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Flowering of strict photoperiodic *Nicotiana* varieties in non-inductive conditions by transgenic approaches

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Abstract The genus Nicotiana contains species and varieties that respond differently to photoperiod for flowering time control as day-neutral, short-day and long-day plants. In classical photoperiodism studies, these varieties have been widely used to analyse the physiological nature for floral induction by day length. Since key regulators for flowering time control by day length have been identified in Arabidopsis thaliana by molecular genetic studies, it was intriguing to analyse how closely related plants in the *Nicotiana* genus with opposite photoperiodic requirements respond to certain flowering time regulators. SUPPRES-SOR OF OVEREXPRESSION OF CONSTANS 1 (SOC1) and FRUITFULL (FUL) are two MADS box genes that are involved in the regulation of flowering time in Arabidopsis. SOC1 is a central flowering time pathway integrator, whereas the exact role of FUL for floral induction has not been established yet. The putative Nicotiana orthologs of SOC1 and FUL, NtSOC1 and NtFUL, were studied in dayneutral tobacco Nicotiana tabacum cv Hicks, in short-day tobacco *N. tabacum* cv Hicks Maryland Mammoth (MM) and long-day *N. sylvestris* plants. Both genes were similarly expressed under short- and long-day conditions in day-neutral and short-day tobaccos, but showed a different expression pattern in *N. sylvestris*. Overexpression of *NtSOC1* and *NtFUL* caused flowering either in strict short-day (*NtSOC1*) or long-day (*NtFUL*) *Nicotiana* varieties under non-inductive photoperiods, indicating that these genes might be limiting for floral induction under non-inductive conditions in different *Nicotiana* varieties.

Keywords *Nicotiana* · Floral induction · MADS box genes · *TobMADS* · *NtSOC1* · *NtFUL*

Introduction

Plants have adapted flowering time for their natural habitats and, therefore, the onset of flowering varies widely among different species and ecotypes. Temperature and day length are the principal environmental cues for plants to track the seasons of the year, which allow flowering to be synchronised for maximum reproductive success. Since the discovery of photoperiodism in soybean and Maryland Mammoth tobacco plants by Garner and Allard (1920), numerous experimental approaches in different plant species have been undertaken in order to study the day-length-dependent flowering in a large variety of photoperiodic plants (Thomas and Vince-Prue 1997). These classical studies, however, were limited in their ability to identify the underlying molecular nature for flowering time control in long- and short-day plants.

In recent years, rapid progress has been made in understanding the molecular mechanisms of floral induction in *Arabidopsis thaliana*, a facultative long-day plant

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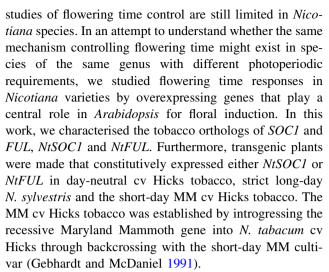
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and, more recently, in rice (Oryza sativa), a short-day plant (reviewed in Searle and Coupland 2004). In Arabidopsis four main pathways that control flowering time have been defined genetically. The photoperiod and the vernalisation pathways are involved in the perception of environmental signals, whereas the autonomous pathway acts independently of environmental cues (Koornneef et al. 1998; Mouradov et al. 2002; Simpson and Dean 2002). Gibberellins (GAs) are limiting for flowering in Arabidopsis (Wilson et al. 1992) and by genetic analysis of double mutant combinations of GA and late-flowering mutants the promotion of flowering by GAs has been shown to be mediated by a fourth independent pathway (Reeves and Coupland 2001). These flowering time pathways converge on pathway integrators, such as LEAFY (LFY) (Blazquez and Weigel 2000), FT (Kardailsky et al. 1999; Kobayashi et al. 1999) and the MADS box protein SOC1 (Borner et al. 2000; Lee et al. 2000; Samach et al. 2000). CON-STANS (CO) encodes a zinc finger protein that is a central regulator for flowering time control in Arabidopsis by long days. CO is expressed in the vasculature where it directly activates FT transcription (An et al. 2004). The FT protein has been shown to be transported to apical meristems (Corbesier et al. 2007; Tamaki et al. 2007), where it interacts with FD (Abe et al. 2005; Wigge et al. 2005) to activate SOC1 expression (Searle et al. 2006), which finally leads to the activation of floral meristem identity genes and the formation of flowers.

In *Sinapis alba* (mustard) plants, the *SOC1* ortholog, *SaMADSA*, is co-expressed with the *FUL* ortholog, *SaMADSB*, in apical meristems and procambial strands during floral transition (Menzel et al. 1996) and in a genome-wide expression analysis, *SOC1* and *FUL* have been found to be similarly up-regulated in response to photoperiodic floral induction in apical meristems in *Arabidopsis* (Schmid et al. 2003). *FUL* was originally identified as a regulator for fruit dehiscence (Ferrándiz et al. 2000b; Gu et al. 1998), but mutant analysis indicated that it also plays a role in flowering time control (Ferrándiz et al. 2000a; S. Melzer, unpublished data). Molecular studies have shown that *FUL* expression is controlled in part by *FT* (Teper-Bamnolker and Samach 2005), but the role of *FUL* for floral induction in *Arabidopsis* is not clear yet.

Nicotiana species and cultivars have different photoperiodic requirements for flowering time control and have been used for many decades in floral induction studies (Lang 1989; McDaniel 1996). These studies have indicated that identical flowering stimulatory and inhibitory substances are formed in different photoperiodic response types under inductive or non-inductive conditions, whereas classical grafting experiments have revealed that these substances can be transferred from one response type to another (Lang 1989). However, molecular and genetic



Overexpression of *NtSOC1* and *NtFUL* caused very early flowering in day-neutral tobacco, and under inductive photoperiods also in the short-day variety MM cv Hicks and in long-day *N. sylvestris* plants. However, under non-inductive conditions the transgenic lines behaved differently. The *35S::NtSOC1* transgene bypassed the photoperiodic requirements in the MM cv Hicks tobacco, but not in long-day *N. sylvestris* plants under non-inductive conditions. Conversely, the *35S::NtFUL* transgene triggered flowering of *N. sylvestris* under non-inductive short days, but had no effect in MM cv Hicks tobacco in long days, suggesting that these genes are limiting in certain *Nicotiana* varieties under non-inductive conditions.

Materials and methods

Plant material and growth conditions

Seeds from *Nicotiana tabacum cv*. Hicks, *N. tabacum* MM cv Hicks and *N. sylvestris* plants were obtained from Dr Susan Singer (Carleton College, Northfield, MN, USA) and were sown on soil. Seedlings were singled out in 16-cm pots after 21 days. Plants were grown either in phytotrons under short-day (8 h light) and long-day conditions (16 h light), under fluorescent tubes emitting a photon flux density of 160 μmol m⁻² s⁻¹ at 20°C, or in greenhouses in long days at 22°C during the day and 18°C during the night.

Isolation and characterisation of SaMADSA/SOC1 and SaMADSB/FUL tobacco orthologs

To identify tobacco orthologs to mustard and *Arabidopsis SaMADSA/SOC1* and *SaMADSB/FUL1* genes, cDNA libraries were constructed from RNA of florally induced apices derived from *N. tabacum* cv Hicks plants with



standard protocols. A λ gt10 library with 5,000,000 plaque forming units (pfu) were screened at low stringency (hybridisation at 45°C and washing in 1× SSC, 0.2% SDS at 37°C) with *Hind*III fragments of *SaMADSA* and *SaMADSB* that did not contain the MADS domain (Menzel et al. 1996). Positive pfu were purified, phage DNA was isolated and *Eco*RI-digested with fragments subcloned into a pBluescript SK⁺ vector (Stratagene, Madison, WI, USA). Ten cross-hybridising clones for each gene were selected and sequenced.

The SOC1 and FUL homologous sequences of Arabidopsis thaliana (At), Sinapis alba (Sa), Petunia x hybrida (Ph), Lycopersicon esculentum (Le), N. tabaccum (Nt) and N. sylvestris (Ns) were extracted from the NCBI database (http://www.ncbi.nlm.nih.gov). Protein sequences were aligned with ClustalW (Thompson et al. 1994), and the alignment was edited with BioEdit (URL: http://www.mbio.ncsu.edu/BioEdit/bioedit.html), resulting in an alignment of the conserved residues of the MADS, I and K domain. Neighbour-Joining (Saitou and Nei 1987) trees for the proteins were constructed with TREECON (Van de Peer and De Wachter 1997) based on Poisson-corrected distances. To assess support for the inferred relationships, 500 bootstrap samples (Felsenstein 1985) were generated.

Transgene construction and plant transformation

By using primers with EcoRI adaptors, the coding regions of NtSOC1 and NtFUL were amplified by standard PCR reactions and fused to the CaMV 35S promoter in pRT101 plasmids (Töpfer et al. 1987). The expression cassette of the pRT101 vector was introduced as a HindIII fragment into pRD400 (Datla et al. 1992) and the binary vector was transformed into the Agrobacterium tumefaciens strain C58C1. For Agrobacterium-mediated leaf disc transformation (Horsch et al. 1985), day-neutral N. tabacum cv. Hicks, N. tabacum MM cv Hicks and N. sylvestris plants were grown in vitro. Regenerating plants were selected on agar plates containing Murashige and Skoog (MS) media (Duchefa, Haarlem, The Netherlands) supplemented with 500 mg/l Timenten and 200 mg/l kanamycin. Seeds from T1 and T2 plants were tested for homozygosity on kanamycin plates and T3 or T4 homozygous lines were used for experiments.

RNA blot and semi-quantitative RT-PCR analysis

Total RNA was isolated according to Melzer et al. (1990). For transgene expression analyses, 30 μg of total RNA from seedlings was separated on formaldehyde agarose gels, transferred to nylon membranes and hybridised with *NtSOC1* or *NtFUL* probes, without the MADS box region, at 65°C according to standard protocols.

For a semi-quantitative RT-PCR analysis, reverse transcription of 3 µg of total RNA was performed with SuperScript II RT (Invitrogen, Carlsbad, CA, USA) as described (Melzer et al. 1999). Gene-specific PCR products were amplified with the following primer pairs: for 5'-NtSOC1: GCATGCGGCAGCAAGTTTGAT; for 3'-NtSOC1: GGAAAATATAATACACATCC; for 5'-NtFUL: GGGAAGCATATCAGAGTAC: for 3'-NtFUL: CAAGG CTGATAAAGATCAG; for 5'-NtNAP1-2: CCTCCTAC AACCACATCCAT; for 3'-NtNAP1-2: TAGGAAATT TACATTCCTCA: for 5'-NFL1: CAAGAAGATGAGTG GAATATTAACGA and for 3'-NFL1: CAGTTACAGAA TTTGCAGAACTGAAT. Semi-quantitative PCR was performed at 58°C annealing temperature for 20 or 25 cycles. The PCR fragments were gel separated, transferred to nylon membranes and hybridised with their corresponding probes. As a control, tobacco eIF4A10 transcripts (Mandel et al. 1995) were amplified with 5'-NteIF4A10 CAATTGCTACCACCAAAGAT and 3'-NteIF4A10 AAA GGAGATCGGCCACATTGG primers.

Microscopy

For microscopic analysis, samples were fixed overnight with 4% formaldehyde in 50 mM phosphate buffer, pH 7, dehydrated with EtOH and embedded in Technovit 7100 resin (Heraeus Kulzer, Wetzlar, Germany). Sections of 6 μ m were cut with a rotary microtome, stained with phloroglucinol/HCL to visualize lignified cells and mounted in DePex medium (British Drug House, UK).

Results

Identification of SOC1 and FUL orthologs in tobacco

Since SOC1 and FUL in Arabidopsis as well as the mustard orthologs, SaMADSA and SaMADSB, have similar expression patterns, both in apical meristems and in procambial strands of the developing inflorescence after floral induction (Borner et al. 2000; Menzel et al. 1996), we analysed the orthologs of both genes in strict photoperiodic tobacco plants. To identify putative SOC1/SaMADSA and FUL/ SaMADSB orthologs in tobacco, we screened a N. tabacum cDNA library, which was made from mRNA of apical buds from induced plants, under low-stringency conditions with mustard SaMADSA and SaMADSB probes. Probes without the conserved MADS box regions hybridised to several phage plaques from which 10 were selected for further characterisation. All clones that cross-hybridised with SaMADSA were identical to the previously identified TobMADS1 gene (Mandel et al. 1994; X76188), which



showed a high sequence homology to *SOC1* and *SaMADSA*, indicating that this is the *SOC1* ortholog of tobacco. Therefore, for more clarity, this gene was re-designated to *NtSOC1*. From cDNAs showing cross-hybridisation with *SaMADSB*, three were identical to *NtNAP1-1* and its homolog *NsMADS1* in *N. sylvestris* (Wu et al. 2000). Seven other cDNAs were identical, had a higher amino acid identity to *SaMADSB* and *FUL* and were subsequently designated *NtFUL* (GenBank: DQ534202).

Phylogenetic relationships of NtSOC1 and NtFUL

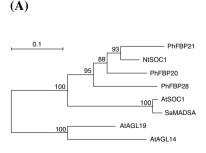
NtSOC1 (CAA53782) belongs to the TM3 (TDR3-X60756) subfamily of MADS-box proteins, of which SOC1/AGL20 (AAG16297) and SaMADSA (AAB41526) are the *Arabidopsis* and mustard representatives, respectively (Borner et al. 2000; Samach et al. 2000). Based on the phylogenetic tree of the closest SOC1 homologs, we can conclude that NtSOC1 is most closely related to the petunia (*Petunia hybrida*) FBP21 protein (AAK21252). The petunia FBP20 or UNSHAVEN (UNS) protein (AAK21252) (Ferrario et al. 2004) also belongs to this subfamily, albeit it is not directly related to the tobacco NtSOC1 protein (Fig. 1A).

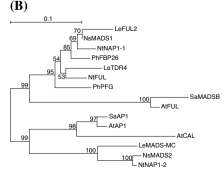
NtFUL (ABF82231) belongs to the SQUAMOSA (SQUA) (X63701) subfamily and is closely related to the *Arabidopsis* FUL (NM_125484) and mustard SaMADSB (U25695) proteins (Fig. 1B). The *FUL* clade has undergone a duplication in the *Solanaceae* lineage (Litt and Irish 2003) and the tobacco protein described here belongs to the clade of the tomato (*Lycopersicon esculentum*) TDR4 (X60757) and the PETUNIA FLOWERING GENE (PFG) (AF176782) (Immink et al. 1999) proteins, whereas NtNAP1-1 (AAD01421) and NsMADS1 (AF068725) (Wu et al. 2000) belong to the clade of LeFUL2 (AY306156) and PhFBP26 (AAF19164) (Fig. 1B).

Expression analysis of NtSOC1 and NtFUL

Expression of NtSOC1 and NtFUL in roots, stems and leaves of N. tabacum plants was analysed by semi-

Fig. 1 Phylogenetic relationships of NtSOC1 and *NtFUL*. The phylogenetic trees of NtSOC1 (**A**) and NtFUL (**B**) are based on protein sequences. The scale represents the evolutionary distance, with 0.1 substitutions per site





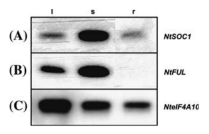


Fig. 2 Expression analysis of *NtSOC1* and *NtFUL* in different tissues. RT-PCR was performed on samples from leaves (1), upper internodes of stems without the apex (s) and roots (r) from 6-week-old vegetative plants with either *NtSOC1* (**A**) or *NtFUL* (**B**) primer pairs. As a control fragments of tobacco *eIF4A10* cDNAs were amplified (**C**)

quantitative RT-PCR. As shown in Fig. 2, both genes were detectable in leaves with 20 cycles of RT-PCR and at higher levels in stems of 9-week-old vegetative plants grown under long days. NtSOC1 was also visible at low levels in roots, in which we did not see any transcript accumulation of NtFUL. SOC1 and FUL are both expressed during vegetative stages in Arabidopsis plants, but are dramatically up-regulated in apical meristems after floral induction (Borner et al. 2000; Mandel et al. 1995; Schmid et al. 2003). Therefore, we analysed by semi-quantitative RT-PCR whether NtSOC1 and NtFUL might also be developmentally regulated in apices of different Nicotiana varieties, grown for 3, 6, 9, and 12 weeks either under short- or long-day conditions. Until 9 weeks after sowing, all *Nicotiana* plants examined remained vegetative, but the 12-week-old plants grown under inductive conditions had started flower development. Expression profiles of NtSOC1 and NtFUL as well as that of the APETALA1 (AP1) ortholog NtMADS5 (Calonje et al. 2004) and that of one of the Nicotiana FLORICAULA (FLO) LEAFY (LFY) orthologs, NFL1 (Kelly et al. 1995), were compared from apices of day-neutral tobacco, short-day cv Hicks MM tobacco and long-day N. sylvestris plants (Fig. 3). The NtSOC1 and NtFUL expression levels were very low in apical buds of day-neutral tobacco plants and short-day cv Hicks MM plants grown for 3 weeks in short- and long-day regimes. The expression levels increased similarly in apical buds of both tobacco varieties under short photoperiods to high



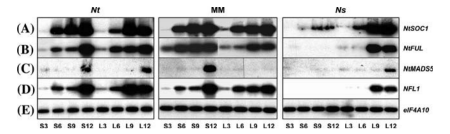


Fig. 3 Development-dependent expression of *NtSOC1* and *NtFUL* in apices of different *Nicotiana* varieties. A semi-quantitative RT-PCR was performed with RNA from apices of plants grown for 3, 6, 9 or 12 weeks under short (S3–S12) or long days (L3–L12) with *NtSOC1*

levels in 12-week-old plants. In long days, transcript levels of both genes increased also continuously with age, but compared to *NtSOC1*, *NtFUL* mRNA accumulated at lower levels in apices of vegetative MM cv. Hicks plants. However, both genes were expressed under these non-inductive conditions. We observed a different expression pattern in long-day *N. sylvestris* plants: expression of both genes was not detectable in non-inductive short days 3 weeks after sowing. Whereas *NtSOC1* mRNA accumulated to very low levels, *NtFUL* was undetectable throughout further development in short days. Under long-day conditions, both genes were expressed at low levels during the early stages of development and mRNA levels gradually increased to high levels after 9 and 12 weeks from sowing (Fig. 3A, B).

NtMADS5 could not be detected above background in apical buds of all vegetative stages, except at very basal levels in N. sylvestris in long days, but increased to high levels at stages in which all Nicotiana varieties had undergone the floral transition (Fig. 3C). Transcripts of NFL1 were already visible in apical buds of vegetative stages in day-neutral and short-day tobacco plants. The expression levels increased during subsequent stages to high levels in 12-week-old plants. In N. sylvestris, expression of NFL1 was not observed in apical buds of plants grown under non-inductive short days. In long days, expression of NFL1 was first detectable in 9-week-old plants at a level similar to that in 12-week-old plants (Fig. 3D).

Modulation of flowering time by constitutive expression of *NtSOC1* or *NtFUL*

Constitutive expression of *SOC1* and *FUL1* in the facultative long-day plant *Arabidopsis* shortened drastically the vegetative phase and caused a photoperiod-independent flowering (Borner et al. 2000; S. Melzer, unpublished result). To analyse whether the orthologous *NtSOC1* and *NtFUL* transgenes might also modulate flowering time in transgenic *Nicotiana* plants, we introduced the coding regions of *NtSOC1* and *NtFUL* under the control of the strong cauliflower mosaic virus 35S promoter into *N. tabacum* cv Hicks, *N. tabacum* MM cv Hicks and

(A), NtFUL (B), NtMADS5 (C) or NFL1 (D) primers on samples from N. tabacum ev. Hicks (Nt), N. tabacum ev. Hicks MM (MM) and N. sylvestris plants (Ns). Amplification of N. sylvestris eIF4A10 is shown as an internal control for the expression in N. sylvestris (E)

N. sylvestris plants. We obtained several independent transgenic lines either expressing the *NtSOC1* or the *NtFUL* transgene. Five transgenic lines of day-neutral tobacco and 10 transgenic lines each from MM cv Hicks and *N. sylvestris* with each transgene were studied in detail. All lines had very high transgene expression levels compared to those of the endogenous genes (Fig. 4).

Since the number of leaf nodes can be used to analyse flowering time in tobacco plants (McDaniel 1996), we compared different genotypes by counting leaf numbers. The NtSOC1 and NtFUL transgenes promoted flowering in each transgenic line under inductive photoperiods (Fig. 5A– E). Day-neutral cv Hicks tobacco wild-type plants flowered after the production of 25 leaves under long days (Fig. 5E, Nt), whereas the different transgenic 35S::NtSOC1 tobacco lines flowered after forming 14–22 leaves (Fig. 5E, T1-1 to T1-5). The transgenic 35S::NtFUL tobacco lines started flowering later after forming 18-22 leaves (Fig. 5E, T2-1 to T2-5). In short days, the wild-type tobacco plants flowered earlier than under long days after forming 18 leaves and the transgenic lines flowered again earlier after the initiation of 9-14 (35S::NtSOC1) or 12-16 (35S::NtFUL) leaves (Fig. 5E).

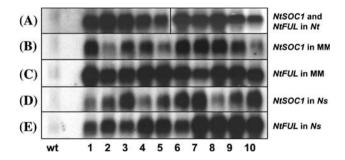
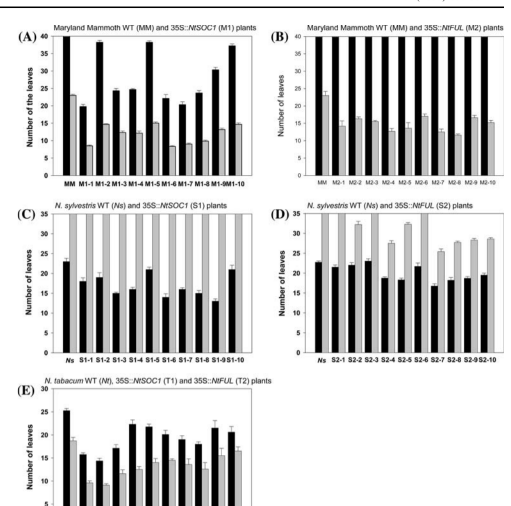


Fig. 4 Expression levels of 35S::*NtSOC1* and 35S::*NtFUL* in different *Nicotiana* varieties. (A) Expression level of 35S::*NtSOC1* (left) and 35S::*NtFUL* (right) transgenes in five analysed lines each from *N. tabacum* cv. Hicks. (B) *N. tabacum* MM cv. Hicks wild-type plants and 10 transgenic lines overexpressing 35S::*NtSOC1*. (C) *N. tabacum* MM cv. Hicks wild-type and 10 transgenic lines overexpressing 35S::*NtFUL*. (D) *N. sylvestris* wild-type plants and 10 transgenic lines overexpressing 35S::*NtSOC1*. (E) *N. sylvestris* wild-type plants and 10 transgenic lines overexpressing 35S::*NtFUL*



Fig. 5 Flowering time analysis of transgenic Nicotiana plants overexpressing NtSOC1 or NtFUL. Comparison of leaf numbers of wild-type and different transgenic Nicotiana lines grown either under long (black bars) or under short days (grey bars). (A) N. tabacum cv. Hicks MM wild-type (MM) and transgenic plants overexpressing 35S::NtSOC1 (M1-1 to M1-10). (B) N. tabacum cv. Hicks MM wild-type (MM) and transgenic plants overexpressing 35S::NtFUL (M2-1 to M2-10). (C) N. sylvestris wild-type (Ns) and transgenic plants (S1-1 to S1-10) overexpressing 35S::NtSOC1. (D) N. sylvestris wild-type (Ns) and transgenic plants (S2-1 to S2-10) overexpressing 35S::NtFUL. (E) N. tabacum ev. Hicks wildtype (Nt) and transgenic plants (T1-1 to T1-5 for 35S::NtSOC1 and T2-1 to T2-5 for 35S::NtFUL lines). Bars reaching the top of the graphs, without an error bar, represent plants that did not flower under the particular conditions

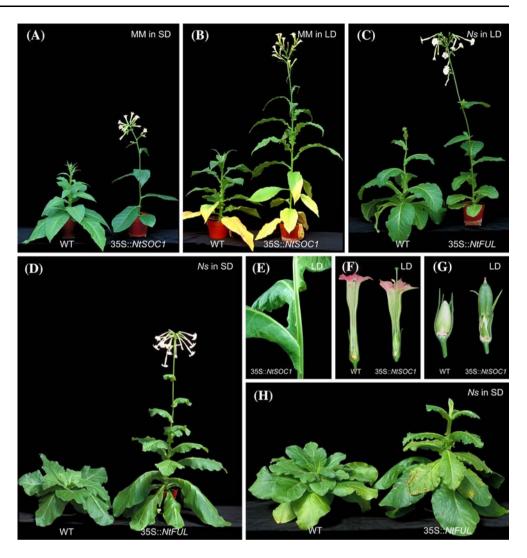


Nicotiana tabacum MM cv Hicks wild-type plants had a short initial rosette stage under inductive short days and flowered after forming 23 leaves, whereas the transgenic lines overexpressing NtSOC1 had no initial rosette stage (Fig. 6A) and flowered after forming 8–15 leaves (Fig. 5A, M1-1 to M1-10). N. tabacum MM cv Hicks wild-type plants never flowered under our experimental conditions in long days. However, all 35S::*NtSOC1* lines flowered under non-inductive conditions after the formation of 20-38 leaves (Figs. 6B and 5A, M1-1 to M1-10). Transgenic MM cv Hicks lines overexpressing NtFUL also flowered earlier under inductive short days after forming 12–17 leaves, but these transgenic lines never flowered in non-inductive long days (Fig. 5B, M2-1 to M2-10). N. sylvestris plants overexpressing 35S::NtSOC1 had no initial rosette stage and flowered earlier under inductive long days (Fig. 6C). The leaf number was reduced from 23 in wild-type plants (Fig. 5C, Ns) to 13–21 leaves in the transgenic lines (Fig. 5C, S1-1 to S1-10). Under non-inductive short days, wild-type N. sylvestris plants remained for up to 9 months in the rosette stage before senescing, whereas the

transgenic lines overexpressing NtSOC1 bolted after a short initial rosette stage, but did not flower under these conditions (Fig. 6H). In addition to flowering time effects, we also observed some other pleiotropic phenotypes in transgenic lines overexpressing NtSOC1. In N. sylvestris lines with strong transgene expression, leaves were fused at the base (Fig. 6E), which was never observed in transgenic tobacco or MM cv Hicks plants, indicating that the NtSOC1 transgene interferes with processes in the apical meristem of vegetative N. sylvestris plants. In all transgenic lines of the different *Nicotiana* varieties overexpressing *NtSOC1*, floral structures were also affected. Flowers had a longer style, which consequently raised the stigma above the anthers and prevented self-pollination (Fig. 6F). In addition to the longer style, the pods developed on twisted petioles that were 10-15 mm long (Fig. 6G). N. sylvestris plants overexpressing 35S::NtFUL (lines S2-1 to 2-10) flowered after 17–22 leaves had been formed, only a bit earlier than the wild-type N. sylvestris plants under inductive long days (Fig. 5D, S2-1 to S2-10). However, seven 35S::NtFUL transgenic lines with strong transgene expression levels



Fig. 6 Phenotypes of transgenic Nicotiana plants overexpressing NtSOC1 or NtFUL. (A) Wild-type MM plant (left) flowered later than a 35S::NtSOC1 MM plant (right) under short days. (B) Under long days, MM wild-type plants (left) never flowered, whereas the 35S::NtSOC1 MM plant (right) flowered. (C) N. sylvestris plant overexpressing NtFUL (right) flowered earlier than control plants (left) under inductive long days. (D) N. sylvestris 35S::NtFUL plants (right) overcame the photoperiodic barrier for flowering in short days. (E) N. sylvestris 35S::NtSOC1 plant showing laminar connections of subsequent leaves. (F) N. tabacum flowers overexpressing 35S::NtSOC1 (right) had shorter tubes and longer styles than wild type plants (left). (G) The capsules of N. tabacum 35S::NtSOC1 plants sitting on a petiole (right), absent in wild-type plants. (H) In short days, N. sylvestris 35S::NtSOC1 plants (right) started to bolt, but did not flower under these non-inductive conditions as the N. sylvestris wild-type plants (left)



actually flowered under otherwise non-inductive short-day conditions (Figs. 5D, 6D).

Constitutive expression of *NtFUL* prevents seed dehiscence in transgenic *Nicotiana* plants

Constitutive expression of *FUL* in *Arabidopsis* has been shown to prevent seed dispersal. Since the orthologous gene also might have the same function in *Nicotiana* plants, we analysed whether transgenic lines overexpressing *NtFUL* were altered in fruit opening. Capsules of wild-type *Nicotiana* plants open at maturity from the top (Fig. 7A) and the seeds are dispersed from the capsules by slight movements. However, the capsules of plants overexpressing *NtFUL* remained closed (Fig. 7B) and had alterations at the cellular level. As seen in Fig. 7C, D, the amount of lignified cells and the degree of lignification at the midrib of a carpel was higher in wild-type plants (Fig. 7C) than that of a transgenic line (Fig. 7D). Together with a larger incision in wild-type plants, physical forces of the drying capsule had created

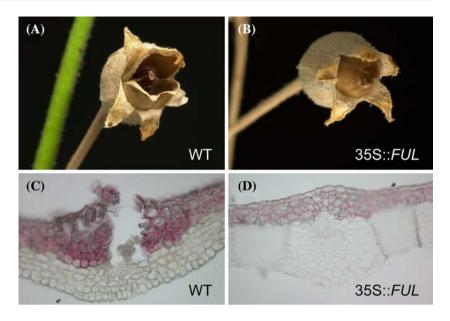
tissue tensions that easily opened the pods, whereas those of the transgenic lines remained closed. This phenotype strongly indicates that *NtFUL* from tobacco is a true orthologs of *FUL* in *Arabidopsis*.

Discussion

The developmental switch to flowering has been studied intensively in different model plants over the last decades. However, only in the last 15 years molecular-genetic studies in *Arabidopsis* have shown that complex networks of genetic pathways converge on integrators to control the transition to reproductive growth. Triple mutants with mutations in key genes of the autonomous, photoperiodic and GA pathways, such as the *fca co ga1* triple mutant, do not flower (Reeves and Coupland 2001), whereas triple *lfy ft soc1* integrator mutants still do (Moon et al. 2005), implying that other genes might act in parallel. *SOC1* and *FUL* have a similar expression pattern during the transition to flowering in apical meristems (Borner et al. 2000;



Fig. 7 Altered pod dehiscence in transgenic *N. sylvestris* plants overexpressing *NtFUL* in long days. (A) Open capsule of a *N. sylvestris* wild-type plant. (B) Closed capsule of a *N. sylvestris* 35S::*FUL* plant. (C) Section through the midrib of a *N. sylvestris* wild-type carpel. (D) Section through the midrib of a *N. sylvestris* 35S::*FUL* carpel

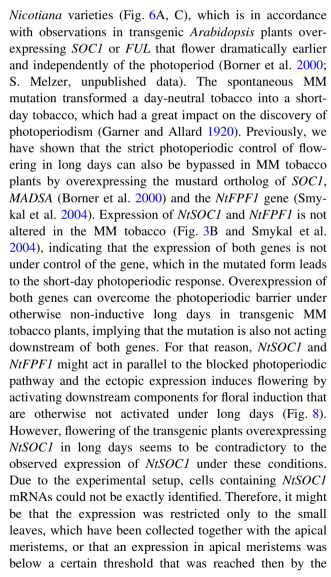


Mandel and Yanofsky 1995) and their proteins interact (de Folter et al. 2005), suggesting that *FUL* might also represent an important factor downstream of flowering time pathways. Therefore, it was intriguing to analyse whether the orthologous genes of *SOC1* and *FUL* also act as flowering time regulators in *Nicotiana* varieties. We identified and characterised the putative orthologs, *NtSOC1* and *NtFUL* from tobacco that were highly expressed in stems of non-flowering plants. Since in situ hybridisations in mustard have revealed that the orthologs, *SaMADSA* and *SaMADSB* are expressed in procambial strands and in inflorescence stem veins (Menzel et al. 1996), these genes as well as *NtSOC1* and *NtFUL* in *Nicotiana* species might also have a physiological function there.

In cv Hicks and MM cv Hicks tobacco plants, expression of *NtSOC1* and *NtFUL* was already observable in vegetative stages under long and short days and the expression increased to high levels until the plants flowered. In *N. sylvestris*, the expression patterns of both genes and that of *NFL1* were different from those in the tobacco varieties. The *SOC1* ortholog was weakly detectable from the 6th week on and increased only slightly during the following weeks in short days. However, no expression of the *NFL1* and *FUL* orthologous genes was detectable under non-inductive short days over the entire growth period, indicating that these genes might be limiting for flowering of *N. sylvestris* in short days.

Modulation of flowering time by *NtSOC1* and *NtFUL* in *Nicotiana* varieties

Overexpression of NtSOC1 and NtFUL shortened the vegetative phase under inductive conditions in all





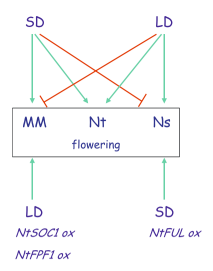


Fig. 8 Diagram for flowering in different *Nicotiana* varieties. *N. sylvestris* (*Ns*) and the MM tobacco flower only under long days or short days (green arrows), respectively, whereas the opposite conditions are inhibitory (red lines) and the day-neutral *N. tabacum* (*Nt*) flowers under both photoperiods. Overexpression of *NtSOC1ox* and *NtFPF1ox* caused flowering under non-inductive long days in the MM tobacco, but not in *N. sylvestris* under short days. Vice versa, overexpression of *NtFULox* leads to flowering in *N. sylvestris* under non-inductive short days, but not to flowering of transgenic MM plants in long days

overexpression of *NtSOC1*, which caused finally flowering under non-inductive long-day conditions.

The FT ortholog from tomato, SINGLE FLOWER TRUST (SFT), also bypassed the photoperiodic requirements of MM tobacco under non-inductive long days (Lifschitz et al. 2006) and the transgenic plants flowered even earlier than those overexpressing NtSOC1 in both photoperiods. FT in Arabidopsis activates in addition to SOC1 also FUL and SEPALLATA3 (SEP3) expression (Abe et al. 2005; An et al. 2004; Schmidt et al. 2003; Teper-Bamnolker and Samach 2005; Wigge et al. 2005). Therefore, the activation of *FUL* and *SEP3* orthologs might also contribute to the earlier flowering phenotype of 35S::SFT MM tobacco plants. However, overexpression of NtFUL alone is not sufficient to induce flowering in the short-day MM tobacco under long days. Similarly, overexpression of NtSOC1 and NtFPF1 (Smykal et al. 2004) did not induce flowering in N. sylvestris under non-inductive short days. But on the contrary, constitutive expression of NtFUL caused flowering in N. sylvestris in short days, in which no expression of the FUL orthologs was observed (Fig. 3B), indicating that FUL is controlled by the photoperiodic pathway and is a limiting factor for flowering under non-inductive conditions. Most obviously, the earliest transgenic line overexpressing NtFUL (S2-7) flowered almost at the same time as did wild-type plants under long days and resembled a wild-type plant flowering under inductive conditions (Fig. 6D). The role of FUL for floral induction in *Arabidopsis* has not been clarified yet. However, the fact that *FUL* is activated by *CO* and *FT* (Schmidt et al. 2003; Teper-Bamnolker and Samach 2005) and overexpression of *CO* in double mutants of *ful* and *soc1* greatly delays flowering compared to overexpression of *CO* in wild-type plants and single mutants (S. Melzer, unpublished results) argues for a specific and highly redundant role of *FUL* for the control of flowering by long days in *Arabidopsis* as well as in *N. sylvestris*.

Like in Arabidopsis, the SOC1 ortholog might also act as a basic integrator in different Nicotiana varieties and might interact with other genes, such as FUL, to execute the signal-integrating function for certain flowering time pathways. Therefore, overexpression of SOC1 alone might be not sufficient to overcome the photoperiodic barrier for flowering time control when other genes are limiting, as observed in N. sylvestris plants in short days, in which the FUL ortholog is not expressed. However, the data so far are also in line with the assumption that NtSOC1 and NtFUL are specifically involved in two different pathways in photoperiodic Nicotiana varieties, which are not activated in non-inductive photoperiods, but can be activated or bypassed, by NtSOC1 or NtFUL overexpression (Fig. 8). Therefore, our results indicate that the behaviour of longand short-day Nicotiana varieties might differ in the requirement for certain MADS box genes to establish floral development in apical meristems after the arrival of the mobile flowering signal FT.

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