

# Structural basis for LEAFY floral switch function and similarity with helix-turn-helix proteins

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**The LEAFY (LFY) protein is a key regulator of flower development in angiosperms. Its gradually increased expression governs the sharp floral transition, and LFY subsequently controls the patterning of flower meristems by inducing the expression of floral homeotic genes. Despite a wealth of genetic data, how LFY functions at the molecular level is poorly understood. Here, we report crystal structures for the DNA-binding domain of *Arabidopsis thaliana* LFY bound to two target promoter elements. LFY adopts a novel seven-helix fold that binds DNA as a cooperative dimer, forming base-specific contacts in both the major and minor grooves. Cooperativity is mediated by two basic residues and plausibly accounts for LFY's effectiveness in triggering sharp developmental transitions. Our structure reveals an unexpected similarity between LFY and helix-turn-helix proteins, including homeodomain proteins known to regulate morphogenesis in higher eukaryotes. The appearance of flowering plants has been linked to the molecular evolution of LFY. Our study provides a unique framework to elucidate the molecular mechanisms underlying floral development and the evolutionary history of flowering plants.**

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## Introduction

Homeotic genes control developmental patterns and organ morphogenesis. In animals, they encode transcription factors of the homeodomain family, such as Hox and paired proteins, which contact DNA through one or several helix-turn-helix (HTH) motifs (Gehring *et al*, 1994; Underhill, 2000). In plants, most homeotic genes determining the identity of floral organs encode MADS-box transcription factors, suggesting that plants and animals have adopted distinct types of homeotic regulators (Meyerowitz, 1997; Ng and Yanofsky, 2001). In addition to organ identity genes, plants also use another class of regulators named 'meristem identity genes', which control floral meristem versus shoot/inflorescence fate. In *Arabidopsis thaliana*, the meristem identity genes *LEAFY* (*LFY*) and *APETALA1* (*API*) induce flower development, whereas *TERMINAL FLOWER1* (*TFL1*) promotes inflorescence development (Blazquez *et al*, 2006). Mutations or ectopic expression of these genes result in complete or partial interconversions between flower and inflorescence meristems.

The *LFY* gene encodes a plant-specific transcription factor, which has a cardinal function in this process, regulating both the transition to flowering and the subsequent patterning of young floral meristems. During the plant vegetative growth, *LFY* expression increases in newly formed leaves until a certain threshold is reached. *LFY* then induces the expression of *API* and *CAULIFLOWER* (*CAL*) genes and triggers the abrupt floral transition (Blazquez *et al*, 2006). Once the floral meristem is established, *LFY* governs its spatial patterning by inducing the expression of the floral homeotic ABC genes, such as *API*, *AP3* or *AGAMOUS* (*AG*), which control the identity of stereotypically arranged floral organs (Coen and Meyerowitz, 1991; Lohmann and Weigel, 2002).

*LFY* is found in all terrestrial plants from moss to angiosperms; its sequence shows a high level of conservation throughout the plant kingdom but no apparent similarity to other proteins (Maizel *et al*, 2005). Unlike many plant transcription factors that evolved by gene duplication to form a multigene family (Riechmann and Ratcliffe, 2000; Shiu *et al*, 2005), *LFY* is present in single copy in most angiosperms and *lfy* mutants available from several species such as snapdragon, petunia, tomato or maize show, as in *Arabidopsis*, partial or complete flower-to-shoot conversions (Coen *et al*, 1990; Souer *et al*, 1998; Molinero-Rosales *et al*, 1999; Bomblies *et al*, 2003). In gymnosperms, a paralogous *NEEDLY* (*NLY*) clade of genes exists. No mutant is available in these species, but *LFY* and *NLY* expression patterns are also consistent with a role in reproductive organ development

(reviewed in Frohlich and Chase, 2007). Because of its central role in determining floral meristem identity, and considering that *NLY* disappeared concomitantly with the appearance of flowers, *LFY* has been put at the centre of different evolutionary scenarios that rationalize the appearance of the successful angiosperm group (Albert *et al*, 2002; Frohlich, 2003; Frohlich and Chase, 2007; Theissen and Melzer, 2007).

*LFY* activates gene expression by recognizing pseudo-palindromic sequence elements (CCANTGT/G) in the promoters of its target genes, including *AP1* (one site) and *AG* (four sites; *AG-I* to *AG-IV*) (Parcy *et al*, 1998; Busch *et al*, 1999; Lohmann *et al*, 2001; Lamb *et al*, 2002; Hong *et al*, 2003). *LFY* has two domains, a partially conserved N-terminal domain that is thought to contribute to transcriptional activation and a highly conserved C-terminal domain responsible for DNA binding (*LFY-C*) (Coen *et al*, 1990; Maizel *et al*, 2005). *LFY* functions synergistically with coregulators such as the *WUSCHEL* (*WUS*) homeodomain protein (Lenhard *et al*, 2001; Lohmann *et al*, 2001) or the *UFO* F-Box protein (Lee *et al*, 1997; Parcy *et al*, 1998; Chae *et al*, 2008).

In this study, we show that *LFY* binds DNA cooperatively as a dimer, a property shown to be essential to trigger developmental switches. The crystal structure of *LFY-C* bound to DNA reveals the molecular basis for sequence-specific recognition and cooperative binding as well as an unexpected similarity of *LFY* with HTH proteins such as homeodomain transcription factors. Our findings enable to formulate new hypotheses on the appearance of angiosperms in evolution.

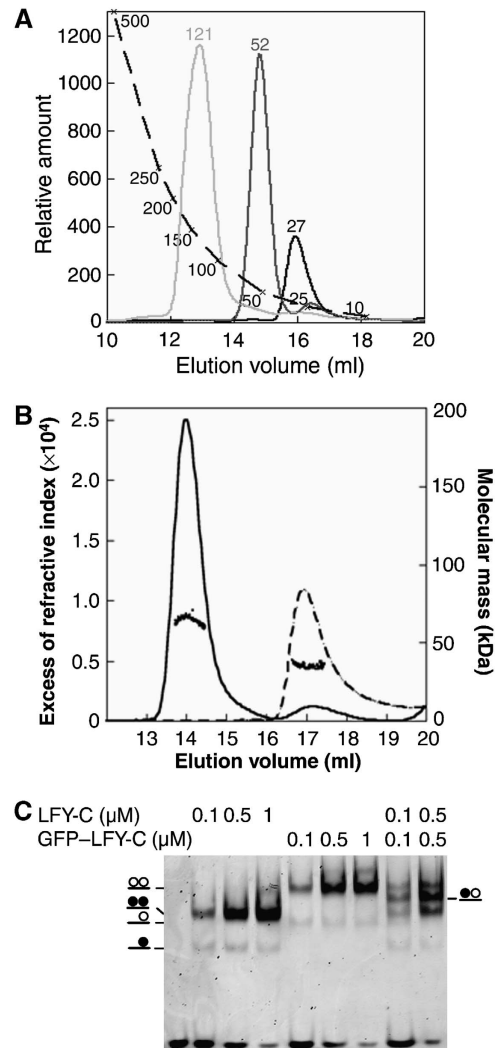
## Results and discussion

### *LFY-C* dimerizes on DNA binding

We produced the recombinant *LFY* DNA-binding domain (*LFY-C*, residues 223–424) and showed by size-exclusion chromatography (SEC) that it is monomeric in the absence of DNA (Figure 1A and B). In electrophoretic mobility shift assays (EMSAs), *LFY-C* recognized a DNA probe bearing an *AP1* site as two distinct species: a major protein–DNA complex and a minor one of higher mobility (Figure 1C). Multi-angle laser light scattering (MALLS) coupled to SEC demonstrated that the major complex contained two *LFY-C* molecules per DNA duplex (Figure 1B). The homodimeric nature of *LFY* in this complex was confirmed by mixing untagged and GFP-tagged *LFY-C* and observing a single new species attributable to the formation of an *LFY-C*/GFP–*LFY-C*/DNA complex (Figure 1C). Using probes mutated in one half-site of the palindrome, we confirmed that the minor, high-mobility species corresponds to a single *LFY-C* monomer bound to DNA (Supplementary Figure 2).

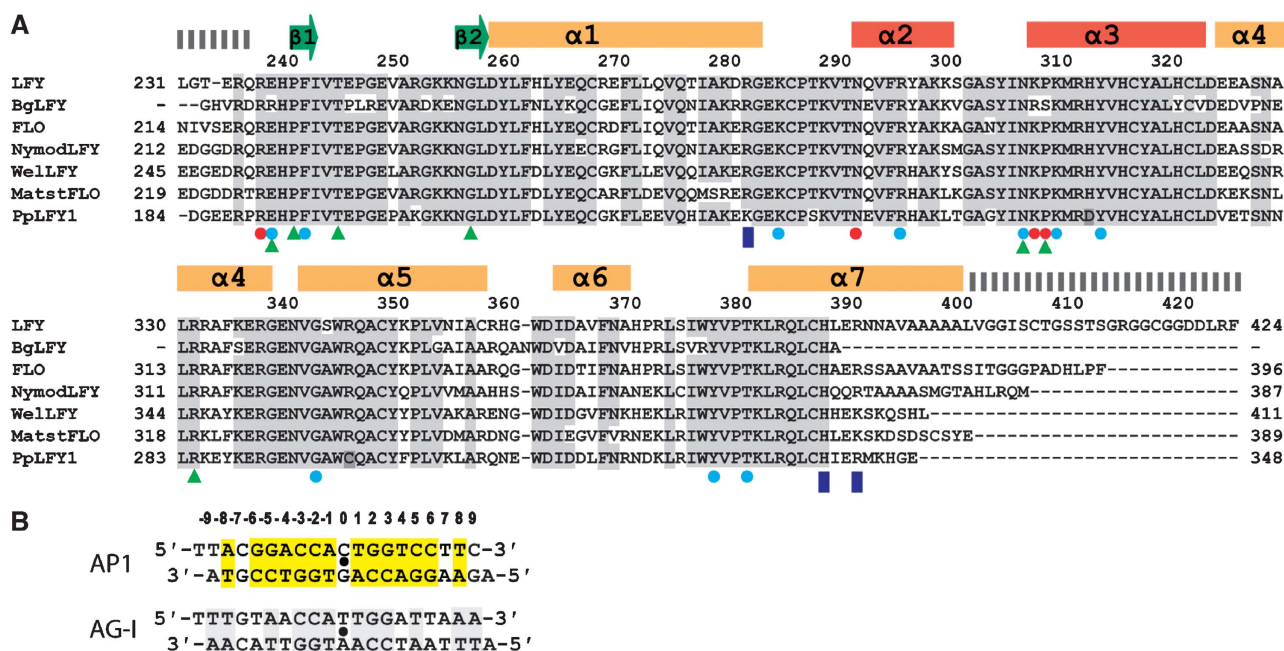
### Structure of the *LFY* DNA-binding domain bound to its DNA recognition site

To understand how *LFY* specifically recognizes its DNA target sequences, we crystallized *LFY-C* in complex with DNA. We solved the structure of *LFY-C* bound to two different *LFY*-binding sites, *AP1* and *AG-I* at 2.1- and 2.3-Å resolution, respectively (Figures 2, and 3A and B; Table I). The overall structure shows an *LFY-C* dimer bound to a pseudo-palindromic DNA duplex, where the *LFY-C* monomers are related by a crystallographic dyad. The DNA duplexes used for co-crystallization deviate from strict two-fold symmetry at the 5' ends and at base pairs (bp)  $\pm 9$ ,  $\pm 7$  and  $\pm 0$  in the *AP1* site



**Figure 1** DNA-dependent dimerization of *LFY-C*. (A) Size-exclusion chromatography. *LFY-C* (40  $\mu$ M, black curve), *AP1* DNA (10  $\mu$ M, dark grey curve), *LFY-C* (40  $\mu$ M) + *AP1* DNA (10  $\mu$ M, light grey curve) were analysed. *LFY-C* elution at a volume corresponding to 28 kDa is consistent with the monomer size (25.7 kDa), the DNA duplex elutes earlier than expected at a volume corresponding to 28 kDa because of its elongated shape. The *LFY*/DNA complex elutes at a volume corresponding to 121 kDa. Molecular weights estimated from the calibration curve (dashed line) are indicated. (B) Molecular mass of *LFY-C* alone (dashed line) or in combination with *AP1* DNA (solid line) determined by multi-angle laser light scattering and refractometry combined with size-exclusion chromatography. Elution profiles were monitored by excess refractive index (left ordinate axis). Dots show the molecular mass distribution (right ordinate axis). Average molecular mass is  $64 \pm 2$  kDa for the *LFY-C*/DNA complex (65 kDa theoretical size for a dimeric complex) and  $35 \pm 1$  kDa for *LFY-C* alone (26 kDa theoretical size for *LFY-C* monomer). (C) Electrophoretic mobility shift assay (EMSA) with 10 nM *AP1* DNA and various *LFY-C* or GFP–*LFY-C* concentrations. Schematic complexes with *LFY-C* (filled circle) and GFP–*LFY-C* (open circle) are depicted.

(5' end, bp  $\pm 7$ ,  $\pm 6$ ,  $\pm 4$  and  $\pm 0$  in the *AG-I* site). Nevertheless, the pseudo-dyads of the DNA duplexes coincide with the crystallographic dyad, probably as a result of the random bimodal orientation of the DNA duplex around the dyad (see Materials and methods). The resulting molecular averaging does not impair our interpretation of the protein–DNA interface. In the final  $2F_o - F_c$  elec-



**Figure 2** Sequence alignments. (A) Aligned C-terminal amino-acid sequences of LFY (*Arabidopsis thaliana*, AAA32826), BgLFY (*Brownea grandiceps*, AAS79888), FLO (*Antirrhinum majus*, P23915), NymodLFY (*Nymphaea odorata*, AAF77609), WellFY (*Welwitschia mirabilis*, AAF23870), MatstLFY (*Matteuccia struthiopteris*, AAF77608) and PpLFY1 (*Physcomitrella patens*, BAD91043). Identical and conservatively substituted residues are depicted on a grey background. Secondary structure elements are indicated. Residues involved in interactions with DNA bases and backbone are labelled with red and blue circles, respectively. Dashed bars indicate disordered regions in the crystal, blue rectangles indicate the residues involved in dimerization. Green triangles indicate the position of *Arabidopsis* mutations and residues divergent in PpLFY1 are highlighted in pink. (B) Two DNA duplexes containing the LEAFY-binding sites from *AP1* and *AG-I* promoters present in the LEAFY-DNA complex crystals are depicted. Base pairs related by a dyad (indicated by a black dot) are highlighted in yellow.

tron density map, but also in the initial solvent-flattened single isomorphous replacement with anomalous scattering (SIRAS) electron density map (Supplementary Figure 1), the sugar-phosphate backbone of the DNA is well defined and shows no evidence of conformational averaging. The density for palindromic DNA bases is clearly defined, whereas the density at non-palindromic positions is consistent with the superposition of two different base pairs. Furthermore, all residues close to the DNA are clearly defined and we do not observe any diffuse density, which suggests that each monomer undergoes only minor changes to adapt to the slightly different half-sites. Despite the differences between the *AP1* and *AG-I* binding sites (Figure 2B), both complex structures are very similar and can be superimposed with an r.m.s. distance of 0.55 Å for 163 C $\alpha$  and 19 phosphate atoms.

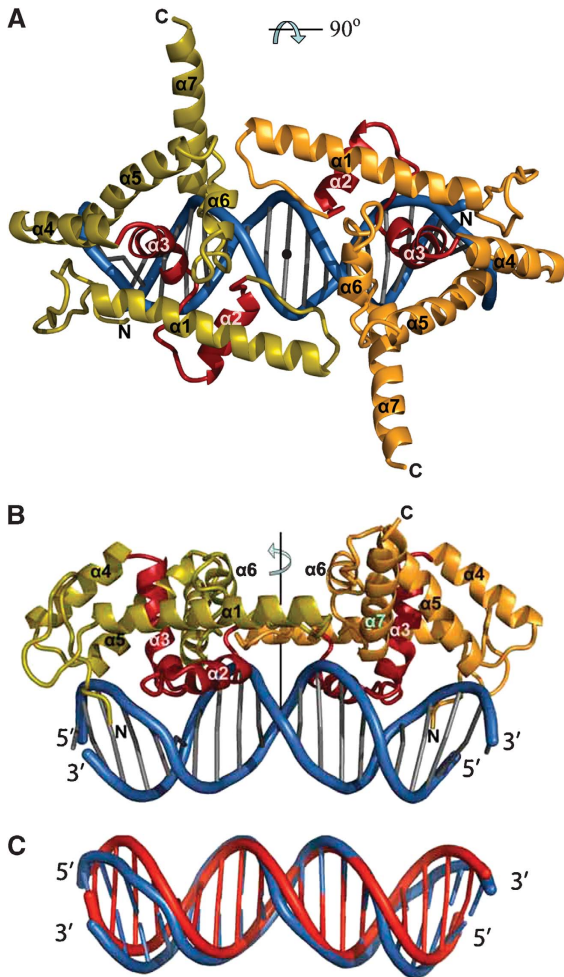
LFY-C (with residues 237–399 ordered in the crystal structure) adopts a compact fold that interacts principally with a single DNA half-site (Figures 3A and B, and 4A and B). The fold is defined by two short  $\beta$ -strands followed by seven helices connected by short loops (Figure 3A and B). The absence of any extended hydrophobic patches at its surface suggests that LFY-C represents an autonomous DNA-binding domain without a large interface to its N-terminal domain. Helices  $\alpha 2$  and  $\alpha 3$  define a HTH motif (Aravind *et al*, 2005), with helix  $\alpha 3$  occupying the major groove and mediating most of the DNA contacts. The DNA in the complex adopts a B-DNA-like conformation exhibiting an overall bend of about 20° (Figure 3C), which can be localized to two kinks of about 10° at base pairs  $\pm 2/\pm 3$ . Both ends of the DNA duplex are AT rich and the minor grooves are narrower compared with

classical B-DNA. Narrowing of the minor groove is slightly more pronounced in the *AG-I* duplex than in *AP1*.

#### DNA recognition in the major and minor grooves

Sequence-specific contacts between LFY and the DNA involve both the minor and major grooves. Base-specific contacts in the major groove are formed by Asn291 and Lys307 in helices  $\alpha 2$  and  $\alpha 3$ , which together specify the two invariant guanines at positions  $\pm 2$  and  $\pm 3$  (Figure 4A and B). Mutating either of these residues into alanine resulted in considerably lower DNA-binding affinity (Figure 4C), whereas previous studies showed loss of binding when the corresponding base pairs were mutated (Parcy *et al*, 1998; Busch *et al*, 1999). The *Arabidopsis lfy-20* mutation (N306D) adjacent to Lys307 also leads to a reduced DNA-binding affinity (Supplementary Figure 3) and a weak *lfy* phenotype *in planta* (Weigel *et al*, 1992; Maizel *et al*, 2005), presumably because the negatively charged aspartate interacts unfavourably with the DNA backbone (Figure 4A and B).

Base-specific recognition in the minor groove is mediated by Arg237, which is the first ordered N-terminal residue in the crystal structure. At the *AP1* site, its side chain points towards A:T base pairs  $\pm 8$  and contacts the exocyclic O2 of thymine 8 and also the O2 of cytosine 7 in one half-site, or the O2 of cytosine-9 in the other half-site (Figure 4A and B; Supplementary Figure 4). In the *AG-I* site, T:A base pair 8 is replaced by A:T, and in the LFY/*AG-I* complex, the Arg237 side chain adopts a different conformation, which allows it to recognize the thymine of the opposite strand (Supplementary Figure 4). The importance of this interaction is underscored by



**Figure 3** Structure of the LFY-C dimer bound to DNA. (A, B) Two orthogonal views of the LFY-C dimer (residues 237–399) bound to DNA. Monomers are coloured in olive and orange with the helix-turn-helix (HTH, helices  $\alpha 2$  and  $\alpha 3$ ) motif in red. The DNA duplex is depicted in blue. Figures 3, 4B, 5A and 6 were produced with program Pymol (Delano, 2002). (C) Superposition of the DNA duplex found in the LEAFY-DNA complex (blue) with regular B-form DNA (red).

the presence of A:T or T:A base pairs at position 8 in all 12 confirmed half-sites (Parcy *et al*, 1998; Busch *et al*, 1999; Lohmann *et al*, 2001; Lamb *et al*, 2002; Hong *et al*, 2003). The consensus LFY-binding site is therefore more accurately defined as T/ANNNNCCANTGT/GNNNNT/A (with the centre of the pseudo-palindrome underlined). The Arg237 side chain is inserted into an AT-rich narrow minor groove (Figure 4A and B), similar to that observed in the Hox homeodomain-Exd-DNA complex, where the narrow minor groove was shown to enhance the electrostatic interaction between DNA backbone and arginine side chain (Joshi *et al*, 2007). The R237A mutation led to a strongly reduced affinity of LFY-C for *API* (Figure 4C). In contrast, changing the adenine 8 into a cytosine in *API* reduced only moderately the LFY-C-binding affinity (Figure 4C; *API* m5), presumably because the arginine side chain can contact the adjacent base (Supplementary Figure 4). Finally, next to Arg237, the two *lfy* mutations (*lfy-4* (E238K) and *lfy-5* (P240L)) result in decreased *in vitro* binding affinities (Supplementary Figure 3) and lead to a mutant phenotype *in planta* (Weigel *et al*, 1992).

An unusual contact with DNA is mediated by Pro308 that points between the guanines in base pairs  $\pm 5$  and  $\pm 6$ , which results in a pronounced propeller twist for base pair  $\pm 5$  and local bending of the DNA at this position (Figure 4B). The mutant *lfy-28* (P308L) is impaired in DNA binding and gives rise to an intermediate to strong phenotype *in planta* (Figure 4C–F), as a likely consequence of a steric clash of the leucine side chain with the guanine bases. In contrast, a small side chain such as alanine perfectly fits in this protein–DNA interface and, indeed, the mutant protein P308A showed a wild-type-binding affinity (Figure 4C). Pro308 is not strictly conserved and is substituted by serine in some *Brownea* species (Figure 2A). This substitution probably modifies DNA binding, because serine can form direct hydrogen bonds to DNA bases at positions  $\pm 4$  and  $\pm 5$  and it replaces P308, which locally distorts DNA. However, the conformational flexibility of serine and its ability to function as hydrogen bond donor or acceptor makes it difficult to predict the preferred binding specificity. Moreover, P308S is systematically associated with the K307R substitution, affecting the base-contacting residue Lys307 (Figure 2A). LFY proteins from *Brownea* species might therefore recognize significantly different DNA target sites.

Similar to most protein–DNA co-crystal structures, not all bases in the consensus LFY site T/ANNNNCCANTGT/GNNNNT/A are specified through direct interactions with the protein. Additional specificity presumably arises from sequence-dependent deformability of the DNA, sometimes referred to as ‘indirect readout’. Dinucleotide steps CA/TG at bp  $\pm 1$ /bp  $\pm 2$  are part of the consensus LFY site and are particularly flexible, which might facilitate the observed kink of the DNA at base pairs  $\pm 2/\pm 3$ . However, these particular sequences are not critically required as they are not conserved in the *AP3-I* binding site (Lamb *et al*, 2002).

Not all LFY mutations directly affect DNA contacts. Mutations *lfy-3* (T244M) and *lfy-9* (R331K) (Weigel *et al*, 1992) disturb two interacting amino acids, both of which contribute to a polar network that connects N-terminal residues with helices  $\alpha 1$  and  $\alpha 4$ . Similarly, two other residues (His312 and Arg345) interact in a typical planar stacking. His312 and Arg345 are conserved except in the LFY protein from *Physcomitrella patens* (PpLFY1) where they are substituted by aspartate and cysteine, respectively (Maizel *et al*, 2005). His312 forms part of helix  $\alpha 3$  and is located just one helical turn above Pro308 at the N-terminal end of helix  $\alpha 3$ . In addition, the preceding residue Lys307 directly contacts the guanine in base pairs  $\pm 2$  (Figure 4A). The loss of the His312/Arg345 stacking interaction in the moss PpLFY1 likely affects the orientation of helix  $\alpha 3$ , explaining the altered DNA-binding properties of this orthologue, whereas reverting the aspartate into histidine restores the binding activity of PpLFY1 to canonical LFY-binding sites (Maizel *et al*, 2005).

#### Structural basis for cooperative DNA binding

The structure of the LFY-C/DNA complex also reveals important monomer–monomer interactions governing its DNA-binding mode (Figure 5A). Our EMSA analysis shows that LFY-C binds DNA in a cooperative manner: the monomeric complex is present only in minor amounts as compared with the dimeric complex, even at low LFY-C concentrations (Figure 5B) and binding of the second monomer occurs with a 90-fold higher affinity than binding of the first one

**Table 1** Structure determination of the LEAFY–DNA complex

Data statistics					
Data set	Resolution (Å) <sup>a</sup>	Reflections measured/unique	$R_{\text{meas}}$ (%) <sup>b</sup>	$I/\sigma$	Completeness (%)
API/LFY: space group P6 <sub>5</sub> 22, unit cell dimensions a = b = 98.8 Å, c = 177.4 Å					
Native	20–2.1 (2.2–2.1)	503 443/29 859	4.8 (54.3)	38.5 (5.0)	97.4 (83.9)
EMTS	20–2.4 (2.5–2.4)	283 416/20 463	11.1 (50.2)	15.5 (4.7)	99.6 (99.3)
AG-I/LFY: space group P6 <sub>5</sub> 22, unit cell dimensions a = b = 98.4 Å, c = 176.4 Å					
Native	20–2.3 (2.4–2.3)	328 932/23 456	7.9 (75.1)	26.7 (3.4)	99.8 (100.0)
Phasing statistics for the EMTS derivative (SIRAS)					
Wavelength (Å)	0.934				
Phasing power <sup>c</sup>	1.68				
Figure of merit	0.468				
$R_{\text{cullis}}$ <sup>d</sup>	0.674				
Number of mercury sites	4				
Refinement statistics					
		LFY/API	LFY/AG-I		
Resolution <sup>a</sup>		19.1–2.1 (2.2–2.1)	20.0–2.3 (2.4–2.3)		
Total number of non-hydrogen protein atoms		1352	1332		
Total number of non-hydrogen DNA atoms		526	567		
Number of water molecules		148	105		
R-factor (%) <sup>a</sup>		21.0 (25.0) for 26 896 reflections	22.1 (29.0) for 22 559 reflections		
$R_{\text{free}}$ (%) <sup>a,e</sup>		23.7 (26.4) for 1521 reflections	24.9 (30.7) for 1183 reflections		
<i>r.m.s. deviations</i>					
Bond lengths (Å)		0.009	0.008		
Bond angles (deg)		1.22	1.21		
<i>Average temperature factors (Å<sup>2</sup>)</i>					
Protein		41.1	39.8		
DNA		48.1	46.4		
Solvent		42.8	48.5		
r.m.s.d. of covalently linked atoms (Å <sup>2</sup> )		2.55	3.45		
<i>Residues in Ramachandran plot<sup>f</sup></i>					
Most favoured regions (%)		93.2 (136)	95.2 (139)		
Additionally allowed regions (%)		6.8 (10)	4.8 (7)		
Generously allowed regions (%)		0 (0)	0 (0)		
Disallowed region (%)		0 (0)	0 (0)		

<sup>a</sup>Values for the highest resolution range are given in parenthesis.

<sup>b</sup> $R_{\text{meas}}$  is a redundancy independent R-factor as defined in Diederichs and Karplus (1997).

<sup>c</sup>Phasing power is the mean value of the heavy-atom structure factor amplitudes divided by the mean lack of closure.

<sup>d</sup> $R_{\text{cullis}}$  is the mean lack of closure divided by the mean isomorphous difference.

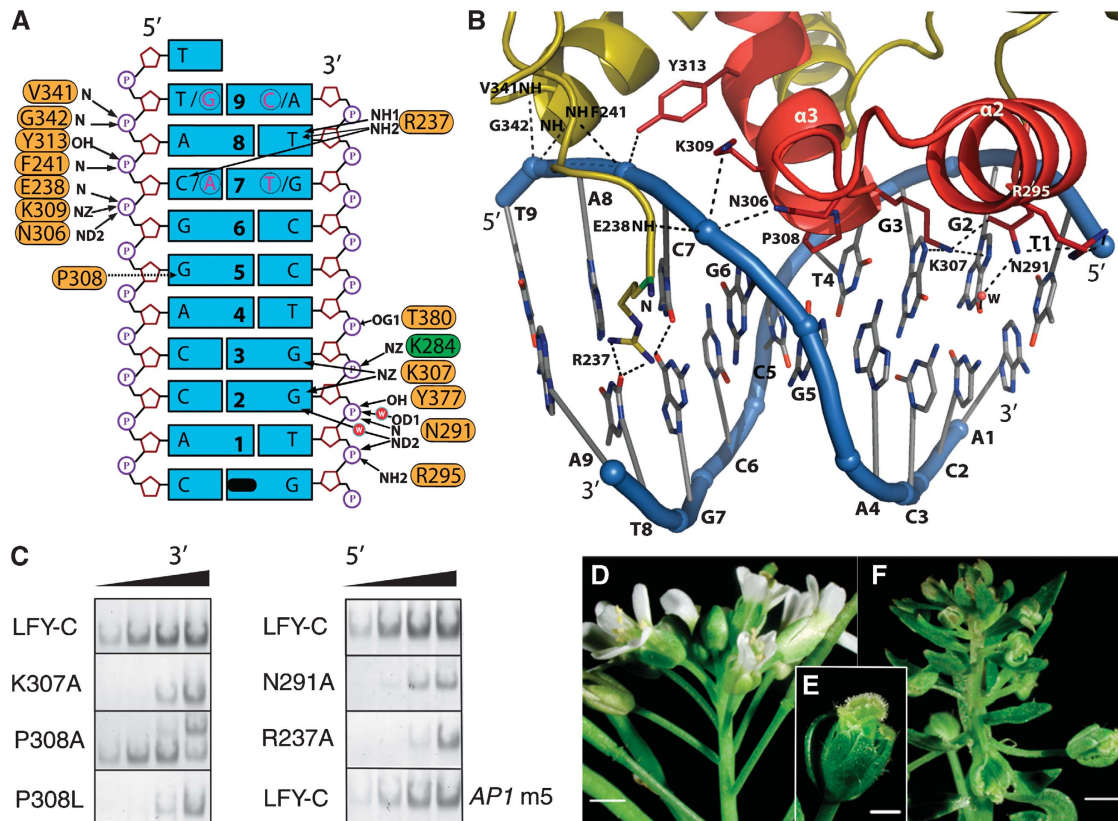
<sup>e</sup> $R_{\text{free}}$  was calculated from a subset of 5% of the data.

<sup>f</sup>Number of residues are given in parentheses.

(Figure 5C; Supplementary Figure 5). This type of cooperative binding can result either from DNA conformability, where binding of one monomer favours the binding of the second monomer, or from protein–protein interactions between DNA-bound monomers (Senear *et al*, 1998; Schumacher *et al*, 2002; Panne *et al*, 2004). Our structure suggests the latter. The LFY dimer comprises a small interface of 420 Å<sup>2</sup> buried surface area formed by loop  $\alpha$ 12 and helix  $\alpha$ 7 in which the two residues His387 and Arg390 form hydrogen bonds with the backbone carbonyl of Asp280 (Figure 5A). We validated the importance of these contacts by mutagenesis: cooperativity of binding is moderately affected in the H387A or R390A single mutants but more strongly reduced in a H387A/R390A double mutant (Figure 5B and C; Supplementary Figure 5). Therefore, the small monomer–monomer interface (with a major contribution of His387 and

Arg390) rather than DNA conformability is responsible for the cooperative binding. Whether the N-terminal domain of LFY also participates in dimerization, in the presence or absence of DNA, will require additional experiments.

A better understanding of LFY's DNA-binding mode also provides insight into its molecular switch function. DNA-binding cooperativity, as well as dimerization, allows transcription factors to work at lower concentrations and to enhance the sigmoidality of their response curves. When combined with feedback loops, it has been shown essential for threshold-dependent genetic switches (Burz *et al*, 1998; Cherry and Adler, 2000). LFY is involved in a positive autoregulation loop through activation of the homologous API and CAL genes, that in turn activate LFY expression (Bowman *et al*, 1993; Liljegren *et al*, 1999). LFY-binding cooperativity combined with the API/CAL feedback loop



**Figure 4** DNA recognition by LEAFY. (A) Protein–DNA interactions in one *AP1* half-site. Dyad-related base pairs 7 and 9 from the other half-site are shown in pink and encircled. Polar and hydrophobic interactions are shown with solid and dashed arrows, respectively. K284 belongs to the other monomer and is depicted in green. The pseudo-dyad coinciding with the crystallographic dyad is depicted in black. (B) Ribbon diagram of one LEAFY monomer bound to its *AP1* half-site. The protein is coloured in olive except for the HTH motif shown in red. Polar interactions are indicated by dashed lines. For clarity, only side chains in contact with DNA are shown. (C) Effect of selected mutations on LFY-C DNA-binding affinity to *AP1* DNA. EMSAs were performed with wild-type and mutant LFY-C (100–250–750–2000 nM from left to right). Only dimeric complexes are shown except for P308A that gave rise to an unknown higher complex. *AP1* m5 mutant DNA contains base pair C:G instead of A:T at position  $\pm 8$  (see Supplementary Table 3 for full DNA sequences). Phenotype of the wild-type *Arabidopsis* inflorescence (D) and *lfj-28* (P308L) mutant inflorescence (F) and flower (E). Scale bar is 1 mm on (D, F) and 0.5 mm on (E).

therefore provides a plausible explanation for the threshold-dependent floral switch triggered by LFY.

Many transcription factors bind DNA as homodimers but also form heterodimers, thereby extending their spectrum of recognized DNA target sequences (Klemm *et al*, 1998; Garvie and Wolberger, 2001). LFY has been shown to activate the *AG* organ identity gene synergistically with the homeodomain protein WUS (Lohmann *et al*, 2001). As adjacent WUS- and LFY-binding sites are present on the *AG* regulatory sequence, it has been suggested that LFY and WUS could bind simultaneously (Lohmann *et al*, 2001; Hong *et al*, 2003). Preliminary model building indicates that LFY homodimers cannot be accommodated with WUS at adjacent LFY- and WUS-binding sites. This observation raises the intriguing possibility that they might either compete for the same binding sites or more likely could form LFY–WUS heterodimers.

#### LFY shows similarities with HTH proteins

The nature and origin of LFY had so far remained elusive: LFY-C's primary sequence shows unusually strong sequence conservation within its family but has no apparent similarity to any described transcription factor. The crystal structure of LFY-C bound to DNA reveals a seven-helix domain with many residues involved in protein–DNA interactions, tightly

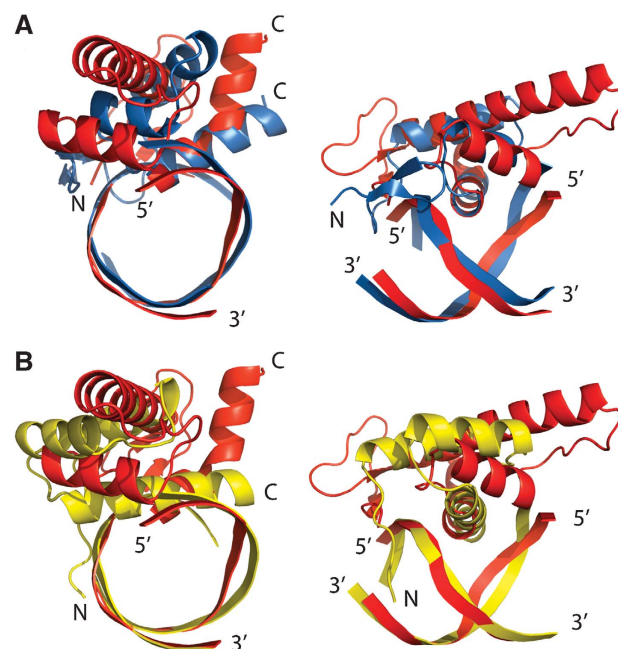
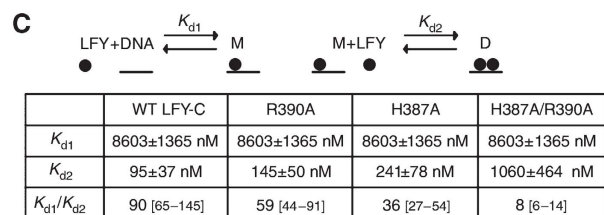
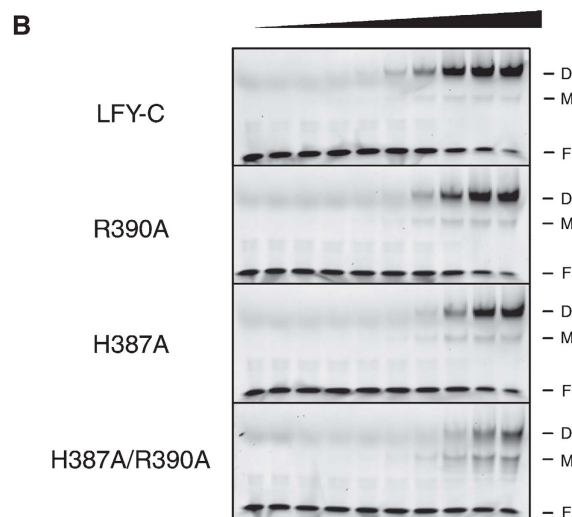
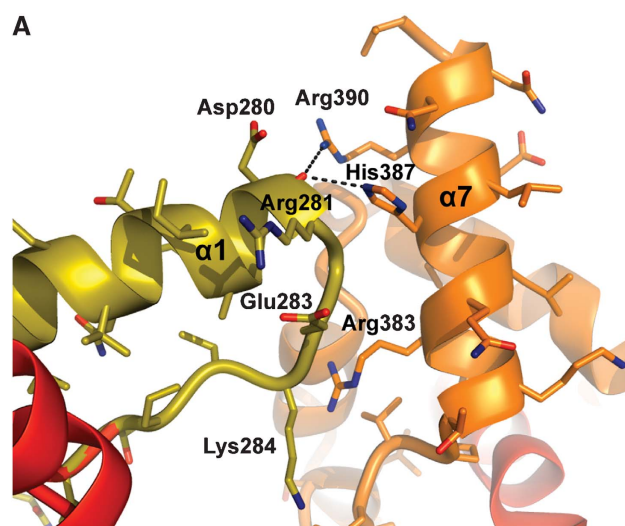
constrained packing interactions in the hydrophobic core and protein–protein interactions with the other monomer. Presumably, these observed tight structural and functional constraints on many residues spread over the entire DNA-binding domain explain the high level of sequence conservation within LFY-C.

The LFY-C structure contains an unpredicted HTH motif formed by helices  $\alpha 2$  and  $\alpha 3$  as part of the overall fold. HTH motifs are present in a wide variety of DNA-binding proteins throughout the three kingdoms of life. They are typically found in a bundle of 3–6  $\alpha$ -helices or combined with  $\beta$ -sheets (winged HTH/fork head domain), which provide a stabilizing hydrophobic core (Weigel and Jackle, 1990; Aravind *et al*, 2005). Comparison of LFY-C against the Protein Data Bank using program DALI (Holm and Sander, 1993) detects similarity of relative short  $\alpha$ -helical segments ( $\sim 60$  amino-acid residues) with different  $\alpha$ -helical proteins including HTH proteins (maximal Dali Z-score 3.0, pairs with  $Z < 2.0$  are structurally dissimilar). A search comprising only the first three N-terminal helices, including the HTH motif, mainly showed similarity to different HTH proteins with slightly higher scores (maximal Dali Z-score: 4.5). When considering just the three helices  $\alpha 1$ ,  $\alpha 2$  and  $\alpha 3$ , LFY aligns well with other three-helix bundle HTH proteins, including the homeodo-

main protein engrailed (r.m.s.d.<sub>40C $\alpha$</sub>  = 2.9 Å), the paired domain (r.m.s.d.<sub>44C $\alpha$</sub>  = 3.5 Å) and the Tc3 transposase (r.m.s.d.<sub>30C $\alpha$</sub>  = 2.4 Å). LFY and partitioning protein KorB (r.m.s.d.<sub>71C $\alpha$</sub>  = 3.7 Å) share some similarity beyond the typical DNA/RNA-binding three-helical bundle core (Russell and Barton, 1992; Khare *et al*, 2004), where five of the seven LFY-C helices, including the HTH motif, roughly superimpose with KorB helices. However, LFY cannot be easily assigned to any of the described classes of HTH proteins (Aravind *et al*, 2005) and it therefore represents a new variant of multi-helical bundle proteins.

The DNA recognition mode of LFY is similar to those observed for the paired domain, Tc3A transposase, Hin recombinase and  $\lambda$  repressor (van Pouderooyen *et al*, 1997; Xu *et al*, 1999). The axis of the recognition helix  $\alpha 3$  in the

HTH of these proteins is oriented parallel to the edges of the nucleotide bases. Only the N terminus of the recognition helix is inserted into the major groove of the DNA, whereas the short helix  $\alpha 2$  has a supporting function. In contrast, in homeodomain proteins, the long probe helix  $\alpha 3$  runs more parallel to the neighbouring DNA phosphate backbone, and mainly the central part of helix  $\alpha 3$  contacts the DNA



**Figure 6** Comparison of LFY-C with paired and homeodomain DNA binding. (A) Two orthogonal views of LFY-C helices  $\alpha 1$ – $\alpha 3$  bound to their DNA target site (red) superimposed with the three-helical bundle core of the N-terminal subdomain of the paired domain of *Drosophila* Prd (blue, PDB-id: 1pdn). (B) Superposition with the homeodomain of *Drosophila* engrailed bound to DNA (yellow, PDB-id: 1hdd), where the centre of recognition helix  $\alpha 3$  inserts into the major groove.

**Figure 5** The LFY-C dimer interface mediates cooperative binding. (A) The dimer interface is viewed perpendicular to the DNA axis. Polar contacts between the two monomers (in orange and olive) are shown with dashed lines. (B) EMSA with increasing concentrations (0, 10, 20, 50, 100, 200, 500, 1000, 2000 and 3000 nM from left to right) of LFY-C wild-type, R390A mutant, H387A mutant, and H387A/R390A double mutant and 50  $\mu$ M *API* DNA. Free DNA (F), monomeric (M) and dimeric (D) complexes are indicated. (C) Estimation of dissociation constants for wild-type LFY-C and three mutant versions (H387A, R390A and H387A/R390A). Binding of LFY-C to *API* DNA was modelled as two equilibrium reactions as detailed in Supplementary data: (1) Binding of a first LFY-C monomer to *API* DNA, leading to the formation of the monomeric complex (M) and characterized by the  $K_{d1}$  dissociation constant; (2) binding of a second LFY-C monomer to M, leading to the formation of the dimeric complex (D) and characterized by  $K_{d2}$ . EMSA signals from (B) were quantified and the corresponding experimental values were fitted with theoretical equations describing the two equilibria. The errors and intervals between square brackets indicated correspond to the 95% confidence interval. An elevated  $K_{d1}/K_{d2}$  ratio reflects a high level of cooperativity, whereas a ratio of 1 would indicate an absence of binding cooperativity. The single mutations resulted in a weak decrease of cooperativity, whereas the H387A/R390A double mutation strongly decreased the cooperativity.

(Figure 6). Similarity between LFY and the paired domain also includes a small two-stranded  $\beta$ -sheet, which precedes the three-helix bundle, and N-terminal residues, which are inserted into the minor groove. However, the minor groove contacting residues are located at the most N-terminal end of LFY-C, whereas in the paired domain they protrude from the loop connecting the two short N-terminal  $\beta$  strands.

Sequence similarities are too weak to suggest a precise evolutionary origin for LFY, although structural resemblances indicate that it might derive from ancestral HTH proteins, including paired and homeodomain proteins (Rosinski and Atchley, 1999; Breitling and Gerber, 2000; Aravind *et al*, 2005). Until now, most plant homeotic genes were found to encode MADS box transcription factors, whereas plant homeodomain proteins rather control meristem homeostasis and cell division (Meyerowitz, 1997; Ng and Yanofsky, 2001). Our study reveals that the LFY master regulator, which determines flower meristem fate and controls the expression of floral organ identity genes, shares structural similarity with other HTH proteins, indicating that this universal DNA-binding motif has also been adopted in plants to trigger major developmental switches.

### Prospects regarding the appearance of angiosperms

The LFY-C structure combined with more than 200 LFY sequences from all types of terrestrial plants offers a unique opportunity to detect key residues in evolution. Some charged LFY-C surface residues (such as Lys253 or Lys254) are strictly conserved, suggesting that they might participate in interactions with other proteins. Other residues are conserved in all angiosperms but not in the non-flowering plants. For example, R390, identified as one of the residues mediating interaction between monomers and cooperative binding, has been conserved in angiosperm LFY proteins, whereas most LFY from non-flowering plants, such as gymnosperms and ferns, show a lysine at this position. This amino-acid change presumably weakens the interaction between monomers and thereby reduces the DNA-binding affinity. The acquisition of R390 might therefore have been important for flower evolution. Because LFY stands at the very centre of the network regulating flower development, it has been proposed that modifications of the LFY gene contributed to the appearance of floral structures in evolution (Albert *et al*, 2002; Frohlich, 2003; Frohlich and Chase, 2007; Theissen and Melzer, 2007). The availability of the LFY-C crystal structure provides a unique framework for generating plausible hypotheses that relate the appearance of angiosperms to specific events during the molecular evolution of LFY. The 'functional synthesis' approach that combines phylogeny, biochemical and structural analyses with functional assays *in vivo* (Dean and Thornton, 2007) can now be applied to LFY to try to solve one of the most puzzling enigmas of plant biology: the origin of flowers.

## Materials and methods

### Plant material

The *lfy-28* mutant allele of *A. thaliana* (accession Landsberg *erecta*) was kindly provided by D Weigel (Max Planck Institute, Tübingen, Germany) and originally isolated by J Fletcher (PGEC, Albany). *lfy-28* mutant had been back-crossed twice with the wild type, and individuals showing a mutant phenotype were selected from segregating populations. Plants were grown at 25°C in long days (16 h light).

### Plasmid constructions

**Expression plasmids.** LFY-C (residues 223–424 from *A. thaliana* LFY cDNA) was amplified from pIL-8 (obtained from D Weigel) with Pfu Turbo Polymerase (Stratagene, France) and primers of pP1242 (5'CTCTCGAGCCCGGGCTAGAAACGCAAGTCGTCGCC3') and of pP1244 (5'CTCTCGAGCCCGGGCTATCCGGTACAGCTAATACCGCC3'), subcloned into pCR-TOPO-BluntII (Invitrogen, Cergy Pontoise, France) and shuttled to pETM-11 (Dummler *et al*, 2005) as *NcoI/XhoI* fragment to yield the pCH28 expression vector. pETM-11 contains an N-terminal 6  $\times$  His tag followed by a tobacco etch virus (TEV) cleavage site.

**LFY-GFP plasmid.** A GFP fragment was amplified from pBS-GLFY plasmid obtained from X Wu (Wu *et al*, 2003) using primers oETH1001 5'CCCACTACTGAGAATCTTTATTTCCAGGGCCAGTTCAG TAAAGGAGAAGAAC3' and oETH1002 5'CCCAAACTACTACCTCCG TTGCCGTATCTGTTTGTATAGTTCATCCAT3'. The amplified fragment was subsequently used as a megaprimer to amplify plasmid pCH28 and yield pETH8 (6His-TEV-GFP-LFY-C).

**Expression plasmids for mutant LFY-C.** pCH45 (K307A), pCH46 (N291A), pCH47 (R237A), pCH48 (P308A), pCH49 (D280K), pCH50 (H387A/R390A), pCH54 (H387A), pEDW127 (R390A), pCH55 (*lfy-28*, P308L), pETH21 (*lfy-4*, E238K), pETH23 (*lfy-20*, N306D) and pCH56 (*lfy-5*, P240L) were derived from pCH28 using the megaprimer strategy with appropriate primers (Kirsch and Joly, 1998). All plasmids were verified by sequencing.

### Protein expression, purification and crystallization

Wild-type and mutant LFY-C domains were expressed using *Escherichia coli* strain RosettaBlue(DE3)pLysS (Novagen, Strasbourg, France). After induction by 0.5 mM IPTG, cells were grown overnight at 22°C. For cell lysis, the pellet of 1 l culture was sonicated in 30 ml lysis buffer A (500 mM NaCl, 20 mM Tris-HCl pH 8, 5 mM imidazole, 5% glycerol, 5 mM Tris(2-carboxyethyl)phosphine hydrochloride), one protease inhibitor cocktail tablet Complete EDTA-free (Roche, Meylan, France) and centrifuged for 40 min at 30 000 g. The clear supernatant was incubated for about 1 h with 1 ml Ni-NTA resin (Qiagen, Courtaboeuf, France). The resin was transferred into a column, washed with 20 column volumes (CVs) of buffer A, buffer A + 50 mM imidazole (10 CV) and eluted with buffer A + 380 mM imidazole. The fractions containing the protein were pooled and applied to a Hi-load Superdex-200 16/60 prep grade column (GE Healthcare, Orsay, France) equilibrated with 200 mM NaCl, 20 mM Tris-HCl pH 8, 5 mM dithiothreitol (DTT) to eliminate aggregated proteins by SEC. Protein concentration was estimated using the Bradford assay (Bradford, 1976).

For crystallographic experiments, after elution on the metal-affinity column, the histidine tag was cleaved at 4°C overnight with TEV protease (0.01% w/w, 16 h, 4°C) during the dialysis step against buffer B (500 mM NaCl, 20 mM Tris pH 7.5, 5 mM DTT). The TEV protease, the histidine tag and the uncleaved protein were removed by re-passing the dialysed sample over the Ni-affinity column. The protein was separated from the remaining DNA contamination using the anion-exchange column MonoQ HR10/10 (GE Healthcare) pre-equilibrated in buffer B. Pure protein was recovered in the flow-through, whereas DNA remained bound to the resin. Aggregated protein was removed by SEC with Superdex S75GL column (GE Healthcare) in 200 mM NaCl, 10 mM Tris pH 7.5 and 5 mM DTT. The protein concentration was adjusted to 7.5 mg/ml. DNA oligonucleotides were chemically synthesized and purified by anion-exchange chromatography following established procedures (Cramer and Muller, 1997).

### EMSAs

Single-stranded oligonucleotides, 5'-labelled with tetra-methylcarboxy-rhodamine (Sigma, Saint Quentin Fallavier, France), were annealed to non-fluorescent complementary oligonucleotides in annealing buffer (10 mM Tris pH 7.5, 150 mM NaCl and 1 mM EDTA). The sequences of oligonucleotides used are indicated in Supplementary Table 1. Binding reactions were performed in 20  $\mu$ l binding buffer (150 mM NaCl, 20 mM Tris-HCl pH 7.5, 1% glycerol, 0.25 mM EDTA, 2 mM MgCl<sub>2</sub> and 1 mM DTT) supplemented with 28 ng/ $\mu$ l fish sperm DNA (Roche) and 10 nM double-stranded DNA probe or 140 ng/ $\mu$ l fish sperm DNA for 50 nM DNA probe (Figure 5). Binding reactions were loaded onto native 6% polyacrylamide gels



0.5 × TBE (45 mM Tris, 45 mM boric acid and 1 mM EDTA pH 8) and electrophoresed at 90 V for 80 min at 4°C. Gels were scanned on a Typhoon 9400 scanner (Molecular Dynamics, Sunnyvale, CA; excitation light 532 nm, emission filter 580 BP 30) and signals were quantified using ImageQuant software (Molecular Dynamics). Estimations of  $K_{d1}$  and  $K_{d2}$  (Figure 5; Supplementary Figure 5) were based on the quantifications of binding experiments shown in Figure 5B. The binding model equations used to calculate these  $K_d$  values are explained in detail in Supplementary data.

### SEC

The molecular size of LFY-C/*API* complex was determined using a Superdex-200 10/300GL column (GE Healthcare), equilibrated with buffer containing 150 mM NaCl, 16 mM Tris–HCl pH 7.5, 0.6 mM EDTA and 1 mM DTT, and calibrated with low and high molecular weight protein standards (gel filtration calibration kit; GE Healthcare). Our samples (LFY-C 40 μM, *API* WT 10 μM and LFY-C 40 μM + *API* WT 10 μM) were analysed in the same buffer as protein standards, and molecular size is deduced from the standard curve.

### Analytical SEC and MALLS-SEC

Separation by SEC was carried out with a S200 Superdex column (GE Healthcare). The column was equilibrated in 20 mM Tris–HCl, 150 mM NaCl buffer at pH 7.5. Separations were performed at 20°C with a flow rate of 0.6 ml min<sup>-1</sup>. Protein solution (50 μl) at a concentration of 5 mg ml<sup>-1</sup> was injected. The elution was monitored by using a DAWN-EOS detector with a laser emitting at 690 nm for online MALLS measurement (Wyatt Technology Corp., Santa Barbara, CA), and with a RI2000 detector for online refractive index measurements (Schambeck SFD). Molecular mass calculation was performed as described using the ASTRA software (Gerard *et al*, 2007).

### Crystallization

For co-crystallization using the hanging drop method, protein and DNA duplexes were mixed in a molar ratio of 2:1. The best crystals were obtained at 4°C with 20-mer oligonucleotides bearing complementary A:T overhangs and with 10% PEG 400, 100 mM KCl, 10 mM CaCl<sub>2</sub>, 50 mM HEPES (NaOH) pH 7.0 as reservoir solution. Single crystals grew to a maximal size of 300 × 300 × 500 μm<sup>3</sup> and were stepwise transferred to reservoir solution containing 30% (v/v) glycerol for cryo-protection. For preparation of the mercury derivative, the crystals were soaked in the reservoir solution supplemented with 0.1 mM ethylmercury thiosalicylate (EMTS) for 2 h.

### X-ray structure determination

The crystals of the LFY-C/*API*/DNA complex belong to space group P6<sub>5</sub>22 ( $a = b = 98.8 \text{ \AA}$ ,  $c = 177.4 \text{ \AA}$ ), diffracted up to 2.1 Å resolution and contain half a complex per asymmetric unit. Crystals of the LFY-C/*AG-I*/DNA complex are isomorphous but diffracted slightly weaker (Table I). Diffraction data collected at ESRF beamlines ID14-1, ID29 and ID23-2 were processed using program XDS (Kabsch, 1993). The structure of the LFY-C/*API*/DNA complex was solved using the SIRAS method with EMTS as derivative. The quality of native and derivative data sets is summarized in Table I. Mercury sites were located using program SOLVE (Terwilliger and Berendzen, 1999) and phases were calculated with program SHARP (de la Fortelle and Brice, 1997). The experimental electron density map (Supplementary Figure 1) allowed us to automatically build the initial model using program ARP/wARP (Perrakis *et al*, 2001) followed by manually adjusting some side chain conformations

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with program COOT (Emsley and Cowtan, 2004) and refinement with program Refmac5 including a TLS refinement with seven groups (Murshudov *et al*, 1997) and later with program Phenix (Adams *et al*, 2002). In space group P6<sub>5</sub>22, the two monomers bound to the pseudo-palindromic DNA duplex are related by a crystallographic dyad. In the crystal, the pseudo-dyad of the DNA coincides with the crystallographic dyad, although the DNA duplexes deviate from strict two-fold symmetry at base pairs 0, ±7, ±9 and the overhanging 5'-end, in the *API* site and at base pairs 0, ±4, ±6 and ±7 and the overhanging 5'-end in the *AG-I* site. To confirm our space group assignment and the underlying assumption that the DNA duplexes used for co-crystallization are randomly distributed in two orientations, the data were reprocessed in the lower symmetry space group P6<sub>5</sub> lacking the dyad, which did not significantly change the  $R_{\text{meas}}$  values. Subsequently, models of the LFY-C dimer bound to the 20-mer DNA duplex were built in space group P6<sub>5</sub> for the *API* and *AG-I* sites and refined in two independent orientations yielding very similar final  $R_{\text{cryst}}$  and  $R_{\text{free}}$  values compared with the refinement in space group P6<sub>5</sub>22. In both orientations (and for both target sites), the final  $F_o - F_c$  electron density maps showed pairs of difference Fourier peaks ( $\sim 7\sigma$ ) of similar height at the non-palindromic bases, indicating that a unique orientation of the DNA duplexes does not correctly describe the situation in the crystals. Finally, simulated-annealing omit maps in space group P6<sub>5</sub> where the non-palindromic bases were omitted showed averaged densities for the omitted bases in both complexes, further confirming the assigned space group P6<sub>5</sub>22.

To account for the two orientations of the DNA in the crystal during the refinement, two nucleotides with 50% occupancy were introduced at the non-palindromic positions. The final model of the LFY-C/*API* complex at 2.1-Å resolution ( $R_{\text{cryst}} = 21.0\%$ ;  $R_{\text{free}} = 23.7\%$ ) comprises residues 237–399 of the LFY DNA-binding domain, whereas the poorly conserved 25 C-terminal residues are disordered. For the refinement of the LFY/*AG-I* complex, the *API* DNA sequence in the LFY/*API* complex was replaced with the *AG-I* sequence. Multiple rounds of refinement (including TLS refinement with seven groups) using program Refmac5 (Murshudov *et al*, 1997) and Phenix (Adams *et al*, 2002) yielded a model with  $R_{\text{cryst}}$  of 22.1% and  $R_{\text{free}}$  of 24.9% using data between 20 and 2.3 Å resolution. The atomic coordinates and structure factors for the LFY/*API* and LFY/*AG-I* complexes have been deposited with the Protein Data Bank under accession codes 2vy1, r2vy1sf and 2vy2, r2vy2sf, respectively.

### Supplementary data

Supplementary data are available at *The EMBO Journal* Online (<http://www.embojournal.org>).

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