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TITLE

Characterisation of the laccase-encoding gene *abr2* of the dihydroxynaphthalene-like melanin gene cluster of *Aspergillus fumigatus*

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Running title: Abr2 laccase of *A. fumigatus*

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ABSTRACT

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24 *Aspergillus fumigatus* is an important pathogen of the immunocompromised host. Previously,
25 it was shown that the polyketide synthase encoded by the *pksP* (*alb1*) gene represents a
26 virulence determinant. *pksP* is part of a gene cluster involved in dihydroxynaphthalene
27 (DHN)-like melanin biosynthesis. Because a putative laccase-encoding gene (*abr2*) is also
28 part of the cluster and a laccase was found to represent a virulence factor in *Cryptococcus*
29 *neoformans*, here, the Abr2 laccase was characterised. Deletion of the *abr2* gene changed the
30 gray-green conidial pigment to a brown color and the ornamentation of conidia was reduced
31 compared with wild-type conidia. In contrast to the white *pksP* mutant, the susceptibility of
32 the Δ *abr2* mutant against reactive oxygen species (ROS) was not increased, suggesting that
33 the intermediate of DHN-like melanin produced up to the step catalysed by Abr2 already
34 possesses ROS scavenging activity. In an intranasal mouse infection model, the Δ *abr2* mutant
35 strain showed no reduction in virulence compared with the wild type. In the Δ *abr2* mutant,
36 overall laccase activity was reduced only during sporulation, but not during vegetative
37 growth. An *abr2p-lacZ* gene fusion was expressed during sporulation, but not during
38 vegetative growth confirming the pattern of laccase activity due to Abr2.

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KEYWORDS:

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42 Dihydroxynaphthalene-like melanin, *Aspergillus fumigatus*, gene cluster, laccase

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INTRODUCTION

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48 The improvement in transplant medicine and the therapy of hematological malignancies is
49 often complicated by the threat of invasive aspergillosis. *Aspergillus fumigatus* accounts for
50 approximately 90% of invasive aspergillosis cases. Specific diagnostics are still limited, as
51 are the possibilities of therapeutic intervention, leading to a high mortality rate of 30 % to 98
52 % for invasive aspergillosis (reviewed in 8, 12, 22). An important question concerning *A.*
53 *fumigatus* is the identification of pathogenicity determinants and their regulation. The group
54 of J. Kwon-Chung and our group had identified a gene that encodes a pathogenicity
55 determinant. It was designated *pksP* (or alternatively *alb1*) for polyketide synthase involved
56 in pigment biosynthesis (14, 15, 19, 20, 38). Conidia of a *pksP* mutant strain are white. Based
57 on genetic and biochemical data the conidial pigment consists of dihydroxynaphthalene
58 (DHN)-like melanin (6, 9, 21, 39, 40).

59 The complete absence of DHN-like melanin, as in the case of *pksP* mutants resulted in a
60 severe reduction in virulence. *PksP* mutant conidia of *A. fumigatus* were significantly more
61 sensitive to hydrogen peroxide and sodium hypochlorite than wild-type conidia. As in other
62 cases, it was shown that melanin-containing conidia are able to quench ROS derived from
63 human granulocytes (14, 15). These results indicated that conidial DHN-like-melanin of *A.*
64 *fumigatus* is involved in protecting conidia from the host immune response in which ROS are
65 important for eliminating fungal conidia (reviewed in 21, 22). However, because *A. nidulans*
66 conidia are also protected by a green pigment, resistance against ROS does not explain why
67 *A. fumigatus* conidia can be pathogenic while this is rarely the case with *A. nidulans* conidia.
68 One attractive hypothesis is that besides the pigment, the *pksP* gene product of *A. fumigatus*
69 is involved in the production of another compound that is immunosuppressive (5). This
70 hypothesis was further supported by the notion that the presence of a functional *pksP* gene in

71 *A. fumigatus* conidia is associated with an inhibition of the fusion of phagosomes and
72 lysosomes in human monocyte-derived macrophages (MDMs) (13, 16, 33). Other pathways
73 involving polyketide synthases have been shown to synthesize two different active products
74 (reviewed in 21).

75 The *pksP* gene is part of a gene cluster which consists of six genes. One of the genes
76 designated *abr2* encodes a putative laccase (39). As mentioned above, our previous data led to
77 the hypothesis that only the *pksP* gene is involved in virulence and not the other genes of the
78 DHN-like melanin pathway. On the other side, a cell wall bound laccase was found to be
79 required for virulence of the human-pathogenic fungus *Cryptococcus neoformans* (27, 28, 30,
80 43, 44). Therefore, we characterised the putative laccase Abr2 of *A. fumigatus* in order to
81 study its impact on virulence.

82

83

MATERIALS AND METHODS

84

Fungal and bacterial strains, media and growth conditions.

85 Fungal strains used in this study are listed in Table 1. *A. fumigatus* ATCC46645 was used to
86 generate an *abr2* knock-out strain. *A. fumigatus* KH Δ *pyrG* is a uracil-auxotrophic mutant of
87 strain ATCC46645. The uracil-auxotrophic strain *A. fumigatus* KH Δ *pyrG* contains a deletion
88 of 144 bp in the 3'-coding region of the *pyrG* gene and therefore, codes for a truncated,
89 nonfunctional orotidine-5'-monophosphate decarboxylase (see below).

90 Strain KH Δ *pyrG* was used to generate strains KH*pksPp-lacZ* and KH*abr2p-lacZ*. *A.*
91 *fumigatus* strains were cultivated at 37 °C in *Aspergillus* minimal medium (AMM) as
92 previously described (41). As solid media, malt extract medium (2 % (w/v) malt extract, 0.2
93 % (w/v) yeast extract, 1 % (w/v) glucose, 5 mM ammonium chloride, 1 mM di-potassium
94

95 hydrogenphosphate) or AMM containing 3 % (w/v) agar were used. Uridine (5 mM) or uracil
96 (5 mM) were added to the media when required. In case the *hph* or *ble* gene were used as a
97 selection marker gene for transformation of *A. fumigatus*, 100 µg hygromycin B and 100 µg
98 phleomycin per ml, respectively, were added to agar plates.

99 For induction of conidiophore formation of *A. fumigatus* (developmental cultures), conidia
100 were used to inoculate liquid cultures which were grown for 24 h at 37 °C. Mycelia were
101 filtered and exposed to air as previously described for *A. nidulans* (41). For transformation of
102 *Escherichia coli*, XL1-Blue (Stratagene, USA), INVαF' or TOP10F' (Invitrogen, The
103 Netherlands) were used. *E. coli* strains were grown at 37 °C in LB medium supplemented,
104 when required, with 100 µg or 50 µg per ml of ampicillin or kanamycin, respectively.

105 **Colony radial growth rate determination.**

106 Colony diameters of *A. fumigatus* were measured twice a day on both malt extract and AMM
107 agar plates over a period of 94 h. At least 10 colonies of each strain were analysed. Agar
108 plates had been point-inoculated centrally with a 2.5 µl drop of a suspension of 1×10^6 spores
109 per ml. Colony radial growth rates (C_r) (37) were calculated from the slope of the line
110 between 40 h and 72 h from a plot of colony radius *versus* time starting from the time of
111 inoculation. Data were processed by least square regression analysis.

112 **Analysis of conidial germination.**

113 50 ml AMM were inoculated with 1×10^7 conidia. Cultures were incubated with 180 rpm at
114 37 °C. Over a period of 16 h, samples were taken and deposited on microscope slides. To
115 determine germination, at least 100 conidia of each sample were counted.

116 **Standard DNA techniques.**

117 Standard techniques in the manipulation of DNA were carried out as described by Sambrook
118 *et al.* (31). Chromosomal DNA of *A. fumigatus* was prepared as previously described for *A.*

119 *nidulans* (1). For Southern blot analysis, chromosomal DNA of *A. fumigatus* was cut by
120 different restriction enzymes, as indicated. DNA fragments were separated on an agarose gel
121 and blotted onto Hybond N⁺ nylon membranes (Amersham Pharmacia Biotech, UK).
122 Labeling of the DNA probe, hybridization and detection of DNA-DNA hybrids were
123 performed using the DIG High Prime Labeling and Detection System (Amersham Pharmacia
124 Biotech, UK) according to the manufacturer's recommendations.

125 **Sequence analysis.**

126 Plasmid DNA was sequenced on both strands by primer walking using the Big Dye™
127 Terminator Cycle Sequencing Kit (Applied Biosystems, UK). Sequencing reactions were
128 separated on an Applied Biosystems ABI 310 sequencer. DNA sequence data were edited by
129 the programs "Sequence Navigator" and "Auto Assembler" (Applied Biosystems, UK). The
130 analysis of sequences was carried out using "Gene Works 2.2" (IntelliGenetics Inc., USA).

131 **Generation of recombinant plasmids.**

132 For generation of plasmid pKH1*pyrGhph* an upstream flanking region (1020 bp) of the *A.*
133 *fumigatus pyrG* gene with an introduced *Xba*I restriction site at its 3'end was generated by
134 PCR using oligonucleotides PyrG1 and PyrG2*Xba*I (Table 2), and genomic DNA from the *A.*
135 *fumigatus* wild-type strain as a template. The downstream fragment of *pyrG* with a size of
136 1032 bp with an introduced *Xba*I restriction site at its 5'end was synthesised by PCR,
137 employing PyrG3*Xba*I and PyrG4 oligonucleotides (Table 2), and genomic DNA from the
138 wild-type strain as a template. The upstream and downstream fragments of the *pyrG* gene
139 were cloned independently into the pCR2.1 vector (TA cloning kit, Invitrogen, The
140 Netherlands), yielding plasmids pTAFR1 and pTAFR2, respectively. After linearisation of
141 pTAFR1 with *Xba*I, the *hph* selection marker gene conferring hygromycin B resistance, was
142 introduced. A DNA fragment encoding the *hph* gene was obtained via double restriction of
143 plasmid pUCh*ph*1 (25) with *Xba*I and *Spe*I. In the *Xba*I site of the resulting vector designated

144 (pTAFR1*hph*), the downstream fragment was transferred from the vector pTAFR2, by
145 excision with *Xba*I restriction endonuclease.

146 To construct the *abr2* deletion plasmid **pUC Δ *abr2***, a 4125 bp PCR product was generated
147 using oligonucleotides *Abr2del_for* and *Abr2del_rev* (Table 2), and chromosomal DNA of the
148 *A. fumigatus* wild-type strain ATCC46645 as a template. The resulting PCR fragment spanned
149 the *abr2* gene and its upstream and downstream flanking region with a size of 1035 bp and
150 1020 bp, respectively. This PCR fragment was cloned into the pCR2.1 TOPO vector (TA
151 Cloning Kit, Invitrogen, The Netherlands) to give plasmid pCR2.1*abr2*. The PCR fragment
152 was re-isolated by restriction digest of plasmid pCR2.1*abr2* with *Eco*RI. The obtained DNA
153 fragment of 4125 bp was ligated into pUC18, also digested with *Eco*RI, yielding plasmid
154 pUC*abr2*. As a selection marker, the *hph* gene was used. It was obtained from plasmid
155 pUC*hph*1 (25), digested with *Nsi*I and *Xmn*I. The resulting DNA fragment of 2792 bp was
156 cloned into plasmid pUC*abr2* after restriction with *Nsi*I, which allowed the replacement of the
157 *abr2*-encoding region by *hph*. Thus, the resulting plasmid pUC Δ *abr2* contained the *E. coli*
158 *hph* gene under control of the *A. nidulans gpdA* promoter, flanked by fragments encoding
159 upstream and downstream sequences of the *abr2* gene. Plasmid pUC Δ *abr2* was linearised by
160 digestion with *Eco*RI. The resulting 4.8 kbp DNA fragment was used for transformation of the
161 *A. fumigatus* wild-type strain ATCC46645.

162 For complementation of the *abr2* deletion mutant, a PCR fragment of 3641 bp comprising the
163 *abr2* gene and 1 kb of promoter region, was generated by the use of primer pair *Abr2P_for*
164 and *Abr2.2_rev* (Table 2). The resulting *abr2*-encoding PCR fragment was used for a co-
165 transformation approach. Co-transformation was carried out by the use of the plasmid pAN8-
166 1 which contains the *ble* gene, conferring resistance to phleomycin (29). The *ble* gene is under
167 control of the *A. nidulans gpdA* promoter. Transcription of *ble* is terminated by the *A.*

168 *nidulans trpC* terminator. For the increase of transformation frequency plasmid HELP1 (19)
169 was simultaneously added to the transformation mix. Transformants were selected on AMM
170 agar plates containing phleomycin. Phleomycin-resistant transformants producing gray-green
171 conidia were checked by PCR for the presence of the *abr2* gene using primers Abr2disr_for
172 and Abr2disr_rev (Table 2) and chromosomal DNA of the transformants as the template.

173 To measure *abr2* expression, an *abr2p-lacZ* gene fusion was generated. For this purpose, a
174 1.23 kbp DNA fragment spanning the putative promoter region of *abr2* was amplified by PCR
175 using the oligonucleotides Abr2P_rev and Abr2P_for (Table 2), each of which encoding
176 *Bam*HI restriction sites. As the template, chromosomal DNA of *A. fumigatus* wild type was
177 used. The PCR fragment obtained was cloned into the pCR2.1 TOPO vector (TA Cloning Kit,
178 Invitrogen, The Netherlands). After restriction with *Bam*HI, the DNA fragment spanning the
179 promoter region was cloned into the *Bam*HI site of plasmid pUC*pyrG2lacZ* (23), which
180 carries the non-functional *pyrG2* allele. The resulting plasmid pUC*pyrG2abr2PlacZ* encoded
181 an in frame *abr2p-lacZ* gene fusion. This was checked by DNA sequence analysis across the
182 junctions. All *lacZ*-containing plasmids carried the *pyrG2* allele as the selection marker. It
183 encodes a nonfunctional *pyrG* gene of *A. fumigatus*, which forced site-specific integration of
184 plasmids into the chromosomal *pyrG* locus (42).

185 **Transformation of *A. fumigatus* and generation of the *pyrG* deletion strain KH Δ *pyrG***
186 **and of strain KH*pksPp-lacZ*.**

187 Transformation of *A. fumigatus* was carried out using protoplasts as previously described
188 (42). The *pyrG* deletion strain KH Δ *pyrG* was generated as follows: the vector pKH1*pyrG**hph*
189 was cut with *Spe*I and *Bgl*II yielding a DNA fragment of 5.9 kbp, which was used for
190 transformation of the *A. fumigatus* wild-type strain ATCC46645. The selection of *pyrG*
191 deletion strains occurred on AMM agar plates containing uracil and hygromycin B. Uracil-

192 auxotrophic, hygromycin B-resistant transformants were checked by Southern blot analysis
193 (data not shown). One of the resulting uracil-auxotrophic, hygromycin-resistant transformants
194 was designated KH Δ *pyrG*. It contained a deletion of 144 bp in the 3' coding region of the
195 *pyrG* gene. When uracil-auxotrophy was used, the *A. fumigatus* strain KH Δ *pyrG* (Table 1)
196 was applied. When selection for hygromycin B resistance was used, the wild-type strain
197 ATCC46645 was employed. Strain KH*pksPp-lacZ* was generated by transformation of strain
198 KH Δ *pyrG* with plasmid pUC*pyrG2pksP-lacZ* (23) encoding a *pksPp-lacZ* gene fusion. By
199 Southern blot analysis, a transformant strain designated KH*pksPp-lacZ* was identified
200 encoding a single copy of plasmid pUC*pyrG2pksP-lacZ* integrated at the chromosomal *pyrG*
201 gene locus (data not shown).

202 **Field emission scanning electron microscopy (FESEM).**

203 FEMES was carried out according to Maerker et al. (26). In brief, the conidia were harvested
204 with sterile water, containing 10 mM MgCl₂ and 10 mM CaCl₂ to a final concentration of 3 x
205 10⁸ conidia per ml. For fixation of the conidia 35% (v/v) formaldehyde was used in a final
206 concentration of 5% (v/v). After incubation on ice for 10 min, glutaraldehyde was added to a
207 final concentration of 2% (v/v). After washing with cacodylate buffer and subsequently with
208 TE buffer, the samples were placed onto poly(L-lysine) coated glass cover slips.

209 **β -Galactosidase (β -GAL) activity assays.**

210 *A. fumigatus* strains were grown in AMM at 37 °C. β -GAL activities were measured in
211 protein extracts obtained from three *A. fumigatus* cultures grown in parallel. Specific activities
212 were calculated as previously described (25).

213 **Laccase activity assay and determination of protein concentrations.**

214 *A. fumigatus* strains were grown in AMM at 37 °C. After harvesting at different time points
215 as indicated, mycelia were frozen in liquid nitrogen and ground to a fine powder. Mycelia

216 were suspended in extraction buffer (Tris/HCl 0.1 M, pH 7.0). Samples were centrifuged at 4
217 °C with 13,000 rpm for 10 min. The supernatant was retained. N,N-Dimethyl-*p*-
218 phenylendiamine (DMP) was used as the substrate. Laccase activity was determined in 900 µl
219 laccase buffer (37 mM citric acid monohydrate, 126 mM Na₂HPO₄, pH 6.0), 50 µl DMP (147
220 mM stock solution) and 50 µl enzyme solution. Extinction was followed at 550 nm at 25 °C
221 and the enzyme activities were calculated using a molar extinction coefficient of 1.8 mM⁻¹
222 cm⁻¹, as previously described (32). Laccase activities were measured in protein extracts
223 obtained from three *A. fumigatus* cultures grown in parallel. Protein concentrations were
224 determined according to Bradford (3).

225 **Sensitivity towards reactive oxygen species.**

226 1×10^8 *A. fumigatus* conidia were mixed with 10 ml AMM top agar and poured onto AMM
227 agar plates. In the center of the agar plate, a hole with a diameter of 10 mm was created,
228 which was filled with a solution of H₂O₂ or diamide. After an incubation of the agar plates at
229 37 °C for 16 h, the diameter of the inhibition zone was measured.

230 **Animal infection model.**

231 An optimised murine low dose model for invasive aspergillosis was applied (24, 35). Mice
232 were intranasally infected with a 25 µl drop of a fresh suspension containing 5×10^4 conidia.
233 Survival was monitored daily, and moribund animals were sacrificed by intraperitoneal
234 injection of 200 µl 3.2 % (v/v) narcoren (Rhone Merieux, Germany). The drinking water was
235 supplemented with 0.5 mg of tetracycline (Sigma) per ml to prevent opportunistic bacterial
236 infections. A control group (inhalation of PBS) remained uninfected to monitor the influence
237 of the immunosuppression procedure on vitality.

238

RESULTS

239

240

241 **Deletion of the laccase-encoding gene *abr2* of *A. fumigatus*.** To analyse the importance of
242 *Abr2* for the overall laccase activity of *A. fumigatus*, the *abr2* gene was deleted. For this
243 purpose, plasmid pUCdel*abr2* (Fig. 1A) was generated (see Materials and Methods). A DNA
244 fragment obtained by digestion of plasmid pUCdel*abr2* with *Xmn*I which encodes the
245 hygromycin B resistance gene *hph* flanked by upstream and downstream sequences of the
246 *abr2* gene (Fig. 1A), was used for transformation of the *A. fumigatus* wild-type strain
247 ATCC46645. Forty two hygromycin B resistant transformants were isolated. Twenty of
248 them were tested by PCR for presence of the hygromycin resistance gene. Seven of those,
249 which showed the presence of the selection marker gene, were analysed by Southern blot
250 analysis. Four of these transformants exhibited the expected gene replacement (Fig. 1B, lanes
251 3, 4, 5, 7; Fig. 1C). One of the mutant strains (lane 3) was designated *AfΔabr2* and used for
252 further studies. Growth of strain *AfΔabr2* on agar plates revealed that the *abr2* deletion
253 affected the pigment formation of conidia (Fig. 2A). The colonies showed a brown color. A
254 similar finding was previously reported for an *abr2* mutant by both Tsai et al. and
255 Krappmann et al. (17, 38). Complementation of the *abr2* deletion mutant using the wild-type
256 *abr2* gene was carried out. For this purpose, the *abr2* gene was amplified by PCR. The
257 generated PCR fragment was applied to a co-transformation experiment, using plasmids
258 pAN8-1 and pHELP1 (see Materials and Methods). After transformation of the *Δabr2* mutant,
259 17 phleomycin-resistant transformants were isolated. Six of them produced gray-green
260 conidia, i.e., were complemented to the wild type (data not shown). The presence of the *abr2*
261 gene in the transformants producing wild-type conidia was shown by PCR analysis (data not
262 shown).

263 The germination of conidia (Fig. 2B), the growth rate measured as diameter of colonies on
264 AMM agar (Fig. 2C) and malt agar (Fig. 2D) were the same for both the *abr2* deletion mutant
265 and the wild-type strain, indicating that *abr2* is not essential for vegetative growth. Radial
266 growth rates (Cr) of colonies of *A. fumigatus* wild-type and Δ *abr2* mutant strain on AMM
267 agar plates were 0.46 and 0.48 mm/h, respectively. Cr on malt extract showed values of 0.5
268 mm/h for *A. fumigatus* ATCC46645 and 0.53 mm/h for the *abr2* deletion strain. SDs were in
269 the range of 0.005-0.02.

270 Conidia of the strain Af Δ *abr2* were analysed by FESEM (Fig. 3). As previously reported (14)
271 and shown here as a control, wild-type conidia display an ornamented surface (Fig. 3A) which
272 is lacking in *pksP* mutant conidia (Fig. 3B). Conidia of the *abr2* deletion strain displayed
273 some ornamentation which was less pronounced than that observed on wild-type conidia. This
274 finding indicates that formation of the DHN-like melanin intermediate present in the Δ *abr2*
275 mutant is sufficient for production of at least some ornamentation on the surface of conidia.

276

277 **In contrast to the *pksP* mutant the *abr2* deletion mutant showed the same sensitivity**
278 **against H₂O₂ and diamide *in vitro* as the wild type.**

279 Previously, we showed that mutation of the polyketide synthase gene *pksP* led to increased
280 sensitivity of the respective mutant against ROS generated by immune effector cells (14). To
281 analyse whether the deletion of *abr2* enhances sensitivity against ROS in the respective
282 mutant or whether the compound produced by the DHN-like melanin pathway to the stage of
283 the Abr2 laccase is sufficient to scavenge ROS, the sensitivity of the Δ *abr2* mutant against
284 ROS was compared with that of both the *pksP* mutant and the wild-type strain. As shown in
285 Fig. 4A, the *pksP* mutant showed increased sensitivity against ROS whereas there was no
286 significant difference between the Δ *abr2* mutant and the wild-type strain. The analysis of the

287 effect of diamide confirmed the results obtained for H₂O₂, but in case of the *pksP* mutant the
288 effects were less prominent (Fig. 4B). Taken together, the Δ *abr2* mutant showed the same
289 sensitivity against ROS as the wild type.

290

291 **Abr2 activity was detectable during sporulation but did not contribute to laccase activity**
292 **during hyphal growth under standard conditions.**

293 To determine the contribution of Abr2 to total laccase activity, cell extracts of both the wild
294 type and the Δ *abr2* deletion strain were analysed for laccase activity. Laccase activity was
295 detectable in both strains during vegetative growth after 24 h (Fig. 5). In contrast to the wild-
296 type strain, laccase activity was reduced in the Δ *abr2* mutant strain in sporulating mycelia.
297 These findings indicate that the Abr2 laccase is mainly active during sporulation under
298 standard conditions. Furthermore, these results imply that additional laccases are active
299 during vegetative growth of the fungus under the conditions tested and that their activities
300 also increase during sporulation as seen from the *abr2* deletion strain.

301

302 **Sporulation-dependent expression of an *abr2p-lacZ* gene fusion.**

303 To analyse the regulation of *abr2* expression and correlate this data with the Abr2 activity
304 during sporulation, *A. fumigatus* strains were generated carrying an *abr2p-lacZ* gene fusion
305 integrated in single copy at the *pyrG* gene locus. For this purpose, plasmid
306 pUCPyrG2*abr2PlacZ* was used (Fig. 6A). It encodes the *abr2* gene promoter fused in frame
307 with the *E. coli lacZ* gene. Transformation of *A. fumigatus* strain KH Δ *pyrG* using this plasmid
308 resulted in the isolation of eight transformants. Southern blot analysis indicated the presence
309 of the gene fusion in single copy at the *pyrG* gene locus (Fig. 6B, C). The 8 kbp band
310 characteristic of the wild-type *pyrG* gene (Fig. 6A, lane 1) had disappeared in the

311 transformants (Fig. 6A, lanes 2 and 3). Instead, they showed a 10 kbp band due to the
312 integration of the plasmid at the *pyrG* gene locus. The transformants shown in Fig. 6C were
313 designated KH*abr2placZ*-2 and KH*abr2placZ*-4 and used in further studies.

314 The expression of the *abr2p-lacZ* gene fusion was determined. Results are shown in Fig. 7A.
315 As expected, the expression of the gene fusion was detectable during sporulation and
316 increased up to 48 h. There was hardly *abr2p-lacZ* expression during vegetative growth of the
317 fungus. These data well agree with the results on the laccase specific activity (Fig. 5)
318 indicating that Abr2 mainly contributes to overall laccase activity during sporulation but not
319 during vegetative growth under the conditions applied. Interestingly, the expression pattern of
320 the *pksPp-lacZ* and *abr2p-lacZ* gene fusions apparently differed. The *abr2p-lacZ* expression
321 increased seventeen-fold during sporulation after 48 hours compared with the expression
322 during vegetative growth, whereas for the *pksPp-lacZ* gene fusion, this increase during
323 sporulation only was 2.5-fold (Fig. 7B).

324

325 **The Δ *abr2* mutant showed no reduction in virulence compared with the wild type.**

326 To assess a possible role of Abr2 in pathogenesis, the corresponding deletion mutant was
327 tested in an intranasal mouse infection model of invasive aspergillosis. Groups of 10
328 immunosuppressed mice were infected by intranasal inhalation with 5×10^4 conidia of the
329 wild-type or the Δ *abr2* mutant strain. Results of a representative experiment are shown in Fig.
330 8. In the groups infected with wild-type conidia (strain ATCC46645), mortality was 80 %
331 after 11 days (Fig. 8). When mice were infected with conidia of the Δ *abr2* mutant strain
332 Af Δ *abr2*, mortality was 80 % after 13 days. Hence, the overall-mortality of the *abr2* deletion
333 strain was similar to that of the wild type. Moreover, at the beginning of the experiment more
334 of the mice infected with conidia of the Af Δ *abr2* strain died compared with mice infected

335 with wild-type conidia. Taken together, these data indicate that Abr2 does not contribute to
336 virulence of *A. fumigatus*.

337

338

DISCUSSION

339

340 Under the conditions applied, laccase activity of *A. fumigatus* was detected during vegetative
341 growth, which strongly increased during sporulation. The analysis of the *abr2* deletion mutant
342 led to the conclusion that Abr2 only contributes to overall laccase activity during sporulation.
343 It is worth to notice that the substrate for determining laccase activity, DMP, can be
344 oxidised not only by laccases but also by other enzymes as peroxidases (32). However,
345 because Abr2 clearly used DMP as the substrate and its amino acid sequence resembles that
346 of a typical fungal laccase, it is very likely that Abr2 represents a laccase. The finding that
347 Abr2 only contributes to overall laccase activity during sporulation well agrees with the
348 analysis of an *abr2p-lacZ* gene fusion which was mainly expressed during sporulation. By
349 contrast, there was expression of a *pksPp-lacZ* gene fusion during vegetative growth which
350 also increased during sporulation, but the increase was less than that observed for the *abr2p-*
351 *lacZ* gene fusion. In fact, the *abr2p-lacZ* expression increased steadily up to seventeen-fold
352 during sporulation compared with the expression during vegetative growth, whereas for the
353 *pksP-lacZ* gene fusion, this increase during sporulation only was 2.5-fold and reached its
354 maximum after 24 hours. This finding implies that differential regulation of genes from the
355 same cluster occurs. A similar observation was made for genes of other clusters in
356 filamentous fungi, e.g., genes belonging to the penicillin biosynthesis cluster in *A. nidulans*
357 are, at least in part, differentially regulated (4, 7). Consistently, the promoter regions of *pksP*
358 and *abr2* apparently contain different *cis*-acting elements. Computer analysis revealed that the
359 *pksP* promoter region encodes a single putative stress response element (STRE) (11, 36),

360 three putative AbaA binding sites (2), a potential cAMP-responsive element originally
361 described in the *S. cerevisiae* SSA3 gene promoter and two putative stunted A (StuA) binding
362 sites (10). By contrast, in the promoter region of *abr2* a single putative AbaA site, a single
363 potential STRE and two putative copper signalling elements (CuSE) were detected. CuSE are
364 defined by the consensus sequence 5'-DWDDHGCTGD-3' (D = A, G, or T; H = A, C, or T
365 and W= T, or A). They are bound by Cuf1 (copper-sensing transcription factor) and were
366 described for *Schizosaccharomyces pombe* (18). Whether these elements contribute to *abr2*
367 expression remains to be tested.

368 Even in liquid medium *abr2p-lacZ* expression increased during later stages of the cultivation
369 (data not shown) when most likely sporulation was induced even in liquid medium (34).
370 Based on the analysis of the *abr2* deletion mutant it can be concluded that the laccase activity
371 measured during vegetative growth was due to other laccases of *A. fumigatus*. In the genome
372 of *A. fumigatus*, there are at least three additional candidate genes (Afu1g15670, Afu4g14280,
373 Afu2g17540; www.tigr.org). *Abr2* activity is mainly, if not exclusively, required for the
374 production of the gray-green spore pigment. Interestingly, in contrast to the *pksP* mutant,
375 conidia of the Δ *abr2* mutant did not show increased sensitivity against ROS or diamide.
376 Therefore, the intermediate of DHN-like melanin produced up to the stage of *Abr2* is
377 apparently sufficient to protect conidia against ROS. This assumption was further supported
378 by the observation that in a low dose intranasal mouse infection model there was no
379 difference in virulence of the *abr2* mutant compared with the wild type. Furthermore, the
380 surface of Δ *abr2* conidia looked more similar to that of wild-type conidia, i.e., it displayed at
381 least some ornamentation, whereas the *pksP* mutant showed a smooth surface (14). Taken
382 together, these data support the model that mainly the *pksP* gene of the DHN-like melanin
383 biosynthesis gene cluster is important for virulence.

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384

385

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528 **TABLE 1.** *Aspergillus fumigatus* strains.

Strain	Genotype and/or phenotype	Reference
ATCC46645	wild type	ATCC
KH Δ <i>pyrG</i>	derived from ATCC46645; <i>pyrG::hph^a</i> , <i>pyrG</i> , Hyg ^R	This study
Af Δ <i>abr2</i>	Δ <i>abr2</i> , Hyg ^R	This study
Af <i>pksP</i>	<i>pksP</i>	14,19
KH <i>pksPp-lacZ</i>	<i>pyrG1::pyrG2</i> , <i>pksPp-lacZ</i> , PyrG ⁺	This study
KH <i>abr2p-lacZ</i>	<i>pyrG1::pyrG2</i> , <i>abr2p-lacZ</i> , PyrG ⁺	This study

529 ^a *hph*: *E. coli* hygromycin B phosphotransferase gene

530

531

532

533 **TABLE 2.** Oligonucleotide primers used in the study

Primer	Sequence ^a
Abr2del_for	TAAGGAATCGCACCATCGCC
Abr2del_rev	GAGTGGTCATATGGCAGTGC
Abr2P_for	GTGGATCCCTCAACCGGTGC
Abr2P_rev	GGATCCCATCCTGTTGTCGTGTAT
Abr2.2_rev	TGCTGGGATCCGACTAGTACG
Abr2disr_for	GGCGAGCAGATACTTATGGG
Abr2disr_rev	GAGTGGTCATATGGCAGTGC
PyrG1	TTGACCCACAGTCGGAGGC
PyrG2XbaI	GGAGTCTAGAATTGCTGTCC
PyrG3XbaI	CCTCTAGAAGCAAAAGTGTAGTGC
PyrG4	TCCTCCCCTCATCTGTTGG

534 ^a All sequences are depicted in the 5'→3' direction

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LEGENDS TO FIGURES

542

543 **Fig. 1.** Deletion of *A. fumigatus abr2* gene.

544 **A.** Schematic map of the generation of the *abr2* knock-out plasmid pUCdelabr2.

545 Abbreviations: amp^R, ampicillin resistance gene; *hph*, hygromycin B phosphotransferase

546 gene used as the selection marker gene in *A. fumigatus*.

547 **B.** Southern blot analysis. Chromosomal DNA of the wild-type strain ATCC46645 (lane 1)

548 and transformant strains (lanes 2-8) was cut by *Sac*II. A 641 bp *abr2*-derived PCR fragment

549 was used as the probe. In the *AfΔabr2* mutant strains, the band characteristic of the wild type

550 (lane 1) of 4344 bp had disappeared. Instead, the band of 2725 bp characteristic of gene

551 replacement at the *abr2* locus was detected (see Fig. 1C).

552 **C.** Schematic representation of the chromosomal *abr2* locus of the wild type (1) and the

553 *AfΔabr2* deletion mutant (2). Restriction endonuclease cleavage sites, the DNA fragments

554 identified by Southern blot analysis (Fig. 1B) and the position to which the probe hybridises,

555 are indicated.

556

557 **Fig. 2.** Phenotypic characterisation of mutant strain *AfΔabr2* and wild-type strain

558 ATCC46645.

559 **A.** Growth and sporulation. Colonies were grown on AMM agar plates for 72 h at 37 °C.

560 **B.** Kinetics of germ tube outgrowth for *A. fumigatus* conidia incubated in AMM at 37 °C.

561 The number of conidia showing a germ tube was recorded at different times of incubation

562 and is presented as the percentage of the total number of conidia. The results are

563 representative of the results of two independent experiments.

564 **C. + D.** Growth of *A. fumigatus* strains on AMM agar plates and malt agar plates,
 565 respectively, at 37 °C. The diameter of colonies was determined. Data for each strain
 566 represent the mean of at least 10 colonies, grown independently. SDs were in the range of 0.2
 567 - 0.7 mm.

568

569 **Fig. 3.** Field emission scanning electron micrographs of conidia. Bar: 2µm. **A.** *A. fumigatus*
 570 wild-type strain ATCC46645. **B.** *A. fumigatus pksP* mutant. **C.** *A. fumigatus abr2* deletion
 571 mutant.

572

573 **Fig. 4.** Sensitivity of *AfΔabr2* towards H₂O₂ and diamide.

574 The strains *A. fumigatus* ATCC46645 (*Af wt*), *AfpksP* and *AfΔabr2* were analysed. Two
 575 concentrations of H₂O₂, 50 µl and 100µl from a 6% (v/v) H₂O₂ solution, were examined.
 576 100µl and 200µl from a 0.1M diamide solution were used in the experiment. Data for each
 577 strain and concentration represent the mean and SDs of five independently performed assays.

578 **A.** H₂O₂ **B.** Diamide

579

580 **Fig. 5.** Laccase activity of mutant strain *AfΔabr2* and wild-type strain ATCC46645 (*Af wt*).
 581 Enzyme activity was monitored during vegetative growth in AMM after 24 h (mycelium) and
 582 sporulation after 24, 34 h and 48 h (sp).

583

584 **Fig. 6.** Integration of an *abr2p-lacZ* gene fusion in single copy at the *A. fumigatus*
 585 chromosomal *pyrG* gene locus.

586 **A.** Schematic map of plasmid pUCPyrG2*abr2PlacZ*. Abbreviations: *abr2P*, promoter region
 587 of the *abr2* gene; Amp^R, ampicillin resistance gene; *lacZ*, *E. coli lacZ* gene; *PyrG*,

588 orotidine 5'-monophosphate decarboxylase gene of *A. fumigatus*, used as the selection
589 marker gene. The asterisks indicate mutations.

590 **B.** Schematic representation of the chromosomal *pyrG* gene locus of strain KH Δ *pyrG*
591 carrying a deletion of part of the *pyrG* gene and of strain KH*abr2placZPyrG* carrying the
592 *abr2p-lacZ* gene fusion integrated at the *pyrG* gene locus. Restriction endonuclease
593 cleavage sites and the position, to which the probe hybridizes, are indicated.

594 **C.** Southern blot analysis. Chromosomal DNA of the *A. fumigatus* wild-type strain
595 KH Δ *pyrG* (lane 1) and strains KH*abr2placZPyrG*-2 and KH*abr2placZPyrG*-4 (lanes 2
596 and 3) was digested by *Bgl*III. A 450 bp PCR fragment encoding a part of the *A. fumigatus*
597 *pyrG* gene (Fig. 6B), was used as the probe.

598

599 **Fig. 7.** Expression of *abr2p-lacZ* (A) and *pksPp-lacZ* (B) gene fusions during vegetative
600 growth and sporulation. The β -GAL activity was measured after 24 hours of growth in liquid
601 culture, and after 24 and 48 hours after induction of sporulation. Data for each condition
602 represent the mean and SDs of three independently grown cultures.

603

604 **Fig. 8.** Virulence of *A. fumigatus* wild type (ATCC46645) and Δ *abr2* mutant strain in mice.
605 Groups of 10 BALB/c mice were analysed, each infected intranasally with 5×10^4 *A.*
606 *fumigatus* conidia, as indicated. Survival was monitored for 17 days.

Fig. 1

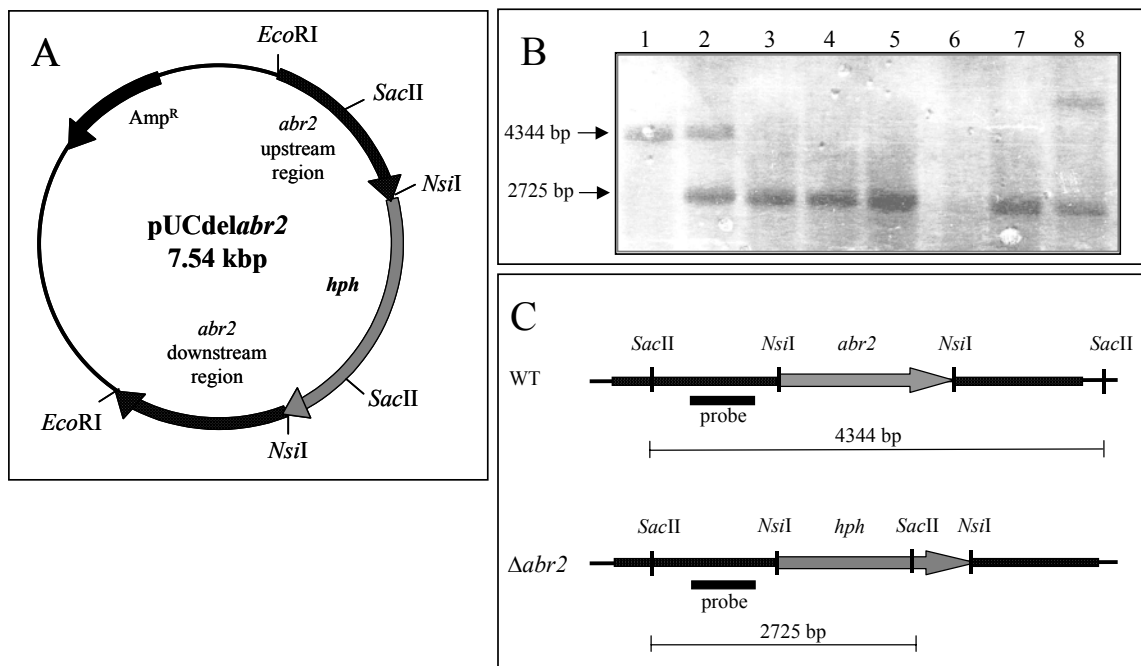


Fig. 2

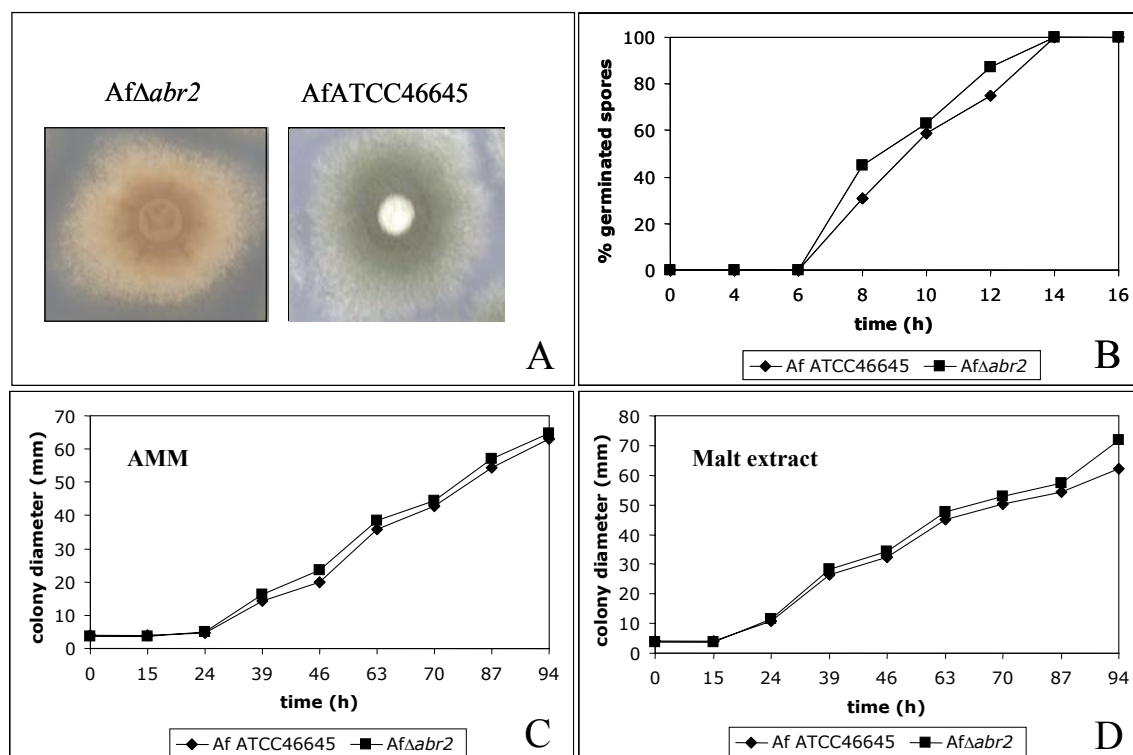


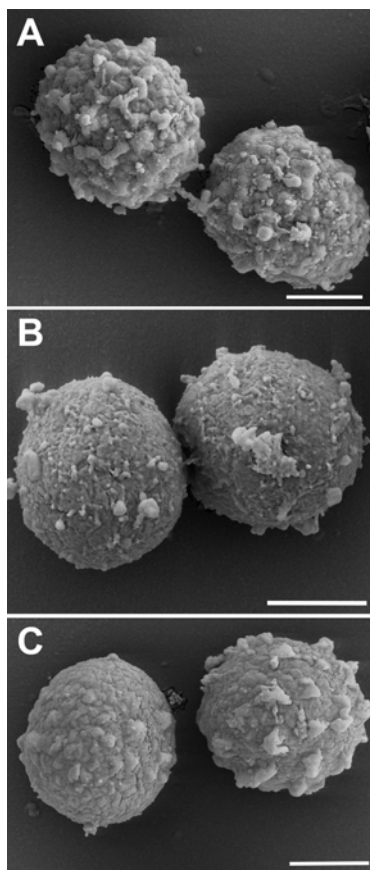
Fig. 3

Fig. 4

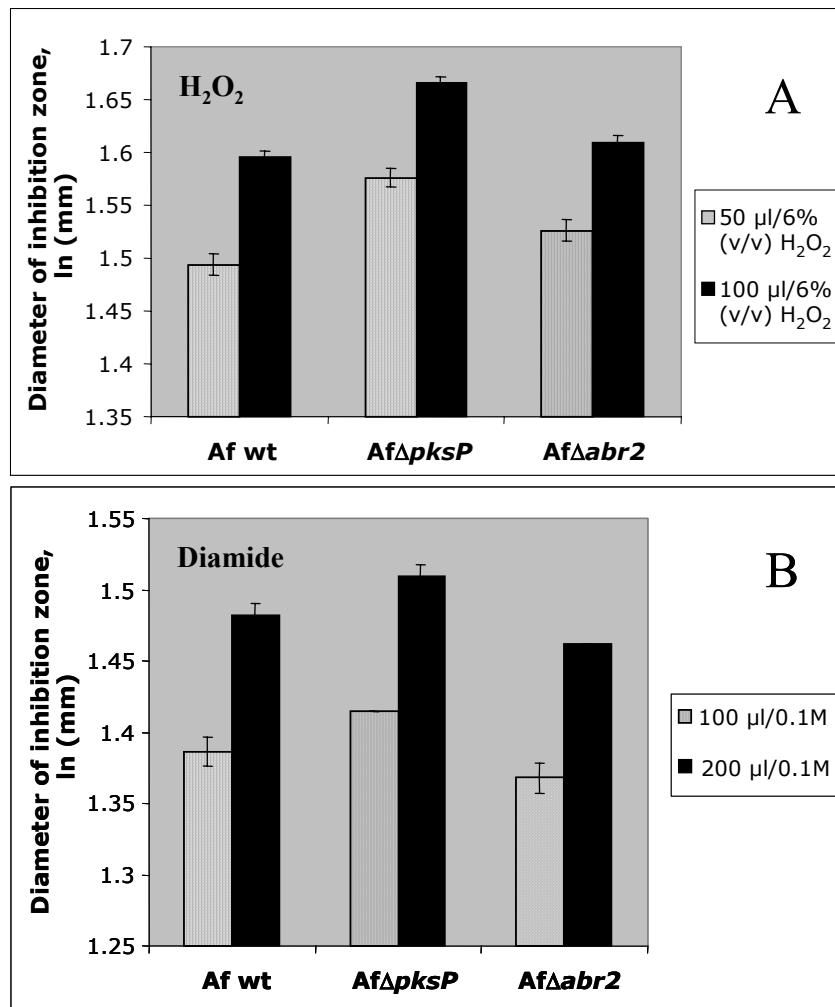


Fig. 5

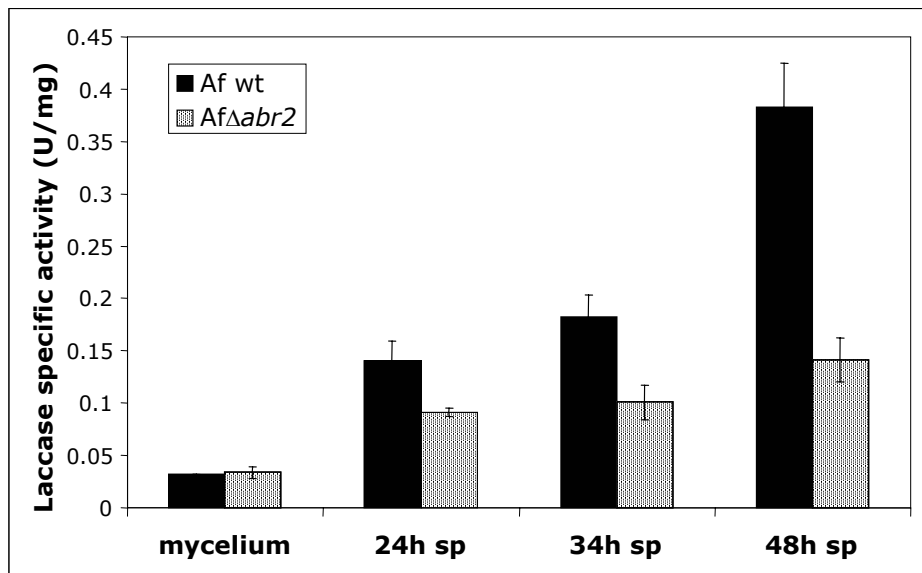


Fig. 6

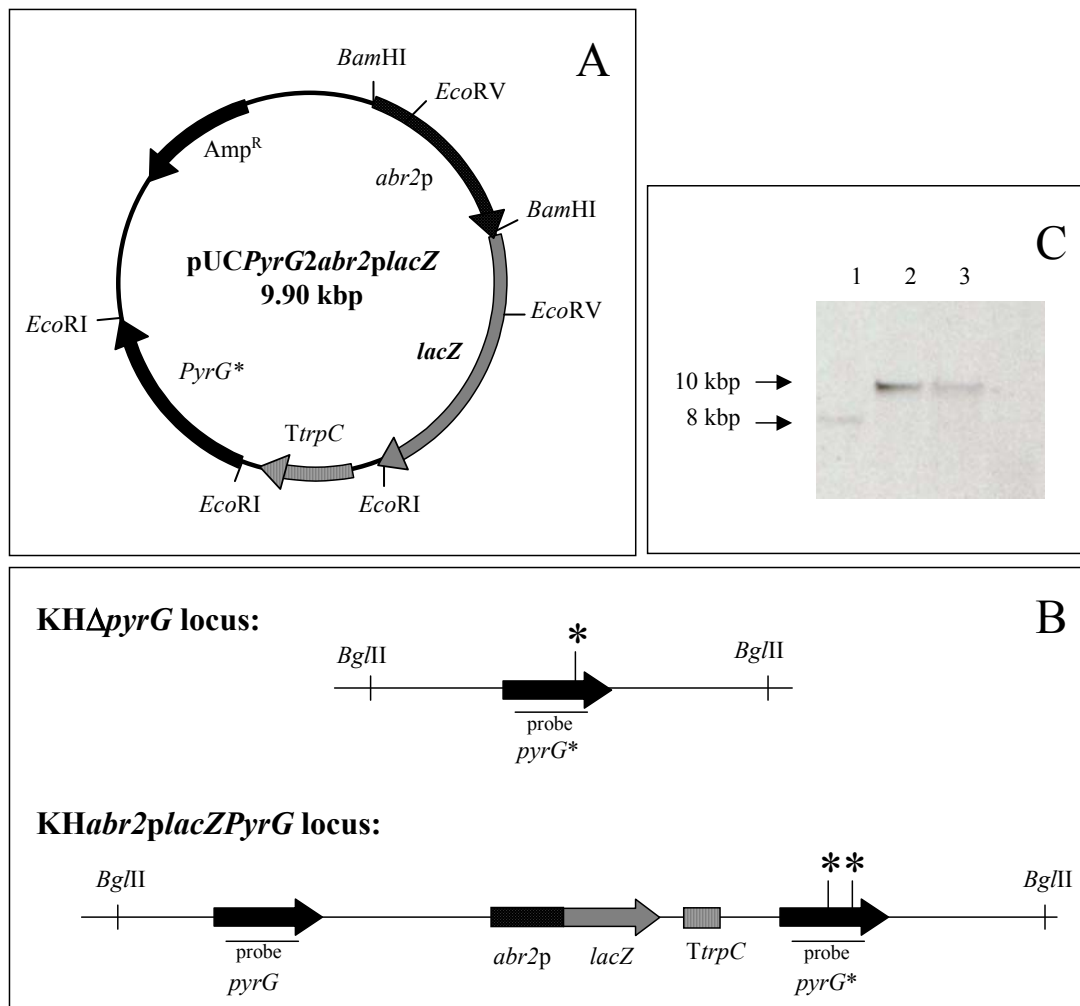


Fig. 7

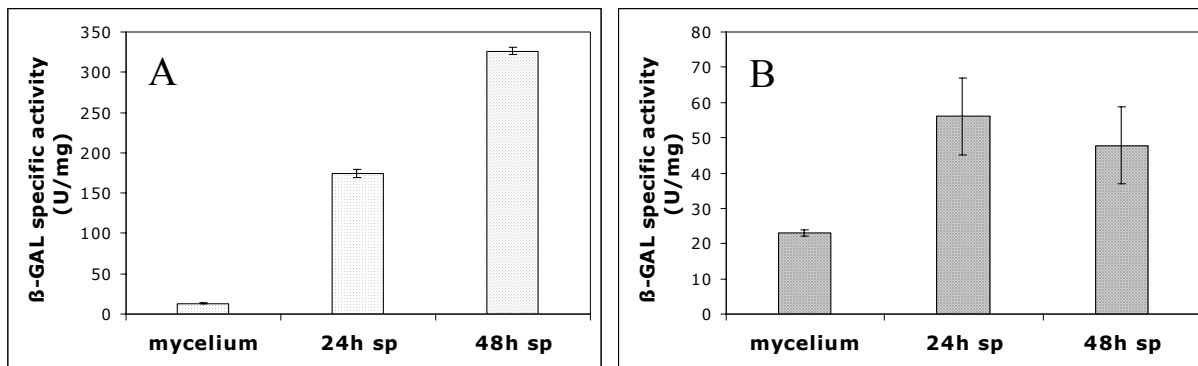


Fig. 8

