**Candida albicans** is the most frequent fungal pathogen, being the causative agent of both superficial and deep-seated and life-threatening mycoses. Many molecular methods have been employed in investigating species evolution and epidemiology. One of them is the genotyping developed by Mercure *et al.* (1993) and McCullough *et al.* (1999a) based on the presence or absence of the self-splicing group I intron in the large subunit (CaLSU) of rRNA genes. Strains can be differentiated into three genotypes: genotype A without the intron, genotype B harboring the intron, and genotype C possessing LSU rDNA with and without the intron in a single genome. In the last decade, several authors reported the distribution of the A, B, and C genotypes in different groups of clinical *C. albicans* isolates (Tamura *et al.*, 2001, Millar *et al.*, 2002; Karahan, 2004; Karahan and Akar, 2005; Qi *et al.*, 2005; Millar *et al.*, 2005; Girish Kumar *et al.*, 2006, Nawrot *et al.*, 2004; Nawrot *et al.*, 2008). Although the data obtained by the particular authors differ in detail, genotype A has been reported as prevalent in most groups of clinical isolates, including those regarded as invasive. It is supposed that intron-containing genotypes can be eliminated because of their high susceptibility to some drugs, for example 5-fluorocytosine, pentamidine, and bleomycin, interfering with the self-splicing process (Mercure *et al.*, 1993; Zhang *et al.*, 2002; Jayaguru and Raghunathan, 2007). On the other hand, some authors observed increased occurrence of genotype C (McCullough *et al.*, 1999b; Gurbuz and Kaleli, 2010). Recently, Karahan and Akar (2005) found significant differences in the LSU rDNA sequences of genotype A isolates and constructed an RFLP-based method for differentiating genotype A into eight subtypes. In this communication we present the results of the RFLP analysis of LSU rDNA of *C. albicans* isolates, which is complementary to our previous study on the distribution of CaLSU among *C. albicans* isolated from blood and the respiratory tract (Nawrot *et al.*, 2008).

The study was performed on 95 isolates of *C. albicans*, including 55 blood isolates obtained from BCCM/IHEM (30 genotype A, 20 B, and 5 C) and 40 isolates (32 genotype A and 8 B) from the laboratory collection of Wroclaw Medical University and originating from different clinical samples (22 from sputum or pharyngeal swabs, 7 from blood, and 8 from body fluids or pus). Genomic DNA of the tested yeasts was extracted using the CTAB method according to O’Donnell *et al.* (1997). The PCR assay was performed with the primer pair CA-INT-L (5’-ATA AGG GAG TGC GGC AAA ATA GAT CCG TAA-3’) and CA-INT-R (5’-GCT TGG CTG TGG TTT CGG TAA GAT TAA-3’), described previously by McCullough *et al.* (1999b). DNA samples were denatured at 94°C for 3 min before 30 cycles of 94°C for 1 min, 65°C for 1 min, 72°C for 1 min, and a final extension step at 72°C for 5 min.

**Key words:** *Candida albicans*, group I self-splicing intron, rDNA genotyping, RFLP

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**Abstract**

The study presents an analysis of the restriction pattern of rDNA fragments of 95 *C. albicans* isolates previously classified on the basis of the presence of the intron in rDNA into genotypes A (62 isolates), B (28), and C (5). Most isolates (61) with genotype A were classified as “subtype a” and one as “subtype d” (Karahan and Akar; 2005). No differences were observed in the restriction patterns of the tested genotype B isolates. Similarly, most genotype C strains (4/5) showed the same restriction pattern. The results indicate low subtyping variations of the analyzed isolates, which is in contrast to published data obtained from a Turkish collection of yeasts.

**Key words:** *Candida albicans*, group I self-splicing intron, rDNA genotyping, RFLP

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for 1 min, and 72°C for 4 min, with a final extension at 72°C for 4 min following the last cycle. The PCR products were digested, separately overnight with the enzymes Hae III (BsuRI) and MspI (HpaII, Fermentas), in accordance with the protocol of Karahan and Akar (2005). The undigested and digested PCR products were analyzed by electrophoresis in 4% agarose gel and visualized in UV after staining with EtBr. The selected DNA samples were additionally analyzed by capillary electrophoresis in the QIAxcel system (Qiagen) using a QIAxcel DNA High Resolution Kit, the QX Alignment Marker 15-bp/3-kb, and the QX DNA Size Marker FX 174. The tests were performed with the help of Biocalculator software and presented as both simulated bands on gel images and peaks in electrophoregrams.

PCR with the primer pair CA-INT-L/CA-INT-R and DNA of genotype A resulted in a single PCR product of ~460 bp (McCullough et al., 1999b),

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Fig. 1. Electrophoretic gel image of PCR products digested with Hae III (A) and MspI (B). M-QX DNA Size Marker FX 17.

Lines 1–5 strains with genotype C (IHEM 19482, 19076, 19265, 19491, 19608), line 6 genotype B (IHEM 19651), line 7 genotype A subtype d (1228, laboratory collection), lines 8 and 9 genotype A subtype a (IHEM 19069, 19144). Performed with the help of Biocalculator software (Qiagen).
which can be slightly different in particular subtypes (Karahan and Akar, 2005). The strains of genotype A tested in this study give PCR products of typical size, except for one isolate (no 1228, laboratory collection), which gave a larger product (~470 bp). Most isolates (61/62, 98%) of genotype A showed the same RFLP pattern, namely three fragments of 294, 92, and 72 bp, after digestion with Hae III and two fragments of 289 and 171 bp after digestion with MspI (Figs. 1A and B). This result corresponds well with “subtype a” described by Karahan and Akar (2005), which was characterized by three bands of 296, 93, and 71 bp after digestion with Hae III. Isolate no 1228 showed a different restriction pattern, consisting of two bands of ~400 and 72 bp with enzyme Hae III and three bands (290, 93, and 84 bp) with MspI, which is in accordance with the subtype classified as “subtype d” by Karahan and Akar.

In their study, Karahan and Akar tested 144 genotype A isolates obtained from different clinical samples from three Turkish hospitals and 52% of them were identified as “subtype a”, whereas the other strains were distributed among seven different subtypes. In a recently published paper, Gurbuz and Kaledi (2010) found 84 (84.8%) “subtype a” samples among 99 genotype A isolates. The genotype A isolates tested in this study were highly homogenous (98% with “subtype a”). Analysis of the results obtained by us and by other authors, indicates that the level of diversity of LSU rDNA varies in particular C. albicans populations and may be geographically related. It is interesting that “subtype d” identified in this study, as well as 9 isolates with “subtype d” described by Karahan and Akar (2005), were obtained from blood, which may suggest a high invasiveness of “subtype d”. A future study performed with a higher number of invasive and non-invasive isolates can be helpful in verifying such an hypothesis.

In this study we also performed the RFLP analysis of the PCR products obtained for genotypes B and C. The public PubMed database (http://www.ncbi.nlm.nih.gov/sites/entrez?db=pubmed) contains only two sequences (accession nos. DQ465844 and DQ465845) of LSU of C. albicans genotype B which include the DNA fragment flanked by the CA-INT-L and CA-INT-R primers. The restriction analysis of these sequences performed with the use of SeqBuilder Lasergene software indicated the same pattern with MspI (four bands of 310, 225, 169, and 131 bp) and two different patterns with Hae III (398, 242, 93, 71, 31 bp for DQ465844, and 248, 242, 182, 93, and 71 bp for DQ465845). This finding suggests sequence diversity in the analyzed LSU fragment of genotype B and the usefulness of the Hae III enzyme in its testing. The results obtained experimentally differed from those predicted on the basis of an analysis of the reference sequences. In our study, all the investigated strains with genotype B showed the same PCR product size (~840 bp) and the same restriction pattern with enzymes Hae III (six bands of 251, 239, 101, 90, 68, and 35 bp) and MspI (540, 170, and 130 bp). It is worth noting that there was some discrepancy between the results obtained by capillary electrophoresis (the above pattern) and classical gel electrophoresis, which cannot easily distinguish the bands of ~239, 90, and 35 bp after digestion with Hae III (data not shown). What is interesting is that replacing the intron present in DQ465844 by the sequence of the intron from another strain (accession no X74272.1) resulted in changing the restriction pattern to one more similar to our finding for genotype B, namely seven bands of 247, 242, 103, 93, 72, 32, and 20 bp for Hae III and three bands of 539, 170, and 131 bp for MspI (Fig 1B). An analysis of the DQ465844 and DQ465845 sequences performed after excluding the intron indicated their high similarity (99.8%) to “subtype a”. This suggests that the strains with genotype B tested in this study can also be related to “subtype a” and their intron to the X74272.1 sequence.

Genotype C generates two amplicons, one of ~460 bp and one of ~840 bp, so the PCR-RFLP result could reflect the sequence diversity of the two products. The five strains of genotype C tested in this study displayed the typical sizes of the PCR products and the same restriction pattern with MspI (four bands of 580, 290, 170, and 130 bp), whereas with Hae III, four strains showed the same pattern (five bands of 290, 251, 239, 90, and 70 bp), but the fifth differed from them by the presence of an additional band of ~400 bp (Fig. 1A). The evolutionary processes which
resulted in the formation of the heteroallelic genotype C were discussed by many authors. McCullough et al. (1999) proposed two hypotheses of genotype C development. The first assumes that genotype C arises after losing the intron by genotype B and the second is that genotype C is formed due to the acquisition of the intron by genotype A, most probably as a result of sexual recombination, which has not yet been detected in C. albicans. Recently, Miletti-Gonzalez and Leibowitz (2008) studied the genetic arrangement of the CaLSU intron in the rDNA of an isolate with genotype C and showed that intron-possessing rDNA and intron-less rDNA copies are arrayed in tandem and adjacent to each other, forming rDNA clusters present in two R chromosomes. The authors observed high variability in the number of rDNA complex copies among clones of genotype C. The simple RFLP experiment performed in this study showed that the diversity of genotype C described above can be wider due to variability in the rDNA sequence.

In summary, the presented results indicate a high homogeneity of the analyzed fragment of LSU rDNA in the clinical isolates of C. albicans, in contrast to published data obtained with a Turkish collection of yeasts. The restriction analysis of the amplicons obtained after A, B, and C genotyping is a simple and reproducible method enabling broader strain characteristics and can be useful in epidemiological and evolutionary studies.

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Literature


