**Graphical Abstract**

Systematic dissection of cis-regulatory logic underlying promoter decoding of transcription factor dynamics

A model incorporating TF binding sites, nucleosome occupancy and TATA box location quantitatively explains promoter threshold and activation timescale

**Highlights**

- *cis*-regulatory logic underlying threshold and activation timescale is elucidated

- Via modulation of TF binding sites, the promoter class can be switched

- A simple model quantitatively accounts for dynamic promoter behavior

- The promoter amplitude threshold can be decoupled from the activation timescale

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**In Brief**

Cells control gene expression in part by regulating the dynamics of transcription-factor activity. Hansen and O’Shea investigate the *cis*-regulatory logic underlying promoter decoding of transcription-factor dynamics for 20 promoter variants and reveal that two properties characterizing the gene expression response—threshold and activation timescale—can be decoupled.

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cis Determinants of Promoter Threshold and Activation Timescale

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SUMMARY

Although the relationship between DNA cis-regulatory sequences and gene expression has been extensively studied at steady state, how cis-regulatory sequences affect the dynamics of gene induction is not known. The dynamics of gene induction can be described by the promoter activation timescale (AcTime) and amplitude threshold (AmpThr). Combining high-throughput microfluidics with quantitative time-lapse microscopy, we control the activation dynamics of the budding yeast transcription factor, Msn2, and reveal how cis-regulatory motifs in 20 promoter variants of the Msn2-target-gene SIP18 affect AcTime and AmpThr. By modulating Msn2 binding sites, we can decouple AmpThr from AcTime and switch the SIP18 promoter class from high AmpThr and slow AcTime to low AmpThr and either fast or slow AcTime. We present a model that quantitatively explains gene-induction dynamics on the basis of the Msn2-binding-site number, TATA box location, and promoter nucleosome organization. Overall, we elucidate the cis-regulatory logic underlying promoter decoding of TF dynamics.

INTRODUCTION

Regulation of gene expression is important for the ability of cells to maintain homeostasis and survive stress. The expression level of a gene depends on cis-regulatory motifs present in promoters that are interpreted by transcription factors (TFs), which control the rate of transcription (Levo and Segal, 2014; Rando and Winston, 2012). A major goal is to quantitatively understand and predict gene expression from knowledge of cis-regulatory DNA sequence and TF activity. Accordingly, how the promoter input-output relationship depends on the number and location of TF binding sites, nucleosome stability, and positioning in the promoter, the affinity of TF binding sites, and the presence of other cis-regulatory motifs have been extensively studied at steady state (Gertz et al., 2009; Lam et al., 2008; Mogno et al., 2013; Rajkumar et al., 2013; Raveh-Sadka et al., 2012; Sharon et al., 2012). Moreover, several studies have explored the relationship between promoter architecture and cell-to-cell variability in expression (noise) at steady state (Hornung et al., 2012; Sharon et al., 2014). However, a significant aspect of gene regulation occurs out of steady state: the kinetics of gene induction crucially determine how cells respond dynamically to signals and stresses, but how gene induction kinetics are influenced by regulatory DNA sequences is poorly understood. Along these lines, recent studies demonstrate that cells transmit gene expression information about external signals and stresses by regulating TF activation dynamics (Behar and Hoffmann, 2010; Castillo-Hair et al., 2015; Levine et al., 2013; Purvis and Lahav, 2013). Yet, despite this, the relationship between promoter cis elements and how the promoter responds to dynamical TF inputs has not been studied.

To investigate the relationship between the architecture of a promoter and how it decodes TF dynamics, we study the SIP18 promoter, which is activated by the budding yeast TF, Msn2. During stress exposure, Msn2 encodes information about stress identity in its nuclear translocation dynamics—for example, Msn2 exhibits brief nuclear pulses with dose-dependent frequency under glucose limitation but a sustained pulse with dose-dependent amplitude under oxidative stress (Hao et al., 2013; Hao and O’Shea, 2012; Petrenko et al., 2013). Msn2 target genes can differentially decode Msn2 dynamics such that stress-relevant target genes are predominantly expressed under the relevant stress (Hansen and O’Shea, 2013, 2015; Hao and O’Shea, 2012). However, at a mechanistic level, we do not currently understand how promoters decode TF dynamics differently.

Conceptually, we can characterize the gene expression response with two parameters: (1) the amplitude threshold (AmpThr), which quantifies how sensitive the promoter is to the nuclear TF concentration, and (2) the activation timescale (AcTime), which quantifies gene induction kinetics, i.e., how long it takes the TF to activate the gene (Hansen and O’Shea, 2013). Based on these two variables, four extreme promoter...
classes exist (high or low AmpThr with slow or fast AcTime, corresponding to the HS, HF, LS, and LF classes). To understand gene induction dynamics and why natural promoters decode the same dynamical TF signals differently, we must understand the mechanistic and cis-regulatory logic that determines AmpThr and AcTime.

Here, we combine high-throughput microfluidics and quantitative time-lapse microscopy to pharmacologically control Msn2 translocation dynamics and measure how 20 SIP18 promoter variants decode Msn2 dynamics in single cells. We find that just three variables—the number of Msn2 binding sites, nucleosome occupancy over Msn2 binding sites, and their distance to the TATA box—are sufficient to quantitatively account for AmpThr and AcTime. Furthermore, we find that AmpThr and AcTime can be decoupled. Although the wild-type SIP18 promoter (WT pSIP18) belongs to the HS class, by modulating the number and location of Msn2 binding sites, we can switch it to the LF or LS class. Additionally, we show that AcTime, but not AmpThr, determines the gene expression noise level. Finally, we propose a mechanistic model that plausibly explains promoter class from promoter cis elements and chromatin organization.

RESULTS

Models for Inferring Promoter AmpThr and AcTime

To systematically investigate how AmpThr and AcTime depend on promoter cis elements, we sought an efficient way of determining AmpThr and AcTime for a large number of promoter variants. Previously, to determine AmpThr and AcTime, we exposed an Msn2 target promoter of interest driving dual YFP/CFP reporter expression in diploid cells to a panel of 30 different Msn2 inputs, fit a mathematical model, and then calculated AmpThr and AcTime based on the fit (Supplemental information; Hansen and O’Shea, 2013). Given that our microfluidic platform enables us to perform five experiments in parallel (Hansen et al., 2015), we sought inference models to quantitatively estimate AmpThr and AcTime from just five experiments in haploid cells. To obtain a training data set for the models, we exposed seven wild-type Msn2 target promoters for which we already know AmpThr and AcTime (Hansen and O’Shea, 2013) to five dynamical Msn2 inputs (the five x-conditions in Figure 1A; see Figure S1 for full data set). To understand why we chose these five experiments, consider the behavior of SIP18, an oxidative stress response gene (Rodriguez-Porrata et al., 2012), and of HXXI, a glucose limitation response gene (Herrero et al., 1995). SIP18 has a high AmpThr and slow AcTime (HS class), whereas HXXI has a low AmpThr and fast AcTime (Hansen and O’Shea, 2013). Because the SIP18 promoter activates slowly, it filters out brief nuclear Msn2 pulses (x(FM4) and x(FM8) in Figure 1A) observed during glucose limitation (Hao and O’Shea, 2012), whereas HXXI strongly induces under these conditions. Similarly, having a high AmpThr allows SIP18 to filter out low-amplitude pulses (x(175 nM) in Figure 1A) and only induce during sustained Msn2 activation (x(3 μM) in Figure 1A) observed during oxidative stress (Hao and O’Shea, 2012). HXX1, on the other hand, has a low AmpThr and induces significantly during low-amplitude input (x(175 nM) in Figure 1A). Thus, YFP expression under x(175 nM), x(690 nM), and x(3 μM) allows us to infer AmpThr, whereas we can infer AcTime from YFP expression under x(690 nM), x(FM4), and x(FM8). Building on this intuition, we developed simple inference models to estimate AmpThr and AcTime with two and three fitted parameters, respectively (Figure 1B; Experimental Procedures). These simple models could account for >98% of the variance in AmpThr and AcTime (Figure 1B). However, a model with a sufficient number of fitted parameters can fit any data set. Overfitting is a particular concern because it reduces the predictive power of the model. To test whether the inference models were overfit, we use leave-one-out cross-validation (LOOCV): we leave out one promoter, fit the model to the remaining six, and use these fit parameters to predict AmpThr and AcTime for the promoter that was left out. We repeat this for all seven promoters. Even when we correct for overfitting using LOOCV, the inference models still account for >96% and >93% of the AmpThr and AcTime variances, respectively (Figure 1B). Thus, the inference models are not overfit and we can use them to calculate AmpThr and AcTime from just five experiments.

Systematic Dissection of How Msn2 STRE Location in the SIP18 Promoter Affects AmpThr, AcTime, and Strength

Msn2 binds the stress-response element (STRE) (5’-CCCT-3’) with sub-micromolar affinity and with limited flanking base preference (Siggers et al., 2014; Stewart-Ornstein et al., 2013). To systematically investigate how AmpThr and AcTime depend on Msn2 STRE number and location, we focus on the SIP18 promoter, which contains three STREs. The “null mutant” without the two STREs at –386 and –367 bp between the –2 and –3 promoter nucleosomes cannot activate expression (Figure 1C), and we therefore consider the STRE at –524 bp to be non-functional on its own. We next developed a combinatorial promoter DNA synthesis method (Figure S2A) and divided the SIP18 promoter into four regions: A, B, C, and D. We added 2, 3, or 4 STREs (A2-4, B2-4, C2-4, and D2-4 in Figure 1C) to each region, mimicking natural Msn2 target genes, which also contain STRE clusters: HXX1, for example, contains five clustered STREs (Figure S2C). We also made “scattered” mutants (S4; SA–SD). Whereas region A and C are in the accessible linker regions between nucleosomes, region B and D are located in sequences within strongly positioned nucleosomes (Figure 1C). We exposed strains containing each of these 16 promoter variants (chromosomally integrated at the SIP18 locus and driving sip18::YFP) to the five dynamical Msn2 inputs (Figure 1A) and measured YFP expression (Figure S1; raw single-cell time-trace data are available as Supplemental Source Data). We used the models (Figure 1B) to infer AmpThr and AcTime for each promoter variant (Figure 1D) from the YFP measurements. Whereas WT pSIP18 has a very high AmpThr and very slow AcTime, most promoter variants have lower AmpThr and faster AcTime (Figure 1D). We find that most binding–site changes cause incremental effects and the variants generally fall along the AmpThr/AcTime diagonal. Thus, AmpThr and AcTime appear to be coupled for these 16 mutants. We observe minor discrepancies: D3, for example, appears to be slightly faster than both D2 and D4 (Figure 1C). We attribute this to slight experimental or measurement error or to a minor effect on nucleosome organization.
Figure 1. Inferring AmpThr and AcTime for \( p\text{SIP18} \) Promoter Variants

(A) Inferring promoter class from five experiments. Top panels: Msn2-mCherry nuclear translocation dynamics in five different dynamic treatments of 1-NM-PP1. 1-NM-PP1 inhibits PKA\(^\text{ss} \) and causes Msn2 to translocate to the nucleus (Hansen et al., 2015). Bottom panels: corresponding \( \text{sip18} \)::YFP (red) and \( \text{hxk1} \)::YFP (blue) gene expression measured using time-lapse microscopy for each 1-NM-PP1 treatment are shown. From the YFP expression ratios, it is possible to estimate the promoter amplitude threshold (AmpThr) and activation timescale (AcTime).

(B) Models for inferring AmpThr and AcTime. For the seven wild-type promoters for which we know AmpThr and AcTime from measurements in diploid cells (Hansen and O'Shea, 2013), we performed the five experiments shown in (A) in haploid cells and fit to models defined in Experimental Procedures. These models account for most of the variance in AmpThr and AcTime even when corrected for overfitting using leave-one-out cross-validation (LOOCV).

(C) Overview of \( p\text{SIP18} \) promoter architecture. Nucleosome occupancy (gray) and nucleosome centers (dyads; black) are plotted using MNase-seq data from Hansen and O'Shea (2013). The wild-type promoter contains three Msn2 binding sites (STREs: 5'-CCCCT-3') and a TATA box (Baihichar et al., 2004). The promoter was divided into four regions: A; B; C; and D. Promoter variants containing two, three, or four STREs in each region or “scattered” among the regions were constructed (Figure S2A). For each mutant, the experiments shown in (A) were performed and the promoter strength (response to 50 min; 3 \( \mu \text{M} \) 1-NM-PP1) relative to the wild-type \( p\text{SIP18} \) promoter is shown in blue bars.
Although previous studies have shown that changing the location of the TF binding site does not measurably affect promoter nucleosome positioning (Lam et al., 2008) and that neither does replacing the SIP18 ORF with a YFP reporter (Hansen and O'Shea, 2013), we cannot fully exclude that minor differences in nucleosome positioning may account for some of the differences observed. Mutant A4 shows the biggest change: A4 entirely switches to the LF class and has a very low AmpThr and a fast AcTime (Figure 2D). Furthermore, A4 shows ~5-fold higher strength than WT pSIP18 (defined as absolute YFP level under x(3 μM); Figure 1C). Thus, both AmpThr and AcTime are tunable in cis.

Three Variables—STRE Number, STRE Distance from TATA Box, and Nucleosome Occupancy of STREs—Suffice to Quantitatively Account for AmpThr, AcTime, and Strength

Next, we sought to mechanistically understand how AmpThr, AcTime, and promoter strength are determined. In the simplest scenario, the number of STREs could simply determine these variables. However, although AmpThr (Figure 2A), AcTime (Figure 2B), and promoter strength (Figure 2C) generally show a monotonic relationship with the number of STREs, the number of STREs alone cannot fully explain this relationship. For example, among the mutants with four STREs, B4, C4, D4, and S4 have similar AcTime, but A4 stands out with a much-lower AcTime (Figure 2A).

There could be several reasons why STRE number alone fails to account for promoter behavior. For example, nucleosomes restrict TF binding and nucleosome remodeling may be required for initiation of transcription (Lam et al., 2008; Rando and Winston, 2012; Raveh-Sadka et al., 2012; Weiner et al., 2012). Furthermore, for some promoters, removal of the nucleosome occupying the TATA box can be sufficient to activate transcription (Adkins and Tyler, 2006; Zhang and Reese, 2007). We therefore hypothesized that, together, the number of STREs (STREs), the level of nucleosome occupancy over the STREs (nuc; Figure S2B), and the average distance from the STREs to the TATA box (TATA)—can account for more than 90% of the variance in AmpThr, AcTime, and strength. Full details on the models and variables are given in Supplemental Information. See also Figure S2B.

(D) AmpThr versus AcTime for pSIP18 mutants. For each mutant in (C), YFP expression was measured for each Msn2 input in (A) and the models in (B) applied to estimate the AmpThr and AcTime for each mutant. Mutants have been colored based on the promoter classification, and intermediate promoters are shown in black.

See also Figures S1 and S2A. Raw time-trace data for all strains in Figure 1 (66,088 single cells) are available as Supplemental Source Data.
Decoupling Promoter AmpThr from AcTime: Mutant D6 Is an LS Promoter

(A–C) Decoupling AmpThr from AcTime in region D. AmpThr (A), AcTime (B), and strength (C) are plotted as a function of the number of STREs in region D. Mutant D6 appears to belong to the LS class.

(D) Systematic experimental dissection of how WT pSIP18, D6, and A4 decode Msn2 dynamics. Each row corresponds to a specific Msn2-mCherry input (left, in red), and the corresponding gene expression response for each of the three promoters is shown in the corresponding rows on the right. The gene expression responses for each promoter are internally normalized to their maximal expression level. Each row is the per-cell average of a few hundred cells from at least two replicates. WT data are from Hansen and O’Shea (2013). Data (15,875 single cells in total) for all 30 experiments for A4 and D6 are shown in Figure S3.

(E) Clustering of promoters. The full 30-experiment data set for A4 and D6 was fit to a previously described differential equations model (Hansen and O’Shea, 2013) and best-fit parameters inferred. Numerically, AmpThr is defined as the nuclear Msn2-mCherry level (AU) required to reach the half-maximal level of promoter activity obtained at 3 μM 1-NM-PP1 (the maximal nuclear Msn2-mCherry level). AcTime is defined as the time (min) it takes to reach the half-maximal level of promoter activity reached at steady state at 690 nM 1-NM-PP1. Both AmpThr and AcTime were obtained from model simulations. Full details are given in Supplemental Information.

(F and G) Total noise (σ²/μ²; F) and intrinsic (G) noise are plotted against the Msn2 AUC (μ) (Figures 1D and 3G), and the experiments are colored based on promoter class: WT (HS, red); D6 (LS, orange); and A4 (LF, blue). Each dot corresponds to the noise (mean across time points after gene expression has reached a plateau) for each of the 30 experiments performed in (D).

See also Figure S3 and Supplemental Source Data for all raw A4 and D6 data.

Mutants A4 and D6 Switch Promoter Class

Based on the five haploid experiments and the inference model estimate of AmpThr and AcTime (Figures 1D, 3A, and 3B), mutants A4 and D6 switch promoter class to LF and LS, respectively. To verify our inference approach and confirm these results, we made diploid strains with dual sip18::CFP and sip18::YFP reporters on the homologous chromosomes driven by the A4 and D6 promoters. Having dual CFP/YFP reporters allows us to also study gene-expression noise (Elowitz et al., 2002). We then exposed each diploid mutant to 30 experiments systematically varying Msn2 pulse duration, amplitude, pulse number, and interval (Figures 3D and S3), fit a previously described model (Hansen and O’Shea, 2013), and calculated the actual AmpThr and AcTime (Supplemental Information). Indeed, these results confirm that A4 is an LF promoter and D6 an LS promoter (Figure 3E). Having a slow AcTime, both WT pSIP18 and D6 filter out oscillatory and short-duration Msn2 input, whereas A4 with a fast AcTime responds strongly (Figure 3D). Similarly, both A4 and D6 have a low AmpThr and
therefore activate strongly to low levels of Msn2 activation, whereas WT pSIP18 largely filters out low-amplitude input (Figure 3D). Taken together, these results confirm that A4 and D6 completely switch the promoter class.

**Gene-Expression Noise Level Is Affected by AcTime Not AmpThr**

The relationship between gene-expression noise level ($\sigma^2 / \mu^2$) and wild-type promoter variants (Bar-Even et al., 2006; Newman et al., 2006) or synthetic promoter variants (Hornung et al., 2012; Sharon et al., 2014) has been extensively studied at steady state, but it is not clear how noise depends on AmpThr and AcTime. Previously, we observed a high negative correlation between noise and AcTime. Based on this, we argued that noise should strongly depend on AcTime—such that slow promoters show dramatically higher noise in gene expression—but that AmpThr should not strongly affect noise (Hansen and O’Shea, 2013). To causally test this, it is necessary to compare noise levels for promoters where only either AmpThr or AcTime are changed. Comparing WT pSIP18 and D6, AmpThr changes from low to high without much of a change in AcTime (Figure 3E). Likewise, comparing D6 and A4, AcTime changes from slow to fast without much of a change in AmpThr. For each experiment in Figures 3D and S3, we calculate the total (Figure 3F) and intrinsic (Figure 3G) noise (Elovitz et al., 2002) and plot the noise as a function of the Msn2 area under the curve (Msn2 AUC) (time-integrated Msn2 activation) such that each dot in Figures 3F and 3G corresponds to a single Msn2 input for a single promoter. We find that, whereas WT pSIP18 and mutant D6 exhibit high total and intrinsic noise, mutant A4 shows lower noise (Figures 3F and 3G). Because mutants A4 and D6 differ only by AcTime, this experimentally demonstrates that the noise level is strongly affected by AcTime, but not much affected by AmpThr.

**DISCUSSION**

Recently, it has become clear that cells transmit information and control cell fate by regulating the dynamics of master TFs (Levine et al., 2013; Purvis and Lahav, 2013). For example, in murine neural progenitor cells, control of TF Asc1 dynamics is sufficient to control cell fate: oscillatory Asc1 activity leads to cell proliferation, whereas sustained Asc1 activity causes neuronal differentiation (Imayoshi et al., 2013). The mechanism is believed to be due to different Asc1 dynamics inducing different downstream gene-expression programs, which requires that target genes show different induction kinetics. The same dynamical signaling logic appears to hold for the budding yeast TF, Msn2 (Hao and O’Shea, 2012). However, how promoter cis elements affect promoter decoding of TF dynamics was not understood.

Here, we systematically investigate how STRE number and location in the SIP18 promoter affects gene-induction dynamics. We find that AmpThr and AcTime can be decoupled and that just three variables—number of STREs, their accessibility, and their distance from the TATA box—suffice to quantitatively explain more than 90% of the variance in AmpThr and AcTime. Whereas the strong dependence of AmpThr on the number of STREs and nucleosome occupancy over the STREs could perhaps be expected from previous steady-state studies (Sharon et al., 2012), the strong dependence of AcTime on the distance from the STREs to the TATA box is surprising.

What is the cause of this relationship? And, mechanistically, how can we explain why A4 and D6 fall into the LF and LS classes? If we assume that (1) remodeling of the nucleosome occupying the TATA box is required for transcription, which is well supported (Adkins and Tyler, 2006; Zhang and Reese, 2007) and (2) remodeling of nucleosomes adjacent to Msn2 binding is fast, but remodeling of nucleosomes distal to Msn2 binding is slow, which is consistent with local recruitment of chromatin remodelers by TFs (Larschan and Winston, 2001; Weake and Workman, 2010), a mechanistic model emerges that explains promoter class from promoter architecture (Figure 4). Wild-type SIP18 promoter (HS class) is slow because, although remodeling of the −2 and −3 nucleosomes adjacent to where Msn2 binds is fast, remodeling of the distal −1 nucleosome occupying the TATA box is slow and remodeling of the −1 nucleosome is required for activation (Figure 4A). Similarly, the additional STREs in mutant D6 (LS class) greatly lowers the AmpThr, but D6 remains slow because Msn2 binds too far upstream of the −1 nucleosome to rapidly remodel it (Figure 4B). Conversely, mutant A4 has both a low AmpThr due to its clustered STREs in a nucleosome-free region and a fast AcTime because Msn2 binds adjacent to the TATA box (Figure 4C). Thus, remodeling the −1 nucleosome is rapid for A4, which therefore belongs to the LF class.

Although this mechanistic model (Figure 4) plausibly explains promoter class and induction dynamics from promoter architecture, it is a simplification of the complex sequence of events taking place during gene activation (Hager et al., 2009). Nevertheless, a direct prediction of the model is that the −2 and −3 nucleosomes should be remodeled faster than the −1 nucleosome occupying the TATA box for the wild-type SIP18 promoter, and this is supported by nucleosome remodeling time course data (Figure S4).

Recent advances in DNA synthesis now make it possible through massively parallel approaches such as FACS-seq to study thousands of promoters (Noderer et al., 2014; Sharon et al., 2012). Through such approaches, it will be interesting to investigate the extent to which the cis-regulatory logic underlying promoter decoding of TF dynamics we uncovered for the SIP18 promoter generalizes to other Msn2-regulated promoters and to promoters regulated by other TFs.

**EXPERIMENTAL PROCEDURES**

**Strains**

All Saccharomyces cerevisiae strains used in this work are in the W303 background. The combinatorial promoter synthesis method is illustrated in Figure S2A, a list of strains is given in Table S1, and further information about how they were constructed is given in Supplemental Information.

**Microfluidics, Time-Lapse Microscopy, and Image Analysis**

Briefly, yeast cells were grown overnight at 30°C with shaking at 180 rpm to an OD$_{600}$ nm of ~0.1 in low fluorescence medium (which exhibits minimal autofluorescence), quickly collected by suction filtration, loaded into the five channels of a microfluidic device pretreated with concanavalin A to
Figure 4. A Mechanistic Model Can Explain Promoter Class from cis Elements

A simplified, mechanistic model that assumes (1) remodeling of the nucleosome occupying the TATA box is required for gene induction and (2) that nucleosome remodeling adjacent to where Msn2 binds is fast but nucleosome remodeling distal to Msn2 binding is slow can explain observed differences in promoter class. Promoter architecture and nucleosome sizes are drawn to scale. (A) Model for wild-type promoter (HS). Msn2 binds to non-clustered STREs with low affinity and rapidly remodels adjacent nucleosomes. After slow remodeling of distal nucleosomes, the TATA box is available and initiation of transcription occurs. (B) Model for D6 promoter (LS). Msn2 binds to clustered STREs with high affinity and rapidly remodels adjacent nucleosomes. After slow remodeling of distal nucleosomes, the TATA box is available and initiation of transcription occurs. (C) Model for A4 promoter (LF). Msn2 binds to clustered STREs with high affinity and rapidly remodels adjacent nucleosomes. Because the TATA box is now available, initiation of transcription immediately occurs. See also Figure S4 for MNase-seq time course data.

Inference Models for Inferring AmpThr and AcTime

The AmpThr and AcTime inference models are given by

\[
\text{AmpThr}(\text{STREs}, \text{nuc}) = c_1 \cdot e^{\frac{f(\text{STREs} - 1) \cdot e^{-\frac{690}{3}} \cdot \text{nuc}}{1}}
\]

\[
\text{AcTime}(\text{STREs}, \text{nuc}) = c_1 \cdot e^{\frac{f(\text{STREs} - 1) \cdot e^{-\frac{690}{3}} \cdot \text{TATA} \cdot e^{-\frac{100}{3}} \cdot \text{nuc}}{1}}
\]

\[
\text{Strength}(\text{STREs}, \text{TATA}, \text{nuc}) = c_1 \cdot \left( \frac{(\text{STREs} - 1)}{\text{TATA} \cdot \text{nuc}} \right)^{c_2}
\]

Further details on how the variables are quantitatively defined, values for best-fit parameters, and the models are given in Supplemental Information.

SUPPLEMENTAL INFORMATION

Supplemental information includes Supplemental Experimental Procedures, four figures, one table, and supplemental source data and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2015.07.035.

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