

# Characterization and Heterologous Expression of *SLF*, a Functional Homolog of the Floral Regulator *LEAFY/FLORICAULA* from *Salix discolor*

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### ABSTRACT

This study was done to contribute to our understanding of the *LFY/FLO* gene activity in willow, a dioecious woody plant. *SLF* is the *Salix discolor* homolog of the *LFY/FLO* gene which was cloned from the reproductive buds of a male individual, clone S365. *In situ* hybridization revealed that *SLF* is strongly expressed in the inflorescence meristems, bracts, and floral meristems, but only weakly expressed in the vegetative meristems and leaf primordia. Since a genetic transformation system coupled with *in vitro* regeneration is currently not available for willow, *Arabidopsis thaliana* was used to analyze the functions of *SLF*. Transformed *A. thaliana* produced flowers more than two weeks earlier than the controls; furthermore, they produced terminal and solitary flowers instead of inflorescence branches. The phenotypes of the transgenic lines were dominant and heritable, demonstrating that *SLF* was functional and participated in the flowering of *A. thaliana*. Many of these phenotypes are being described for the first time from a *LFY/FLO* homolog from a dioecious plant. Complementation test showed that *SLF* was able to restore the wild-type phenotype of the *lfy-6* mutant. This study revealed that *SLF* affected various aspects of floral development in transgenic *A. thaliana* and therefore, suggested that *SLF* is the functional homolog of the *LFY/FLO* gene.

Keywords: Arabidopsis thaliana, flowering, leafy, willow, woody plant

# INTRODUCTION

The switch from vegetative to reproductive stage is one of the remarkable processes in the development of plants. Of the genes that are involved in this event, the LEAFY/ FLORICAULA (LFY/FLO) gene performs one of the most crucial functions. Therefore, functional homologs of this master regulatory gene have been isolated and characterized from a variety of plants ranging from bryophytes to flowering plants (Himi *et al.* 2001; Carlsbecker *et al.* 2004; Dornelas and Rodriguez 2005; Tanahashi et al. 2005; Dornelas et al. 2006; Allnutt et al. 2007). The homologs from seed plants have been almost exclusively obtained from bisexual species, which are either herbaceous (Kelly et al. 1995; Hofer et al. 1997; Bomblies et al. 2003; Busch and Gleissberg 2003; Allnutt et al. 2007) or woody (Southerton et al. 1998; Rottman et al. 2000; Walton et al. 2001; Wada et al. 2002; Carmona et al. 2002; Carlsbecker et al. 2004; Dornelas and Rodriguez 2005; Dornelas et al. 2006). To date, the only LFY/FLO homolog described from a dioecious plant is PTLF from Populus trichocarpa (Rottmann et al. 2000). Therefore, there is a need for more studies in this area to expand our knowledge of the activities of the LFY/ FLO gene in this type of plants.

Although willow and poplar belong to the same family (*Salicaceae*), there are marked differences in the morphology and development of their reproductive structures (Zhang and Fernando 2005; Fernando and Zhang 2006). A willow flower is typically composed of a nectary and two anthers or a single ovary that arises from the axil of the bract; the unisexual flowers are clustered in an upright catkin (Zhang and Fernando 2005). Willow flowers are initiated one to two years after seed germination (Gullberg 1993; Taylor 2002). In the flowers of poplars, a periant cup exists,

there are three carpels or 40 to 60 anthers per flower, flowers are clustered in a hanging catkin, and flowering generally occurs after five years (Boes and Strauss 1994). These morphological and developmental differences suggest that there may also be some differences at the molecular level as regards the mechanism of flowering in willow and poplar.

Except for our report on *SAP1*, the *Salix discolor* homolog of the *AP1* gene (Fernando and Zhang 2006), there is no other information currently available on the pattern of gene expression during the initiation of the inflorescences and flowers in willow. The present study is based on a male plant and therefore, represents the first step towards our understanding of the overall genetic and molecular mechanisms of flowering in willow. This study aims to: 1) isolate the homolog of the *LFY/FLO* gene from a male willow individual, 2) analyze the sequence and structure of the *SLF* gene, 3) characterize the temporal and spatial expression patterns of *SLF*, and 4) analyze the functions of *SLF* through over-expression using a heterologous host, *Arabidopsis thaliana*.

# MATERIALS AND METHODS

#### Plant materials and growth conditions

Twigs of *S. discolor* (clone S365) bearing vegetative and male reproductive buds at various developmental stages were collected from late July to early August in 2003 and 2004. The collections were done from SUNY-ESF's Genetics Field Station in Tully, NY. The vegetative and reproductive bud scales were removed, the buds were either immediately frozen in liquid nitrogen, and stored at -80°C for RNA isolation, or fixed in 4% (v/v) paraformaldehyde in 1X phosphate buffered saline (PBS) buffer for *in situ* hybridization. Young leaves were also collected from the field in July and August 2004, and frozen and stored as above for DNA extraction. Wild-type *A. thaliana* (ecotype Columbia) and 35S::*SLF A. thaliana* transgenic lines were sown on an enriched potting mix (Miracle-Gro, Marysville, OH). The seeds were grown in the greenhouse at 22°C with 16 h of light exposure. The plants were illuminated with a mixture of cool white and plant growth fluorescent lamps.

#### Isolation, cloning, and sequencing

Total RNA was isolated from 0.3 g reproductive buds at inflorescence meristem stage using an improved RNA isolation method by Salzman et al. (1999). Total RNA was resuspended in 25 µL of diethyl pyrocarbonate (DEPC)-treated water. First-strand cDNA was obtained through reverse transcription PCR using CLON-TECH SMART PCR cDNA Synthesis Kit (Clontech, San Jose, CA). A pair of degenerate primers (LFY101 and LFY306) specific to the two highly conserved regions of various LFY/FLO homologs (Coen et al. 1990; Weigel et al. 1992; Southerton et al. 1998; Rottmann et al. 2000) was designed (Table 1). Gradient PCR was done and a 551-bp PCR product was obtained and cloned into pCR2.1-TOPO vector using TOPO TA Cloning Kit (Invitrogen, Carlsbad, CA). DNA sequencing was done at the BioResource Center, Cornell University, Ithaca, NY. The DNA sequence obtained was analyzed by submission to BLAST (http://www.ncbi. nlm.nih.gov/BLAST/) followed by manual inspection of sequences. To obtain the complete coding sequence of SLF, rapid amplification of cDNA ends (RACE) technique was used. Two genespecific primers (SLF213GSP1 and SLF165GSP2) were designed based on the partial sequence of SLF (Table 1). 5'-RACE and 3'-RACE PCR products were obtained using SMART RACE cDNA Amplification Kit (BD Biosciences Clontech, Palo Alto, CA). The PCR product was sequenced and analyzed as above. The fulllength cDNA was amplified by PCR with primers 5SLF1 and 3SLF1 (Table 1) using the first strand cDNA as the template. To confirm the sequence of the full-length cDNA, DNA sequencing was repeated from five different pure colonies.

#### SLF genome structure analysis

To identify the introns in *SLF*, a 2.2 kb DNA fragment was obtained by PCR with primers 5SLF1 and 3SLF1 (**Table 1**) using genomic DNA as the template. The amplified product was cloned into pCR2.1-TOPO vector and sequenced separately with primers 5SLF1, 3SLF1, SLF213GSP1 and SLF165GSP2 (**Table 1**). The sequences were analyzed and compared with the full-length cDNA to determine the location and length of the introns in the *SLF* genome structure.

#### Sequence alignment

The coding sequence of *SLF* was aligned with the nucleotide sequences of *LFY/FLO* homologs from several species. The alignments were done through BLAST. Multiple alignments based on deduced amino acid sequence were done using UNIPROT (http:// www.pir.uniprot.org/search/blast.shtml).

#### **Constructs and plant transformation**

The primers 5SLF3 and 3SLF3 (**Table 1**) containing *Xho*I restriction enzyme site were used to amplify *SLF* for integration into the pCAMBIA3300 construct. The primers 5SLF3 and TER1 (**Table 1**) were used to confirm the orientation of *SLF* in the pCGSLF construct. Another construct (pCGFP) containing only *mgfp5-er* driven by the CaMV 35S promoter was made and used as the negative control for plant transformation. The plasmid DNAs were introduced into *Agrobacterium tumefaciens* strain EHA105 by CELL-Porator Electroporation System (GibcoBRL, Gaithersburg, MD). *A. tumefaciens*-mediated transformation of *A. thaliana* was performed by floral dip technique according to Clough and Bent (1998) and Desfeux *et al.* (2000). T<sub>1</sub> seeds on a moist filter paper were screened under a fluorescence microscope (Leica DMLB, San Jose, CA) based on GFP expression. GFP expressing seeds were sown in potting mix and grown in the greenhouse under

long-day condition. All organs including leaves, roots, and flowers were further screened through GFP expression.  $T_2$  and  $T_3$  seeds were screened from the transgenic lines based on GFP expression. To further examine the function of *SLF*, pollen grains from homozygous  $T_3$  lines were crossed with the strong *lfy-6* mutant allele. Early-formed flowers of *lfy-6* have pistils but do not have petals and stamens, while late-formed flowers are normal. Therefore, early-formed flowers were pollinated and other flowers that developed later were cut out. The resulting seeds were harvested and grown to observe the floral phenotypes. In  $F_2$  individuals, the number of plants showing normal and *lfy-6* phenotypes was counted. The ratio of these two floral phenotypes was calculated and compared with Mendel's segregation patterns.

#### **DNA isolation and Southern analysis**

Genomic DNA was extracted from young leaves of S. discolor using DNeasy Plant Mini Kit (Qiagen, Valencia, CA). For Southern blot analysis, the genomic DNA was purified three times with phenol/chloroform and digested separately with EcoRI and HindIII at 37°C for 15 h. Approximately 10 µg of the digested DNA was subjected to electrophoresis through 0.8% agarose gel and blotted to positively charged nylon membranes (Roche, Indianapolis, IN) (Sambrook et al. 1989). The primers RNSLF3 and RNSLF4 (Table 1) were designed to amplify a 500-bp DNA fragment (Fig. 1A), which is one of the highly conserved regions of SLF (Fig. 1B). The amplified PCR product was used as a probe and labeled with digoxigenin-11-UTP using DIG High Prime DNA Labeling and Detection Starter Kit I (Roche, Indianapolis, IN). Hybridization and immunological detection were done according to the manufacturer. The membranes were incubated with gentle agitation in the probe/hybridization solution at 40°C for 12 h. Stringency washes with  $0.5 \times SSC/0.1\%$  SDS were performed after hybridization. Anti-digoxigenin-alkaline phosphatase (AP) was used as the antibody. Nitro blue tetrazolium/5-bromo-4chloro-3-indolyl phosphate (NBT/BCIP) color substrate solution was used to visualize the hybridization on the membrane. The membrane was scanned using the GS-710 Calibrated Imaging Densitometer (Bio-Rad, Hercules, CA).

#### In situ hybridization

The 500 bp RNSLF3/RNSLF4 PCR product (which is the same fragment used as probe in Southern blot) was cloned into pSPT18 and pSPT19 using the DIG RNA Labeling Kit (Roche, Indianapolis, IN). The digoxigenin-labeled antisense and sense RNA probes were obtained with T7 and SP6 RNA polymerases, respectively. Vegetative and reproductive buds of S. discolor were fixed in 4% (v/v) paraformaldehyde in 1 × PBS buffer (pH 7), dehydrated through a graded series of ethanol, infiltrated, and embedded in paraffin. Sectioning and slide pretreatment, in situ hybridization, and post-hybridization were performed as described by Jackson (1991). Using thin sections from 6 different buds, RNA in situ hybridizations with nonradioactive probes were incubated at 53°C for 18 h. A series of washes with  $0.2 \times SSC$  and NTE were performed after hybridization. The samples were treated with antidigoxigenin-AP antibody and NBT/BCIP color detection was performed in the dark for 2-3 days. The slides were examined under a light microscope (Leica DMLB, San Jose, CA) and representative images were obtained using a DEI-7500 CE digital video camera (Optronics, Goleta, CA).

#### Northern analysis

Total RNA was extracted from leaves of six 35S::*SLF A. thaliana* transgenic lines and one transgenic control line (sans the *SLF* gene) using RNeasy Plant Mini Kit (Qiagen, Valencia, CA). Twenty  $\mu$ g of total RNA for each sample was subjected to electrophoresis through 1.2% formaldehyde agarose gel. Total RNA was blotted on positively charged nylon membranes (Roche, Indianapolis, IN). The 500-bp PCR product amplified from *SLF* cDNA, which is the same fragment used in Southern blot, was used as a probe and labeled with digoxigenin-11-UTP using DIG High Prime DNA Labeling and Detection Starter Kit I (Roche, Indianapolis, IN). Hybridization and immunological detection were done



В

PRFLL

NEEDLY

SLF EAUTAS UTKWDTRAMVPHPNR-LLEMVAPPO-----OPPAAAFAVRPRE-LCGLEELFOAYGIRY 61 -EARTAS - INFAMETRAMVPHERAL BELEVERTY -EARTAS - INFAMETRAMVPHERAL LEMVPPPQ-----QPPAAAFAVRPRE------LCGLEELFQAYGIRY -DARTAS - INFAMETRAMVPHERALLEAVAPPPGAAAPAPVAAAYAIRPRE------LGGLEDLFQE -DARSAN - IFKWDIRGMVVPTNRVQLEAAVPPAAT--AGGAAAGYTLRPSREL----CLGGLEELFQAYGVRY -DARSAN - LFKWDIRTALPQPNR-LLDAVAPPPP----PPQAPSYSMRPRE------CLGGLEELFQAYGIRY PTLF **I**DP 61 VFL 67 AFL1 68 **IDP** FLO 60 ſDΡ MDP-EGFTSG-LERWNPTRALVQAPP-----PVPPPLQQQPVTPQTAAFGMR-----LGGLEGLFGPYGIRE MDPNDAFSAAHPSRWDLGPPAHAAPA-----PAPPP------PPLAPLLLPPHAP-----RELEDLVACYGVRY MDP-ESFSAA-FFKWDQRPPALAPPQMQRSAGLEAQRIFHDFGVPNAAAMAASNNSSSCRKELNCLEELFRNYGVRY MDA-EHFPVG-FFRWDQRP----APVVAAAAAPTTTVFNKDHGRPLEVILPMNG----RKDLKSLEDLFKEYGVRY LFY 60 ZFL1 58 PRFLL 75 NEEDLY 66 YTAAKIAELGFTVNTLLDMKDEELDEMMNSLSQIFRWDLLVGERYGIKAAVRAERRRLDEED SLF PRRROLLSGD- 133 YTAAKIAELGFTVNTILLDMKDEELDEMMNSLSQIFRWDLLVGERYGIKAAVRAERRRLDEED YTAAKIAELGFTVSTILLDMKDEELDDMMNSLOQIFRWDLLVGERYGIKAAVRAERRRLDEEE YTAAKIAKLGFTVNTILLDMKDDELDDMMSSLSQIFRWELLVGERYGIKAAVRAERRREEEED YTAAKIAELGFTVNTILLDMRDEELDEMMNSLOQIFRWDLLVGERYGIKAAVRAERRRIDEEE -PRRROLLSGDN 134 -SRRRHLLSADT 140 PTLF VFL AFL1 RRRNPVSGDT 141 FLO RRRHLLLGDT 133 TTAAKIABLGFTASTLVCMKDEELEEMMNSLSHIFRWELLVGERYGIKAAVRAERRRLOEBE STVARISELGFTASTLLCMKDEELEEMMNSLSHIFRWELLVGERYGIKAAVRAERRRLOEBE ITLTKMVDMGFTVNTLVNMTEOELDDLVRTLVEIYRVELLVGEKYGIKSAIRAEKRLEEAER---KRMEQLFVDVD 149 VTLAKMTEMGFTANTLVNMTEEEIEELDMKTLVEIYHMDLLIGERYGIKSAIRAEKKRLODSLE---MORLEILSEAE 140 LFY ZFL1 PRFLL NEEDLY -----TNTLDALSOE GFSEEPVQQD--KEAAGSGGR---GTWEAVTAGER-KKOPGRK--KGHRKVVDL 189 SLF 

The second se PTLF VFL AFL1 FLO LFY LD-----AASOEALSDERDAAASGGGMAEGEAGRRMVTTTAG---KKGKKGVVGTRKGKKARRK--KELRPLNVL 194 GKRKIDEN---ALDTLSOE----CLSVEEP\_GDNAIILSONNTSANFPLNLNAGMDPVLILONSGHLGTTVSGLIGM 219 ZFL1 ----CLSVDEPOGDNAIILSQNNTSANFPLNLNAGMDPVLILONSGHLGTTVSGLIGM 219 ---CTSK-ELRANDPLIFPESTSADHAPMNIASCKDSTLILONSNQAQFCGSGLIGV 212 PRFLL NEEDLY RKRILHDDONTFAAAMAS DG----DDEHGG------AICEROREHPFIVTEPGEVARGKKNGLDYLFHLYEQCRDFLIQVQ DG----DDEHGG------AICEROREHPFIVTEPGEVARGKKNGLDYLFHLYEQCRDFLIQVQ DDNMNEDDNEGGDEDDD-------KGSGERGSEROREHPFIVTEPGEVARGKKNGLDYLFHLYEQCRDFLIQVQ SLF 242 PTLF 244 VFL 263 GGGHDNDHNEGVDDKDDDMDNMNGQGNGGGGGLLGERQREHPFIVTEPGEVARGKKNGLDYLF<mark>Y</mark>LYELCRDFLIQVQ EEDDDDDDDETEGAEDD------ENIVS<mark>ERQREHPFIVTEPGEVARGKKNGLDYLFHLYEQCRD</mark>FLIQVQ AFL1 276 FLO 259 DEDVNEGEDDDGMDNGN------GGSGLGTERQREHPFIVTEPGEVARGKKNGLDYLFHLYEQCREFLLQVQ LFY 275 DDENDGDEYGGGSESTES-----SAGGSGERQREHPF<mark>V</mark>VTEPGEVARAKKNGLDYLFHLYEQCRVFLLQVQ PDTNYGSEQTKACK-KQKRRR-----SKDSGEDGEERQREHPFIVTEPGE<mark>L</mark>ARGKKNGLDYLF<mark>D</mark>LYEQCGKFLLDVQ ZFL1 260 290 PRFLL PEHSSESDERKADTNKQKRRR----SKEPGEDGEDRPREHPFIVTEPGELARGKKNGLDYLFDLYEQCGKFLL NEEDLY 284 NIAKERGEKCPTKVTNQVFRYAKKAGASYINKPKMRHYVHCYALHCLDEDASNALRRAFKERGENVGAWR SLF 319 SIAKERGEKCPTKVTNQVFRYAKKAGASYINKPKMRHYVHCYALHCLDEDASNALRRAFKERGENVGAWRQACYKPL PTLF 321  ${\tt NIAKERGEKCPTKVTNQVFRYAKKAGASYINKPKMRHYVHCYALHCLDEEASNALRRAFKERGENVGAWRQACYKPL}$ VFL 340 NIAKERGEKCPTKVTNQVFRYAKKSGASYINKPKMRHYVHCYALHCLD<mark>V</mark>EASNVLRRAFKERGENVGAWRQACYKPL AFL1 353 TIAKERGEKCPTKVTNQVFRYAKKAGANYINKPKMRHYVHCYALHCLDEAASNALRRAFKERGENVGAWRQACYKPL FLO 336 IAK<mark>D</mark>RGEKCPTKVTNQVFRYAKKSGASYINKPKMRHYVHCYALHCLDEEASNALRRAFKERGENVGSWRQACYKPL LFY 352 SIAK<mark>I</mark>GG<mark>H</mark>KSPTKVTNOVFRYANKOGASYINKPKMRHYVHCYALHCLDEEASNALRRAYKSRGENVGAWROACYAPL HIAKERGEKCPTKVTNOVFRHAKHSGAGYINKPKMRHYVHCYALHCLDIEOSNRLRRAYKERGENVGAWROACYYPL RIAKEKGEKCPTKVTNOVFRHAKHN<mark>GAV</mark>YINKPKMRHYVHCYALHCLDS<mark>EO</mark>SNHLRR<mark>I</mark>YKERGENVGAWROACYYPL ZFL1 337 367 PRFLL NEEDLY 361 <u>RQGWDIDSIFNAHPRLAIWYVPTKLRQLC<mark>YAER</mark>N-----GATAS-----SSVSGTG----VHLPF-</u> 375 SLF VAIAS<mark>RQGWDIDSIFNAHPRLAIWYVPTKLRQLCY</mark>AERN-----SATSS-----SSVSGTG----GHLPF-VALAARQGWDIDAIFNAHPRLAIWYVPTRLRQLCH<mark>S</mark>ER<mark>SNAAAAAAAAAS-----SCISGGA----DHLPF-</mark> PTLF 377 VFL 402 VIAAAQGWDIDAIFNSHPRLSIWYVPTKLRQLCHAERHN----ATASS-----SASGGGG----EHLPY-AFL1 410 /<u>AIAARQGWDIDTIFNAHPRISIWYVPTKIRQICHABR</u>SS----AAVAAT----SSITGGGPA--DHLPF-FLO 396 LFY 424 391 ZFL1

Fig. 1 Genome structure, cDNA and deduced amino acid sequences of SLF. (A) Genome structure of SLF (top) and SLF cDNA (bottom). The SLF locus has three exons (boxes) and two introns (lines). Gray box (from nucleotides 550 to 1044) in the cDNA region indicates location of the probe used in Southern and northern blots. Arrows indicate the HindIII restriction sites in the genomic SLF clone. There is no EcoRI restriction site in the genomic SLF clone. (B) Alignment of the deduced amino acid sequences of SLF, LFY, FLO, and eight other LFY/FLO homologs (accession numbers in parentheses): PTLF from Populus trichocarpa (U93196), VFL from Vitis vinifera (AF450278), FLO from Antirrhinum majus (M55525), AFL1 from Malus × domestica (AB056158), LFY from Arabidopsis thaliana (M91208), ZFL1 from Zea mays (AY179882), PRFLL from Pinus radiata (U92008), and NEEDLY from Pinus radiata (U76757). Black shadows indicate identical amino acids; dashes indicate gaps introduced to optimize the alignment. The acidic central domain is located between the two arrows.

411

404



Fig. 2 Southern blot of genomic DNA from *Salix discolor* probed with *SLF* partial sequence. Ten  $\mu$ g genomic DNA was loaded per lane. Stringency washes with 0.5 × SSC/0.1% SDS were performed. NBT/BCIP color substrate solution was used for 12 h to visualize the hybridization on the membrane. MW was determined through comparison with the 1 kb DNA ladder.

according to the manufacturer. The membranes were incubated with gentle agitation in the probe/hybridization solution at 50°C for 12 h. Stringency washes with  $0.1 \times SSC/0.1\%$  SDS were performed after hybridization. Anti-digoxigenin-AP was used as the antibody. NBT/BCIP color substrate solution was used to visualize the hybridization on the membrane. The membrane was scanned using the GS-710 Calibrated Imaging Densitometer (Bio-Rad, Hercules, CA).

#### RESULTS

#### SLF cDNA and amino acid sequence analyses

Using degenerate PCR primers designed from the conserved regions of *LFY*, a 551-bp partial sequence of *SLF* cDNA was obtained. The coding region of *SLF* was 1125 bp (GenBank Accession No. AY230817). The *SLF* genome structure was made up of three exons and two introns (**Fig. 1A**). The introns were 582 bp and 495 bp long. The deduced amino acid sequence of *SLF* was compared with the LFY and FLO proteins, LFY/FLO homolog proteins from several woody species, and other recently reported LFY/ FLO homologs from flowering plants (**Fig. 1B**). The alignment confirmed two highly conserved regions, one containing 77 residues from amino acid 47 to 123, and the other containing 157 residues from amino acid 201 to 357. There were three regions with lower similarity (**Fig. 1B**). The proline-rich region in SLF had six proline residues between amino acid 20 and 36. As compared to other LFY/ FLO homologs, SLF had a smaller number of acidic residues. FLO had 14 acidic residues but SLF had only four in the acidic central domain (between the two arrows in Fig. 1B). The alignment of sequences revealed that SLF had 97% identity with PTLF (*P. trichocarpa*), 78% with VFL (*Vitis vinifera*), 76% with FLO (*Antirrhinum majus*), 72% with AFL1 (*Malus × domestica*), 67% with LFY (*A. thaliana*), 56% with ZFL1 (*Zea mays*), 52% with PRFLL (*Pinus radiata*), and 49% with NEEDLY (*P. radiata*).

Southern hybridization using a PCR product amplified from the second highly conserved region of *SLF* revealed a single hybridizing *Eco*RI band of 3.4 kb. A single 1.9-kb *Hind*III band was also detected (**Fig. 2**). In the genomic *SLF* clone, there is no *Eco*RI restriction site but there are two *Hind*III restriction sites. However, the probe used is between the two *Hind*III restriction sites (**Fig. 1A**). This suggests that there is only one copy of *SLF* in *S. discolor*.

#### In situ expression patterns of SLF in S. discolor

The expression patterns of *SLF* in *S. discolor* were analyzed using male reproductive buds bearing inflorescence meristems and developing male flowers. Strong *SLF* expression was localized in the inflorescence meristems as well as in the developing flowers along the flanks of the inflorescence meristems (**Fig. 3A**). *SLF* was strongly detected in all the layers of the inflorescence meristems and bract primordia (**Fig. 3B**), young bracts (**Fig. 3D**), and floral meristems (**Fig. 3D**, **3E**). Using lateral vegetative buds, *SLF* RNA was also detected, but in low levels in the vegetative meristems and leaf primordia (data not shown). Using the sense probe, no detectable hybridization was observed in any part of the male reproductive and vegetative buds (**Fig. 3C**).

#### Functional analysis of SLF in A. thaliana

To test the functions of *SLF*, a construct (pCGSLF) to overexpress the gene in *A. thaliana* was made. In addition to the *SLF* gene, the construct contains *mgfp5-er* driven by the CaMV 35S promoter. Several 35S::*SLF A. thaliana* transgenic lines were successfully screened based on GFP expression (**Table 2**). To analyze the phenotypes of homozygous transgenic lines, two negative controls were used including wild type and 35S::*GFP A. thaliana* (**Table 2**). The T<sub>1</sub> seeds were collected in separate pools, germinated,



# Fig. 3 *In situ* hybridization analysis of *SLF* expression in male reproductive buds of *Salix discolor*. All images are longitudinal

Sanx anscoror. All images are iongluturnal sections. (A) Early male inflorescence, antisense probe. (B) Magnified inflorescence meristem, antisense probe. (C) Early male inflorescence, sense probe. (D) Young bracts and floral meristems, antisense probe. (E) Magnified floral meristem, antisense probe. BP, bract primordia; FM, floral meristem; IM, inflorescence meristem; YB, young bract. Bars = 100 µm.

Table 2 Phenotypes of various T<sub>3</sub> homozygous 35S::SLF transgenic lines of Arabidopsis thaliana and controls under long-day condition.

Genotypes	n	Days to flowering	Rosette leaves at flowering	Inflorescence branches after six weeks	Solitary flowers	Terminal flowers
35S::SLF1	40	$13.2 \pm 0.7*$	$5.9 \pm 0.5*$	1	+	+
35S::SLF2	39	$13.3 \pm 1.2*$	$6.2 \pm 0.7*$	1	+	+
35S::SLF3	40	$11.7 \pm 0.9*$	$4.0 \pm 0.0*$	1	+	+
35S::SLF4	37	$13.4 \pm 1.0*$	$5.7 \pm 0.9*$	1	+	+
35S::SLF5	38	$10.3\pm0.8*$	$2.0 \pm 0.0*$	$7.4 \pm 1.3$	-	-
35S::SLF6	39	$13.2 \pm 0.8*$	$5.8 \pm 1.1*$	1	+	+
35S::GFP	36	$29.6 \pm 1.1$	$14.4 \pm 0.7$	$3.3\pm0.9$	-	-
Wild type	40	$28.7 \pm 1.3$	$13.9 \pm 1.2$	$3.2\pm0.7$	-	-

\* indicates significance at 5% level (P value < 0.05) compared to the controls



Fig. 4 GFP screens of different developmental stages and organs in  $T_3$  35S::*SLF Arabidopsis thaliana* transgenic line. (A) Seeds. (B) Leaf. (C) Roots. (D) Flower.

and screened using GFP expression. A total of 45 positive transgenic T<sub>1</sub> lines were obtained, and all these were considered as independent transformation events. A total of ten  $T_2$  lines were chosen based on the lines with brightest GFP expression and phenotypes such as early flowering, formation of terminal flowers, and conversion of inflorescence branches to solitary flowers. A total of six homozygous T<sub>3</sub> lines were isolated based on the number of seeds expressing GFP and their segregation ratios. For each of the six homozygous T<sub>3</sub> lines, 40 seeds were planted and grown under long-day condition. All organs including leaves, roots, and flowers were further screened for GFP expression (Fig. 4). There was no difference observed in the phenotypes of the wild-type and transgenic control line (35S::GFP, trans-formed with GFP only). This demonstrated that GFP was not responsible for the flowering response that was observed from the SLF transgenic lines. Compared to wild-type and the transgenic control line (Fig. 5A), the transgenic SLF lines displayed early flowering, formation of terminal flowers, conversion of inflorescence branches to solitary flowers, and formation of more inflorescence branches (Fig. **5B-E**), indicating reiteration of *SLF* function.

Under long-day condition, floral buds in lines

35S::*SLF*1, 2, 3, 4, 5 and 6 were produced much earlier and with smaller number of rosette leaves compared to the controls (**Fig. 5B, Table 2**). Conversions of inflorescence branches to solitary flowers were also observed in these lines (**Fig. 5E, Table 2**), except in 35S::*SLF*5 where more inflorescences branches were produced compared to the controls (**Fig. 5D, Table 2**). Early flowering in lines 35S::*SLF*3 and 35S::*SLF*5 occurred with only four and two rosette leaves, respectively (**Fig. 5C, Table 2**).

To determine whether *SLF* can rescue the phenotype of a lfy mutant of A. thaliana, pollen grains were collected from each of three homozygous  $T_3$  lines (35S::SLF) and used to pollinate several lfy-6 individuals. The early-formed flowers of *lfy-6* have pistils but no petals and stamens (Fig. 5F). Compared to the wild-type, the shape of rosette leaves in lfy-6 plants was relatively round with smooth margins (Fig. 5G). The phenotypes produced by the  $F_2$  individuals demonstrated that SLF was able to rescue the floral phenotype of *lfy-6*. In F<sub>2</sub> plants, there were 11 out of 200 individuals showing the phenotype of lfy-6. All other plants had normal flowers (Fig. 5H, 5I). As we have predicted, many of the individuals form terminal flowers or more inflorescence branches. These results confirm that almost 1/16 individuals still display the phenotype of lfy-6 because SLF gene is not expressed in their progenies. This is consistent with Mendel's rule of independent assortment.

To determine the *SLF* expression level in  $T_3$  transgenic lines, northern analysis was done using a digoxigenin-11-UTP labeled probe. *SLF* was expressed at similar levels in all six  $T_3$  transgenic lines (**Fig. 6**). In addition, based on the segregation of the  $T_2$  generation (**Table 3**), the ratio of GFP expression was 3:1 in these transgenic lines. This confirms that only a single T-DNA insertion locus was present in each transgenic line.

#### DISCUSSION

#### Sequence analysis of SLF

The LFY/FLO protein has a proline-rich terminus region and an acidic central domain, which are involved as transcriptional activators (Coen *et al.* 1990; Weigel *et al.* 1992). Sequence analysis of LFY/FLO homologs from various plants also shows the presence of these two regions. PRFLL and NEEDLY from *P. radiata* have fewer proline residues in their proline-rich terminus region than any of the LFY/ FLO proteins from angiosperms. Furthermore, the acidic region is absent in PRFLL and NEEDLY (Mellerowicz *et al.* 

Table 3 Segregation ratios of GFP expression in the T<sub>2</sub> generation of six 35S::SLF transgenic lines

Table 5 Segregation latios of Off expression in the 12 generation of six 555.527 transgenie lines.									
Transgenic line	GFP positive	GFP negative	Segregation ratio	95% CI	p-value*				
35S::SLF1	476	159	2.99:1	0.714, 0.782	1.000				
35S::SLF2	602	199	3.03:1	0.720, 0.781	0.935				
35S::SLF3	634	211	3.00:1	0.720, 0.779	1.000				
35S::SLF4	491	162	3.03:1	0.717, 0.785	0.928				
35S::SLF5	512	170	3.01:1	0.716, 0.783	0.965				
35S::SLF6	487	161	3.02:1	0.716, 0.784	0.928				

\*Test of p = 0.75 vs p not = 0.75

The test is for the proportion of positive GFP expression that is equal to 0.75 vs not equal, and the results in all transgenic lines are not significant (p-value>0.888), which indicates that the data are consistent with the hypothesized 3:1 segregation ratio.



Fig. 5 Heterologous expression of SLF in Arabidopsis thaliana transgenic lines. (A) wild-type Arabidopsis. (B) 35S::SLF1 showing the formation of terminal flower and 5 rosette leaves before flowering. (C) Transgenic line 35S::SLF3 showing the formation of terminal flower and 4 rosette leaves before flowering. (D) Transgenic line 35S::SLF5 showing more number of inflorescence branches and early flowering. (E) The conversion of inflorescence branch to solitary flower. (F) The early formed flowers of lfy-6 with no petals and stamens. (G) The rosette leaves of lfy-6 showing round shape with smooth margins. (H) Comparison of floral phenotypes between lfy-6 (left) and rescued plant (right) at F2. (I) Early formed flowers of F2 bearing petals and stamens. IB, inflorescence branch; TF, terminal flower; SF, solitary flower.).



**Fig. 6 Northern analysis in six T<sub>3</sub>** *Arabidopsis thaliana* transgenic lines. Lane: 1, 35S::*SLF*1; 2, 35S::*SLF*2; 3, 35S::*SLF*3; 4, 35S::*SLF*4; 5, 35S::*SLF*5; 6, 35S::*SLF*6; 7, 35S::*GFP* (transgenic control line). Twenty µg total RNA was loaded per lane. Non-radioactive probe was used. NBT/ BCIP color substrate solution was used for detection. The lower panel displays the rRNA of each sample to indicate equal loading amounts.

1998; Mouradov *et al.* 1998). The proline-rich region of LFY, FLO, and other angiosperm LFY/FLO homolog proteins has nine residues (Coen *et al.* 1990; Weigel *et al.* 1992; Kelly *et al.* 1995; Busch and Gleissberg 2003). However, SLF only has six proline residues. Seven proline residues were reported in PTLF (Rottmann *et al.* 2000). SLF and PTLF also both have a smaller number of acidic residues as compared to LFY/FLO. The alignment of the predicted amino acid sequences of SLF and PTLF shows that they share 97% similarity. SLF has a total of 375 residues, while PTLF had 377 residues. The difference is due to the presence of two asparagine residues at amino acid 134 and 135 in PTLF. Based on gene structure, the number and location of introns in *SLF* and *PTLF* are similar. However, the sequences and sizes of the introns are different.

#### SLF expression patterns in S. discolor

The localization of SLF expression in various parts of the male reproductive buds of willow shows that this gene is strongly involved in the formation of the inflorescence meristems, bracts, and floral meristems. PTLF is also strongly expressed in developing flowers of poplar (Rottmann et al. 2000). The expression of SLF was detected at low levels in the vegetative meristems and leaf primordia from lateral vegetative buds. PTLF was also detected at low levels in the vegetative meristem from terminal buds (Rottmann et al. 2000). It appears that gene expression between the lateral and terminal buds are similar. Both LFY and FLO are strongly expressed in the floral meristems prior to the initiation of floral organ primordia (Coen et al. 1990; Weigel et al. 1992). In leaf primordia, expression of LFY/FLO homologs has also been detected at low levels in A. thaliana, Nicotiana tabacum, and Petunia × hybrida (Weigel et al. 1992; Kelly et al. 1995; Souer et al. 1998), but not in A. majus (Coen et al. 1990) and Cedrela fissilis (Dornelas et al. 2006). In P. radiata, NEEDLY and PRFLL are expressed in the vegetative meristems, but not in the female cones (Mellerowicz et al. 1998; Mouradov et al. 1998). These studies show and confirm that there is variation in the expression patterns of LFY/FLO homologs in the vegetative and reproductive meristems among species.

#### SLF expression in A. thaliana

The functions of *LFY/FLO* homologs from several woody plants have been assessed through heterologous expression using *A. thaliana* or *N. tabacum* (Mellerowicz *et al.* 1998; Mouradov *et al.* 1998; Southerton *et al.* 1998; Rottmann *et al.* 2000; Wada *et al.* 2002; Carlsbecker *et al.* 2004; Dornelas and Rodriguez 2005; Dornelas *et al.* 2006). In willow, an *in vitro* regeneration protocol that is coupled with genetic transformation is currently not available. Therefore, *A. thaliana* was used to generate transgenic lines to characterize the functions of *SLF*. Overexpression of *SLF* in *A. thaliana* produced a total of six homozygous T<sub>3</sub> transgenic lines. The phenotypes of these transgenic lines are dominant and heritable. These demonstrate that *SLF* is functional and affects various aspects of flower development in transgenic *A. thaliana*.

Arabidopsis thaliana transformed with PTLF flowered an average of five days earlier with one fewer rosette leaf than the wild-type (Rottmann et al. 2000). Other than these, no marked difference was observed in the phenotypes of the transformed A. thaliana and the wild type (Rottmann et al. 2000). Using SLF, transformed A. thaliana lines showed dramatic changes in their flowering response, including the formation of terminal flowers and conversion of inflorescence branches to solitary flowers. Also, in all six 35S::SLF A. thaliana transgenic lines, flowering occurred more than two weeks earlier than the control plants. The discrepancy in the results between SLF and PTLF is surprising considering their very high sequence similarity. However, in a few *LFY/FLO* homologs with a high degree of amino acid sequence similarity different phenotypic effects have also been reported. AFL1 and AFL2 are 90% similar, but fewer solitary flowers occurred in AFL1 transgenic A. thaliana than in AFL2 (Wada et al. 2002). It is also possible that the presence of two 35S promoters in the pCGSLF binary vector increased SLF expression. Furthermore, it was reported that the 35S promoter driving the selectable marker gene could alter the expression of transgenes (Yoo et al. 2005).

Conversion of inflorescence branches to solitary flowers and formation of terminal flowers were produced in 35S::SLF1, 2, 3, 4, and 6 transgenic lines. These floral phenotypes were not observed in A. thaliana transformed with PTLF according to the report of Rottmann et al. (2000). However, transgenic A. thaliana expressing NEEDLY from P. radiata and ELF1 from Eucalyptus globulus produced solitary flowers from axils of rosette leaves and terminal flowers from primary shoots (Mouradov et al. 1998; Southerton et al. 1998). Therefore, SLF appears to produce phenotypes typical of a LFY/FLO gene. More branches of primary inflorescence and inflorescence branches were produced in the 35S::SLF5 transgenic line. Since SLF is involved in inflorescence and flower development, overexpression of SLF can produce more inflorescences and flowers. Although these transgenic lines show different phenotypes, northern analysis demonstrated that SLF is expressed at similar levels in all these six transgenic lines. Based on the segregation analysis of the T<sub>2</sub> generation, the 3:1 ratio of seeds expressing GFP implies that the T-DNA was inserted in a single locus in all these lines; and such an approach is typically used in A. thaliana (Berbel et al. 2001; Honda et al. 2002). Although this still needs to be pursued, it is possible that the location of the T-DNA integration in the A. thaliana genome might be responsible for such a dramatic phenotype in the 35S::SLF5 transgenic line.

Introduction of SLF from T<sub>3</sub> homozygous A. thaliana transgenic lines into lfy-6 mutants resulted in the rescue of the lfy-6 floral phenotypes. The mutant lfy-6 lacks both petals and stamens in the early-formed flowers. The rescue experiment showed a high degree of functional conservation between LFY/FLO and SLF. These results, and those of Dornelas and Rodriguez (2005) on rubber tree, are the only reports that described LFY/FLO homologs that are able to rescue the floral phenotype of a *lfymutant*. On the other hand, various complementation tests using LFY homologs from herbaceous flowering plants (Weigel *et al.* 1992; Wada *et al.* 2002) and conifers (Mouradov *et al.* 1998; Carlsbecker *et al.* 2004; Dornelas *et al.* 2006) showed similar positive results when crossed with the *lfy-6* null allele.

This study has revealed that *SLF* from a male individual of *S. discolor* affects various aspects of flower development in transgenic *A. thaliana*. In fact, many of these phenotypes have not been reported from the study using a *LFY/FLO* homolog from a closely related dioecious species, *P. trichocarpa*. These results expand our understanding of the functions of *LFY/FLO* homologs and suggest that the *LFY/FLO* homolog from a male dioecious plant is equally functional to those from bisexual species. Unfortunately, female *S. discolor* was not available during our study.

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