Genome analysis of five recently described species of the CUG-Ser clade uncovers *Candida theae* as a new hybrid lineage with pathogenic potential in the *Candida parapsilosis* species complex

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Abstract

*Candida parapsilosis* species complex comprises three important pathogenic species: *Candida parapsilosis sensu stricto*, *Candida orthopsilosis* and *Candida metapsilosis*. The majority of *C. orthopsilosis* and all *C. metapsilosis* isolates sequenced thus far are hybrids, and most of the parental lineages remain unidentified. This led to the hypothesis that hybrids with pathogenic potential were formed by the hybridization of non-pathogenic lineages that thrive in the environment. In a search for the missing hybrid parentals, and aiming to get a better understanding of the evolution of the species complex, we sequenced, assembled and analysed the genome of five close relatives isolated from the environment: *Candida jiufengensis*, *Candida pseudojiufengensis*, *Candida oxycetoniae*, *Candida margitis* and *Candida theae*. We found that the linear conformation of mitochondrial genomes in *Candida* species emerged multiple times independently. Furthermore, our analyses discarded the possible involvement of these species in the mentioned hybridizations, but identified *C. theae* as an additional hybrid in the species complex. Importantly, *C. theae* was recently associated with a case of infection, and we also uncovered the hybrid nature of this clinical isolate. Altogether, our results reinforce the hypothesis that hybridization is widespread among *Candida* species, and potentially contributes to the emergence of lineages with opportunistic pathogenic behaviour.
1. Introduction

Candida species are a non-monophyletic group of yeasts that include important human pathogens. Although the majority of Candida infections (candidiasis) are caused by Candida albicans, Candida glabrata and Candida parapsilosis, the epidemiology of this disease is shifting with the emergence of other pathogenic lineages. The mechanisms involved in the emergence of pathogenicity are still unknown, but hybridization has been suggested as a possible evolutionary path for the appearance of novel yeast pathogens. Indeed, in the C. parapsilosis species complex, at least six natural independent hybridization events have occurred, leading to four C. orthopsilosis and two C. metapsilosis hybrid lineages with the potential to infect humans. While one of the two parental lineages for C. orthopsilosis hybrids has already been isolated, also in the clinical setting, for C. metapsilosis none of the three inferred parents has been found thus far. The absence of these parents from clinical isolates has led to the hypothesis that they are possibly non-pathogenic and may thrive in the environment, and that these hybridization events contributed to the emergence of the pathogenic lineages identified as a source for candidiasis.

Although most research studies tend to focus on Candida clinical isolates, Candida species, including the strains affiliated to the C. parapsilosis species complex, are commonly isolated from various environmental niches, such as soil, plants and the digestive tract of phytophagous insects. Therefore, it is not surprising that these yeasts are able to assimilate various hydroxyaromatic compounds. Aromatic compounds, such as hydroxybenzenes and hydroxybenzoates, are metabolized in the 3-oxoadipate or in the gentisate pathways. However, even closely related species can grow on different hydroxyaromatic substrates and vary in their inventory of corresponding biochemical pathways. For example, C. parapsilosis utilizes dihydroxybenzenes (hydroquinone, resorcinol), mono- and dihydroxybenzoates (4-hydroxybenzoate, 2,4-dihydroxybenzoate, protocatechuate) in the hydroxyhydroquinone (HHQ) variant of the 3-oxoadipate pathway. In addition, this species also metabolizes 3-hydroxybenzoate and gentisate in the gentisate pathways.

In contrast, the growth of the other two members of the C. parapsilosis species complex (C. orthopsilosis and C. metapsilosis) on 2,4-dihydroxybenzoate and the gentisate pathway substrates is impaired. Similarly, C. albicans does not possess the gentisate pathway and it is unable to assimilate hydroxybenzoates. However, this yeast metabolizes phenol and dihydroxybenzenes including catechol using two branches (catechol and a shorter version of HHQ) of the 3-oxoadipate pathway. Recently, another CUG-Ser clade species, Candida subhashii, has been shown to possess the gentisate pathway as well as both branches of the 3-oxoadipate pathway, which likely corresponds to the ancestral inventory of biochemical pathways for hydroxyaromatic substrate utilization.

Another peculiarity of the C. parapsilosis species complex is the molecular architecture of the mitochondrial DNA. Unlike most other yeasts, C. parapsilosis, C. metapsilosis and C. orthopsilosis contain a linear mitochondrial genome, although circularized mutants formed by end-to-end fusions of the linear DNA molecules were identified in several strains of C. metapsilosis and C. orthopsilosis. The linear mitochondrial DNA molecules terminate at both ends with arrays of long tandem repeats (i.e. n × 738 bp in C. parapsilosis sensu stricto). These mitochondrial telomeres are replicated via a telomerase-independent pathway, reminding the alternative mechanism of telomere maintenance operating in a subset of human cancer and immortal cell lines.

Candida jiufengensis, Candida pseudojiufengensis, Candida oxyjetonae, Candida margitis and Candida theae are five recently described species isolated from the environment. While C. jiufengensis, C. pseudojiufengensis, C. oxyjetonae and C. margitis were isolated from the gut of flower beetles in China, C. theae was described from isolates obtained from modern tea drinks and ancient chicha fermentation vessels in Taiwan and Ecuador, respectively. As preliminary phylogenetic analyses placed all of these species as close relatives to the C. parapsilosis species complex, we hypothesized that some could correspond—or be closely related—to the missing parental lineages of the above-mentioned hybrids. Moreover, we considered that their study could provide additional insight into the metabolic diversity of the C. parapsilosis species complex and into the evolutionary emergence of the linear genome in mitochondria and the mechanism for maintenance of its telomeres. As none of their genomes were available at the start of this project, we sequenced, assembled and analysed the genome of the type strains of these species. Our results showed that these lineages do not correspond to any of the parents of C. orthopsilosis and C. metapsilosis hybrids but, interestingly, revealed an additional hybrid lineage in the C. parapsilosis species complex: C. theae, which was recently shown to have pathogenic potential. We also show that similarly to C. metapsilosis, C. orthopsilosis and C. parapsilosis, the mitochondrial genomes of C. margitis and C. theae are linear and terminate with arrays of long tandem repeats making this mitochondrial genome architecture a typical feature of the C. parapsilosis species complex.

2. Materials and methods

2.1. Genomic DNA sequencing

Genomic DNA paired-end sequencing was performed for the type strain of C. jiufengensis (CBS10846), C. pseudojiufengensis (CBS10847), C. oxyjetonae (CBS10844), C. margitis (CBS14175) and C. theae (CBS12239), as described in Mixão et al. For C. theae a mate-pair library was also sequenced. To this end, DNA was fragmented to sizes between 1 and 20 kb using a transposase that binds to the ends of the linear DNA molecules. Strand displacement was performed to ‘repair’ the nicks left by the transposase. Fragment sizes of 3–6 kb were then selected on a 0.8% agarose gel and were then circularized. Non-circularized DNA was removed by digestion. The circular DNA was then mechanically sheared to fragments of ~100 bp to 1 kb and the fragments containing the biotinylated ends were pulled down using magnetic streptavidin beads and submitted to a standard library preparation. A final size selection on 2% agarose gel was done and fragments of 400–700 bp were selected for the final library. Final libraries were analysed using Agilent High
Sensitivity chip to estimate the quantity and check size distribution and were then quantified by qPCR using the KAPA Library Quantification Kit (ref. KK4835, Kapa Biosystems) prior to amplification with Illumina’s cBot. Libraries were sequenced 2 × 125 bp on Illumina’s HiSeq 2500.

2.2. Candida theae public data
For the analysis of the C. theae clinical isolate, publicly available data were retrieved from the NCBI database. Specifically, we downloaded the Illumina raw sequencing reads and the genome assembly under the accession number PRJNA600777.24 For loaded the Illumina raw sequencing reads and the genome assembly data were retrieved from the NCBI database. Specifically, we down-

Illumina’s HiSeq 2500.

Read mapping was performed with BWA-MEM v0.7.15.36 Picard 2.4. Read mapping and variant calling v0.36.25 Mate-pair reads were filtered with NxTrim v0.4.1-size below 31 bp and for the presence of adapters with Trimmomatic sections/fastqc/). Paired end reads were filtered for quality below 10 or FastQC v0.11.5 (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/). Paired end reads were filtered with IGV version 2.0.30.37 Mapping coverage was deter-

index file and obtain the mapping statistics. The mapping was v2.1.1 (http://broadinstitute.github.io/picard/) was used to sort the as model organism. 33 KAT and BUSCO v3 (Ascomycota database)27,34 were used to estimate the assembly completeness. The best assembly for each species was chosen based on the assembly completeness, genome size, N50 and number of scaf-

dreds. Functional annotation was performed with eggNOG-mapper web-server using the default settings.35

2.3. Genome assembly
Raw sequencing data from all studied strains were inspected with FastQC v0.11.5 (http://www.bioinformatics.babraham.ac.uk/proj ects/fastqc/). Paired end reads were filtered for quality below 10 or size below 31 bp and for the presence of adapters with Trimmomatic v0.36.25 Mate-pair reads were filtered with NxTrim v0.4.1-53c2193,26 After filtering, only reads clearly identified as paired-end or mate-pair by the respective programs were used for subsequent analysis. The K-mer Analysis Toolkit (KAT27) was used to count k-mer frequency and estimate the expected genome size.SOAPdenovo v2.0428 and SPAdes v3.9 in both SPAdes and dipSPAdes modes29,30 were used separately to perform the genome assembly. Redundant contigs were removed with Redundans v0.13c.31 The quality of the different assemblies was inspected with Quast v4.5.35 Genome annotation was performed with Augustus v3.5 using C. albicans as model organism. 33 KAT and BUSCO v3 (Ascomycota database)27,34 were used to estimate the assembly completeness. The best assembly for each species was chosen based on the assembly completeness, genome size, N50 and number of scaf-

folds. Functional annotation was performed with eggNOG-mapper web-server using the default settings.35

2.4. Read mapping and variant calling
Read mapping was performed with BWA-MEM v0.7.15.36 Picard v2.1.1 (http://broadinstitute.github.io/picard/) was used to sort the resultant file by coordinate, as well as, to mark duplicates, create the index file and obtain the mapping statistics. The mapping was inspected with IGV version 2.0.30.37 Mapping coverage was deter-

mined with SAMtools v0.1.18.38 Samtools v0.1.1838 and Picard v2.1.1 (http://broadinstitute. github.io/picard/) were used to index the reference and create its dictionary, respectively, for posterior variant calling, GATK v3.639 was used to call variants with HaploxyreCaller set with –genotypings_-mode DISCOVERY -stand_emit_conf 10 -stand_call_conf 30 -ploidy 2 -nct 8. The tool VariantFiltration of the same programme was used to filter the vcf files with the following parameters: -clusterSize 5 -clusterWindowSize 20 -genotypeFilterName ‘heterozygous’ -genotypeFilterExpression ‘isHet == 1’ -filterName ‘bad_quality’ -filter ‘MQ < 20’ || MQ < 40 || FS > 60.0 || HaploxyreScore > 13.0 || MQRankSum < -12.5 || ReadPosRankSum < -8.0’ -filterExpression ‘DP <= 20’ -filterName ‘DepthOfQuality’. In order to determine the number of Single Nucleotide Polymorphisms per Kilobase (SNPs/ kb), a file containing only SNPs was generated with the SelectVariants tool. Moreover, for this calculation, only positions in the reference with 20 or more reads were considered for the genome size, and these were determined with bedtools genomewc v2.25.0.40 Ploidy estimation was calculated with nQuire histoest,41 Average depth of coverage was calculated with Mosdepth v0.3.1.42

2.5. Loss of heterozygosity block definition
To determine for each heterozygous strain the presence of loss of heterozygosity (LOH) blocks, heterozygous and homozygous variants were separated. Then, the procedure applied and validated by Pryszcz et al.6 was used. Briefly, bedtools merge v2.25.040 with a window of 100 bp was used to define heterozygous regions, and by opposite, LOH blocks would be all non-heterozygous regions in the genome. Minimum heterozygous and LOH block size was estab-

lished at 100 bp.

2.6. Phylome reconstruction
PhylomeDB pipeline43 was used for phylome reconstruction as de-
scribed in Mixão et al.,34 considering the 28 species specified in Supplementary File S1, and using each of the five target species of this study as seed. Gene gain and loss analysis in the seed branch was performed based on the phylome results. An enrichment analysis was done using Fatigo.35 Species-tree reconstruction was based on the final concatenated alignment of 460 single genes, comprising 283,609 amino-acid positions, with RAxML v8.2.446 using the PROTGAMMALG substitution model. Gene phylogenies were used to determine the orthologs of C. albicans and C. parapsilosis genes involved in metabolic pathways.

2.7. MAT locus analysis
The MAT locus alleles of C. orthopsilosis were retrieved from the reference genome assembly (PRJEA172256).47 A BLASTp using these genes as a query was performed on a database containing the predicted protein-coding genes of C. theae and C. margitis. A ‘reciprocal best-hit’ approach was used to confidently identify the MAT loci of these species.

2.8. Mitochondrial DNA analysis
For mitochondrial DNA analysis, we retrieved the mitochondrial sequences available at NCBI for C. jiufengensis (GU13639748), C. pseudojiufengensis (KC99317949), C. oxycetoniae (KC99318749) and C. theae (KC99319548). The mitochondria of C. margitis was assessed in this study with NOVOPlasty v4.250 using C. parapsillosis COX2 as seed (NC_005253.2: c8960-8166). The mitochondrial DNA sequences were annotated using the MFannot tool (http://mega sun.bch.umontreal.ca/cgi-bin/mfannot/mfannotInterface.pl) and manually curated with the Geneious package (version R 11.1.5). Terminal sequences (mitochondrial telomeres) of the type 2 linear mitochondrial DNA were symmetrically reconstructed on both sides of the mitochondrial contigs according to the model of

C. oxycetoniae AS 2.3656 (CBS12239) were investigated in this study. In addition, for the

2.9. Yeast strains and carbon substrate assimilation tests
The type strains of C. jiufengensis AS 2.3688 (CBS10846), C. pseudojiufengensis AS 2.3693 (CBS10847), C. oxycetoniae AS 2.3656 (CBS10844), C. margitis (CBS14175) and C. theae G-17 (CBS12239) were investigated in this study. In addition, for the
screening of hydroxyaromatic substrate assimilation, we also used the type strains of C. parapsilosis CLIB214 (CBS604), C. orthopsilosis MCO457 (CBS10906), C. metapsilosis MCO448 (CBS10907) and the reference strain of C. albicans SC5314. In these tests, the yeast cultures were grown overnight at 28°C in complex liquid media [1% (w/v) yeast extract, 2% (w/v) peptone] containing 2% (w/v) glucose or 1.5% (w/v) glycerol (for cells spotted on the medium with resorcinol). The cells were washed with water and the diluted suspensions were spotted onto the plates with synthetic media [0.17% (w/v) Yeast Nitrogen Base without Amino Acids and Ammonium Sulphate (Difco), 0.5% (w/v) ammonium sulphate, 2% (w/v) agar] containing either 2% (w/v) glucose or 10 mM hydroxyaromatic compound as a sole carbon source [i.e. 3-hydroxybenzoate, gentisate (2,5-dihydroxybenzoate), phenol, catechol (1,2-dihydroxybenzene), 4-hydroxybenzoate, 2,4-dihydroxybenzoate, protocatechuic (3,4-dihydroxybenzoate), hydroquinone (1,4-dihydroxybenzene) and resorcinol (1,3-dihydroxybenzene)]. Hydroxyaromatic compounds were dissolved in dimethyl sulphoxide as 0.5 M stocks prior addition to the medium.

2.10. Data availability

Sequencing reads, genome assemblies and the respective annotation of C. jiujiufengensis (CBS10846), C. pseudojiufengensis (CBS10847), C. oxycetoniae (CBS10844), C. margitis (CBS14175) and C. theae (CBS12239) are available at NCBI database under the BioProject accessions PRJNA726921, PRJNA726922, PRJNA726927, PRJNA748886 and PRJNA726928, respectively. The phylogeny of these species, and the corresponding orthology and paralogy relationships, are available for browsing or download at PhylomeDB53 with the IDs: 16 (C. jiujiufengensis), 404 (C. pseudojiufengensis), 960 (C. oxycetoniae), 423 (C. margitis) and 866 (C. theae).

3. Results and discussion

3.1. Genome assembly and phylome reconstruction

In this study, we performed whole-genome sequencing, assembly and annotation of the type strains of C. jiujiufengensis (CBS10846), C. pseudojiufengensis (CBS10847), C. oxycetoniae (CBS10844), C. margitis (CBS14175) and C. theae (CBS12239). We applied different assembly strategies in parallel (see Materials and methods), and the final assembly for each species was chosen based on several quality parameters, such as genome completeness, percentage of mapped reads, N50 and assembly fragmentation. The metrics of the final assembly for each species was chosen based on several quality parameters, such as genome completeness, percentage of mapped reads, N50 and assembly fragmentation. The metrics of the final assembly for each species were summarized in Table 1. Importantly, estimations based on predicted protein-coding genes and k-mer composition reported > 94% and > 99% assembly completeness, respectively, for each of the genomes (Table 1), thus showing that these are good representatives of the respective isolate and can be used for further analysis.

We next reconstructed the phylome (i.e. the complete collection of gene phylogenies51) for each of the five target species in the context of 23 additional ones (Fig. 1, Supplementary File S1). Our results revealed the existence of 194, 282, 221, 109 and 33 orphan genes in C. jiujiufengensis, C. pseudojiufengensis, C. oxycetoniae, C. theae and C. margitis, respectively. Functional enrichment analyses revealed that genes specifically duplicated in C. jiujiufengensis are enriched in aspartic-type endopeptidase activity and transmembrane transport activity, while those specifically duplicated in C. pseudojiufengensis are enriched in aspartic-type endopeptidase activity, and the ones specifically duplicated in C. margitis are enriched in transmembrane transport activity (Supplementary File S2). No enrichment was observed in C. oxycetoniae and C. theae. The tree species obtained (Fig. 1, see Materials and Methods for details) shows that, although being close-relatives, C. jiujiufengensis, C. pseudojiufengensis and C. oxycetoniae are not part of the C. parapsilosis species complex, and hence they cannot correspond to any of the parental lineages of the above-mentioned hybrids. By contrast, C. theae and C. margitis are closely related to C. orthopsilosis and C. metapsilosis and further analysis was necessary to assess their putative role in the hybridizations. The genome-wide collections of gene phylogenies and corresponding alignments, as well as the corresponding orthology and paralogy predictions with other species, constitute a valuable resource for future investigations of these species and are available for browsing or download at PhylomeDB and MetaPhoRs databases.43,53

3.2. K-mer frequency and genomic variability

We analysed the k-mer frequency patterns of each of these genomes (see Materials and methods). In this type of analysis, the k-mer frequency across read coverage is used as an indicator of the amount of genomic variability. For instance, when a haploid or highly homozygous genome is analysed, all the k-mers have similar coverage and are expected to be represented in the assembly, which is translated as a single density peak in the k-mer plot. On the other hand, when a highly heterozygous diploid genome is analysed, such a plot presents two peaks, as one corresponds to regions where both chromosomes have the same k-mer, and the other one to the regions where both chromosomes differ (two different k-mers, resulting in half coverage for each). Moreover, in this last scenario, if the assembly includes only one of the haplotypes of the heterozygous regions (as it is the case of the assemblies of this study), we expect that only half of the k-mers of the heterozygous peak are represented in the assembly.54 Our results show that C. jiujiufengensis, C. pseudojiufengensis and C. oxycetoniae have a single k-mer frequency peak, while C. margitis and C. theae seem to have two (Supplementary Fig. S1). Consistently with this finding, we identified heterozygous SNPs in all five species, but at a much higher level in the last two (Table 1). These findings support a diploid nature for all the five type isolates and they also suggest that C. margitis and C. theae are highly heterozygous and present a higher genomic variability as compared with the other species. Of note, the k-mer comparison between the five species analysed in this study and the hybrids described in the C. parapsilosis species complex, did not reveal any shared k-mer, completely excluding their role in the hybridization events previously reported in this clade.

To obtain a better understanding of the heterozygous nature of C. margitis and C. theae, we performed a closer inspection of their genomes. Contrary to what was expected for a diploid configuration, the k-mer patterns of C. margitis revealed that the first peak represents four-fifths of the coverage of the second one, and that all the k-mers are present in the assembly (Supplementary Fig. S1). Moreover, the attempt to determine the ploidy of this species with nQuire41 was inconclusive (nQuire histotest $p^2 = 0.50$). Altogether, these results suggest that C. margitis harbours chromosomes at different copy numbers and not highly heterozygous ones, otherwise, the process of removing redundant contigs would have led to the absence of some k-mers from the assembly. To confirm this hypothesis, we inspected the average coverage of each scaffold. Although we found that indeed some scaffolds have much higher read coverage than the others, the proportion of their coverage was not consistent with the proportion found in the k-mer frequency analysis.
Therefore, we are uncertain whether these two peaks in the \(k\)-mer plot may represent real copy number variations among \(C.\ margitis\) chromosomes, or a sequencing artefact resulting in coverage irregularities.

Regarding the \(k\)-mer profile of \(C.\ theae\), the two peaks of the \(k\)-mer plot are clearly associated to a diploid genome because: (i) the first peak has half coverage of the second one, and (ii) half of the \(k\)-mers of the first peak (heterozygous) are absent of the assembly.
(Supplementary Fig. S1), consistently with the assembly reduction we expect for a diploid genome. Indeed, our estimates suggest that C. theae is diploid (nQuire histotest $r^2 = 0.97$), and the genomic variability of C. theae type strain is 6.34 heterozygous SNPs/kb. Importantly, these heterozygous variants are not homogeneously distributed along the genome, but rather form blocks of high heterozygosity separated by homozygous regions (Supplementary Fig. S3). The estimated sequence divergence in the heterozygous blocks has a normal distribution (Supplementary Fig. S4), resembling the patterns previously described for Candida hybrids.

3.3. Comparison of C. theae environmental and clinical isolates

In this study, we assembled the genome of the type strain of C. theae, which, as above-mentioned, was isolated from a tea can. However, a recent publication reported the case of C. theae infection in an immunocompromised child. Taking advantage of the whole-genome sequencing data publicly available for a clinical isolate from that infection, we performed a comparative genomics analysis between the two isolates. Our results show that 98.4% of the Illumina sequencing reads of the clinical isolate aligned to the genome assembly described in this work, and that the clinical isolate presents 4.59 heterozygous SNPs/kb, thus showing an overall lower level of heterozygosity as compared with the environmental isolate (6.39 SNPs/kb). As shown in Supplementary Fig. S4, the patterns of genomic variability of the clinical isolate were also consistent with a hybrid origin for this isolate. However, in this case, we estimated 3.6% sequence divergence between haplotypes, which is higher than the 3.15% found for the type strain. Two possible scenarios could explain these discrepancies between the two isolates of the same species: (i) both C. theae isolates descend from the same hybridization event but the clinical isolate accumulated more mutations in the heterozygous blocks and experienced additional events of LOH; or (ii) the isolates descend from independent hybridization events. To determine which was the most plausible scenario, we compared the heterozygous positions in the intersection of heterozygous blocks of the two isolates (i.e. where both have the two ancestral haplotypes) and determined the number of shared alleles. Our results revealed that only 39.5% of the heterozygous SNPs of the environmental isolate and 19.7% of the heterozygous SNPs of the clinical isolate are shared between the two. On the other hand, still in regions that correspond to heterozygous blocks in both isolates (ancestral regions), 59.1% and 79.7% of the heterozygous positions of the environmental and the clinical isolates, respectively, are exclusive of one isolate, i.e. correspond to homozygous positions in the other one. These results are in stark contrast with what was previously found in C. albicans where even with a high intra-species variability, different strains shared ~80% of their heterozygous SNPs in the ancestral block. These results suggest that, although the ITS region of both C. theae isolates classify them as the same species, in fact they originated from independent hybridization events between different closely related lineages (Fig. 2). This scenario resembles what was previously described in the other hybrids of the C. parapsilosis species complex, where multiple independent hybridization events occurred in C. orthopsilosis and C. metapsilosis, showing a remarkably high hybridization propensity in this clade.

3.4. Analysis of the MAT locus

As the Mating-type (MAT) locus can provide a better understanding of the origin of the two isolates of C. theae, we inspected this region in their genome assemblies. The MAT loci of Candida species of the CUG-Ser clade can harbour a and/or alpha idiomorphs surrounded by a conserved set of genes (Fig. 2). Although the mechanism by which hybridization occurs in these species is not fully understood, hybridization can occur through mating between cells harbouring opposite mating types. This scenario is supported by the finding of both mating type idiomorphs in hybrids of C. orthopsilosis and C. metapsilosis. In accordance with this expectation is the MAT locus of the C. theae clinical isolate, which harbours both a and alpha idiomorphs, but not the one of C. theae type strain, which only has MAT a idiomorph (Fig. 2 and Supplementary Fig. S5). Further inspection of the region surrounding C. theae MAT locus shows evidence for the occurrence of a recombination event in the type strain that led to LOH in this region. Therefore, we hypothesize that both hybridizations occurred between cells harbouring different MAT idiomorphs and that a subsequent introgression event erased the alpha alleles of the type strain. Interestingly, this is not the first instance in which such a recombination event is described in hybrids of the C. parapsilosis species complex. Genomic analysis of C. metapsilosis hybrids also revealed the existence of recombination in the MAT locus, with MAT a being partially replaced by MAT alpha alleles. To the best of our knowledge, the relevance of these recombination events leading to LOH in the MAT locus of Candida hybrids is not known. Nevertheless, previous studies have shown that the disruption of the MAT locus may contribute to restoring hybrids’ fertility. Further studies, exploring the mechanisms by which Candida species hybridize and the genomic aftermath of hybridization will be essential to clarify if the same is happening in these species.

3.5. Linear mitochondrial genomes

Linear mitochondrial genomes occur in several yeast species sparsely distributed in the subphylum Saccharomycotina (e.g. in the families Debaryomycesaceae, Phaffiomycetaceae, Pichiaceae, Saccharomycodaceae, Trichomonasaceae) indicating their recurrent emergence in evolution presumably from closely related circular ancestors. This conclusion is further supported by three different types of telomeric structures present at the ends of linear mitochondrial DNAs, i.e. covalently closed single-stranded hairpins (Type 1), arrays of tandem repeats (Type 2) and a protein covalently bound to the S’ termini of DNA molecules (Type 3). The Type 2 mitochondrial telomeres found in the C. parapsilosis species complex are of particular interest as they resemble to the telomeres at the ends of eukaryotic nuclear chromosomes and their investigation provides an insight into telomerase-independent means of telomere maintenance. However, the evolutionary origin of the Type 2 linear mitochondrial genomes remains elusive. Here we show that, similarly to other three members of the C. parapsilosis species complex, C. margitis and C. theae contain linear mitochondrial DNAs of about 25 kp long terminating with tandem repeat arrays
The mitochondrial genomes of all five species from the *C. parapsilosis* species complex share the same set of genes arranged in the same order, yet they differ by the size of the mitochondrial telomere repeat unit (Table 2). *Candida jiufengensis* and *C. pseudojiufengensis* belonging to a sister lineage of the *C. parapsilosis* species complex (Fig. 1) possess circular mitochondrial genomes with rearranged gene order compared with the *C. parapsilosis* species complex. In this study, we identified the linear mitochondrial genome also in *C. oxycetoniae*, which represents a lineage branching off before the split between the *C. parapsilosis* species complex and the lineage formed by *C. jiufengensis* and *C. pseudojiufengensis* (Fig. 1). The *C. oxycetoniae* mitochondrial genome (62.6 bp) is almost twice as long as the mitochondrial DNAs of the remaining species and, in addition to canonical mitochondrial genes, it also codes for DNA (*dpo*) and RNA (*rpo*) polymerases homologous to those carried by linear mitochondrial plasmids, which possess a protein covalently bound to the 5’ ends such as pPK2 of *Pichia kluyveri*. The presence of inverted terminal repeats as well as the *dpo* and *rpo* genes located on the opposite subtelomeric regions of the

(i.e. 2n × 549 and 2n × 553 bp, respectively) on both ends (Table 2, Supplementary Fig. S6). The mitochondrial genomes of all five species from the *C. parapsilosis* species complex share the same set of genes arranged in the same order, yet they differ by the size of the mitochondrial telomere repeat unit (Table 2). *Candida jiufengensis* and *C. pseudojiufengensis* belonging to a sister lineage of the *C. parapsilosis* species complex (Fig. 1) possess circular mitochondrial genomes with rearranged gene order compared with the *C. parapsilosis* species complex. In this study, we identified the linear mitochondrial genome also in *C. oxycetoniae*, which represents a lineage branching off before the split between the *C. parapsilosis* species complex and the lineage formed by *C. jiufengensis* and *C. pseudojiufengensis* (Fig. 1). The *C. oxycetoniae* mitochondrial genome (62.6 bp) is almost twice as long as the mitochondrial DNAs of the remaining species and, in addition to canonical mitochondrial genes, it also codes for DNA (*dpo*) and RNA (*rpo*) polymerases homologous to those carried by linear mitochondrial plasmids, which possess a protein covalently bound to the 5’ ends such as pPK2 of *Pichia kluyveri*. The presence of inverted terminal repeats as well as the *dpo* and *rpo* genes located on the opposite subtelomeric regions of the
C. oxycetoniae linear mitochondrial DNA indicate that these terminal structures are derived from a linear plasmid. Although we did not investigate if a protein is present at the 5' termini of DNA molecules, we assume that the linear mitochondrial DNA of C. oxycetoniae has Type 3 telomeres. Differences in the molecular architectures of the linear mitochondrial genomes occurring in C. oxycetoniae and the members of the C. parapsilosis species complex indicate that they emerged independently in evolution. Therefore, the search for ancestral form of the Type 2 linear mitochondrial genomes will require screening of mitochondrial genomes in additional closely related species from this clade.

3.6. Assimilation of hydroxybenzenes and hydroxybenzoates

Taking advantage of the orthology and paralogy relations that were inferred with the phylome reconstruction, we identified the homologs of the genes coding for the enzymes and plasma membrane transporters of the 3-oxoadipate and the gentisate pathways, as well as the transcription factors involved in their control, in C. jiufengensis, C. pseudojiufengensis, C. oxycetoniae, C. margitis and C. theae using the queries from C. parapsilosis and C. albicans (Supplementary Fig. S7, Supplementary File S3). Our results indicate that, contrary to the catechol branch of the 3-oxoadipate pathway, the HHQ branch of this pathway occurs in all these species (Fig. 2 and Supplementary Fig. S7). However, the genes coding for key enzymes of the catechol branch of this pathway (i.e. CDX1/HQD2, MCI1, MLII1, OEL1) are present only in C. albicans. The homologs of C. parapsilosis genes coding for the components of the gentisate pathway were identified in C. margitis and C. theae. Our searches also revealed that, except for MNX2 and GTF1, C. metapsilosis holds multiple copies of these genes (Supplementary File S3). Next, we examined the growth of these species in the synthetic media containing a hydroxybenzene or a hydroxybenzoate as a sole carbon source to verify if these pathways are functional (Fig. 3). For comparison, C. albicans, C. metapsilosis, C. orthopsilosis and C. parapsilosis were also included in this experiment. Except C. albicans, all examined yeasts grow in medium with protocatechuate, which confirms that the HHQ branch of

Figure 3. Assimilation of hydroxybenzenes and hydroxybenzoates. The cells were grown overnight in a complex liquid medium, washed with water and diluted to a concentration $6 \times 10^6$ cells/ml. Serial 5-fold dilutions were spotted onto plates with synthetic media containing indicated carbon sources (see Materials and methods for details). The plates were incubated for 7 days at 28°C. The metabolic pathways involved in catabolism of corresponding substrates are indicated (i.e. the gentisate pathway and the catechol and hydroxyhydroquinone (HHQ) branches of the 3-oxoadipate pathway [3-OAP]).
the 3-oxoadipate pathway is active in these species. However, the growth of some of them is impaired on dihydroxybenzenes (C. oxy-
 cetoniae), 4-hydroxybenzoate (C. jiufengensis, C. oxyctetoniae) and
2,4-dihydroxybenzoate (C. jiufengensis, C. margitis, C. metapsilosis, C.
orthopsilosis, C. oxyctetoniae, C. pseudojiufengensis, C. theae) pointing to a possibility that the uptake of corresponding substrates is
absent or less efficient, yet functional studies need to be performed
to address this hypothesis. In contrast to C. albicans, the other exam-
ined yeasts do not utilize phenol or catechol, which confirms the con-
fusion that the catechol branch is absent in these species. Our
experiment also shows that, similar to C. parapsilosis, C. margitis and
C. theae assimilate 3-hydroxybenzoate and gentisate as carbon
sources, demonstrating that the gentisate pathway is active in these
species. Since C. metapsilosis does not grow on the gentisate path-
way substrates, we assume that at least some of the identified homo-
logs are not functional in this yeast.

3.7. Concluding remarks
In recent years, the emergence of new pathogenic lineages has shifted
the epidemiology of candidiasis. Species which so far were not con-
sidered as medically relevant are now emerging as new pathogens.
Our understanding of how new pathogens emerge is very limited.
In this context, the study of genomes from recently emerged species and
the inference of their past evolution provides a promising research
avenue. Recent studies have pointed to a role of hybridization on the
emergence of new pathogens, with different hybrid lineages being
described in clinical isolates, such as C. orthopsilosis and C. meta-
psilosis.

Importantly, most of the parental lineages of these hybrids remain unidentified, suggesting they are never associ-
ated with infections. This has led to the hypothesis that hybridization
between non-pathogenic parental species may have originated hybrid
lineages with the ability to associate with and infect humans.
As such, a hypothesis can only be corroborated with a comparative ge-
nomics and transcriptomics analysis between the hybrids and their
respective parents, we attempted to identify these lineages by se-
quencing five recently identified and closely related species, which
did not correspond to clinical isolates, namely, C. oxyctetoniae, C.
jiufengensis, C. pseudojiufengensis, C. theae and C. margitis.

Contrary to our expectations, none of these species corresponded to
any of the unknown parental lineages, thus hampering the identifica-
tion of the mechanisms underlying the possible role of hybridization
on the emergence of Candida pathogens. However, surprisingly, the
genomic patterns of C. theae type strain indicated that this species
has a hybrid origin. Furthermore, the comparative analysis of the
genome of a clinical isolate of the same species indicated that it has an
independent hybrid origin from related but different parental line-
ages. These findings reinforce the idea that lineages of the C. parapsil-
osis clade are prone to hybridize, and that many opportunistic
pathogens within this species complex are hybrids. To understand
what characteristics provide these hybrids with the ability to infect
humans, as opposed to their parents, the identification of parental
species and of hybrids non-associated to humans is necessary. For
this, it is crucial to increase our capacity to isolate and sequence new
species, including those isolated from environmental sources.

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Accession number
Sequencing reads, genome assemblies and the respective annotation of C. jiufengen-
sis (CBS 10846), C. pseudojiufengensis (CBS10847), C. oxyctetoniae
(CBS10844), C. margitis (CBS14175) and C. theae (CBS12239) are available at
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Conflict of interest
None declared.

Supplementary data
Supplementary data are available at DNASAES online.

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