MYCOLOGY

Correlation of SfiI macrorestriction endonuclease fingerprint analysis of Candida parapsilosis isolates with source of isolation

E. PONTIERI, L. GREGORI, M. GENNARELLI*, T. CEDDIA, G. NOVELLI†‡, B. DALLAPICCOLA*‡, FLAVIA DE BERNARDI§ and G. CARRUBA

Department of Experimental Medicine, University of L’Aquila, L’Aquila, *Department of Public Health and Cell Biology, Tor Vergata University of Rome, †Chair of Human Genetics, Catholic University of Rome, ‡Chair of Genetics, University of Urbino and §Laboratory of Bacteriology and Medical Mycology, Istituto Superiore di Sanità, Rome, Italy

SfiI macrorestriction digests from whole chromosome DNA preparations of 46 isolates of Candida parapsilosis from vaginal (20 isolates), blood (23 isolates) and soil (three isolates) sources were examined by CHEF-MAPPER pulsed-field electrophoresis. The isolates were grouped into nine macrorestriction endonuclease fingerprint (MEF) classes according to the number or size of the macrorestriction fragments, or both. The electrophoretic karyotype (EK) was also examined and found to contain 18 karyotypic classes (named A–R). A comparison between SfiI MEF and EK demonstrated that the former correlated much better than the latter with the source of C. parapsilosis isolates. Five SfiI classes (I–V) contained only vaginal isolates (or vaginal and three soil isolates, class I), and the blood isolates were distributed between four classes (VI–IX). This relationship was less evident with the EK classes as several of these were composed of both vaginal and blood isolates (B, G, L and M). The three soil isolates were in class A which also included one vaginal isolate. We conclude that SfiI macrorestriction endonuclease patterns seem to be useful in discriminating among C. parapsilosis isolates, with apparent association with source of isolation.

Introduction

Candida parapsilosis has become increasingly important as a prevalent yeast species in fungaemic episodes [1–3] affecting neutropenic and severely debilitated patients receiving parenteral nutrition [4, 5]. Recent evidence also points to a role for C. parapsilosis in human vaginitis [6]. Vaginal isolates of this species produce one or more aspartylproteinase enzymes in vitro and in vivo. It has been suggested that these are involved in the pathogenesis of vulvovaginitis [6]. Experimental pathogenicity for cyclophosphamide-immunodepressed mice demonstrated the ability of vaginal and blood isolates, but not soil isolates, to cause systemic infections in the mouse [7, 8]. The above-mentioned differences suggest that different biotypes exist among C. parapsilosis isolates, whose identification could be important for pathogenicity, diagnostic and epidemiological studies.

In C. albicans, molecular approaches for epidemiological studies based on DNA analysis have been extensively developed. Restriction fragment length polymorphism (RFLP) analysis [9] particularly concerning mitochondrial [10] and ribosomal DNA [11], and analysis of various repetitive, dispersed genomic DNA sequences have been reported [12–14]. Electrophoretic karyotype analysis (EKA) techniques have also been developed and applied [15–17] by means of several pulsed-field gel electrophoresis (PFGE) systems [18–20]. All these techniques demonstrated the heterogeneity and genetic diversity of C. albicans isolates, but they did not demonstrate any correlation with isolation source or pathogenicity. In C. albicans, DNA translocations have been suggested to be involved in chromosome-length polymorphism (CLP) [21]. In the absence of a sexual cycle, these translocations are hypothesised to be a potential source of genetic variability.

Although considerably less genetic information is available concerning C. parapsilosis isolates, available data point to genomic intraspecies diversity. In a previous study, whole-chromosome DNA preparations of 16 isolates of C. parapsilosis from different sources (soil, vaginitis and candidaemia) were examined [22].
The patterns of bands obtained after digestion with the restriction endonucleases Eco RI, Hind III, Bam HI, Bgl II, Hpa II, Pvu II, Hinf I did not discriminate among isolates. In contrast, electrophoretic karyotypes generated by CHEF showed remarkable heterogeneity, resulting in the definition of seven karyotypic classes on the basis of the number or size of chromosomes, or both [22], in accord with the findings of other authors [18, 23, 24]. Clinical isolates were found in separate karyotypic classes.

In the present study, Sfi I and Not I (rare cut-site endonucleases) were tested as an approach to DNA fingerprinting of human isolates of C. parapsilosis (vaginal and blood) and from soil.

Materials and methods

Yeasts

Forty-six isolates of C. parapsilosis obtained from soil or from vaginitis or candidemic patients were analysed [8, 23]. All isolates were identified according to established laboratory procedures as follows: morphological analysis on corn meal agar, germ-tube formation in serum, carbon source assimilation (API) profile, and agglutination tests with anti-Candida sera (Candida check Kit, Iatron Laboratories Inc., Tokyo, Japan). Vaginal isolates were from patients with vaginitis or from asymptomatic carriers.

Sample preparation

Several colonies from each C. parapsilosis isolate were grown separately to stationary phase (as assessed by measurements of optical density of cultures) in 5 ml of YPD medium (dextrose 2% w/v, Bacto-peptone 2% w/v, yeast extract 1% w/v, Difco) at 30°C in 15 ml Falcon 2099 tubes in a rotatory shaker (Gallenkamp) set at 200 rpm. The cells (in pure yeast form) were harvested and washed in solution A (1.2 M D-sorbitol and 20 mM EDTA, pH 8). The pellet was resuspended at a concentration of 10⁵ cells/ml in solution B (solution A with the addition of 20 mM β-mercaptoethanol) and incubated at 37°C for 10–15 min. Then low-melting-point agarose (BioRad) 1.5% w/v, dissolved at 65°C in solution B, was added at a final concentration of 0.75% w/v and samples were solidified at 4°C. Then, the agarose blocks were overlaid for 2 h at 37°C with solution C (1.2 M D-sorbitol, 20 mM EDTA, 10 mM Tris-HCl, pH 7.5, and Lyticase, Sigma, 0.1 mg) for spheroplast formation. After discarding solution C, the agarose blocks were lysed in solution D (sodium dodecyl sulphate, 1% w/v 100 mM EDTA and 10 mM Tris-HCl, pH 7.5) at 37°C overnight. Finally, solution D was replaced by solution E (N-lauroylsarcosine 1% w/v, 100 mM EDTA and 10 mM Tris-HCl, pH 7.5) and the samples were stored at 4°C.

Karyotype analysis

Chromosome-sized bands of C. parapsilosis were separated by the CHEF-MAPPER system (BioRad) under either of two conditions as follows. Condition A: 72 h with a pulse-time from 60 s to 500 s at 120 V constant voltage, in agarose 1% w/v gel. Electrophoresis buffer was 0.5 × TBE 45 mM Tris base, 45 mM boric acid and 1 mM EDTA maintained at a constant temperature of 14°C. Condition B: 72 h with a pulse-time from 60 s to 300 s and 150 V constant voltage. Agarose gel concentration, electrophoresis buffer and its working temperature were the same as for condition A. After electrophoresis, gels were stained in a solution containing ethidium bromide 0.5 μg/ml for 30 min, destained in distilled water for 1 h and photographed with Polaroid 667 film (Polaroid Corporation).

Sfi I and Not I digestion

Agarose blocks were incubated overnight in 0.5 M EDTA, pH 8, at room temperature, then washed for 30 min with sterile distilled water three or four times. Then the samples were digested with Sfi I or Not I restriction endonucleases (Boehringer Mannheim GmbH) under the following conditions: one agarose block (82 μl volume), was placed in a 1.5-ml Eppendorf tube containing 20 μl of Sfi I or Not I 10 × digestion buffer and 40–60 units of the appropriate enzyme and adjusted to a final volume of 200 μl with sterile distilled water. Digestion was carried out at 50°C for Sfi I and 37°C for Not I for 6 h [25].

Sfi I and Not I digestion pattern analysis by PFGE

The BioRad CHEF-MAPPER system [20] was used to analyse Sfi I- or Not I-generated DNA fragments by the following parameters: two electrophoresis periods of 24 h, with switch times of 90 s and 120 s, respectively, at 150 V constant voltage in agarose gel 1% w/v. All the gels were electrophoresed in 0.5 × TBE buffer (45 mM Tris base, 45 mM boric acid and 1 mM EDTA) at a constant temperature of 14°C. Gels were stained in a solution of ethidium bromide 0.5 μg/ml for 30 min after electrophoresis, destained in distilled water for 1 h and photographed with Polaroid 667 film (Polaroid Corporation).

Results

Electrophoretic karyotype

The electrophoretic karyotypes (EK) of the 46 isolates of C. parapsilosis were examined. Several electrophoretic runs from different preparations of agarose plugs for each isolate were compared with the CHEF-MAPPER system. Electrophoresis was performed under the two sets of conditions described in Materials and methods, because of the wide range of sizes of
chromosomal bands. In fact, high- and low-range chromosomal bands were hardly separated by only one set of electrophoretic conditions. As found previously by the TAFE pulsed-field electrophoresis technique [26], the five chromosomal bands which define the high range (3–1.5 Mb) of chromosome-sized DNA bands were similar in size in all isolates of C. parapsilosis tested. By use of the CHEF-MAPPER technique these data have been confirmed for all isolates except five belonging to classes F, G, K and R (Fig. 1a). Marked isolate-dependent variations were found for smaller sized chromosomal bands (in the range 1.45–0.65 Mb). These variations allowed the grouping of isolates into 18 classes (Figs. 1b and 2). A complete association was not found between any particular karyotype class and source of isolation. Classes B, G, L and M all contained vaginal and blood isolates.

Fig. 1. Electrophoretic karyotype patterns (A and B) of the 46 C. parapsilosis isolates. The karyotype of only one isolate is shown for each class. Sizes (kb) of the most important chromosomal bands are indicated on the right.
Fig. 2. Putative sizes (kb) of chromosomal bands obtained by karyotyping of the 46 C. parapsilosis isolates grouped into 18 classes (A-R).

SfiI and NotI digestion patterns

As found in preliminary experiments, NotI digestion pattern analysis did not give discriminatory results, as all isolates tested (except HEM18) showed an identical macrorestriction endonuclease fingerprint (MEF) (data not shown). On the contrary, SfiI digestion proved to be a useful discriminator. Forty-two of the 46 isolates studied showed four similar bands in the range 2–1.6 Mb (in Fig. 3 they are compressed into a unique band, the first, depending on the electrophoretic conditions) accompanied by a number of restriction fragments below 450 kb virtually identical in all strains except the isolates in class V. However, remarkable polymorphisms were seen in the region 1.2–0.75 Mb. These polymorphisms consisted of four bands with markedly different sizes among isolates, thus allowing grouping of the 42 isolates into six classes (designated as I, II, IV, V, VI and VII) (Fig. 4). Two classes (VI and VII) included 21 blood isolates and none from other sources. Classes II, IV and V contained only vaginal isolates (eight). Finally, class I included 10 vaginal and three soil isolates. Only four isolates could not be assigned to any major class as above. They showed a very peculiar SfiI digestion pattern, characterised by c. 25–28 restriction fragments, spread all over the size range of mol. wt standards. They were grouped in class III (two vaginal isolates) and in classes VIII and IX (one blood isolate each).

Discussion

Previous studies by several groups of workers have shown that only the determination of the electrophoretic karyotype has a discriminatory power for biotyping C. parapsilosis isolates [18, 22]. DNA digestion with several restriction endonucleases (such as EcoRI, HindIII, etc.) did not generate useful biotyping patterns [22, 27]; each enzyme tested produced, for all isolates examined, an identical, highly reproducible digestion pattern. Lott et al. [23] used RAPD (randomly amplified polymorphic DNA) and TAFE (transverse alternating field gel electrophoresis) pulsed-field techniques to biotype both clinical (only blood) and non-clinical (stock collection) isolates of C. parapsilosis. They found genomic heterogeneity among their isolates, and found them characterised by phenotypic switching generating variability in electrophoretic karyotype. Branchini et al. [24] analysed blood and catheter isolates of C. parapsilosis by the CHEF pulsed-field technique and the digestion of whole-chromosomes with BssHII, a rare cut-site restriction endonuclease, together with slime production. This digestion generated polymorphisms, and the authors claimed that MEF was useful for biotyping C. parapsilosis. All these studies showed remarkable heterogeneity among isolates, confirming the widely accepted view about the biotype diversity of C. parapsilosis isolates [8, 18, 22].

In the present study 46 C. parapsilosis isolates from two clinical sources (vaginal and blood) and from soil were analysed, including isolates endowed with highly diverse morphotype and virulence characteristics [8]. To biotype these isolates, a comparative analysis of the electrophoretic karyotype and SfiI digestion pattern was made. As shown by several authors [18, 22–24] the determination of the electrophoretic karyotype provided an efficient, highly discriminatory biotyping system but had the disadvantage that, as shown here, most isolates had a unique pattern, unrelated to the source of isolation or pathogenicity [8]. In fact eight karyotypic classes had only one
Fig. 3. Representatives of nine proposed SfiI MEF classes obtained with the 46 isolates of C. parapsilosis. The restriction pattern of only one member for each class is shown. The classes are indicated with a roman number. The most useful sizes (kb), are indicated on the left.

Fig. 4. SfiI MEF classes of the 46 C. parapsilosis isolates for soil (■), vaginal samples (□) and blood cultures (■).

C. PARAPSILOSIS SfiI FINGERPRINTING AND ISOLATION SOURCE 177

isolate. Furthermore, the five isolates of classes F, G, K and R with an image of the karyotypic pattern were very different from the remaining isolates. Only nine classes were found by SfiI digestion, in contrast with the 18 karyotypic classes, and the two clinical sources (vaginal and blood) were completely separated from each other. Although SfiI generated 20–28 fragments, ranging in size from 50 kb to 2.2 Mb, only the four fragments in the range 1.2–0.75 Mb, under these experimental conditions, were useful for typing, and their different size alone allowed the grouping of six out of nine classes. The other three classes (III, VIII and IX) showed many more SfiI macrorestriction fragments. The isolates of these classes were the same as those showing a very distinct electrophoretic karyotype or morphotype class [8]. They were also peculiar by RAPD analysis (data not shown). It may be that these isolates are a variety of C. parapsilosis. Of particular interest is the fact that the SfiI restriction-based classes were closely associated with the clinical source of the isolates. All vaginal isolates belonged to classes (I–V) that did not include any
blood isolates (classes VI–IX). There was only a limited overlap between the three soil isolates and 10 vaginal isolates (class I). Of additional interest is the fact that, of the 23 blood isolates, 17 (74%) belonged to one class (VII) whereas the vaginal isolates showed a much broader distribution (into five classes).

Branchini et al. [24] grouped their 31 blood isolates into four classes by BssHII MEF. They also found a major BssHII class (A) with 23 (74%) isolates of 31. Similar results were obtained with SfI enzyme in the present study for the 23 blood isolates, assigned to a total of four classes and with class VI containing 17 isolates, i.e., 74% of total blood isolates. This suggests that the blood isolates may be genetically more homogeneous than the vaginal strains. However, no additional comparisons between these data and those of Branchini et al. [24] are possible because 20 vaginal and three soil isolates were also included in the analysis, whereas no isolates from such sources were studied by Branchini et al. [24]. Moreover, our analysis was based on the use of SfI endonuclease whereas a different restriction endonuclease was used by Branchini et al. [24]. Overall, the approach described here seems to give more comprehensive coverage of the various DNA biotype characteristics of C. parapsilosis from different clinical sources.

Further studies are in progress to define the virulence properties of the different isolates of C. parapsilosis by both mucosal and systemic models of experimental infections, so as to assess possible correlations between the different SfI MEF classes, source of isolation and pathogenicity.

This work was supported by the Progetto Finalizzato CNR (Italy) ‘Biotecnologia e Biostrumentazione’ and Progetto Nazionale AIDS (Contract No 820/U). We are grateful to A. Cassone MD (Director of Istituto Superiore di Sanità, Roma), who critically read the manuscript and provided constructive suggestions, and to Professor P. Martino who supplied the blood isolates.

References


