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<tr>
<td>Publisher</td>
<td>National Academy of Sciences (U.S.)</td>
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<tr>
<td>Version</td>
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Gene duplication and the evolution of ribosomal protein gene regulation in yeast

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Edited* by Eric S. Lander, Broad Institute of MIT and Harvard, Cambridge, MA, and approved February 1, 2010 (received for review October 15, 2009)

Coexpression of genes within a functional module can be conserved at great evolutionary distances, whereas the associated regulatory mechanisms can substantially diverge. For example, ribosomal protein (RP) genes are tightly coexpressed in Saccharomyces cerevisiae, but the cis and trans factors associated with them are surprisingly diverged across Ascomycota fungi. Little is known, however, about the functional impact of such changes on actual expression levels or about the selective pressures that affect them. Here, we address this question in the context of the evolution of the regulation of RP gene expression by using a comparative genomics approach together with cross-species functional assays. We show that an activator (Ifh1) and a repressor (Crf1) that control RP gene regulation in normal and stress conditions in S. cerevisiae are derived from the duplication and subsequent specialization of a single ancestral protein. We provide evidence that this regulatory innovation coincides with the duplication of RP genes in a whole-genome duplication (WGD) event and may have been important for tighter control of higher levels of RP transcripts. We find that subsequent loss of the derived repressor led to the loss of a stress-dependent repression of RPs in the fungal pathogen Candida glabrata. Our comparative computational and experimental approach shows how gene duplication can constrain and drive regulatory evolution and provides a general strategy for reconstructing the evolutionary trajectory of gene regulation across species.

stress response | comparative functional genomics | regulatory modules | expression profiling

The coordinated expression of modules of functionally related genes, such as ribosomal proteins or oxidative phosphorylation enzymes, is often conserved at great evolutionary distances (1). This is consistent with a selective pressure to conserve coordinated transcript levels to maintain functional cellular modules. Recent studies have shown that the regulatory mechanisms controlling conserved modules can diverge, most notably by switching from one regulatory system to another while preserving coregulation (1, 2). However, because previous studies have typically relied on functional expression data from only one or two species, it is unknown whether these changes in regulatory mechanisms affect the expression levels of a module’s genes at all or whether both coexpression and expression levels are conserved.

A prominent example of a conserved regulatory module is the ribosomal protein (RP) module. Genes encoding RPs are tightly coexpressed in organisms from bacteria to humans (3, 4), consistent with a selective pressure to conserve coordinated transcript levels to maintain a stoichiometric balance in ribosome assembly. The transcription factors controlling RP gene expression have changed several times since the last common ancestor of the Ascomycota fungi, which span Saccharomyces cerevisiae and Schizosaccharomyces pombe (4–6). These dramatic changes include the loss of the ancestral regulator Tbf1, the emergence of Rap1 as a key activator among the Hemiascomycota (4, 5), as well as the addition of Mem1 as a regulator in Kluyveromyces lactis (7). This phenomenon of regulatory substitutions, first identified in the RP module (4, 7), has since been recognized as a general feature of module evolution in Ascomycota, suggesting that the RP module can serve as a general model for regulatory evolution.

It remains unclear whether this regulatory overhaul has affected RP gene expression. Under normal growth conditions, RP mRNA transcripts constitute a large fraction (>30%) of the total mRNA in S. cerevisiae cells (8, 9) (Fig. S1). These genes have been shown to be strongly coexpressed in S. cerevisiae under environmental stress or nutrient limitation conditions (10, 11). In contrast, a previous study in Candida albicans failed to show similar repression under environmental stress (12), suggesting that RP expression levels may have diverged between species. However, all of the RP regulatory circuits characterized to date have a similar functional organization with some components that are constitutively localized on the RP promoter (Tbf1 in C. albicans and Rap1 in S. cerevisiae; both are also associated with telomere maintenance), and some that are regulated by nutrient- and environmental-response pathways (e.g., Ihf1 and Sfp1) (6, 13–16).

Here, we use a combined computational and experimental strategy to trace the evolution of gene regulation within the RP module. We find that an activator and a repressor that are known to control RP gene regulation in normal and stress conditions in S. cerevisiae are derived from the duplication of a single ancestral protein, followed by divergence of the repressor. This regulatory innovation coincides with the duplication of RP genes in a whole-genome duplication (WGD) event and may have been important for tighter control of higher levels of RP transcripts. To test this hypothesis, we used comparative expression profiling across six yeast species and found evidence for functional specialization between these regulators. We also show that subsequent loss of the derived repressor coincides with the loss of a stress-dependent repression of RPs in the fungal pathogen Candida glabrata and with the loss of duplicate RP genes in this species. Our study is an early example of a systematic, hypothesis-driven, functional phylogenomic study. This approach can be adopted for the study of gene regulation in a wide range of organisms and regulatory modules.

Results

Two Key Regulators of RP Gene Expression Are Paralogs Derived from a WGD Event. We hypothesized that changes in trans factors could conserve RP coexpression while diverging module expression. We therefore examined the Ascomycota gene orthologs (17) of all transcription factors previously implicated in RP gene regulation in S. cerevisiae (13–16, 18, 19) (Fig. L4 and Fig. S2). We discovered that two of these regulators, Ihf1 and Crf1 (Fig. 1B), are in fact paralogs that date to a WGD that occurred ~150 million years ago (20, 21). These genes encode two transcriptional cofactors that affect RP expression levels to maintain a stoichoimetric balance in ribosome assembly.

The authors declare no conflict of interest.

*This Direct Submission article had a prearranged editor.

Freely available online through the PNAS open access option.

Data deposition: Gene expression data is available at http://www.broadinstitute.org/~ilan/
PNAS2010.

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This article contains supporting information online at www.pnas.org/cgi/content/full/0911905107/DCSupplemental.
Evolutionary History of Ifh1/Crf1 Traces That of RP Genes. We found that Crf1 orthologs are present in most post-WGD species, such as the other sensu stricto Saccharomyces and Saccharomycodes castellii, but that it was lost from the genome of C. glabrata, another post-WGD species (unrelated to the pre-WGD species C. albicans) (22) (Fig. 1B and Fig. S2B). Remarkably, duplicate copies of the RP genes themselves are present in the same species as the repressor Crf1 (17, 23); a large number (55, or 69%) of RP genes remain in duplicate copies in S. cerevisiae and S. castellii, but very few (4, or 5%) duplicate RPs were retained in C. glabrata (17, 21–23) (Fig. 1C). When present, paralogous RP genes are highly conserved in function, protein-coding sequence (21), and regulatory program (Table S1).

Taken together, our analysis is consistent with a potential divergence in the regulation of RP gene expression post-WGD. In this model, the retained paralogous RPs in post-WGD lineages result in higher total RP transcript content, which requires a more complex control to coordinate RP repression under stress because of their increased transcriptional burden (8). Crf1 can fulfill this role by competing with Ifh1 for Fhl1 binding, thus rapidly substituting a transcriptional activator with a repressor. We further hypothesized that Crf1’s role as a repressor of RP genes in stress arose after the WGD but before the divergence between the S. cerevisiae and S. castellii lineages. If this hypothesis is correct, then we expect that, in stress conditions, S. cerevisiae and S. castellii RPs will be coordinately repressed, but C. glabrata’s RPs will not.

Comparative Expression Profiling in Three Post-WGD Species Shows That Loss of Crf1 Is Associated with Lack of RP Repression in C. glabrata. To test our hypothesis, we next compared the regulation of RP genes in stress conditions in three species from three post-WGD clades: S. cerevisiae, C. glabrata, and S. castellii. In each species, we used species-specific microarrays (Materials and Methods) to measure genome-wide mRNA expression responses under comparable growth and stress conditions (Fig. 2 and Figs. S4 and S5). We verified the presence of a stress response in each organism by three criteria: (1) change in growth rate upon treatment; (2) induction of known induced environmental stress response genes (Fig. S4A); and (3) repression of ribosome biogenesis (RiBi) genes (8, 11) (Fig. 2 and Fig. S4B). We found that heat shock resulted in the most consistent and prominent stress response across all species; for C. glabrata, we tested shock from 30 °C to both 37 °C and 42 °C, since it is a commensal human pathogen adapted to 37 °C. In normal growth conditions, total RP gene transcripts (Materials and Methods) contributed a higher fraction (∼45–50%) in S. cerevisiae and S. castellii, than in C. glabrata (∼25–30%), as expected given the presence of paralogous RP genes in these species (Fig. S1).
We found that RP mRNA levels are markedly repressed upon stress in both *S. cerevisiae* and *S. castellii*, but are unaffected by any stress condition tested in *C. glabrata* (Fig. 2 and Fig. S5). The lack of RP repression in *C. glabrata* is not merely a consequence of low basal RP expression levels under normal growth conditions (Fig. S1B). Furthermore, under stress, both *S. cerevisiae* and *S. castellii* repress IFH1 expression and induce CRF1 expression (Fig. 2). These findings are consistent with the model that Crf1 became a repressor of RP gene expression following the WGD but before the divergence between *S. cerevisiae* and *S. castellii*. Loss of this repressor in *C. glabrata* resulted in its inability to repress RP transcription levels in response to stress.

Expression Profiling of Pre-WGD Species Suggests a Model for Ifh1/CRF1 Evolution. What was the evolutionary trajectory of these changes? One possibility is neofunctionalization: the preduplication ancestor of Ifh1/Crf1 performed the role of an activator, but, following duplication, Crf1 lost its trans-activating domain and function, resulting in a repressor (Fig. 1D). An alternative is sub-functionalization either via complementary degenerative mutations or by enabling an escape from adaptive conflict within a multifunctional protein (24–27). In this scenario, the preduplication ancestor had both activator and repressor functions, which were separately assumed by the Ifh1 and Crf1 paralogs after their duplication and before the divergence of the post-WGD clades (Fig. 3B). This could have occurred either neutrally or due to selective pressure, e.g., in response to the duplication of RP genes. The fact that the pre-WGD species *C. albicans* was previously reported to lack substantial RP repression under stress (12) led us to hypothesize that the ancestral protein was only an activator.

To distinguish between the neofunctionalization and subfunctionalization models, we measured genome-wide mRNA expression profiles during stress in three pre-WGD species: *K. lactis*, *Kluyveromyces waltii*, and *C. albicans*. We found a coordinated stress-dependent repression of RP gene expression in all three species (Fig. 2). In *C. albicans*, this repression was not observed at 37 °C heat shock [consistent with a previous study (12)], but was very robust at 42 °C, consistent with the behavior in other species. Overall, these data strongly suggest that there is a conserved transcriptional pro-
gram for RP repression that predates the WGD and supports a subfunctionalization model for the evolution of Crf1.

Cross-Species Analysis of cis-Regulatory Programs Supports Our Model of Evolution Through Changes in trans Regulation. To further test our model of evolution through changes in trans factors and to explore the role of additional factors, we next examined the organization of cis-regulatory elements in RP gene promoters in each species. We first searched for overrepresented sequence motifs within the promoters of RP genes of these six species (Materials and Methods). Our analysis shows that IFHL sites (4, 5), known to be bound by Fhl1-Crf1 in S. cerevisiae, are significantly enriched in the RP gene promoters of both pre- and post-WGD species, including K. lactis, K. waltii, S. castellii, and C. glabrata (Fig. 3A and Fig. S6). Furthermore, these are colocated with the RAP1 binding site, suggestive of cooperative interactions [as previously reported (4)]. This supports the role of Ifh1 in regulating RP gene expression in each of these species. As previously shown, the outgroup species C. albicans has a different cis-regulatory organization, dominated by binding sites for Tbf1 and Cbf1 and lacking directly discernible IFHL sites (5).

Because the Ifh1/Fhl1 complex is physically associated with RP gene promoters in C. albicans (6), there are two possibilities for this discrepancy: (1) Fhl1 and Ifh1 bind indirectly through the Tbf1 protein (6), or (2) the Fhl1 protein recognizes a variant site, similar to the Tbf1 consensus (1, 4). Both cases are consistent with Ifh1’s role as a regulator of RP gene expression in C. albicans.

Fig. 3. cis- and trans-regulatory evolution of RP genes across pre- and post-WGD species. (A) Representative cis-regulatory motifs (columns) found to be enriched in RP promoters in at least one species (rows) and the enrichments of RP promoters associated with the motif in each species (dark purple, $P < 10^{-10}$; black, $P = 1$). The total number of annotated RPs in each species is denoted next to the species name. (B) A model of an evolutionary trajectory of IFH1/CRF1 functional roles. The pre-WGD ortholog of IFH1/CRF1 performed dual roles, both inducing and repressing RP expression levels in a condition-dependent manner. After duplication, each paralog specialized to act as a separate activator (IFH1) and repressor (CRF1). The repressor was subsequently lost in the lineage leading to C. glabrata.
RP promoters in all species (except the outgroup C. albicans) were also enriched for three additional sites: RAP1, HomD, and RGE (4, 5) (Fig. 3A and Fig. S6). It is not known which proteins, if any, bind to the HomD ID element (4). The MCM1 element (7) was enriched in the C. glabrata and K. lactis promoters. Because K. lactis RP-optimized repressors are strongly repressed in stress, the MCM1 element alone cannot account for the lack of RP repression in C. glabrata. This overall conserved organization in species that have diverged before and after the WGD suggests that it is unlikely that the lack of RP repression in C. glabrata results from cis-regulatory changes, rather highlighting the role of trans changes in this evolutionary event.

Finally, we examined the possible role of SpI, another transcription factor that was suggested to impact both RP (18, 19) and RIBi (18, 29) gene expression under nutrient limitation in S. cerevisiae. A recent in vitro study showed that S. cerevisiae SpI binds a motif highly similar to the RGE element (30). SpI orthologs are present in all pre- and post-WGD species (with WGD paralogs retained in C. glabrata and S. castellii). Furthermore, we found that the RGE motif is enriched in RP gene promoters across all species in our study, including C. glabrata (Fig. 3B and Fig. S6). The presence of the RGE site and the SpI proteins may explain why C. glabrata RPs are substantially repressed following glucose depletion in the diauxic shift (Fig. S7).

Discussion

In this study, we used a combined computational and experimental approach to study the evolution of RP gene regulation following a WGD event. Our results support a model of trans specialization (through subfunctionalization) in RP gene regulation in the post-WGD lineages. In this model (Fig. 3B), the pre-WGD IFH1/CRF1 ancestor was both an activator of RP expression under rich growth conditions and a repressor under stress. Following the WGD, the paralogous genes may have specialized, resulting in a separate activator (Ifh1) and repressor (Crfl). The loss of the Crfl ortholog eliminated the stress-induced repressor function in C. glabrata, thus accounting for the lack of RP repression under stress treatments in this species. Ifh1 is still functional in all species, as indicated by the enrichment of IFH1 sites in all species’ RP promoters.

What were the evolutionary factors affecting this process? One possibility is that, after the duplication of the IFH1/CRF1 ancestor, each new copy retained the function of the other copy, even though the RGE site was lost in one copy. The RGE site was lost in C. glabrata, but it is still present in all other species, as indicated by the enrichment of IFH1 sites in all species’ RP promoters.

Materials and Methods

Gene Orthologies and Phylogenetic Profiles. All gene orthologies and gene trees were calculated using the Synergy algorithm, as previously described (17). Orthologies are available from http://www.broadinstitute.org/regen/orthogroups/. Notably, many RP genes are missing from the K. warlil genome annotations but are in fact present in the genome sequences.

Strains and Growth Conditions. We used the following strains for each species: S. cerevisiae Bb32 (3), C. glabrata CBS 138, S. castellii CLIB 592, K. lactis CLIB 209, K. warlil NYCC 2644, and C. albicans SC 5314. Cultures were grown following the rich medium: yeast extract (1.5%), peptone (1%), dextrose (2%), SC amino acid mix (Sunrise Science) 2 g/L, adenine 100 mg/L, tryptophan 100 mg/L, uracil 100 mg/L, at 200 rpm in a New Brunswick Scientific Edition, New Jersey air shaker model I26R and water bath model C76. The medium was chosen to minimize cross-species variation in growth. Following the experimental treatments described below, stressed and mock-treated cultures were transferred to shaking water baths.

Heat Shock. Overnight cultures for each species were grown in 650 ml of media at 22 °C to between 3 × 10⁸ and 1 × 10⁹ cell/ml (OD600 = 1.0 for S. cerevisiae, S. castellii, and K. lactis; 0.7 for C. glabrata and 0.85 for C. albicans). The shift to the heat-shock temperatures was carried out as follows: First the overnight culture was split into two 300-ml cultures and cells from each were collected by removing the media via vacuum filtration (Nanoport). The cell-containing filters were resuspended in prewarmed media to either control (22 °C) or heat-shock temperatures (37 °C or 42 °C). Density measurements were taken approximately 1 min after cells were resuspended to ensure that concentrations did not change during the transfer from overnight media. A total of 12 ml of culture was harvested 5, 15, 30, 45, and 60 min after resuspension by quenching them in liquid methanol at −40 °C, which was later removed by centrifugation at −9 °C and stored overnight at −80 °C. Cell density measurements were repeated every 5–15 min for the first 2 hr after treatment. Harvested cells were later washed in RNAse-free water and archived in RNA-later (Ambion) for future preparations. Cells were also harvested from cultures just before treatment for use as controls.

Salt. Overnight cultures for each species were grown in 600 ml of media at 30 °C until reaching a final concentration of 3 × 10⁸ and 1 × 10⁹ cell/ml. The culture was split into two parallel cultures of 250 ml and sodium chloride was added to one culture for a final concentration of 0.3 M NaCl. Cells were harvested by vacuum filtration 5, 15, 30, and 60 min after the addition of sodium chloride and from cultures immediately before the addition of sodium chloride for use as controls (time 0 min). Filters were placed in liquid nitrogen and stored at −80 °C and were later archived in RNAlater for future use.

Hydrogen Peroxide. Cultures were grown exactly as for salt stress, except that hydrogen peroxide (H2O2) was added to a final concentration of 5 mM.

RNA Preparation, Probe Preparation, and Microarray Hybridization. Total RNA was isolated using the RNAeasy midi or mini kits (Qiagen) according to the provided instructions for mechanical lysis. Samples were quality controlled with the RNA 6000 Nano II kit of the Bioanalyzer 2100 (Agilent). Total RNA samples were labeled with either Cy3 or Cy5 using a modified version of the protocol developed by Joe DeRisi (University of California at San Francisco) and Rosetta Inpharmatics that can be obtained at http://www.microarrays.org.

Microarray Data Analysis. Between two and four biological replicates for each time point were hybridized against the mock T = 0 control on two-color Agilent 55- or 60-mer oligo-arrays in the 4 × 44 K format for the S. cerevisiae strain (commercial array; four to five probes per target gene) or the custom 8 × 15 K format for all other species (two probes per target gene). After hybridization and washing per the manufacturer’s instructions, arrays were scanned using an Agilent scanner and analyzed with Agilent’s Feature Extraction software (release 10.5.1.1). The median relative intensities across probes were used to estimate the expression values for each gene, and these median values across replicates were used to estimate the overall expression response per gene per time point.
Estimation of Absolute Transcript Abundance. To assess the absolute levels of each gene’s mRNA transcript, we summed each gene probe’s raw processed signal from the control channel of the microarray. We then confirmed that this procedure renders consistent and accurate estimations by comparing its values across multiple biological replicates and by checking its correlation to the transcript levels from recent mRNA sequencing data (9). The values were highly consistent across replicates and correlated well (R² = 0.75–0.85) with RNA-seq data (9).

Promoter Sequence Analysis. RP genes were identified for each non-S. cerevisiae species by orthologous projection using orthologs available from http://www.broadinstitute.org/regev/orthogroups (17). Promoter sequences for each RP gene were defined as 600 bases upstream of ATG and truncated when neighboring ORFs overlapped with this region. cis-regulatory motifs were discovered using the Amadeus software package (28), searching for up to 5 motifs of lengths 8–12 that are significantly enriched as compared with the background set of promoters. Motif targets were identified via the TestMOTIF software program (32) using a three-order Markov background model estimated from the entire set of promoters per genome. Motifs were then clustered according to their targets, and nonredundant motif sets were determined according to maximal coverage of the RP gene set.

ACKNOWLEDGMENTS. We thank Oliver Rando and Audrey Gasch for helpful discussions and comments on previous versions of this manuscript. We thank Leslie Gaffney for assistance with preparing figure graphics. I.W. is the Howard Hughes Medical Institute Fellow of the Damon Runyon Cancer Research Foundation and was supported by a Lawrence Summers Fellowship. This work was supported by the Human Frontiers Science Program, the Howard Hughes Medical Institute, a Career Award at the Scientific Interface from the Burroughs Wellcome Fund, a National Institutes of Health PIONEER award, the Broad Institute, and a Sloan Fellowship (A.R.).