# **Cooperative action of separate interaction domains promotes high-affinity DNA binding of** *Arabidopsis thaliana* ARF transcription factors

Mattia Fontana<sup>a,b</sup>, Mark Roosjen<sup>b</sup>, Isidro Crespo García<sup>c</sup>, Willy van den Berg<sup>b</sup>, Marc Malfois<sup>c</sup>, Roeland Boer<sup>c</sup>, Dolf Weijers<sup>b,\*</sup>, and Johannes Hohlbein<sup>a,d,\*</sup>

<sup>a</sup>Laboratory of Biophysics, Wageningen University and Research, Stippeneng 4, 6708 WE Wageningen, The Netherlands; <sup>b</sup>Laboratory of Biochemistry, Wageningen University and Research, Stippeneng 4, 6708 WE Wageningen, The Netherlands; <sup>c</sup>ALBA synchrotron Light Source, Carrer de la Llum 2-26, Cerdanyola del Vallès, 08290, Barcelona, Spain; <sup>d</sup>Microspectroscopy Research Facility, Wageningen University and Research, Stippeneng 4, 6708 WE Wageningen, The Netherlands; <sup>\*</sup>Correspondence: dolf.weijers@wur.nl, johannes.hohlbein@wur.nl

- The signaling molecule auxin is pivotal in coordinating many growth and development processes in plants mainly through the modulation of gene expression. The transcriptional response to auxin is mediated by the family of auxin response factors (ARF). Monomers of this family recognize a DNA motif (TGTC[TC]/[GG]) called the auxin-response element (AuxRE). ARFs can homodimerize through their DNA binding domains (DBD) thereby enabling cooperative binding for a bipartite inverted AuxRE (IR7). In addition to the DBD, most ARFs contain a C-terminal Phox and Bem1p (PB1) domain both capable of homotypic interactions, and mediating interactions with Aux/IAA repressors. Given the dual role of the PB1 domain, and the ability of both DBD and PB1 domain to mediate dimerization, a key question is how each of these domains contributes to conferring DNA-binding specificity and affinity. So far, ARF-ARF and ARF-DNA interactions have mostly been approached using qualitative methods that do not provide a quantitative and dynamic view on the binding equilibria. Here, we utilize a DNA binding assay based on single-molecule Förster resonance energy transfer (smFRET) to study the affinity and kinetics of the interaction of several *Arabidopsis thaliana* ARFs with an IR7 AuxRE. We show that both DBD and PB1 domains of AtARF2 contribute toward DNA binding, and we identify ARF dimer stability as a key parameter in defining affinity and kinetics seen for the DBDs of different AtARFs. Lastly, we derived an analytical solution for a four-state cyclic model that explains both the kinetics and the affinity of the interaction between AtARF2 and IR7. Our work demonstrates that the affinity of ARFs towards composite DNA response elements can be tuned by small changes of their dimerization equilibrium suggesting that this effect has major
- <sup>16</sup> implications for ARF-mediated transcriptional activity.

#### Introduction

2

6

7

8

9

10

11

12

13

14

15

The plant signaling molecule auxin plays a major role in many 2 cellular and developmental processes. Auxin triggers both non-3 transcriptional and transcriptional responses with the latter being controlled by the nuclear auxin pathway (1-5). This 5 pathway involves three main players: the transcription factor 6 ARF, its repressor Aux/IAA and the ubiquitin ligase complex SCF<sup>TIR1/AFB</sup>. Binding of auxin to TIR1/AFB enables the recognition and ubiquitination of Aux/IAA. Upon degradation 9 of Aux/IAA, ARF is able to modulate the expression of its 10 downstream target genes. 11

The interaction between ARFs and Aux/IAAs is mediated by
the C-terminal Phox and Bem1p (PB1) domain present in both
proteins. The PB1 domain features two oppositely charged
surfaces (type I/II or AB [acid basic] PB1 domain) that can

<sup>16</sup> undergo head to tail oligomerization(6–9). Remarkably, this

17 structural characteristic enables scenarios of homo- and hetero-

oligomerization among and between Aux/IAAs and ARFs. In 18 addition to the PB1 domain, ARFs consist of two other do-19 mains, the Middle Region (MR) and the N-terminal DNA 20 Binding Domain (DBD). The MR domain is predicted to be 21 intrinsically disordered(5) and its amino acid sequence dif-22 fers between the three phylogenetically separated ARF clades 23 (A, B and C)(10, 11). When tested for their effect on gene 24 expression, some ARFs activate auxin-responsive genes while 25 other repress them. In general, class A ARFs (e.g., A. thaliana 26 ARF5) act as activators while class B (e.g., A. thaliana ARF1 27 and 2) and C ARFs act as repressor(10). The DBD domain 28 physically interacts with its DNA response element called 29 AuxRE (auxin-responsive element)(12). This cis-regulatory 30 element was first identified in promoters of an auxin-responsive 31 genes in pea(13) and solution solution solution (14, 15) and was found to be 32 essential for their auxin inducibility. The canonical TGTCTC 33 recognition sequence was later shown to be bound by differ-34

ent members of the ARF family (12, 16). More recently, the 35 TGTCGG recognition sequence was found to have an even 36 higher affinity for ARFs in vitro(17-19) and was used to cre-37 ate an enhanced artificial auxin response reporter (20). Single 38 AuxREs are bound by single ARF monomers but ARF DBDs 39 can dimerize in solution and bind cooperatively to composite 40 response elements bearing two AuxRE in inverted configura-41 tion (IR)(17); moreover, ARF dimerization through its DBD 42 is necessary for ARF function in vivo(17, 21). Interestingly, 43 the PB1 domain seems to have diverse effects on different class 44 A ARFs as its deletion in *M. polymorpha* ARF1 generates a 45 loss-of-function mutant(22) whereas in A. thaliana ARF5 the 46 mutant maintains its function and is hyperactive (23). The 47 effect of the homotypic interaction of ARF PB1 domains of 48 another class A ARF, AtARF19, has been studied using syn-49 thetic auxin response circuits in yeast, showing that mutating 50 either the positive or the negative side of the PB1 domain 51 reduces its ability to promote transcription(24). 52 Although many structures and relevant interactions among 53 the various components of the auxin nuclear pathway have 54 been identified, quantitative data on the affinity and kinetics 55 of these interactions have remained scarce. In particular, the 56 effects of the dimer/monomer equilibrium on the interaction 57 between ARFs and between ARFs and AuxREs, or the effect 58 of mutations on the DBD and PB1 domains on ARF dimer-59 ization have not yet been systematically studied, obscuring 60 which interactions might be relevant in a cellular context. Par-61 ticularly, it is unclear if and how both interaction domains 62 (DBD and PB1) contribute to DNA binding, and what their 63

relative contributions are. Furthermore, it is unclear whether
oligomerization of ARF PB1 domains contributes to DNA
binding.

67 Here, we employed a DNA-binding assay based on singlemolecule Förster resonance energy transfer (smFRET) to 68 quantitatively assess the binding affinities between different 69 A. thaliana ARFs and a response element composed of two 70 AuxREs in an inverted repeat configuration with a spacing of 71 7 base pairs (IR7). We found that, while the DNA-binding 72 domain alone can bind DNA, the presence of the PB1 domain 73 increases the affinity of AtARF2 towards the tested compos-74 ite response element. In fact, this effect can be ascribed to 75 increased stability of the dimer, whereas AtARF2 oligomeriza-76 tion has no sizable effect. We introduce a general four-state 77 cyclic model to quantify the mechanisms of ARF interaction 78 with the bipartite DNA response element; the simultaneous 79 analysis of the equilibrium and kinetics data using this model 80 revealed that the increase in affinity can be completely pinned 81

to the shift in the dimer-monomer equilibrium. Further analysis of variants of AtARF5-DBD and other AtARF-DBDs showed that changes in dimer stability generated by changes in the DBD domain displays the same pattern on the kinetics as the ones generated by changes in the PB1 domain, highlighting that stable protein dimers ensure high-affinity DNA binding, no matter the source of their stability.

89

#### Materials and Methods

Protein expression and purification. Protein expression and pu-90 rification was carried out as described previously(17). Briefly, 91 the genomic regions corresponding to the DNA binding do-92 main (DBD) of Arabidopsis thaliana ARF1, ARF2, ARF5 and 93 full-length ARF2 were amplified and cloned in an modified 94 expression vector pTWIN1 (New England Biolabs) to gen-95 erate fusions with the Chitin Binding Domain (CBD) and 96 Intein. ARF-CBD fusion proteins were expressed in E. coli 97 strain Rosetta DE3 (Novagen). Cells were inoculated in Difco 98 Terrific Broth (BD), supplemented with ampicillin and grown 99 to an  $OD_{600}$  of 0.5 to 0.7, protein expression was induced by 100 adding IPTG and the temperature was switched from 37 °C to 101 20 °C; the growth was continued for 20 h. Cells were harvested 102 by centrifugation and resuspended in 50 mL extraction buffer 103 (20 mM Tris, 500 mM NaCl, 1 mM EDTA, 0.1 % NP-40 and 104 2 mM MgCl<sub>2</sub>, pH 7.8, 10 mg of DNase and 0.2 mM PMSF). 105 Cells were then lysed by passing the suspension twice through 106 a French Pressure cell press and cell-free extract was generated 107 by centrifugation. The supernatant was loaded onto a chitin 108 column (New England Biolabs) and washed with 10 column 109 volumes washing buffer (20 mM Tris, 500 mM NaCl, pH 7.8) 110 using an AKTA explorer 100 (GE Healthcare). ARF-DBD 111 proteins were eluted by 1 h incubation with 40 mM DTT in 112 washing buffer. Proteins were concentrated using Amicon 113 ultra-15 10K spin filters, and next passed over a Superdex 114 200PG size-exclusion chromatography column. ARF-DBD 115 proteins were eluted using washing buffer with 1 mM DTT, 116 concentrated using Amicon ultra-15 10K spin filters and stored 117 until use at -80 °C. 118

DNA constructs. Single strand DNA oligonucleotides were or-119 dered from Eurogentec. Each strand contained a 5-C6-amino-120 dT modification at the desired position for labelling. Some 121 of the strands were purchased biotinylated at their 5'- end to 122 allow for surface immobilization using a Neutravidin bridge. 123 Strands were labelled with the desired dye (Cy3 or ATTO 124 647N NHS-ester) following a modified version of the protocol 125 provided by the dye manufacturer and purified using polyacry-126 lamide gel electrophoresis (20% Acrylamide). DNA constructs 127 were annealed by heating complementary single strands to 128 95 °C in annealing buffer (250 mM NaCl, 10 mM Tris HCl pH 8, 129

1 mM EDTA) followed by cooling down to room temperature
overnight.

Single-molecule FRET. Imaging was carried out on a home-132 built TIRF microscope, described previously(25). The mea-133 surements were performed using alternating-laser excitation 134 (ALEX)(26); in this excitation scheme, each frame during 135 which the donor is excited is followed by a frame in which the 136 acceptor is directly excited. The emission of the fluorophores 137 is spectrally divided into two different detection channels on 138 the emCCD camera sensor (Andor iXon 897 Ultra). This 139 approach creates four photon streams, three of which are rel-140 evant; (1) donor emission after donor excitation (DD), (2) 141 acceptor emission after donor excitation (DA, arising from142 FRET) and (3) acceptor emission after acceptor excitation 143 (AA). The three photon streams can be used to calculate the 144 raw FRET efficiency  $(E^* = DA/(DD + DA))$  and stoichiome-145 try (S = (DD + DA)/(DD + DA + AA))(27). E<sup>\*</sup> contains the 146 information about the relative distance of the two fluorophores 147 whereas S contains information about the photophysical state 148 of a given molecule (allowing to filter out molecules missing an 149 active donor or an active acceptor). The camera acquisition 150 time and the excitation time were set to 250 ms per frame; 151 laser powers were set to 3 mW for green ( $\lambda = 561$  nm) and 152  $0.5 \,\mathrm{mW}$  for red ( $\lambda = 638 \,\mathrm{nm}$ ) lasers. The PBS-based imaging 153 buffer (pH 7.4) contained 137 mM NaCl, 2.7 mM KCl, 10 mM 154 phosphate, and 1 mM Trolox, 1% gloxy and 1% glucose to 155 decrease the rate of photobleaching (28, 29). 156

Single-molecule titration experiments. Labelled dsDNA oligos 157 were immobilized on a PEGylated glass coverslip as described 158 previously(30). In particular, the PEGylation was carried 159 out inside the wells of silicone gaskets placed on the coverslip 160 (Grace Bio-labs). Each protein titration was performed using 161 a single well, washing it between data points with 600 µL of 1x 162 PBS buffer. The final washing step consisted of three washings 163 separated by 15 minutes. Typically, each data point consisted 164 of four movies (1000 frames each). 165

Binding isotherms analysis. The fit of the FRET efficiency 166 distribution with the two Gaussian distributions pertaining 167 to the free and ARF-bound DNA populations returns an un-168 corrected fraction of ARF-bound DNA for each tested protein 169 concentration i:  $F_{\rm B}^{u}(i)$ . Even when no protein is added, the 170 double Gaussian fit returns an uncorrected fraction bound 171  $F_{\rm B}^u(0) > 0$  (typically  $\approx 0.1$ ). This value is an indication of the 172 error connected to the two-population fit and can be used to 173 renormalize the entire titration under the assumption that, 174 in case the DNA would be completely bound by ARF, the 175 expected uncorrected free fraction would have the same value 176

 $(F_{\rm B}^u(0) = F_{\rm F}^{\rm u}(\infty))$ . Then, the corrected fraction bound for 177 each data point can be calculated as 178

$$F_{\rm B}^c(i) = \frac{F_{\rm B}^u(i) - F_{\rm B}^u(0)}{1 - 2F_{\rm B}^u(0)}.$$
 [1] 179

The corrected fraction bound (henceforth  $F_{\rm B}$ ) can be fitted 180 with the appropriate mathematical model for the interaction. 181

Time traces analysis. First, the time traces from individual 182 DNA molecules were filtered to remove sections in which either 183 the donor or the acceptor were inactive due to fluorophore 184 bleaching or blinking. Each molecule was allowed to take values 185 of  $E^*$  and S outside the thresholds (typically 0 to 0.85 for  $E^*$ 186 and 0.5 to 0.9 for S) for a maximum of three consecutive data 187 points; longer stays outside the thresholding range resulted in 188 the trace being interrupted. In case the molecule reentered the 189 allowed range for  $E^*$  and S, the data points were saved as a new 190 trace. The minimum length of traces was set to 50 data points 191 (100 frames). The filtered time traces were then loaded in the 192 software package ebFRET to perform and empirical Bayesian 193 Hidden Markov Modelling(31). The analysis was performed 194 assuming two states, with two restarts and a convergence 195 threshold of  $10^{-6}$ . The results of the analysis were exported as 196 '.csv' text files and the transition matrix was used to calculate 197  $k_{\rm on}$  and  $k_{\rm off}$ . The  $K_{\rm d}$  was calculated as  $k_{\rm off}/k_{\rm on}$ . 198

SAXS. Different concentrations of AtARF1 and AtARF5 rang-199 ing from  $17 \,\mu\text{M}$  to  $170 \,\mu\text{M}$  ( $0.7 \,\text{mg}\,\text{mL}^{-1}$  to  $7 \,\text{mg}\,\text{mL}^{-1}$ ) were 200 tested to record ARF dimerization depending on protein con-201 centration. All the samples were prepared in a final buffer 202 consisting of 20 mM Tris-HCl pH 8.0, 150 mM NaCl, 1 mM 203 DTT. SAXS data was collected at NCD-SWEET beamline 204 (BL11, ALBA Synchrotron, Barcelona)(32, 33). The buffer was 205 collected for subtraction of protein samples. Measurements 206 were carried out at 293 K in a quartz capillary of 1.5 mm outer 207 diameter and 0.01 mm wall thickness. The data (20 frames 208 with an exposure time of 0.5 sec/frame) was recorded using 209 a Pilatus 1M detector (Dectris, Switzerland) at a sample-210 detector distance of  $2.56 \,\mathrm{m}$  and a wavelength of  $1.0 \,\mathrm{\AA}$ . 211

Buffer subtraction and extrapolation to infinite dilution were 212 performed by using the program package primus/qt from the 213 ATSAS 2.8.4 software suite(34). The forward scattering I(0)214 and the radius of gyration  $(R_g)$  were evaluated by the Guinier 215 approximation, and the maximum distance  $D_{\text{max}}$  of the par-216 ticle was also computed from the entire scattering patterns 217 with AutoGNOM. The excluded volume  $V_{\rm p}$  of the particle 218 was computed from the Porod invariant. The scattering from 219 the crystallographic models was computed with CRYSOL(35). 220 The volume fractions of the oligomers were determined with 221 OLIGOMER(36), using as probe the available PDB structures. 222

### 223 Results

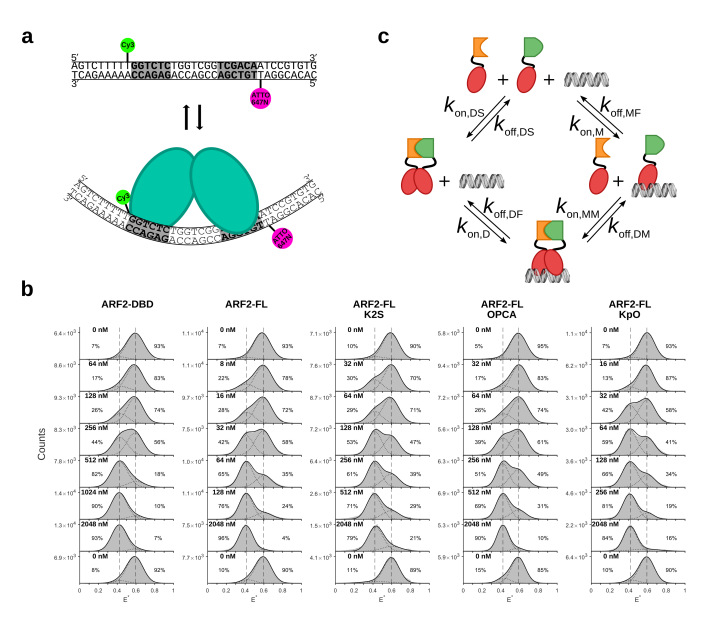
The AtARF2 PB1 domain promotes DNA binding through sta-224 bilization of the AtARF2 dimer. A ChIP experiment on Ara-225 bidopsis Thaliana ARF19 expressed in yeast(24) suggested 226 that PB1 mutations affect DNA-binding affinity. However, 227 other yeast proteins may confound the differences observed in 228 this assay. We therefore focused on the minimal system of the 229 purified ARF protein and its DNA target in vitro and asked 230 whether the interactions between the PB1 domains modulate 231 affinity of ARFs towards a composite AuxRE. We designed 232 smFRET experiments in which the binding of ARFs to a small 233 doubly labelled dsDNA oligo containing two AuxREs in an 234 inverted configuration spaced by seven base pairs (IR7) leads 235 to a decrease of FRET efficiency (Fig. 1a, Supporting Informa-236 tion Fig. S1). We then performed titrations with increasing 237 concentrations of different ARF2 variants (Fig. 1b). The ARF 238 DBD alone is sufficient for cooperative DNA binding to the 239 IR7 element(17); to explore the influence of regions outside 240 the DBD, we first compared binding of AtARF2-DBD and full-241 length AtARF2 (FL; DBD-MR-PB1). The FRET efficiency 242 distributions of the DNA sensor show the free DNA population 243 (93% occupancy) centered at  $E^* = 0.59$  in absence of ARF 244 proteins. With increasing concentration of AtARF2-DBD 245 or AtARF2-FL, the low FRET population representing the 246 ARF-bound DNA fraction (centered at  $E^* = 0.42$ ) becomes 247 progressively more populated. 248

To demonstrate that the shift seen during the titration is 249 generated by specific binding of ARF to the DNA and that 250 the binding is reversible, we performed a washing step at 251 the end of each titration that reverted the FRET efficiency 252 253 distributions to the ones seen in absence of ARF. When comparing AtARF2-DBD and AtARF2-FL, the FRET efficiency 254 distributions clearly show the effect of the PB1 domain on 255 the interaction between ARF2 and the IR7 response element; 256 the shift between the response element being mostly free to 257 mostly bound occurs at a protein concentration almost one 258 order of magnitude lower with the full-length protein (256-259 512 nM ARF2-DBD vs 32-64 nM ARF2-FL). This finding is 260 consistent with the PB1 domain promoting DNA binding. 261 However, the full-length ARF protein also contains an ex-262 tended MR region. To address the role of the PB1 domain 263 specifically, we engineered mutations in the PB1 domain that 264 prevent head-to tail interaction: AtARF2-FL K2S (K737S) 265 and AtARF2-FL OPCA (D797-8S) carry mutations of amino 266 acids on the positive (K2S) and negative (OPCA) side of 267 the PB1 domain respectively, both of which were shown to 268 impair the interaction between PB1 domains(6). In both mu-269 tant ARF2 versions, we see equal percentages of DNA bound 270 and free at concentrations close to the ones of ARF2-DBD 271

(64-128 nM K2S and 128-256 nM OPCA). Thus, PB1 domain 272 interactions contribute to efficient DNA binding. PB1 domains 273 could potentially oligomerize through head-to-tail interactions. 274 To address if oligomerization stabilizes DNA binding, we com-275 pared a mixture of ARF2-FL K2S and ARF2-FL-OPCA in a 276 1:1 ratio (henceforth ARF2 FL KpO) with the wildtype version 277 (ARF2-FL). While the latter allows for oligomerization, the 278 former should only be able to dimerize. Both ARF2 FL KpO 279 and ARF2-FL show high affinity towards the IR7 (32-64 nM). 280 Taken together, these observations indicate that the PB1 do-281 main stabilizes the binding of ARF2 towards an IR7 response 282 element through stabilization of the protein dimer. 283

A four-state cyclic model for describing ARF-DNA interactions. One 284 way of quantifying the effect of the PB1 domain on the affinity 285 between ARF and its RE is to fit the increase of the fraction 286 of DNA bound to ARF as the concentration of protein in-287 creases, with a single apparent  $K_d^*$ . This approach can reliably 288 summarize the strength of the interaction providing a single 289 numerical value that exemplifies at which endogenous protein 290 concentration the interaction becomes relevant(22, 37), but 291 fails to properly describe the underlying system, which results 292 in a lack of predictive power. 293

The interaction between a protein that can dimerize and a 294 bipartite response element on the DNA can be described using 295 a four-state cyclic model. This model allows for monomers 296 or dimers to bind the DNA, for monomers and dimers to 297 exist in solution and for dimers to form or dissociate both 298 in solution or on the DNA. Figure 1c depicts the model for 299 ARF2-FL KpO; the two ARF2-FL variants are characterized 300 by the same DBD (red) and MR (black) but are mutated on 301 the two opposite surfaces of the PB1 domain (K2S and OPCA 302 mutants in orange and green respectively); this allows for the 303 formation of PB1 domain dimers but hinders the formation 304 of oligomers. The system is defined by four  $k_{on}s$  and four 305  $k_{\text{off}}$ s or alternatively by four equilibrium constants (Ks). The 306 presence of the PB1 domain should not change the contacting 307 interface between the DBDs and the DNA; hence, the  $k_{\text{off}}$  of 308 the dimer from the DNA  $(k_{\text{off,DF}})$  should have the same value 309 for all AtARF2-variants; the same holds for the  $k_{\text{off}}$  of the 310 monomer from the DNA  $(k_{\text{off,MF}})$ . Moreover, the PB1 domain 311 has limited influence on the  $k_{on}s$  of the system which stay 312 diffusion limited. The only constants that are expected to be 313 influenced by changes in the stability of the dimer induced by 314 the PB1 domain are the ones associated with the separation 315 of two monomers: the equilibrium dissociation constant of 316 the dimer in solution  $(K_{\rm I} = k_{\rm off,DS}/k_{\rm on,DS})$  and  $k_{\rm off,DM}$ , which 317 encompasses the stability of the dimer on the DNA. As the 318 model is a closed cycle, microscopic reversibility (38) implies 319 that only one of these two parameters is a free parameter; then, 320



**Fig. 1.** SmFRET binding assay and cyclic four-state model. (a) Schematic representation of the DNA-binding assay used to evaluate ARF binding; the dsDNA is labelled with Cy3 and Atto647N on the opposite sides of the response element (RE). Upon protein binding, the increased distance between the dyes leads to a decrease in FRET efficiency. (b) Titrations of the dsDNA with several ARF variants. The dsDNA alone has a FRET efficiency  $E^* = 0.59$ ; as the protein concentration increases the population of bound DNA (centered at  $E^* = 0.42$ ) increases until all the DNA is bound (saturating condition). A washing step suffices to reset the system proving that the bound population is generated by specific and reversible binding of ARF. Vertical dashed lines are added for visual guidance. (c) Schematic representation of the four-state cyclic model for ARF2-FL KpO-IR7. Note that the dsDNA containing the DNA response element can be found in three states: free (F), bound to a monomer (M) and bound to a dimer (D). The two ARF2 full length variants (K2S and OPCA) have the same DBD (in red) and MR domains (in black) but their PB1 domain (in orange and green, respectively) carry a mutation on either one of the two different surfaces; this hinders oligomerization but allows dimerization. The binding of a dimer to a bipartite response element can occur either through two successive binding events of a monomer or through direct binding of a dimer formed in solution. The dissociation can occur either by the loss of a monomer form the DNA or by direct dissociation of a dimer from the DNA.

the different variants tested are characterized in this model by the single variant-specific parameter  $K_{\rm I}$ , which encompasses the stability of the ARF dimer. Fitting experimental results to determine  $K_{\rm I}$  and the other relevant shared kinetic constants provides a deeper understanding of the system allowing to

 $_{\rm 326}$   $\,$  make prediction for other ARF-AuxRE interactions.

ARF-IR7 interaction follows a four-state cyclic interaction 327 mechanism. To obtain the kinetics of AtARF2/IR7 interaction, 328 we analyzed relevant datapoints of the titrations for the series 329 of variants using ebFRET(31), a MATLAB suite for Hidden 330 Markov Model (HMM) analysis of single-molecule time traces 331 332 (see Materials and Methods). As the DNA oligonucleotides are immobilized, their interaction with the proteins in solution can 333 be monitored for several minutes (250s in our experiments). 334 The analyzed FRET traces returned the most probable hidden 335 state sequence for each trace (Fig. 2a) according to the set 336 of kinetic parameters that best explains the transitions and 337 states seen in the entire dataset. 338

To facilitate the identification of transitions by ebFRET, we 339 selected three concentrations of ARF  $([ARF]_T)$  that returned 340 close to equal populations of ARF-bound and free DNA for 341 each AtARF2 variant. Each AtARF2 variant was tested in at 342 least three independent titrations; each datapoint of each titra-343 tion was analyzed independently using ebFRET and returned 344 a value for  $k_{\rm on}$ , a value for  $k_{\rm off}$  and, from their ratio, a value 345 for  $K_{\rm d}$  (Fig. 2b, colored markers). The observed  $k_{\rm on}$  show a 346 trend in which ARF variants with higher affinity show faster 347 association. On the other hand, the  $k_{\text{off}}$  show similar values for 348 all the FL variants whilst AtARF2-DBD has a faster  $k_{\text{off}}$ . The 349 resulting  $K_{ds}$  show the expected trend, with AtARF2-DBD 350 having the lowest affinity, AtARF2-FL KpO and wt showing 351 the tightest binding and AtARF2-FL K2S and OPCA having 352 an affinity in between these. The trends seen in the  $k_{on}$  and 353  $k_{\rm off}$  suggest that the analysis of the kinetics using HMM is 354 capturing the interaction between the ARF dimer and the 355 DNA and that the interaction between the ARF monomer 356 and the DNA occurs on a timescale shorter than the 500 ms 357 acquisition time used in our experiments. 358

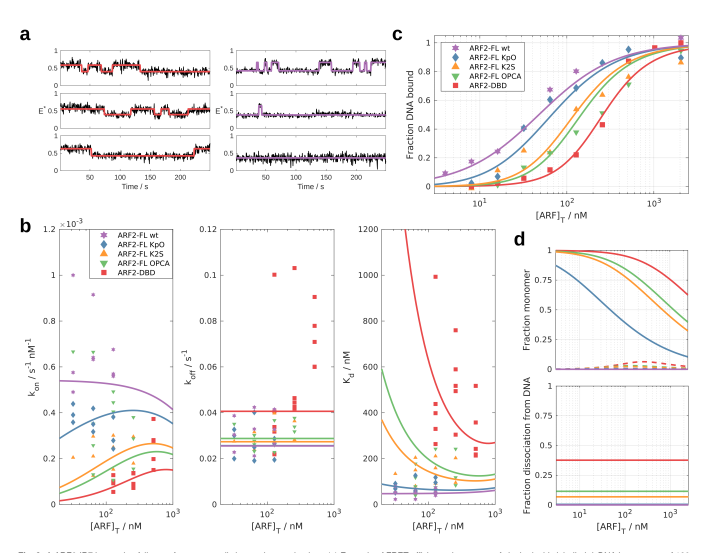
For a four-states cyclic model the binding isotherms (Fig. 2c, 359 colored markers) and observed kinetic constants (Fig. 2b, col-360 ored markers) can be fitted with a system of equations con-361 taining a set of three parameters shared across all AtARF2 362 variants  $(k_{\text{on,mic}}, k_{\text{off,DF}} \text{ and } k_{\text{off,MF}})$  and the variant-specific 363 parameter  $K_{\rm I}$  (see supporting note 1 and 2). Here,  $k_{\rm on,mic}$  is 364 the microscopic  $k_{\rm on}$  that a monomer displays when binding a 365 single AuxRE and hence it is equal to  $k_{\text{on,MM}}$  but it is half 366 the value of  $k_{\text{on,M}}$  and  $k_{\text{on,D}}$ . The global fit (Fig. 2b-c, colored 367 lines) returned the values of  $k_{\text{on,mic}}$ ,  $k_{\text{off,DF}}$ ,  $k_{\text{off,MF}}$  and  $K_{\text{IS}}$ 368 that best explain the experimental data (see Table 1). The 369

global fit converged to a  $K_{\rm I}$  of 0 nM for AtARF2-FL wt; in this 370 situation the equation of the fraction bound for the four-states 371 system simplifies to a simple binding isotherm for the dimer 372 (see supporting note 1). On the other binding isotherms, the 373 fit captured the shift of the binding to higher  $[ARF]_T$  thanks 374 to increasing values of  $K_{\rm I}$ , which corresponds to a decrease in 375 dimer stability (Fig. 2c). The fit of the binding isotherm of 376 AtARF2FL KpO is still very close to the one of AtARF2FL wt 377 but because of the decrease in dimer stability  $(K_{\rm I} = 0.016 \,\mu{\rm M})$ 378 its steepness is increased. The two AtARF2-FL mutants, K2S 379 and OPCA, show similar values of  $K_{\rm I}$  (0.23 µM and 0.41 µM 380 respectively). Lastly, the fit returned a value of  $K_{\rm I}$  of  $1.9\,\mu{\rm M}$ 381 for AtARF2-DBD. 382

Looking at the observed binding kinetics, the global fit cap-383 tures the trends of the observed  $k_{\rm on}$  and  $k_{\rm off}$  (Fig. 2b). Here, 384 AtARF2 variants with higher dimer stability display higher 385 values of observed  $k_{on}$  as their lower  $K_{I}$  increases the effec-386 tive concentration of ARF dimer in solution. The fits for the 387 observed  $k_{\text{off}}$  of AtARF2-FL wt and KpO converge to the 388 value of the dissociation kinetic of the dimer from the DNA 389  $(k_{\text{off},\text{DF}} = 0.026 \,\text{s}^{-1})$ . For the other datasets (AtARF2-FL 390 K2S, AtARF2-FL OPCA and AtARF2-DBD) the dissociation 391 of the dimer from the DNA caused by the loss of a monomer 392 plays a role and becomes almost as likely as the dissociation 393 of the dimer from the DNA in the case of AtARF2-DBD 394  $(k_{\text{off,DM}} = 0.016 \,\text{s}^{-1}).$ 395

The kinetic and equilibrium constants obtained from the global 396 fit show that the monomer is the predominant species in 397 solution for most of AtARF2 variants and for most of the 398 tested concentration range (Fig. 2d top, solid lines). Strikingly, 399 the fraction of DNA bound by a monomer never exceeds 400 10% (Fig. 2d top, dashed lines). The complex consisting of 401 the AtARF2 dimer bound to the DNA can split by either 402 a monomer or the dimer dissociating from the DNA. Our 403 results show that the dissociation of the dimer from the DNA 404 is the route used by AtARF2-FL and AtARF2-FL KpO, while 405 the dissociation from the DNA through loss of a monomer 406 becomes viable for AtARF2-FL K2S and OPCA and accounts 407 for approximatively 40% of the splitting events in the case 408 AtARF2-DBD (Fig. 2d bottom). 409

Dimer stability determines the binding kinetics of ARF-DBDs. 410 We showed that AtARF2 dimer stability induced by the pres-411 ence of the PB1 domain influences the kinetics of the binding 412 of ARF towards its DNA response element. We next asked 413 whether dimer stability is a more generic parameter defin-414 ing DNA binding affinity across the ARF protein family. To 415 this end, we purified the DNA-binding domains of two other 416 ARFs (class B AtARF1-DBD, class A AtARF5-DBD) and two 417 mutant versions (AtARF5-DBD G279N and AtARF5-DBD 418



**Fig. 2.** AtARF2-IR7 interaction follows a four-state cyclic interaction mechanism. (a) Example of FRET efficiency time traces of single doubly labelled dsDNA in presence of 128 nM of either AtARF2-DBD (left) or AtARF2-FL wt (right). The FRET efficiency is reported in black and the most probable sequence of hidden states returned by ebFRET is represented in red (AtARF2-DBD) and purple (AtARF2-FL wt). (b) Kinetics parameters obtained from ebFRET. For each ARF variant three concentrations closest to having half of the DNA bound to ARF were measured in at least three independent titrations. Each repeat of each concentration is analyzed independently using ebFRET obtaining a value of observed  $k_{con}$  and  $k_{off}$  (and, from their ratio the  $K_d$ ) and plotted using colored markers. (c) Fraction of DNA bound by ARF as function of ARF concentration (binding isotherm). The fractions bound were obtained from the histograms and plotted using colored markers (Fig. 1b, Materials and Methods). (b-c) The result of the global fit of the kinetics of binding and the fraction of DNA bound is reported as colored lines. (d) Features of the four states system as solved by the global fit. Top: In solution, the monomer is the most abundant species (solid lines). On the DNA, the monomer accounts for less than 10 % of the bound DNA (dashed lines). Bottom: Fraction dissociation of the AtARF2 monomer. Direct unbinding of the dimer is the predominant route for all AtARF2 monomer. Direct unbinding of the dimer is the predominant route for all AtARF2 monomer. Direct unbinding of the dimer is the predominant route for all AtARF2 monomer. Direct unbinding of the dimer is the predominant route for all AtARF2 monomer. Direct unbinding of an AtARF2-DBD.

Protein	$k_{on,mic}$ $[\mathrm{nM}^{-1}\mathrm{s}^{-1}]$	$k_{off,MF}$ $[\mathrm{s}^{-1}]$	$k_{off,DF}$ $[\mathrm{s}^{-1}]$	<i>K</i> ι [μΜ]
AtARF2-DBD AtARF2-FL OPCA AtARF2-FL K2S AtARF2-FL KpO AtARF2-FL wt	5.4 [3.6:7.3]×10 <sup>-4</sup>	1.7 [0.6:2.8]	2.6 [2.2:2.9]×10 <sup>-2</sup>	1.9 [0.9:2.9]×10 <sup>0</sup> 4.1 [1.0:7.1]×10 <sup>-1</sup> 2.3 [0.4:4.2]×10 <sup>-1</sup> 1.6 [-1.5:4.7]×10 <sup>-2</sup> 0.0 [-0.4:0.4]×10 <sup>-6</sup>

Table 1. Global fit: values and uncertainty of the fitting parameters reported as mean  $[95\,\%\,\text{Cl}]$ 

<sup>419</sup> R215A) and quantified their DNA binding affinity.

Experiments with AtARF1-DBD showed similar values 420 of  $k_{\rm on}$  and  $k_{\rm off}$   $(1.3 \times 10^{-4} \,\mathrm{nM^{-1}s^{-1}} \,95\%$  CI [0.8:1.8] 421  $0.080\,{\rm s}^{-1}$ 95 % CI [0.062:0.098], respectively) as AtARF2-422 DBD  $(1.1 \times 10^{-4} \text{ nM}^{-1} \text{s}^{-1} 95\% \text{ CI } [0.7:1.4], 0.066 \text{ s}^{-1} 95\%$ 423 CI [0.045:0.087], respectively; Fig. 3). On the other 424 hand, AtARF5-DBD showed a 5-fold increase in  $k_{on}$ 425  $(5.9 \times 10^{-4} \,\mathrm{nM^{-1}s^{-1}} \,95 \,\% \,\mathrm{CI} \,[2.3:9.4])$  and an 8-fold reduc-426 tion in  $k_{\text{off}}$  (  $0.0085 \,\text{s}^{-1} 95 \%$  CI [0.0051:0.0118]) compared to 427 AtARF2-DBD; which lead to a  $K_d$  of 15 nM (95% CI [12:18]) 428 In analogy with the considerations made for AtARF2-FL, the 429 increase in  $k_{on}$  and part of the decrease in  $k_{off}$  can be explained 430 with AtARF5-DBDs forming a tighter protein dimer compared 431 to AtARF1 and AtARF2 DBDs. 432

433 To test this hypothesis directly, we tested AtARF5-DBD G279N, a single amino acid mutation known to reduce 434 AtARF5-DBD dimerization(17). Strikingly, the kinetics 435 of the interaction between AtARF5-DBD G279N and the 436 IR7 became similar to the ones of AtARF1 and AtARF2 437 DBDs  $(1.2 \times 10^{-4} \text{ nM}^{-1} \text{s}^{-1} 95\% \text{ CI} [1.0:1.5], 0.10 \text{ s}^{-1} 95\%$ 438 CI [0.04:0.17]) validating our hypothesis. Finally, we tested 439 AtARF5-DBD R215A, a mutant in which a key amino acid for 440 the interaction with the DNA is mutated(17). This mutant 441 showed a 13-fold reduction of  $k_{\rm on}$  compared to the wild-type 442  $(0.46 \times 10^{-4} \,\mathrm{nM^{-1}s^{-1}} \,95 \,\% \text{ CI} [0.41:0.51]$  as well as a 39-fold 443 increase of  $k_{\text{off}}$  (0.33 s<sup>-1</sup> 95 % CI [0.14:0.52]) which translates 444 in a reduction of affinity of three orders of magnitude. We 445 note that the magnitude of the reduction of  $k_{on}$  is consistent 446 with the effect of charge neutralization of DNA-contacting 447 residues seen in other protein-DNA interactions(39) and is a 448 reminder of the importance of charged residues in defining 449 association kinetics (40, 41). 450

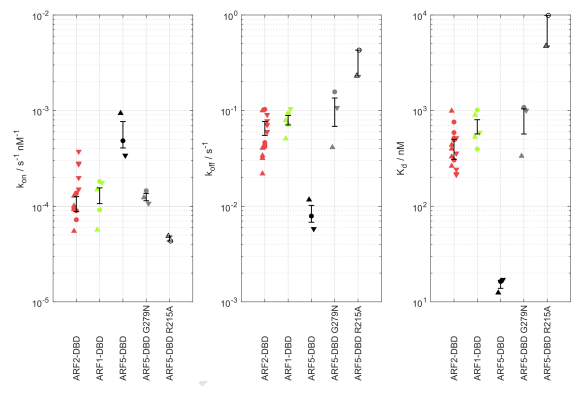
To directly measure ARF dimer stability, we measured SAXS
(Small-angle X-ray scattering) intensity profiles of AtARF5DBD and AtARF1-DBD (Fig. 4). The difference in dimer
stability is clear with AtARF5-DBD exhibiting higher dimer

prevalence at all tested concentration. The results on AtARF5-DBD are consistent with a dimerization  $K_d$  in the order of the tenth of  $\mu$ M while for AtARF1-DBD the  $K_d$  is in the order of few  $\mu$ M. These results confirm the expectation set by the analysis of the binding kinetics that AtARF5-DBD forms relatively stable dimers even in absence of the PB1 domain.

## Discussion

The ARF PB1 domain mediates the binding of Aux/IAA to 462 ARFs, allowing for inhibition of ARF activity(3-5). In ARFs 463 from Arabidopsis, deleting or mutating the PB1 domain leads 464 to hyperactive ARFs(23), consistent with a role in suppressing 465 activity. In contrast, the Marchantia ARF1 PB1 domain is 466 required for function, a deletion of this domain renders the pro-467 tein inactive (22). Given that the minimal set of ARF proteins 468 found in Marchantia qualifies these as likely representatives of 469 ancestral protein functions(42), an open question is what the 470 actual roles of ARF PB1 domains are. Here, we explored the 471 role of this domain in modulating the DNA binding affinity of 472 AtARF2 towards an IR7 response element. We found that full-473 length ARF2 protein has a strongly increased DNA-binding 474 affinity, which can be ascribed to interactions between the 475 PB1 domains. Interestingly, our results show that oligomer-476 ization does not further enhance the affinity towards bipartite 477 response elements. This behaviour is consistent with the fact 478 that additional ARF monomers (beside the initial two) do 479 not have any AuxRE left to further stabilize the binding to 480 the DNA. This said, the effect of the PB1 domain seen in our 481 experiments predicts that oligomerization should be relevant 482 on response elements comprising of more than two AuxREs 483 as the PB1 domain would enable cooperative binding beyond 484 the dimer. Given the short consensus sequence in AuxRE 485 motifs, these may occur in close proximity in promoters, in 486 which case oligomerization could generate additional cooper-487 ativity of ARF-DNA interaction. Regardless, the use of a 488 C-terminal head-to tail oligomerization domain (with two in-489

461



**Fig. 3.** Kinetics of the interaction between AtARF-DBDs and IR7. Kinetic parameters obtained from HMM analysis using ebFRET. The datapoints are marked with  $\Delta$ ,  $\Box$  and  $\nabla$  in order of increasing ARF concentration. The ARF concentrations were 128, 256, 512 nM for AtARF2-DBD and AtARF1-DBD, 8, 16 and 32 nM for AtARF5-DBD, 64, 128 and 256 nM for AtARF5-DBD G279N and 128 and 512 nM for AtARF5-DBD R215A. The error bars represent the standard deviations of the mean values. AtARF2-DBD and AtARF1-DBD behaved similarly while AtARF5-DBD showed increased  $k_{on}$  and decreased  $k_{off}$ . Consistent with a model in which part of the difference in kinetic can be explained by an increased stability of AtARF5-DBD mer. A weakening of AtARF5-DBD dimerisation (G279N mutant) leads to kinetic parameters that resemble the ones of AtARF1-DBD and AtARF2-DBD. In addition, AtARF5-DBD R215A mutant in a key amino acid for the interaction with DNA showed a  $k_{on}$  reduced by one order of magnitude and a  $k_{off}$  increased by almost two orders of magnitude compared to the wild-type.

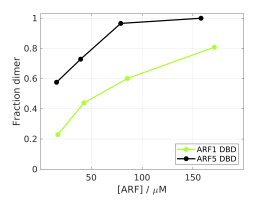


Fig. 4. Fraction of dimer measured using SAXS. The stability of the dimer of AtARF5-DBD is higher than the one of AtARF1-DBD for all tested concentration.

teraction faces) can be considered an efficient means of flexibleinteraction.

By simultaneously fitting affinity and kinetic data with the 492 analytical solution of a four-state cyclic model, we showed that 493 the increase of ARF DNA binding affinity can be completely 494 attributed to a shift in the dimer/monomer equilibrium of 495 ARFs. It follows that the effect of the PB1 domain on ARFs 496 affinity towards an IR7 response element is to shift the dimer-497 ization equilibrium towards the dimer. Strikingly, the fit allows 498 to obtain quantitative information about the protein dimer-499 ization  $K_{\rm d}$  in solution (i.e.,  $K_{\rm I}$ ) although the protein was not 500 directly observed in the experiment. 501

The kinetic parameters for AtARF2-DBD show that DNA 502 binding and unbinding is almost equally probable to happen 503 through a monomer-bound DNA intermediate or through di-504 rect binding of the dimer (Fig. 5). Moreover, the monomer 505 is the most common species in solution even at ARF concen-506 trations that saturate the DNA (i.e., DNA fully bound by an 507 ARF dimer); despite this, the percentage of DNA bound by 508 an ARF monomer never exceeds 10% as this intermediate is 509 short lived and quickly proceeds to either forming a dimer or 510 dissociate. 511

The kinetics of AtARF2-FL KpO-IR7 interaction is remarkably different (Fig. 5); here, the association between the DNA and a dimer follows almost exclusively the pathway where the ARF dimer is formed in solution. Moreover, the unbinding of a single monomer from a dimer bound to the DNA is unlikely  $(K_{\rm d} < 1 \text{ nM})$ .

In general, the importance of dimerization for stable DNA binding clearly emerges from our analysis as a reminder of the importance of cooperativity in protein-DNA interaction. Moreover, cooperativity is symmetric: a protein that can dimerize on a bipartite response element will bind it with higher affinity but also a bipartite element will stabilize the dimer of the protein that is bound to it. In particular, the dimer of AtARF2 is  $\approx 60$  times more stable when bound to the DNA compared to being in solution. 526

The analysis of the kinetics of the interaction between dif-527 ferent ARF DBDs and the IR7-RE suggests that the tighter 528 binding of AtARF5-DBD compared to AtARF1 and AtARF2 529 DBDs is in part due to the higher stability of its dimer. This 530 prediction is further corroborated by SAXS data showing that 531 AtARF5-DBD forms more stable dimers in solution compared 532 to AtARF1-DBD. The stable DNA binding that AtARF5-533 DBD achieve even in absence of the PB1 domain could explain 534 why AtARF5PB1 is a gain-of-function mutant that can ac-535 tivate auxin-responsive genes even in absence of auxin(23). 536 Then, the role of the PB1 domain of AtARF5 appears to be 537 mainly to bind the PB1 domain of Aux/IAAs coupling the 538 transcriptional output of ARF with the presence of auxin. 539 This hypothesis is confirmed by the fact that the PB1 domain 540 of AtARF5 has a homodimerization  $K_{\rm d}$  of 870 nM but an 541 heterodimerization  $K_d$  with the PB1 domain of Aux/IAA17 542 of 73 nM(8). Moreover, AtARF5 and other A-class ARFs have 543 been found to interact with many different Aux/IAA in a series 544 of protein-protein interaction assays(24, 43–53). A different 545 scenario is seen in case of AtARF2 (a class B ARF), where 546 our data suggest that the interaction between PB1 of different 547 AtARF2 monomers might be required to achieve the stable 548 DNA binding that enables protein function. This behavior of 549 the PB1 domain might be a common feature of other class 550 B/C ARFs and could explain why this class of ARFs has 551 been seen to interact with fewer members of the Aux/IAA 552 family(43, 51-54). 553

The picture emerging is that the PB1 domain has different 554 functions in the two main ARF classes in A. thaliana. In 555 class A ARFs, it serves as a mediator of auxin-responsiveness 556 whereas it stabilizes the DNA binding in class B (and perhaps 557 C) ARFs. This model of action for the PB1 domain is similar 558 to the one found in *M. polymorpha* as part of the recently 559 published minimal auxin response system(22) with one key 560 difference: MpARF1 (the only class A ARF in this species) 561 cannot function without its PB1 domain. Therefore, the PB1 562 domain of class A ARFs in *M. polymorpha* probably has the 563 double function of stabilizing the binding to the DNA and 564 interacting with MpAux/IAA. This double function opens 565 the possibility of a double repression by Aux/IAA, where, in 566 addition to the recruitment of the co-repressor TOPLESS(55)567 (TPL), a destabilization of ARF-DNA interaction might also 568 play a role. 569

Binding of ARF to bipartite AuxREs in the other two possible orientations (directed repeat DR and everted repeat ER), 571 should resemble the one seen for the IR with the difference that the dimerization through the DBD domain should not 573

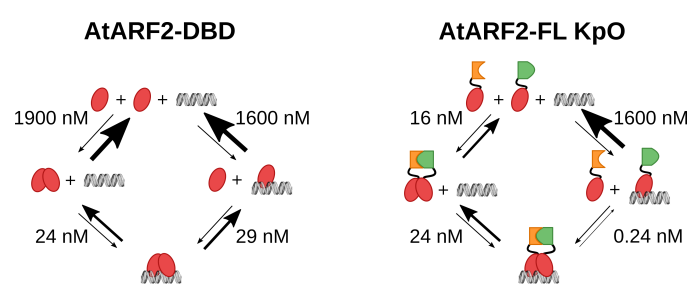


Fig. 5. Dissociation constants for the four-state cyclic model as determined from the fit in figure 2. ARF2-DBD binds the DNA via dimerization on the DNA and dimerization in solution with similar probability. ARF2-FL KpO mostly dimerizes in solution and then binds the DNA. The presence of the PB1 domain reduces all Kds making all interactions more stable (aside from the monomer-DNA interaction). This results in the higher affinity of ARF towards the RE when the PB1 domain is present.

be possible. In this scenario the analytical solution of the 574 4-state model for ARF-DBDs simplifies to a binding isotherm 575 for independent binding of the monomers characterized by low 576 steepness (no cooperativity, see also supporting Note 1). Strik-577 ingly, titrations presented in a recent publication(37) confirmed 578 this prediction; the binding of AtARF1-DBD and AtARF5-579 DBD to a bipartite DR5 was compatible with a simple binding 580 isotherm, whereas binding to an IR8 showed steeper response, 581 similar to the one seen here for AtARF2-DBD. Since stable 582 DNA binding arises from stable dimerization/oligomerization, 583 the topology of composite AuxREs dictates the affinity to-584 wards distinct ARF members differentiating them based on the 585 relative strength of the homotypic interaction through their 586 DBD and PB1 domains. Tweaking the affinities of different 587 ARFs towards the same DNA sequence can be achieved by af-588 fecting their dimerization properties and opens the possibility 589 for an evolutionary pathway of complex interactions between 590 members of the family. 591

Lastly, it is interesting to speculate on the biological signifi-592 cance of the dual, cooperative dimerization mode we identified 593 here. Effectively, the double-check mechanism would favor 594 dimerization of ARFs, when bound on DNA. ARF monomers 595 have limited sequence specificity of DNA binding. In the 596 hexanucleotide binding site, only 2 nucleotides are invariant, 597 and four are conserved (17, 19, 56). Thus, one would expect 598 a monomer to find binding sites frequently in the genome. 599 Dimerization adds two constraints that dramatically increase 600 specificity: a second, symmetric DNA element as well as a 601 fixed optimal space between the elements. This strongly limits 602 the probability of a random occurrence of the response element 603 and explains why dimerization is such a common features of 604

transcription factors across all domains of life(57, 58). Unfor-605 tunately, probing genome-wide ARF binding has been chal-606 lenging, and no comparisons between monomers and dimers 607 have yet been made. However, ARF2 and ARF5 have both 608 been used in DAP-seq binding site mapping on the Arabidop-609 sis genome(19). If dimerization limits the number of genomic 610 binding sites, one would predict that ARF5 – with a higher 611 propensity to dimerize (as shown here) binds fewer sites. This 612 is exactly what was found: ARF2 appears more promiscuous 613 in its binding  $\operatorname{profile}(19, 56)$ . A hypothesis, to be tested in the 614 future, is therefore that dimerization is the primary mechanism 615 for defining ARF-DNA binding specificity in vivo. 616

## Authors' Contributions

The project was initiated and supervised by Dolf Weijers 618 (D.W.) and Johannes Hohlbein (J.H.). Mattia Fontana (M.F.) 619 developed the methodology, defined the experimental design, 620 wrote and adapted the software for data analysis. Mark Roos-621 jen and Willy van den Berg purified the proteins. M.F. per-622 formed smFRET experiments, analyzed the data, derived the 623 analytical solutions for the system and implemented the global 624 fit. Isidro Crespo García performed SAXS experiments and 625 data analysis, supported by Marc Malfois and under super-626 vision by Roeland Boer. M.F., J.H. and D.W discussed the 627 content of the manuscript. M.F. wrote the draft manuscript 628 and produced the figures. All authors provided feedback on 629 the draft. 630

## **Competing Interests**

None to declare.

617

631

632

## 633 Funding Statement

This work was supported by a PhD fellowship (M.F.) from the Graduate School Experimental Plant Sciences to J.H.

- and D.W., the Ministry of Economy and Competitiveness
- of the Spanish Government grants PID2020-117028 GB-I00
- 638 (AEI/FEDER, EU), BIO2016-77883-C2-2-P (AEI/FEDER,
- <sup>639</sup> EU) and FIS2015-72574-EXP (AEI/FEDER, EU) to R.B.
- $= 10^{\circ} \text{ and } 10^{\circ} \text{ and } 10^{\circ} \text{ for the last of the$
- and a VICI grant (no. 865.14.001) from the Netherlands
- <sup>641</sup> Organization for Scientific Research (NWO) to D.W.

#### 642 Data availability

643 The experimental data is available on Zenodo:

644 https://doi.org/10.5281/zenodo.7249508.

#### 645 Acknowledgements

<sup>646</sup> We would like to thank all our colleagues at the Laboratory

of Biophysics and the Laboratory of Biochemistry for helpful
 discussions.

- P Grones, J Friml, Auxin transporters and binding proteins at a glance. J. Cell. Sci. 128, 1–7 (2015).
- 2. M Lavy, M Estelle, Mechanisms of auxin signaling. *Development* 143, 3226–3229 (2016).
- D Weijers, D Wagner, Transcriptional responses to the auxin hormone. Annu. Rev. Plant Biol.
   67, 539–574 (2016).
- F Parcy, T Vernoux, R Dumas, A glimpse beyond structures in auxin-dependent transcription.
   *Trends Plant Sci.* 21, 574–583 (2016).
- M Roosjen, S Paque, D Weijers, Auxin response factors: output control in auxin biology. J.
   *Exp. Bot.* 69, 179–188 (2018).
- 6. MH Nanao, et al., Structural basis for oligomerization of auxin transcriptional regulators. *Nat. Commun.* 5, 3617 (2014).
- DA Korasick, et al., Molecular basis for AUXIN RESPONSE FACTOR protein interaction and the control of auxin response repression. *Proc. Natl. Acad. Sci.* 111, 5427–5432 (2014).
- M Han, et al., Structural basis for the auxin-induced transcriptional regulation by Aux/IAA17.
   *Proc. Natl. Acad. Sci.* 111, 18613–18618 (2014).
- DC Dinesh, et al., Solution structure of the PsIAA4 oligomerization domain reveals interaction modes for transcription factors in early auxin response. *Proc. Natl. Acad. Sci.* 112, 6230–6235 (2015).
- T Ulmasov, G Hagen, TJ Guilfoyle, Activation and repression of transcription by auxin-response factors. *Proc. Natl. Acad. Sci.* 96, 5844–5849 (1999).
- SB Tiwari, G Hagen, T Guilfoyle, The Roles of Auxin Response Factor Domains in Auxin-Responsive Transcription. *The Plant Cell* 15, 533–543 (2003).
- T Ulmasov, G Hagen, TJ Guilfoyle, ARF1, a transcription factor that binds to auxin response
   elements. *Science* 276, 1865–1868 (1997).
- N Ballas, LM Wong, A Theologis, Identification of the Auxin-responsive Element, AuxRE, in
   the Primary indoleacetic Acid-inducible Gene, PS-IAA4/5, of Pea (Pisum sativum). J. Mol. Biol.
   875 233, 580–596 (1993).
- Y Li, ZB Liu, X Shi, G Hagen, TJ Guilfoyle, An Auxin-Inducible Element in Soybean SAUR
   Promoters. *Plant Physiol.* **106**, 37–43 (1994).
- T Ulmasov, ZB Liu, G Hagen, TJ Guilfoyle, Composite structure of auxin response elements
   The Plant Cell 7, 1611–1623 (1995).
- T Ulmasov, G Hagen, TJ Guilfoyle, Dimerization and DNA binding of auxin response factors.
   *The Plant J.* **19**, 309–319 (1999).
- DR Boer, et al., Structural basis for DNA binding specificity by the auxin-dependent ARF transcription factors. *Cell* **156** (2014).
- JM Franco-Zorrilla, et al., DNA-binding specificities of plant transcription factors and their potential to define target genes. *Proc. Natl. Acad. Sci.* 111, 2367–2372 (2014).
- R O'Malley, et al., Cistrome and epicistrome features shape the regulatory DNA landscape
   *Cell* 165, 1280–1292 (2016).
- 688 20. CY Liao, et al., Reporters for sensitive and quantitative measurement of auxin response. *Nat.* 689 *Methods* 12, 207–210 (2015).

- J Liu, et al., Natural variation in ARF18 gene simultaneously affects seed weight and silique length in polyploid rapeseed. Proc. Natl. Acad. Sci. 112, E5123–E5132 (2015).
- H Kato, et al., Design principles of a minimal auxin response system. Nat. Plants 6, 473–482 (2020).
- NT Krogan, W Ckurshumova, D Marcos, AE Caragea, T Berleth, Deletion of MP/ARF5 694 domains III and IV reveals a requirement for aux/IAA regulation in arabidopsis leaf vascular 695 patterning. *New Phytol.* **194**, 391–401 (2012). 696
- E Pierre-Jerome, BL Moss, A Lanctot, A Hageman, JL Nemhauser, Functional analysis of molecular interactions in synthetic auxin response circuits. *Proc. Natl. Acad. Sci.* 113, 11354–11359 (2016).
- S Farooq, J Hohlbein, Camera-based single-molecule FRET detection with improved time resolution. *Phys. Chem. Chem. Phys.* 17, 27862–27872 (2015).

702

703

- J Hohlbein, TD Craggs, T Cordes, Alternating-laser excitation: single-molecule FRET and beyond. *Chem. Soc. Rev.* 43, 1156–1171 (2014).
- AN Kapanidis, et al., Fluorescence-aided molecule sorting: Analysis of structure and interactions by alternating-laser excitation of single molecules. *Proc. Natl. Acad. Sci.* 101, 8936–8941 705 (2004). 706
- I Rasnik, SA McKinney, T Ha, Nonblinking and long-lasting single-molecule fluorescence 707 imaging. Nat. Methods 3, 891–893 (2006). 708
- T Cordes, J Vogelsang, P Tinnefeld, On the Mechanism of Trolox as Antiblinking and Antibleaching Reagent. J. Am. Chem. Soc. 131, 5018–5019 (2009) Publisher: American Chemical Society.
   GW Evans, J Hohlbein, T Craggs, L Aigrain, AN Kapanidis, Real-time single-molecule studies 712
- GW Evans, J Hohlbein, T Craggs, L Aigrain, AN Kapanidis, Real-time single-molecule studies 712 of the motions of DNA polymerase fingers illuminate DNA synthesis mechanisms. *Nucleic 713 Acids Res.* 43, 5998–6008 (2015). 714
- JW van de Meent, J Bronson, C Wiggins, R Gonzalez, Empirical bayes methods enable advanced population-level analyses of single-molecule FRET experiments. *Biophys. J.* 106, 1327–1337 (2014).
- J González Fernández, et al., NCD-SWEET Beamline Upgrade. Proc. Mech. Eng. Des. Synchrotron Radiat. Equip. Instrumentation MEDSI2018 (2018).
- N Gonzalez, et al., Beam Conditioning Optics at the ALBA NCD-SWEET Beamline. Proc. 720 Mech. Eng. Des. Synchrotron Radiat. Equip. Instrumentation MEDSi2018 (2018). 721
- 34. D Franke, et al., ATSAS 2.8: a comprehensive data analysis suite for small-angle scattering from macromolecular solutions. J. Appl. Crystallogr. 50, 1212–1225 (2017).
   722
- D Svergun, C Barberato, MHJ Koch, CRYSOL– a Program to Evaluate X-ray Solution Scattering of Biological Macromolecules from Atomic Coordinates. J. Appl. Crystallogr. 28, 768–773 (1995).
- PV Konarev, VV Volkov, AV Sokolova, MHJ Koch, DI Svergun, PRIMUS: a Windows PC-based 727 system for small-angle scattering data analysis. J. Appl. Crystallogr. 36, 1277–1282 (2003). 728
- A Freire-Rios, et al., Architecture of DNA elements mediating ARF transcription factor binding and auxin-responsive gene expression in *Arabidopsis. Proc. Natl. Acad. Sci.* **117**, 202009554 (2020).
- D Colquhoun, KA Dowsland, M Beato, AJR Plested, How to impose microscopic reversibility in complex reaction mechanisms. *Biophys. J.* 86, 3510–3518 (2004).
- SP Hancock, DA Hiller, JJ Perona, L Jen-Jacobson, The Energetic Contribution of Induced
   Electrostatic Asymmetry to DNA Bending by a Site-Specific Protein. J. Mol. Biol. 406, 285–312
   (2011).
- G Schreiber, G Haran, HX Zhou, Fundamental aspects of proteinprotein association kinetics. 737 Chem. Rev. 109, 839–860 (2009). 738
- HX Zhou, X Pang, Electrostatic Interactions in Protein Structure, Folding, Binding, and Condensation. Chem. Rev. 118, 1691–1741 (2018).
- SK Mutte, et al., Origin and evolution of the nuclear auxin response system. *eLife* 7, e33399 741 (2018).
   742
- F Ouellet, PJ Overvoorde, A Theologis, IAA17/AXR3: Biochemical Insight into an Auxin Mutant Phenotype. *The Plant Cell* 13, 829–841 (2001). 744
- T Hamann, E Benkova, I Bäurle, M Kientz, G Jürgens, The Arabidopsis BODENLOS gene encodes an auxin response protein inhibiting MONOPTEROS-mediated embryo patterning.
   *Genes & Dev.* 16, 1610–1615 (2002).
- K Tatematsu, et al., MASSUGU2 Encodes Aux/IAA19, an Auxin-Regulated Protein That Functions Together with the Transcriptional Activator NPH4/ARF7 to Regulate Differential Growth Responses of Hypocotyl and Formation of Lateral Roots in Arabidopsis thaliana. *The Plant Cell* 16, 379–393 (2004).
- 46. CS Hardtke, et al., Overlapping and non-redundant functions of the Arabidopsis auxin response factors MONOPTEROS and NONPHOTOTROPIC HYPOCOTYL 4. *Development* 131, 1089–1100 (2004).
- D Weijers, et al., Developmental specificity of auxin response by pairs of ARF and Aux/IAA 755 transcriptional regulators. *The EMBO J.* 24, 1874–1885 (2005). 756

- 757 48. H Fukaki, Y Nakao, Y Okushima, A Theologis, M Tasaka, Tissue-specific expression of
- stabilized SOLITARY-ROOT/IAA14 alters lateral root development in Arabidopsis. *The Plant J.* 44, 382–395 (2005).
- H Muto, et al., Fluorescence Cross-Correlation Analyses of the Molecular Interaction between an Aux/IAA Protein, MSG2/IAA19, and Protein–Protein Interaction Domains of Auxin Response Factors of Arabidopsis Expressed in HeLa Cells. *Plant Cell Physiol.* 47, 1095–1101 (2006).
- T Uehara, Y Okushima, T Mimura, M Tasaka, H Fukaki, Domain II Mutations in CRANE/IAA18
   Suppress Lateral Root Formation and Affect Shoot Development in Arabidopsis thaliana. *Plant*
- Cell Physiol. 49, 1025–1038 (2008).
   C Shen, et al., Functional analysis of the structural domain of ARF proteins in rice (Oryza
- Shen, et al., Functional analysis of the structural domain of AHF proteins in rice (Oryza sativa L.). *J. Exp. Bot.* **61**, 3971–3981 (2010).
- 768 52. T Vernoux, et al., The auxin signalling network translates dynamic input into robust patterning
   769 at the shoot apex. *Mol. Syst. Biol.* **7**, 508 (2011).
- S Piya, SK Shrestha, B Binder, CNJ Stewart, T Hewezi, Protein-protein interaction and gene
   co-expression maps of ARFs and aux/IAAs in arabidopsis. *Front. Plant Sci.* 5 (2014).
- T Ulmasov, J Murfett, G Hagen, TJ Guilfoyle, Aux/IAA proteins repress expression of reporter genes containing natural and highly active synthetic auxin response elements. *The Plant Cell* 9, 1963–1971 (1997).
- 55. H Szemenyei, M Hannon, JA Long, TOPLESS Mediates Auxin-Dependent Transcriptional
   Repression During Arabidopsis Embryogenesis. *Science* **319**, 1384–1386 (2008).
- 777 56. A Stigliani, et al., Capturing auxin response factors syntax using DNA binding models. *Mol.* 778 *Plant* 12, 822–832 (2019).
- 57. GD Amoutzias, DL Robertson, Y Van de Peer, SG Oliver, Choose your partners: dimerization
   in eukaryotic transcription factors. *Trends Biochem. Sci.* 33, 220–229 (2008).
- 781 58. L Strader, D Weijers, D Wagner, Plant transcription factors being in the right place with the
- 782 right company. *Curr. Opin. Plant Biol.* **65**, 102136 (2022).