgg ctc ttc tat gaa gaa aat ggg			409
	Pro Pro Asn Arg Trp Leu Phe Tyr Glu Glu Asn Gly		
	80 85		
cga aca tat cat gga tat cgc aga gga gtt tac ccg ctg cca tgc gat			457
Arg Thr Tyr His Gly Tyr Arg Arg Gly Val Tyr Pro Leu Pro Cys Asp			
90 95 100 105			
gaa cag gaa cag gac cgt ctc gat atc ttc cat aaa ctg ttc aca gta			505
Glu Gln Glu Gln Asp Arg Leu Asp Ile Phe His Lys Leu Phe Thr Val			
110 115 120			
gca cgg atg tcc gag agc tta atc tac gca cct cac ccc cca aat ggt			553
Ala Arg Met Ser Glu Ser Leu Ile Tyr Ala Pro His Pro Pro Asn Gly			
125 130 135			
cga ttc cta gat ctg ggg tgc ggc act ggg atc tgg gcc att gat gta			601
Arg Phe Leu Asp Leu Gly Cys Gly Thr Gly Ile Trp Ala Ile Asp Val			
140 145 150			
gcc cac aag tat ccc aat gct ttc gtt gct gga gta gat cta gca cct			649
Ala His Lys Tyr Pro Asn Ala Phe Val Ala Gly Val Asp Leu Ala Pro			
155 160 165			
ata cag cct ccc aac cac ccc gat aac tgc gag ttc tat gca cct ttt			697
Ile Gln Pro Pro Asn His Pro Asp Asn Cys Glu Phe Tyr Ala Pro Phe			
170 175 180 185			
gac ttt gag gcg cca tgg acg ctt ggg gaa aat tct tgg gat ctc att			745
Asp Phe Glu Ala Pro Trp Thr Leu Gly Glu Asn Ser Trp Asp Leu Ile			
190 195 200			
cat cta cag atg ggt tgc ggc agt gtt ctg ggc tgg cag aat ctc tac			793
His Leu Gln Met Gly Cys Gly Ser Val Leu Gly Trp Gln Asn Leu Tyr			
205 210 215			
aag cga atc tta agg cat ctt cag cct ggg gca tgg ttt gaa cag gtg			841
Lys Arg Ile Leu Arg His Leu Gln Pro Gly Ala Trp Phe Glu Gln Val			
220 225 230			
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Glu Ile Asp Phe Glu Pro Arg Cys Asp Asp Arg Ser Leu Asn Gly Leu			
235 240 245			
gca ctc cgg gag tgg tac cag tac ctg aag cag gcg aca caa gat aca			937
Ala Leu Arg Glu Trp Tyr Gln Tyr Leu Lys Gln Ala Thr Gln Asp Thr			
250 255 260 265			
atg cga ccc ata gcg cac agc tcg cgg gat acc atc aga cac ctt gag			985
Met Arg Pro Ile Ala His Ser Ser Arg Asp Thr Ile Arg His Leu Glu			
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gag gca ggc ttt acc cag atc gac cat cag atg gtg ggg ctg cct ctc			1033
Glu Ala Gly Phe Thr Gln Ile Asp His Gln Met Val Gly Leu Pro Leu			
285 290 295			
aac cct tgg cac cgt gat gaa cat gag cag aag gta gcc cgt tgg tat			1081
Asn Pro Trp His Arg Asp Glu His Glu Gln Lys Val Ala Arg Trp Tyr			
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Asn Leu Ala Ile Ser Glu Ser Ile Glu Thr Leu Ser Leu Ala Pro Phe			
315 320 325			
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Ser Arg Ile Phe His Trp Asp Leu Asp Arg Ile Arg Gln Ile Thr Ala			
330 335 340 345			
gag gtc aag tca caa gcc ttc aac aag gaa atc cac gct tac aat atc			1225
Glu Val Lys Ser Gln Ala Phe Asn Lys Glu Ile His Ala Tyr Asn Ile			
350 355 360			
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Leu His Ile Tyr Gln Ala Arg Lys Pro Gly Gly Pro Ser Leu
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<210> SEQ ID NO 59
 <211> LENGTH: 375
 <212> TYPE: PRT
 <213> ORGANISM: *Aspergillus niger*

<400> SEQUENCE: 59

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 20 25 30

Gly Arg Ser Arg Gln Asn Ser Asp Ala Met Asp Ile His Val Ile Thr
 35 40 45

Ala Gln Glu Pro Pro Arg Glu Pro Pro Asp Asn Asn Asp Pro Tyr Asp
 50 55 60

Gly His Gly Gly Pro Ala Gly Thr Ser His Tyr Ser Lys Pro Pro Asn
 65 70 75 80

Arg Trp Leu Phe Tyr Glu Glu Asn Gly Arg Thr Tyr His Gly Tyr Arg
 85 90 95

Arg Gly Val Tyr Pro Leu Pro Cys Asp Glu Gln Glu Gln Asp Arg Leu
 100 105 110

Asp Ile Phe His Lys Leu Phe Thr Val Ala Arg Met Ser Glu Ser Leu
 115 120 125

Ile Tyr Ala Pro His Pro Pro Asn Gly Arg Phe Leu Asp Leu Gly Cys
 130 135 140

Gly Thr Gly Ile Trp Ala Ile Asp Val Ala His Lys Tyr Pro Asn Ala
 145 150 155 160

Phe Val Ala Gly Val Asp Leu Ala Pro Ile Gln Pro Pro Asn His Pro
 165 170 175

Asp Asn Cys Glu Phe Tyr Ala Pro Phe Asp Phe Glu Ala Pro Trp Thr
 180 185 190

Leu Gly Glu Asn Ser Trp Asp Leu Ile His Leu Gln Met Gly Cys Gly
 195 200 205

Ser Val Leu Gly Trp Gln Asn Leu Tyr Lys Arg Ile Leu Arg His Leu
 210 215 220

Gln Pro Gly Ala Trp Phe Glu Gln Val Glu Ile Asp Phe Glu Pro Arg
 225 230 235 240

Cys Asp Asp Arg Ser Leu Asn Gly Leu Ala Leu Arg Glu Trp Tyr Gln
 245 250 255

Tyr Leu Lys Gln Ala Thr Gln Asp Thr Met Arg Pro Ile Ala His Ser
 260 265 270

Ser Arg Asp Thr Ile Arg His Leu Glu Glu Ala Gly Phe Thr Gln Ile
 275 280 285

Asp His Gln Met Val Gly Leu Pro Leu Asn Pro Trp His Arg Asp Glu
 290 295 300

His Glu Gln Lys Val Ala Arg Trp Tyr Asn Leu Ala Ile Ser Glu Ser
 305 310 315 320

Ile Glu Thr Leu Ser Leu Ala Pro Phe Ser Arg Ile Phe His Trp Asp
 325 330 335

Leu Asp Arg Ile Arg Gln Ile Thr Ala Glu Val Lys Ser Gln Ala Phe
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Asn Lys Glu Ile His Ala Tyr Asn Ile Leu His Ile Tyr Gln Ala Arg
 355 360 365

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Lys Pro Gly Gly Pro Ser Leu
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<210> SEQ ID NO 60
<211> LENGTH: 6411
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: transgene to complement alg3delta mutant

<400> SEQUENCE: 60

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aatgccaaca acttcagcca ttttttgaag agcaaagcaa ccgaagtaaa cgatcctgta      360
ccatcagcgc tcctcacaat ggaatctttt agatgtttct gttccattca tcttgcttac      420
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cattctcccc atgaagtctg atgctaacta tcctgcagct ctgtcattgc cgtcgtctctg      780
gggtgccatcc cgtcctgag cttcgtccat ggcgcgctcg tgtctcgtct ccgcaaggaa      840
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ggcagtcgac ccacccggtg ctctgcactc gacctgctga ggteccctcag tccctggtag     1860
gcagctttgc cccgtctgtc cggccgggtg gtcggcgggg ttgacaaggt cgttgcgtca     1920
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ttcttttccc atcttcagta tattcatctt cccatccaag aacctttatt tcccctaagt	2040
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ttcgagcttt cccacttcat cgcagcttga ctaacagcta ccccgcttga gcagacatca	2160
ccatggccaa gttgaccagt gccgttccgg tgctcaccgc gcgcgacgtc gccggagcgg	2220
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cagacagttc atgtttaaat ggacagtcaa ttggagattt gttggcgaag aagtattctt 4500
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gatgcagacg aggcactggc ggtcttcgaa gacctccatg gtgaagtcac cacacttgat 6360
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<210> SEQ ID NO 61
<211> LENGTH: 899
<212> TYPE: DNA
<213> ORGANISM: Aspergillus niger

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<400> SEQUENCE: 61

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tcgatcgtec tcccccgatc catggttggt ctttagtcgc ctctctctct gtaacgcctg	240
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tccactctcc tcttctctcc aatcctttta cccctctctg gccactgatt cattccgcgg	660
aggatacccc cagttagtag ttttgtggtt gccgtctttt cgtctgatca gcttttcaat	720
ccaatcattg gctcaatttg cggcgccgaa ctccaacttc ctctatgtgc cactgactt	780
acgattcccc gatatacct gcagcctgca tacaccctgt tggctaacat tggcgtttta	840
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<210> SEQ ID NO 62
 <211> LENGTH: 1012
 <212> TYPE: DNA
 <213> ORGANISM: *Aspergillus niger*

<400> SEQUENCE: 62

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agccggggcgg cccatcactt tgaagataga gagaaaatta cggcagtgcc ggctgtaatg	180
cagatttcat tccggatacc catatgcagc cttttgtggg agggccatcc tacctgttct	240
cttctttttg ttcattccat tttttctctg atgaggatta gtgacgacca attccatctc	300
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actggacttg aactcgtacc aacactgcta gtttgacgca agaatgccag gaacgcagac	420
aggcttatct tcattgtgtt attgcctact ttagtgcaga cagctgaacg gcacaaaggg	480
acgcgaagta tattatcaat cacaccatgt ggattgttac cgtcgcaaaa gatactgctc	540
gaggttttaa ataatagaag cacttcaaga agatgaggtg ccgttgctcg gactcgaaga	600
agtggccgac gactataaag taccagattt gccaggatcc aaaagaagct accaatctca	660
tgttgcatth ttgaaaagcc ctgtaattca ggcgggtgta actgcggaga tcccagaatc	720
atggcagttg actgagtact tgcgtgacg ggccttaaac atcagccatt ccatccgaac	780
tcaaaactct ctgatgaccg tggcggtctt gtggcacagc caccgataat ttttttgtt	840
ggcacttatt aatgaattag atgagaaaga agcgaagaac cgagggcttg aacaaacgca	900
aagccctcca aatggttact attgaacttt ttgctctccc accgaaactc cactgatca	960
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<210> SEQ ID NO 63
 <211> LENGTH: 1360
 <212> TYPE: DNA
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
 <223> OTHER INFORMATION: Synthetic polynucleotide (hph expression cassette)

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<400> SEQUENCE: 63

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tgtaagcagt agcggcggtg ctgaagtgt gactcttatt agcagacagg aacgaggaca   180
ttattatcat ctgctgcttg gtgcacgata acttggtcgt ttgtcaagca aggtaagtga   240
acgacccggt cataccttct taagttcgcc cttcctccct ttatttcaga ttcaatctga   300
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ttctgatcga aaagttcgac agcgtctccg acctgatgca gctctcggag ggcaagaat   420
ctcgtgcttt cagcttcgat gtaggagggc gtggatatgt cctgcgggta aatagctgcg   480
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cggtcgcgga ggccatggat gcgatcgtg cggcgatct tagccagacg agcgggttcg   720
gcccattcgg accgcaagga atcggtaaat aactacatg gcgtgatttc atatgcgcga   780
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tcattgactg gacgagggcg atgttcgggg attcccaata cgaggtcgcc aacatcttct  1020
tctggaggcc gtggttggtt tgtatggagc agcagacgcg ctacttcgag cggaggcatc  1080
cggagcttgc aggatcgcg cgcctccggg cgtatatgct ccgcattggt cttgaccaac  1140
tctatcagag cttggttgac ggcaatttcg atgatgcagc ttgggcgcag ggtcgatgcg  1200
acgcaatcgt ccgatccgga gccgggactg tcgggcgtac acaaatcgcc cgcagaagcg  1260
cggccgtctg gaccgatggc tgtgtagaag tactcgccga tagtggaac cgacgcccc  1320
gcactcgtcc gagggcaaag gaatagagta gatgccgacc  1360

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<210> SEQ ID NO 64

<211> LENGTH: 2340

<212> TYPE: DNA

<213> ORGANISM: *Aspergillus niger*

<400> SEQUENCE: 64

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ctctccccc cgaagggttc ctgtccttgc ggggaggaga ctttttccc cctacctatt   180
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gtctgatcag cttttcaatc caatcattgg ctcaatttgc ggcgcgaac tccaacttcc   540
tctatgtgcc acctgactta cgattcccc atatcacctg cagcctgeat acaccctgtt   600
ggctaacatt ggcgttttac gttaccgtca cgaagaacgc cggctctctt gattgaacga   660

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accctgtat ctctaccata ccttgagcag gaaaagtcga atctcttccc agcgaaccga	720
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tggtctcgcc gaatcgcaat aactacagct accaagggat agaatcctat gattccggcc	840
gttcacagca aaactcggat gctatggaca ttcacgtcat tacggcccaa gaacctctc	900
gagaaccccc ggacaacaac gatccttatg atggccatgg gggccagct gggactagcc	960
attatagcaa gtacttctcc cttctcatac tctgcacccc acgtacccc caaaatccct	1020
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cagatcgacc atcagatggt ggggctgcct ctcaaccctt ggcaccgtga tgaacatgag	1800
cagaaggtag cccgttggtg taacctcgca atctctgaga gtatcgagac gctcagcctc	1860
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ccatctcctt gacgggatca tactgaaatg cttatacacc aaagcgagca aagcccacaa	2220
aaccatcact ggacttgaac tcgtaccaac actgctagtt tgacgcaaga atgccaggaa	2280
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<210> SEQ ID NO 65

<211> LENGTH: 3306

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: Synthetic polynucleotide (laeA deletion cassette)

<400> SEQUENCE: 65

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acttgcaatt ccgggatcgg attttccctt tctgatccga tgatecagtc caattgacct	180
aactctcgat cgtcctcccc cgatccatgg ttgtttctta gtcgcctctc tcctcgtaac	240
gcctgacacc tgctgctggt atcgataccc tatttattat ttattattta ttttttttcc	300
ctattttcct atgccgctgc cgtgacgct acttttctct ttgtttctct ccccccgac	360

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gggttctctgt ccttgccggg aggagacttt tttcccccta cctatttagt atcctcattc	420
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tgcgcatgca cgggtgtgcg tgcctctttg ctctcataca actctcctgg gttggggcaa	540
atttacacca ttgtggaatt tggatcattc ttacttacta tccttggett ttgtctctcc	600
ggctttccac tctcctcctt ctccaatcc ttttaccctt tctcgccac tgattcattc	660
cgcgaggat acccccagtt agtagttttg tgggtgccgt cttttcgtct gatcagcttt	720
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The invention claimed is:

1. A method of making citric acid, comprising culturing an isolated *Aspergillus* species fungus transformed with at least one *Aspergillus* LaeA (a loss of aflR expression A) gene under conditions that permit the fungus to make citric acid, thereby making citric acid, wherein said LaeA gene encodes a protein having at least 80% sequence identity to the polypeptide of SEQ ID NO: 41 or SEQ ID NO: 59.

2. The method of claim 1, wherein the *Aspergillus* LaeA gene encodes a protein comprising SEQ ID NO: 41 or SEQ ID NO: 59.

3. The method of claim 1, wherein the fungus has been transformed with at least one *Aspergillus* LaeA gene, wherein said LaeA gene has at least 80% sequence identity to nucleotides 1-236 and 367-1252 of SEQ ID NO: 40, or to nucleotides 1-230 and 373-1267 of SEQ ID NO: 58.

4. The method of claim 3, wherein the *Aspergillus* LaeA gene comprises nucleotides 1-236 and 367-1252 of SEQ ID NO: 40, or nucleotides 1-230 and 373-1267 of SEQ ID NO: 58.

5. The method of claim 1, wherein the fungus further comprises a genetic inactivation of an endogenous dolichyl-P-Man:Man(5)GlcNAc(2)-PP-dolichyl mannosyltransferase (Alg3) gene, wherein the Alg3 gene is genetically inactivated by complete or partial deletion mutation or by an insertional mutation, wherein the endogenous Alg3 gene prior to the

genetic inactivation encodes a protein having at least 80% sequence identity to the amino acid sequence of SEQ ID NO: 2 or SEQ ID NO: 4.

6. The method of claim 5, wherein prior to genetic inactivation, the Alg3 gene encodes a protein comprising the amino acid sequence of SEQ ID NO: 2 or SEQ ID NO: 4.

7. The method of claim 1, wherein the fungus further comprises a genetic inactivation of an endogenous Alg3 gene, wherein the Alg3 gene is genetically inactivated by complete or partial deletion mutation or by insertional mutation, wherein the endogenous Alg3 gene prior to the genetic inactivation has at least 80% sequence identity to the nucleotide sequence of SEQ ID NO: 1 or SEQ ID NO: 3.

8. The method of claim 7, wherein prior to genetic inactivation, the Alg3 gene comprises the nucleotide sequence of SEQ ID NO: 1 or SEQ ID NO: 3.

9. The method of claim 1, wherein the *Aspergillus* species is *Aspergillus niger* (*A. niger*).

10. The method of claim 9, wherein the *A. niger* is *A. niger* strain American Type Culture Collection (ATCC) 11414 or *A. niger* strain 11414KusA.

11. The method of claim 1, wherein the fungus is cultured in media comprising at least 50 g/l sugar and 8 to 15 ppb manganese.

12. The method of claim 1, further comprising isolating the citric acid made by culturing the fungus.

* * * * *