FLC-mediated flowering repression is positively regulated by sumoylation

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Abstract

Flowering locus C (FLC), a floral repressor, is a critical factor for the transition from the vegetative to the reproductive phase. Here, the mechanisms regulating the activity and stability of the FLC protein were investigated. Bimolecular fluorescence complementation and in vitro pull-down analyses showed that FLC interacts with the E3 small ubiquitin-like modifier (SUMO) ligase AtSIZ1, suggesting that AtSIZ1 is an E3 SUMO ligase for FLC. In vitro sumoylation assays showed that FLC is modified by SUMO in the presence of SUMO-activating enzyme E1 and conjugating enzyme E2, but its sumoylation is inhibited by AtSIZ1. In transgenic plants, inducible AtSIZ1 overexpression led to an increase in the concentration of FLC and delayed the post-translational decay of FLC, indicating that AtSIZ1 stabilizes FLC through direct binding. Also, the flowering time in mutant FLC (K154R, a mutation of the sumoylation site)-overexpressing plants was comparable with that in the wild type, whereas flowering was considerably delayed in FLC-overexpressing plants, supporting the notion that sumoylation is an important mechanism for FLC function. The data indicate that the sumoylation of FLC is critical for its role in the control of flowering time and that AtSIZ1 positively regulates FLC-mediated floral suppression.

Key words: AtSIZ1, FLC, flowering, post-translational modification, SUMO, sumoylation.

Introduction

In eukaryotic cells, protein function and stability are post-translationally regulated by small and large molecules such as phosphates, carbohydrates, lipids, and small proteins (Castro et al., 2012). The post-translational modification of target proteins by small ubiquitin-like modifier (SUMO) is an important regulatory mechanism (Wilkinson et al., 2010). The reversible covalent attachment of SUMO to a lysine residue in a target protein is catalysed by E3 SUMO ligases, although conjugation of SUMO to target proteins can occur without the help of an E3 SUMO ligase (Wilkinson et al., 2010). As in other eukaryotes, SUMO modification in plants has been implicated in numerous basic cellular processes, such as stress and defence responses, nitrogen metabolism, and the regulation of flowering (Hotson et al., 2003; Kurepa et al., 2003; Lois et al., 2003; Murtas et al., 2003; Miura et al., 2005, 2007; Catala et al., 2007; Lee et al., 2007; Conti et al., 2008; Yoo et al., 2006; Park et al., 2011).

AtSIZ1, a Siz/PIAS (SP)-RING-finger protein, regulates plant responses to nutrient deficiency and environmental stresses, and controls vegetative growth and development (Miura et al., 2005, 2007, 2010; Catala et al., 2007; Lee et al., 2007; Yoo et al., 2006; Park et al., 2011; Garcia-Dominguez
et al., 2008; Jin et al., 2008; Miura and Ohta, 2010). Due to its important roles in a wide range of physiological processes, sumoylation has been the subject of a growing number of studies in the past decades. Recently, two separate studies have identified a significant number of SUMO conjugates using proteomics methods and yeast two-hybrid screening in Arabidopsis under non-stress and stress conditions (Elrouby and Coupland, 2010; Miller et al., 2010). The results indicate that sumoylation can regulate diverse biological processes, although the functional consequences of this modification have not been fully characterized. Only a few Arabidopsis proteins, such as COOLAIR and COLDAIR, although their regulation is tightly controlled by long non-coding RNAs (Amasino, 1999). Two recent reports have shown that FLC transcription is positively regulated by RNA-binding or processing proteins such as MADS-box transcription factor flowering locus C (FLC) plays an important role in phase transition (Samach et al., 2000; Simpson and Dean, 2002). The expression of FLC is negatively regulated by vernalization and by components of the autonomous pathway (Michaels and Amasino, 1999; Sheldon et al., 1999). Vernalization-induced histone modifications are mediated by VRN1, VRN2, VRN5, and VIN3 (He and Amasino, 2005; Greb et al., 2007), leading to the repression of FLC expression. In addition, FVE, FLD, AtSWP1, and AtCZS, which participate in the autonomous pathway, modulate the histone deacetylation of FLC chromatin (He and Amasino, 2005; Krichevsky et al., 2006), repressing the transcription of the FLC gene. FLC transcription is also repressed by DNA-binding or processing proteins such as FCA, FY, FPA, FLK, and LD (Michaels and Amasino, 1999). Two recent reports have shown that FLC transcription is tightly controlled by long non-coding RNAs such as COOLAIR and COLDAIR, although their regulatory roles differ (Swiezewski et al., 2009; Heo and Sung, 2011). In addition, FLC transcription is positively regulated by FRI and EFS, an Arabidopsis PAF1 homologue (He and Amasino, 2005; Kim et al., 2005; Zhao et al., 2005). Although several factors affecting the transcription of FLC have been described, the post-translational regulation of FLC stability and function has not been clearly characterized.

A recent study has shown that FLC is polyubiquitinated by SINAT5 in vitro (Park et al., 2007), indicating that its stability may be regulated by a specific E3 ubiquitin ligase. This result suggests that the regulation of the floral transition by FLC involves a post-translational mechanism.

In the present study, it is shown that sumoylation plays a role in the regulation of flowering time by modulating the activity of FLC. AtSIZ1 stabilizes FLC through direct interaction, and it inhibits FLC sumoylation in vitro. Overexpression of mFLC, a sumoylation site mutant gene, had no effect on flowering time. These findings indicate that FLC is stabilized by the E3 SUMO ligase AtSIZ1, and FLC-mediated flowering repression is stimulated by sumoylation.

**Materials and methods**

**Plant materials and growth conditions**

The wild-type Arabidopsis thaliana plants used in this study were of the Columbia-0 (Col-0) ecotype. For plants grown in medium, seeds were surface-sterilized in commercial bleach that contained 5% sodium hypochlorite and 0.1% Triton X-100 solution for 10 min, rinsed five times in sterilized water, and stratified at 4 °C for 2 d in the dark. Seeds were planted on agar plates containing Murashige and Skoog (MS) medium, 2% sucrose, and 0.8% agar, buffered to pH 5.7. For plants grown in soil, seeds were directly sown into sterile vermiculite. All plants including seedlings were grown at 22 °C under a 16 h light/8 h dark cycle in a growth chamber.

**Construction of recombinant plasmids**

To produce His$_6$-FLC, the cdNA encoding full-length FLC was amplified by PCR and inserted into the pET28a vector (Novagen). To produce glutathione S-transferase (GST)-AtSIZ1 or its deletion mutants, the cdNAs encoding either the full length or the deletion mutants of AtSIZ1 cdNA were inserted into the pGEX4T-1 vector (Amersham Biosciences). GST–AtSIZ1 (D1), GST–AtSIZ1 (D2), and GST–AtSIZ1 (D3) contained amino acids 90–470, 300–470, and 1–100 of AtSIZ1, respectively. For the maltose-binding protein (MBP)–AtSIZ1-ahaemagglutinin (HA) fusion, a cdNA encoding full-length AtSIZ1 was amplified by PCR using a primer tagged with HA and inserted into the pMALc2 vector (New England Biolabs).

For His$_6$-FLC-Myc and GST–FLC-Myc production, cdNA encoding full-length FLC was amplified by PCR using primers tagged with Myc and inserted into pET28a and pGEX4T-1, respectively.

To produce the FLC mutant proteins GST–FLC(K5R)-Myc, GST–FLC(K135R)-Myc, GST–FLC(K154R)-Myc, and His$_6$-FLC(K154R)-Myc (the numbers indicate the positions of the lysines in FLC that were mutated to arginine), GST–FLC-Myc and His$_6$-FLC-Myc were subjected to site-directed mutagenesis using overlapping primers (Supplementary Table S1 available at JXB online). The double mutants GST–FLCm1(K5R, K135R)-Myc, GST–FLCm2(K5R, K154R)-Myc, and GST–FLCm3(K135R, K154R)-Myc were also generated by site-directed mutagenesis of GST–FLC(K5R)-Myc, GST–FLC(K135R)–Myc, and GST–FLC(K154R)-Myc using overlapping primers (Supplementary Table S1).

The Arabidopsis SUMO1 full-length cdNA was amplified by PCR with gene-specific primers and inserted into pET28a to produce the His$_6$-AtSUMO1-GG, containing full-length FLC extended with GG at the 3′ end. To produce GST–IAA4 (INDOLEACETIC ACID 4), the cdNA encoding full-length IAA4 was amplified by PCR with gene-specific primers and inserted into the pGEX4T-1 vector.

**Production of transgenic Arabidopsis plants**

To produce FLC- or mFLC (K154R)-overexpressing plants, the corresponding full-length cdNAs were amplified by PCR using a forward primer and a reverse primer tagged with FLAG3 and inserted into the pRS316 and picoplast vectors (Supplementary Table S1 at JXB online). All the constructs were verified by automatic DNA sequencing to ensure that no mutations were introduced.
5 mM dithiothreitol (DTT), 2 mM phenylmethylsulphonyl fluoride (PMSF), and a proteinase inhibitor cocktail (Roche), and purified sumoylation was performed in 30 μM TRIS-HCl (pH 7.5), 100 mM NaCl, 1% Triton X-100. Absorbed proteins were analysed as described above.

**Purification of recombinant proteins**

All of the recombinant proteins were expressed in *E. coli* strain BL21 and were purified in accordance with the manufacturer’s instructions. Briefly, for His6-AtSAE1b, His6-AtSAE2, His6-AtSCE1, His6-AtSUMO1, His6-FLC, His6-mFLC, His6-FLC-Myc, and His6-mFLC-Myc purification, bacteria were lysed in 50 mM NaH2PO4 (pH 8.0), 300 mM NaCl, 1% Triton X-100, 1 mM imidazole, 5 mM dithiothreitol (DTT), 2 mM phenylmethylsulphonyl fluoride (PMSF), and a proteinase inhibitor cocktail (Roche), and purified on Ni2+–nitrilotriacetate (Ni2+–NTA) resins (Qiagen). For GST–AtSIZ1 (GS), GST–AtSIZ1 (D1), GST–AtSIZ1 (D2), GST–AtSIZ1 (D3), GST–AtSUMO1, GST–FLC-Myc, GST–FLCm1–Myc, GST–FLCm2–Myc, GST–FLCm3–Myc, and GST–IAA4 purification, bacteria were lysed in PBS buffer (pH 7.5) containing 1% Triton X-100, 2 mM PMSF, and a proteinase inhibitor cocktail (Roche), and purified on glutathione resins (Pharmacia). For MBP–AtSIZ1 purification, bacteria were lysed in 20 mM TRIS-HCl (pH 7.4), 200 mM NaCl, 1 mM EDTA, 1% Triton X-100, and 2 mM PMSF containing a proteinase inhibitor cocktail (Roche), and purified on amylose resins (New England Biolabs). Protein concentrations were determined by the Bradford assay (Bio-Rad). For MBP–AtSIZ1-HA, bacteria were lysed in 50 mM TRIS-HCl pH 7.5, 200 mM NaCl, 1% Triton X-100, 5 mM dithiothreitol (DTT), 2 mM PMSF, and a proteinase inhibitor cocktail (Roche), and purified on amylose resins (New England Biolabs).

**In vitro binding assay**

To examine the *in vitro* binding of GST–AtSIZ1 to His6-FLC, 2 μg of full-length GST–AtSIZ1 or deletion mutant baits and 2 μg of full-length His6-FLC prey were added to 1 ml of binding buffer [50 mM TRIS-HCl (pH 7.5), 100 mM NaCl, 1% Triton X-100, 0.2% glycerol, 0.5 mM β-mercaptoethanol]. After incubation at 25 °C for 2h, the reaction mixtures were incubated with a glutathione resin for 2h before washing six times with buffer [50 mM TRIS-HCl (pH 7.5), 100 mM NaCl, 1% Triton X-100]. Absorbed proteins were analysed by 11% SDS–PAGE and detected by western blotting using an anti-His antibody (Santa Cruz Biotechnology).

To examine the dimerization of the FLC protein, 2 μg of full-length GST–FLC bait and 2 μg of full-length His6-FLC or His6-mFLC prey were added to 1 ml of binding buffer as described above. After incubation at 25 °C for 2h, the reaction mixtures were incubated with a glutathione resin and absorbed proteins were analysed as described above.

For determination of the *in vitro* binding of the FLC mutant protein His6-mFLC to MBP–AtSIZ1, 2 μg of full-length MBP–AtSIZ1 bait and 2 μg of full-length His6-mFLC or His6-mFLC prey were added to 1 ml of binding buffer as described above. After incubation at 25 °C for 2h, the reaction mixtures were incubated with an amylose resin for 2h before washing six times with buffer [50 mM TRIS-HCl (pH 7.5), 100 mM NaCl, 1% Triton X-100]. Absorbed proteins were analysed as described above.

**Sumoylation assays**

*In vitro* sumoylation was performed in 30 μl of reaction buffer [200 mM HEPES (pH 7.5), 5 mM MgCl2, 2 mM ATP] with 50 ng of His6-AtSAE1b, 50 ng of His6-AtSAE2, 50 ng of His6-AtSCE1, 8 μg of His6-AtSUMO1-GG, and 100 ng of His6-FLC-Myc (or GST–FLC-Myc) with or without 500 ng of MBP–AtSIZ1-HA. After incubation for 3h at 30 °C, the reaction mixtures were separated on 10% SDS–polyacrylamide gels. Sumoylated His6-FLC-Myc or GST–FLC-Myc was detected by western blotting using an anti-Myc antibody (Santa Cruz Biotechnology).

To identify the sumoylation site on FLC, GST–FLCm1–Myc, GST–FLCm2–Myc, GST–FLCm3–Myc, and GST–mFLC-Myc were added to the reaction mixtures instead of His6-FLC-Myc or GST–FLC-Myc, respectively. The reaction and the subsequent steps were as described above.

To confirm the identity of the sumoylated FLC band, the sumoylation reaction was performed with GST–AtSIZ1-HA instead of His6-AtSUMO1-GG under the reaction conditions described above.

**Bimolecular fluorescence complementation of AtSIZ1 and FLC**

To generate constructs for the bimolecular fluorescence complementation (BiFC) protein interaction assay, the cDNAs for AtSIZ1 and FLC were cloned into the pDONR201 vector. Next, the cDNAs for AtSIZ1 and FLC were transferred from their respective entry clones to the gateway vector pSAT4-DEST-n[174]YEFP-C1 (ABRC stock number CD3-1089) or pSAT5-DEST-c[175-end]YEFP-C1(b) (ABRC stock number CD3-1097), which contained the N-terminal 174 amino acids of enhanced yellow fluorescent protein (EYFPN) or the C-terminal 64 amino acids of EYFP (EYFP). The fusion constructs encoding nEYFP–SIZ1 and cEYFP–FLC proteins were mixed at a 1:1 ratio and co-bombarded into onion epidermal cells using a helium biostatic gun. Bombarded tissues were incubated at 25 °C in the dark for 16h and YFP signals were observed by confocal laser scanning microscopy.

**Effects of AtSIZ1 overexpression on FLC concentration in vivo**

Fourteen-day-old light-grown (16h light/8h dark) plants carrying 35S-FLC-FLAG3 and XVE-HA3-AtSIZ1 or 35S-mFLC-FLAG3 and XVE-HA3-AtSIZ1 transgenes on MS medium were treated in the light with or without β-oestradiol for 15h. Samples were grown in liquid nitrogen and lysates were separated by SDS–PAGE. FLC–FLAG3 and mFLC–FLAG3 levels were examined by western blotting with anti-FLAG antibody. HA3-AtSIZ1 induction was analysed by western blotting with anti-HA antibody. Post-translational degradation of FLC was examined using double transgenic plants of 35S-FLC-FLAG3 and XVE-HA3-AtSIZ1 or 35S-mFLC-FLAG3 and XVE-HA3-AtSIZ1. Transgenic plants were incubated in liquid medium with β-oestradiol for 15h for the induction of AtSIZ1 expression, washed, and then transferred to MS medium with 100 μM cycloheximide (CHX). Treated plants were then incubated for 4h. Proteins were extracted at the indicated time points and analysed by western blotting using anti-HA or anti-FLAG antibodies as described above.

**Investigation of flowering time**

To examine the effect of sumoylation on FLC-mediated flowering, transgenic plants carrying 35S-FLC-FLAG3 or 35S-mFLC-FLAG3 were generated. After selection of FLC–FLAG3- or mFLC–FLAG3-overexpressing transgenic plants, wild-type (WT) and transgenic plants were grown in soil under long-day conditions (16h light/8h dark). Flowering time was assessed by counting the number of rosette leaves present at the time of appearance of inflorescences or was also determined by counting the days to flowering.

**Yeast two-hybrid assays**

Yeast two-hybrid assay was performed using the GAL4-based two-hybrid system (Clontech). Full-length AtSIZ1 and IAA4 cDNAs were cloned into pGAD424 and pGBT8 (Clontech) to generate the constructs AD-AtSIZ1 and BD-IAA4. The constructs were
transformed into the yeast strain AH109 with the lithium acetate method. The yeast cells were grown on minimal medium (–Leu/–Trp). Transformants were plated onto minimal medium (–Leu/–Trp/–His) to test the interactions between AtSIZ1 and IAA4.

Results
AtSIZ1 physically interacts with FLC

It was recently reported that FLC directly interacts and co-localizes with the Arabidopsis E3 ubiquitin ligase SINAT5 in the nucleus (Park et al., 2007). Since the SP-RING motif protein AtSIZ1 also localizes to the nucleus (Miura et al., 2005), the possible physical interaction between AtSIZ1 and FLC was examined using a BiFC assay system. Arabidopsis FLC tagged with the C-terminus of EYFP and AtSIZ1 tagged with the N-terminus of EYFP were transiently expressed in onion epidermal cells. It is not known to what extent onion cells reflect the situation in Arabidopsis cells. Nevertheless, yellow fluorescence was detected (Fig. 1), indicating the direct interaction of these proteins in vivo. To confirm the interaction between FLC and AtSIZ1 in an in vitro system, pull-down assays were performed by overexpressing the recombinant proteins in E. coli and purifying them with affinity columns (Fig. 2B). Figure 2C shows that GST–AtSIZ1, but not GST alone, was able to pull down Arabidopsis FLC. Experiments using deletion mutants showed that the N-terminal region containing the SAP domain of AtSIZ1 [GST–AtSIZ1 (D3)] interacts with FLC (Fig. 2C). Therefore, these in vitro results suggest that the co-localization of FLC and AtSIZ1 in the nucleus probably reflects their direct interaction in vivo.

FLC is sumoylated without AtSIZ1

The direct interaction of FLC and AtSIZ1 indicated by the in vivo and in vitro results led to the hypothesis that AtSIZ1 may function as an E3 SUMO ligase for FLC. Therefore, the recombinant proteins GST–AtSIZ1-HA3 and His6-FLC-Myc were produced to determine whether AtSIZ1 is the E3 SUMO ligase for FLC. In the in vitro sumoylation experiments, purified His6-FLC-Myc was sumoylated in the presence of E1 and E2 activities (Fig. 3A). However, the sumoylation of His6-FLC-Myc was not induced by AtSIZ1. It was also tested whether another AtSIZ1-interacting protein, IAA4, could be sumoylated by AtSIZ1 (Fig. 3B). The result showed that IAA4 was not sumoylated under the reaction conditions employed, including the presence of E1, E2, and E3 (Fig. 3C).

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Fig. 1. FLC interacts with AtSIZ1 in vivo. The interaction between AtSIZ1 and FLC was examined by a bimolecular fluorescence complementation (BiFC) assay in onion epidermal cells. AtSIZ1 and FLC cDNAs were fused with YFP at the N-terminal (N) and C-terminal (C) ends, respectively. Each combination of YFP(N)/35S–YFP(C)-FLC, 35S-YFP(N)–AtSIZ1/YFP(C), and 35S-YFP(C)–FLC/35S-YFP(N)–AtSIZ1 was introduced into onion epidermal cells by particle bombardment, and fluorescence signals were detected by confocal microscopy. Bar=50 μm.
Regulation of FLC function by SUMO conjugation

AtSIZ1 inhibits FLC sumoylation

Despite the interaction between AtSIZ1 and FLC shown in Figs 1 and 2, the results indicate that AtSIZ1 has no E3 SUMO ligase activity for FLC (Fig. 3A). Therefore, experiments were carried out to examine whether AtSIZ1 could block or inhibit the sumoylation of FLC. The addition of increasing amounts of AtSIZ1 protein to the reaction mixture resulted in the gradual inhibition of FLC sumoylation (Fig. 4A). However, AtSIZ1 was sumoylated under the reaction conditions used here (Fig. 4A), indicating that AtSIZ1 is active and that it has self-sumoylation activity under the reaction conditions used. Since all purified proteins used in this experiment were dialysed prior to the reaction, to confirm the effect of AtSIZ1 on FLC sumoylation, an equal volume of dialysis buffer was added to the reactions; this buffer had no effect on FLC sumoylation (Fig. 4B). Therefore, these results

Fig. 2. Interaction of AtSIZ1 with FLC. (A) Schematic diagram of bait [GST (G), GST–AtSIZ1 (GS), GST–AtSIZ1 (D1), GST–AtSIZ1 (D2), and GST–AtSIZ1 (D3)] and prey (His6–FLC) proteins. In vitro pull-down of FLC with AtSIZ1. (B) His6–FLC, full-length AtSIZ1, or its deletion mutants were overexpressed in *E. coli* and purified with Ni²⁺–NTA or glutathione affinity columns. (C) The His6–FLC protein was pulled down with full-length AtSIZ1 or its deletion mutant proteins, separated on 11% SDS–polyacrylamide gels, and analysed by western blotting with an anti-His antibody. I, input (His6–FLC).

Fig. 3. FLC is sumoylated in vitro. (A) *Arabidopsis* His6–AtSAE1b, His6–AtSAE2, His6–AtSCE1, MBP–AtSIZ1, His6–AtSUMO1, and His6–FLC-Myc were overexpressed in *E. coli* and purified with Ni²⁺–NTA, glutathione, and amylose affinity columns, respectively. Sumoylation of His6–FLC-Myc was assayed in the presence or absence of E1 (His6–AtSAE1b and His6–AtSAE2), E2 (His6–AtSCE1), E3 (MBP–AtSIZ1), and His6–AtSUMO1. After the reaction, sumoylated FLC was detected by western blotting with an anti-Myc antibody. GST–IAA4 was also used for the sumoylation assay as a negative control. (B) AtSIZ1 directly interacts with GST–IAA4 in yeast. Full-length AtSIZ1 and IAA4cDNAs were fused to sequences encoding the Gal4 activation domain (AD) and the Gal4 DNA-binding domain (BD) in pGAD424 and pGBT8, respectively. The constructs were transformed into yeast strain AH109. Each number indicates the yeast cells transformed with a combination of only pGAD424 and pGBT8 vectors or recombinant plasmids. Transformants were plated onto minimal medium –Leu–Trp or –Leu–Trp–His and incubated for 4 d. (C) Sumoylation of GST–IAA4 was assayed using the same reaction conditions as above. After the reaction, IAA4 was detected by western blotting with an anti-GST antibody.
Son et al. indicate that FLC sumoylation is blocked by the AtSIZ1 protein.

**Identification of sumoylation sites on FLC**

The deduced amino acid sequences of FLC showed three putative sumoylation sites (ΨKXE) located at Lys5 (K5), Lys135 (K135), and Lys154 (K154; Fig. 5A, B). To identify the sumoylation sites on the FLC protein, single or double mutant derivatives were generated with the mutations K154R, K5R/K135R, K5R/K154R, and K135R/K154R. The proteins were overexpressed in E. coli, purified with glutathione affinity columns, and used for in vitro sumoylation assays. In vitro sumoylation with the double mutant proteins GST–FLCm1-Myc (K5R/K135R), GST–FLCm2-Myc (K5R/K154R), GST–FLCm3-Myc (K135R/K154R), and GST–mFLC-Myc (K154R) showed that GST–FLCm1-Myc was sumoylated, whereas GST–FLCm2-Myc, GST–FLCm3-Myc, and GST–mFLC-Myc were not (Fig. 5C). In vitro sumoylation assays including the single mutant protein GST–mFLC-Myc (R) showed that this protein was not modified with SUMO (Fig. 5D), indicating that K154 is the principal site of SUMO conjugation on FLC.

**FLC is stabilized by AtSIZ1**

The AtSIZ1–FLC interaction and the inhibition of FLC sumoylation by AtSIZ1 imply that the concentration of FLC may be regulated by the amount of AtSIZ1 present in vivo. FLC concentrations were therefore measured in transgenic plants carrying a 35S-FLC-FLAG3 transgene and an oestradiol-inducible XVE-HA3-AtSIZ1 transgene. Induction of the expression of AtSIZ1 increased the FLC concentrations up to 1.5- and 3.3-fold in two independent transgenic plants, respectively (Fig. 6A). However, the two independent transgenic plants carrying a 35S-mFLC-FLAG3 transgene and an oestradiol-inducible XVE-HA3-AtSIZ1 transgene showed no changes in mFLC concentration in response to AtSIZ1 induction (Fig. 6B). It may be possible that the transcript levels of FLC or mFLC can affect the levels of FLC and mFLC proteins in transgenic plants. Thus FLC and mFLC transcript levels were examined by real-time reverse transcription–PCR (RT–PCR) and quantitative real-time RT–PCR after induction of AtSIZ1 in FLC- or mFLC-overexpressing double transgenic plants. The result showed that the transcript levels of FLC and mFLC were comparable under these conditions (Fig. 6A, B; Supplementary Fig. S1 at JXB online).

The effect of AtSIZ1 on FLC decay was examined by treating the transgenic plants described above with CHX to block new protein synthesis. The results showed that the degradation of FLC was delayed in plants co-expressing AtSIZ1 (Fig. 6C, E). However, the rate of decay of mFLC was not significantly altered by the expression of AtSIZ1 (Fig. 6D, F).

**FLC modification by SUMO is necessary for flowering repression**

FLC overexpression causes late flowering, and FLC mutants are characterized by early flowering in Arabidopsis (Sanda and Amasino, 1996). Based on these known effects of FLC and the present sumoylation data, the effect of sumoylation on the activity of FLC as a repressor of the transition to flowering was next examined. FLC- and mFLC-overexpressing transgenic Arabidopsis plants were generated using 35S-FLC-FLAG3 and 35S-mFLC-FLAG3 constructs, respectively. After selecting homozygous lines (Supplementary Fig. S2 at JXB online), the recombinant protein levels of FLC-FLAG3 and 35S-mFLC-FLAG3 were first examined and then the transgenic plants were investigated for vegetative growth and flowering time (Fig. 7A, B). The relative flowering time of each transgenic plant was assessed by counting the number of rosette leaves. The number of rosette leaves in WT plants was 14.75 ± 0.71, and that of mFLC-overexpressing plants was 14.63 ± 1.16, which was comparable with that of the WT. However, in FLC-overexpressing plants, the number of days before the appearance of inflorescences was 14.75 ± 0.71, and that of mFLC-overexpressing plants was 14.63 ± 1.16, which was comparable with that of the WT. However, in FLC-overexpressing plants, the number of days...
before the appearance of inflorescences was $52.31 \pm 1.57$, which represented an ~1.85-fold increase (Fig. 7D). As a result, the flowering time was significantly delayed in FLC-overexpressing Arabidopsis plants, while no changes were detected in mFLC-overexpressing plants (Fig. 7C, D). However, vegetative growth was not affected in FLC- or mFLC-overexpressing plants (Fig. 7A), suggesting that sumoylation is an important modification for the regulation of FLC function.

**Mutant FLC can interact with AtSIZ1 and FLC**

The observation that AtSIZ1 stabilizes FLC but not mFLC suggests that mFLC does not interact with AtSIZ1. Therefore, the possible interaction between AtSIZ1 and mFLC was examined using an in vitro pull-down assay. His$_6$-FLC, His$_6$-mFLC, and full-length MBP–AtSIZ1 were purified with N$t^{2+}$-NTA or glutathione affinity columns and it was determined whether or not His$_6$-FLC or His$_6$-mFLC proteins could be pulled down with AtSIZ1. The mutant proteins mFLC, FLCm1, FLCm2, and FLCm3 have amino acid substitutions at residues that are predicted to be SUMO conjugation sites in FLC, as indicated. After the reaction, sumoylated FLC protein was detected by western blotting with an anti-Myc antibody.

![Fig. 5. In vitro identification of the sumoylation site on FLC. (A) Deduced amino acid sequences of the FLC protein. Three putative sumoylation sites (ΨKXE) identified using the SUMOplot Analysis Program are indicated in bold type. (B) Schematic diagram of the recombinant GST–FLC-Myc protein. The MADS-box and putative sumoylation sites (K5, K135, and K154) are indicated. (C and D) In vitro sumoylation assays. Recombinant GST–FLC-Myc, GST–mFLC-Myc, GST–FLCm1-Myc, GST–FLCm2-Myc, and GST–FLCm3-Myc were overexpressed in E. coli and purified using a glutathione affinity column. The reaction mixture contained E1 (His$_6$-AtSAE1b and His$_6$-AtSAE2), E2 (His$_6$-AtSCE1), E3 (GST–AtSIZ1), and His$_6$-AtSUMO1 without (−) or with (+) a substrate protein. The mutant proteins mFLC, FLCm1, FLCm2, and FLCm3 have amino acid substitutions at residues that are predicted to be SUMO conjugation sites in FLC, as indicated. After the reaction, sumoylated FLC protein was detected by western blotting with an anti-Myc antibody.](image-url)
Fig. 6. FLC is stabilized by AtSIZ1 in vivo. Double transgenic plants of 35S-FLC-FLAG, and XVE-HA3-AtSIZ1 (A) or 35S-mFLC (K154R)-FLAG, and XVE-HA3-AtSIZ1 (B) were incubated in liquid medium with β-oestradiol for the induction of AtSIZ1 expression. After incubation for 15 h, HA3-AtSIZ1, FLC-FLAG, and mFLC-FLAG levels were assessed by western blotting with anti-HA or anti-FLAG antibodies. Tubulin was used as a loading control. Numbers under lanes indicate relative intensities. Protein levels were normalized to a value of 1.00 for FLC or mFLC levels in the ‘−’ inducer in both panels. RNA concentrations for FLC-FLAG and mFLC-FLAG were determined by real-time RT–PCR using a FLAG primer and a gene-specific primer. For HA3-AtSIZ1, RNA concentration was measured by real-time RT–PCR using an HA primer and a gene-specific primer. Tubulin RNA was used as a loading control. To assess the degradation of FLC, double transgenic plants of 35S-FLC-FLAG, and XVE-HA3-AtSIZ1 (C) or 35S-mFLC (K154R)-FLAG, and XVE-HA3-AtSIZ1 (D) were incubated in liquid medium with β-oestradiol for the induction of AtSIZ1 expression, washed, and transferred to MS medium with 100 μM cycloheximide (CHX). At the indicated times, protein was extracted and analysed by western blotting with anti-HA or anti-FLAG antibodies. Tubulin was used as a loading control. FLC or mFLC levels during degradation were also expressed in graph form. The relative protein levels of FLC (E) or mFLC (F) were normalized to numerical values based on a value of 1.0 for the protein levels at 0 h using the data shown in both C and D. Open circles indicate FLC (or mFLC) with AtSIZ1 and filled circles indicate FLC (or mFLC) without AtSIZ1.
or His₆-mFLC proteins could be pulled down with GST or GST–FLC proteins was examined. As shown in Fig. 9, GST–FLC formed a complex with both His₆-FLC and His₆-mFLC.

**Discussion**

In the present study, it was shown that FLC-mediated flowering repression is activated by sumoylation and that AtSIZ1 stabilizes FLC.

Eukaryotic cells express SP-RING finger proteins, SAP and Miz-finger domain (Siz) proteins, and protein inhibitor of activated STAT (PIAS) proteins (Hochstrasser, 2001). Recently, SIZ1-type proteins with a SP-RING domain were also identified in plants and were shown to be involved in diverse biological processes (Ishida et al., 2012; Novatchkova et al., 2012).

The function and stability of transcription factors are modulated by various post-translational modifications. The conjugation of SUMO (a protein modifier) to a target protein regulates its function and stability. FLC is modified by ubiquitin (Park et al., 2007), indicating that other post-translational modifications, such as sumoylation, may play a role in the regulation of FLC activity. Experiments were therefore carried out to examine whether AtSIZ1 has E3 SUMO ligase activity for FLC. The results of pull-down and BiFC assays showed a strong interaction between FLC and AtSIZ1 (Fig. 1), and in vitro sumoylation assays showed that FLC is modified by SUMO (Fig. 3). However, the results showed that
the attachment of SUMO to FLC occurred independently of AtSIZ1 in vitro (Figs 3, 4).

The covalent attachment of SUMO to a lysine residue in the target protein is generally mediated by E3 SUMO ligases. However, direct transfer from the SUMO-conjugating enzyme Ubc9 can occur through at least two ligase-independent mechanisms. First, Ubc9 can directly recognize the sumoylation motif $\Psi$-K-x-[D/E] ($\Psi$, an aliphatic branched amino acid; x, any amino acid) and conjugate the lysine residue (Bernier-Villamor et al., 2002). Secondly, some SUMO substrates contain SUMO-interacting motifs (SIMs) that promote their own conjugation (Meulmeester et al., 2008; Zhu et al., 2008). These SIMs bind to the SUMO moiety to which Ubc9 is attached, thereby increasing its local concentration and facilitating sumoylation. The results of the present study indicate that FLC is sumoylated by one of these mechanisms in the absence of an E3 SUMO ligase.

Since FLC sumoylation was inhibited by AtSIZ1 (Fig. 4), the mechanisms underlying the binding of AtSIZ1 to FLC and its effect on FLC activity and stability were further examined. For this purpose, double transgenic Arabidopsis plants were generated through transformation with a 35S-FLC-FLAG3 transgene and an oestradiol-inducible XVE-HA 3-AtSIZ1 transgene to examine the effect of AtSIZ1 on the stability of FLC. AtSIZ1 induction with oestradiol increased the concentration of FLC but not that of mFLC (Fig. 6A, B). Furthermore, AtSIZ1 overexpression retarded the degradation of FLC, whereas that of mFLC was not affected (Fig. 6C, D). To confirm these results, the biological effect of AtSIZ1 on FLC and mFLC function and stability is also currently being investigated using double transgenic plants that constitutively overexpress AtSIZ1 and FLC or mFLC.

In any case, based on the present findings, these data suggest that AtSIZ1 stabilizes FLC through direct binding to FLC before or after FLC sumoylation in vivo (Supplementary Fig. S3 at JXB online). Furthermore, the inhibitory effect of AtSIZ1 on FLC sumoylation suggests the possible existence of another E3 SUMO ligase for FLC in Arabidopsis (Supplementary Fig. S3).

However, there may be many factors affecting FLC conjugation with SUMO in vivo. For example, in vivo concentrations of proteins comprising the sumoylation machinery, including Arabidopsis SUMO-activating enzyme E1 (SAE1+2) and conjugating enzyme E2 (AtUBC9), AtSUMO, and AtSIZ1, may differ from the concentrations of the proteins used in the in vitro system used here, and the expression of each of these components may vary according to developmental stage, thereby affecting FLC sumoylation. In addition, AtSIZ1 can form complexes with various proteins in vivo (Novatchkova et al., 2012), which affects AtSIZ1 conformation and activity, and, thus, FLC sumoylation. In addition, the timing and localization of FLC expression can also be controlled by changes in chromatin structure through histone modifications and DNA methylation (He, 2012). FLC can form complexes with other proteins as well, which can lead to changes in FLC concentration and conformation, thereby leading to increases or decreases in the sumoylation of this protein. Therefore, the possibility that AtSIZ1 enhances FLC sumoylation as an E3 SUMO ligase in vivo still cannot be ruled out.

Since FLC is a central regulator of flowering, extensive research has been conducted to elucidate the mechanisms regulating FLC expression at the transcriptional and post-transcriptional levels in association with flowering time (He and Amasino, 2005; Kim et al., 2005; Zhao et al., 2005; Krichevsky et al., 2006; Greb et al., 2007; Park et al., 2007; Swiezewski et al., 2009; Heo and Sung, 2011). In the present study, the role of FLC in the transition to flowering...
was examining using the sumoylation site mutant mFLC. To characterize the function of FLC in the control of flowering time, FLC- or mFLC-overexpressing transgenic Arabidopsis plants were generated and their flowering time was examined by counting the number of rosette leaves. FLC overexpression delayed flowering, whereas mFLC overexpression had no notable effect on flowering time (Fig. 7A, B), indicating that sumoylation is critical for FLC to exert its floral repressor function.

The lack of an effect of mFLC overexpression on flowering time may have resulted from an impaired interaction of mFLC with AtSIZ1 or a defect in complex formation with FLC. However, in vitro pull-down analysis showed that mFLC interacted with AtSIZ1 and with FLC. From these results, several possible mechanisms explaining why mFLC overexpression does not affect flowering time are proposed. First, sumoylation of the FLC protein may be necessary for its activation. As mFLC cannot be modified with SUMO, this protein may not have an effect on flowering time despite its overexpression. Secondly, mFLC may inactivate endogenous FLC. Transgenic mFLC may form a complex with endogenous FLC and act in a dominant-negative form. Thus, a possible reason for the observation that flowering time in mFLC-overexpressing plants is comparable with that of WT plants is that the FLC level is originally low in WT plants, although this protein could be scavenged by the overexpressed mFLC through complex formation.

It is believed that if sumoylated FLC can be detected in vivo, it may also be possible to find an answer for why FLC overexpression delayed flowering, whereas mFLC overexpression had no effect on flowering time. However, to date, it has not been possible to detect sumoylated protein in vivo, perhaps due to its low level or presence at specific stages. Recently, Robertson et al. (2008) showed that endogenous FLC can be detected by western blot analysis with anti-FLC antibody, but the FLC band intensities were quite weak, even in C24 WT plants. It is well known that FLC protein levels are much lower in the Col background than in the C24 background. Thus, there appear to be specific challenges in detecting FLC in the Col background using antibodies. Production of a specific anti-FLC antibody which works well in vivo will be a solution.

DET1 (De-etiolated 1), a SINAT5-interacting partner, blocks the ubiquitination of LHY (Long Hypocotyl) by SINAT5 through direct interaction with SINAT5 (Park et al., 2007). The present data show that AtSIZ1 inhibits the sumoylation of FLC through direct interaction with SINAT5 (Fig. 4). However, AtSIZ1 increased the level of FLC in transgenic plants (Fig. 6A). Furthermore, the degradation of FLC was delayed in the presence of AtSIZ1 (Fig. 6C). These findings suggest that direct binding of AtSIZ1 to FLC protects the protein from degradation induced by its ubiquitination by SINAT5, as shown for DET1, which blocks the ubiquitination of LHY by SINAT5. AtSIZ1 may thus have a protective effect on FLC by antagonizing its ubiquitination (Supplementary Fig. S3 at JXB online).

In conclusion, the present results indicate that AtSIZ1 controls the stability of FLC by directly binding to FLC, but not through its E3 SUMO ligase activity, and that the FLC-mediated floral transition is negatively regulated by SUMO conjugation. In addition, it was shown that proteolytic turnover of flowering-associated proteins can be regulated by sumoylation. The biochemical mechanisms underlying the regulation of FLC function and stability by sumoylation were also elucidated. Together with previous findings, the data suggest that both of the post-translational modification systems, ubiquitination and sumoylation, can regulate flowering by direct modulation of FLC stability and activity.

**Supplementary data**

Supplementary data are available at JXB online.

- Figure S1. The effect of AtSIZ1 on FLC transcript levels.
- Figure S2. Selection of FLC- and mFLC-overexpressing plants.
- Figure S3. Possible regulatory modes of FLC stability.
- Table S1. List of primers used for this study.

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