# The Impact of a Codon Reassignment Event on the Evolution of the Yeast C. cylindracea

Rita Coimbra<sup>1,2</sup>, Miguel Pinheiro<sup>1</sup>, Andreia Reis<sup>1</sup>, Ana R. Bezerra<sup>1</sup>, José Luís Oliveira <sup>2</sup>, Jean-Luc-Souciet <sup>3</sup>, Manuel A.S. Santos<sup>1</sup> and Gabriela Moura<sup>1</sup>

<sup>1</sup> Department of Medical Sciences & iBiMED, University of Aveiro; <sup>2</sup> Department of Electronics and Telematics, IEETA, University of Aveiro; <sup>3</sup> Institute de Botanique, Strasbourg, France.

The genetic code is degenerate allowing for introduction of species specific codon utilization biases. These has been intensively studied but it is not yet clear whether they arise from mutational pressure alone or whether other forces, such as the translational efficiency, also influence it To address this question we took advantage of a unique genetic code alteration that occurred in the fungal Saccharomycotina subphylum, the so called CTG clade, which translate leucine CTG codons as serine due to a novel transfer RNA tRNACAGSer The atypical serine CTG codons are rarely used among the CTG clade species except for Candida cylindracea which uses CTGs at high level, allowing to gain further insight on how they were remodeled after the reassignment event For this, we conducted de novo whole genome sequencing, and annotation of C cylindracea. Alongside, we performed systematic alignment of yeast orthologous genes, phylogenetic reconstruction and codon evolution analysis The results showed that C cylindracea's genome is different from the other Saccharomycotina genomes and occupies a basal position with respect to the reassignment event. Our findings support the hypothesis that CTG codon reassignment, and not only G+C pressure, exerted a strong influence on serine/leucine codon evolution in yeast.

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Institute for Research in Biomedicine – iBiMED & Department of Medical Sciences, University of Aveiro

cInstitute de Botanique, Strasbourg, France



## Abstract

allowing for introduction of speciesspecific codon utilization biases. advantage of a unique genetic code Saccharomycotina subphylum, the so-called CTG clade, which translate leucine CTG codons as serine due transfer RNA codon evolution in yeas

#### Introduction

Now we know that the yeast CTG reassignment event has a polyphyletic nature [1] and erased thousands of codons from the genome of the CTG clade ancestors [2]. As a consequence, these alpipical serine CTG codons are rarely used among the CTG clade species, and the RIRNA<sub>cons</sub>Ser that decodes it is also a low abundance RRNA, normally encoded by one single gene copy. Candida cylindracea is the exception to the rule as it uses CTGs at high level (Fig.1), and has at least 3 genes for this tRNA (data not shown). We have sequenced the entire genome of C. cylindracea to gain further insight on how and why the usage of this codon diverged so dramatically between C. cylindracea the other CTG clade species.

### Methodology

Whole genome sequencing and assembling was conducted at Genolevures Program, scaffolds annotation was done using MAKER [3]. To create the phylogenelic tree, we used 28 proteomes from PhylomedB together with Candida cylindraces. We ran OrthoMCL algorithm to identify the orthologous, chosing the ones with more than 50% match between pairs of proteins. Within orthologous sets we isolated groups that had hits in all species using Saccharomyces cerevisiae as root. With this method we obtained 357 groups of orthologous that were present in 29 species. We aligned these 357 groups separately with TcOffee (11.08) using Martin, Muscle and Kalign methods. We ran the Trimal (1.2rev59) in all alignments to trim regions with less than 80% match. The trimmed alignments were concatenated yelding a total sequence of 184718p. Smart Model Selection in PhylML (SMS 1.8.1) was then used to create the phylogenetic tree. SMS selects the substitution model LG '4G+I+P' with AIC=12423869.84956. The codons present in each aligned position for all species, at CTG positions, were computed through in-house made Python scripts.

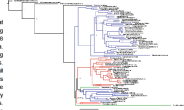
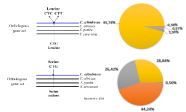


Figure 2 | The evolutionary history of yeasts, including C. cylindracea (last branch of the tree). All clades that suffered CTG reassignment are identified according to [4].



CTG ■ Serine ■ Conserved ■ Othe

Figure 3 I.CTG remodeling in C. cylindracaa. The alignments of orthologous proleins were analyzed focusing on the degree of conservation of CTG codons along the species. Interestingly, only less than 1% of C. cylindracae extant CTG codons were conserved as CTG codons in the other species (bellow), suggesting a major remodeling of CTG useape among yeasts. Neverthelesis, suggesting a major remodeling of CTG useape among vests. Neverthelesis, are conserved for serine residues (44%). When CTG positions were fixed in the conserved for serine residues (44%), When CTG positions were fixed in the code of the CTG positions were fixed in the code of the CTG positions were fixed in the code of the CTG positions were fixed in the code of the CTG positions were fixed in the code of the CTG positions were fixed in the code of the CTG positions were fixed in the code of the CTG positions were fixed in the CTG positions were compared to the CTG positions were fixed in the CTG positions were

### Conclusion

C. cylindracea prefers GC-rich codons and uses CTG as the preferred serine codon (Fig.1)

According to the phylogeny reconstruction, C. cylindracea has a basal position in relation to the other non-standard yeast decoders (Fig.2). The phylogenetic tree places C. cylindracea at the margin of the CTG-Ser decoders, closer to the Ser2 clade [4]. Although highly abundant, practically none of the CTG codons of C. cylindracea are conserved at the codon level, due to the reassignment of this codon from leucines to serines (relative to the standard decoders) and to the ambiguity of its decoding (still present in many CTG clade species).

Ongoing lines of research aim at confirming the evolutionary origin of C. cylindracea's IRNACAGSer genes and if CTG usage influences gene expression, which could explain intriguing generalized use in C. cylindracea's genome.



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