The \(akuB^{KU80}\) Mutant Deficient for Nonhomologous End Joining Is a Powerful Tool for Analyzing Pathogenicity in \(Aspergillus fumigatus\)

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To increase the frequency of homologous recombination, we inactivated the \(KU80\) homologue in \(Aspergillus fumigatus\) (named \(akuB^{KU80}\)). Homologous integration reached about 80% for both calcineurin A (\(calA\)) and polyketide synthase \(pksP\) (\(alb1\)) genes in the \(akuB^{KU80}\) mutant to 3 and 5%, respectively, when using a wild-type \(A. fumigatus\) strain. Deletion of \(akuB^{KU80}\) had no influence on pathogenicity in a low-dose murine infection model.

Double-strand breaks are the most severe form of DNA damage. Eukaryotes have two main pathways to deal with this type of DNA damage: (i) homologous recombination, involving interaction between homologous sequences, and (ii) nonhomologous end joining, involving direct ligation of the strand ends independently of DNA homology (17). The most important double-strand break repair pathway in \(Saccharomyces cerevisiae\) is homologous recombination (18, 20), while other organisms, such as humans, preferentially use nonhomologous end joining. The nonhomologous end-joining process is mediated by the DNA-dependent protein kinase catalytic subunit, the Ku70-Ku80 heterodimer, and the DNA ligase IV-Xrcc4 complex (17). Recently, Ninomiya et al. (13) disrupted \(Neurospora\ crassa\ genes homologous to human \(KU70\) and \(KU80\). Transformation of the \(N. crassa\ Ku80\) disruption strains yielded 100% transformants exhibiting integration at the homologous site, compared to 10 to 30% for a wild-type recipient.

\(Aspergillus fumigatus\) causes life-threatening pulmonary disease in severely immunocompromised patients (3). The completion of the \(A. fumigatus\) genome sequence has offered an opportunity to investigate complete pathways and classes of genes and do high-throughput gene tagging (12). New methodologies for rapid construction of gene replacements in \(Aspergillus\) have been proposed (21, 22). The rate-limiting step for further progress in this area is the percentage of homologous integration in \(A. fumigatus\), which in some cases can be below 5%. Thus, we decided to delete the \(A. fumigatus\ KU80\) gene and verify whether, as in \(N. crassa\) (13), the absence of nonhomologous end joining could increase the integration of introduced exogenous DNA fragments by homologous recombination.

To evaluate whether the deletion of these genes could increase homologous recombination, we selected \(calA\) and \(pksP\) (\(alb1\)), which encode the calcineurin A catalytic subunit and a polyketide synthase, respectively. Calcineurin is a serine-threonine-specific phosphatase heterodimer consisting of a catalytic subunit A and a \(Ca^{2+}\)-binding regulatory B subunit. The association of the two subunits is essential for activity (7). The polyketide synthase is involved in the melanin biosynthesis required for conidial pigmentation (9, 19), so that the conidia of the \(\Delta pksP\) mutant obtained by gene replacement will be white instead of green like the parental strain (11).

\(A. fumigatus\) strain CEA17 (pyrG) (2) was used throughout this work. The media used were YAG containing 2% (wt/vol) glucose, 0.5% (wt/vol) yeast extract, 2% (wt/vol) agar, and trace elements; YUU (YAG medium supplemented with 1.2 gliter each of uracil and uridine), YG medium (same composition, but without agar) or minimal medium (MM: 1% glucose, nitrate salts, trace elements, 2% agar, pH 6.5). Trace elements, vitamins, and nitrate salts are described in the appendix to Kafer (5). DNA manipulations were according to Sambrook et al. (15). DNA fragment probes for Southern blots were labeled with [\(\alpha^{-32}\)P]dCTP using the RTS Rad Prime DNA labeling system kit (Gibco-BRL).

PCR primers were designed for amplifying each DNA fragment necessary for the PCR-mediated technique (8). In all the deletion constructs, the \(A. fumigatus\) pyrG gene was amplified from the pCDA21 plasmid (1) and is referred to as the \(zeo-pyrG\) cassette because the amplified fragment also contains the zeocin resistance gene. For DNA fragments containing the gene-flanking regions, genomic DNA was used as the template. For a description of the oligonucleotides used in this work and the strategy for PCR-mediated amplification, see Table 1.

The \(KU80\) gene (here named \(akuB^{KU80}\), Afu2g02620, accession number EAL87359) was identified in the \(A. fumigatus\) genome (http://www.tigr.org/tdb/e2k1/afu1/) by similarity to \(N. crassa\) mus-52 (42% identity, 60% similarity, e-value = 7e-117). Routinely, in PCR-mediated deletions for \(A. fumigatus\), flank-
ing regions of about 1.5 to 2.0 kb were used. This size increased the frequency of homologous integration (data not shown). Flanking sequences shorter than 500 bp did not yield any homologous integration (data not shown).

Transformation of A. fumigatus strain CEA17 (mutant pyrG recipient strain) was according to the procedure of Osmani et al. (14) using 5 μg of linear DNA fragments. A. fumigatus was transformed with the PCR-mediated deletion cassette for the aYKUS08 gene. Several transformants were obtained and identified by their ability to grow in YAG. These transformants were screened for sensitivity to hydroxyurea, camptothecin, 4-nitroquinoline oxide, and methane methyl sulfonate (MMS). Only one transformant in 100 displayed decreased growth in the presence of MMS (a frequency of 1%) (Fig. 1A). The N. crassa mus-52 mutant also showed mild sensitivity to MMS (13). The pyrG+ strain KU80Δ was point inoculated on YUU plates containing 5-fluoroorotic acid in a subinhibitory concentration of 0.55 mg/ml, resulting in the recovery of a sector that was auxotrophic to uridine and uracil and resistant to up to 0.75 mg/ml of 5-fluoroorotic acid. This strain was named

| Primer | Sequence | PCR conditions |
|--------|----------|----------------|
| ALB-1  | 5'-CTGACTGATTCGAGCCAGG-3' | (i) 5'-Flanking region about 2,000 bp (ALB-1 × ALB-2) |
| ALB-2  | 5'-CTGGAAATCTTGCAAGGAGTAGTGG-3' | (ii) zeo-pyrG cassette (ALB-1 × ALB-2) |
| ALB-3  | 5'-AAGAGATTGGAAGAGAG-3' | (iii) 3'-Flanking region about 2,000 bp (ALB-3 × ALB-4) |
| ALB-4  | 5'-TGAGGCAATTCTGGTCATACACITTGGAAT-3' | (iv) Fusion PCR fragment about 6.4 kb (ALB-1 × ALB-4). |
| ALB-A  | 5'-CTCTGAGCTGAGAATTCGAGTTGCGG-3' | (i) zeo-pyrG cassette (ALB-A × ALB-B) |
| ALB-B  | 5'-AGTTATGCACAAAAATTGCCCTCAAAAT-3' | (ii) Fusion PCR fragment about 6.4 kb (ALB-1 × ALB-4). |

*The 50-μl amplification mixture included 1× Platinum Taq DNA polymerase high-fidelity buffer (Invitrogen), 20 pmol of each primer, 0.4 mM deoxynucleotide triphosphate mix, 1.0 U of Taq DNA HiFi platinum polymerase (Invitrogen), and 500 ng of genomic DNA or 100 ng plasmid. PCR amplification was carried out in a PTC100 96-well thermal cycler (MJ Research) at 94°C for 2 minutes and 30 cycles of 94°C for 1 minute, 52 to 60°C (depending on the fragment) for 1 minute, and 68°C for 7 minutes, followed by an extension step at 68°C for 7 minutes. After the reaction, the PCR products were purified with a Qagen PCR cleanup kit according to the manufacturer’s instructions.

FIG. 1. ΔakaB<sub>KLUS0</sub> strain displays increased sensitivity to MMS. (A) Growth phenotype of the A. fumigatus wild-type and KU80ΔpyrG strains grown for 48 h at 37°C in YUU medium in the presence or absence of different concentrations of MMS. (B) Southern blot analysis of the KU80ΔpyrG strain. At the left panel, genomic DNA from the wild-type and KU80ΔpyrG strains was isolated and cleaved with the enzyme KpnI and the pyrG gene was used as a hybridization probe. The akaB<sub>KLUS0</sub> gene probe recognized a single band of approximately 14.0 kb only in the wild-type strain, indicating the akaB<sub>KLUS0</sub> gene was deleted in the KU80ΔpyrG strain. In the right panel, genomic DNA from the wild-type and KU80ΔpyrG strains was isolated and cleaved with the enzyme KpnI and the pyrG gene was used as a hybridization probe. The pyrG gene recognized one band in the wild-type strain (approximately 4.0 kb) and two bands in the akaB<sub>KLUS0</sub> deletion strain (around 4.0 and 9.0 kb). This additional band in the KU80ΔpyrG strain indicates a single event of integration of the transforming deletion cassette.
KU80ΔpyrG. The allelic replacement of the \( akuB^{KU80} \) gene in this transformant (KU80ΔpyrG strain) and the presence of a single integration event were confirmed by Southern blot analysis (Fig. 1B).

Next, the effects of deleting the \( akuB^{KU80} \) gene on \( A. fumigatus \) homologous recombination integration of the \( calA \) and \( pksP \) genes were investigated. Therefore, the \( A. fumigatus \) \( calA \) and \( pksP \) deletion cassettes were amplified by using PCR-mediated deletion (Table 1). One hundred transformants were obtained and 15 of them were evaluated by Southern blotting for the absence of the \( calA \) gene; 80% of them showed homologous integration of the deletion cassette at the \( calA \) locus and a single event of integration, as shown by four representative transformants in Fig. 2A. The transformants that displayed nonhomologous integration at the \( calA \) locus were also not integrated at the \( pyrG \) locus (data not shown), suggesting that there are other mechanisms of nonhomologous recombination in \( A. fumigatus \) besides nonhomologous end joining. When the wild-type strain was transformed with the \( calA \) deletion cassette, the frequency of homologous integration was 3%.

The primary transformants already displayed a pronounced phenotypic defect, i.e., small sporulating colonies that, when purified, showed comparable growth at different temperatures (Fig. 2B). The morphology of the \( \Delta calA \) germ tubes showed increased branching compared to the corresponding wild-type strain (Fig. 2C). Calcineurin A deletion was also found to affect colony morphology in several media known to modulate the \( C. albicans \) dimorphic switch (16). These results suggest that the \( calA \) gene is not essential in \( A. fumigatus \), but its deletion confers a severe growth defect that is probably due to abnormal branching.

To further confirm homologous integration in a high frequency in the \( \Delta akuB^{KU80} \) strain, the \( pksP \) deletion cassette was introduced into strain KU80ΔpyrG. Transformants were
FIG. 3. Virulence of A. fumigatus KU80Δ mutant strain. Survival of mice infected intranasally with conidia of A. fumigatus CEA17pyrG+ and KU80Δ was determined. As shown by D’Enfert et al. (3) uracil auxotrophic A. fumigatus strains show attenuated virulence. Therefore, the uracil prototrophic strain CEA17pyrG+ was isolated from strain CEA17 by spontaneous reversion of the point mutation in the pyrG locus. CEA17pyrG+ shows the same virulence in comparison to the A. fumigatus wild-type strain ATCC 46465 (data not shown). For the same reason the uracil prototrophic strain KU80Δ was used instead of KU80ΔpyrG. In brief, female BALB/c mice (Harlan Winkelmann, Borchsen, Germany) of 18 to 20 g body weight were immunosuppressed by intraperitoneal injection of cyclophosphamide (100 mg/kg; Sigma, Taufkirchen, Germany) on days −4, −1, 2, 5, 8, 11, and 14 prior to and after infection on day 0. A single dose of cortisone acetate (200 mg/kg of body weight; Sigma) was injected subcutaneously on day −1. Suspensions of A. fumigatus conidia were harvested with phosphate-buffered saline containing 0.1% (vol/vol) Tween 80 (Merck) and filtered through a 40-μm nylon cell strainer (BD Biosciences Europe, Belgium). Mice were anesthetized by intraperitoneal injection of 200 μl of 1% (vol/vol) ketamine (Velonarcon, Berlin Chemie, Germany) and 0.02% (vol/vol) xylazine (Rompun, Bayer Leverkusen, Germany) and intranasally infected with a 25-μl drop of a fresh suspension containing 2 × 10^7 conidia. Survival was monitored daily, and moribund animals were sacrificed by intraperitoneal injection of 200 μl of 3.2% (vol/vol) nacarone (Rhone Mérieux, Laupheim, Germany). The drinking water was supplemented with 0.5 mg of tetracycline (Sigma) per ml to prevent opportunistic bacterial infections. Mice were tested in cohorts of 10 animals. A control group (inhalation of phosphate-buffered saline [PBS]) remained uninfected to monitor the influence of the immunosuppression procedure on vitality. Survival was monitored for 16 days. Data are representative of several independent experiments.

checked for the percentage with white spores. Flanking regions of about 0.25, 0.5, 1.5, and 2.0 kb were used. Flanking sequences shorter than 500 bp did not yield any homologous integration in either the wild-type or akuB deletion strains (about 30 transformants each were observed for flanking regions of 0.25 and 0.5 kb). However, flanking regions of 1.5 to 2.0 kb yielded about 80 and 5% transformants that produced flanking regions of 1.5 and 2.0 kb). Taken together, these results suggest that deletion of the akuB ΔKU80Δ gene in A. fumigatus provides a much higher frequency of homologous recombination.

To determine a possible influence on pathogenicity resulting from deletion of the akuB ΔKU80Δ gene, the corresponding deletion mutant was tested in an animal model. The virulence of A. fumigatus strain KU80Δ in comparison to strain CEA17pyrG+, derived from its parental strain CEA17(pyrG) by reversion to uracil prototrophy, which was used as the wild type, was determined in a murine low-dose model for invasive aspergillosis as optimized by Liebmann et al. (10). The results of a representative experiment are shown in Fig. 3. They reveal no differences in virulence between the wild-type and the akuB ΔKU80Δ deletion strain. This is true for both the overall mortality and the curve shape representing the time course of survival after infection. Infection with either the wild-type or the akuB ΔKU80Δ deletion strain resulted in a mortality rate of 80% 10 days after inoculation of the conidia. These results indicate that the akuB ΔKU80Δ mutation is not causing any loss of virulence in the KU80Δ strain compared to the corresponding wild-type strain.

Taken together, our results strongly indicate that the akuB ΔKU80Δ deletion strain could be a powerful tool for high-throughput gene replacement in A. fumigatus. Once the target gene has been deleted, the wild-type akuB gene might be restored by complementation studies to allow analysis of the deletion strain in an otherwise wild-type background. The recognition of the complementation of the akuB ΔKU80Δ deletion mutation might be simple considering that the complemented strain will be resistant to MMS. Moreover, we have used a homologous auxotrophic marker pyrG gene for our transformation studies, but it is also possible to use dominant markers such as hygromycin resistance, as shown in an A. fumigatus akuB ΔKU80Δ deletion mutation background (6).

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