- 1 Short title: Reporter system for transcription factor complex DNA interactions
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9 10	DIMR, a Yeast-Based Synthetic Reporter System for Probing Oligomeric Transcription Factor DNA Binding
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17 18 19 20 21	<u>One sentence summary:</u> The DIMR system provides an accessible and easy-to-use platform to elucidate DNA binding and transcriptional regulatory capacity of oligomeric transcription factor complexes
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29 30 31 32	<u>Funding information:</u> Funding was provided by an EAGER grant from the National Science Foundation (MCB 1747539)
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39 40 41	Abstract
42 43 45 46 47 48 49 50 51	Transcription factors (TFs) are fundamental components of biological regulation, facilitating the basal and differential gene expression necessary for life. TFs exert transcriptional regulation through interactions with both DNA and other TFs, ultimately influencing the action of RNA polymerase at a genomic locus. Current approaches are proficient at identification of binding site requirements for individual TFs, but few methods have been adapted to study oligomeric TF complexes. Further, many approaches that have been turned toward understanding DNA binding of TF complexes, such as electrophoretic mobility shift assays, require protein purification steps that can be burdensome or scope-limiting when considering more exhaustive experimental design. In order to address these shortfalls and to facilitate a more streamlined approach to
52 53	understanding DNA binding by TF complexes, we developed the DIMR ( <u>Dynamic</u> , <u>Interdependent TF binding Molecular Reporter</u> ) system, a modular, yeast-based synthetic

 $\underline{I}$  interdependent IF binding <u>M</u>olecular <u>R</u>eporter) system, a modular, yeast-based synthetic transcriptional activity reporter. As a proof of concept, we focused on the NUCLEAR 55 FACTOR-Y (NF-Y) family of obligate heterotrimeric TFs in *Arabidopsis thaliana*. The 56 DIMR system was able to reproduce the strict DNA-binding requirements of an 57 experimentally validated NF-Y<sup>A2/B2/C3</sup> complex with high fidelity, including recapitulation of 58 previously characterized mutations in subunits that either break NF-Y complex 59 interactions or are directly involved in DNA binding. The DIMR system is a novel, 60 powerful, and easy-to-use approach to address questions regarding the binding of 61 oligomeric TFs to DNA.

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### 63 Introduction

Transcription factors (TFs) are a fundamental component of biological control, facilitating 64 differential gene expression in response to environmental stimuli and making complex life 65 possible. The mechanism through which differential gene expression manifests is 66 inherently complex, requiring coordination of stimulus perception, transcriptional and 67 68 translational responses, and alteration of relevant protein activity (Crick 1970). The transcription-level integration of stimuli is of particular interest, as this process supports 69 context-specific recruitment of RNA polymerase to a specific genomic locus (Diamond et 70 al. 1990). This integration is accomplished through the independent, competitive, and 71 72 cooperative functions of TFs and the effects of these relationships on DNA-binding and RNA polymerase recruitment (Amoutzias et al. 2008; Lickwar et al. 2012). A better 73 74 understanding of the mechanisms through which transcriptional activity are modulated, particularly how TF interactions influence regulatory capacity, could facilitate the design 75 76 of more precise and predictable molecular tools to address long-standing issues in the 77 fields of agriculture, health care, and manufacturing, among others (Khalil and Collins 78 2010).

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80 Current approaches to identify TF-DNA interactions are effective at *de novo* identification of binding site preferences of individual TFs in vitro and at identifying global binding 81 82 patterns of individual TFs in vivo. In many cases, a combination of approaches can be used for a more complete understanding of TF targeting. In particular, many researchers 83 have found great success in combining approaches that effectively identify binding 84 85 specificity (such as Protein Binding Microarrays (PBMs), 1-Hybrid-based library 86 screening, or DNA Affinity Purification (DAP-seq)) and lower-resolution, global-scale 87 binding site identification (such as Chromatin Immunoprecipitation (ChIP-seq) or Assay for Transposase-Accessible Chromatin (ATAC-seq)). 88

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While we will not focus on the general strengths of each approach or combination of approaches (see reviews, (Mahony and Pugh 2015; Jayaram et al. 2016)), one aspect of TF function that remains less explored is the impact that TF oligomerization exerts on DNA binding (Jolma et al. 2015). Most approaches, including those introduced above,

94 have focused on understanding some aspect of DNA-binding of an assumed individual TF: however, a much more complex interplay of TF function exists than can be readily 95 and methodically addressed with current techniques. Transcriptional regulation through 96 97 multi-component TF complexes is a common theme, but our understanding of how these complexes identify and interact with specific genomic loci remains less explored (Jolma 98 99 et al. 2015; Morgunova and Taipale 2017). For example, a recent study identified dramatic shifts in cognate binding sequences of bZIP homodimers compared to 100 101 heterodimers (Rodriguez-Martinez et al. 2017); however, the experimental approach 102 used in this study required purification of a large suite of individual transcription factors. 103 An approach enabling rapid and straight-forward testing of complex TF units, with the ability to make targeted mutations in any component of the TF complex or its potential 104 105 binding site, would create a new lens through which we could explore more nuanced 106 aspects of TF-DNA interactions.

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108 Traditional definitions of transcription factors often highlight the presence of both a DNA-109 binding and a transcriptional-regulation domain; however, many TFs function as obligate protein complexes, where individual components contain partial DNA-binding or 110 111 transcriptional-regulation domains that are only fully-reconstituted within the context of a functional complex. One example of such an arrangement can be found in the NUCLEAR 112 113 FACTOR-Y (NF-Y) transcription factors, a complex that functions as an obligate heterotrimer of three distinct subunits, NF-YA, NF-YB, and NF-YC, and physically interacts with 114 115 the DNA sequence CCAAT. Although this complex was initially identified over 30 years ago (Dorn et al. 1987; Jones et al. 1985), guestions remain regarding the molecular forces 116 117 driving specificity of the NF-Y complex to CCAAT. Several key observations and studies 118 have led to persistent interest in the NF-Y, particularly in plant lineages, including: (1) the ubiquitous nature of the CCAAT box and, subsequently, NF-Y regulated genes, (2) the 119 120 expansion of the NF-Y subunit families in plants compared to animals, where animals 121 encode 1-2 members of each subunit, while higher plants often encode 10+ of each 122 subunit (Siefers et al. 2009), and (3) the identification and molecular characterization of the first non-canonical NF-Y complex in plants, which replaces an NF-YA subunit for a 123

plant-specific CCT (<u>CONSTANS</u>, <u>CONSTANS-LIKE</u>, and <u>TOC1</u>) domain-containing
 protein, leading to altered DNA binding properties (Gnesutta et al. 2017).

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Published crystal structures of various NF-Y complexes, particularly of the full Homo 127 128 sapiens NF-Y complex bound to DNA (Nardini et al. 2013), have raised increasingly 129 complex questions regarding the specificity of the NF-Y complex for a particular CCAAT box. While all three NF-Y subunits have been shown to make physical contact with DNA, 130 131 it appears that only NF-YA makes sequence-specific contact, inserting directly into the 132 minor groove of the CCAAT box, while NF-YB and NF-YC make contacts with the DNA 133 backbone, appropriately positioning both the DNA and NF-YA subunit for tight physical 134 interaction (Nardini et al. 2013). Despite this understanding, the NF-Y complex appears 135 to show more selectivity than has been experimentally derived. For example, most CCAAT boxes in human cell lines are not consistently bound by NF-Y complexes (Encode 136 137 Project Consortium et al. 2007; Zambelli and Pavesi 2017), meaning that the mere presence of a CCAAT box is not predictive of NF-Y binding. Factors such as chromatin 138 139 accessibility and repressive epigenetic modifications regularly limit the binding landscape of a given TF (John et al. 2011); however, the NF-Y are thought to function as 'pioneer' 140 141 TFs that are able to bind less-accessible DNA and promote further TF complex formation (Tao et al. 2017; Oldfield et al. 2014; Donati et al. 2008). Alternatively, while the only strict 142 143 requirement on NF-Y binding is the presence of the pentanucleotide CCAAT, the flanking 144 nucleotides could serve as a fine-tuning mechanism for complex binding and stability. 145 Ultimately, the source of this CCAAT box selectivity is not well understood and is made 146 significantly more complex when considering the expanded family size and combinatorial 147 complexity of plant NF-Y subunits.

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In particular, we hypothesized that different NF-YB/NF-YC dimers might contribute to *CCAAT* box selectivity through interactions with the nucleotides flanking the *CCAAT* pentamer; however, comparing all possible NF-YB/NF-YC combinations with even a single NF-YA component would be a significant undertaking, with 100 possible combinations in *Arabidopsis thaliana* (Petroni et al. 2012) and 208 possible combinations in *Oryza sativa* (rice, (Hwang et al. 2016)). *In vitro* approaches requiring protein 155 purification, such as Electrophoretic Mobility Shift Assay (EMSA), become time- and cost-156 prohibitive at this scale. More importantly, many proteins are recalcitrant to protein 157 purification, and can only be isolated if appropriately truncated or co-expressed with other 158 factors. Despite these technical limitations, EMSA analysis is well-suited for these types 159 of investigations, and any approach intended to address similar questions would need to 160 match or complement its capabilities. A reporter system with a large dynamic range and 161 high sensitivity would allow scientists to address mutations that have much more modest effects on TF-DNA interactions, and could facilitate examination of more precise or 162 163 biologically-relevant questions.

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To address these issues and to better facilitate these types of research, we designed the 165 166 DIMR (Dynamic, Interdependent TF binding Molecular Reporter) system (Figure 1), a modular yeast-based transcriptional activity reporter system developed through repeated 167 168 application of the synthetic biology approach of build-test-learn. The DIMR system allows 169 for dose-dependent induction of a suite of transcription factors (effectors) and subsequent 170 monitoring of transcriptional regulation. We validated our approach by testing the DNA binding and transcriptional activation capabilities of the Arabidopsis NF-Y<sup>A2/B2/C3</sup> complex 171 172 and found that the DIMR system was able to faithfully recapitulate both wild-type CCAAT 173 box binding and previously-described mutations impacting DNA binding and complex 174 formation.

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### 176 **Results**

### 177 DIMR system components, design philosophy, and functional description

178 The DIMR system is composed of three modules: (1) an Activator Module, encoding an 179 inducible artificial transcription factor (ATF), (2) an Effector Module, encoding a single ATF-driven cassette containing viral 2A 'cleavage' sites between effector components. 180 181 and (3) a Reporter Module, containing a dual-luciferase reporter composed of a 182 constitutively transcribed Renilla luciferase and a conditionally transcribed Firefly 183 luciferase (Figure 1A). Each module is carried on a yeast shuttle vector with different 184 auxotrophic selection, and each has been designed to be as modular as possible, with 185 restriction enzyme sites strategically placed to facilitate swapping individual effector or

reporter components with minimal laboratory effort. As currently implemented, point 186 187 mutations in existing effectors can be easily accomplished through commercially available 188 site-directed mutagenesis kits, novel effector modules can be synthesized de novo at 189 affordable rates, and permutations of binding sites can be cloned in as little as 3 days at 190 a relatively low cost and with very minimal active time, resulting in a streamlined, quick, 191 and hands-off assay to investigate TF-DNA interactions (Figure 1B). Finally, because of 192 the synthetic biology-based approach taken during the design and implementation of the 193 system, each component of the modules can easily be further iterated upon for improved 194 function or to accomplish different goals.

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The Activator module (Figure S1) utilizes the chimeric Z<sub>4</sub>EV artificial transcription factor, 196 197 which contains three functional domains: (1) an engineered zinc finger with specificity to a DNA sequence not found in the genome of Saccharomyces cerevisiae, (2) an estrogen 198 199 receptor that modulates activity and localization, and (3) a VP16 activation domain 200 (McIsaac et al. 2013). Z<sub>4</sub>EV is constitutively expressed under the ACT1 promoter 201 (Flagfeldt et al. 2009) but remains inactive and restricted to the cytosol. Activation of Z<sub>4</sub>EV 202 with the hormone  $\beta$ -estradiol initiates translocation to the nucleus and activation of the 203 Effector cassette. This activation was previously shown through transcriptome profiling to 204 induce remarkably few unintended effects, either through the action of  $\beta$ -estradiol itself or 205 through off-target binding of Z<sub>4</sub>EV (McIsaac et al. 2013).

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207 The Effector module (Figure S2) is designed to emulate a polycistronic message through 208 the incorporation of viral 2A 'self-cleaving' sequences. Our initial designs included 209 cassettes expressing both 2 and 3 genes, with a translationally-fused flexible linker and 210 unique epitope tag on each (Sabourin et al. 2007). The linkers between components A – B and B – C also encode previously-validated T2A peptide sequences (Beekwilder et al. 211 212 2014), facilitating translation of individual proteins from a single mRNA. Restriction 213 enzyme recognition sites were embedded into the coding sequences of the T2A linkers 214 through the introduction of silent mutations. This allows easy swapping of effectors through either (1) the inclusion of the appropriate flanking linker sequence through gene 215

synthesis (recommended, along with codon optimization (Kotula and Curtis 1991;
Gustafsson et al. 2004)) or (2) PCR amplification to produce the appropriate over-hangs.

219 The Reporter module (Figure S3) encodes a dual luciferase reporter system, including a 220 Renilla luciferase variant constitutively expressed under the ACT1 promoter, and a 221 conditionally expressed Firefly luciferase variant. Design of the Firefly luciferase promoter 222 drew heavily on previously characterized yeast promoters bound by yeast NF-Y 223 complexes. Specifically, we chose native, experimentally validated promoters of 224 Saccharomyces cerevisiae whose activation required the presence of a specific CCAAT 225 box (implying direct regulation by the NF-Y), and replaced the required CCAAT box binding site with the various permutations described below. Cloning of binding site 226 227 permutations is quick and simple, requiring only a pair of appropriately designed, 5'phosphorylated oligonucleotides and simple restriction enzyme digestion and ligation 228 229 reactions.

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### 231 Validation of effector induction and cleavage

As a starting point, we tested our ability to induce accumulation of Arabidopsis NF-Y<sup>A2/B2/C3</sup> effectors through the activation of Z<sub>4</sub>EV. After a 6-hour induction period, we were able to visualize accumulation of NF-YA2:HA (Figure 2A), NF-YB2:MYC (Figure 2B), and NF-YC3:FLAG (Figure 2C). Differences were seen in the level of protein accumulation across samples, and several instances of failed T2A-mediated cleavage were clear; however, we were able to clearly and consistently observe individual NF-Y subunits in repeated experiments.

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Further experiments to explore the inducibility of the effector cassette identified detectable levels of individual effector proteins with around 5 nM  $\beta$ -estradiol (Figure S4), with no obvious differences in accumulation between the different NF-Y subunits. A short timecourse of effector accumulation at 10 nM  $\beta$ -estradiol identified fairly stable accumulation of individual effector proteins at 3 and 6 hours, though we often observed a peak accumulation at 6 hours and a drop-off in signal when extending to 9 hours (Figure S5). From these and other preliminary data, we established standard low- and high-induction

- ranges of ~1 nM and ~10 nM  $\beta$  -estradiol, respectively, and a standard induction length of 6 hours. While incomplete cleavage of individual effectors remains to be addressed,
- the induction scheme of the DIMR system is effective and robust.
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### 251 Validation of reporter activity following effector induction

252 Our initial system design and testing were built upon a previously-characterized NF-Y regulated CCAAT box from the FLOWERING LOCUS T (FT) promoter in Arabidopsis. 253 254 This CCAAT box, as well as 20 flanking base pairs upstream and downstreat, was 255 positioned immediately upstream of the minimal NF-Y regulated yeast promoter of CITRATE SYNTHASE1 (CIT1). Upon induction of the NF-Y<sup>A2/B2/C3</sup> effector cassette, we 256 observed a significant, dose-dependent increase in relative luminescence (Figure 3A). 257 258 Normalization of the data into a fold change value, relative to mock induction, found an average ~3.5-fold increase in high induction conditions (10 nM, Figure 3B). Despite the 259 260 successful recapitulation of NF-Y DNA binding in our initial designs, we were not able to 261 statistically distinguish between low- and high-induction conditions, and the maximum fold 262 changes observed were lower than desired. A larger dynamic range of reporter activation 263 would facilitate addressing more nuanced questions, such as the DNA binding impact of 264 individual effector mutations or changes in cognate binding sites.

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### 266 <u>Refinement of binding site design</u>

267 From these initial tests, we next sought to refine the DIMR system for increased reporter 268 dynamic range. We focused our system tuning on two approaches: reducing mock-level 269 signal and increasing maximum activation. To decrease mock-level signal, we designed 270 and tested different NF-Y regulated minimal promoter architectures to identify reporters 271 with lower basal level activation. We focused on promoters for the genes CIT1 and ASPARAGINE SYNTHETASE1 (ASN1), driven by the above-described FT CCAAT box 272 273 footprint (Figure 4A). Luciferase activity levels were lower in mock-treated samples with 274 the ASN1-based promoter compared to the originally-tested CIT1-based promoter 275 (Figure 4B). Notably, this reduction in mock-level activation translated to a larger fold 276 change of ~6x in ASN1-based reporters when comparing mock and high induction 277 conditions.

### 278

279 To increase our maximum signal, we additionally tested the effects of different numbers 280 of binding sites and the spacing between them. Multimerization of available binding sites 281 can result in an increase in observed transcriptional activity (Khalil et al. 2012), but this 282 larger DNA footprint can be more difficult to accommodate in cloning. To abrogate this, 283 we also tested the effects of reducing the length of each individual binding site footprint. First, we attempted to cut the CCAAT box footprint roughly in half by including only 10 284 285 base pairs of flanking sequences in single, double, and triple binding site configurations 286 (Figure 4C), and found that multimerization of the binding site increased total system 287 activation. Finally, we combined the original, larger CCAAT box footprint into single, double, and guadruple binding site configurations (Figure 4D). In this set, the guadruple 288 289 binding site configuration stands out at a much-improved ~15-fold increase in reporter activity from mock. The larger DNA footprint of the quadruple binding site approach 290 291 required a slightly modified cloning approach, where two pairs of annealed, 292 phosphorylated oligos were simultaneously ligated into the Reporter module; however, 293 we were able to preserve the flexibility of the DIMR system and avoided the need for 294 further gene synthesis for individual binding site permutations.

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### 296 Validation and assessment of refined binding site setup

297 With an increased signal level upon system activation, we began testing the activation requirements of the DIMR system. First, we examined dose-dependent system activation 298 levels of the NF-Y<sup>A2/B2/C3</sup> complex on the quadruple *CCAAT* box over a wide range of  $\beta$ -299 300 estradiol induction levels (Figure 5A). This induction gradient was analyzed through 4-301 parameter logistic regression (4PLR), with an R<sup>2</sup> value of 0.942. No system activation 302 was observed at induction levels below 1 nM, while activation peaked between 10 and 100 nM β-estradiol. The dose-dependent activity of the system was most pronounced 303 304 between 1 nM and 10 nM, with a half maximal effective concentration (EC<sub>50</sub>) of ~2.2 nM 305 and total span of ~10.5-fold change. 4PLR is a widely used metric in chemistry, 306 biochemistry, and computational biology to describe cooperative action, and in this case, 307 statistically supports the idea that the individual NF-Y subunits are working 308 interdependently in an induction-dependent manner to bind the CCAAT box.

### 309

310 To further solidify the correlation between effector induction and system activation, we 311 tested the impacts of loss of individual DIMR modules, previously described mutations in 312 NF-Y effectors, and changes in the CCAAT box (Figure 5B). Replacing either the  $Z_4EV$ 313 activator cassette or the NF-Y effector cassette with empty vectors resulted in a complete 314 loss of system activation upon induction. Importantly, we found that mutation of the CCAAT box to CCAGC eliminated system activation. These three pieces of data 315 316 collectively suggest that the observed induction responses are accomplished through the 317 action of the NF-Y complex on the CCAAT box driving the firefly luciferase reporter.

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We also performed further system tests with the NF-YB2 E65R point mutation, a 319 320 previously characterized mutation shown to prevent interaction of the NF-YB/NF-YC dimer with NF-YA (Sinha et al. 1996; Siefers et al. 2009). As expected, this point mutation 321 322 abolished system activation (Figure 5B). While many highly-conserved residues in each 323 of the NF-Y subunits have been previously identified and characterized to completely 324 break DNA binding (Sinha et al. 1996; Kim et al. 1996; Sinha et al. 1995), we wanted to 325 examine novel mutations that could help reveal the evolutionary constraints acting on the 326 NF-Y complex. To this end, we examined the effects of one particular point mutation at a residue thought to be directly involved in CCAAT box binding, NF-YA2 H183A. With the 327 328 native histidine residue replaced with an alanine at this location, we still observed a 329 moderate level of system activation (Figure 5B). This alanine-replacement approach is 330 traditionally used to replace residues that contribute to sequence-specificity with an 331 unobtrusive and relatively-inert alanine that is unlikely to mediate sequence-specific 332 interactions (Luscombe et al. 2001).

333

Because we had previously observed instances of failed 2A site function in the form of fused effector components, we directly tested whether these fused effectors were functional by creating point mutations that break the existing 2A sites. We found that fusion of the NF-YA and NF-YB components did lower maximum reporter activity compared to a 2A-cleaved trimeric system or a fusion between NF-YB and NF-YC, but each instance of broken 2A cleavage was still able to significantly activate the reporter

system (Figure S6). While not all transcription factor complexes will be able to effectively
form and regulate DNA while fused together, this should be tested for each group of
transcription factors being tested.

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Finally, we tested whether yeast NF-Y orthologs might be able to compete with or take 344 345 the place of our effector module-encoded Arabidopsis NF-Y proteins. Sequentially removing one NF-Y subunit from the otherwise-complete effector module induced no 346 347 system activation (Figure S7), suggesting that native NF-Y orthologs are not responsible for any significant activation of the reporter. While interactions between NF-Y orthologs 348 349 of different species has been demonstrated (Calvenzani et al. 2012), we hypothesized 350 that the level of induction through the activator and effector modules would guickly 351 saturate any unintended interactions. Further, not all yeast NF-Y orthologs are thought to 352 be constitutive expressed, with at least one ortholog only expressed in response to non-353 fermentable carbon sources (Bourgarel et al. 1999).

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### 355 Validation of CCAAT-based NF-Y binding

356 To further explore the affinity of the NF-Y complex to the CCAAT box, we tested 357 permutations of the CCAAT box with variation at the 3' end, corresponding to all possible combinations of CCANN (Figure 6). We focused on variation at these locations because 358 359 the crystal structure of the DNA-bound human NF-Y complex identified many more direct 360 interactions with NF-YA and the first three bases of the CCAAT box than the fourth and 361 fifth bases. Beyond this, our earlier exploration of the NF-YA2 H183A mutation suggests 362 possible differences in the way human and plant NF-Y complexes bind DNA, an idea 363 further supported by key differences in the otherwise-conserved linker region of NF-YA2. 364 This linker region was proposed to be important for precise positioning of flanking  $\alpha$ helices for proper NF-Y complex stabilization and DNA binding, and NF-YA2 is the only 365 366 family member in Arabidopsis with this unusually long linker region.

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368 Unsurprisingly, the highest-activated binding site of the CCA<u>NN</u> suite was CCA<u>AT</u> at ~14-

fold over mock, with no other binding site permutation activating greater than 2.2-fold over

mock (Figure 6A). Among these non-CCAAT binding sites, we saw a small increase in

371 reporter activity in *CCA<u>N</u>T*, but not in *CCAA<u>N</u>* (Figure 6B). A targeted comparison of these 372 two binding site permutations identified significant increases relative to mock in all 373 *CCA<u>N</u>T* binding sites, but none other than *CCAAT* in the *CCAA<u>N</u>* variants (Figure 6C). 374 While further investigation is necessary, this observation runs counter to what is currently 375 understood about DNA binding by the NF-Y complex in humans, as a direct physical 376 interaction between NF-YA has been identified at the fourth position, but not the fifth 377 position, of the CCAAT box.

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### 380 Discussion

### 381 <u>Using DIMR to investigate TF complex-DNA interactions</u>

382 In spite of the importance and ubiquitous nature of multi-component transcription factor assemblies, relatively few molecular or biochemical approaches have been devised to 383 384 explore the intricacies of DNA binding by these multimeric complexes. Current and widely 385 adopted methods are not well-suited for many questions, particularly those that could 386 benefit from extensive mutagenesis-based analyses, such as alanine scanning mutagenesis to probe the relative contributions of DNA binding amino acids within a given 387 388 TF. After fine-tuning and optimizing our induction and binding site schemes, we ultimately produced a sensitive, straightforward, and versatile reporter system for assaying 389 390 transcription factor complex DNA interactions. As one example of this type of study, we 391 examined the effect of permutations of the CCAAT box and identified a slightly higher 392 level of system activation when altering bases at position 4 than position 5 (CCANT vs. 393 CCAAN). Our system by passes the need for protein purification or deep sequencing, two 394 common requirements for other approaches. The proposed system can easily be scaled 395 to simultaneously include dozens of effector or binding site permutations with relative simplicity, and requires only relatively basic lab equipment. The injector-fitted, 396 397 luminescence-based plate reader is likely the only equipment not commonly found in a 398 standard molecular lab, and these types of equipment are often housed in core facilities 399 available to most universities or institutes.

400

### 401 Incomplete cleavage at T2A sites in effector module

402 Our initial designs sought to leverage viral 2A peptide sequences to generate equimolar 403 amounts of individual effectors through a cleavage mechanism during translation, an 404 important goal when considering obligate oligomeric TF complexes. We use the term 405 cleavage throughout; however, strictly speaking, the end result is an inability to form the peptide bond between two adjacent amino acids, not true cleavage of an existing peptide 406 407 bond (Kim et al. 2011). Because eukaryotes do not utilize the polycistronic mRNA approach so common in bacteria, researchers have turned to 2A sites and other 408 409 approaches to emulate this effect. While not a perfect solution to this problem, we found that viral T2A sites worked sufficiently well for our needs here. It is important to note, 410 however, that incomplete cleavage of the effector cassette does alter the interpretation of 411 412 system activation. Therefore, with the shortcomings of this design, we can test whether a 413 transcription factor complex can bind a particular consensus sequence, but we cannot directly address whether a particular complex *can or cannot form*. While this distinction is 414 415 fairly minor in many cases, particularly when working with previously-described complexes, it is an important limitation of the system that should be addressed in future 416 417 iterations. Unfortunately, no completely effective eukaryotic-based system has been 418 described to recreate the polycistronic mRNA system so widely employed in bacteria 419 (Blount et al. 2012). However, new 2A peptide sequences are still being identified and 420 have been seen to vary in efficiency from organism to organism (Luke et al. 2010), raising 421 the possibility that a 2A site more efficient in yeast remains to be identified.

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To circumvent this issue, we have begun designing effector cassettes with individual components driven by separate inducible promoters. These separate promoters each contain the Z<sub>4</sub>EV -bound consensus sequence, but utilize different minimal promoters to reduce recombination within yeast (Broach et al. 1982). Whether this modification will lead to relatively-equal protein accumulation is unknown; however, it should at least provide a framework to begin further modifying, tweaking, and improving the system for our specific needs.

430

# 431 Considerations and improvements on the DIMR system

While much of the parameter refinement presented above should help inform novel 432 433 design considerations, some applications will require more significant alterations. In 434 particular, issues are likely to arise where effector complexes do not possess intrinsic 435 transcriptional activation potential or when a binding site is bound and autoactivated by endogenous yeast proteins. In cases where complexes completely lack activation 436 437 potential, an activation tag might be fused to one or more effectors (Knop et al. 1999); however, whether this approach could overcome active transcriptional repression is 438 439 unclear. Similarly, instances of reporter autoactivation might be addressed through fusion of one or more effector components to a transcriptional repressor domain (Edmondson 440 et al. 1996) and measuring inducible *reductions* of reporter activity. While we have not 441 442 yet worked through either of these concerns, the modular design of the system should 443 allow for relatively straight-forward modification and testing.

444

### 445 <u>Teasing apart the details of NF-Y DNA binding: a case for DIMR</u>

Despite consistent investigations into the mechanistic function of the NF-Y in animals 446 447 over the past 30 years, many aspects of complex formation and DNA binding remain relatively unexplored in plant NF-Y orthologs. First, a significant amount of plant NF-Y 448 449 research, including most DNA-binding assays, has been conducted using only conserved 450 domains of each NF-Y subunit. In fact, the survey presented above represents one of the 451 first comprehensive examinations of DNA binding and transcriptional regulation by a 452 complete, full-length plant NF-Y trimeric complex. While the core domains of each NF-Y 453 subunit show significant conservation across all eukaryotic lineages, the flanking termini 454 show remarkable divergence from one another. The biological significance of these 455 flanking regions has remained elusive in most cases, though several protein-protein 456 interactions with NF-Y complexes are thought to be mediated through NF-Y terminal domains (Cao et al. 2011). A better understanding of the functions of the less-conserved 457 458 termini is critical for plant NF-Y research in particular, as the vast majority of differences 459 between individual members of expanded NF-Y gene families are found in these flanking 460 regions. The DIMR system is capable of facilitating a systematic domain-swap approach of the flanking regions of NF-YB or NF-YC paralogs, and could potentially uncover 461

462 changes in NF-Y complex affinity or specificity achieved through modulation of the463 specific NF-Y subunits within a functional complex.

464

Like NF-YB and NF-YC, the conserved domains of NF-YA family members are 465 466 remarkably similar to one another. One critical exception to this observation is a 4-5 amino 467 acid elongated linker sequence encoded in the NF-YA2 subunit. This linker is positioned between two highly-conserved alpha helices that are each responsible for NF-Y complex 468 formation or DNA binding of the mature complex, and in addition to influencing the 469 470 positions of the flanking alpha helices, this linker also makes physical contact in several 471 places with the sugar-phosphate backbone of the CCAAT box (Nardini et al. 2013; Romier et al. 2003). While experimental evidence is necessary to understand the impact and 472 473 significance of this atypical linker, it is possible that NF-Y complexes containing NF-YA2 have slightly different DNA binding profiles or NF-Y complex dissociation constants. It 474 475 should be noted, however, that NF-YA2 is likely the best-characterized plant NF-YA 476 subunit, as it has been a major focus of research through its regulation of photoperiodic 477 flowering. While the majority of our observations regarding NF-YA2 DNA binding and complex formation have supported the crystal structure of the DNA-bound human NF-Y 478 479 complex, many important aspects of NF-Y form and function have not been thoroughly investigated and developed. 480

481

482 In fact, many of the most exciting and significant advances in NF-Y research have 483 occurred in the last two years through the identification and characterization of the first non-canonical NF-Y complex. This complex utilizes the NF-YB2/NF-YC3 dimer to 484 485 stabilize interactions between CONSTANS (CO) and its consensus binding site. This 486 complex, termed NF-CO, recognizes a CCACA motif found in the proximal promoter of the FLOWERING LOCUS T (FT) gene and is critical for proper photoperiodic floral 487 488 induction (Gnesutta et al. 2017). The interaction between CO and the NF-YB/NF-YC 489 dimer is mediated through a conserved, plant-specific CCT (CONSTANS, CONSTANS-490 *LIKE*, and *TOC1*) domain that is found in over 40 genes in Arabidopsis (Griffiths et al. 2003; Farré and Liu 2013), raising the possibility of a significantly-expanded pool of 491 492 potential NF-Y or NF-CCT complexes. Excitingly, several families of CCT domain

493 containing proteins have been extensively studied and shown to be of critical importance 494 for photoperiodic flowering (Griffiths et al. 2003), circadian clock entrainment and 495 maintenance (Mizuno 2004), and seedling environmental responses (Reyes et al. 2004). 496 Unfortunately, testing hypotheses concerned with these non-canonical NF-CCT 497 complexes is far from straightforward, with over a decade passing between the initial 498 suggestions that CO might function through NF-YB/NF-YC to a complete set of work describing the NF-CO complex. Systematic testing of the ability of different NF-CCT 499 500 complex to form and bind DNA is a massive undertaking, with over 4,000 possible NF-501 CCT complexes to test. Further, while the two are clearly related, the CCT domain shows 502 key differences from the DNA-binding domain of NF-YA, and variation within CCT 503 members occurs at residues predicted to be important for DNA binding in aligned NF-YA 504 sequences. A systematic mutational analysis focusing on the differences between important residues of aligned NF-YA and CCT members could create a map of critical 505 506 DNA-binding residues, and inform the search for cognate binding sequences of various 507 NF-CCT complexes.

508

509 The DIMR system is well-suited to address these types of questions on structure and 510 function, as supported by our analysis of the NF-YA2 H183A point mutation and our comparison of NF-Y<sup>A2/B2/C3</sup> system activation in CCANT and CCAAN permutations. The 511 512 *NF-YA2* histidine residue at position 183 is thought to make sequence-specific contact 513 with the CCAAT box (Gnesutta et al. 2017), but reduced DIMR system activation is still 514 observed in the H183A point mutation. Importantly, when aligning NF-YA and CCT DNA 515 binding domains, this residue diverges between, but not among, many clades and sub-516 clades. Considering this pattern of divergence, it is intriguing to consider that this 517 particular residue might contribute to DNA binding specificity of different NF-CCT complexes. The identification of an NF-YA mutation at this residue that retains some 518 519 activity could be relevant to this observation in an evolutionary context, as a sub-optimal 520 variant is thought to facilitate further functionalization by providing a novel evolutionary 521 trajectory (i.e., without such a permissive intermediate, purifying selection cannot be 522 overcome to reach a different high-fitness state (Poelwijk et al. 2007; Anderson et al. 523 2015)). Further, the relative tolerance of wild-type system activation at CCANT compared 524 to CCAAN is interesting to consider in light of the evolutionary constraints acting on the 525 functionalization of NF-Y and NF-CCT complexes. This permissiveness could provide 526 space for evolution to act, allowing suboptimal interactions to lead to novel functions of the complex. Direct testing of these types of evolutionarily significant hypotheses is 527 technically challenging, requiring more exhaustive mutagenic approaches such as 528 529 alanine scanning mutagenesis and subsequent functional validation (Cunningham and Wells 1989). The DIMR system presented here was designed to bridge the gap in tools 530 531 necessary to more directly address these types of more nuanced questions.

532

### 533 Acknowledgements

The authors would like to thank Dr. Scott Russell for assistance with grant administration. Additionally, we thank Dr. Daniel Jones, Dr. Marielle Hoefnagels, Dr. Laura Bartley, and Andrew Willoughby for their insightful and constructive comments on the preparation of this manuscript. Finally, we thank the National Science Foundation for funding this research through MCB EAGER 1747539.

539

### 540

### 541 Methods

### 542 Yeast strain selection

The tests presented here all utilized the BY4735 (ATCC 200897) strain (*MATa* ade2 $\Delta$ ::hisG his3 $\Delta$ 200 leu2 $\Delta$ 0 met15 $\Delta$ 0 trp1 $\Delta$ 63 ura3 $\Delta$ 0), a derivation of the S288C laboratory strain. BY4735 was obtained and is available from the American Type Culture Collection (https://www.atcc.org/).

547

### 548 Yeast growth, transformation, and induction conditions

549 Yeast strains were uniformly grown at 30C. Wild type strains were grown with constant 550 agitation in YAPD liquid medium and transformed through the traditional lithium-acetate 551 based approach, as previously described. Transformants were selected after 2-3 days 552 growth on synthetic medium lacking Leucine, Tryptophan, and Uracil (L<sup>-</sup>W<sup>-</sup>Ura<sup>-</sup>).

553

For DIMR system induction, colonies were first grown to saturation in liquid L<sup>-</sup>W<sup>-</sup>Ura<sup>-</sup> media. Saturated cultures were then diluted 1:1 to a total sample volume of 500µL with fresh liquid L<sup>-</sup>W<sup>-</sup>Ura<sup>-</sup> media containing either ethanol (mock treatments) or one of a range of  $\beta$ -estradiol concentrations. These 500µL induction samples were cultured on deep, 2mL 96-well plates for 6 hours before sample collection for dual luciferase reporter testing. Plates were covered with Breathe Easy strips during induction.

560

564

561 Unless directly stated otherwise, each system test was conducted after 6 hours of 562 induction, with treatments corresponding to mock induction (ethanol only), low induction 563 (1-1.5nM  $\beta$ -estradiol), and high induction (10nM  $\beta$ -estradiol).

# 565 *Module design and construction*

The individual modules were each incorporated and carried on a different yeast shuttle vector – pRS314 and pRS315 for the effector and activator constructs, respectively, and a modified pRSII316 removing an *Ncol* recognition site within the *URA3* coding sequence through site directed mutagenesis (described below) for the reporter construct. Initial module designs were synthesized by Biomatik<sup>®</sup>, while individual effector drop-in components were synthesized by Genewiz<sup>®</sup>.

572

The activator module encodes the Z<sub>4</sub>EV artificial transcription factor, driven by the 573 constitutive ACT1 promoter (Flagfeldt et al. 2009) and flanked by the CYC1b terminator 574 575 (Curran et al. 2013). The effector module uses the Z<sub>4</sub>EV-bound promoter to drive 576 expression of the 'polycistronic' effector mRNA. Each component has been translationally fused to a 5x-glycine flexible linker (Sabourin et al. 2007) and a unique epitope tag 577 578 (component A: HA, component B: MYC, component C: FLAG). Viral T2A sites were incorporated after the HA and MYC tags to facilitate cleavage into individual effector 579 580 components. The entire cassette is flanked by the CYC1b terminator. The reporter 581 module situates the two luciferase variants in a tail-to-tail fashion. The constitutively-582 active Renilla luciferase is expressed under the ADH1 promoter (McIsaac et al. 2013) and is flanked by the ADH1 terminator (Curran et al. 2013), while the conditionally-583 584 expressed Firefly luciferase is driven by a modified yeast minimal promoter (ASN1 and 585 *CIT1* presented here, (Sundseth et al. 1997; Nevoigt et al. 2006)) that includes previously

described NF-Y bound CCAAT box sequences (Cao et al. 2014; Gnesutta et al. 2017;
Siriwardana et al. 2016) and is flanked by the CYC1b terminator. All plasmids used in this
study are freely available, and are described in Table S1.

589

# 590 Binding site and promoter architecture cloning, site directed mutagenesis

591 Drop-in Binding Site (DIBS) cloning was accomplished through annealing of 5' 592 phosphorylated oligos and subsequent ligation into the reporter module. DIBS primers 593 were designed and ordered as pairs of complementary oligos that were then annealed together by boiling for 5 minutes in annealing buffer (10mM Tris HCI, pH 8.0, 1mM EDTA, 594 595 50mM NaCl) and slowly cooling to room temperature. The reporter module was restriction 596 enzyme digested (Sacl/BamHI, New England Biolabs) and dephosphorylated (Quick Dephosphorylation Kit, New England Biolabs), then purified and concentrated through the 597 Zymo DNA Clean and Concentrator kit (PN). Ligations were set up with 3:1 ratios of 598 599 insert:backbone free ends with ~50-150ng of purified backbone.

600

601 Site directed mutagenesis was conducted on effector entry clones through New England 602 Biolabs Q5<sup>®</sup> Site Directed Mutagenesis Kit, following manufacturer's instructions. Primers 603 for mutagenesis were designed with guidance of the NEBaseChanger<sup>™</sup> tool, and 604 mutagenesis was verified through restriction enzyme digestion (where appropriate) and 605 Sanger sequencing through the University of Oklahoma's Biology Core Molecular Lab.

606

# 607 Western blots

Total yeast proteins were extracted using Y-PER<sup>™</sup> Yeast Protein Extraction Reagents 608 (Thermo Scientific, Cat no: 78991) following manufacturer's protocol. The protein 609 concentrations were measured using Pierce<sup>™</sup> BCA Protein Assay Kit (Thermo Scientific, 610 611 Cat no: 23225) following manufacturer's protocol, and was quantified on a Synergy HTX Multi-Mode Reader (BioTeK, USA) at 562 nm wavelength. The protein concentrations 612 were calculated through generation of a BSA standard curve. A total of 3 ug of protein 613 was loaded to Mini-PROTEAN TGX Stain-Free Gels (10% gel, BIO-RAD). Proteins were 614 615 transferred to standard PVDF membranes through an OWL semi-dry transfer apparatus. 616 and the presence of NF-YA2:HA, NF-YB2:MYC, or NF-YC3:FLAG was probed with high affinity anti-HA primary antibodies (Roche, catalog no. 11 867 423 001), anti-MYC 617 618 (Abcam, catalog no. ab9106), and anti-FLAG (Abcam, catalog no. F3165) followed by 619 rabbit anti-rat (Abcam, catalog no. ab6734), goat anti-rabbit (Abcam, catalog no. 620 ab205718), and goat anti-mouse (Abcam, catalog no. ab6789) HRP-conjugated 621 secondary antibodies, respectively. A Bio-Rad ChemiDoc XRS imaging system was used for visualizing the protein blot after incubations with ECL plus reagent (GE Healthcare, 622 623 catalog no. RPN2132).

624

# 625 Dual luciferase assays

Dual Luciferase assays were performed on the BioTek<sup>®</sup> Synergy<sup>™</sup> HTX multi-mode plate
reader fitted with dual reagent injectors, using the Illumination<sup>™</sup> Firefly & Renilla
Luciferase Enhanced Assay Kit (Goldbio<sup>®</sup>, cat# I-920) per manufacturer instructions, with
the following alterations: (1) we used only 5µL of each culture, (2) samples were not spun
down and/or washed, and (3) we used half-volume injections of both luciferase buffers.
Our initial system tuning followed the manufacturer instructions more strictly, and we

632 could determine either no functional difference or minor improvements between633 manufacturer instructions and our modified instructions (data not shown).

634

# 635 Statistical approaches

636 Statistics were calculated through Graphpad Prism. One-way ANOVA on relative 637 luminescence values was used for comparisons of DIMR system activation to mock 638 levels. When denoting significance in fold change-based metrics, the relative 639 luminescence data was used for statistical analysis.

640

# 641 *Figure and model construction*

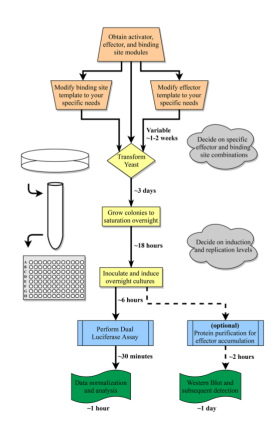
Individual graphs were generated through Graphpad Prism, while full figures were
 composed in Adobe Photoshop CC2018. The model in Figure 1A was constructed in
 Inkscape, while the flowchart in Figure 1B was constructed through draw.io v10.4.5
 (<u>https://draw.io/</u>).

646



# Figures a

В



# **Figures and Tables**

Figure 1. Model illustrating fundamental concepts and practical application of the DIMR system. (A) Circled numbers follow the flow of DIMR system activation: (1)  $\beta$  -estradiol activates the Z<sub>4</sub>EV artificial transcription factor by competing with and replacing Hsp90, allowing for nuclear accumulation of Z<sub>4</sub>EV; (2) Z<sub>4</sub>EV induces transcription of the effector cassette; (3) translation of the effector cassette mRNA produces individual effector components 2A-mediated through translational cleavage; (4) effectors translocate to the nucleus and form functional complexes; (5) transcriptional activation and is measured as the ratio of conditionallyexpressed firefly luciferase activity and constitutively-expressed Renilla luciferase activities from the reporter module. (B) Practical application of the DIMR system, including approximate timelines for design and implementation of an individual experiment.

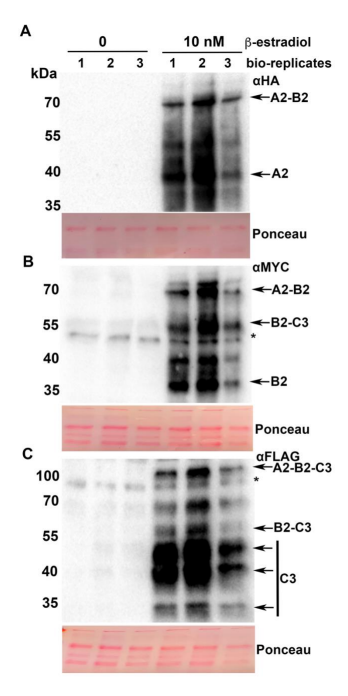
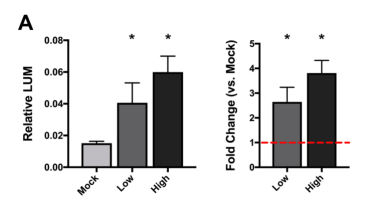


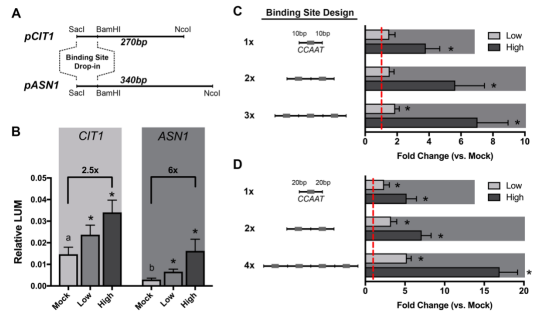
Figure 2. Induction and cleavage of NF-YA2, NF-YB2 and NF-YC3 effectors in yeast. Western blot analysis of (A) NF-YA2:HA, (B) NF-YB2:MYC, and (C) NF-YC3:FLAG accumulation in response to 10 nM βestradiol induction for 6 hours. Three independent biological replicates, corresponding to lanes 1, 2, and 3, were tested for protein accumulation using anti-HA, anti-Myc and anti-Flag antibodies. Ponceau S staining of the PVDF membrane prior to transfer was used to test loading control. The experiment was repeated three times with similar result.



688

Figure 3. DIMR system test of NF-YA2/B2/C3 on FT CCAAT box before system optimization. (A) Relative luminescence of mock, low, and high concentration β-estradiol (0, 1uM, and 10uM, respectively) treated samples, and fold change levels (compared to mock) of low and high treated samples. Each condition includes at least 6 biological replicates. Asterisks indicate a significant increase between mock and treated samples, as determined through two-way ANOVA (p < 0.01). Error bars represent 95% confidence intervals.

697



### 698

Figure 4.

Parameter refinement of the DIMR system to optimize reporter dynamic range. (A) 699 Model of two tested promoter architectures, derived from the CIT1 and ASN1 native yeast 700 promoters. Binding site permutations were cloned into the reporter module through 701 702 flanking Sacl / BamHI restriction sites. (B) Effects of different promoter architectures of CIT1 and ASN1 on DIMR system activation of NF-Y<sup>A2/B2/C3</sup> on FT CCAAT box. (C. D) 703 Effects of binding site multimerization and altered spacing between binding sites. In (C), 704 each binding site footprint was 25bp, while in (D), each footprint was 45bp. Both sets 705 were testing binding of NF-Y<sup>A2/B2/C3</sup> on FT CCAAT box permutations in the ASN1-based 706 707 promoter architecture. Error bars represent 95% confidence intervals. Asterisks above individual bars indicate statistical significance over matched mock-treated samples, as 708 determined through two-way ANOVA (p < 0.01). Asterisks above brackets connecting two 709 710 bars indicate significance between the two conditions, determined similarly. Letters above mock-induced bars indicate significance groups between the different promoter 711 architectures, also determined through two-way ANOVA (p < 0.01). 712

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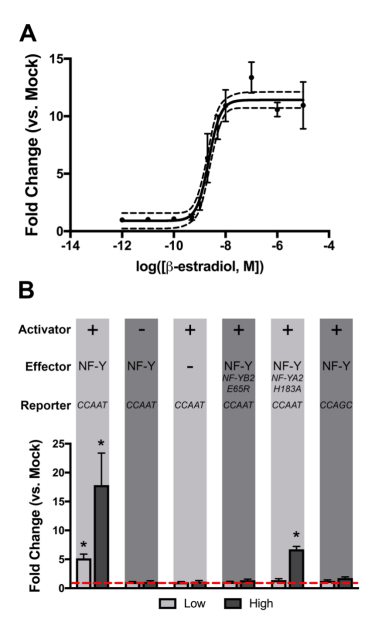
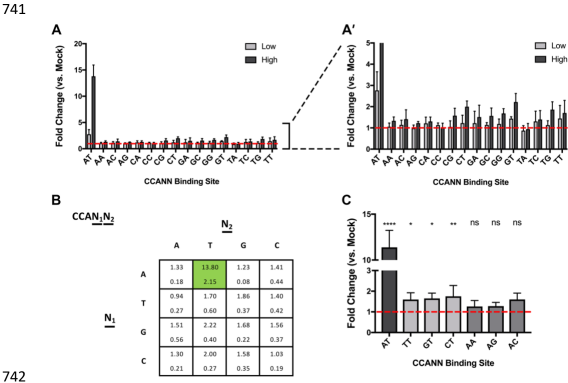
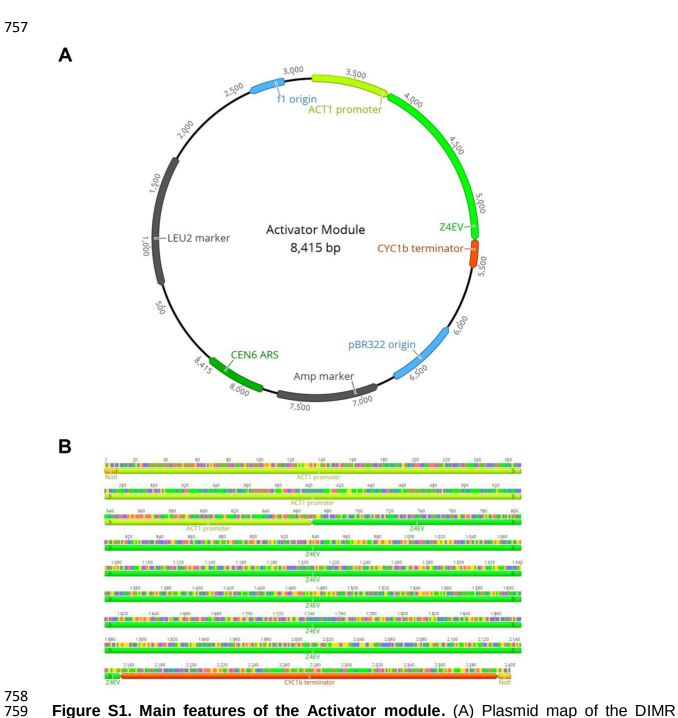


Figure 5. Functional assessment of DIMR system activation. (A) Dose response curve of NF-Y<sup>A2/B2/C3</sup> on the 4x FT CCAAT box in the ASN1 promoter architecture. The solid line fits the dataset to a 4parameter logistic regression, with 95% confidence intervals plotted with dashed lines ( $R^2 = 0.942$ ). (B) DIMR system requirements for NF-Y/CCAAT-mediated activation. Module components are generally described above each condition: +, present; -, absent; NF-Y, NF-Y<sup>A2/B2/C3</sup>. Error bars represent 95% confidence intervals. Asterisks individual bars indicate above statistical significance over matched mock-treated samples, as determined through two-way ANOVA (p < 0.01); all bars lacking asterisks were found to be nonsignificant compared to matched mock-treated samples.

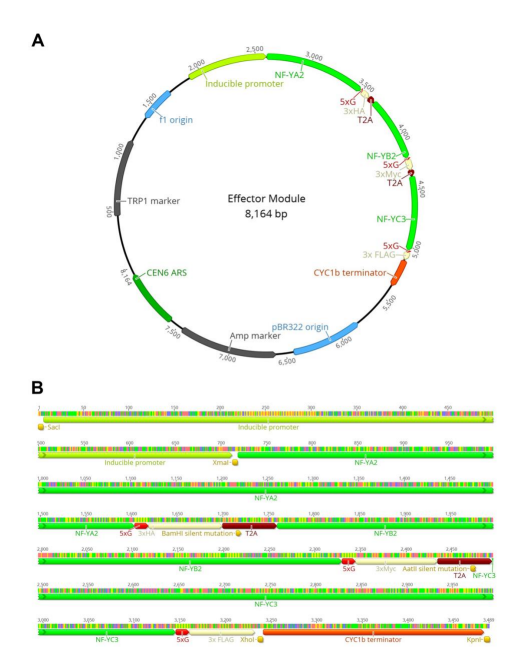




744 Figure 6. DIMR-based validation and investigation of NF-YA2/B2/C3 CCAAT binding. (A) System activation of NF-YA2/B2/C3 on the FT CCAAT box, with all possible 745 binding site permutations corresponding to CCANN. Error bars represent 95% confidence 746 intervals. Asterisks above individual bars indicate statistical significance over matched 747 mock-treated samples, as determined through two-way ANOVA (p < 0.01); all bars 748 lacking asterisks were found to be nonsignificant compared to matched mock-treated 749 samples. (B) Graphical representation of the data presented in panel A. Each cell 750 751 corresponds to a specific permutation of CCANN, and contains the average fold change 752 (top) and standard deviation (bottom). (C) A targeted comparison of CCANT and CCAAN binding site permutations. Asterisks indicate significance compared to mock induction as 753 754 determined by two-way ANOVA with Sidak's multiple comparison adjustment (\*\*\*\*, p < 755 0.001; \*\*, p < 0.01; \*, p < 0.05). 756



- 759 **Figure S1. Main features of the Activator module.** (A) Plasmid map of the DIMR 760 Activator (pDIMR\_A###) module showing important features and their orientation. (B)
- 761 Zoomed-in view of the relevant Activator components.
- 762
- 763

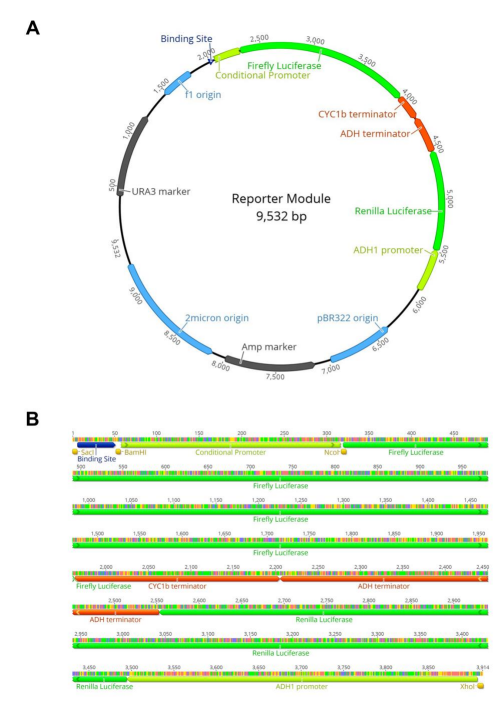


764

**Figure S2. Main features of the Effector module.** (A) Plasmid map of the DIMR Effector (pDIMR\_E###) module showing important features and their orientation. (B) Zoomed-in view of the relevant Effector components, including important restriction enzymes used for swapping new components into or out of the system.

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**Figure S3. Main features of the Reporter module.** (A) Plasmid map of the DIMR Reporter (pDIMR\_R###) module showing important features and their orientation. (B) Zoomed-in view of the relevant Reporter components, including important restriction enzymes used for Drop-In Binding Site (DIBS) cloning.

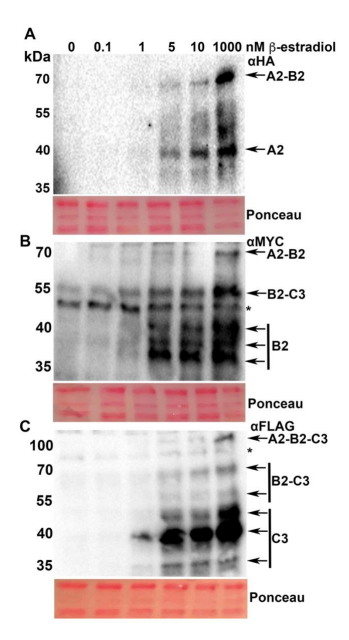


Figure S4. Induction gradient of NF-YA2, NF-YB2 and NF-YC3 effectors in yeast. Western blot analysis of (A) NF-YA2:HA, (B) NF-YB2:MYC, and (C) NF-YC3:FLAG accumulation in response to varying levels of  $\beta$ estradiol induction for 6 hours. A single biological replicate was tested for protein accumulation using anti-HA, anti-Myc and anti-Flag antibodies. Ponceau S staining of the PVDF membrane prior to transfer was used to test loading control. The experiment was repeated twice with similar results.

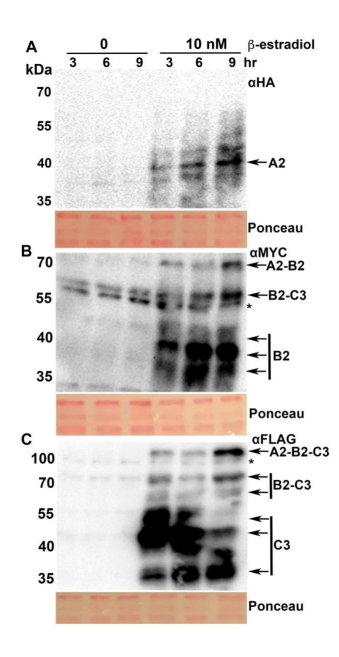
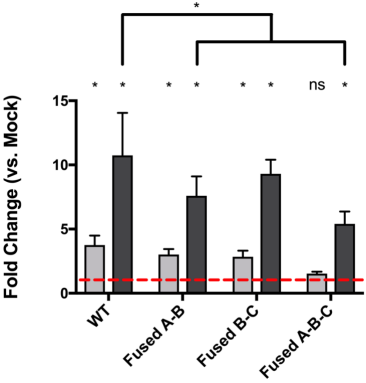
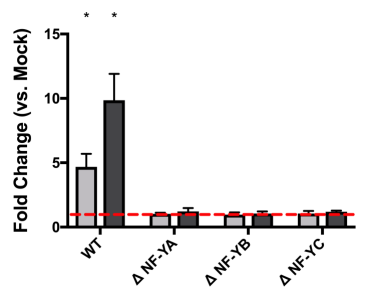


Figure S5. Induction time-course of NF-YA2, NF-YB2 and NF-YC3 effectors in yeast. Western blot analysis of (A) NF-YA2:HA, (B) NF-YB2:MYC, and (C) NF-YC3:FLAG accumulation in response to 10nMf ßestradiol induction for 3, 6, and 9 hours. A single biological replicate was tested for protein accumulation using anti-HA, anti-Myc and anti-Flag antibodies. Ponceau S staining of the PVDF membrane prior to transfer was used to test loading control. The experiment was repeated twice with similar results.



808

Figure S6. Activation of the DIMR system in effector cassettes with broken T2A 809 sites. Point mutations were generated in the T2A sites between NF-YA / NF-YB, between 810 NF-YB / NF-YC, and in both T2A sites simultaneously. T2A sites were broken through a 811 proline to alanine mutation near the end of each T2A site, as previously described 812 (Beekwilder et al. 2014). Error bars represent 95% confidence intervals. Asterisks above 813 individual bars indicate statistical significance over matched mock-treated samples, as 814 determined through two-way ANOVA (p < 0.01). Brackets with asterisks indicate 815 significance between induced samples. ns, not significantly different. 816 817



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Figure S7. Activation of the DIMR system in the absence of individual NF-Y effectors. NF- $^{YA2/B2/C3}$  Effectors with empty components for NF-YA ( $\Delta$ NF-YA), NF-YB ( $\Delta$ NF-YB), or NF-YC ( $\Delta$ NF-YC), were tested for ability to activate the 4x *FT CCAAT* box in the *ASN1*-based promoter. Error bars represent 95% confidence intervals. Asterisks above individual bars indicate statistical significance over matched mock-treated samples, as determined through two-way ANOVA (p < 0.01); all bars lacking asterisks were found to be nonsignificant compared to matched mock-treated samples.

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