Evolutionary rewiring of regulatory networks is an important source of diversity among species. Previous evidence suggested substantial divergence of regulatory networks across species. However, systematically assessing the extent of this plasticity and its functional implications has been challenging due to limited experimental data and the noisy nature of computational predictions. Here, we introduce a novel approach to study cis-regulatory evolution, and use it to trace the regulatory history of 88 DNA motifs of transcription factors across 23 Ascomycota fungi. While motifs are conserved, we find a pervasive gain and loss in the regulation of their target genes. Despite this turnover, the biological processes associated with a motif are generally conserved. We explain these trends using a model with a strong selection to conserve the overall function of a transcription factor, and a much weaker selection over the specific genes it targets. The model also accounts for the turnover of bound targets measured experimentally across species in yeasts and mammals. Thus, selective pressures on regulatory networks mostly tolerate local rewiring, and may allow for subtle fine-tuning of gene regulation during evolution.

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Introduction

Changes in gene regulation have been postulated to play a key role in generating the vast phenotypic diversity observed across species (King and Wilson, 1975; Wittkopf et al, 2004; Prud’homme et al, 2007). Evolutionary rewiring of regulatory networks is often driven by the gain or loss of cis-regulatory elements in gene promoters bound by sequence-specific transcription factors or chromatin remodelers (Khaitovich et al, 2006; Wilson et al, 2008; Wittkopf et al, 2008; Tirosh et al, 2009), as observed for specific factors in yeasts (Borneman et al, 2007; Doniger and Fay, 2007; Tuch et al, 2008), flies (Moses et al, 2006; Bradley et al, 2010), and mammals (Odum et al, 2007; Schmidt et al, 2010). Changes in cis-regulatory elements in genes’ promoters are associated with fine-grained regulatory ‘tinkering’ of individual genes (Borneman et al, 2007; Lavoie et al, 2010). For example, the individual target genes of the yeast regulatory factor Mcm1 have diverged significantly between three related species (Tuch et al, 2008), while maintaining its regulation of the cell cycle and mating in all three species. On the other hand, there are cases where changes in cis-regulatory elements lead to dramatic rewiring of the regulation of entire gene modules (Hogues et al, 2008; Tuch et al, 2008). For example, the transcription of ribosomal protein encoding genes in yeasts is regulated by distinct transcription factors in Candida albicans (Tbf1 and Cbf1) and Saccharomyces cerevisiae (Rap1), primarily through changes in cis-regulatory elements in promoter regions (Tanay et al, 2005; Hogues et al, 2008). Although such individual examples are instructive, they represent only anecdotal evidence of the role that cis-regulatory divergence plays across evolution. It is thus of great interest to quantitatively and qualitatively assess the extent of cis-regulatory plasticity of different regulatory DNA motifs and their associated target genes and its functional implications.

For example, consider the yeast transcription factor Gcn4. According to the occurrences of its predicted binding sites in gene promoters (motif targets), we estimate that Gcn4 potentially regulates an average of 470 target genes in each of the 20 Ascomycota fungi in which we can identify it (see below; Materials and methods). In each of these species, the set of predicted Gcn4 target genes is enriched for genes encoding enzymes involved in amino-acid metabolism,
consistent with Gcn4’s known role in the model organisms *S. cerevisiae* and *C. albicans* (Marchenko et al., 2007). Naively, this observation suggests that Gcn4 would mostly target orthologous genes across the different species. To our surprise, however, the orthologs of only 8% of all the Gcn4 targets in *S. cerevisiae* are also targeted by Gcn4 in all five closest relatives of *S. cerevisiae* in this study. Furthermore, for only 8 of the 307 genes targeted by Gcn4 in *S. cerevisiae* are the orthologs across 16 or more species also targeted by Gcn4.

Such an analysis is interesting but suffers from several potential caveats. First, computational predictions of regulatory targets based on occurrences of DNA motifs in gene promoters are noisy (Maclusa et al., 2006; Hannenhalli, 2008; Zhu et al., 2009) and have only limited reliability, as compared with experimentally measured binding (Capaldi et al., 2008). However, directly measuring the binding of a host of transcription factors across dozens of species and in all relevant conditions is still very challenging. Second, there is a lack of known regulatory DNA motifs in non-model organisms, and determining targets in one species with motifs from another (possibly distant) species can lead to both false positive and negative predictions.

To address these challenges, we developed CladeoScope (Figure 1; Materials and methods)—an unbiased phylogenetic approach to reconstruct cis-regulatory evolution. CladeoScope reconstructs regulatory networks, associating DNA motifs with putative target genes across species in a phylogeny. It starts with experimentally determined DNA motifs from a model organism, and computationally adapts them to each species. To control for noisy predictions in individual species, CladeoScope predicts reliable targets in ancestral genomes based on phylogenetic support from multiple extant species. We used CladeoScope to trace the evolutionary history of regulatory interactions of 88 regulatory DNA motifs associated with transcription factors (or groups of paralogous factors) across 23 *Ascomycota* fungi, spanning >300 million years of evolution, showing that most have conserved their cognate motifs over large evolutionary distances. While most motifs show widespread gain and loss of individual target genes, the biological processes associated with them are typically highly conserved. We reconcile these trends by a model that assumes a strong selection to conserve the overall function of a motif, and a much weaker selection for its specific target genes. The model is consistent with the number of highly conserved target genes and with observed turnover of bound target genes as determined by protein–DNA binding experiments across species, thus revealing a unifying principle of cis-regulatory evolution.

**Results**

**CladeoScope: a framework for reconstructing cis-regulatory evolution**

We developed CladeoScope (Figure 1), a computational framework for reconstructing cis-regulatory networks and their evolution across a phylogeny of species. CladeoScope relies on two assumptions. First, we assume that the binding specificities of transcription factors, represented as DNA motifs, are largely conserved, even when their specific target genes and functional roles may have substantially diverged (Wapinski et al., 2007; Tuch et al., 2008; Schmidt et al., 2010). We therefore initiate our reconstruction with DNA motifs of known transcription factors that have been experimentally determined, but without any further assumptions about conservation of their individual targets or their global functional roles. We do allow for relatively small changes in binding affinities across evolution, and thus refine those motifs in a species-specific manner (see below). Second, although predicting the target genes for a motif (motif targets) across the genome is prone to errors (Hannenhalli, 2008), we assume that targets that are conserved across several related species within a monophyletic clade provide a reliable and conservative estimate for the targets in the ancestor of the clade. Thus, for each motif associated with a known transcription factor in *S. cerevisiae* (e.g., Gcn4), CladeoScope finds its ancestral target genes in various ancestors in the phylogeny. A gene is considered to be targeted by a motif in the ancestor of a clade of species only if evolutionary analysis of the orthologous targets across the species in the clade indicated that the ancestral gene (Wapinski et al., 2007) of that clade was a target of the motif (see Materials and methods). CladeoScope then compares between the ancestral targets of different clades, allowing us to reliably track evolutionary changes across the phylum by considering the evolutionary changes between clades while filtering out spurious targets within a clade.

CladeoScope consists of four steps (Figure 1B): In step 1—Initialization—CladeoScope is initialized with known DNA motifs (position weight matrices (PWM)) from one or more model organisms in the phylogeny. It uses these initial motifs to find a set of provisional target genes for each initial motif in each species, according to the motif’s occurrences in a gene’s promoter. We do not require these provisional target sets to be evolutionarily conserved. In step 2—Species-Specific Motifs—CladeoScope takes each initial motif and its provisional target sets, and learns species-specific motifs and targets in an iterative manner. In step 3—Network Refinement—CladeoScope uses a parsimony-based algorithm to reconstruct the set of each motif’s ancestral targets for the last common ancestor (LCA) of each clade in the phylogeny (Figure 2). These inferred ancestral targets within a clade are considered reliable (Figure 2; Supplementary Figure 1). In step 4—Filtration—CladeoScope filters motifs and target genes based on their phylogenetic conservation. In particular, we define a motif as detectable in an ancestor and in each of its descendant extant species if the number of the targets in the ancestor and in each extant species is statistically significant (see details below). The algorithm iterates between steps 3 and 4 until it converges. CladeoScope’s output includes for each motif, its weight matrix in each species, the ancestors and extant species in which it is detectable, and the targets in each ancestor.

**Parsimionous phylogenetic filtering of motifs and targets**

To infer the ancestral motif targets in step 3, CladeoScope traces motif-target relations across orthologous loci. This is done separately for each ancestral gene at each ancestral position in the tree (Figure 2). To determine if an ancestral...
gene is a motif target, CladeoScope uses a parsimonious phylogenetic reconstruction approach to minimize the number of target gain and loss events (Fitch, 1971). This reconstruction explicitly considers each gene paralog derived from the same ancestor by duplication, and distinguishes a lost gene from a present gene that is not a target (see Materials and methods).

Phylogenetic filtering addresses both noisy predictions of target genes as well as DNA motifs that are ‘non-functional’ in a species or a clade (i.e., no longer act as a functional regulatory element bound by a cognate transcription factor). CladeoScope tests each motif in each species independently, based on the overlap between the motif’s putative target genes in that species and the motif’s ancestral targets in any relevant ancestor. Only motifs where the overlap is statistically significant (hypergeometric $P$-value $<0.001$, see Materials and methods) are termed ‘detectable’ in the species. Since filtering the motifs and the reconstruction of ancestral targets are dependent, our algorithm iterates between both steps. If any insignificant motifs are found in the clade (step 4), the most insignificant one is removed, and CladeoScope returns to step 3. After convergence, CladeoScope filters the motifs at the clade level, requiring that the number of inferred targets for a motif in the clade’s ancestor is statistically significant (empirical $P$-value computed by 1000 reconstructions of ancestral targets for random sets of motif targets of the same size for each species, see Materials and methods).

**CladeoScope is robust to noise in target prediction and to different parameters**

Using simulated data we confirmed that CladeoScope is highly robust to noise in target prediction for individual species and to other input variations. To assess robustness, we used hundreds of simulated evolved motif-target sets, where each simulation varied the extent and type of noise in target prediction, the size of the ancestral target set, the degree of target turnover and the topology of the species tree (960 different combinations of parameters, see Materials and methods, Supplementary Note 1). For example, when 30%
of the true targets were removed from the set of target genes provided to CladeoScope, CladeoScope has >85% sensitivity (percent predicted targets among true targets), and when 80% false targets were added in each species, CladeoScope has >80% specificity (percent true targets among predicted targets) (Supplementary Figure 1; Supplementary Note 1).

CladeoScope’s predictions are also highly robust to variation in its various parameters (Supplementary Note 1). For example, varying the threshold for the significance of a motif in a species between $10^{-5}$ to $5 \times 10^{-2}$ had little or no effect on the number of ancestral targets reconstructed per clade. Similarly, varying the threshold for conservation of a motif in a clade between 0.05 and 0.001 had little impact on the number of significant motifs per clade. Thus, evolutionary conservation within a clade—rather than parameter fine-tuning—is the main determinant of CladeoScope’s results and performance.

To examine the possibility that our relatively strict motif-target detection threshold excludes weak, yet functional, binding sites, we compared the score distribution of functional
Figure 3  Motif detectability corresponds to the phylogenetic profile of the cognate transcription factor. (A) The phylogenetic tree for species in this study. Shown is the phylogenetic tree of the 23 Ascomycota species in this study (see Materials and methods). A–L: clades in which ancestral target sets are defined; clade names are denoted next to their letter in dark blue. (B) Motif detection and transcription factor presence across the species. Shown are 88 motifs (rows) across 23 species (columns) along with a phylogenetic tree (as in a, but not shown to scale). Yellow—motif is detectable and transcription factor is conserved; yellow–gray—the motif is detectable but the transcription factor is not conserved; blue—the motif is not detectable but the transcription factor is conserved, black—the motif is not detectable and the transcription factor is absent. The fraction of the motifs inferred to be detectable up to clades D, H, and L is marked on top of the respective clades. Red line denotes the most ancestral clade in the species tree where a motif is detectable. Motif names in red denote motifs that are further discussed in the text.

but weak binding sites to non-functional sites. We identify potential candidates for weak functional sites as ones with conserved target genes in the sister species within the same clade, which are classified as non-target ('lost') in the reference species. Indeed, in 85% of the cases we tested, such 'lost' targets have a distribution of scores similar to genes that are not targets throughout the clade. Hence, lowering the threshold would not have increased our sensitivity to such weak sites (Supplementary Note 2). Nonetheless, as an additional validation, we tested the main findings using a lower threshold for motif-targets detection and found our results to be robust (Supplementary Notes 1 and 3).

Finally, as a negative control, we provided CladeoScope an input set of randomly generated motifs. Although in each species we do find targets for such motifs, CladeoScope’s phylogenetic filtering found that these motifs are not conserved (Supplementary Note 1). The only exception is in the closely related sensu-stricto Saccharomyces, where intergenic sequences have not yet had enough time to acquire sufficient mutations. We therefore do not report motifs found to be conserved only in this clade.

Systematic reconstruction of the regulatory history of 23 Ascomycota species

We applied CladeoScope to 88 DNA motifs associated with known transcription factors or groups of paralogous factors from S. cerevisiae (MacLisaac et al., 2006; Matys et al., 2006; Zhu et al., 2009) across 23 Ascomycota species, defining motif-target genes in 12 clades (A–L, Figure 3A; Supplementary website). As points for reconstruction of ancestral targets we chose clades with a large evolutionary distance between them and relatively small distances within each (Figure 3A). These clades include: the sensu-stricto Saccharomyces (four species, clade A), the Kluyveromyces (four species, clade C), the
Candida (seven species, clade G), the Hemiascomycota (Kluyveromyces, Saccharomyces, Candida clades, and Yarrowia lipolytica), 18 species, clade I), and the full Ascomycota clade (23 species, clade L). The resulting ancestral network contains 190 689 reliable motif-target connections (conserved in at least one clade), compared with 996 476 connections prior to phylogenetic filtering. For example, of the 307 predicted Gcn4 targets in S. cerevisiae, 195 pass our phylogenetic filter.

To assess CladeoScope’s performance in this phylogeny, we compared its predicted targets to those measured in vivo by chromatin immunoprecipitation (ChIP) in S. cerevisiae (Maclsaac et al, 2006) and four other species (Borneman et al, 2007; Tuch et al, 2008) (Supplementary Note 1). In most cases, using CladeoScope’s in-clade conservation increases the precision of the predicted motif targets. For instance, for the Cbf1 motif, CladeoScope reaches 80% precision rate and 50% sensitivity using the ancestral motif targets in clade A, compared with 55% and 10%, respectively, in the predicted motif targets in S. cerevisiae that are not conserved (Supplementary Note 1). These improved predictions are consistent for different thresholds for motif-targets detection in each species (Supplementary Note 1).

**Regulatory motifs are detectable across large evolutionary distances**

For most regulatory DNA motifs we could detect ancestral target genes within clades across the phylogeny (Figure 3B; Supplementary Table 1). This is consistent with our assumptions that transcription factors retain their binding specificities and that many of their target genes are conserved in closely related species. For example, ~83% of the motifs were detectable in clade D (Kluyveromyces and post-whole genome duplication (post-WGD) clades) and ~68% were detectable in clade H, including in species as remote from each other as S. cerevisiae and C. albicans. The latter include motifs involved in central metabolic and cellular processes (Figure 3B, red highlights), such as Gcn4 (amino-acid biosynthesis), Rpn4 (proteasome), and Mig1 (glucose repression). In all, 39% of motifs were detectable up to the LCA of the entire Ascomycota phylum (clade L), including those involved in cell-cycle regulation (Fkh1, Swi6-MBP1, Figure 3B) and stress response factors (Hsf1, STRE, Figure 3B, red highlights). The number of motifs detectable across the phylogeny is particularly remarkable given the substantial evolutionary distances, the large intra-species divergence within the Schizosaccharomyces (Rhind et al, 2011), and the fact that as many as 25% (102 of 392) of the transcription factors in S. cerevisiae do not have a clearly identifiable ortholog in Schizosaccharomyces pombe (Wapinski et al, 2007).

The phylogenetic profiles of transcription factors largely correspond to the detectability of their cognate motifs, supporting our reconstruction. In most cases (73%), detectable motifs and factors are co-conserved (Figure 3B): when a motif is detectable in a species, the ortholog of its known cognate factor is present in the same species, and vice versa. The few cases where there is discrepancy are due to either evolutionary innovations or limitations in ortholog mapping or motif detection (Supplementary Note 2).

**Rapid target turnover for conserved motifs during evolution**

To assess changes during the evolution of regulatory networks, we first calculated the amount of turnover events for each of the 88 regulatory motifs as the number of target genes gained or lost at each clade since its direct ancestral clade. Overall, there is an extensive and rapid turnover of motif-target genes. This high turnover of targets is apparent even for broadly conserved motifs with ancient ancestral targets, such as Gcn4 and Fkh1 (Figure 4A and B). For example, less than half of the targets of Gcn4 in clade D (the LCA of pre- and post-WGD species) remained as Gcn4 targets in its two daughter clades B (post-WGD species) and C (pre-WGD, Kluyveromyces species). This plasticity at the clade level is consistent with our initial analysis of Gcn4’s target turnover at the species level.

For many of the regulatory motifs (72%), the targets are substantially changed at a specific point in the phylogeny. For example, the Mig1 motif, involved in glucose repression in S. cerevisiae (Nehlin and Ronne, 1990), is detectable in species across the phylum (up to the LCA, clade L), including a set of ancestral targets in clade D (Kluyveromyces and post-WGD, spanning S. cerevisiae and Kluyveromyces lactis) and in clade G (Candida), but with no statistically significant set of shared ancestral targets between these two clades (Figure 4C). Thus, although the motif likely existed in their shared ancestor (clade H), its targets have diverged significantly between the two descendant clades, precluding reconstruction of the ancestral state. This suggests substantial plasticity in the targets associated with many regulatory DNA motifs.

To quantify the extent of plasticity of motif targets, we developed a model of motif-targets turnover, which handles the gains and losses of a target gene as a stochastic continuous-time Markov process (see Materials and methods). This model is akin to standard models of sequence character evolution (Felsenstein, 1981). We found that motif targets are globally gained and lost at fast rates (Supplementary Figure 2), with a median loss rate per target of 5.2 losses/tU (time unit) and a median gain rate per target of 0.24 gains/tU (Supplementary Table 2; Supplementary Figure 2, see Materials and methods). This discrepancy in the rates is due to differences in the pool of targets versus non-targets in the genome. The typical gain rate is ‘lower’ than the loss rate since it is calculated as a fraction of a larger number of non-target genes (~4000), whereas the loss rate is calculated out of ~100 ancestral target genes.

An instructive measure for the target turnover rates is the number of targets we expect to be retained at different branch lengths, computed by averaging simulations over the expected gain and loss rates of all regulatory motifs (Figure 5A, see Materials and methods). Turnover rates vary substantially among individual motifs. For example, the Hsf1 (heat shock factor) motif exhibits low rates of target gain and loss (Figure 5B), while variants of the CACGTG motif (bound by Pho4, Tye7, and Met28) have very high turnover rates (Figure 5C). On average we found that only 7% of a given motif’s targets in the sensu-s stricto clade (clade A, Figure 3A) are expected to be conserved in the LCA of the phylogeny (clade L, Figure 3A), and only 16% of the targets were conserved since the LCA with the Candida clade (clade H, Figure 3A).
To assess the functional implications of target turnover, we next associated each motif in each clade with a regulatory function, based on the functional categories to which its targets in the clade belong. The substantial redundancy between functional annotations can lead to many overlapping ‘functions’, which are hard to compare across clades (Supplementary Note 3). We therefore developed a method to create functional modules that contain genes that share functional annotations and are all ancestral targets of the same regulatory motif (see Materials and methods, Supplementary Figure 3a). For example, consistent with our initial analysis, Gcn4 targets in each clade are associated only with the amino-acid metabolism module (Figure 6A). This module includes several overlapping gene sets, such as amino-acid biosynthetic process (Ashburner et al, 2000), amino-acid metabolism (Segal et al, 2003), amino-acid nitrogen metabolism (Segal et al, 2003), or pyridoxal phosphate binding (Ashburner et al, 2000). Notably, each motif can be associated with one or more such modules in each clade, and possibly with different modules in different clades (Supplementary Tables 2 and 3; Supplementary website).

Compared with direct enrichment of individual gene sets, functional modules are a more concise and non-redundant representation that can be easily compared across the clades.
phylogeny. In addition, they are more robust to the choices of detection threshold, to the threshold over enrichment of functional categories with motif targets, and to the threshold for merging functional categories (Supplementary Note 3; Supplementary Table 7). Furthermore, supporting our procedure and CladeoScope’s predictions, our functional assignments are consistent with known functions of the associated transcription factors in *S. cerevisiae*, *C. albicans*, and *S. pombe*, for most motifs with a known function (75%) (with another 12% of the motifs with a partial match; Supplementary Note 3; Supplementary Table 8).

**Extensive functional conservation of regulatory DNA motifs**

We observed functional conservation for a large fraction of the regulatory DNA motifs. In all, 48% of motifs are associated only with the same functions in all clades in which the motif is detectable, even across large phylogenetic distances. Examples include the Gcn4 motif (Figure 6A), the Hsf1 motif with a heat shock module (Supplementary Figure 3b), and the Mbp1 motif with cell-cycle and DNA replication modules (Supplementary Figure 3b). Furthermore, although in other cases the motif might gain or lose an association to functional modules during evolution, 82% of all the motifs have at least one conserved function across all clades (Figure 6B).

**Innovations through expansion and switch of functions**

In some cases, turnover of target genes does contribute to evolutionary innovation, by either expanding or switching the scope of functions ascribed to a regulatory DNA motif (Figure 6A). For 34% of the motifs, we observed clade-specific expansion: a motif gains a new function in a specific clade in addition to maintaining its ancestral function. In such cases, the motif is identified in genes from the same functional module(s) in all clades where it was detected, and is also associated with an additional module unique to a specific clade.

![Figure 6](image-url)
We find various innovations in different clades (Figure 6A; Supplementary Figure 4). For example, the Rpn4 motif is associated with the proteasomal module in all clades (Mannhaupt and Feldmann, 2007), while in clade C (the Kluyveromyces species) it is also identified in genes of a cytoskeletal module (Figure 6A). There are several cases of highly conserved motifs exhibiting innovations in the remote Schizosaccharomyces clade (K). For example, the cell-cycle motif Fkh1 regulates genes involved in meiosis specifically in this clade, and the Hap4 motif associated with oxidative phosphorylation in all clades also regulates extracellular matrix and GTP binding genes in the Schizosaccharomyces species (Supplementary Figure 4a). This latter example involves a regulatory switch, as the regulation of GTP binding genes in all other clades is regulated by the Sfp1 motif (Supplementary Figure 4b).

For 18% of the regulatory motifs we observed a functional switch between clades: the same motif has target genes from distinct functional modules in different clades, thus losing one function while gaining another. For example, the Mig1 motif is associated in the Candida (G) clade with modules such as peroxisome and fatty acid metabolism, whereas in the Kluyveromyces (C), the ‘post-WGD’ (B), and the Schizosaccharomyces (K) clades it is associated with other carbon metabolism modules (Figure 6A). An additional example is the motif bound by the factor Ynr063w (Zhu et al, 2009). This motif is associated with general metabolic processes in all clades where it is detected, but switches its specific function; it is associated with the TCA cycle in the Candida clades (E–G), glycolysis in Schizosaccharomyces clade (K), but with the peroxisome and aerobic metabolism in the ‘post-WGD’ clade (A–B) (Supplementary Figure 4a).

Conservation of regulatory function despite high turnover rate of targets

The observations of substantial target turnover and extensive functional conservation are seemingly contradictory. One possible way to reconcile this contradiction would be if the rapid turnover of motif targets is mainly restricted to motifs that exhibit functional changes, but not to those with conserved functions. However, we find rapid target turnover for most regulatory DNA motifs, including those associated with conserved functional modules (Supplementary Tables 2 and 4), such as Gcn4.

Moreover, we observed extensive turnover of motif targets within the functional modules themselves. Specifically, in 80% of modules associated with the same motif in more than one clade, we observed substantial turnover of the motif targets between those clades (Figure 7A). On average, 62% of a module’s genes are associated with the regulatory motif in only a minority of the relevant clades. For example, the Fkh1 motif is consistently associated with a cell-cycle regulation module across the entire phylum (12 clades), but its individual targets substantially turnover, with ~90% of genes detected as Fkh1 targets in only one or two clades (Figure 7B).

A functional selection turnover model

The observed conservation of regulatory function despite high target turnover suggests that the global functional roles associated with a regulatory motif are under stronger selection than the individual regulatory interactions. To formalize this notion we propose the Functional Selection Turnover Model, where selective pressure acts differentially to conserve motif-target relations within the same biological process (compared with outside of the process), but not particular target genes within that process (Figure 7C).

To test this hypothesis, we used a likelihood ratio test (LRT) to compare two alternative evolutionary models (see Materials and methods): (1) a ‘neutral’ turnover model, where targets are gained and lost at the same rates regardless of the functional module to which they belong and (2) a ‘module-specific’ turnover model (described above), where turnover rates—both gain and loss—are different for targets in the functional module compared with those outside. We applied this test to all functional modules in all associated clades (a total of 745 tests).

In nearly all cases (96%, 715 tests), target turnover is significantly constrained by the genes’ function (P-value <0.05 after Bonferroni correction, Figure 7D). Most notably, the probability to gain an additional target gene within the same functional module is typically at least two orders of magnitude higher than the probability to gain a new target from genes outside of the module (Supplementary Table 4). Thus, gain and loss of target genes are highly constrained by their function, resulting in conservation of the motif’s functional role despite turnover at individual sites. These results are not sensitive to the choice of parameters used in the process of target prediction or in defining functional modules, and hence are not an artifact of specific threshold choices made in our computational analysis (Supplementary Note 3).
**Figure 7** Target turnover and the functional selection turnover model. (A) Extent of target turnover within functional modules. The distribution of the percent of conserved targets (x-axis, defined as targets of a motif in the majority of the clades associated with the functional module), for all functional modules with targets in at least two clades. The bin pertaining to the Fkh1 cell-cycle module is marked with a red arrow. (B) Fkh1 target turnover within the cell-cycle module. Target genes (rows) for the Fkh1 motif within the cell-cycle module across the clades (columns). Blue: target in a clade; black: non-ancestral target in the clade. (C) Functional Selection Turnover Model. Cartoon illustration of the model (white: gene in functional module; gray: gene not in module; blue border: target gene of motif; blue node: motif/transcription factor) with two alternative scenarios for target genes turnover from the ancestral Network 1 (top). Both scenarios (bottom) show extensive turnover of target genes. The functional selection scenario (Network 2, bottom right) has selection on the genes’ function, as reflected by the module to which they belong, but not on individual targets, and leads to enrichment of targets within the module along with turnover of individual targets. The module-neutral scenario (Network 3, bottom left) has random turnover of targets and thus leads to loss of target enrichment within the module genes. (D) Testing the functional selection model for cis-regulatory site turnover. The distribution of the log \( P \)-value of the LRT between the two models for target turnover, a module-neutral turnover model (\( H_0 \), Figure 3C, bottom left) and a functional selection turnover model (\( H_1 \), Figure 3C, bottom right), for 745 functional modules and each of their associated clades. Red line—threshold of \( P \)-value 0.05 after the Bonferroni multiple hypothesis correction for rejecting the \( H_0 \) model. (E) The number of highly conserved Gcn4 target genes is as expected given the functional selection turnover model. The distribution of the number of expected highly conserved Gcn4 targets from 1000 simulations, according to the functional selection turnover model. The observed number of Gcn4 targets conserved up to clade H is 35 (red line), with an empirical \( P \)-value \( > 0.5 \). Thus, we cannot reject the null hypothesis that the number of highly conserved genes is as expected by the functional selection turnover model.
The functional selection turnover model is consistent with transcription factor binding data measured across yeast and mammalian species

To examine the generality of our results, we tested whether they hold at the level of individual species as well as clades, when targets are determined experimentally rather than computationally. We thus examined published in vivo transcription factor binding data (from ChIP-chip or ChIP-seq experiments) (Borneman et al, 2007; Tuch et al, 2008; Schmidt et al, 2010). Recent functional studies of transcription factor binding to DNA reported substantial divergence in the bound targets of conserved transcription factors in Ascomycota yeast species (Borneman et al, 2007; Tuch et al, 2008) and between mammalian species (Schmidt et al, 2010). These include Mcm1 binding measured across three relatively distant species (S. cerevisiae, K. lactis, and C. albicans) (Tuch et al, 2008), Ste12 and Tec1 binding in three closely related Saccharomyces species (S. cerevisiae, Saccharomyces mikatae, and Saccharomyces bayanus) (Borneman et al, 2007), and HNF4alpha measured across three mammalian species (human, mouse, and dog) (Schmidt et al, 2010).

Consistent with our cis-regulatory analysis, the binding profiles of all four factors demonstrate high turnover of targets within conserved functional modules (see Materials and methods, Supplementary Table 6; Figure 8), in addition to some species-specific innovations. Applying the two tests described above, we find that the Functional Selection Turnover Model fits the binding data of these four factors in all species (P<10^{-12}), and that the number of highly conserved targets of these factors is as expected by the model (P>0.2). Notably, in mammals the results are not sensitive to the specific threshold for associating an upstream binding site with a target gene (see Materials and methods). Overall, this analysis demonstrates the generality of our findings at different evolutionary distances, measurement methods (sequence analysis and ChIP assays), phylogenetic resolution (species and clades), and group (yeast and mammals).

Discussion

We applied a novel approach for reconstructing cis-regulation to 23 Ascomycota species to study the evolution of their cis-regulatory networks. Using this approach, we systematically identified cis-regulatory interactions for 88 known regulatory DNA motifs across the 23 species, their conserved target genes in each clade and their functional annotations. We exploited this resource to study the regulatory history of...
specific transcription factors and to reach general principles of regulatory evolution. In addition, this constitutes a rich public resource (http://www.compbio.cs.huji.ac.il/OrthoMotifs/) that will facilitate future studies of regulatory evolution of individual clades or species, including human and plant pathogens.

We find and quantify a pervasive gain and loss of motif targets at high evolutionary rates. The high turnover rates that we estimate from data for all motifs (average ~7% expected conserved targets from the LCA of the phylum, Figure 5A) are reflected by the small number of highly conserved targets, and by the complete switch of targets for 72% of motifs in at least one point in the phylogeny. These high turnover rates generalize previous binding studies of few individual transcription factors across three yeast species (Borneman et al., 2007; Tuch et al., 2008), four flies (Moses et al., 2006; Bradley et al., 2010) and five mammals (Schmidt et al., 2010).

The seemingly contradictory trends of a broad conservation of the functions associated with a motif, and the pervasive gain and loss of the motif in individual targets within the module, are reconciled by our proposed Functional Selection Turnover Model, implying that it is a general principle of regulatory evolution. Such conservation of a transcription factor’s cellular function but high turnover of its individual targets had been previously indirectly implicated in the comparison of cell-cycle genes between two yeast species (S. cerevisiae and S. pombe) (de Lichtenberg et al., 2007) and for liver-specific transcription factors across five vertebrates (Schmidt et al., 2010). Our analysis suggests that it is a broad and general phenomenon and our model shows that it can be explained by a strong selection to conserve the function of the motif, but a weaker selection over the specific target genes within this function. This evolutionary model accounts for patterns of turnover from direct measurements of transcription factors across individual species in yeast and mammals, suggesting that the same principle applies at different evolutionary distances, measurement methods, phylogenetic resolution (clades and species), and remote phyla.

There are several alternative potential explanations for the observed conservation of regulatory function despite high target turnover. First, determination of transcription factor targets based on cis-regulatory elements (rather than on limited experimental ChIP data) is challenging and noisy (Gasch et al., 2004; Tanay et al., 2005; Wohlbach et al., 2009), resulting in many false positives and false negatives, which may lead to low overlap in target genes between species. To exclude this option, we compared evolutionary conserved target genes at the clade level instead of target genes for individual species. Second, the transcription factor may target additional genes in some species, thus expanding the scope of functions it regulates, as has been previously shown in yeasts for various factors, such as Mcm1 (Tuch et al., 2008). Although we detect such expansions in several cases (Figure 5; Supplementary Figure 4), we detect high turnover within the large majority of functional modules (Figure 7), including highly conserved modules.

Finally, the high degree of target turnover within a module may be facilitated by the fact that many target genes are co-regulated within dense overlapping regulons (Alon, 2007), where multiple factors have overlapping roles. In ‘single input modules’ (Alon, 2007) (e.g., the galactose utilization pathway in yeast), all the genes in a module are co-regulated by one factor, and we expect strong target conservation. Conversely, in a Dense Overlapping Regulon (e.g., Ribosomal Protein genes), multiple transcription factors regulate the module’s genes, and are partly redundant, such that loss of one regulator might be compensated for by another. For example, amino-acid metabolism genes are commonly regulated by Gcn4 and Leu3, with loss of the regulation by one transcription factor compensated for through gain of regulation by the other (Tanay et al., 2005; Tsong et al., 2006; Hugues et al., 2008; Tuch et al., 2008; Wohlbach et al., 2009; Weirauch and Hughes, 2010) (Supplementary Figure 5b and c). This would be consistent with the conserved co-expression of many functional modules in yeast (Tanay et al., 2005; Hugues et al., 2008) and mammals (Odum et al., 2007). More broadly, transcriptional regulation is only one of many regulatory layers, and control of one or a few members of a complex or pathway may determine the activity level of the whole complex (de Lichtenberg et al., 2007). Thus, a transcription factor may influence the activity of a cellular process by targeting a few genes, and loss of regulation of one target can be compensated by gain of regulation by another transcription factor.

Overall, the function-centered model of targets turnover provides an important insight into the use of conservation as a filter for functional elements in comparative genomics studies (such as ChIP experiments that rely on evolutionary conservation to filter out noise in transcription factor target genes (Harbison et al., 2004)). Moreover, by taking a function rather than a gene-centered view of cis-regulatory evolution, our findings suggest that selection forces are more permissive than has been previously assumed. At the module and transcription factor levels, although turnover within a module may not affect the overall regulatory role of a factor, it may allow for more subtle fine-tuning of gene regulation, facilitating adaptation while controlling against dramatic changes in phenotype.

Materials and methods

CladeoScope algorithm: phylogenetic reconstruction of cis-regulatory networks

The CladeoScope algorithm reconstructs cis-regulatory networks across species: It learns species-specific DNA motifs (including in species lacking any functional annotations and known motifs), using prior knowledge about known PWMs in a model organism, and computationally adapting them to each species (see species-specific motifs). For each motif it then assigns a set of ancestral target genes in the LCA of each clade of species across the phylogeny (Figure 1), inferred using a maximal parsimonious phylogenetic reconstruction (see Phylogenetic reconstruction of ancestral targets). Initially, a gene is predicted to be targeted by a regulatory DNA motif in a species if it contains a binding site of the motif in its promoter (see Motif scanning for putative targets). These predicted targets are used in the reconstruction to find the ancestral set of target genes. In addition, after the reconstruction of ancestral targets, we use them as an input to the motif refinement per species, and choose the optimal motif (see species-specific motifs). The DNA motifs in each species and the ancestral targets in each clade are filtered based on evolutionary conservation within clades of species by their statistical significance (see Phylogenetic filter for noisy motifs and statistical significance). The resulting resource of species-specific motifs, ancestral target sets and functional modules per clade of species are available for download.
at our Supplementary website: http://www.compbio.cs.huji.ac.il/
OrthoMets.

Pseudocode:
Caldesosope (PWM, promoter sequences, gene trees):
1. Find provisional motif targets in each species by scanning promoters.
2. Learn species-specific motifs:
   a. Apply motif discovery algorithm on the provisional target set initialized by the PWM.
   b. Rescan for putative targets using new motif.
   c. Repeat steps a–b using the provisional targets defined in b.
3. Reconstruct ancestral motif targets for each clade in the phylogeny from the provisional species sets using maximum parsimonious dynamic programming algorithm.
4. Repeat step 2 starting with the ancestral targets in each clade, and choose the motif conforming to the higher enrichment threshold.
5. Filter motifs:
   a. Filter motifs in each species by enrichment of motif targets with ancestral targets.
   b. If any motifs are removed go back to step (3).
   c. Filter motifs in the clade level based on statistical significance of the number of ancestral targets.

We now describe the procedure involved in each step of this pseudocode.

Motif scanning for putative targets (steps 1 and 2b)
To identify the putative targets of a motif in the genome, we score each gene’s promoter by summing over all possible positions of the promoter on both strands (as in Tanay, 2006), taking into account the nucleotide background distribution in the promoters of the relevant genome:

\[
\text{score} = \log \left( \sum_{n=0}^{N} \left( \prod_{i=1}^{M} \frac{P_{\text{PWM}}(n_{i})}{P_{\text{BG}}}(n_{i}) \prod_{i=1}^{M} \frac{P_{\text{PWM}}(n_{i+1})}{P_{\text{BG}}}(n_{i+1}) \right) \right)
\]

Where \( N \) is the length of the promoter; \( M \) is the length of the motif; \( n_{i} \) is the nucleotide at the \( i \)th position of the promoter; \( P_{\text{BG}} \) is the background distribution of nucleotides in all promoters of the genome; \( P_{\text{PWM}} \) is the probability vector for nucleotides in position \( i \) of the motif and similarly \( P_{\text{PWM}} \) for the reverse motif (equivalent to searching the reverse strand of the DNA). We considered the 600 base region upstream of each gene’s ATG as its promoter, truncating this region whenever it overlapped a neighboring gene.

We define the target set of the motif as those genes whose promoters have a score above a threshold \( T = 0.8 \) (mean of 20 highest scoring promoters in the genome). The threshold and scanning method where determined by optimizing the precision rate and sensitivity of predictions of in vivo transcription factor target genes from ChiP-chip (Harbison et al., 2004) assays in S. cerevisiae of two different transcription factors: Hsf1 and Rpn4 (Supplementary Note 4). Arguably, this might bias our choices to levels of binding that are significantly detectable by these assays. However, perturbation analysis of this threshold show that our result are mostly robust to this choice (Supplementary Notes 1–3). Since this score is relative to each genome and to each motif, we exclude motifs that do not have any occurrences in the genome by filtering out motifs whose highest score in the genome is < 50% of the maximum possible score of this motif in a single location. Additionally, we removed motifs from the collection if the number of inferred targets was > 1500 (the upper bound was chosen to exceed the maximal number of promoters bound by any transcription factor in S. cerevisiae, as measured by ChiP-chip (Harbison et al., 2004)).

Species-specific motifs (step 2)
Our underlying assumption in this step is that the binding specificities of transcription factors, represented as DNA motifs, are largely conserved, even when their specific target genes and functional roles may have substantially diverged (Wapinski et al., 2007; Tuch et al., 2008; Schmidt et al., 2010). We therefore initiate our reconstruction with DNA motifs for known transcription factors that have been experimentally determined in model organisms. CladoScope uses the MEME motif discovery algorithm (Bailey and Elkan, 1994) on the promoters of the putative targets of each initial motif in each species, with the initial motif’s consensus sequence as the initialization point to the algorithm. MEME is parameterized to identify motifs on either strand, of length within two bases from the input consensus, and it is given the species-specific nucleotide distributions as background models for learning. Of the top two motifs reported by MEME, the highest scoring motif in this species is then used to rescan the species’ genome and to identify a revised set of targets. CladoScope repeats this process of motif discovery and rescanning to refine the motif and its target set once; typically the motif is not altered after the first iteration. To allow more variation in the motifs, we repeat the process, initializing the refinement with the conserved ancestral targets. This allows us to find motifs in species where the first iteration did not succeed. CladoScope chooses the motif with the highest enrichment score between these iterations.

Phylogenetic reconstruction of ancestral targets (step 3)
To infer the ancestral motif targets we trace regulatory events across orthologous loci. CladoScope handles genes derived from a common ancestor gene in the root of the phylogeny as related (‘orthogroup’ in the terminology of Wapinski et al., 2007), and defines an orthogroup as a target of a motif if it is predicted as a target in at least one of the ancestors in the phylogeny. The reconstruction is done using maximum parsimony (Fitch, 1971) to minimize the number of target gain and loss events along the branches of the tree. This is done separately for any potential ancestral target gene by a dynamic programming algorithm. The inputs to the algorithm are (1) the phylogenetic gene trees for each set of orthologous genes (Wapinski et al., 2007) and (2) a binary classification denoting whether each gene in the tree is a predicted target of the motif in each species. This reconstruction accounts for gene duplications and losses, distinguishing a lost gene from a present gene that is not a target, and operates independently on each paralogous lineage following gene duplication events, by utilizing gene trees when reconstructing ancestral targets. Given that most of these species have diverged sufficiently to lose sequence similarity at the promoters, paralogs will not necessarily be co-targets of a motif due to spurious conservation of their promoters. Thus, paralogs can be in different motif-target sets in the CladoScope output.

Phylogenetic filter for noisy motifs and statistical significance (step 4)
We use phylogenetic conservation within a clade of species as a filter for noisy predictions of target genes (described above) as well as the DNA motifs themselves. CladoScope filters the motifs for each species independently by enrichment of their putative target genes in that species with ancestral targets in any relevant clade (hypergeometric \( P \)-value < 0.001). Since filtering the motifs (step 4) and the reconstruction of ancestral targets (step 3) are dependent, we solve this problem by iterating between the two steps. If any insignificant motifs are found in the clade, the most insignificant one is removed, and CladoScope returns to step 3 of reconstructing the ancestral targets. This filtration per species allows for a motif to be detected in a clade of species, although it is not functional in a single species but is functional in all other species in the clade. We then filter the motifs at each clade, requiring that the number of inferred targets for a motif in the clade’s ancestor be statistically significant (\( P \)-value < 0.005) against the null hypothesis that the targets predicted in the individual species are independent. We compute an empirical \( P \)-value by simulating target sets of the relevant size for each species in the clade, and reconstructing ancestral targets from these random sets. This process is repeated 1000 times to estimate the probability of getting a set of ancestral targets of a certain size or larger by chance. A motif is detectable in a clade if it has a statistically significant (\( P \)-value < 0.005)
set of ancestral targets in the clade. Finally, we exclude motifs that are found to be significant only in clade A (sensu-stricto), since in this clade promoter sequences have not evolved enough for random occurrences of a motif have a non-trivial chance to be conserved.

Validating CladeoScope’s performance using synthetic data

We generated simulated target sets in extant species for evaluating CladeoScope’s robustness (Supplementary Figure 1) by evolving targets from an ancestral set of targets using turnover (gain and loss) rates of target genes, with several variations. First, we used two types of noise factors: (1) the proportion of erroneous target genes relative to the species true motif targets (false positives, ranging between 0% and 200% of erroneous targets within each species set) and (2) the proportion of missing (true) targets not included in the species motif target set (false negatives, ranging between 0% and 60% of removed targets from each original species set). Second, we varied the size of the ancestral target set. Using ancestral motif targets in clade A (Figure 3A): 22 targets of Hsf1, 198 targets of Mbp1, 297 targets of Fkh1. Third, we varied the degree of targets turnover (the fast turnover is the average frequency measured in clade E (Figure 3A) over all motives: Fast $F_{\text{gain}} = 0.002, F_{\text{loss}} = 0.3$. Medium $F_{\text{gain}} = 0.002, F_{\text{loss}} = 0.03$. Slow $F_{\text{gain}} = 0.00002, F_{\text{loss}} = 0.003$). Fourth, we estimated the gain and loss frequencies for each of the three motifs in each relevant species directly from the data (as in the LRT described below) (Supplementary Note 1). Fifth, we used two topologies of the species tree: the topology in the sensu-stricto clade (clade A, Figure 3A), and the asymmetrical topology in the Candida clade (clade E, Figure 3A). Overall, we considered 960 combinations of these parameters. For each set of parameters, we executed CladeoScope and calculated sensitivity and specificity measures averaged over 100 independent simulations.

Assessing performance on random motifs

We created random motifs by concatenating randomly sampled positions from all known motifs from the literature (using all motifs from S. cerevisiae, as described below). We confirmed that the random motifs we constructed were not similar to any known motifs, compared the random motifs to all known motifs using BLiC (Habib et al., 2008). For each random motif, we scanned for targets in each species (as described above), and ran CladeoScope to reconstruct the ancestral sets. We then computed an empirical P-value for each motif in all clades using random targets (as described above).

Assessing CladeoScope’s robustness to parameters and comparison to the literature

We tested CladeoScope’s robustness to variations in different parameters including (as described in Supplementary Note 1): (1) The P-value threshold for detection of a motif in a species—We ran the algorithm on nine different motifs across all clades, using seven different thresholds, ranging between 5e – 2 and 1e – 5, and compared the number of ancestral targets reconstructed per clade. (2) The P-value threshold for conservation of a motif in a clade—We tested different P-value thresholds ranging between 0.05 and 0.001, and compared the number of statistically significant ancestral motifs predicted per clade. (3) The motif-targets detection threshold—We tested three different thresholds (80, 75, or 70% out of the best score per motif and species). In each case, we compared the ancestral and species targets determined by CladeoScope to those from in vitro Chip-chip data in S. cerevisiae and four other species (Supplementary Note 1).

Targets turnover rates and expected number of changes in target genes

For each motif we computed the turnover rate of its target genes based on the following model. The model treats each pair (motif, gene) as a binary character denoting whether the gene is a target of the motif or not. We model changes in this character (gain or loss events) as a stochastic continuous-time Markov process parameterized by motif-specific rates, one for gain, and another for loss. This model is akin to standard models of character evolution (Felsenstein, 1981). The rates are expressed in terms of expected number of events per time unit (t), where a time unit corresponds to the time in which one amino-acid substitution per site is expected on average. The model assumes a constant turnover rate of targets along the phylogeny, which is reflected by two parameters for each motif: its gain rate (a) and its loss rate (b), given by the following rate matrix R:

$$ R = \begin{pmatrix} a & b \\ a & b \end{pmatrix} $$

Given this rate matrix we can compute the probabilities for target gain and loss for a given evolutionary distance t using the following equations:

$$ P(\text{Gain} | t) = \frac{1}{a+b} \left(1 - e^{-\frac{a+b}{a+b}t} \right) $$

$$ P(\text{Loss} | t) = \frac{1}{a+b} \left(1 - e^{-\frac{a+b}{a+b}t} \right) $$

We use a maximum-likelihood estimator to infer the parameters in R for each motif. The likelihood is computed based on sufficient statistics for each clade relative to its immediate ancestral clade, including the branch length (t), the observed number of gained ($N_{\text{gain}}$) and lost ($N_{\text{loss}}$) motif targets of ancestral (ancestral) target genes, and the probabilities described above, as:

$$ \text{In Likelihood (Motif)} = \sum_{b \text{branches}} \left[ N_{\text{gain}} \ln P(\text{Gain} | t^b) + N_{\text{loss}} \ln P(\text{Loss} | t^b) + N_{\text{initTarget}} \ln (1 - P(\text{Gain} | t^b)) + N_{\text{initTarget}} \ln (1 - P(\text{Loss} | t^b)) \right] $$

The maximum-likelihood estimator is found by a gradient descent algorithm using Matlab’s fminunc function. We assume the tree topology and branch length are known (see Species phylogeny).

Annotating motifs with functional modules

To associate motifs with regulatory functions, we cluster functional gene sets together by the fraction of associated motif targets shared between them, creating sets of functional modules containing genes that share functional annotations and are all ancestral targets of the same regulatory motif in at least one clade (Supplementary Figure 3A).

The method is applied to each motif separately. As input we provide the ancestral target genes of the motif in each clade, and gene sets of functional annotations from various sources. In step 1 (Initialization), we identify all functional annotations enriched in each set of ancestral targets in each clade using Fisher’s exact test ($P < 0.01$ after correction; however, the results presented here are robust to various thresholds), and define a functional module as genes from each enriched category that are ancestral targets in any clade. In step 2 (Merge functional modules), we merge modules according to the fraction of associated motif targets shared between them. In this greedy procedure, we start from the most enriched module, choose another one that is most highly overlapping with it (at least 60% gene membership overlap; however, the results presented here are robust to various thresholds) and unite them into a new functional module, eliminating the two daughter modules from the collection. In step 3 (Recalculate enrichments), we recalculate the enrichment of the ancestral targets in each clade with this new functional module. We repeat steps 2 and 3 until no further functional modules are merged. Following the automatic assignment of modules, we manually annotated each functional module with a biologically meaningful label based on its underlying annotations.

The functional modules assigned to each motif, and the target genes assigned to each module can be found in Supplementary Table 3. Modules and genes created by different sets of parameters can be found in Supplementary Table 9.
Note that the assignment to functional modules is based on phylogenetic projection from S. cerevisiae, C. albicans, and S. pombe annotations. As a consequence, the function assignment often cannot distinguish between paralogs. Moreover, in a previous study of functional evolution (Wapinski et al., 2007), we showed that when we can evaluate such divergence, most paralogs maintain the same functional category. Thus, we expect some functional modules to be enriched for paralogs (e.g., the Ribosome, due to the massive duplications of genes encoding ribosomal proteins). This, however, reflects a real phenomenon.

The classification of motifs to three functional categories

The classification of motifs to three functional categories is based on the explicit definition of each category: For conserved motifs, all associated modules are enriched across (the relevant) clades. Expanded motifs have at least one module conserved across clad"es, and an additional module specific for a subset of these clades. Switched motifs have modules enriched in different subsets of clades. We applied these definitions with one exception, in which we excluded small modules (with <5 genes uniquely classified to the module). To increase confidence in these results, we subsequently examined each classification individually, and compared the module enrichment pattern to the conservation pattern of target genes across species. We marked in Supplementary Table 2 cases where this examination raised doubts in the classification. To further ensure that these classifications are not an artifact of specific thresholds in module construction, we selected three representative parameter sets and applied this process separately to each one of them (Supplementary Table 2). For 18 motifs (including all the ones mentioned in the text and figures), we performed a more extensive comparison of module-construction parameters (Supplementary Table 7). Notably, the examples throughout the text and figures include only robust examples across all parameters. For the analysis presented in the manuscript and in Figure 6B, we constructed a consensus classification for each motif. We examined the three classifications described above and defined the consensus/majority classification for each motif (Supplementary Table 2). In virtually all cases there was a clear-cut consensus classification.

Functional annotation resources

We used functional annotations from several sources. GO annotations (Ashburner et al., 2000) were assembled from the genome databases of S. cerevisiae (SGD), C. albicans (CGD), and S. pombe (GeneDB). Other S. cerevisiae-based annotations include expression clusters (Chen et al., 2003), MIPS (Mewes et al., 2011), KEGG (Kanehisa and Goto, 2000; Kanehisa et al., 2006), and mutant phenotypes (Hughes et al., 2000). Other S. pombe-based annotations include expression clusters (Chen et al., 2003). We projected each set of annotations from genes to their orthologs (Wapinski et al., 2007) to test gene set enrichments across all clade core-sets, as previously described (Wapinski et al., 2007).

Assessing robustness of functional modules and their comparison to the literature

To test the robustness of the functional modules, we applied the algorithm with different parameters and inputs, including (see Supplementary Note 3): (1) Enrichment thresholds for functional modules with motif targets (HyperGeometric P-value threshold ranging between: 1e − 3 and 1e − 6), (2) Threshold for merging gene sets (overlap threshold ranging between 40% and 75%), (3) Threshold for initial predictions of target genes (80% or 75% out of the best score per motif and species). For each set of parameters we tested several characteristics: (1) the number of modules; (2) the fit of our functional selection turnover model; (3) the classification of motifs to functional classes (functional conservation, Clade-specific innovation or Functional switch), where we examined in detail 18 motifs including those discussed specifically in the manuscript; and (4) robustness of the functional annotations of motifs by the functional modules, where we examined in detail 18 motifs including those discussed in the manuscript.

In addition, we compared the resulting functional modules to known motif and transcription factor annotations from the literature in SGD, CGD, and GeneDB.

The functional selection turnover model

We used an LRT to determine if the observed functional conservation with widespread turnover occurs by chance or according to our functional selection model. We defined two alternative hypotheses: H0: Module-Neutral turnover: Targets turnover at the same (‘neutral’) rate regardless of the functional module to which they belong (implying that the functional conservation may be a byproduct of the insufficient evolutionary distance between species).

H1: Functional selection: There is selective pressure on the targets to be gained or lost within modules of genes sharing the same function. Turnover rates—both gain and loss—are different for targets in the functional module compared to those outside.

We applied the LRT to each functional module testing separately each associated clade (total of 745 tests), by computing the likelihood of the observations under each hypothesis and calculating a P-value ($\chi^2$ distribution with one degree of freedom). The likelihood computations were based on maximum-likelihood estimates of gain and loss probabilities in the clade relative to its immediate ancestral clade. The required sufficient statistics are the observed number of gained ($N_{gain}$), lost ($N_{loss}$), and conserved ($N_{cons}$) target genes. In the functional selection model (H1 hypothesis), we computed the gain and loss probabilities separately for genes within the module and genes outside of the module.

Description of the equations:

\[
LRT(Motif) = \log(P(Gain | H_0) + \log(P(Loss | H_0)) + \log(1 - P(Loss | H_0)) + \log(1 - P(Gain | H_0))
\]

\[
LRT(b | H_0) = \frac{N_{gain}^{IN} \log(P(Gain^{IN} | H_1)}{N_{total}^{IN} - T_{ Cp}} - \frac{N_{loss}^{IN} \log(P(Loss^{IN} | H_1)}{T_{ Cp}}
\]

\[
LRT(b | H_0) = \frac{N_{gain}^{OUT} \log(P(Gain^{OUT} | H_1)}{N_{total}^{OUT} - T_{ Cp}} - \frac{N_{loss}^{OUT} \log(P(Loss^{OUT} | H_1)}{T_{ Cp}}
\]

The maximum-likelihood estimation for the probability of gain and loss of each motif’s target genes in the current clade $C_1$, compared with the immediate ancestral clade $C_p$:

\[
P(Gain | H_0) = \frac{N_{gain}}{N_{total} - T_{ Cp}}
\]

\[
P(Loss | H_0) = \frac{N_{loss}}{T_{ Cp}}
\]

\[
P(Gain^{IN} | H_1) = \frac{N_{gain}^{IN}}{N_{total}^{IN} - T_{ Cp}}
\]

\[
P(Loss^{IN} | H_1) = \frac{N_{loss}^{IN}}{T_{ Cp}}
\]

Where $N_{total}$ = total number of genes in the genome, $T_{ Cp}$ = total number of targets in clade $C_p$, IN = genes belonging to the functional module, OUT = genes not belonging to the functional module, $N_{gain}^{IN}$ = total number of genes in the functional module, $N_{loss}$ = total number of genes in the functional module that are targets in clade $C_p$. © 2012 EMBO and Macmillan Publishers Limited Molecular Systems Biology 2012 15
Simulating the number of highly conserved targets

To test whether the number of highly conserved targets is explained by the functional selection model, we computed the probability of observing the inferred number of these targets under the functional selection model (the $H_1$ hypothesis defined in the LRT, above). We then simulated targets in the two subclades that share the same direct ancestral clade, and computed the overlap between these simulated target sets. The simulations were initialized with the target set of the ancestral clade, and we simulated the targets in each subclade according to its probability of gain or loss of target genes, within and outside of the functional module (computed as described above for the LRT, using a maximum-likelihood estimator). We repeated this process 1000 times, counting the number of times in which the number of simulated ancestral targets was equal to or greater than the number observed in our data. We ran the test on motifs conserved at least up to clade H (29 motifs), and computed these empirical probabilities of the intersection between the target sets at clade D and clade G. In general, for two target sets $C_1$ and $C_2$ of clades that share an immediate ancestral clade, we computed the empirical probability for the number of genes in the intersection between the two target sets, denoted as $I_{p}$:

$$P(I \geq I_{p}) = \frac{\#(I_{simulated} \geq I_{p})}{\#simulations}$$

Where $\#(I_{simulated} \geq I_{p})$ is the number of simulations where $C_1 \cap C_2 \geq I_{p}$ and $\#simulations$ is the total number of simulations (1000).

Experimental transcription factor binding data

We used Ste12 and Tc11 binding in three closely related Saccharomyces species by ChIP-chip (Borneman et al., 2007); Mcm1 binding measured across three more distant yeast species by ChIP-chip (Tuch et al., 2008); and HNF4α binding measured across three mammalian species (human, mouse, and dog) by ChIP-seq (Schmidt et al., 2010). For the yeast studies, we used target genes defined in the original manuscripts. For the mammals, where regulatory elements can reside far from their target genes, we had to assign each bound regulatory element with the gene(s) it controls. We focused on binding events in the proximity of the gene, and used five alternative definitions of promoters, ranging from 1000 bases up to 600 bases upstream from the first codon, truncated at the poly-A/T sequences longer than nine bases and replaced stages. We filter out stretches of poly-A or poly-T sequences of five bases or longer and poly-A/T sequences longer than nine bases and replaced them with poly-N of the same length.

Motifs were assembled from TRANSFAC (Matys et al., 2006), protein microarray data (Zhu et al., 2009), and previous analysis of ChIP-chip data (MacIsaac et al., 2006). All motifs were transformed to a PWM format (a $n \times 4$ matrix, where each $i,j$ cell contains the count of nucleotide $j$ in position $i$ of the motif), and clustered (using BLiC Habib et al., 2008) to unite highly identical motifs.

Species phylogeny

The CladeoScope algorithm as well as the maximum-likelihood estimators described above, assume the species phylogeny is known. Thus, to reconstruct the phylogenetic relationship between the species, we first identified all the orthologous genes with exactly one copy in each of the species (Wapinski et al., 2007) and aligned their orthologous protein sequences. We concatenated all of these alignments to produce a meta-alignment of over 300,000 positions. We sampled 10,000 residues from this alignment, giving us an artificial protein from which we reconstructed the phylogeny using the PhyML (Guindon et al., 2010) software package with its default parameter settings. We repeated this process 10 times, rendering the same phylogeny at all branches except for the post-WGD clade of species, in which Candida glabrata and S. castellii were sometimes found to be inverted. Recent work (Scannell et al., 2006) has shown that it requires fewer genomic rearrangements to place S. castellii as the outgroup of this clade and that the longer branch length leading to C. glabrata may be due to increased selective pressure as it became a pathogenic species. Thus, we fixed the branches at this location of the tree. In order to re-estimate the branch lengths with this fixed tree topology, we repeated the same process to construct an artificial protein and ran the SEMPHY software package (Friedman et al., 2002) to optimize branch lengths with default parameters. We repeated this process 10 times and found branch length correlations of over 0.99 between replicates. We then averaged the branch lengths among the 10 replicates to obtain branch length estimates for the given species phylogeny.

Supplementary information

Supplementary information is available at the Molecular Systems Biology website (www.nature.com/msb).

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Author contributions: NH, IW, HM, AR, and NF conceived and designed the study; NH and IW developed and applied the CladeoScope framework; NH developed the models and performed the analyses; NH, IW, HM, AR, and NF wrote the paper.

Conflict of Interest

The authors declare that they have no conflict of interest.

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