How duplicated transcription regulators can diversify to govern the expression of nonoverlapping sets of genes

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The duplication of transcription regulators can elicit major regulatory network rearrangements over evolutionary timescales. However, few examples of duplications resulting in gene network expansions are understood in molecular detail. Here we show that four Candida albicans transcription regulators that arose by successive duplications have differentiated from one another by acquiring different intrinsic DNA-binding specificities, different preferences for half-site spacing, and different associations with cofactors. The combination of these three mechanisms resulted in each of the four regulators controlling a distinct set of target genes, which likely contributed to the adaption of this fungus to its human host. Our results illustrate how successive duplications and diversification of an ancestral transcription regulator can underlie major changes in an organism’s regulatory circuitry.

Results and Discussion

Successive duplications of a LYS regulator homolog in the Candida clade

The C. albicans genome contains four homologs of LYS14, the key transcriptional regulator of lysine biosynthesis genes in the free-living, model yeast Saccharomyces cerevisiae [Ramos et al. 1988; Feller et al. 1994]. When LYS14 is deleted, S. cerevisiae can no longer up-regulate the genes coding for the lysine biosynthetic enzymes, and, as a result, the mutant strain grows poorly on media lacking lysine [Ramos et al. 1988]. Phylogenetic reconstructions (Supplemental Fig. 1) indicate that the ancestor of the LYS14 regulator underwent several successive duplications in the Candida clade lineage, which includes numerous human commensal and pathogenic species [Fig. 1A]. Indeed, extant species in the Candida clade contain two to five homologs of LYS14. In contrast, there appears to be a single copy, that of LYS14 and its orthologs, in all of the extant species that belong to the Saccharomyces clade [Fig. 1A]. The Candida and Saccharomyces clades last shared a common ancestor ~300 million years ago (Taylor and Berbee 2006).

C. albicans LYS regulators control largely nonoverlapping sets of target genes

A priori, there are two plausible explanations as to why C. albicans has four homologs of LYS14 while S. cerevisiae has a single gene: Either [1] the duplicated genes underwent “subfunctionalization” whereby each C. albicans gene now regulates a subset of lysine metabolic genes and the combined action of all four is equivalent to the one or both genes need to occur. Duplicated transcription regulator genes could undergo changes that modify the activities of the encoded proteins and/or their regulation.

To deduce the mechanisms whereby duplicated transcription regulators acquire new target gene repertoire and DNA-binding specificities, we examined a group of four closely related transcription regulators that arose by successive duplications in the lineage leading to the human commensal and pathogenic yeast Candida albicans. Using a combination of in vivo genome-wide molecular biology approaches and large-scale in vitro biochemical measurements, we show that these regulators have differentiated from one another by a combination of three mechanisms: [1] small changes in the intrinsic, monomer DNA-binding specificities; [2] different preferences for half-site arrangements; and [3] association with cofactors. It is the summation of these three mechanisms that confer on the proteins the specificity to regulate independent sets of target genes and control different aspects in the biology of the fungus. These changes in the duplicated transcription regulators led to large expansions in transcription circuitry and evolutionary novelty as they contributed to the ability of C. albicans to survive as part of the human microbiota.

Gene duplication is a major contributor to the emergence of new genetic functions in all three domains of life [Ohno 1970; Conant and Wolfe 2008]. For this process to generate new gene functions, the two resulting duplicates, which start as identical copies right after duplication, must differentiate from each other. For instance, immediately after duplication of a transcription regulator [i.e., a sequence-specific DNA-binding protein], the two resulting copies will have identical DNA-binding profiles; hence, the two will control a common set of target genes. Population genetics predicts that this redundant state will be evolutionarily short-lived [Lynch 2007]. Thus, if both copies are to be retained over evolutionary timescales, changes in

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function of the single ancestral gene or [2] at least one of the duplicated genes in C. albicans acquired new functions and no longer regulates the lysine biosynthetic enzymes. To distinguish between these two possibilities, we determined the set of genes regulated by each of the four \( LYS14 \) homologs in \( C. \) albicans by full-genome chromatin immunoprecipitation (ChIP) followed by array hybridization (ChIP–chip). Because some of the \( C. \) albicans \( LYS14 \) homologs are expressed only in a mammalian host (Perez et al. 2008; Perez et al. 2013), we ectopically expressed each regulator by placing its coding region under the control of the strong promoter \( T D H 3 \) (Nobile et al. 2008; Perez et al. 2013). Using this strategy, we determined that each of the four transcription regulators binds to ~50 target regions (Supplemental Table 1), with only partial overlap among them (Fig. 1B). None of the ChIP peaks was located in the intergenic region upstream of any lysine biosynthesis gene, consistent with our previous finding that none of the four \( C. \) albicans homologs of \( LYS14 \) is required for growth in the absence of lysine (Homann et al. 2009). Although none of them regulates lysine biosynthesis, we nevertheless retain the names originally assigned to them in the \( C. \) albicans genome annotation (Boram et al. 2005): \( LYS144, LYS143, LYS142, \) and \( LYS14 \). This nomenclature thus reflects the shared ancestry with the \( S. \) cerevisiae \( LYS14 \) gene rather than the functions in \( C. \) albicans.

**Figure 1.** Recently duplicated \( C. \) albicans \( LYS \) transcription regulators bind to largely nonoverlapping sets of target genes. (A) Cladogram depicting the phylogenetic relationships among extant species of the *Candida* and *Saccharomyces* clades. The arrows to the right of the tree represent the homologs of *S. cerevisiae* \( LYS14 \) bound in each species’ genome. Gene orthology assignments (represented by arrows of the same color) are based on synteny and the reconstructed phylogeny of the gene family (Supplemental Fig. 1). Red ovals in the branches of the cladogram represent the inferred single-gene duplication events that gave rise to the four *C. albicans* homologs. No “strict orthology” can be inferred between a particular *Candida* \( LYS \) gene and *Saccharomyces* \( LYS14 \) based on phylogenetic reconstructions (Supplemental Fig. 1) or synteny (therefore, the annotation of one of the *Candida* genes as \( LYS14 \) is misleading in this respect). The similarity in color between \( LYS144 \) and *Saccharomyces* \( LYS14 \) depicts simply the fact that the DNA sequences recognized by these homologous proteins most closely resemble one another. (B) Inferred relationships among the four \( LYS \) regulators (to the left) in *C. albicans* and the gene network (to the right) formed by the four regulators (purple, orange, blue, and green circles) and their target genes (small black circles) as mapped by ChIP. The distances separating the four Lys proteins are inversely proportional to the number of shared target genes (fewer shared targets, greater separation). Although some target genes are bound by more than one regulator, most of the targets are specific to only one of the four LYS regulators.

Candida \( LYS \) regulators show different intrinsic DNA-binding specificities

Despite their recent expansion, the ChIP results show that the four *C. albicans* \( LYS \) regulators bind to largely independent sets of target genes, raising the question of how this differentiation in target gene recognition occurred. We considered three nonexclusive models: [1] The proteins have different intrinsic (i.e., monomer) DNA-binding specificities. [2] The proteins function as dimers and have different preferences for the arrangement of their half-sites. [3] The proteins have acquired different cofactor interactions.

To address the first possibility, we used a microfluidics-based approach (MITOMI 2.0) (Fordyce et al. 2010, 2012) that measured the DNA sequence preferences of each of the four \( LYS \) regulators in vitro. The proteins belong to the zinc cluster class of regulators (for review, see MacPherson et al. 2006), which typically consist of three domains—DNA-binding, dimerization, and a large “activation” domain—all connected by linker regions of variable size (Supplemental Fig. 2A). The core 35-amino-acid DNA-binding domain (based on crystal structures of other members of the family) (e.g., see Fitzgerald et al. 2006) shows 60% amino acid identity across the four \( LYS \) proteins (Supplemental Fig. 2B) and includes six cysteine residues, all of them conserved, which are predicted to interact with two zinc atoms (MacPherson et al. 2006). For MITOMI experiments, we used truncated versions of the \( LYS \) proteins that included either the DNA-binding domain alone (\( LYS143 \)) or the DNA-binding domain in conjunction with the putative dimerization domains (\( LYS14, LYS142, \) and \( LYS144 \)); full-length proteins were not efficiently transcribed/translated in the assay. By measuring binding affinities for a library containing all possible DNA 8-mers with each of the four proteins, the MITOMI experiments revealed that each Lys protein binds to a set of DNA sequences that were similar, but not identical, to one another (Supplemental Fig. 3; Supplementary Data Set 1). For each protein, we then analyzed this pattern of oligonucleotide binding to determine a consensus binding specificity, represented here as a position-specific affinity matrix (PSAM) (Fig. 2A). The consensus sequence that we identified for each protein, GCGCA\(^T\)\(_{p}\), represents the monomer-binding sequence and resembles the monomer-binding sequences described for other regulators of the zinc cluster class (MacPherson et al. 2006). While broadly similar, these PSAMs also revealed small but significant differences in binding preferences. The MITOMI-derived motifs were significantly overrepresented in the regions bound by the \( LYS \) proteins in vivo (ChIP peaks compared with a set of random intergenic sequences) (Supplemental Fig. 4); however, differences between these MITOMI motifs per se were insufficient to explain why each of the regulators bound in vivo to a unique set of target genes (Supplemen-
We note that Lys144p shows the most distinct MITOMI-derived motif compared with the other three (Fig. 2A; Supplemental Figs. 3, 4); this motif most closely resembles the DNA sequence bound by S. cerevisiae Lys14p (Becker et al. 1998). These results indicate that there must be determinants in addition to the monomer-binding specificity that contribute to the distinct in vivo DNA-binding profiles of each Lys protein.

Candida LYS regulators bind to different arrangements of DNA-binding sites

To address the second model (i.e., the potential contribution of different arrangements of DNA monomer-binding sites to the overall affinity of the proteins), we more closely analyzed the DNA sequences enriched in each ChIP data set. The DNA sequences occupied in vivo by Lys14p and Lys144p were clearly composed of inverted repeats arranged in specific configurations that for Lys14p consisted of inverted repeats located directly adjacent to each other or overlapping by 1 nucleotide [nt] [Fig. 2A]. In contrast, the sequences occupied by Lys144p were comprised of inverted repeats separated by 5 nt [Fig. 2A]. We refer to each repeat as a half-site, which is the sequence bound by a monomer of the protein. The half-site motifs independently derived from the ChIP data closely resembled the corresponding MITOMI-generated motifs. Additional DNA-binding experiments using gel mobility shift assays confirmed that the regulators preferentially bound to the predicted arrangement of DNA sequences, the introduction of point mutations in key positions of the DNA motifs abolished this binding [Fig. 2B]. In contrast to the specific configurations of the Lys14p and Lys144p half-sites, the sequences occupied in vivo by Lys142p and Lys143p indicated that these proteins bind to flexible arrangements between a “strong” and a “weak” half-site separated by 20–30 nt for Lys142p and 3–25 nt for Lys143p [Fig. 2A]. The motifs representing the “strong” sites are in close agreement with the corresponding MITOMI-derived motif.

To experimentally determine whether the Lys proteins are able to discriminate among their different half-site arrangements by themselves, we carried out competition binding assays in which we incubated each one of the four proteins with its “preferred” arrangement of DNA-binding sequences. Upon binding, we competed each reaction with unlabeled DNA fragments containing the other regulator’s DNA-binding sites. In general, Lys14p, Lys142p, and Lys143p showed strong preferences for their specific arrangements, as described above (twofold to 20-fold higher affinity) [Fig. 3; Supplemental Figs. 5, 6]. In contrast, Lys144p showed no strong preference for its “own” site arrangement compared with the others [Fig. 3C; Supplemental Fig. 7], suggesting that additional factors must contribute to the binding profile of Lys144p in vivo.

Lys144p binds to DNA cooperatively with Mcm1p

As described above, a third model that could account for the discrimination of Lys144p in binding in vivo is the interaction with one or more cofactors. To test this possibility, we searched the Lys144p ChIP data set for additional overrepresented sequences. This analysis revealed a DNA sequence [Fig. 4A] that was reminiscent of the DNA motif recognized by Mcm1p in S. cerevisiae Lys14p (Tuch et al. 2008; Askew et al. 2011). A significant proportion of these sites is occupied by Mcm1p in vivo ($P = 3.4 \times 10^{-5}$) [Fig. 4B; Tuch et al. 2008]. Moreover, purified Mcm1 protein binds to one of these sites in vitro [Supplemental Fig. 8]. The Lys144p and Mcm1p sites are arranged in a strict configuration [Fig. 4B], suggesting that binding of one protein may facilitate or, conversely, interfere with the binding of the other. We tested this possibility and found that Lys144 protein preferentially binds to its DNA sites when the Mcm1 protein is present [Fig. 4C]; thus, the two proteins bind cooperatively. The binding of Mcm1p to promoters in combination with a second regulator is a common mechanism to regulate transcription of specific sets of genes in S. cerevisiae [for example, see Carr et al. 2004] and C. albicans (Tuch et al. 2008; Tuch et al. 2011).
Our findings illustrate the prominent role that changes in the binding specificity of transcription regulators can play in the rewiring of gene regulatory networks. While
Figure 4. Mcm1p promotes Lys144p binding to DNA. (A) Putative Mcm1p motif identified in the set of DNA sequences occupied in vivo by Lys144p. (B) Distribution of Lys144p and Mcm1p DNA-binding sites in a subset of the sequences occupied by Lys144p. Putative Lys144p sites are shown in purple (half-sites), whereas the predicted Mcm1p sites are underlined. Check marks to the right indicate whether Mcm1p has been found to bind in vivo to the respective target gene [Tuch et al. 2008]. (C) Gel shift assays carried out with one of the sequences shown in B [ORF190777] and purified Lys144 and Mcm1 proteins. The open arrow corresponds to the DNA+Lys144p-bound complex, whereas the solid arrow indicates the location of the DNA+Lys144p+Mcm1 tripartite complex.

the mechanisms uncovered here have been invoked to account for observed differences in DNA-binding preferences among members of monophyletic groups of transcription regulators [Ptashne 2004; Jolma et al. 2013; Nakagawa et al. 2013], our results demonstrate [largely through direct biochemical experiments] that it is the interplay of all three mechanisms—not any single mechanism on its own—that is responsible for the diversification of the LYS regulators. This diversification took place during the evolution of the C. albicans lineage and is closely linked to the ability of this clade to proliferate in mammalian hosts. Since a significant proportion of transcription regulators found in eukaryotic genomes arose through gene duplications [Weirauch and Hughes 2011], we propose that similar combinations of biochemical changes underlie the diversification of duplicated transcription regulators, the consequent expansion of transcription networks, and the generation of novel phenotypes.

Materials and methods

C. albicans strains

All strains used in this study are derivatives of the clinical isolate SC5314 (Noble and Johnson 2005) and are listed in Supplemental Table 2. Gene tagging and construction of TDH3 promoter-driven overexpression strains followed standard fusion PCR strategies [Noble et al. 2008; Hernday et al. 2010]. Oligos used in the study are listed in Supplemental Table 3.

Phylogenetic reconstructions

Three maximum likelihood phylogenies were generated by aligning [1] full-length amino acid sequences, [2] activation domain sequences, and [3] DNA-binding domain sequences using the software MUSCLE (Edgar 2004). Multiple phylogenetic models were then tested using RAxML (Stamatakis 2006) in order to find the best-fitting phylogenetic model according to the Akaike Information Criteria (Akaike 1973). The best-fitting models were [1] PROTGAMMAJTT, [2] PROTGAMMALG, and [3] PROTCATLG, respectively. Approximate likelihood ratio (aLR) branch supports were computed using PhyML [Anisimova et al. 2011] by first computing the aLR test statistics and then manually converting the statistic to aLRs using the following relationship: aLR = exp[aLRT/2.0]. The Candida Gene Order Browser [Maguire et al. 2013] was used to assess synteny.

Full-genome ChIP

Each transcription regulator was tagged with a GFP tag at the N-terminal end of the protein in a wild-type reference strain background. The tagged strains along with untagged controls were grown to mid-logarithmic phase in YPD at 30°C, and ChIP was carried out as described [Hernday et al. 2010] with the following modifications: GFP-tagged regulators were immunoprecipitated with an anti-rGFP polyclonal antibody (Clontech), the DNA recovered after cross-link reversal was purified with QIAquick PCR purification columns (Qiagen) and amplified using the GenomePlex complete whole-genome amplification kit (Sigma). Input and immunoprecipitated DNA were fluorescently labeled and competitively hybridized to custom full-genome oligonucleotide tiling microarrays (Agilent) as previously described [Perez et al. 2013]. MochiView (Homann and Johnson 2010) was used for data visualization, identification of binding events, and DNA motif analysis.

ChIP–chip data analysis

The microarray data were normalized using the global lowess method. A full description of the data analysis can be found in the Supplemental Material.

ChIP DNA motif analysis

Sequences of 500 nt centered on the midpoint of ~15–20 of the top-scoring peaks for each regulator were used to derive motifs in Scope (Chakravarty et al. 2007) or MochiView. The software’s default parameters were used.

MITOMI 2.0

Experiments for transcription regulator target site discovery were performed largely as published previously [Fordyce et al. 2010, 2012] and are described in the Supplemental Material.

Protein purification

The N-terminal portions of the C. albicans Lys144, Lys143, Lys142, and Lys14 proteins [amino acids 1–279, 1–142, 1–280, and 1–236, respectively] were N-terminally fused to 6His and to the maltose-binding protein, respectively, in Escherichia coli as described [ Fitzgerald et al. 2006], and purified with Ni-NTA columns (Qiagen). Protein concentrations were estimated in Coomassie blue-stained gels using known concentrations of bovine serum albumin as standards.

Gel mobility shift assays

EMSA were carried out as described previously [Cain et al. 2012].

White-to-opaque switching assay

Plate-based, quantitative white–opaque switching assays were performed as described [Miller and Johnson 2002] with the following modifications. Either three [in case of the mutant deletion strains] or five [reference strain] biological replicates were performed for each strain. Each biological replicate used cells taken from three to five independent colonies.

Accession numbers

The ChIP-chip data reported in this study have been deposited in the NCBI Gene Expression Omnibus (GEO) database under accession number GSE52203.
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