Computational Comparative Study of Transcription Regulation in Eukaryotes

Thesis submitted for the degree of “Doctor of Philosophy”

by

Naomi Habib

Submitted to the Senate of the Hebrew University
August 2012
This work was carried out under the supervision of Prof. Nir Friedman and Prof. Hanah Margalit
Abstract

Transcription regulation plays a central role in the activity of living cells and in their response to internal or external stimuli. This complex regulatory process is mediated by multiple interacting mechanisms. These mechanisms include both sequence specific binding proteins called transcription factors, and epigenetic mechanisms, including DNA methylations and chromatin modifications, as well as non-coding regulatory RNAs. Changes in gene regulation have been postulated to play a key role in generating the wide phenotypic diversity observed across species. Yet the evolutionary driving forces and the dynamics of this evolutionary process are largely unknown. I studied this process from two different perspectives.

Cis-regulatory evolution

Regulation of gene expression can evolve through mutations in the DNA sequence leading to altered activity of trans acting factors such as transcription factors, or to changes in cis-regulatory sequences at promoter and enhancer regions, which affect the binding of regulatory proteins. While trans acting factors are largely conserved, large-scale changes in cis-elements were observed for specific factors in organisms as diverse as yeasts, flies, and mammals. Rigorously studying regulatory evolution has been hampered by the lack of large-scale and systematic experimental studies, and by the noisy nature of computational predictions.

We developed an unbiased computational scheme to study the evolution of transcription regulation across large phylogenies, and used it to trace the regulatory history of more than 90 transcription factors across 23 yeast species. Our analysis revealed general principles in the evolution of transcription regulation both in yeasts and mammals. We found that: (1) The regulatory network of transcription factors and their target genes is highly plastic (i.e. transcription factors gain and lose target genes at a fast rate). (2) Transcription factors tend to conserve their functions. (3) A functional selection turnover model can reconcile these two trends, suggesting that the global functional roles associated with a transcription factor are under stronger selection than the individual target genes. In our model, selective pressures act differentially to conserve target genes within the same biological process (compared to outside of the process), but not particular target genes within that process. The model is sufficient to explain the observed number of highly conserved targets (across all species), and fits the variation in measured transcription factor binding profiles across species, both in yeasts and mammals. Our findings suggest that selection
forces are more permissive than has been previously assumed. We show that selective pressures on regulatory networks tolerate massive local rewiring, facilitating adaptation of gene-expression while controlling against dramatic changes in phenotype.

**Epigenetic inheritance**

Epigenetic regulatory mechanisms provide an additional potential driving force in the evolution of transcription regulation, which can lead to transgenerational reprogramming of gene-expression. Epigenetic inheritance implies that information about the environment experienced by parents could be transferred to offspring by non-Mendelian inheritance. Whether or not organisms can inherit characters induced by ancestral environments in mammals is unclear and has far-reaching implications. To test whether such transgenerational inheritance occurs, we carried out an expression-profiling screen for genes in mice that responded to paternal diet. We focused on paternal diet to rule out simple plastic responses of offspring to the in-utero environment, as fathers often contribute little more than sperm to offspring.

Relative to the offspring of males fed a control diet, the offspring of males fed a low-protein diet increased the expression of many genes involved in lipid and cholesterol biosynthesis, and had increased levels of cholesterol esters, triglycerides, and free fatty acids, lipids. Extensive epigenetic profiling and computational analysis of offspring livers, as well as whole genome characterization of cytosine methylation patterns and RNA content in sperm, revealed numerous modest (20%) changes in cytosine methylation of offspring liver depending on paternal diet, including reproducible changes in methylation over a likely enhancer for the key lipid regulator Ppara. Our work is one of the first to provide a systematic evidence that: (1) **Paternal diet affects metabolic gene expression in the offspring of mice.** (2) **Epigenetic information carriers in sperm respond to environmental conditions.** These results, in conjunction with recent human epidemiological data, indicate that parental diet can affect cholesterol and lipid metabolism in offspring and define a model system to study environmental reprogramming of the heritable epigenome. Moreover, these results suggest **rethinking basic practices in epidemiological studies of complex diseases such as diabetes, heart disease, or alcoholism.**

**Taken together,** our results shed light on two different selection forces driving evolution of transcription regulation, and emphasize the need for an extended evolutionary theory, integrating both genetic and non-genetic inheritance.
Contents

Chapter 1 – Introduction

1.1 From DNA to RNA................................................................. 1
1.2 Transcription factors and regulatory networks................................ 2
1.3 Epigenetic factors - DNA and chromatin modification...................... 3
1.4 Regulatory non-coding RNAs.................................................. 4
1.5 Evolution of transcription regulation.......................................... 5
1.6 Overview............................................................................. 7

Chapter 2 - A Functional Selection Model Explain Robustness Despite Plasticity in cis-Regulatory Networks................................................. 8-59

2.1 Introduction.......................................................................... 8-15
2.1.1 Rewiring of regulatory networks through changes of cis-regulatory elements.......................................................... 8
2.1.2 Experimental methods to study cis-regulatory evolution............. 9
2.1.3 Computational methods to study cis-regulatory evolution........... 10
2.1.4 Current studies of cis-regulatory evolution............................. 12
2.1.5 Yeast as a model for cis-regulatory evolution........................ 13

2.2 Results................................................................................. 16-43
2.2.1 CladeoScope: a framework for reconstructing cis-regulatory evolution.......................................................... 16
2.2.2 Systematic reconstruction of the regulatory history of 23 Ascomycota species................................................................ 20
2.2.3 Plasticity of regulatory networks in Ascomycota fungi.................. 27
2.2.4 Functional evolution of transcription factors in Ascomycota fungi... 30
2.2.5 Functional Selection Turnover Model – A general Principle of Regulatory Evolution.................................................. 38

2.3 Methods................................................................................. 44-59
2.3.1 CladeoScope algorithm: Phylogenetic reconstruction of Cis-regulatory networks.................................................. 44
2.3.2 Resources for phylogenetic reconstruction in Ascomycota fungi... 50
2.3.3 Evaluations of CladeoScope in Ascomycota fungi....................... 51
2.3.4 Targets turnover rates and expected number of changes in target genes... 53
2.3.5 Annotating motifs with functional modules and their evaluation..... 54
2.3.6 The functional selection turnover model.................................. 56
**Chapter 3 - Paternally Induced Transgenerational Environmental Reprogramming of Metabolic Gene Expression in Mammals**

**3.1 Introduction**
- Epigenetic Inherence and the Environment ........................................ 60
- Evidence for Trans-Generational Effects of the Environment ............... 61
- Evidence for Heritable Epigenetic Effects of Environmental Perturbations 61

**3.2 Results**
- Experimental paradigm ................................................................. 63
- Upregulation of proliferation and lipid biosynthesis genes in low protein offspring ................................................................. 66
- Transgenerational effects on lipid metabolism ................................ 68
- MicroRNAs in offspring ................................................................. 70
- Cytosine methylation in offspring .................................................. 71
- Cytosine methylation, RNA, and chromatin in sperm ....................... 73

**3.3 Methods**
- Experimental procedure ............................................................... 76
- Micro-Arrays data processing and differentially expressed genes in the liver ................................................................. 77
- MicroRNA identification from deep sequencing data and analysis ....... 78
- Comparison to public murine liver microarray data ......................... 79
- Percent variance explained by *Ppara* RNA levels ......................... 80
- Analysis of sperm RNA data .......................................................... 80

**Chapter 4 - Discussion**
- Robustness in the face of plasticity – *cis*-regulatory evolution .......... 81
- Epigenetic inheritance - Transgenerational environmental reprogramming of gene expression .................................................. 86
- An extended evolutionary theory ...................................................... 90

**References** .................................................................................. 92

**Appendices** .................................................................................. i-xvii
Chapter 1 - Introduction

1.1 From DNA to RNA

The genome of a living organism contains the hereditary instructions for its development and function. This information is encoded in DNA molecules that are found inside each cell, and are built of nucleotides (A,C,G,T). Segments of the DNA sequence (genes) are transcribed to RNA molecules, and can then be translated to proteins (gene expression). Proteins perform a variety of functions in the cell. The collection of RNAs and proteins expressed in a cell determine its morphology and how it functions. The DNA sequence does not change in different stages throughout the life of a cell and is identical in different cell types of multicellular organisms. However, the function and structure of cells are not constant, but change in response to internal or external stimuli (e.g. while cells differentiate; in single cellular organisms in response to changes in the environment). Generating diverse outputs from a single set of instructions (DNA sequence), requires a tight and highly specific regulation on the content of RNA molecules and active proteins in the cell.

Transcription regulation is the first layer of this regulation, which plays a critical role in the activity of living cells and in their response to internal or external stimuli. Complex regulation is required to determine which genes would be expressed at any given time and to what extent. This regulation responds to changes in the environment, as well as to the internal state of the cell, and is mediated by multiple interacting mechanisms (Figure 1). These mechanisms include regulatory proteins that bind to specific DNA sequences, as well as diverse epigenetic regulatory mechanisms, including the chromatin state, DNA modification and regulation by non-coding RNAs. I review each of these mechanisms in more detail.
1.2 Transcription factors and regulatory networks

The information regarding which genes will be expressed at any given time is encoded in the DNA sequence. This information is mostly separate from the sequence encoding the protein - and appears in *regulatory regions* of the DNA (Figure 1), primarily located upstream to genes in *promoter* regions. Such regulatory sequences are recognized by proteins, called *transcription factors*, that bind to specific DNA sequences (Figure 1). This sequence specificity is important for the expression of specific genes under every condition. Once a factor binds to the DNA it modulates the RNA level of specific (typically nearby) genes, by activating or repressing their transcription.

There are multiple transcription factors in every organism (estimated at several hundreds in yeasts (Wapinski et al., 2007) and at least two thousand in mammals (Babu et al., 2004; Messina et al., 2004; Vaquerizas et al., 2009)). Each factor is activated by different stimuli, and mediates the expression of genes relevant to specific conditions. The genes that are regulated by a specific factor are considered its *target genes*. A transcription regulation network is a map of all transcription factors and their target genes, and provides a general view of transcription regulation, enabling us to infer which genes

---

**Figure 1. Transcription regulation machinery**

Transcribing genes at the right time and level involves multiple regulatory mechanisms, including: *trans* acting regulatory proteins, such as transcription factors, that bind to specific *cis*-regulatory sequence elements on the DNA; Nucleosomes position and histone modifications; DNA methylations and regulatory non-coding RNAs (ncRNA).
will be expressed under diverse conditions. Previous studies, focusing on specific systems across different organisms (Amit et al., 2009; Benfey and Chua, 1990; Capaldi et al., 2008; Davidson, 2001; Novershtern et al., 2011; Parker et al., 2011) showed that a gene is often controlled by several transcription factors and each factor regulates multiple genes, implying a complex transcription regulation network.

An example of a combinatorial regulatory system is the response to osmotic stress in yeast. In this system, five different transcription factors were found to coordinately activate and repress hundreds of genes (Capaldi et al., 2008). Induced genes can be divided to eight sets, each regulated by a different combination of these factors (Capaldi et al., 2008). Interestingly, among these factors Capaldi et al found that the Sko1 factor both represses and activates the same genes before and after the stress, respectively (Capaldi et al., 2008). This complex network allows a sensitive activation of specific combination of genes in responses to different stresses, such as osmotic stress by salt versus sugar.

### 1.3 Epigenetic factors - DNA and chromatin modification

The traditional view of transcription regulation was based on the acknowledgement that all the regulatory information is encoded in the DNA sequence, and this information is "read" by sequence specific transcription factors. More recently, other layers of information beyond the DNA sequence have been recognized as central players in transcription regulation. These layers include chemical modifications of the DNA or chromatin (Figure 1). These include:

1. **DNA Methylation** - Addition of methyl groups to the DNA, usually occurs at a cytosine nucleotide immediately followed by a guanine (CpG dinucleotide). This is a stable modification that can be inherited through cell divisions (Wigler et al., 1981). In animals, methylation near gene promoters varies considerably depending on cell type. The degree of methylation in the promoter correlates with low transcription of the downstream gene (Colot and Rossignol, 1999; Jones and Taylor, 1980; Suzuki and Bird,
In addition, the DNA methylation pattern is primarily determined early during development of the organism, and is required for maintaining the specific transcriptional program in different cell types.

(2) **Chromatin modification** - In eukaryotic cells, the DNA is condensed within the nucleus, packed around proteins to form chromatin (Kornberg, 1999). The basic packaging units are nucleosomes, complexes of histone proteins wrapped with DNA. This packing serves not only to condense DNA within the nucleus, but also plays an important role in transcription regulation. First, changes in position of the nucleosomes on the DNA can inhibit or enable access to specific DNA regions and as a result modulate gene expression (Almer et al., 1986; Bergman and Kramer, 1983; Tirosh and Barkai, 2008). Second, the histone proteins are subjected to modifications, such as addition of methyl and acetyl groups, which are associated with repression and activation of genes (Koch et al., 2007; Kooistra and Helin, 2012; Liu et al., 2005).

Importantly, several of these modifications were shown to cause heritable changes in gene expression, since they may remain through cell divisions for the remainder of the cell's life and may also last for multiple generations (Bird, 2002; Colot and Rossignol, 1999; Groth et al., 2007; van der Heijden et al., 2006).

### 1.4 Regulatory non-coding RNAs

Some RNA molecules are not translated to proteins, but rather have important functions in the cell. Among these *non-coding RNAs (ncRNA)*s are several classes of regulatory RNAs that affect the levels of other RNA molecules, by either regulating gene expression, or post-transcriptionally regulating the degradation or translation of RNAs (Figure 1). For example, there are short RNA sequences of ~21 nucleotides called *microRNAs*, that bind to complementary sequences on target mRNA transcripts, resulting in inhibition of translation and/or destabilization of the target mRNA. Another example is of the Piwi-interacting RNAs (*piRNAs*), which silence specific genes by mediating DNA methylation (Aravin et al., 2008; Rajasethupathy et al., 2012) and also cause the
degradation of target RNA molecules (O'Donnell and Boeke, 2007). Overall, regulatory RNAs preform a wide range of important function in the cell (Costa, 2007; Eddy, 2001; Mattick and Makunin, 2006; Storz, 2002). Moreover, recent evidence suggest that they can trans-generationally reprogram gene-expression, since not only that they are transferred through cell divisions, but they can also be inherited to offspring through the germ cells (Ashe et al., 2012; Burton et al., 2011; Cuzin and Rassoulzadegan, 2010; Rechavi et al., 2011).

1.5 Evolution of Transcription Regulation

Changes in gene regulation have been postulated to play a key role in generating the wide phenotypic diversity observed across species (King and Wilson, 1975; Prud'homme et al., 2007; Wittkopp et al., 2004). For example, comparison of humans to our closest living primate relatives shows that despite the vast phenotypic differences between humans and other primates, we all share a remarkable amount of DNA sequence (King and Wilson, 1975). These apparent phenotypic differences are mostly explained by significant changes in gene expression patterns among primate species (Caceres et al., 2003; Enard et al., 2002; Gilad et al., 2006). Moreover, it was recently shown that these transcriptional differences are caused mainly by changes in non-coding regions of the DNA, and specifically in transcription factor binding sites (McLean et al., 2011; Shibata et al., 2012), indicating that changes in transcription regulation are driving the phenotypic changes.

Transcription regulation evolves through two types of changes. The first type follows the classical evolutionary theory, where random mutations in the DNA sequence can lead to rewiring of the transcription regulation network. These changes are inherited, exposed to selection pressures and eventually might fixate within the population. The mutations cause rewiring of the network when they lead to changes in cis or trans: Changes in trans-acting transcription factors can alter their DNA binding specificity, which might lead to recognition of different sets of target genes (Doniger and Fay, 2007;
Konopka et al., 2009; Ravasi et al., 2010; Yvert et al., 2003), as shown for the AP-1 transcription factor in yeasts (Kuo et al., 2010). Changes in cis-regulatory sequences at promoter and enhancer regions affect the binding affinities of proteins at a specific genomic position, and can lead to binding of different sets of transcription factors, chromatin remodelers or even change the chromatin structure (Borneman et al., 2007; Bradley et al., 2010; Gasch et al., 2004; Schmidt et al., 2010; Tanay et al., 2005; Tuch et al., 2008). The current view is that changes in cis are more common, although this is still a subject of much debate (Tirosh et al., 2009; Wang et al., 2007). It is possible that, compared to changes in trans, rewiring of the regulatory network through changes in cis-elements allows for subtler fine-tuning by local changes in the network. While regulatory proteins and their DNA binding domains are often highly conserved (Schmidt et al., 2010; Tuch et al., 2008; Wapinski et al., 2007), many rewiring events in regulatory networks occur through changes in cis-regulatory elements (Khaitovich et al., 2006; Tirosh et al., 2009; Wilson et al., 2008; Wittkopp et al., 2008). Such large scale changes in cis-elements were observed for specific factors in organisms as diverse as yeasts (Borneman et al., 2007; Doniger and Fay, 2007; Tuch et al., 2008), flies (Bradley et al., 2010; Moses et al., 2006), and mammals (Odom et al., 2007; Schmidt et al., 2010).

Epigenetic regulatory mechanisms, such as DNA methylation, chromatin modifications and non-coding RNAs, provide an additional potential driving force in the evolution of transcription regulation, which can lead to transgenerational reprogramming of gene-expression. Unlike the transcription factors-mediated regulation described above, such reprogramming does not require changes in the DNA sequence. This evolutionary scheme requires variation and stable inheritance of epigenetic traits, however, it differs from the classical evolutionary view that is based on random mutations in the DNA sequence as the carrier of information. In the past few decades there has been an important expansion of our understanding of inheritance, as a wide variety of epigenetically inherited traits have been described. Interestingly, since the environment has a direct effect on epigenetic factors, epigenetic inheritance implies that information about the environment experienced by parents could be transferred to their offspring by non-Mendelian mechanisms (Jablonka and Lamb, 2007; Jablonka et al., 1995). Thus, this
provides a highly effective mechanism to modulate gene-expression in the short term of just one or two generations.

1.6 Overview

In this work I studied the evolution of transcription regulation from these two different perspectives. In the first chapter I discuss cis-regulatory evolution, and describe a cross species comparative study in yeasts that addresses central questions regarding the evolution of cis-regulatory networks. This work was part of a collaborative effort, in which Dr. Ilan Wapinski (while doing his PhD in The Broad Institute of MIT/Harvard) and I developed a novel computational scheme and applied it to yeasts. I then conducted extensive data analysis and derived models (Habib et al., 2012). In addition, I discuss the analysis I have done within a collaborative effort lead by Prof. Nicholas Rhind (University of Massachusetts) to study fission yeasts (Rhind et al., 2011b). Throughout these works I received guidance from my two advisors, Prof. Hanah Margalit and Prof. Nir Friedman, and from Prof. Aviv Regev (The Broad Institute of MIT/Harvard). In addition, I took part in other collaborative works, not detailed in this dissertation, where my focus was on developing computational methods for analysis of regulatory networks and dynamic gene expression data both in yeasts and in mammals (Capaldi et al., 2008; Habib et al., 2008; Novershtern et al., 2011; Sivriver et al., 2011).

In the second chapter I discuss epigenetic inheritance, and describe an experiment in mice aimed to test the existence of transgenerational environmental reprogramming of gene-expression and a search for the epigenetic ‘carrier’ of the environmental information. This work was done in collaboration with Prof. Oliver Rando (University of Massachusetts) and his experimental lab. Oliver designed the experiment and conducted, with several students in his lab, large-scale and extensive experiments. My contribution to this work was in the computational analysis and interpretation of the results, under the guidance of my advisor Prof. Nir Friedman (Carone et al., 2010).
Chapter 2 –

A Functional Selection Model Explains Robustness Despite Plasticity in cis-Regulatory Networks

2.1 Introduction

The first part of this dissertation focuses on evolution of transcription regulation, driven by mutations in the DNA sequence. Specifically it regards cis-regulatory evolution, which refers to changes in cis-regulatory elements in the DNA that affect binding of transcription factors and can lead to rewiring of the regulatory network.

2.1.1 Rewiring of regulatory networks through changes of cis regulatory elements

Changes in cis-regulatory elements in genes’ promoters can have diverse effects on the regulatory network. On the one hand, such changes can lead to fine-grained regulatory ‘tinkering’ of the regulation 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 yeasts species (Tuch et al., 2008). However, the factor still regulates the cell cycle and mating processes 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 sets of gene (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 (Hogues et al., 2008; Tanay et al., 2005).

The connection between rewiring of regulatory networks and changes in gene expression is unclear. Previous studies on gene modules in bacteria (Isalan et al., 2008) and yeasts (Hogues et al., 2008; Tanay et al., 2005; Tsong et al., 2006; Tuch et al., 2008; Weirauch and Hughes, 2010c; Wohlbach et al., 2009) showed that while some regulatory
changes (e.g., in the control of mitochondrial ribosomal protein encoding genes (Tsong et al., 2006)) can be coupled to a transcriptional and phenotypic change, many other dramatic re-wiring events (e.g., in ribosomal proteins (Hogues et al., 2008; Tanay et al., 2005) or mating genes (Tuch et al., 2008)) have little or no apparent impact (reviewed in (Wohlbach et al., 2009; Weirauch and Hughes, 2010)). For example, over 40% of the binding events of four orthologous liver-specific transcription factors in mouse and human are species-specific (Odom et al., 2007), but the liver-associated function of the factors and the liver-specific expression of their target genes are highly conserved (Odom et al., 2007). This demonstrates the complexity of the regulatory system and raises important questions regarding the implications of the plasticity in regulatory networks, and specifically the implications on the functions of transcription factors (i.e. which cellular processes do they control).

The mechanism leading to a coordinated loss or gain of a transcription factor’s binding sites in many functionally related genes is unclear, especially when the gene expression does not change. Analysis of specific regulatory programs led to different suggested mechanisms for this dynamic evolutionary process (Dermitzakis et al., 2003; Gasch et al., 2004; Ihmels et al., 2005; Tanay et al., 2005), calling for a comprehensive study of this question. While individual examples of cis-regulatory evolution 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, its functional implications and the underlying selection forces. A large-scale unbiased study of the evolutionary history of regulatory networks, by a cross-species comparative analysis of regulatory networks in extant species, will advance us toward this goal.

2.1.2 Experimental methods to study cis-regulatory evolution

Rigorously studying cis-regulatory evolution has been hampered by the lack of large-scale and systematic experimental studies (Wohlbach et al., 2009). One major
obstacle is the limited data on transcription factor-target interactions in non-model organisms. Genome-wide experimental determination of factor-target interactions can be conducted by a couple of approaches. One approach is location analysis (chromatin immunoprecipitation followed by micro-arrays assay or DNA sequencing) (Ren et al., 2000), measuring directly where a transcription factor binds to the DNA. Another approach regards genetic perturbations, finding direct and indirect targets by measuring changes in expression levels of genes in response to a knockout or over-expression of a transcription factor (Amit et al., 2009; Capaldi et al., 2008; Chua et al., 2006; Horton et al., 2003). The main caveat of the first approach is that it may lead to false targets due to spurious and non-functional binding to the DNA. The major limitation in the second approach is that direct and indirect target genes are indistinguishable. Overall, in all experimental methods, measuring targets of dozens of factors across dozens of species is prohibitively expensive and labor intensive. In addition, since the regulation of a gene by a transcription factor is specific to the cell’s state, a complete characterization of target genes requires many experiments under different environmental conditions.

There are few studies that measured the binding of one or a few transcription factors across two or three yeast species (Borneman et al., 2007; Hogues et al., 2008; Tuch et al., 2008), flies (Bradley et al., 2010; Moses et al., 2006), or mammals (Konopka et al., 2009; Odom et al., 2007; Schmidt et al., 2010), showing in all cases extensive rewiring of the regulatory networks, even within closely related species. These intriguing anecdotal examples on the role that cis-regulatory divergence plays across evolution, call for extending such studies to the entire repertoire of transcription factors across dozens of species.

2.1.3 Computational methods to study cis-regulatory evolution

A possible alternative is to computationally predict regulatory interactions of transcription factors and their target genes from widely available genome sequences of many species. Such predictions require a DNA motif model of the sequence binding preferences of each transcription factor. This model can then predict the factor’s potential
binding sites across the genome. This initial mapping indicates which factors can bind to
the DNA at a given location and consequently are potential regulators of proximal genes.
The association between DNA motifs and target genes is the basic computational scheme
for constructing a full regulatory network from DNA sequence data and a catalogue of
DNA motif models. Thus, DNA motifs can be viewed as a compact and informative
representation of the building blocks of the regulatory network.

Modeling the sequence preferences of DNA-binding proteins with DNA motif
models, can be done in several different ways, most of which rely on a set of known
binding sites (Benos et al., 2002; Bulyk et al., 2001; Day and McMorris, 1992; Osada et
al., 2004; Stormo, 2000). A common representation, which benefits from being relatively
simple yet flexible, is a matrix of positions in the binding site versus nucleotides. In the
matrix each row represents one residue (A, C, G or T), and each column represents a
position in a set of aligned binding sites. All matrix representations assume that the
choice of nucleotides in each position of the motif is independent of all other positions.
Such a matrix representation that is widely used is a Position Weight Matrix (PWM),
which contains nucleotide frequencies in each position of the motif.

To learn DNA motif model of a specific transcription factor requires an aligned
set of its known binding site. Due to the lack of known sites for many factors, different
algorithms were developed for the identification of transcription factor DNA motifs (e.g.
(Bailey and Elkan, 1995; Hughes et al., 2000a; Liu et al., 2002; Siddharthan et al., 2005)).
Most algorithms identify statistically significant overrepresented sequence patterns in the
promoters of co-regulated genes, which are presumably binding sites of a specific
transcription factor, and require as input only promoter DNA sequences. Several analysis
pipelines were developed for such tasks, which output a non-redundant set of statistically
significant motifs (Gordon et al., 2005; Habib et al., 2008; Mahony et al., 2007). An
alternative approach is to use protein binding microarray technology to characterize in
vitro the transcription factor sequence specificities in a high-throughput manner
(Mukherjee et al., 2004). This method might suffer from artifacts since it is done in-vitro,
usually using only the DNA-binding domain of the factor.
To associate between motifs and target genes in the construction of a regulatory network, we computationally infer each transcription factor putative binding sites across a genome by scanning the genome for the corresponding binding motifs. The genes containing a motif instance in their promoters are termed here *motif targets*. Different scoring schemes have been used for such scans (Barash et al., 2005; Hughes et al., 2000a; Tanay, 2006). After inferring motif targets, we can determine the functional role of a transcription factor (or equivalently its DNA motif) according to the known functional annotation of its target genes. This requires functional annotations of genes, and improves our understanding of the regulatory network.

2.1.4 Current studies of cis-regulatory evolution

Computational methods can be used to conduct a large-scale study of *cis*-regulatory evolution. However, there are several drawbacks in the computational scheme described above: (1) Motif discovery algorithms have limited success rate and are not entirely robust to noisy inputs (Li and Tompa, 2006; MacIsaac and Fraenkel, 2006). Moreover, the co-regulated gene sets used as input are both noisy and incomplete. (2) Networks derived by computational methods are notoriously noisy, with both spurious and missing connections between transcription factors and their target genes. This is primarily because not all instances of the motifs in the genome are bound by the relevant factor and the bound instances are not necessarily functional (Capaldi et al., 2008). (3) Inferring the function of a transcription factor or a motif is affected by the noisy targets and is limited due to missing gene annotations. (4) This approach is limited to model organisms, due to the lack of known DNA regulatory motifs in non-model organisms.

The common approach to address these problems is to leverage evolutionary conservation to filter out spurious predictions of motifs, target genes and functions (Gasch et al., 2004; Kellis et al., 2003; Tanay et al., 2005). Evolutionary conservation is also used to find regulatory motifs in non model organisms with missing annotations (Gasch et al., 2004; Tanay et al., 2005). However, the conservation assumption is especially problematic when attempting to study divergence across species. Previous
studies (Cliften et al., 2003; Gasch et al., 2004; Ihmels et al., 2005; Kellis et al., 2003; Lavoie et al., 2010; Marino-Ramirez et al., 2006; Tan et al., 2007; Tanay et al., 2005) overcame these obstacles by focusing on at least one conserved feature, and tested divergence in the others. These conserved features include studying transcription factors with strongly conserved functions and target genes (Gasch et al., 2004), modules of orthologous genes with conserved expression patterns (Tanay et al., 2005), or binding sites whose relative positioning is conserved in individual promoters (Cliften et al., 2003; Gasch et al., 2004; Kellis et al., 2003) or classes of genes (Lavoie et al., 2010).

These studies (Cliften et al., 2003; Gasch et al., 2004; Ihmels et al., 2005; Kellis et al., 2003; Lavoie et al., 2010; Marino-Ramirez et al., 2006; Tan et al., 2007; Tanay et al., 2005) have found both conserved and diverged motifs associated with specific functions. Overall, they uncovered substantial plasticity in regulatory networks, with extensive turnover of motif targets and diverged location of binding sites within promoters of target genes. This is consistent with the experimental studies described above (Borneman et al., 2007; Hogues et al., 2008; Tuch et al., 2008) (Bradley et al., 2010; Moses et al., 2006) (Konopka et al., 2009; Odom et al., 2007; Schmidt et al., 2010). However, each computational work has made strong conservation assumptions to overcome noisy predictions, resulting in crude snapshots of a complex evolutionary process and biasing the results by the underlying assumptions of the computational method. Thus, an unbiased computational approach to reconstruct cis-regulatory evolution across large phylogenies for dozens of transcription factors is needed.

2.1.5 Yeast as a model for cis-regulatory evolution

The comparative studies described above were done in the Ascomycota fungi phylogeny (yeasts), which includes the known model organism Saccharomyces cerevisiae, the human pathogen Candida albicans and the remote Schizosaccharomyces pombe. Yeasts have proven to be an ideal model for studying transcription regulation and regulatory evolution. On the one hand, these are simple single cells eukaryotic organisms, easy to grow in the lab and manipulate genetically. They have a condensed genome with
4,000-7,000 genes and relatively short intergenic regions, simplifying computational analysis and models. Several yeasts species, mainly *S. cerevisiae*, have been extensively studied, and thus a lot of information is available, including fully sequenced and well-annotated genomes. On the other hand, yeasts share the same complex internal cell structure as higher eukaryotes, including similar transcriptional machinery and transcription regulation mechanisms. An extreme example is the Hsf1 transcription factor, which is highly conserved, including its DNA binding domain, from yeasts to mammals (Liu et al., 1997). Thus, yeasts are suitable for developing and testing new methodologies, and many of the principles discovered in them are potentially relevant to higher organisms as well. For a comparative study across species this phylogeny provides an optimal setting, since it includes dozens of fully sequenced genomes of highly diverse organisms, both in sequence and phenotype, spanning more than 800 million years of evolution.

In mammalian systems, transcription regulation is much more complex compared to yeasts. First, the intergenic regions are much longer, and transcription factors can bind to remote regulatory sequences, enhancers, which can be more than 100kb away from the genes they are modulating (Bejerano et al., 2006). Second, the number of transcription factors regulating a single response or biological process is large (Amit et al., 2009; Novershtern et al., 2011). An example for a complex regulatory system in mammals is the transcriptional response to inflammation in immune system cells in mice, which is regulated by at least a dozen transcription factors, operating through different modes of activation, including fast responding factors (e.g. NFkB) and secondary response factors synthesized de-novo during the response (e.g. Irf8), resulting in a wide range of dynamical transcriptional responses (Amit et al., 2009; Hoffmann et al., 2006; Medzhitov and Horng, 2009; Sivriver et al., 2011).

Thus, model organisms can be used to develop methodologies relevant to higher eukaryotes, but these require adjustments to account for the increased complexity of the system. Specifically when considering DNA motifs, the long intergenic regions can introduce an enormous amount of noise that will be difficult to overcome. A possible
alternative approach is to derive models based on observations in yeasts, and then directly
test these models in higher eukaryotes, to deduce general principles of transcription
regulation and regulatory evolution.

Here, we conduct a large-scale study of cis-regulatory evolution for dozens of
transcription factors across large phylogenies of yeast species. To this end we developed
an unbiased computational method and used it to address several questions: (1) What is
the extent of plasticity in regulatory networks? (2) What is the impact of the network’s
plasticity on the function of transcription factors? (3) What are the underlying selection
pressures driving this evolutionary process? (4) Can we find a general model relevant to
yeasts and mammalian species?
2.2 Results

2.2.1 CladeoScope: a framework for reconstructing cis-regulatory evolution

We developed CladeoScope (Figure 2), a computational framework for an unbiased reconstruction of 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 (Schmidt et al., 2010; Tuch et al., 2008; Wapinski et al., 2007). 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 of that clade was a target of the motif (Wapinski et al., 2007) (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.

An overview of the CladeoScope method

CladeoScope consists of four steps (Figure 2b): In Step 1- Initialization- CladeoScope is initialized with known DNA motifs (Position Weight Matrices) 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 of each clade in the phylogeny (Figure 3). These inferred **ancestral targets** within a clade are considered reliable (Figure 3). 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.

**Parsimonious 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 3). 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 (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, 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 1,000 reconstructions of ancestral targets for random sets of motif targets of the same size for each species, Methods).
2.2.2 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 (Maclsaac 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 4a, 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 4a). These clades include: the sensu stricto Saccharomyces (four species, clade A), the Kluyveromyces (four species, clade C), the 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 to 996,476 connections prior to phylogenetic filtering. For example, of the 307 predicted Gcn4 targets in S. cerevisiae, 195 pass our phylogenetic filter.

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 4b). 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-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 4b, red highlights), such as Gcn4 (amino acid biosynthesis), Rpn4 (proteasome), and Mig1 (glucose repression). 39% of motifs were detectable up to the last common ancestor (LCA) of the entire Ascomycota phylum (clade L), including those involved in cell cycle regulation (Fkh1, Swi6-MBP1, Figure 4b) and stress response factors (Hsf1, STRE, Figure 4b, 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., 2011b), 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 *S. 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: when a motif is detectable in a species,
the ortholog of its known cognate factor is present in the same species, and vice versa (Figure 4b). In a minority of cases (15%), a factor is present in a species, but its cognate motif is not detectable, possibly due to lack of conserved targets within this species, or to substantial changes in the factor’s sequence specificity (Baker et al., 2011). For example, the Zap1 motif is detectable only up to clade D, despite the presence of its ortholog up to clade J, suggesting a possible change of its DNA binding specificity or a lack of any significant target conservation within the relevant clades. These cases demonstrate the limitations of our approach in tracing regulatory evolution when the factor’s binding specificity has diverged substantially, or when target turnover rate within a clade is very high. This can be alleviated if more binding profiles are measured in non-model organisms. In a few cases (12%), a motif is detectable in a species lacking a clear orthologous cognate factor. This may indicate a relic ‘pseudomotif’ that is present in a genome but no longer functional. However, in our case this is not very likely, since we require the conservation of the motif and its targets across a clade of species in which the promoter sequences diverged significantly. More likely, we detect a DNA motif without its factor due to faulty orthology resolution (e.g. the Sko1 motif in Schizosaccharomyces) or to multiple members of a transcription factor family with similar binding specificities (e.g. factors binding the CACGTG motif).

**Evaluation of the CladeoScope algorithm**

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, Methods, Appendix Note 1). For example, when 30% of the true targets were removed from the set of target genes provided to CladeoScope, CladeoScope has greater >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) (Figure 5, Appendix Note 1).
CladeoScope’s predictions are also highly robust to variation in its various parameters (Appendix 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 to 0.001 had little impact on the number of significant motifs per clade (Figure 6). 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 but weak binding sites to non-functional sites. We identify potential candidates for weak functional sites as ones with a 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 (Appendix 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 (Appendix Note 1 & 3).

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 (Appendix 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. We tested the contribution of the species-specific motif refinement process to the quality of CladeoScope. This refinement step generates a species-specific motif based on an input motif in the model organism S. cerevisiae (see Methods). While the distances between the motifs are small (measured by BLiC (Habib et al., 2008)), they increase with the distance between species (Methods, Correlation = -0.58, p-value=0.005, for the mean distance of all motifs in each species). Since the distances are small, we next compared motif targets predicted for the S. cerevisiae motif to prediction for the species-specific

![Figure 7. Comparison of motif targets predicted with the known S.cervisiae motif and the refined motif. A plot of the fraction of the shared predicted motif targets (blue), targets predicted by S. cerevisiae motif only (red), and targets predicted by the species-specific refined motif only (green), for different motifs and species (from left to right): Fkh1 motif in S.bayanus, K.lactis, C.albicans and S.pombe; Hsf1 motif in S.bayanus, K.lactis, C.albicans and S.pombe; Rpn4 motif in S.bayanus, K.lactis and C.albicans; Gcn4 motif in S.bayanus, K.lactis and C.albicans; Mbp1 motif in S.bayanus, K.lactis, C.albicans and S.pombe; Mig1 motif in S.bayanus, K.lactis and C.albicans.](image)
We find that even in species within the sensu-stricto clade that refinement of motifs does change the predicted targets by adding and removing targets. Finally, 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* (*MacIsaac et al., 2006*) and four other species (*Borneman et al., 2007*; *Tuch et al., 2008*) (*Appendix 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 to 55% and 10%, respectively in the
predicted motif targets in *S. cerevisiae* that are not conserved (Figure 8). These improved predictions are consistent for different thresholds for motif targets detection in each species (Figure 8).

### 2.2.3 Plasticity of regulatory networks in Ascomycota fungi

The vast majority of *cis*-regulatory elements in genes’ promoters are rapidly gained and lost across species. As a result, even at relatively short evolutionary distances, transcription factors both gain and lose a substantial portion of their targets.

**Widespread 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 9a,b). For example, less than half of the targets of Gcn4 in clade D (the LCA of pre- and post-whole genome duplication (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* & post-WGD, spanning *S. cerevisiae* and *K. lactis*) and in clade G (*Candida*), but with no statistically significant set of shared ancestral targets between these two clades (Figure 9c). Thus, although the motif likely existed in the 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 as associated with many regulatory DNA motifs.

**Fast turnover rates of motif targets**

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 (Methods). This model is akin to standard models of sequence character evolution (Felsenstein, 1981). The rates are expressed in terms of expected number of events per time unit (tU), where a time unit corresponds to the time in which one amino-acid substitution per protein coding sequence is expected on average. We found that motif targets are globally gained and lost at fast rates (Figure 10), 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 (Figure 10, Methods). This discrepancy in the rates is due to differences in the pool of targets.
versus non-targets (~4,000) 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 (~4,100), 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 11a, Methods). As an illustrative example, consider a motif with 150 targets in clade B. We expect, on average over all motifs, that in the descendant clade A this motif will have 210 targets, but only 38% of those targets will be ancestral ones (conserved from B). Turnover rates vary substantially among individual motifs. For example, the Hsf1 (heat shock factor) motif exhibits low rates of target gain and loss (Figure 11b), while variants of the CACGTG motif (bound by Pho4, Tye7, and Met28) have very high turnover rates (Figure 11c). On average we found that only 7% of a given motif’s targets in the sensu-stricto clade (clade A, Figure 4a) are expected to be conserved in the LCA of the phylogeny (clade L, Figure 4a), and only 16% of the targets were conserved since the LCA with the Candida clade (clade H, Figure 4a).
2.2.4 Functional evolution of transcription factors in Ascomycota fungi

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. Due to the large redundancy between functional annotations, such simple enrichment testing leads to numerous overlapping “functions”, which are hard to interpret and even more challenging to compare across clades. For example, examining Rpn4, a known regulator of the proteasomal genes in *S. cerevisiae*, we find more than 45 gene-sets enriched in the motif targets across clades, with different sets having different degrees of enrichment with motif targets across clades (conservation). These sets include categories such as: *Proteasome complex, Stress* and
Cytoskeleton, each found to be enriched with Rpn4 targets in different clades (Figure 12). A closer examination of the motif targets within these categories shows many overlaps (e.g., 76% of the targets annotated as Proteasome are also annotated as Stress genes).

Figure 12. From gene sets to functional modules. (a) Shown are all functional gene sets (rows) that are enriched (red) or not enriched (black), in each clade (columns) with Rpn4 motif targets. (b) The resulting functional modules and their enrichment with Rpn4 motif targets across clades.
To obtain a non-redundant description of motif functions, we developed an algorithm that clusters functional gene-sets by the fraction of associated motif targets shared between them. This procedure defines functional modules, each containing genes that share functional annotations and are ancestral targets of the same regulatory motif in at least one clade (Methods). Revisiting the Rpn4 example, we see that we have two functional modules: stress response and proteasome module conserved across species, and cytoskeleton module conserved mainly in clade C (Figure 12).

An additional example, Gcn4 targets in each clade are associated only with the amino acid metabolism module (Figure 13a). 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.

Compared to direct enrichment of individual gene sets, functional modules are a more concise, non-redundant and robust representation that can be easily compared across the phylogeny. We extensively evaluated the robustness and correctness of the functional module assignments. We show that the method is robust to different parameters in the algorithm, including the choices of motif detection threshold, the threshold over enrichment of functional categories with motif targets, and to the threshold for merging functional categories (Appendix Note 3). Furthermore, in support of 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; Appendix Note 3).

**Innovations through expansion and switch of functions**

In some cases turnover of target genes contributes to evolutionary innovation, by either expanding or switching the scope of functions ascribed to a regulatory DNA motif
Figure 13. Patterns of Functional Evolution of DNA motifs across clades.
(a) Examples of functional conservation and innovation patterns. In each case, the enrichment of motif target genes with different functional modules is shown across the clades (Blue: targets enriched in module; Black: not enriched; Grey: no ancestral targets in clade), demonstrating functional conservation of the Gcn4 motif (top), clade-specific innovation of the Rpn4 motif (middle), and functional switch of the Mig1 motif (bottom). Additional examples are shown in Figures 14-16. (b) Distribution of functional conservation patterns for cis-regulatory motifs. Pie chart of the fractions of motifs associated with complete functional conservation (dark grey), clade-specific innovation (light grey), or a functional switch (white).

(Figure 13). For 36% 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 14. Examples for functional innovations of DNA motifs across clades.

(a) Examples of functional conservation patterns for different motifs (as in Figure 13, the enrichment of target genes with different functional modules is shown across clades. Blue: clade targets enriched in module; Black: not enriched; Grey: no ancestral targets in clade). The motif in a representation of a sequence logo is shown next to the motif’s name. Demonstrating clade specific innovation in various clades (from top to bottom): the Hap4 motif with extracellular matrix and GTP binding; the Gln3 motif with nitrogen metabolism; the Skn7 motif with ribosome; the Fkh1 motif with the meiotic and ascospore formation modules; and functional switch in the uncharacterized Ynr063w motif, between peroxisome/aerobic metabolism, TCA cycle/aerobic respiration and glycolysis. (b) Example of turnover in the motif regulating a functional gene-set between clades. In each case, the enrichment of motif target genes with the relevant gene-set is shown across clades (Blue: clade targets enriched in module; Black: not enriched; Grey: no ancestral targets in clade). The name of the motif associated with the gene-set is indicated on the left. Demonstrated is a regulatory switch in the GTP-binding genes from Sfp1 motif in most clades, to Hap4 motif in clade K.
We find various innovations in different clades (Figure 13a, Figure 14-15). For example, the Rpn4 motif is associated with the proteasomal module in all clades (Mannhaupt and Feldmann, 2007), while in clade C (the Kluyvermyces species) it is also identified in genes of a cytoskeletal module (Figure 13a). 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 (Figure 14). This latter example involves a regulatory switch, as the regulation of GTP binding genes in all other clades is regulated by the Sfp1 motif (Figure 14b).

Another interesting innovation is of motifs expanding to new functions associated with clade specific genes. Such an example is the motif bound by Rtg3 in S. cerevisiae, associated with amino acid metabolism genes across the phylum. In fission yeast however, it is also enriched in genes responsive to various stresses (Figure 15a). Of the stress genes that have Rtg3 motifs in S. pombe, 36% are found only in the Schizosaccharomyces clade, and many are also associated with the Atf1 motif, a conserved regulator of the stress response (Figure 15b). Rtg3 does not have a detectable ortholog in the Schizosaccharomyces clade (Wapinski et al., 2007), but the motif recognized by Rtg3 in S. cerevisiae is clearly identifiable in fission yeast, suggesting that these regulatory motifs are more conserved than their binding proteins. We also found a similar acquisition of Schizosaccharomyces-specific genes by the Fkh1- and MBF-associated motifs, which regulate meiotic transcription in S. pombe (Abe and Shimoda, 2000; Lowndes et al., 1992). In particular, these two motifs were enriched with genes with antisense transcripts (Figure 15c). Antisense transcripts are RNAs transcribed from the antisense strand compared to a known coding gene and are thus complementarity to it. They have been identified in multiple eukaryotes, and there is evidence suggesting they have a regulatory role (Brunskill and Steven Potter, 2012; Guttman et al., 2010; Rhind et al., 2011a; Yassour et al., 2010). Most of the Fkh1/Mei4 target genes with antisense
transcripts (80%, 47 genes) are meiotic genes (Rhind et al., 2011a), the majority of which are specific to the Schizosaccharomyces clade (Figure 15c).

For 20% 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

Figure 15. Conserved regulatory motifs with clade specific target genes
A) The enrichment of gene functional modules regulated by the Rtg3-binding motif in 23 Ascomycota. This motif is enriched upstream of amino acid metabolism genes in all Ascomycota. However, in fission yeast, it is specifically enriched upstream of stress-response genes. S. cerevisiae (Scer), S. paradoxus (Spar), S. mikatae (Smik), S. bayanus (Sbay), C. glabrata (Cgla), S. castellii (Scas), K. waltii (Kwal), A. gossypii (Agos), K. lactis (Klac), D. hansenii (Dhan), C. guilliermondii (Cgui), C. lusitaniae (Clus), C. albicans (Calb), C. tropicais (Ctro), C. parapsillosis (Cpar), C. elongosporus (Celo), Y. lipolytica (Ylip), N. crassa (Ncra), A. nidulans (Anid), S. japonicus (Sjap), S. octosporus (Soct), S. pombe (Spom). B) Enrichment of Rtg3- and Aft1-binding sites in the promoters of stress response genes. Each row represents a gene. The strength of the strongest regulatory site upstream of the gene is indicated in the blue heat map. The expression of the gene in glucose depletion (gd) and early-stationery phase (es) relative to log phase is indicated in the blue-yellow heat map. Genes specific to the fission yeast clade are indicated in orange. C) Enrichment of Fkh2/Mei4- and MBF-binding sites in front of antisense-transcribed genes. As in B, but each row represents a gene with greater antisense than sense transcription. Gene associated with meiosis (Mata et al., 2002) are indicated in magenta.
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 13a**). 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) (**Figure 14a**).

**Extensive functional conservation of regulatory DNA motifs**

We observed functional conservation for a large fraction of the regulatory DNA motifs. 44% 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 with Amino-Acid biosynthesis module (described above, **Figure 13a**), the Hsf1 motif with a heat shock module across the entire phylogeny (**Figure 16**), and the Mbp1 motif with cell-cycle and DNA replication modules across the entire phylogeny (**Figure 16**). Furthermore, although in other cases the motif might gain or lose an association to functional modules during evolution, 80% of all the motifs have at least one conserved function across all clades (**Figure 13b**).

**A resource for studying regulatory evolution in Ascomycota**

The regulatory history of the 88 transcription factors across the *Ascomycota* phylum, including their specific target genes in each clade, their turnover rates and their functions, constitutes a valuable resource for future studies of regulatory evolution and of individual species, including human and plant pathogens. We provide this resource as a website (http://www.compbio.cs.huji.ac.il/OrthoMotifs), where a user can query individual motifs, or genes, and trace their evolutionary relationship at the species and clade level.
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, 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 17a). 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 17b).

Figure 17. Target turnover and the functional selection turnover model
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 to outside of the process), but not particular target genes within that process (Figure 17c).

To test this hypothesis, we used a likelihood ratio test to compare two alternative evolutionary models (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 to 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 17d). 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. 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 (Appendix Note 3).

The fit of our model is good in all thresholds and parameters. More specifically, we find a fit to the model (p-value<0.01 after Bonferoni correction for multiple hypothesis) for at least 91% of the functional modules when changing the different thresholds in the algorithm and input (as described in section 2). More specifically, ranging between 91% - 95% when changing the enrichment threshold, ranging between 94% - 96% when changing the merge threshold, and between 92% - 96% when changing the motif detection threshold.

The functional selection turnover model explains the number of highly conserved targets

Against the backdrop of rapid turnover, some motif targets remain highly conserved. For example, 25 of the Gcn4 targets have Gcn4 binding motifs in their promoters in every clade (out of an average of 130 Gcn4 targets per clade). Such conservation may reflect an important specific function of these particular genes; alternatively, a few conserved genes may be expected by chance, given the functionally constrained turnover rate of the motif and the size of the functional module. To distinguish between these possibilities, we performed simulations to estimate the probability of the observed number of highly
conserved targets under our functional selection model (Methods), assuming a differential turnover rate for the targets, based on their function but not based on their individual identity. We examined 20 regulatory DNA motifs that have ancestral targets broadly conserved across clades, such as Gcn4 (Figure 17e), Mbp1 and Rpn4.

For all motifs tested, we could not reject the null hypothesis that the observed number of highly conserved targets is consistent with the overall turnover rates according to the Functional Selection Turnover Model (p ≥ 0.5). Thus, even the number of highly conserved targets is consistent with selection at the module level rather than selection towards the individual function of each gene within the module.

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; Schmidt et al., 2010; Tuch et al., 2008)). 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, S. mikatae and S. bayanus) (Borneman et al., 2007), and HNF4α 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 (Methods, Figure 18), 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 (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).
2.2 Methods

2.3.1 CladeoScope algorithm: Phylogenetic reconstruction of Cis-regulatory networks

General Overview
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 position weight matrices (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 last common ancestor (LCA) of each clade of species across the phylogeny (Figure 2), 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/OrthoMotifs.
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 psuedocode.

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:

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

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; \( P_{BG} \) is the background distribution of nucleotides in
all promoters of the genome; $P_{PWM}^j$ is the probability vector for nucleotides in position $j$ of the motif and similarly $P_{r-PWM}^j$ 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 $T = 0.8*(\text{mean of 20 highest scoring promoters in the genome})$. 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 (Appendix 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 shows that our result are mostly robust to this choice (Appendix 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 less than 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 greater than 1,500 (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)).

**Optimization of the scanning method and threshold**

To find the optimal scanning method and threshold we compared two different methods for scoring putative instances of a DNA motif (binding sites):

1. **Max** – maximum log-likelihood scores over all possible positions along the promoter, where the log likelihood ratio score is the ratio between the probability of an individual K-mer within the promoter given the motif model and its probability in the background distribution of nucleotides across all promoters (Stormo, 2000).

2. **Sum** – Summing the likelihood scores over all possible positions on the promoter (Tanay, 2006):
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 in the promoter, \( P_{BG} \) is the background distribution of nucleotides in all promoters in the genome, \( P_{PWM} \) is the probability vector for nucleotides in position \( j \) of the motif and similarly \( P_{r-PWM} \) for the reverse motif (equivalent to searching the reverse strand of the DNA).

We compared different thresholds for target prediction:

1. Threshold on the \( p \)-value of the score (when using the \textit{Max} score): the probability of finding a score as high as this in random K-mers sampled from the background distribution (using compound importance sampling (Barash et al., 2005)). We tested \( p \)-values ranging between 0.01-0.001

2. Using a relative threshold per motif as the percent of the highest possible score for the specific motif in a specific genome. Here we use thresholds ranging between 70%-90% identity. The estimate of the highest achievable score for the motif is specific per genome, and defined as the mean of 20 highest scoring instances (or promoters) in the genome (this is important since we want to scan new genomes with a motif originating from a different species).

To optimize the choice of threshold, we compared the predicted targets to ChIP-chip assays in \textit{S. cerevisiae} (MacIsaac et al., 2006). We note that the two scoring methods (\textit{Max} and \textit{Sum}) are highly correlated in their assessment of individual promoters (R>0.95 for all motifs tested), and the resulting predicted target sets are similar as well. However, the choice of threshold does behave differently. For example, both for the Rpn4 and the Hsf1 motif we get an identical set of targets predicted based on 100% of the best \textit{Max} score or 75% of the \textit{Sum} score.

Since there is substantial evidence for transcription factors binding several weak binding sites in the promoter (\textit{e.g.} (Parker et al., 2011)), we preferred the \textit{Sum} method that takes such cases into account. We chose the threshold to be 80% of the best possible
Sum score (equivalent to ~70% of the Max score, which is similar to the threshold used in Harbison, et. al.(Harbison et al., 2004)). This is the lowest threshold that still had a high precision in the predictions. We see that for 75% threshold there is already a big decrease in accuracy. Additional tests for the effects of lowering this threshold on our predictions of ancestral targets and assignments of motifs to functional modules are included in Appendix Notes 1&3.

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 (Schmidt et al., 2010; Tuch et al., 2008; Wapinski et al., 2007). We therefore initiate our reconstruction with DNA motifs for known transcription factors that have been experimentally determined in model organisms. CladeoScope 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 re-scan the species’ genome and to identify a revised set of targets. CladeoScope 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. CladeoScope 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. CladeoScope 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 CladeoScope 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. CladeoScope 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 CladeoScope 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 1,000 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.

2.3.2 Resources for phylogenetic reconstruction in Ascomycota fungi

Gene, promoter annotations and DNA motifs

We acquired the genome sequences and annotations of the 23 Ascomycota species from the online Fungal Orthogroups (Butler et al., 2009; Cherry et al., 1997; Cliften et al., 2003; Dietrich et al., 2004; Dujon et al., 2004; Kellis et al., 2004; Kellis et al., 2003) (Arnaud et al., 2007; Galagan et al., 2005; Rhind et al., 2011b; Wood et al., 2002). Promoters were defined as the 600 bases upstream from the first codon, truncated at the neighboring coding sequence. To avoid bias in the motif discovery stage, we filter out stretches of poly-A or poly-T sequences of 5 bases or longer and poly-A/T sequences longer than 9 bases and replaced them with poly-N of the same length.

Motifs were assembled from TRANSFAC (Matys et al., 2006), protein microarrays (Zhu et al., 2009), and previous analysis of ChIP-chip data (MacIsaac et al., 2006). All motifs were transformed to a Position Weight Matrix (PWM) format (a n×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 in 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 *C. glabrata* and *S. castellii* were found to be inverted in some cases. 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.

### 2.3.3 Evaluations of CladecoScope in Ascomycota fungi

**Validating CladecoScope’s performance using synthetic data**

We generated simulated target sets in extant species for evaluating CladecoScope’s robustness 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 4a**): 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 4a) over all motifs: Fast $F_{\text{gain}}=0.002$ $F_{\text{loss}}=0.3$, Medium $F_{\text{gain}}=0.0002$ $F_{\text{loss}}=0.03$, Slow $F_{\text{gain}}=0.00002$ $F_{\text{loss}}=0.003$). Forth, we estimated the gain and loss frequencies for each of the three motifs in each relevant species directly from the data (as in the Likelihood Ratio Test described below) (Appendix Note 1). Fifth, we used two topologies of the species tree: the topology in the sensu-stricto clade (clade A, Figure 4a), and the asymmetrical topology in the Candida clade (clade E, Figure 4a). 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, comparing 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 Appendix 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-vivo ChIP-chip data in S. cerevisiae and four other species (Appendix Note 1).
2.3.4 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 (tU), where a time unit corresponds to the time in which 1 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:

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

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

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

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

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 ($N_{gain}$), lost ($N_{loss}$) and conserved ($N_{cons}$) target genes, and the probabilities described above, as:

$$\text{lnLikelihood(Motif)} = \sum_{b \in \text{branches}} \left[ N_{gain}^b \ln P(Gain \mid t^b) + N_{loss}^b \ln P(Loss \mid t^b) + N_{cons}^b \ln(1 - P(Loss \mid t^b)) + N_{non-target}^b \ln(1 - P(Gain \mid t^b)) \right]$$

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).
2.3.5 Annotating motifs with functional modules and their evaluation

**The functional modules algorithm**

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.

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.

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*, Nature 2007), it was shown that when it is possible to 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.

**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 transcription-modules (Segal et al., 2003), MIPS (Mewes et al., 2010), KEGG (Kanehisa and Goto, 2000; Kanehisa et al., 2006), and mutant phenotypes (Hughes et al., 2000b). 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 Appendix 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 bests 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 in this chapter; and (4) Robustness of the functional annotations of motifs by the functional modules, where we examined in detail 18 motifs including those discussed in this chapter. In addition, we compared the resulting functional modules to known motif and transcription factor annotations from the literature in *S.cerevisiae (SGD)*, *C.albicans (CGD)* and *S.pombe* (GeneDB).
2.3.6 The functional selection turnover model

Likelihood Ratio Test

We used a Likelihood Ratio Test (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:

$H_0$: 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).

$H_1$: 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 ($N_{\text{gain}}$), lost ($N_{\text{loss}}$) and conserved ($N_{\text{cons}}$) target genes. In the functional selection model ($H_1$ hypothesis) we computed the gain and loss probabilities separately for genes within the module and genes outside of the module.

Description of the equations:

$LRT(\text{Motif}) = ll(\text{Motif} \mid H_0) - ll(\text{Motif} \mid H_1)$

$ll(\text{Motif} \mid H_0) = \sum_{b \in \text{branches}} ll(b \mid H_0)$

$ll(b \mid H_0) = N_{\text{gain}} \ln P(Gain \mid H_0) + N_{\text{loss}} \ln P(Loss \mid H_0) + N_{\text{cons}} \ln [1 - P(Loss \mid H_0)] + N_{\text{nonTarget}} \ln [1 - P(Gain \mid H_0)]$

$ll(b \mid H_1) = N_{\text{gain}}^{\text{in}} \ln P(Gain^{\text{in}} \mid H_1) + N_{\text{loss}}^{\text{in}} \ln P(Loss^{\text{in}} \mid H_1) + N_{\text{cons}}^{\text{in}} \ln [1 - P(Loss^{\text{in}} \mid H_1)] + N_{\text{nonTarget}}^{\text{in}} \ln [1 - P(Gain^{\text{in}} \mid H_1)]

+ N_{\text{gain}}^{\text{out}} \ln P(Gain^{\text{out}} \mid H_1) + N_{\text{loss}}^{\text{out}} \ln P(Loss^{\text{out}} \mid H_1) + N_{\text{cons}}^{\text{out}} \ln [1 - P(Loss^{\text{out}} \mid H_1)] + N_{\text{nonTarget}}^{\text{out}} \ln [1 - P(Gain^{\text{out}} \mid H_1)]$
The maximum likelihood estimation for the probability of gain and loss of each motif’s target genes in the current clade $C_1$, compared to the immediate ancestral clade $C_p$:

$$P(Gain \mid H_0) = \frac{N_{gain}}{N_{total} - T_{C_p}}$$

$$P(Loss \mid H_0) = \frac{N_{loss}}{T_{C_p}}$$

$$P(Gain^{IN} \mid H_1) = \frac{N_{gain}^{IN}}{N_{total}^{IN} - T_{C_p}^{IN}}$$

$$P(Loss^{IN} \mid H_1) = \frac{N_{loss}^{IN}}{T_{C_p}^{IN}}$$

Where,

$N_{total}$ = Total number of genes in the genome

$T_{C_p}$ = Total number of targets in clade $C_p$

$IN$ = Genes belonging to the functional module

$OUT$ = Genes not belonging to the functional module

$N_{total}^{IN}$ = Total number of genes in the functional module

$T_{C_p}^{IN}$ = Total number of genes in the functional module that are targets in clade $C_p$

**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 sub-clades that share the same direct ancestral clade, and computed the overlaps between these simulated target sets. The simulations were initialized with the target set of the ancestral clade, and we simulated the targets in each sub-clade according to its probability of gain or loss of target genes, within and outside of the functional module (computed as described above for the Likelihood Ratio Test, using a maximum likelihood estimator). We repeated this
process 1,000 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 (1,000).

**Experimental transcription factor binding data**

We used Ste12 and Tec1 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 between 1kp to 5kp upstream of the transcription start site (sequences taken from UCSC genome Browser versions hg19 (Lander et al., 2001), canFam2 (Lindblad-Toh et al., 2011), mm10 (Waterston et al., 2002)). The specific list of target genes changes when we modify this parameter, but the fit to the functional turnover model does not.

To find functional modules we conducted the same analysis as described above, using the target gene enrichments from the individual species instead of the targets at the clades. For the LRT and simulation tests, we conducted the same analysis as described
above, but comparing targets of each individual species to the ancestral target set of all three species, defining the highly conserved targets as targets conserved in all three species. For mammals, we used gene functional annotations from MsigDB (Liberzon et al., 2011) (Release 3.0).
Chapter 3 -

Paternally Induced Transgenerational Environmental Reprogramming of Metabolic Gene Expression in Mammals

3.1 Introduction

My second focus in this work is on evolution of transcription regulation driven by epigenetic changes. Specifically, on transgenerational reprogramming of gene expression by epigenetic inheritance, and the interplay between the environment and such reprogramming.

3.1.1 Epigenetic Inherence and the Environment

Inheritance of epigenetic regulatory factors, such as DNA methylations, chromatin modifications and non-coding RNAs can lead to transgenerational reprogramming of gene-expression. Epigenetic inheritance mechanisms are potential carriers of information about the environment experienced by parents to their offspring (Jablonka and Lamb, 2007; Jablonka et al., 1995). Theoretical studies imply that environmental regimes exist for which “carryover” epigenetic memory would be adaptive (Jablonka and Lamb, 2007; Jablonka et al., 1995). In other words, mechanisms exist that could allow organisms to “inform” their progeny about prevailing environmental conditions. Under certain historical circumstances – for example, repeated exposure over evolutionary time to a moderately toxic environment that persists for tens of generations – such non-Mendelian information transfer would be adaptive (Jablonka et al., 1995; Rando and Verstrepen, 2007). Whether or not organisms can inherit characters induced by ancestral environments has far-reaching implications, and this type of inheritance has come to be called “Lamarckian” inheritance after the early evolutionary theorist J. Lamarck. However, there is scant evidence for trans-generational effects of the environment in mammals.
3.1.2 Evidence for Trans-Generational Effects of the Environment

A small number of cases have been described in which phenotype of an organism differs depending on the environment experienced by the parents. This is most commonly seen as a maternal effect. In a variety of rodents, information about photoperiod can be passed on to offspring by mothers, and cross-fostering experiments show that this information is transmitted in utero (Horton, 2005). In worms, osmotic stress applied in one generation results in offspring with increased resistance to osmotic stress, but decreased resistance to anoxia (Frazier and Roth, 2009). In this case, the maternal effect appears to function via altered sugar metabolism – offspring of osmotically-stressed worms have less glycogen, but more glycerol, than offspring of unstressed worms. In human populations, epidemiological data, particularly from the Dutch “Hunger Winter” of World War II, suggests that children whose mothers went hungry during pregnancy have significantly increased rates of diabetes, obesity, and cardiovascular disease (Hales and Barker, 2001; Lumey et al., 2007).

The existence of effects of the maternal environment on phenotype is not particularly surprising, as the womb is a baby’s first environment – for example, fetal alcohol syndrome is a phenotypic consequence of excessive maternal alcohol intake, and conceptually does not require epigenetic information to mediate the phenotype. Thus, demonstration of multi-generational changes is important in maternal effects to rule out simple plastic responses of offspring to the in utero environment.

3.1.3 Evidence for Heritable Epigenetic Effects of Environmental Perturbations

Some studies have demonstrated heritable epigenetic effects of environmental perturbations on offspring. For example, treatment of gestating rat mothers with the endocrine disruptor vinclozolin results in decreased fertility and behavioral changes in several generations of offspring (Anway et al., 2005; Crews et al., 2007; Skinner et al., 2008). The authors suggested that the mechanism for inheritance was induction of
changes in cytosine methylation patterns, although genetic alterations to the Y chromosome were not ruled out. In another study, withholding methyl donors (such as folate) from pregnant female mice resulted in decreased cytosine methylation across a transposable element inserted in the agouti gene (Waterland and Jirtle, 2003). While an effect in the first generation might simply reflect a direct environmental influence, the altered cytosine methylation profile persisted well beyond the first generation (Cropley et al., 2006) and was even transmissible through the male germ line. In neither of these cases is the detailed mechanism of transgenerational heritability understood.

While in principle these effects may be passed through the maternal or paternal germline, in practice it is quite difficult in females to separate epigenetically heritable effects from plastic responses of the progeny to its environment. Fathers, on the other hand, often have very little direct influence on their offspring’s environment, especially in mice. If the paternal environment affects the progeny, such effects are likely to act through the germ line. A handful of paternal effects have been documented in the literature – preconception fasting of male mice has been reported to affect serum glucose in offspring (Anderson et al., 2006), and epidemiological data from human populations links hunger in paternal grandfathers to obesity and cardiovascular disease two generations later (Kaati et al., 2002; Pembrey et al., 2006).

It is therefore of great interest to determine what environmental conditions have transgenerational effects in mammals, and to characterize the mechanisms that mediate these effects. To test whether such transgenerational inheritance occurs in mammals, we carried out a screen for genes and epigenetic modifications in mice that responded to paternal diet. Relative to the offspring of males fed a control diet, the offspring of males fed a low-protein diet increased the expression of many genes involved in lipid and cholesterol biosynthesis, and had increased levels of cholesterol esters, triglycerides, and free fatty acids, lipids.
3.2 Results

3.2.1 Experimental paradigm

Male mice were fed control or low protein diet (11% rather than 20% protein, with the remaining mass made up with sucrose. While the relevant dietary change in this experiment could be protein content, sucrose content, fat/protein ratio, etc., for simplicity we refer to the diet as low protein throughout the text) from weaning until sexual maturity. They were then mated to females reared on control diet (Figure 19A, 20A). Fathers were removed after one or two days of mating, limiting their influence on their progeny to the mating itself. All mothers were maintained on control diet throughout the course of the experiment. After birth, the offspring were reared with their mothers until three weeks old, at which point their livers were harvested for RNA isolation. DNA microarrays were used to profile global gene expression differences in the livers of the offspring from the two types of crosses.

Testing for differences between 26 matched pairs of mice from the two F1 groups, we found a significant overabundance of differentially-expressed genes, relative to the null hypothesis that the parental treatment does not affect offspring (1,595 genes at false discovery rate – FDR – of 0.001, Figure 20B-C). We also identified a more robust (t-test with null hypothesis of mean change 0.2, FDR of 0.01) group of 445 genes whose expression strongly depended on the diet consumed by their fathers (Figure 19B). In our analysis we focus on this more robust group of genes, however, all the phenomena described below are true for the larger group as well. These gene expression changes were observed in 13 (7 low protein, 6 control) litters in experiments spanning several years, carried out in three different animal facilities. In principle, random factors should be distributed equally between our two groups given the numbers of offspring examined, but we directly address a number of potential artifacts nonetheless, including changes in cell populations, circadian cycle, litter size, order of sacrifice, and cage location.
Figure 19. A screen for genes regulated by paternal diet

(A) Experimental design. Male mice were fed control or low (11%) protein diet from weaning until sexual maturity, then were mated to females that were raised on control diet. Males were removed after 1 or 2 days of mating. Livers were harvested from offspring at 3 weeks, and RNA was prepared, labeled, and hybridized to oligonucleotide microarrays. (B) Overview of microarray data, comparing offspring of sibling males fed different diets—red boxes indicate higher RNA levels in low-protein than control offspring, green indicates higher expression in controls. Boxes at the top indicate comparisons between two male (purple) or two female (yellow) offspring. Each column shows results from a comparison of a pair of offspring. Only genes passing the stringent threshold for significant change are shown. Data are clustered by experiment (columns) and by genes (rows). (C) Validation of microarray data. Quantitative RT-PCR was used to determine levels of Squalene epoxidase (Sqle) relative to the control gene Vitronectin (Vtn), which showed no change in the microarray dataset. Animals are grouped by paternal diet and by sex, and data are expressed as DCT between Sqle and Vtn, normalized relative to the average of control females. p values were calculated using t test.
We confirmed our results by q-RT-PCR (Figures 19C, 20A). Squalene epoxidase (Sqle), which catalyzes the first oxygenation step in sterol biosynthesis, exhibited a ~3-fold increase in the low protein cohort in our microarray data, and q-RT-PCR showed a similar average expression difference across over 25 animals, gathered in crosses carried out several years apart (Figure 19C). The differences we observe occur in both male and female progeny (Figures 19C), though these dietary history-dependent differences are superimposed on a baseline of differential expression between the sexes.

Figure 20. Validation and identification of differentially expressed genes
(A) Microarray data and q-RT-PCR results are shown for the indicated genes, for two offspring comparisons. (B and C) Evaluating the statistical significance of the number of genes that are differentially expressed between offspring of low-protein diet fathers and control diet fathers. Blue line, the number of differentially expressed genes that separate the two sets of offspring (y axis) that were scored a given p value (x axis) in a t test; black line, the number of genes expected by chance with that p value from 1000 simulations with random reshuffling of subject labels; light gray or red line, the range of numbers of differentially expressed genes in the 95th percentile of 1000 random simulations. Overabundance of differentially expressed genes is observed when using both tests: (B) combination of two one-tailed t tests; (C) combination of two one-tailed t tests using a null hypothesis with mean change of 0.2. In this case the random reshuffling of the data corresponds to a null hypothesis with mean 0 rather than 0.2 and thus is an upper bound on the number expectancy by chance.
3.2.2 Upregulation of proliferation and lipid biosynthesis genes in low protein offspring

To help define the physiological differences between our cohorts, we calculated enrichments of various Gene Ontology (GO) processes in the differentially expressed genes. Genes upregulated in our treatment group’s offspring were enriched for a number of categories of genes involved in fat and cholesterol biosynthesis, including lipid biosynthesis (p < 9 x 10^{-26}), steroid biosynthesis (p < 3 x 10^{-19}), cholesterol biosynthesis (p < 2 x 10^{-12}), and oxidation-reduction (p < 4 x 10^{-10}). Another major group of upregulated genes are annotated to be involved in S phase, such as DNA replication (p < 2 x 10^{-9}) and related annotations. Downregulated genes were enriched for GO annotations such as sequence specific DNA binding (p < 6 x 10^{-6}) and ligand-dependent nuclear receptor activity (p < 6 x 10^{-5}), although the number of genes matching these annotations was small (14 and 5, respectively).

The increase in S phase genes likely indicates a hyperproliferative state, while the metabolic expression differences suggest that lipid metabolism is altered in these animals. To explore the mechanisms responsible for these altered gene expression programs, we asked whether the observed gene expression differences might reflect altered regulation of a small number of pathways. We checked for significant overlaps of the gene expression profile observed in our low protein offspring with a compendium of 120 publicly-available murine liver gene expression datasets (Experimental Procedures). Our low protein offspring gene expression profile significantly (p < .05 after Bonferroni correction) overlapped gene expression changes from 28 published profiles (Figure 21), including gene expression profiles associated with perturbation of transcription factors that regulate cholesterol and lipid metabolism (SREBP (Horton et al., 2003), KLF15 (Gray et al., 2007), PPARα (Rakhshandehroo et al., 2007), and ZFP90 (Yang et al., 2009)). Our gene expression dataset also significantly matched hepatic gene expression in a variety of mice with mutations affecting growth hormone (GH) and insulin-like growth factor 1 (IGF-1) levels (Boylston et al., 2004; Madsen et al., 2004; Tsuchiya et al., 2004). Hierarchical clustering according to the enriched public profiles revealed two types of
prominent gene functions in our data: DNA replication ($p < 6 \times 10^{-14}$) and lipid or cholesterol biosynthesis ($p < 2 \times 10^{-27}$) (Figure 21). The partial overlap observed with each of many different transcription factor and growth factor profiles suggests that the altered gene expression profile observed in low protein offspring is likely related to reprogramming of multiple distinct pathways.

To assess whether the reprogrammed state in offspring reproduces the paternal response to low protein diet, we measured global gene expression changes in the livers of pairs of animals weaned to control or low protein diet as in Figure 19A. Genes that change in offspring are not the same as the genes induced in the parental generation by these protocols (Figure 22). Instead, males fed the low protein diet upregulate immune response and apoptosis-related genes, and downregulate genes involved in carboxylic acid metabolism (analysis not shown).
3.2.3 Transgenerational effects on lipid metabolism

We further focused on cholesterol biosynthesis genes. Coherent upregulation of genes involved in cholesterol metabolism is observed in the offspring of low protein fathers (Figure 23A). Figure 23B shows a more detailed comparison between our upregulated dataset and published data (Schneider et al., 2003) for genes activated by a major transcriptional regulator of cholesterol metabolism, SREBP. Many of the genes upregulated in low protein offspring have previously been shown to be upregulated by overexpression of SREBP-1a or SREBP-2 or downregulated by loss of the SREBP-activating gene, Scap.

Figure 22. Analysis of Paternal Response to Low-Protein Diet. (A) Males were fed control or low-protein diet from weaning until sexual maturity, and then were sacrificed and livers were harvested for gene expression profiling as in Figure 19. Here, genes on both panels have the same order, showing gene expression differences as low-protein/control in fathers and offspring. Gene expression differences in offspring do not reflect the paternal response to the dietary regimes (note that these males were not fathers of the offspring analyzed in Figure 19, but were treated equivalently). (B) Scatterplot of average gene expression in offspring (x axis) versus in males treated with LP or C diet (y axis). Only genes were chosen with fewer than 30% missing spots in each experiment (26 arrays each). R = -0.129.
Figure 23. Altered Cholesterol Metabolism in the Low-Protein Cohort
(A) Cholesterol biosynthesis. Genes annotated as cholesterol biosynthesis genes are shown, with colors indicating average difference in expression in low-protein versus control comparisons. (B) Many genes upregulated in the low-protein cohort are SREBP targets. Upregulated cluster from Figure 19 is shown, along with data from Horton et al. (2003). Genes scored as up in both replicates from Horton et al. (2003) are shown as yellow, genes scored as down are blue. Columns show data from transgenic mice overexpressing SREBP-1a or SREBP-2 or from Scap knockout mice. (C) Cholesterol levels are decreased in livers of low-protein offspring. Data from lipidomic profiling of liver tissue from three control and three low-protein animals are shown as mean ± standard deviation. Red line indicates no change. p values were calculated using a paired t test on log-transformed lipid abundance data. Cholesterol esters, CE; phosphatidylethanolamine, PE; free cholesterol, FC; triacylglycerol, TAG; phosphatidylcholine, PC; cardiolipin, CL; phosphatidylserine, PS; free fatty acid, FA; lysophosphatidylcholine, LYPc; and diacylglycerol, DAG.
To explore the correspondence between hepatic gene expression and physiology, we measured lipid levels in three pairs of control and treatment livers to determine whether increased levels of lipid biosynthesis genes resulted from changes in lipid levels (Figure 23C). Livers in the cohort with low protein diet fathers were depleted of cholesterol and cholesterol esters (whose levels were reduced more than two-fold). Additional differences were found in specific lipid classes, such as substantial increases in relative levels of saturated cardiolipins, saturated free fatty acids, and saturated and mono-unsaturated triacylglycerides in low protein offspring. Together, these results demonstrate that paternal diet affects metabolites of key biomedical importance in offspring.

3.2.4 MicroRNAs in offspring

Small (19-35) RNAs such as microRNAs (miRNAs) have recently been implicated in epigenetic inheritance in mice (Wagner et al., 2008). To determine whether altered small RNA populations might drive our reprogramming effect, we characterized the small (19–35 bp) RNA population from control and low protein offspring livers by high throughput sequencing (Ghildiyal et al., 2008), and mapped reads to known microRNAs. A number of miRNAs changed expression in the offspring from low protein diet fathers (Figure 24). Changes were often subtle in magnitude (~50%), but were reproduced in four control vs. low protein comparisons (paired t-test), and given the number of sequencing reads obtained for these RNAs this magnitude of difference is well outside of counting error. Offspring of low protein cohort upregulated miR-21, let-7, miR-199, and miR-98, and downregulated miR-210. Many of these upregulated miRNAs are associated with proliferation in liver, with miR-21 and miR-199 both associated with hepatocellular carcinoma (Jiang et al., 2008), while let-7 is well-known as a tumor suppressor (Jerome et al., 2007). The increase in growth-associated miRNAs is consistent with the hyperproliferative gene expression profile observed in the offspring of low protein diet fathers.
We found no statistically-significant overlap (p > 0.05) between the predicted targets of the miRNAs here and the gene expression changes we observe, though the subtle (~50%) changes in miRNA abundance we observe might be expected to have little effect on mRNA – even when specific miRNAs are artificially introduced in cells, downregulation of target mRNAs is less than 2-fold for the majority of predicted targets (Hendrickson et al., 2008). Our results therefore suggest that miRNAs are likely to be additional targets of the reprogramming pathway, yet are likely not the direct upstream regulators of the entire response (but see (Yotova et al., 2008)).

3.2.5 Cytosine methylation in offspring

How are offspring reprogrammed by paternal diet? Cytosine methylation is a widespread DNA modification that is environmentally-responsive, and carries at least some heritable information between generations (Bartolomei et al., 1993; Cropley et al., 2006; Holliday, 1987; Rakyen et al., 2003; Waterland, 2003). We performed reduced
representation bisulfite sequencing (RRBS, (Meissner et al., 2008)) to characterize cytosine methylation at single nucleotide resolution across ~1% of the mouse genome. RRBS was performed for livers from a pair of control and low protein offspring, and fraction of methylated CpGs was calculated for a variety of features such as promoters, enhancers, and other nongenic CpG islands. In general, we found that cytosine methylation was well-correlated between control and low protein offspring. However, we did observe widespread modest (~10-20%) changes in CpG methylation between the two samples, consistent with many observations indicating that environmental changes tend to have small quantitative effects on cytosine methylation (Blewitt et al., 2006; Heijmans et al., 2008; Ng et al., 2010; Weaver et al., 2004). Importantly, changes in promoter methylation did not globally correlate with changes in gene expression in offspring, indicating that the gene expression program in offspring is unlikely to be epigenetically specified at each individual gene. Of course, widespread gene expression differences can be caused by changes to a small number of upstream regulators, and a number of differentially-methylated regions are associated with cholesterol or lipid-related genes.

Most interestingly, we found a substantial (~30%) increase in methylation at an intergenic CpG island ~50 kb upstream of Ppara. This locus is likely an enhancer for Ppara, as it is associated with the enhancer chromatin mark H3K4me1 (Heintzman et al., 2007) in murine liver (F. Yue and B. Ren, personal communication). Ppara is downregulated in the majority (but not all) of offspring livers, and the overall gene expression profile in our offspring livers significantly matches the gene expression changes observed in ppara knockout mice (Figure 21), suggesting that epigenetic regulation of this single locus could drive a substantial fraction of the observed gene expression changes in offspring. Indeed, variance of Ppara mRNA levels alone can be used to explain ~13.7% of the variance in the entire gene expression dataset (although this of course does not determine causality).

We therefore assayed the methylation status of this locus by bisulfite sequencing in an additional 17 offspring livers (8 control and 9 low protein), finding average differences of up to 8% methylation between low protein and control livers at several
CpGs in this locus. Importantly, these pooled data underestimate the potential role of this locus in reprogramming as they include animals exhibiting a range of changes in Ppara gene expression – individual animals exhibit differences of up to 30% at various cytosines across this locus. Taken together, these results identify a differentially-methylated locus that is a strong candidate to be one of the upstream controllers of the hepatic gene expression response.

3.2.6 Cytosine methylation, RNA, and chromatin in sperm

To globally investigate effects of paternal diet on sperm cytosine methylation, we isolated sperm from four males – two consuming control diet, one consuming low protein diet, and one subjected to a caloric restriction regimen. We then surveyed cytosine methylation patterns across the entire genome via MeDIP-Seq (immunoprecipitation using antibodies against 5me-C followed by deep sequencing (Jacinto et al., 2008; Weber et al., 2005)). Notably, global cytosine methylation profiles were highly correlated between any pair of samples, indicating that the sperm “epigenome” is largely unresponsive to these differences in diet. This lead us to consider alternative epigenetic information carriers including RNA (Rassoulzadegan et al., 2006; Wagner et al., 2008) and chromatin (Arpanahi et al., 2009; Brykcynska et al., 2010; Hammoud et al., 2009; Ooi and Wood, 2007).

We analyzed RNA levels for three pairs of males and for two matched epididymis samples by Affymetrix microarray (Figures 25A). Curiously, low protein and caloric restriction samples consistently exhibited more “sperm-like” RNA populations (as opposed to epididymis RNA) than did control samples (Figure 25B). Whether this reflects systematic contamination issues or biological differences in sperm maturity or quality is presently unknown, although we note that we confirmed consistently-higher levels of the sperm-specific Dnahc3 by q-RT-PCR in an additional 7/8 low protein sperm samples (Figure 25E). We note that control sperm samples were routinely >99.5% sperm as assayed by microscopy, but nonetheless we cannot completely rule out systematic
contamination issues. With this possibility in mind, we identified genes were differentially-packaged in control vs. low protein sperm by correcting for potential...
epididymal contamination (Figures 25B-F). Interestingly, we observed downregulation of a number of transcription factors and chromatin regulators such as Smarcd3 and Ppard, although q-RT-PCR validation was not statistically significant due to high inter-animal variability (Figure 25F).

Although the downregulation of Smarcd3 was not significantly confirmed by q-RT-PCR, this could reflect the variable penetrance described above. Given that heterozygous mutants in chromatin remodelers can affect offspring phenotype even when the mutant allele segregates away (Chong et al., 2007), we used an initial genome-wide mapping (not shown) of overall histone retention (pan-H3 ChIP) abundance and the key epigenetic histone modification H3K27me3 in sperm to identify targets for single locus analysis. We observed a consistent decrease in H3K27me3 in low protein sperm at the promoter of Maoa (Monoamine oxidase) in 5/5 pairs of sperm samples, and a decrease in H3K27me3 at Eftud1 in 4/5 paired samples. These results demonstrate proof of principle that the sperm epigenome is regulated by dietary conditions, although the biological implications of these observations are not yet clear.
3.3 Methods

3.3.1 Experimental procedure for the epigenetic inheritance experiment in mice

The experimental work was led by Prof. Oliver Rando, and were carried out by various people all acknowledged in Carone et al., 2010. The extended experimental procedures can be found in the supplementary section of Carone et al.

Mouse husbandry and diets

All experiments were performed with mice which had been raised for at least two generations on control diet. Male mice were weaned from mothers at 21 days of age, and sibling males were put into cages with low protein or control diet (moistened with water to allow mice to break the hard pellets). Females were weaned to control diet. Males were raised on diet until 9-12 weeks of age, at which point they were placed with females for one or two days. At three weeks of age offspring were sacrificed, and median lobe of liver was rapidly dissected out and flash-frozen in liquid N2. Diets were obtained from Bio-serv, and sterilized per standard protocol.

RNA extraction and Microarray hybridization

Liver samples were ground with a liquid N2-cooled mortar and pestle. Total RNA for microarray analysis was extracted from liver powder using Trizol. For the microarray hybridization, 30 mg of total RNA was labeled for 2 hours at 42 C with Superscript II reverse transcriptase using 4 mg of random hexamer and 4 mg of oligo dT. Cy3 and Cy5-labeled samples were hybridized to home-printed “MEEBO” microarrays. Microarrays were hybridized at 65 C for 16 hours, washed as previously described (Diehn et al., 2002), and scanned using Axon Genepix 4000B microarray scanner.

Lipid measurements, Small RNA cloning and sequencing

For lipid measurements ~50-100 mg of ground liver tissue from six animals (three paired sets) was sent to Lipomics for “Truemass” mass spectrometry characterization of 450 lipid levels. Total RNA was isolated from ground liver tissue using mirVana
(Ambion). 18–35 nt small RNA was purified from 100 µg of total RNA, ligated to adaptors, amplified, gel-purified, and sequenced using a Solexa Genome Analyzer (Illumina) (Ghildiyal et al., 2008).

**Sperm isolation**

Caudal epididymis was dissected from sacrificed animals, punctured, and incubated for 30 minutes in M2 media at 37 C. Supernatant was removed, pelleted (10,000g for 10 minutes), and washed 2X with PBS, then 1X in water. Sperm preparations were only used that were >99.5% pure as assessed by microscopy, and q-RT-PCR was also used to reject any sperm samples with detectable epididymis-specific genes Actb or Myh11 compared to sperm-specific genes Smcp or Odf1.

**RRBS and MeDIP**

Reduced representation bisulfite sequencing (RRBS) was carried out as previously described (Meissner et al., 2008). Methyl-DNA immunoprecipitation (MeDIP) was carried out essentially as described (Weber et al., 2005; Weber et al., 2007). 4 mg of purified genomic DNA was fragmented to a mean size of 300bp using a Covaris machine, denatured, and immunoprecipitated with 5mC antibody (Eurogentec). ChIP material was Solexa sequenced, with ~21 million uniquely mappable reads per library.

### 3.3.2 Micro-Arrays data processing and differentially expressed genes in the liver

Array features were filtered using the autoflagging feature. The remaining features for each array were then block normalized by calculating the average net signal intensity for each channel in a given block, and then taking the product of this average and the net signal intensity for each filtered array feature in the block. Afterwards, all block-normalized array features were normalized using a global average net signal intensity as the normalization factor (Liu et al., 2005). Differential genes were determined based on a t-test, at FDR correction of 0.001 (Figure 20B). The second more stringent set of differential genes was determined using a
combination of two one-tailed t-test, testing for genes with mean fold change significantly higher than 0.2 or lower than -0.2, at FDR correction of 0.01 (Figure 20C). Enrichment of differential genes observed in our low protein offspring and of the two distinct cluster of up regulated genes (described above) with GO categories was computed using the Hyper-Geometric \( p \)-value \( \leq 0.05 \) after Bonferoni correction for multiple hypothesis testing.

3.3.3 MicroRNA identification from deep sequencing data and analysis

Deep sequencing reads where mapped to the mouse genome (July 2007 assembly of the mouse genome - mm9, NCBI Build 37, (Waterston et al., 2002)), where at least the first 18 bases of the read matches to the genome and the remainder matches to the 3′ adaptor sequence. In this mapping we allow up to 2 mismatched positions. Reads that had six or more matching locations in the genome were discarded. We retained reads with five or less matches to allow situations of micro-RNA families with highly similar sequence. We then cluster the mapped locations from all lanes to find “islands” of close location with a gap of less than 100 bases. Each island is a putative small RNA gene. We filter out islands that contain less than 10 mapped reads or are represented by less than 3 different unique read sequences. Determining the differentially expressed small RNAs (islands) was done using a paired t-test comparing pairs of lanes corresponding to age and sex-matched low protein and control offspring. We tested for islands with mean log₂ fold change significantly higher than the null hypothesis of 0.2 or lower than -0.2.

MicroRNAs were identified using the known locations of microRNAs in the genome. Novel microRNAs were predicted using a modified version of the MirDeep package (Friedlander et al., 2008). The first stage of the prediction in which we identifying the candidate precursors was based on the mapping to the genome and clustering to islands described above. The rest of the prediction stages where done as in (Friedlander et al., 2008) using the MirDeep package. Briefly, this procedure selects islands that the position and frequency of the sequenced reads are compatible with the
secondary structure of the miRNA precursor. We predict an island is a microRNA if it has a prediction score above 0. Prediction of miRNA targets were taken from the TargetScan (Friedman et al., 2009) database (Release 4.2), and were then checked for enrichment in differentially expressed genes using the Hyper-Geometric $p$-value.

3.3.4 Comparison to public murine liver microarray data

We built a compendium of public microarray data consisting of 120 gene-expression profiles in the murine liver under various conditions and genetic perturbations. To build this compendium we retrieved 113 datasets from GEO (http://www.ncbi.nlm.nih.gov/projects/geo/), manually chose the relevant labels for comparison, and created signatures of differentially expressed genes. The signatures were determined using a combination of two one-tailed t-test, with FDR correction of 0.1. In addition, we added further published sets of differentially-expressed genes (Horton et al., 2003; Yang et al., 2009) to the compendium. Expression-profiles significantly enriched with up or down regulated genes in our low protein offspring, were defined by a Hyper-Geometric $p$-value <= 0.05 after bonferoni correction for multiple hypotheses ($p<0.00025$). In this enrichment test we included only genes which were measured both in our expression arrays and in at least one other public murine liver profile. Our up regulated set of genes was clustered according to their overlap with the significantly enriched signatures, using a hierarchical clustering algorithm (with default parameters, (de Hoon et al., 2004; Eisen et al., 1998)).

While most of the overlaps described in Figure 21 are with gene expression measured upon perturbation of transcription factor activity, for HNF-4a not only do our differential genes significantly overlap those genes affected by a deletion of this factor ($p < 4 \times 10^{-6}$), but our differentially-expressed genes were also significantly enriched for genes previously shown to be directly bound by HNF-4a (Odom et al., 2007) ($p < 0.01$, data not shown).
3.3.5 Percent variance explained by *Ppara* RNA levels

We performed PVE (percent variance explained) analysis to determine how much of the overall gene expression profile is explained by *Ppara*. In this analysis we learn for each gene a linear prediction based on the *Ppara* expression values (standard linear-regression in Matlab), and then compare the gene's variance in expression across samples (Vtotal), to the variance after we remove the prediction (Vpred). The PVE is the percent of the total variance (without prediction) explained by the prediction, or in formula $100 \times (1 - V_{pred}/V_{total})$. Note that this does not determine whether *Ppara* is causal for the genes that it “explains”, as *Ppara* could be part of a regulon with the genes it correlates with.

3.3.6 Analysis of sperm RNA data

We first normalized the eight Affymetrix arrays using *rmasummary* method in Matlab (RMA normalization for the probe intensity and quantile normalization between arrays). We computed log-ratio values per gene in the six sperm RNA arrays, using matched pairs of low-protein diet compared to control diet arrays (2 pairs), and caloric-restriction diet compared to control diet (1 pair). To correct the sperm data for possible contamination by epididymis RNA, we used the two additional epididymal RNA data, and applied two different methods. In the first method we correct each sperm sample separately, by learning the maximal possible fraction of epididymal RNA within each sperm RNA sample (maximization is done by requiring all genes except 2% should have a non negative expression value after subtracting the epididymal fraction). The resulting fractions vary between 10%-30%. In the second method we correct each log-ratio value compared to the log-ratio of sperm RNA and Epididymis RNA (mean values across all sperm or epididymal samples), by using a standard lowess normalization (*malowess* function in Matlab).
Chapter 4 - Discussion

The evolution of transcription regulation is a major driving force generating the wide phenotypic diversity observed across species. Overall in this work I unraveled basic selection forces underlying this evolutionary process. First, we focused on evolution driven by random mutations in the DNA sequence, focusing on changes in DNA binding sites of transcription factors (cis), showing that: (1) The regulatory network of transcription factors and their target genes is highly plastic. (2) The transcription factors tend to conserve their functions. (3) A functional selection turnover model can reconcile these two trends, unlike traditional gene-centered models. Second, we focused on epigenetic inheritance mechanism, directly affected by previous parental environmental exposures. The evidence we present indicates that: (1) Paternal diet affects gene expression in the offspring of mice. (2) Epigenetic information carriers in sperm respond to environmental conditions. Each of these studies raises numerous mechanistic questions (discussed below).

4.1 Robustness in the face of plasticity – cis-regulatory evolution

To study regulatory evolution we applied to 23 Ascomycota species a novel approach for reconstructing regulatory networks based on cis-elements. Using this approach, we systematically built regulatory networks from 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, relevant to yeasts and mammals. In addition, we established a rich public resource (http://www.compbio.cs.huji.ac.il/OrthoMotifs/) that will facilitate studies of regulatory evolution of individual clades or species, including human and plant pathogens.

Our computational pipeline, CladeoScope, provides a major advance toward the computational reconstruction of cis-regulatory networks across many transcription factors
and species - a notoriously challenging problem (Weirauch and Hughes, 2010). CladeoScope relies on two assumptions. **First**, it assumes that transcription factor binding specificities, as reflected in DNA motifs, are largely conserved, even when their specific targets and functional roles have diverged. Thus, unlike previous approaches (Gasch et al., 2004; Kellis et al., 2003), CladeoScope does not assume any conservation of the known motif function or target genes. However, CladeoScope cannot track transcription factors whose binding specificity has significantly changed (as previously reported for some transcription factors (Baker et al., 2011; Kuo et al., 2010; Ravasi et al., 2010; Yvert et al., 2003)). This can be alleviated if more binding profiles are measured in non-model organisms. **Second**, we use conservation within a clade to focus on a reliable set of ancestral targets, improving the quality of the motif-target prediction for each clade. This minimizes many of the spurious interactions present in inferred *cis*-regulatory networks in a single species, while allowing us to trace regulatory divergence between clades, albeit at the cost of studying species-specific innovations.

While the conservation profiles of most DNA motifs match those of their cognate transcription factors, 27% of pairs show some discrepancy, more typically in the species most distant from *S. cerevisiae*. In some cases, a motif is not detectable despite the conservation of the related transcription factor (*e.g.* Zap1). This demonstrates the limitations of our approach in tracing regulatory evolution when the factor’s binding specificity has diverged substantially, or when target turnover rate within a clade is very high. These problems can be partially alleviated by using additional binding profiles from non-model organisms. In other cases, a motif is detectable but the related transcription factor is not identified as conserved, due to faulty target predictions, faulty orthology resolution (*e.g.* Sko1), or the presence of a family of related transcription factors with very similar binding specificities (*e.g.* the CBF1 family).

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 11a**) 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 (Bradley et al., 2010; Moses et al., 2006) 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 was 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 evolutionarily 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 13-15), we detect high turnover within the large majority of functional modules (Figure 17), 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 (Hogues et al., 2008; Tanay et al., 2005; Tsong et al., 2006; Tuch et al., 2008; Weirauch and Hughes, 2010; Wohlbach et al., 2009) (e.g., Figure 26). This would be consistent with the conserved co-expression of many functional modules in yeast (Hogues et al., 2008; Tanay et al., 2005) and mammals (Odom et al., 2007). More broadly, transcriptional regulation is only one of many regulatory layers, thus loss of the regulation of some genes might not have a functional effect, since other 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 determining 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.

Figure 26. Target turnover and compensatory regulation within a functional module
(a) Compensatory regulation by Leu3 and Gcn4 at leucine & isoleucine biosynthesis genes. Shown are selected target genes (rows) and their connection to Leu3 and Gcn4 motifs across the different clades (columns). (b) The position of the target genes in the Leucine-Isoleucine biosynthesis pathway.
4.2 Epigenetic inheritance - Transgenerational environmental reprogramming of gene expression

To test the existence of transgenerational environmental reprogramming we developed an experimental system in which we can directly test the effects of paternal exposures on their offspring, and on the germline. Our results demonstrate that paternal diet in mice reprograms gene-expression in the offspring and perturbs epigenetic information carriers in sperm. These results have numerous implications for human health.

We clearly identify a set of physiological pathways whose expression is sensitive to paternal diet. Specifically, we find that hepatic expression of genes involved in proliferation and cholesterol biosynthesis can be regulated by paternal diet, and these changes are reflected in levels of several lipid metabolites. Combined with data showing that offspring glucose levels are affected by paternal fasting in mice (Anderson et al., 2006), these results demonstrate that paternal diet has wide-ranging effects on the metabolism of offspring in rodents. Interestingly, a very recent study from Ng et al (Ng et al., 2010) reported that chronic exposure of male rats to high fat diet was associated with pancreatic beta cell dysfunction in female offspring. It will naturally be of great interest in the near future to compare the transgenerational effects of high fat and low protein diets, although one clear difference is that in our system a transgenerational effect is observed in both sex offspring. Whether the effects we observe on cholesterol metabolism prove advantageous in low protein conditions remains to be tested, but it will be important to investigate ecologically-relevant diets in order to speculate more firmly about adaptive significance of any observed transgenerational effects. For example, at present we cannot say with certainty what aspect of the low protein regimen is sensed by males – it is possible that offspring metabolism is affected by overall protein consumption, or high sucrose, or fat/protein ratio, or even levels of micronutrients, as our males consumed diets ad libitum and thus might have over consumed the low protein diet.
What is the mechanistic basis for the reprogrammed gene expression state? Genome-scale analyses of cytosine methylation in offspring livers identified several lipid-related genes that were differentially-methylated depending on paternal diet. Most notably, a putative enhancer for a major lipid regulator, PPARα, exhibited generally higher methylation in low protein offspring than in control offspring. Methylation at this locus was variable between animals, consistent with the partial penetrance of Ppara down-regulation in our data. The overall gene expression profile observed in low protein offspring significantly overlaps gene expression changes observed in ppara/- mice (Rakhshandehroo et al., 2007), leading to the hypothesis that epigenetic Ppara downregulation via enhancer methylation is an upstream event that affects an entire downstream regulon in reprogrammed animals. Note that while the hepatic downregulation of Ppara suggests a liver-autonomous epigenetic change, we cannot rule out that hepatic gene expression changes result from global physiological changes resulting from downregulation of Ppara in some other tissue. Interestingly, Ppara expression in liver is also regulated by maternal diet – offspring of female mice consuming a high fat diet exhibit altered hepatic Ppara expression, with increased expression at birth but decreased expression at weaning (Yamaguchi et al., 2010). Together with our data, these results suggest that Ppara is a key nexus that integrates ancestral dietary information to control offspring metabolism.

Paternal diet could potentially affect offspring phenotype via a number of different mechanisms. While we focus here on epigenetic inheritance systems, it is important to note that parental information can also be passed to offspring via social or cultural inheritance systems (Champagne and Meaney, 2001; Jablonka et al., 1995; Meaney et al., 2007; Weaver et al., 2004). While such maternally-provided social inheritance is unlikely in our paternal effect system – males were typically only in females’ cages for one day – it is known that in some animals females can judge mate quality and allocate resources accordingly (Pryke and Griffith, 2009), and that seminal fluid can influence female postcopulatory behavior in Drosophila (Fricke et al., 2008; Wolfner, 2002). These and other plausible transgenerational information carriers cannot be excluded at present – ongoing artificial insemination and in vitro fertilization
experiments will determine whether sperm carry the relevant metabolic information in our system.

We focused on the hypothesis that paternal dietary information does indeed reside in sperm epigenetic information carriers. First, a subset of cytosine methylation patterns in sperm are known to be heritable (Chong et al., 2007; Cropley et al., 2006; Rakyan et al., 2003; Waterland and Jirtle, 2003). Second, several reports suggest that RNA molecules packaged in sperm can affect offspring phenotype (Rassoulzadegan et al., 2006; Wagner et al., 2008). Third, chromatin structure has been proposed to carry epigenetic information, as sperm are largely devoid of histone proteins but retain them at a subset of developmentally-important loci (Arpanahi et al., 2009; Brykczynska et al., 2010; Chong et al., 2007; Hammoud et al., 2009). Finally, it is conceivable that additional or novel epigenetic regulators (such as prions) are packaged into sperm, or that sperm quality is affected by diet, or that genetic changes are directed by the environment (although it is important to emphasize that inbred mouse strains were used in this study).

Here, we report whole genome characterization of cytosine methylation patterns and RNA content in sperm obtained from mice maintained on control, low protein, and caloric restriction diets. Globally, cytosine methylation patterns are similar in all three conditions, indicating that the sperm epigenome is largely unaffected by these diets. Nonetheless, changes in relatively few loci can have profound effect in the developing animal, and our data do not rule out the possibility of inheritance through sperm cytosine methylation, especially given that MeDIP is unlikely to identify ~10-20% differences in methylation at a small number of cytosines. Importantly, the putative enhancer of Ppara was not differentially-methylated in sperm. It will therefore be of great interest in the future to determine when during development the differential methylation observed in liver is established, and to identify the upstream events leading to differential methylation (Blewitt et al., 2006).

Interestingly, we did identify effects of diet on RNA content and chromatin packaging of sperm. For example, sperm from control animals were consistently depleted
of the highly sperm-specific \textit{Dnahc3} gene relative to sperm from low protein animals. We cannot presently determine whether this represents reproducible differences in contamination, differences in sperm maturity, or something else. Finally, based on our observation that low protein sperm tended to be depleted of genes encoding a number of chromatin regulators, we have begun to search for dietary effects on sperm chromatin structure. Interestingly we found that the \textit{Maoa} promoter was consistently depleted of the key Polycomb-related chromatin mark H3K27me3, demonstrating as a proof of concept that chromatin packaging of the sperm genome is responsive to the environment, and motivating genome-wide investigation into dietary effects on sperm chromatin. Given the common behavioral changes observed in other transgenerational inheritance paradigms, the possibility that H3K27me3 at \textit{Maoa} affects offspring behavior will be of great future interest.

These results are likely to be relevant for human disease, because not only is \textit{maternal} starvation in humans correlated with obesity and diabetes in children (Lumey et al., 2007), but, remarkably, limited food in fathers and grandfathers has also been associated with changed risk of diabetes and cardiovascular disease in grandchildren (Kaati et al., 2002; Pembrey et al., 2006). Interestingly, in these studies paternal access to food and disease risk was not associated with disease risk in the next generation, but was only associated with F2 disease risk. However, it is important to note that the transgenerational effects of food availability for paternal grandfathers depend on the exact period during childhood of exposure to rich or poor diets (Pembrey et al., 2006), whereas our experimental protocol involved continuous low protein diet from weaning until mating. Thus, future studies are required to define when and how paternal exposure to a low protein diet affects epigenetic programming of offspring metabolism.

Together, these results suggest rethinking basic practices in epidemiological studies of complex diseases such as diabetes, heart disease, or alcoholism. We believe that future environmental exposure histories will need to include parental exposure histories as well as the exposure histories of the patient, to disentangle induced epigenetic effects from the currently-sought genetic and environmental factors underlying complex
diseases. Our observations provide an inbred mammalian model for transgenerational reprogramming of metabolic phenotype that will enable dissection of the exposure history necessary for reprogramming, genetic analysis of the machinery involved in reprogramming, and suggest a number of specific pathways likely to be the direct targets of epigenetic reprogramming.

4.3 An extended evolutionary theory

Taken together, our results shed light on two different selection forces driving evolution of transcription regulation, and emphasize the need for an extended evolutionary theory, integrating both genetic and non-genetic inheritance (Danchin et al., 2011; Jablonka et al., 1998; Jablonka and Raz, 2009; Shea et al., 2011).

To understand evolutionary processes inclusively, it is necessary to account for all forms of inheritance and selection pressures. Specifically, adaptation to the environment involves changes in the expression pattern of genes, which can be caused by both genetic and non-genetic inheritance, yet on different time scales. Genetic inheritance involves the slow process of natural selection and fixation of random mutations, and non-genetic inheritance involves the direct inheritance of acquired characteristics immediately effecting the variation in the population (Jablonka et al., 1998). Notably, the evolutionary implications of epigenetic inheritance mechanisms might be different for unicellular and multicellular organisms (Shea et al., 2011), and should be accounted for. In the integrated evolutionary theory (Danchin et al., 2011; Day and Bonduriansky, 2011), variation in both genetic and non-genetic factors drive evolution, with a strong interplay between them. In which, the DNA sequence encodes potential epigenetic carriers of information, such as non-coding RNAs or positions of CpG dinucleotide potentially subjected to DNA methylations. Moreover, the DNA sequence directs environmental reprogramming of these information carriers, for example, through cis-regulatory elements controlling the expression of non-coding RNAs in the germline. At the same time, these non-genetic
factors can potentially shape the selection landscape on the DNA sequence, affecting for example the rewiring of regulatory networks.

An example demonstrating such a causal relationship between non-genetic inheritance and the evolution of the DNA sequence is of nucleosome occupancy in human sperm. Recent evidence suggests that transmission of paternal nucleosomes and their modifications influences gene expression in the early embryo (Vavouri and Lehner, 2011). Vavouri and Lehner show that nucleosome retention in human sperm is related to base composition variation, indicating that chromatin organization in the male germline may be an important selective pressure on GC-content evolution in mammalian genomes. Taken more broadly, this example strengthens the hypothesis that a requirement to propagate paternal epigenetic information to the embryo may be an important selective pressure on sequence evolution in mammalian genomes. Another example for interaction between different inheritance mechanisms is of maternal care in rodents (Champagne, 2008). The level of maternal care affects the level of DNA methylation of genes in offspring’s brain, which is maintained throughout their life, and lowers the level of maternal care they provide (Champagne, 2008). The extended evolutionary theory, combining both genetic and non-genetic inheritance and the interplay between them, has implications for diverse areas, from the question of missing heritability in human complex-trait genetics (Maher, 2008) to the basis of major evolutionary transitions.
References


Weirauch, M. T., and Hughes, T. R. (2010). Conserved expression without conserved regulatory sequence: the more things change, the more they stay the same. Trends Genet 26, 66-74.
Appendices

1. Appendix Note 1 - Validations of the CladeoScope Method
   1.1. Evaluations with synthetic data
   1.2. Robustness
   1.3. Evaluation of the phylogenetic filtering of non functional motifs
   1.4. Comparing motif targets to bound target genes measured by ChIP

2. Appendix Note 2 - Characteristics of Motifs and Target Genes
   2.1. Distances between species-specific motifs
   2.2. Motif score distribution of lost targets vs. non-targets

3. Appendix Note 3 - Validations of the Functional Modules
   3.1. Robustness of the functional modules
   3.2. Comparison of functional modules to the literature

Appendix Note 1 – Validations of the CladeoScope Method

1.1 Evaluations with synthetic data

We evaluated CladeoScope on synthetic data, where the ground truth is known. We defined a set of true ancestral targets (‘original set’) and evolved them into extant targets with a given turnover rate. We then introduced different levels of noise to the “predicted” targets in the extant species (false positives and false negatives) and applied CladeoScope to this noisy input. Finally, we compared the reconstructed ancestral set to the original (true) set, to test CladeoScope’s sensitivity and specificity. We conducted these tests multiple times, varying the extent of noise, the size of the ancestral target set, the degree of target turnover and the topology of the species tree. For each set of parameters (a total of 720 different sets), our results are averaged over 100 independent simulations.

CladeoScope is highly robust to noisy target predictions and other variations in the input (the percent of false positives we tested ranges between 0%-200%, and the percent of
false negatives ranges between 0%-60%, Figure 5 in main text). For example, on average, when 30% true targets were removed in each species set, CladeoScope has 86% sensitivity (Figure 5b in main text), and when 80% false targets were added in each species, CladeoScope has 81% specificity (Figure 5b in main text). Moreover, the percent of noisy targets added hardly influences the sensitivity (Figure 5b in main text), and the percent of true targets removed hardly influences the specificity (Figure 5a in main text).

CladeoScope error rate increases proportionally to the increase in the rate of target turnover (Figure 5 in main text). While for slow turnover (probability for target gain and and loss of 0.00002 and 0.003, respectively) and medium turnover (gain, and loss 0.0002 and 0.03, respectively) the errors are almost identical, we get a decreased sensitivity and specificity when the turnover is very fast (gain and loss of 0.002 and 0.3, respectively). In this case, when 30% true targets are removed we reach an average of 53% sensitivity (Figure 5c in main text), which is equivalent to removing almost 60% true targets for the slowest turnover rate. The specificity is less sensitive to the turnover rate (Figure 5d in main text). For the fast turnover, for 80% noisy targets added in each species we get on average 69% specificity as opposed to 81% and 82%, for the medium and slow turnover rates, respectively.

As a more realistic simulation, we estimated the gain and loss frequencies (Methods in main text) per species in all species in clades A (Sensu-stricto) and F (Candidas), for three different motifs, Hsf1, Fkh2 and Mbp1. We then ran the simulations with these parameters, as described above. Here again the percent of noisy targets added hardly influenced the sensitivity, and the percent of true targets removed hardly influences the specificity. Overall we observe that CladeoScope is very robust. For example, for the slow turnover motif Hsf1, when 30% true targets were removed in each set of motif targets per species, CladeoScope has 87% sensitivity in clade A and 97% sensitivity in clade F, and when 80% false targets were added in each species, CladeoScope has more than 90% specificity in both clades. The Fkh1 motif, with a fast turnover, demonstrated a similar sensitivity. However, the specificity of this motif was not as good, with only 43% specificity in the clades when 80% false targets were added in each species (in addition to the estimated fast gain and loss rates).
Other parameters we tested had a smaller effect on the reconstruction error. The number of genes in the original ancestral set does not affect the sensitivity of the prediction (Figure 5e in main text), but the specificity increases when the ancestral set size is substantially smaller (Figure 5f in main text). The topology of the species tree, which is a parameter of the CladeoScope algorithm, also has a minor effect on the reconstruction error. We compared two topologies, the topology in the sensu-stricto clade (clade A), and a symmetrical topology as in the Candida clade (clade E). In the symmetrical topology, the sensitivity is slightly higher and the specificity is slightly lower (Figure 5g,f in main text), since according to the parsimony rule there are assignments that will be inferred as ancestral targets in this topology and not in the asymmetrical one.

Overall we see that our method is highly robust to noise in the target predictions. Thus, we consider the ancestral target sets per clade as a reliable prediction of motif targets that can be used in future analyses.

1.2 Robustness

Different thresholds for significance of motifs in a species

We tested the robustness of the CladeoScope method to different significance thresholds for a motif in a species. The significance is computed by the HyperGeometric p-value over the overlap between the motif targets in the species and the ancestral targets in the relevant clade. This is an iterative process, where for each clade the CladeoScope algorithm first reconstructs the ancestral sets for all genes in the clade, and then computes the significance of each species based on this set. After filtering out insignificant species, it reconstructs again the ancestral motif targets in the clade, and uses the new set to re-evaluate the species significance. This is repeated until the ancestral set does not change (Methods in the main text).

We ran the algorithm on nine different motifs across all clades, using seven different thresholds, ranging between 5e-2 – 1e-3. For all motifs, the choice of threshold had little effect (if at all) on the number of ancestral targets reconstructed per clade. In clades where
the motif is not statistically significant there was a larger variation in the number of targets, but it was found to be insignificant in all cases, indicating that the algorithm is robust to the threshold.

**P-values for significance of a motif in a clade (significance of ancestral target sets)**

To test the significance of an ancestral set of motif targets in a clade, we compute an empirical p-value by simulating target sets of respective sizes for each species in the clade, and reconstruct ancestral targets from these random sets. This process is repeated 1,000 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.

To test the robustness of our results to this p-value threshold, we examined the number of statistically significant ancestral motif target sets across all clades and motifs for different p-value thresholds (Figure 7 in main text), ranging from 0.05 to 0.0005. We show that changing the threshold has a small effect on the number of significant motifs per clade. There are clades where the threshold hardly makes a difference, as in clade A & B (Figure 6 in main text), and most clades have a mild effect only. The most significant effect is seen in clade K for the highest p-value threshold of 0.05 (Figure 6 in main text).

**1.3 Evaluation of the phylogenetic filtering of non-functional motifs (random targets & random motifs)**

The CladeoScope procedure controls for non-functional motifs by requiring the motif in each clade to have a set of ancestral motif targets. Since it might be possible to reconstruct ancestral targets from random sets of target genes in each species, we validate the statistical significance of each ancestral set by computing an empirical p-value over the number of ancestral targets when using randomized targets (see Section 2 in this note).

To further test that our phylogenetic filter controls for non-functional motifs, we tested the
algorithm 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 in the **Methods** in the main text). We then ensured that the random motif we constructed is not similar to any known motif. For each such random motif, we scanned for targets in each species, and ran the CladeoScope algorithm to reconstruct the ancestral sets. In most clades the random motifs are insignificant, thus showing that the phylogenetic filter is good for filtering functional motifs. The only exception is clade A, where all random motifs were found to be statistically significant. Most of those (7 out of 10) were significant in clade B as well, but excluded due to insufficient coverage of the clade since they all were conserved outside of clade A only in the *S. glabrata* species. This reflects the promoter sequence conservation between species in clade A and *S. glabrata*. Therefore, in our results we do not include motifs found to be conserved only in clade A (and in clade B when including only the *S. glabrata* species).

### 1.4 Comparing motif targets to bound target genes measured by ChIP

To evaluate whether CladeoScope filtering improves target prediction, we computed the Sensitivity (fraction of true positive predictions out of the experimentally determined targets) and Precision Rate (fraction of true positive predictions out of the total predicted motif targets), using ChIP-chip data in *S. cerevisiae* (MacIsaac et al., 2006) as a source of experimentally determined targets. We compare the ancestral targets in the immediate clade (A) to motif targets that are not ancestral. This comparison shows a consistent high precision among ancestral targets without affecting the Sensitivity for three different thresholds for motif targets detection in a species (**Figure 8a,b** in main text). We then tested the contribution of ancestral targets to ChIP-chip data measures in other Yeast species (Borneman et al., 2007; Tuch et al., 2008), and show that in most cases the use of ancestral targets improves the predictions (**Figure 8c,d** in main text).

We conducted this test for three different thresholds over motif targets detection. Overall, we see that the precision rate is optimized when using the high threshold for motif
targets detection (80% of the best score for the motif in the species). Nonetheless, to further test the effects of lowering the threshold we ran CladeScope with a lower threshold for motif targets (75% of the best score for the motif in the species). We then assigned functional modules to the resulting ancestral set, and show that our *Functional selection turnover model* as well as the functional classification are robust to this threshold, (see details in Appendix Note 3).

**Appendix Note 2 – Characteristics of Motifs and Target Genes**

2.1 **Distance between species-specific motifs**

We tested the contribution of the species-specific motif refinement process to the quality of CladeoScope. This refinement step generates a species-specific motif based on an input motif in the model organism *S. cerevisiae* (see Methods in main text). We first tested the distance between each refined motif to the original motif in *S. cerevisiae*, and their correlation with the distances between species. We see that the distances between motifs are small as expected (as measured by BLiC (Habib et al., 2008)). However, the distance is anti-correlated with the distance between species (branch length are computed based on amino acid substitutions, see Methods in main text): Correlation=-0.58, p-value=0.005, for the mean distance of all motifs in each species.

A measure for the contribution of the species-specific motif refinement process to CladeoScope output is their effect on motif targets. We compared a genomic scan for motif targets with the *S. cerevisiae* motif to a scan for targets with the species-specific motif, and observed that the refinement of motifs both adds additional targets and removes other targets (Figure 7 in main text). In most motifs there are more targets added than removed (Figure 7 in main text). We note that the target genes change even in species close to the sensu-stricto clade (Figure 7 in main text).
2.2 Motif score distribution of lost targets vs. non-targets

When predicting motif targets, genes with weak binding sites in their promoters are not considered as targets, however in some cases these sites might be functional. Thus to explore the detectability of such weak targets we wished to compare the scores of non-target genes to those of functional but weak targets. To do so we reasoned that likely candidates for weak and functional sites would be genes that were predicted as ancestral targets, but lost in the reference species (i.e. strong targets in sister species within the same clade). Thus, we compared the distribution of motif scores of such lost targets (lost in a species compared to the clade) to non-target genes. The results show that in 85% of cases tested, the lost targets have the same score distributions as the ancestral targets (p-value<0.001, Kolmogorov-Smirnov test), in the remaining cases the “lost targets” genes are an intermediate between “conserved targets” and “non-targets”. Since we show here that functional but weak binding sites have similar score distributions to non-functional sites, then lowering the threshold of significance would not improve the sensitivity of our results.

To rule out the possible concern that our procedure suffers from a detectability problem in remote clades that biases our results, we conducted a similar analysis, but this time comparing between neighboring clades. We reason that if we have a detectability problem, we might find in a specific clade the same targets as in a different clade only with weaker binding sites. Here again we divided all genes to sets of “missing targets” (not targets in a species and in its direct ancestor but targets in the neighboring clade) and “non targets” (not targets in both clades). In this case we see that the “missing” and “non” targets are identical to each other. Indicating that between neighbor clades we do not suffer from a detectability problem.
Appendix Note 3 – Validations of the Functional Modules

3.1 Robustness of the functional modules

To test the robustness of the functional modules, we applied the algorithm with different parameters and inputs, including:

Enrichment thresholds for functional modules with motif targets
When creating functional modules our procedure selects for the relevant functional annotations by selecting only categories that are enriched with motif targets. We tested the robustness of our method to the HyperGeometric p-value threshold ranging between: 1e-3 - 1e-6.

Threshold for merging gene-sets
The procedure for building functional modules merges gene-sets (in the context of a given motif across all clades) by comparing the set of motif targets in each gene set and checking that the fraction of motif targets in the intersection is above the threshold. We tested the robustness of our method to this threshold by comparing results with thresholds ranging between: 40% – 75%.

Thresholds for initial predictions of target genes
We tested the robustness of our method to different motif target detection thresholds. Testing two alternative thresholds (80% and 75% out of the best possible score for each motif in each species). The change in threshold results in different number of ancestral targets per clade (see Appendix Note 1).

For each set of parameters we tested several characteristics:

a) The number of modules
b) The fit of our Functional Selection Turnover model
c) The classification of motifs to functional classes (Functional conservation, Clade specific innovation or Functional switch) – we examined in detail 18 motifs including those
discussed explicitly in the main text.

d) Robustness of the functional annotations of motifs by the functional modules - we examined in detail 18 motifs including those discussed in the main text.

**Results of the robustness tests of the functional modules**

**a) Number of functional modules**

Most parameters tested do not affect significantly the number of functional modules. Specifically, the number does not change significantly when changing the motif detection threshold. For example, when comparing the lower threshold to the higher one, the number of modules changes from 197 modules to 155 modules (with enrichment threshold of 2e-5). When changing the enrichment and merge thresholds, the number of functional modules increases as the enrichment threshold becomes more permissive (**Appendix Figure 1**), and as the merge threshold increases (**Appendix Figure 2**). This is most noticeable for the most extreme thresholds, while for all other thresholds we see minor differences. In addition, the effect is mainly a result of adding small modules, consisting of one gene-set, which have less than 10 motif targets within them.

**Appendix Figure 1. Number of modules per enrichment threshold.** Shown are the numbers of functional modules for all motifs at different enrichment thresholds (-log p-value, Fisher’s exact test), plotted with different colors from yellow to blue. (a) The number of modules in three categories: all functional modules (>0), modules with more than one gene-set (>1), and modules with more than two gene-sets (>2). (b) The number of modules in three categories: all functional modules (>0), modules with at least 10 motif targets (>10), and modules with at least 15 motif targets (>15).
Appendix Figure 2. Number of modules per merging threshold. Shown are the number of functional modules for all motifs at different thresholds for merging modules (percent overlap in motif targets), plotted with different colors from yellow to blue representing the percent overlap. (a) The number of modules in three categories: all functional modules (>0), modules with more than one gene-set (>1), and modules with more than two gene-sets (>2). (b) The number of modules in three categories: all functional modules (>0), modules with at least 10 motif targets (>10), and modules with at least 15 motif targets (>15).

b) Fit of the functional selection turnover model
The fit of our model is good in all thresholds and parameters. More specifically, we find a fit to the model (p-value<0.01 after Bonferoni correction for multiple hypothesis) for at least 91% of the functional modules across all thresholds (ranging between 91% - 95% when changing the enrichment threshold, ranging between 94% - 96% when changing the merge threshold, and between 92% - 96% when changing the motif detection threshold).

c) The classification of motifs to functional classes
In this work we classified motif to three functional classes: Functional conservation, Clade specific innovation or Functional switch. We now repeated this process for different thresholds of motif enrichment (0.001 and 2e-6), merge threshold (45%,55%,65%) and motif targets detection threshold (80%, 75%), for 18 motifs. In most cases the classification of the motifs did not change as a result of changing the threshold. This implies that our conclusion regarding the distribution of the motifs to classes is robust to this parameter in the algorithm. The enrichment threshold changed the classification in two cases (Cat8 and Stb5 motifs). The merge threshold and motif detection threshold, each effected the classification of one motif (Swi6/Mbp1 motif and YNR063W respectively).
**d) Robustness of the functional annotations of motifs**

The functional annotations of motifs are largely robust to the choice of threshold, although many gene-sets are not found to be enriched when lowering the threshold. The robustness of the modules is due to compensation by overlapping gene-sets. Thus, by creating modules we overcome sporadic non-enrichment of specific gene sets. However, we do find that small modules (usually with a small number of motif targets and only one gene-set) enriched in a single clade are sensitive to the choice of threshold and tend to be excluded as the threshold is stricter.

**3.2 Comparison of functional modules to the literature**

The CladeoScope algorithm itself does not take functional annotation into consideration. Moreover, we do not require the motif targets in *S. cerevisiae* to be conserved in remote clades and species. Thus, we can use the known functional annotations as an indication for the functionality of the DNA motifs. For each motif we compared the functional annotations to the known function of the motif (when available) in *S. cerevisiae, C. albicans* and *S. pombe*. Overall we find a very good fit between the functional modules and the known function. From all motifs with known annotations we find 75% that have an exact match, 12% that have a partial match and 13% with no match.

**References**


של ליפידים של Ppara. העובדות שלנו היא האחת העובדות הראשות המספרות ראות חותכות
לרכ: (1) התוכנה האבחנה המְשְפּועַת על ביוטו גוֹנִים מִטָּבֻּוליים בֶּזֶּאצְאִים של עכברים (2) גורמים אפיגנטים בגרשים, הַיְכֵלוֹמִיָּה לְעָבָר מִדָּעַד בַּרְבּוּר, מַשְׁחַטְיָה בַּתְּנָבָגָה לָתְמוֹאֵת הסְבִיבָה. חָצְאוּ אֵלָה, בְּשִׁילָבָהּ עַל נַנוֹּיֵה מַמַּחֲקֶרֶים אפִידֶמִיוֹרְגִיִּים בֵּאָדָם, מְצָבִיעֵה עַל כּוֹרַהֲתָנֹהַהּ של ההַרְמוֹנָה יִכּוֹל לַזְהָבֶּיִּי על מֵטָבֻּוליים של קְולֶשְׁנָדָל וְשְׁמוֹנְיַיִם בְּזֶאצְאִים. כְּמוּ כּוֹרַהֲבָדוּ זֶה הָגְרְדוֹ מְדּוֹלָל לָמְעָרְכָהּ המַמַּחֲקֶרֶים לְחָכְרָיִים של הַרְרוֹנָה שֶל סְבִיבָה הָתוּרַה עַל די הָורְשָה של גורָמִים אפִיְגַנְטִיִּים. יִתְּרַע עַל כּוֹרַהֲנָתָהּ אַלּ מְצָבִיעֵה עַל הָזָרְוִר בְּחַשִּׁיבָהּ מַחֲזֶהָתָהּ בַּגֶּבֶר מַמַּחֲקֶרֶים אפִידֶמִיוֹרְגִיִּים שֶל מַחֲזֶהָתָהּ מְרוֹבֶּכֶבֶר בֵּאָדָם כְּנוֹן סְוֹכָרָה, מַחֲזֶהָתָהּ לָבָא, אִלָּאָכּוּלוֹיִיָּה.

לַסְיָם, הָחַזֶּהָתָה של שְׁתֵּי הָעֲבְדָוֹת הַמְּנוּבָדוֹת כָּאִי שְׁפִיקָת פָּרָו לע שֶׁיָּמֶנֶנֶגְוִי הָורָשָה שֶוָנֵי הָחוֹזֵהָתָוַיִּפְּלִיךְ הָפְעָלִיִּים בְּצֶאצְאִים של בֶּקֶרָה שָׁפַעְתוּ. עַבְדָוֹת אַלּ מְזָהְבָּתָה הַזָּרְוִר בֵּיתְאָרִיִּים בְּצֶאצְאִים מְרָוְחְבָּת, הָמְשָׁלַבְּתָּ מַמַּחֲקֶרֶים הָורָשָה גְּטִיטִיָּוַיִּיָּוַיִּי. 
The goal is to functionally maintain the regulatory activity of the clock. This function is achieved by a selection model according to a selection function that operates on the replacement of genetic targets. In this model, the regulatory tasks of the clock tend to be more strongly selected for compared to individual genetic targets. In our model, the selective pressures operate in different ways to maintain the general genetic targets belonging to the same biological process, but not to maintain specific genetic targets within the process.

The results of our research explain the number of genetic targets that are maintained in evolution (with a high number of species), and also explain the changes in the genetic targets measured for specific effectors that are studied for different numbers of species, both in mammals and in their environments.

We show that selective pressures are more stringent than those observed in the past. We show that selective pressures on networks allow many local changes to occur, which lead to adaptation of gene expression to changing environmental conditions, despite resistance to major changes in phenotype that are likely to be lethal.

Epigenetic inheritance may provide a mechanism for additional evolution in the regulation of the clock, as the inheritance of epigenetic factors can lead to changes in the expression profiles of genes in the offspring generation.

In our research, we performed a search for genes in mice that showed changes in expression in the offspring due to maternal diet changes. We focused on maternal diet because in mice, the fathers do not contribute significantly to the offspring, and thus we could rule out direct responses of the offspring to changes in their environment, particularly to the first environment they experience.

We found that the offspring of mothers who ate a diet rich in proteins, compared to mothers who were fed a normal diet, showed an increase in the expression of many genes involved in lipid and cholesterol metabolism, and also showed high levels of cholesterol, triglycerides, lipids and free fatty acids. In addition, a comprehensive mapping and thorough analysis of epigenetic factors in the liver of the offspring, as well as changes in the transcription of the "H3 methylase" gene throughout the genome in the liver of the offspring, led to the discovery of over 0.2% changes in nutritional epigenetic inheritance in the "H3" gene in many regions of the genome in the liver of the offspring.

Between these changes lies a region of distant regulation linked to a gene encoding a central regulator.
בקטרת שעתוק ממלא תפקיד מרכזי בפעילות של תאים חיים ובתגובתם לאותות פנימיים או חיצוניים. בקרה מורכבת תזווית מתווכת על ידי מנגנונים שונים ומסומנים במאות. הנמצאים באנטרקציה זה עם זה. מנגנונים אלה כוללים בקרת שעתוק "י" של חלבונים קושרי DNA המזהים רצפים ספציפיים (פקטורי שעתון), בקרה אפיגנטית של الكرموسום, תמצות של תאים ו/change של DNA. הנחה היא ששיווקי בקרת השעתון מה גורמים ביצירת הגיוון הפונטי הרחב הנצפה בין מינים. עם זאת, התפקיד והדינמיקה של התהליך לרוב אינן ידועות. הנחיה היא ששינויים בבקרת השעתון מופיעים מרכזי ביצירת הגיוון הפונטי הרחב הנצפה בין מינים. עם זאת, התפקיד והדינמיקה של התהליך לרוב אינן ידועות.

ב industria של בקרת שעתון דזר הרזרה גנטית

בקטר בとにかく גלגלים מתוך רחבת מבט שונים של הורשה גנטית ולא-גנטית

נזורות של חלקי הקורקוניל הקלים הבקרת lenders היא המבוקד, וול יעים. בкрат השעתון, שיניים אלה יוצרים להורשה עלי ידי שיני הורשה שוב שיניים ומפעלים במאות. המודדים יוצרים "רוכל" להשפעת שיניים, וול חלופי של חלובים (כנון פקטורי שעתון).

המשטח של הלוגו "רוכל" וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקמח לשיניים, וול בקまま innovation

בכר הבקרת שעתון במהלך המかる

נמצאות במ conexión של "י" של חלבונים קושרי DNA המזהים רצפים ספציפיים (פקטורי שעתון), בקרה אפיגנטית של الكرموسום, תמצות של תאים ו/change של DNA. הנחה היא ששיווקי בקרת השעתון מה גורמים ביצירת הגיוון הפונטי הרחב הנצפה בין מינים. עם זאת, התפקיד והדינמיקה של התהליך לרוב אינן ידועות. הנחיה היא ששינויים בבקרת השעתון מופיעים מרכזי ביצירת הגיוון הפונטי הרחב הנצפה בין מינים. עם זאת, התפקיד והדינמיקה של התהליך לרוב אינן ידועות.

אנו פיתחנו שיטת חיוויית לherited האבולוציון של בקרת שעתון על פי פמינו ריבים, והשמדתיי ב כ ידי תעבון עבור הביטוי של בקרת השעתון של זירר מתضرورة פקטורי שעתוןIGHL(CancellationTokentencial) והשמדתיי ב כידי תעבון עבור הביטוי של בקרת השעתון של זירר מתضرورة פקטורי שעתוןIGHL(CancellationTokentencial) והשמדתיי ב כידי תועבון עבור הביטוי של בקרת השעתון של זירר מתضرورة פקטורי שעתוןIGHL(CancellationTokentencial) והשמדתיי ב כידי תועבון עבור הביטוי של בקרת השעתון של זירר מתضرورة פקטורי שעתוןIGHL(CancellationTokentencial) והשמדתיי ב כידי תועבון עבור הביטוי של בקרת השעתון של זירר מתضرورة פקטורי שעתוןIGHL(CancellationTokentencial) והשמדתיי ב כידי תועבון עבור הביטוי של בקרת השעתון של זירר מת بصورة מאוזר (כלומר, פקטורי השעתון מופיעים והם

המעטים שלומ ברקץ מהיר. שיניים אלה חלקו על הבזר וגי טור שדישים והן שזינו הגי טור נהוג ובין

קציר

תקציר

בקרת שעתוק ממלא תפקיד מרכזי בפעילות של תאים חיים ובתגובתם לאותות פנימיים או חיצוניים. בקרה מורכבת תזווית מתוּכת על ידי מנגנונים שונים ומסומנים במאות. הנמצאים באנטרקציה זה עם זה. מנגנונים אלה כוללים בקרת שעתוק "י" של חלבונים קושרי DNA המזהים רצפים ספציפיים (פקטורי שעתון), בקרה אפיגנטית של الكرموسום, תמצות של תאים ו/change של DNA. הנחה היא ששיווקי בקרת השעתון מה גורמים ביצירת הגיוון הפונטי הרחב הנצפה בין מינים. עם זאת, התפקיד והדינמיקה של התהליך לרוב אינן ידועות. הנחיה היא ששינויים בבקרת השעתון מופיעים מרכזי ביצירת הגיוון הפונטי הרחב הנצפה בין מינים. עם זאת, התפקיד והדינמיקה של התהליך לרוב אינן ידועות.
עבודה זו עשתה בהדרככתו של פרופ' ניר פרידמן
פרופ' חנה מרגלית
פרופ' חנה מרגליית
מחקר חישובי השואות של בקרת שעון
במאלי"ט

חיבור לשם קבלת תואר דוקטור לפילוסופיה

מאית

נעמי חביב

הוגש לטencent האוניברסיטה העברית, ברושליימ
אוגוסט 2012