Synthetic gene expression perturbation systems with rapid, tunable, single-gene specificity in yeast

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ABSTRACT

A general method for the dynamic control of single gene expression in eukaryotes, with no off-target effects, is a long-sought tool for molecular and systems biologists. We engineered two artificial transcription factors (ATFs) that contain Cvs₂His₂ zinc-finger DNA-binding domains of either the mouse transcription factor Zif268 (9 bp of specificity) or a rationally designed array of four zinc fingers (12bp of specificity). These domains were expressed as fusions to the human estrogen receptor and VP16 activation domain. The ATFs can rapidly induce a single gene driven by a synthetic promoter in response to introduction of an otherwise inert hormone with no detectable off-target effects. In the absence of inducer, the synthetic promoter is inactive and the regulated gene product is not detected. Following addition of inducer, transcripts are induced >50-fold within 15 min. We present a quantitative characterization of these ATFs and provide constructs for making their implementation straightforward. These new tools allow for the elucidation of regulatory network elements dynamically, which we demonstrate with a major metabolic regulator, Gcn4p.

INTRODUCTION

A current goal of synthetic biology is the development of modular tools for programming genetic circuits. This includes the ability to control gene expression in a fast, quantitative fashion without otherwise affecting cellular physiology. Rapid perturbation of a protein's abundance followed by monitoring of the genome-wide

transcriptional response has previously been used to dissect the architecture of complex regulatory networks *in vivo* (1,2). This kinetic information can directly inform quantitative models of gene regulation and elucidate novel design principles.

In yeast, the most commonly used expression systems rely on nutritional perturbations such as the addition of copper, sugar or methionine to modulate levels of a target gene's expression (3,4). These nutritional perturbations can be quite severe. For example, a gene whose expression is driven by the *MET25* promoter can be selectively repressed, but this requires the addition of 1 mM extracellular methionine (5). For overexpression of a target gene, the transcription factor (TF) Gal4p is commonly used. A target gene placed under the control of a promoter containing Gal4p recognition sequences (UAS_{GAL}) can be selectively induced in the presence of galactose (5). Since Gal4p activity is attenuated in glucose, this approach requires growing cells on alternative carbon sources like raffinose or glycerol.

Improvements to expression systems using otherwise inert inducers such as the tetracycline analog doxycycline or the mammalian hormone β -estradiol have reduced the perturbations of the physiological cell state due to introduction of the inducer (6–8). Although these previous systems have provided useful tools, demonstration of an expression system that satisfies the criteria of being (i) fast-acting, (ii) tightly regulated, (iii) nearly gratuitous (i.e. relatively few off-target effects) and (iv) graded over a range of inducer concentrations was only recently published for yeast (7).

Here, we provide a full characterization of a next-generation β -estradiol inducible expression system for yeast that displays true single-gene precision today as well as the potential for multiplex regulation in the future. This system utilizes a constitutively expressed artificial transcription factor (ATF) from the *ACT1* promoter.

The authors wish it to be known that, in their opinion, the first two authors should be regarded as joint First Authors.

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The ATF consists of a DNA-binding domain (DBD), the human estrogen receptor (ER) and the VP16 activation domain (Figure 1). In the absence of inducer, the ER interacts with the Hsp90 chaperone complex, sequestering the ATF to the cytoplasm (9). Introduction of β -estradiol displaces Hsp90, revealing a nuclear localization signal, and the ATF translocates to the nucleus (Figure 1). The ATF as described provides a strong transcriptional activator that is dependent on the presence of β -estradiol. By using synthetic DBDs that bind only a cognate DNA sequence in place of the Gal4p DBD, residual off-target effects have been completely eliminated.

Previously, the DBD from the yeast transcriptional activator Gal4p was used in the chimeric activator Gal4dbd.ER.VP16 (GEV) (7,10). Yet, as a DBD from a yeast TF, and a factor with only 6 bp of specificity, this DBD has many potential off-target sites in the yeast genome. DBDs of bacterial TFs (e.g. LacI and tetR) have been repurposed for use in the development of gene expression systems that are orthogonal to the native eukaryotic regulatory machinery. However, as discussed elsewhere (11), bacterial DBDs have numerous limitations (oligomerization issues, cooperative binding, etc.) that restrict their use for engineering novel TFs for use in eukaryotes.

We constructed a set of ATFs that utilize the modular and designable Cys₂His₂ zinc-finger DBD. A single zinc-finger domain contains ~30 amino acids and will offer three to four bases of specificity, while multiple domains can be assembled as zinc-finger arrays where each additional finger adds three bases of specificity [reviewed in (12)]. This domain has previously shown to be engineer-able to take on novel specificity and function in a somewhat modular fashion while high-quality zinc-finger arrays have been selected by multiple methods (13–15). Through decades of research some basic rules of specificity have been gleaned and specificity can sometimes be designed (16,17). ATFs have previously been constructed by fusing a designed Cys₂His₂ zinc-finger DBD to a hormone-responsive domain and transcriptional activation domain and were functional in mammalian cells (18,19). A repressive ATF utilizing an engineered Cys₂His₂ zinc-finger DBD has been reported to achieve single-target precision (also in mammalian cells) (20).

In this study, we replaced the Gal4dbd of GEV with either the three-fingered DBD of the mouse TF Zif268 or a rationally designed four-fingered zinc-finger array (Z_4). We refer to these constructs as Z_3 EV and Z_4 EV, respectively. The design of the Z_4 array leaned heavily on previously defined design principles and targets that have been shown to be most effective (i.e. GNN fingers). The Z_4

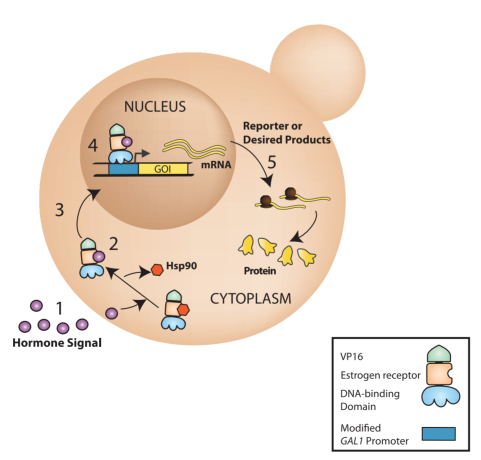


Figure 1. Schematic of hormone-based gene expression system. ATFs contain a DNA-binding zinc-finger array, the ligand binding domain of the human estrogen receptor and the VP16 activation domain. In the presence of β-estradiol (1), ATFs dissociate from Hsp90 (2), translocate to the nucleus (3) and activate transcription of a gene of interest (GOI) (4). Once produced (5) the gene products can be detected using a variety of methods.

array was designed to target not only a sequence that is not found in the yeast genome but also a sequence that accommodated a likely to be successful assembly of GNN-binding zinc fingers.

We demonstrate that the ATFs Z_3EV and Z_4EV are fast-acting and achieve a graded output response over a range of inducer concentrations, as did the GEV system. Using gene expression microarrays, we find that while GEV activation results in the induction or repression of several hundred genes within 30 min, these new ATFs have no significant off-target expression response. As a result, these ATFs eliminate the growth defect incurred by strong induction of GEV. Finally, we demonstrate the power of these new expression systems by inducing GCN4 and following the genome-wide expression responses dynamically.

MATERIALS AND METHODS

Media and growth conditions

For batch culture experiments, cells were grown in either synthetic complete medium lacking uracil or YPD (1% yeast extract, 2% bacto-peptone and 2% dextrose), unless noted otherwise. Induction of GEV, Z_3EV or Z_4EV by β -estradiol (Tocris Biosciences, Ellisville, MO, USA) was performed in cells during log-phase growth (culture absorbance = 50–100 Klett units). Chemostat cultures were maintained in phosphate-limited growth medium (20 mg/l potassium phosphate). When needed to complement an auxotrophy, phosphate-limited media was supplemented with excess L-Leucine (200 mg/l). The components of chemostat medium are carefully prepared to mimic the composition of standard yeast nitrogen base medium and have been detailed previously (21).

Strain and plasmid construction

Transformations were performed with a standard lithium acetate method. To construct Z_3EV and Z_4EV strains, the Gal4dbd of GEV was deleted by PCR-mediated disruption with URA3. Zif268 and Z_4 DNA-binding domains with homology to the ER and ACTI promoter were then transformed into cells and selected via 5-Fluoroorotic Acid (5-FOA) counter selection of URA3. Single colonies were isolated and sequenced to verify the presence of the proper DNA-binding domain (Supplementary Figure S1A).

The reporter plasmid was created from the base plasmid pRS416 (gift from Megan McClean), a CEN plasmid containing the *URA3* selectable marker. The *GAL1* promoter region was amplified from genomic DNA. Overlap-extension PCR was used to add the restriction enzyme sites for XbaI and NotI, respectively, on either side of the region of the three continuous Gal4p-binding sites 5'-CGG-N₁₁-CCG-3' (22). This promoter fragment was then cloned into pRS416 in front of green fluorescent protein (GFP) using the restriction enzymes NheI and XmaI. Before zinc-finger-binding sites were added the XbaI site in GFP was re-coded using a silent mutation to remove this restriction site. Finally, the three canonical Gal4p-binding sites were removed via digestion with XbaI and NotI and triplets of dimeric Z₃EV and Z₄EV-binding

sites were cloned into respective plasmids (Supplementary Data and Supplementary Figure S1B).

RNA extraction, labeling and hybridization

RNA extraction, labeling and hybridization were performed as described previously (7). Five milliliters of cells were harvested from chemostat cultures by vacuum filtration onto 0.45-um nylon membranes (Millipore, HNWP02500) and flash frozen in liquid nitrogen. Crude RNA was extracted with a standard acid-phenol method and subsequently cleaned with RNeasy (QIAGEN, Valencia, CA, USA). Cleaned RNA was labeled using the Agilent Quick-Amp Labeling Kit (#5190-0447). Reference RNA was extracted from a laboratory wild-type strain (DBY12001) grown to steady state in phosphate-limited growth medium with a doubling time of 3.9 h. Microarrays were hybridized for 17 h at 65°C on a rotisserie at 20 rpm. Hybridized microarrays were washed, scanned and raw data were extracted with Agilent Feature Extraction Software version 9.5.

Microarray analysis

Sample and reference channel intensities were first floored to a value of 350 (23,24). Once \log_2 ratios were computed between samples and reference, the data were time-zero transformed. Genes that were flagged by Agilent Feature Extraction Software or equal to 0 at every time point were removed in R (25). Data were hierarchically clustered in the Cluster 3.0 software package (26) with average linkage using the Pearson correlation distance as the metric of similarity between genes (27). *K*-means clustering and figures of merit analysis were carried out with the MeV software suite (28,29). Enrichment of TF binding to promoters in each cluster was determined with Fisher's exact test. *P*-values were Bonferroni-corrected to account for multiple comparisons.

Quantitative real-time PCR

RNA was converted to cDNA with MultiScribeTM Reverse Transciptase using random hexamer priming (Applied Biosystems). Quantitative real-time PCR (qRT–PCR) of 100 ng cDNA was performed in triplicate with SYBR® Green PCR Master Mix on an ABI 7900HT series PCR machine. PCR conditions were 10 min at 95°C (one cycle) followed by 15 s at 95°C and 1 min at 60°C (40 cycles). Analysis was performed using SDS2.3 software. C_T values were computed using the standard curve method. GFP expression levels were normalized to TDH2 expression. Primers for amplification from GFP were 5'-TTTCTGTCTCCGGTGAAGGT-3' and 5'-GACTAAGGTTGGCCATGGAA-3'. Primers for

amplification from the housekeeping gene *TDH2* were 5'-TGACTCCACTGGTGTTTTCAAG-3' and 5'-ACCT TCTTCAATACCGAAAGCA-3'.

Flow cytometry

Approximately 10⁷ cells were harvested by centrifugation and subsequently washed and resuspended in DPBS+0.1% Tween-20. Measurements of GFP fluorescence were performed with a BD LSRII Multi-Laser Analyzer with HTS (BD Biosciences, Sparks, MD, USA). Mean fluorescence values were determined from at least 50 000 cells.

In Figure 3, dose response curves were fit to a Hill function of the form $G(D) = G_{\min} + (G_{\max} - G_{\min}) \frac{D^n}{K^n + D^n}$, where G is the level of GFP, D is the amount of β -estradiol, n is the Hill coefficient and K is the dose yielding half of $(G_{\max} + G_{\min})$. G_{\min} and G_{\max} are constants that are the minimum and maximum mean GFP levels from the data. Data were fit using least-squares regression with the *cftool* function in MATLAB (MathWorks, Natick, MA, USA).

Zinc-finger array design

The Z₄ array was designed to bind a sequence that does not exist in the yeast genome while keeping design principles biased toward GNN sequences that have been shown to be more functional and modular (30,31). The zinc fingers were designed with four primary contacts that have been described: Arg's at position -1 and 6 of each recognition helix to offer G specificity, Asp at position 3 of the helix to offer C specificity and Asn at position 3 to offer A specificity. To avoid high affinity that may be offered by cross-strand contacts, and therefore bias binding by a subset of zinc fingers as opposed to the whole four-fingered array, small amino acids such as Ala and Gly were employed at position 2 of each helix. At the more flexible positions 1 and 5, residues were either chosen because of size or charge. Furthermore, as a fourfingered protein, the backbone of one finger of the threefingered Zif268 was duplicated and recoded. We chose to repeat finger 2 and extend the linker between the second and third fingers (TGSQKP) as has been described (32).

The final assembly used the backbones of Zif268 fingers 1–2–2–3. As a result, the modest design principles noted above were used to design the recognition helices on each

of the four fingers, while the backbone residues were copied from the Zif268 fingers in a finger 1–2–2–3 pattern (the exact coding sequence can be found in the Supplementary Data). Similar finger helices to those designed here have been shown to be functional on these targets when engineered with the OPEN system, giving us confidence that this would be a functional finger array with the desired specificity (31).

Bacterial one-hybrid assay

Characterization of the DNA-binding specificity of the Z₄ protein was performed by bacterial one-hybrid assay as previously described (33,34). The protein was assembled by overlapping PCR using the Zif268 scaffold as a template and cloned into the pB1H2w2 plasmid (34). Using this system, functional binding sites are found by selecting for sequences that are able to activate the *HIS3* reporter gene in bacteria from a 28-bp library of randomized DNA sequences upstream of the promoter (Supplementary Figure S2). Functional binding sites were recovered by sequencing this promoter region from surviving clones (see Supplementary Data for sequences) and analyzed by MEME to discover an over-represented motif that represents the DNA-binding specificity of the protein (Supplementary Figure S2) (33).

RESULTS

Identifying non-yeast DNA-binding domains with few potential genomic-binding sites

In order to produce a synthetic system that has no off-target induction, we sought synthetic target sequences extremely rare or absent from the yeast genome. The Gal4p DNA-binding domain of GEV dimerizes and binds to the sequence 5'-CGG-N₁₁-GCC-3' (Figure 2). Scanning the *Saccharomyces cerevisiae* genome for instances of this motif, we identified more than 500 potential binding sites. The canonical motif recognized by Zif268, the zinc-finger domain used in Z₃EV, is GCGTGGGCG (Figure 2), for which we only find 11 predicted binding sites. The motif recognized by the rationally designed zinc-finger array used in Z₄EV is GCGGCGGAGGAG (Figure 2). The Z₄ zinc-finger array was specifically designed to target this sequence because it is not present in the yeast genome. While the motifs for Gal4p and

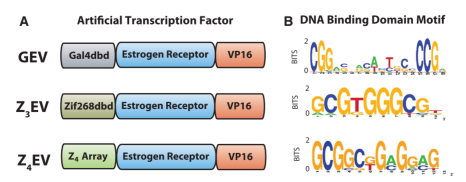


Figure 2. (A) The structure of the ATFs Z₄EV, GEV and Z₃EV. (B) The DNA-binding motifs of Gal4p, Zif268 and the Z₄ array.

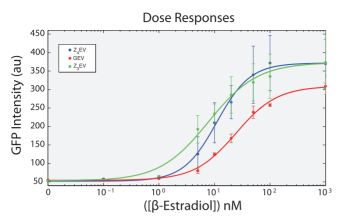


Figure 3. Dose response curves. Strains yMN5 (GEV), yMN7 (Z₃EV) and yMN14 (Z₄EV) were grown to early log-phase and then different amounts of \beta-estradiol were added to the growth medium. GFP was quantified following 12h of induction by flow cytometry. Error bars represent ± 1 SD from three independent cultures.

Zif268 have been published previously, we confirmed the DNA specificity of the Z₄ array with a bacterial onehybrid system (33,34).

Measuring transcriptional output

To characterize the responsiveness of Z_3EV and Z_4EV , we constructed synthetic promoters based on the zinc-finger DNA-binding specificity (see Supplementary Data for sequences). Each promoter mimics the GEV promoter, containing six monomeric binding sites, which allows for three dimerizations of the ER as one would see in the GEV system. These promoters are referred to as Z₃EVpr and Z₄EVpr, respectively. Gal4p recognition sequences used in the previous GEV system were replaced with either the Z_3EV or Z_4EV targeting sequences within the GAL1 promoter (Supplementary Data and Supplementary Figure S1B). Control experiments confirmed that the correct combination of DBD and promoter, as well as introduction of β-estradiol, were required for activation of a target gene (Supplementary Figure S3). When the UAS_{GAL} sequences are removed, Gal4p can no longer induce transcription (Supplementary Figure Furthermore, with the appropriate DBD-promoter combinations, the dose response of each construct was characterized (Figure 3). No induction of GFP was observed for any ATF at concentrations of β-estradiol <1 nM. Interestingly, a similar amount of GFP intensity is achieved with \sim 10-fold less induction of Z₃EV or Z₄EV in comparison to GEV (Figure 3). We determined that the Hill coefficient for each ATF-promoter pair is ~ 1 , consistent with non-cooperative activation transcription.

The effect of ATF induction on gene expression

To assess off-target expression directly, we measured genome-wide gene expression of cells grown to steady state in chemostats containing GEV, Z₃EV or Z₄EV and cognate reporter constructs (Figure 4A), and assessed global gene expression 0, 0.5 and 3h after addition of 1 μM β-estradiol to the steady-state cultures. In response to GEV induction, 155 genes are induced >2-fold and 263 genes are repressed >2-fold 3 h following β-estradiol addition. In contrast only one gene is induced >2-fold in response to Z₃EV activation at 3 h. This gene is YOR343C, which is a dubious open reading frame (ORF) unlikely to encode a protein according to the Saccharomyces Genome Database (35). Similarly, zero genes are induced >2-fold in response to Z₄EV activation 3 h following β-estradiol addition; however, two genes, RPS8A (a ribosomal protein) and YDR133C (a dubious ORF), were induced 2-fold in the Z₄EV time course at 30 min, but returned to 0-fold at 3 h.

Each of the chimeric activators induced a GFP reporter over the course of the experiment. Cells were visualized by microscopy to confirm GFP expression (data not shown), and the level of GFP transcript was measured by gRT-PCR, revealing that each of the chimeric activators induced GFP mRNA >50-fold by 15 min following addition of β-estradiol (Figure 4B). The maximum level of induction for the chimeric activators ranges from \sim 100 - to 300-fold.

The effect of ATF induction on cell growth

We next examined whether these differences in gene expression might be related to the growth defect we previously reported to be conferred by the GEV system. We compared growth between Z₄EV-, GEV- and Z₃EVcontaining strains in the presence of different amounts of β-estradiol (Figure 5A). Between 10 nM and 10 μM β-estradiol, the growth rate of GEV-containing cells decreased ~70% in rich medium (Figure 5B). In contrast, induction of either Z₃EV or Z₄EV with 10 nM-10 μM β-estradiol has no effect on growth (Figure 5B).

Tuning the transcriptional output of Z₃EV

The above results demonstrate the precision with which the Z₃EV or Z₄EV systems are able to activate a single target. However, they do little to indicate how sensitive the system is to the strength of the protein–DNA interaction. To this end, we tested the Z₃EV system against four targets with known differences in binding affinity in comparison to the consensus target of Zif268 (36). Within a 9-fold range of dissociation constants, relative to the consensus sequence, activation of GFP is correlated to the strength of the protein-DNA interaction (Figure 6). Furthermore, a binding-site that offers a 20-fold reduction in affinity is not above background after 6h of induction. These results illustrate that our system is tunable and is unlikely to function on binding sites that offer >20-fold reduction of binding affinity. Future research might investigate the perturbation of networks with finely tuned levels of transcription based on the use of one of these or other characterized binding sites.

Using Z₄EV to probe the Gcn4p transcriptional regulatory network

To test whether this system is able to provide single-gene precision on a genomic target, we decided to target GCN4.

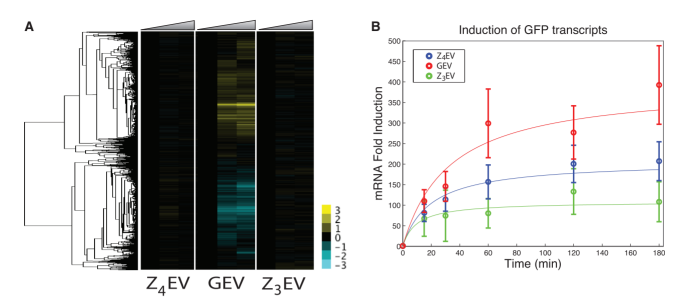


Figure 4. (A) Hierarchically clustered gene expression responses of continuous cultures containing GEV, Z_3 EV or Z_4 EV upon addition of 1 μM β-estradiol (data are time-zero normalized in each experiment). Time points in each experiment are 0, 0.5 and 3 h. Each culture was maintained at a doubling time of 4.07 h. (B) Induction of GFP reporters in cultures from (A) measured by qRT-PCR. Error bars represent ± 1 SEM of three technical replicates. Strains are yMN5 (GEV), yMN7 (Z_3 EV) and yMN14 (Z_4 EV).

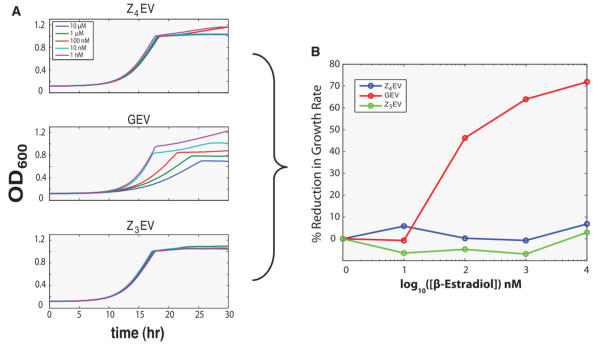


Figure 5. Non-yeast DNA-binding domains eliminate growth defect conferred by GEV induction. (A) Growth curves of GEV-, Z_3 EV- or Z_4 EV-containing strains in the presence of different amounts of β-estradiol. (B) The level of growth reduction conferred by ATFs.

To simplify constructing inducible genomic alleles, we created plasmids (pMN9 and pMN10) with the KanMX cassette, which confers resistance to G418, fused to either Z_3EVpr or Z_4EVpr (template sequences can be found in the Supplementary Data). These synthetic promoters can be inserted into the genome by homologous recombination to make an allele that is conditionally expressed in the presence of β -estradiol (Figure 7A). Gcn4p is a strong

transcriptional activator of a wide range of enzymes required for the production of amino acids (Supplementary Figures S5–S14). Cells lacking GCN4 grow slowly in minimal medium (37), a phenotype we were able to recapitulate in a strain in which GCN4 is under the control of KanMX-Z₄EVpr (Figure 7B). This growth defect is fully alleviated by adding 5 nM β -estradiol (Figure 7B).

Reducing binding affinity reduces expression output 1200 2-fold 5-fold 1000 9-fold No Promot GFP Intensity (au) 800 600 400 200 0 2 3 4 5 time (hr)

Figure 6. Induction of GFP reporters driven by promoters of different affinity by Z₃EV following addition of 1 μM β-estradiol. Error bars represent ± 1 SD of three independent cultures.

GCN4 is rapidly induced by Z₄EV in response to 1 µM β-estradiol (Figure 7C) and in continuous cultures we were able to quantify the dynamic transcriptional impact of GCN4 expression on cells. By 2h, induction of GCN4 resulted in >2-fold induction of 327 genes (\sim 5% of the genome; Figure 7D and Supplementary Dataset S1), which are strongly enriched for cellular amino acid biosynthesis processes (corrected *P*-value = 1.81×10^{-50}). There are 255 genes repressed >2-fold. These genes are enriched for processes involved with cytoplasmic translation (corrected *P*-value = 1.41×10^{-11}), ATP synthesis coupled proton transport (corrected P-value = 6.1×10^{-4}) nucleotide biosynthesis (corrected $= 3 \times 10^{-3}$) (Figure 7D and Supplementary Dataset S1). The presence of Gcn4p binding based on previously published ChIP-Chip data (38) is noted next to the hierarchically-clustered gene expression (Figure 7D), and clearly correlates with induced expression. Based on ChIP-Chip, of the induced genes, 116 have previously been shown to be direct Gcn4p targets. Only five of the repressed genes (DLD3, ICY1, MET3, RFA3 and TIM10) (Supplementary Figure S15) have been shown to be direct targets. Repression is typically mediated at late time points, consistent with an indirect response to Gcn4p production.

To perform an unbiased search for enriched regulatory motifs corresponding to particular expression responses (determined via K-means clustering) we used the FIRE algorithm (39). The number of K-means clusters was determined using figures of merit analysis (Supplementary Figure S16). Two clusters were identified as having been strongly induced by Gcn4p (clusters 1 and 2; Figure 7E). Based on FIRE, promoters of genes in these clusters are enriched for the presence of Gcn4p-binding sequences (Figure 7F; z-scores = 123.7 and 27.1, respectively). Furthermore, cluster 1 genes are enriched for

Gcn4p, Rtg3p, Yap7p, Gln3p and Leu3p binding based ChIP-Chip (corrected *P*-values = 1.47×10^{-55} . 4.77×10^{-5} , 0.006, 0.031 and 0.06, respectively; Fisher's Exact Test). Cluster 2 genes are enriched only for Gcn4p binding (corrected *P*-value = 1.50×10^{-30} ; Fisher's Exact Test). The remaining clusters show no enrichment for Gcn4p binding.

These results, in combination with those detailed above, demonstrate the utility of these expression systems for dissecting complex regulatory networks in vivo. They offer the ability to maintain precise levels of an individual gene's expression. We plan in the future to adapt other DBDs, target sequences or hormone receptors to make it possible to control several inputs simultaneously or in sequence with the same level of precision.

DISCUSSION

The synthetic biology and broader experimental communities have been hampered by a lack of quantitative expression systems for both engineering novel and probing native regulatory circuitry in vivo. In this article, we developed quantitative expression systems for yeast that address this problem. These expression systems achieve single-gene specificity: in an appropriately modified strain a single gene of interest can be selectively activated in an inducer-dependent fashion. Output expression can be tuned to different levels by either changing the amount of inducer or the identity of the binding sites upstream of a target gene. We have taken advantage of the functionality of hormone receptors as effective post-translational switches in S. cerevisiae. With this methodology, we anticipate that by utilizing multiple hormone receptors with orthogonal-binding domains a combinatorial library of tightly-regulated, inducible expression systems can be used to design and program more sophisticated regulatory functions. These may also be combined with IPTG- and doxycycline-based expression systems, the most recent of which have been characterized in (11).

Here, we have used the Cys₂ His₂ zinc-finger DBD to provide the necessary affinity and specificity to site-specifically activate the target of interest. However, transcription activation-like (TAL) effector domains have emerged as promising tools for engineering DBDs in eukaryotes. TAL effectors are virulence factors utilized by pathogenic bacteria in the genus *Xanthomonas* (40). Specificity is determined by repeat variable di-residues (RVDs) at positions 12 and 13 while RVD regions that recognize each of the four nucleotides have been identified (41,42). Their modularity and relative simplicity suggests that TAL domains can be stitched together to form novel DNA-binding domains that recognize any DNA sequence of interest. Synthetic TAL effectors have been constructed that function as activators or repressors of eukaryotic transcription (43). Therefore, if applications would benefit from the use of multiple inducers used simultaneously, a conversion to TAL-hormone receptor system might be beneficial. Still, limitations in the TAL assembly may result in the continued use of zinc fingers

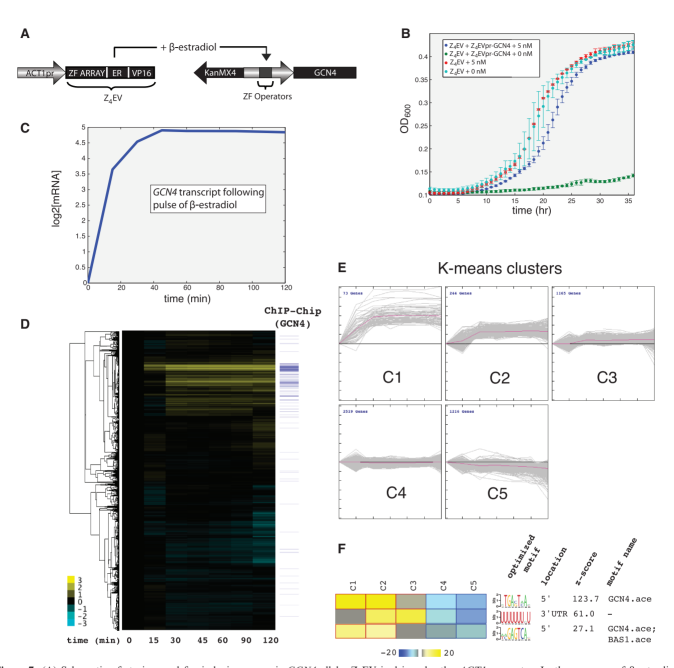


Figure 7. (A) Schematic of strains used for inducing genomic GCN4 allele. Z_4EV is driven by the ACT1 promoter. In the presence of β-estradiol, Z_4EV activates transcription of the native GCN4 allele driven by the synthetic promoter Z_4EV pr. KanMX is expressed in the reverse orientation. (B) Measuring growth Z_4EV -containing haploids containing GCN4 or Z_4EV pr-GCN4 in the presence or absence of 5 nM β-estradiol. Cells were grown in low phosphate chemostat medium. (C) The transcriptional response of Z_4EV pr-GCN4 (strain = DBY12423) in response to 1 μM β-estradiol in a continuous controlled maintained under phosphate limitation. The culture was maintained at a doubling time of 6.3 h. (D) The global transcriptional response of cells in (C) measured out to 2 h. We denote the presence of Gcn4p binding to a promoter by a blue line [(38) P-value <0.001]. (E) Expression data from (D) were divided into five expression clusters using K-means clustering with the Euclidean distance similarity metric. The optimal number of clusters over-represented and under-represented motifs within these clusters was determined using the FIRE algorithm (F).

(size, repetitive coding sequence, etc.). Moreover, as we have demonstrated in this manuscript, extensive biochemical characterization such as knowledge of Cys₂His₂ DNA-binding affinity can be used to predictably tune output expression of distinct targeting sequences. The wealth of literature on the zinc-finger protein–DNA interaction may prove advantageous toward this end.

Different inputs can be used to stimulate a network and reveal its relevant components and architecture. A 'transfer function' relates a prescribed input to a measured output. In the described case, we provided a step increase of Gcn4p to identify and quantify the responsive outputs of Gcn4p activation. It is, however, often desirable to provide alternative stimuli. For example, to determine the bandwidth of a signaling pathway, the pathway input can be varied at different frequencies and the output can be measured using fluorescently tagged reporters. This approach has been

used previously to show that the Hog1p MAPK-signaling cascade acts as a low-pass filter in response to fluctuations in extracellular salt concentration and quantify its bandwidth (44). Future work may investigate how to directly provide cells with varying amounts of protein at desired frequencies. While varying extracellular concentrations of stimulus is relatively straightforward with microfluidic-based approaches, varying intracellular levels of protein over time presents a significant technical challenge. One solution may be the implementation of light-responsive expression systems, which can induce expression of a target gene in the presence of a particular frequency of light, and ablate expression in its absence (45).

We previously demonstrated that by inducing the TEV protease with GEV, we could rapidly degrade an N-degron-tagged protein with a half-life of <15 min (7). By combining two orthogonal induction systems (one for inducing an N-degron-tagged allele and a second for inducing TEV) one could provide cells with different size and/or length protein pulses. In response to the first inducer the protein would be synthesized, and in response to the second inducer, the protein would be destroyed by the protease. This approach could also utilize the TIR1 ubiquitin ligase from Arabidopsis thaliana, which is post-translationally induced by the auxin indole-3-acetic acid and is functional in yeast (46). This may be advantageous because TIR1 can be expressed constitutively and induced by auxin without having to be first transcribed and translated as in the case of TEV.

In conclusion, synthetic approaches can aid in the understanding of native regulatory machinery and provide the toolkit to design and implement rationally designed genetic programs. We demonstrated the utility of a perturbative approach in quantifying the downstream effects of a major transcriptional regulator of amino acid biosynthetic genes. More broadly our work provides a new set of expression systems for a variety of experimental and potentially industrial applications. These systems serve as a flexible platform that can be used to tune, integrate and re-purpose eukaryote functions.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online: Supplementary Tables 1 and 2, Supplementary Figures 1–16 and Supplementary Dataset 1.

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