Move on up, it’s time for change—mobile signals controlling photoperiod-dependent flowering

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Plants do not bloom randomly—but how do they know when and where to make flowers? Here, we review molecular mechanisms that integrate spatial and temporal information in day-length-dependent flowering. Primarily through genetic analyses in two species, Arabidopsis thaliana and rice, we today understand the essentials of two central issues in plant biology: how the appropriate photoperiod generates an inductive stimulus based on an external coincidence mechanism, and the nature of the mobile flowering signal, florigen, which relays photoperiod-dependent information from the leaf to the growing tip of the plant, the shoot apex.

We all love springtime, not least because flowers are everywhere. But how do plants know it is the right time to bloom? Just like us, plants can recognize the seasons. Among different indicators such as temperature and light, the increase in day length is the most robust tell-tale sign that spring is upon us, at least for those of us who do not live in the tropics. The ability to recognize and respond to changes in day length is known as photoperiodism and is common to both plants and animals. In this review, we will focus on events downstream from the circadian clock that relay the photoperiodic signal from the site of perception, the leaves, to the shoot apex, where flowers are formed. Recent advances in this area have finally provided a molecular underpinning for two long-standing tenets of plant biology, the external coincidence model and the florigen hypothesis.

The discovery of photoperiodic control of flowering

One of the first to recognize the importance of day length in flowering was the English botanist Arthur Henfrey (1852), who suggested that summer day length, which changes with latitude, is an important factor in determining where plants could grow. Some 50 years later, the French scientist Tournois (1912) found that both Humulus (hop) and Cannabis (hemp) flower precociously when kept in winter in a greenhouse supplemented with artificial light, in agreement with Henfrey’s postulate. The German Klebs made similar observations with Semprevivum (house leek). Importantly, he realized that it was unlikely to be light quantity [as a nutritive factor, as he called it] but rather light duration [acting as a catalytic factor] that was critical in this process [summarized in Klebs 1913].

Despite Tournois’ and Klebs’ contributions, the discovery of day length as a crucial factor in flowering control is generally accredited to Wightman Garner and Harry Allard, scientists at the US Department of Agriculture. Garner and Allard (1920, 1923) were working on two practical problems. One was the question of why certain strains of soybeans often would initiate flowers more or less simultaneously, typically during the height of summer, even if farmers had spread out the sowing dates. The other related to a tobacco variety that had spontaneously arisen in the field. This strain, appropriately called “Maryland Mammoth,” would grow very tall, yielding nearly 100 leaves, whereas normal tobacco plants switch to making flowers with only ~20 leaves. The issue here was that this variety did not live up to its full commercial potential, because the delay in flowering often caused Maryland Mammoth plants to succumb to frost before they had produced any seeds, making it difficult to propagate the strain.

The reason that Garner and Allard are remembered for being the first to clearly demonstrate the effect of photoperiod is that they performed controlled experiments. Starting with Biloxi soybean and Maryland Mammoth tobacco plants raised in pots, they exposed one set to natural outdoors conditions, but artificially shortened the day for the other group by moving them into a windowless shed in the afternoon, returning them to the field in the morning. This simple trick greatly accelerated the flowering response of both soybean and tobacco, leading Garner and Allard (1920, 1923) to deduce that these strains only flowered when day length fell under a certain threshold. This neatly explained why the exact...
time point of sowing has not much influence on when Biloxi soybeans start to make flowers or why Maryland Mammoth tobacco produce flowers only late in fall.

It did not take much longer for scientists to realize that flowering plants generally belong to three categories: those in which short days promote flowering or long days prevent it, those in which long days promote flowering or short days prevent it, and a group that does not care either way. These categories are known as short-day, long-day, and day-neutral plants, and are often distinguished further by the adjectives obligate and facultative, depending on whether the corresponding day length requirement is absolute (for review, see Thomas and Vince-Prue 1997). Furthermore, the terms “short days” and “long days” are relative—they merely indicate that one plant will flower when day length falls below its critical photoperiod, while another flowers once it rises above its own specific threshold. Hence, the short day of one plant may actually not be different in length from the long day of another plant.

How plants (and other organisms) measure the length of day and night

The discovery of photoperiodism raised several questions, including which molecules the plant uses to sense light and how these molecules help the plant to measure day length. Perhaps the simplest mechanism one can envision is modeled on an hourglass. For example, every time the light is switched on, a substance that prevents flowering above a certain level begins to accumulate. If this substance is in addition degraded again every night, it will only stay under the critical threshold and hence permit flowering if days are sufficiently short.

Erwin Bünning was the first to propose that plants might use a more sophisticated mechanism. His model was based on the known fact that bean seedlings show a pronounced daily rhythm in leaf movement and that this rhythm is maintained even after plants have been moved to permanent darkness. Key insights of Bünning and his collaborator Kurt Stern (Bünning and Stern 1930) were that the red component of the light spectrum causes the movement of different individual plants to be synchronous and that without the red light cue the period would slowly diverge from the normal 24 h. Bünning correctly concluded that the “biological clock” of plants is partially independent of the daily light/dark cycle. He furthermore found that different strains of beans have endogenous rhythms that can be either shorter or longer than 24 h, and, using crosses, he demonstrated a genetic basis for these differences.

Initially, Bünning thought that the movement of bean leaves had no adaptive value, but rather was just a convenient indicator of what we call today the circadian clock, to reflect its role in time measurement on a 24-h scale. He did, however, believe that an interaction of the endogenous rhythm with external rhythms would be important for plant development. Bünning (1936) subsequently came to appreciate that photoperiodic control of flowering could be a perfect example for where such an interaction might occur. According to the Bünning hypothesis, a 24-h day is divided into a photophile (lightsensitive) and scotophile (dark-sensitive) phase, and the rhythmic alternation of the two is prescribed by an endogenous circadian oscillator. The presence or absence of external light during either phase would tell a plant whether it was exposed to a short or a long day (Fig. 1). The circumstances of the time prevented the ideas laid out by Bünning, an avowed Social Democrat opposed to the Nazis, from gaining immediate and wide acceptance.

From the late 1950s, Colin Pittendrigh, Professor at Princeton and Stanford, became a great advocate of Bünning’s work. Pittendrigh, who was working with insects, formalized and expanded the Bünning hypothesis, coin- ing the term “external coincidence,” since the model posits that the photoperiodic signal is only generated when a specific external phase [light or dark] coincides with the appropriate internal phase of the 24-h cycle. Importantly, light has a dual role in this model: It entrains the circadian oscillation of light- and dark-sensitive phases, and it is directly required for the production of the signal (Pittendrigh 1960). In contrast to a simple hourglass mechanism, what matters in the external co-
incidence model is not the duration of light, but rather the presence of light at specific times during the 24-h cycle. This was shown as early as the 1930s both by using light/dark cycles that were longer or shorter than the normal 24 h and by complex cycles in which dark periods were interrupted by brief light pulses (Fig. 1; Hamner and Bonner 1938; Harder and Bode 1943; Pittendrigh and Minis 1964). In some cases, night breaks as short as 1 min are effective in preventing flowering of short-day plants, a finding incompatible with a simple hourglass model (Hamner and Bonner 1938).

Despite the success of the external coincidence model in explaining the effects of different light treatments, Pittendrigh (1972) realized that a related mechanism, relying only on internal rhythms, might be exploited as well in photoperiod measurement. In the internal coincidence model, external cues—e.g., lights on and lights off—entrain two different oscillations. Only when these oscillations are in synchrony is the inductive or repressive signal triggered (Fig. 1).

Where plants measure the length of day and night

Having discussed insightful studies from the premolecular era that allowed many inferences about how plants measure day length, we now turn to the question of where this process occurs. Flowers form at the growing tip of the plant, the shoot apex. Because the delicate shoot apex often has a protective covering of leaves, it might not be the best place for plants to interpret ambient light conditions. Julius Sachs—Professor of Botany in Bonn, Freiburg, and Würzburg, and one of the mentors of Klebs—had conducted experiments in which he darkened parts of a plant, and from this he concluded that leaves in the light produce flower-forming substances (summarized in Sachs 1865). This was demonstrated directly in the 1930s by exposing only leaves, but not the shoot apex, to flower-inducing photoperiods (Knott 1934). The Russian botanist Mikhail Chailakhyan formalized this as the florigen hypothesis, with florigen being defined as a substance that is generated in leaves under inductive photoperiods and translocated from leaves to the shoot apex (Chailakhyan 1936a,b, 1937). The florigen hypothesis was soon confirmed experimentally, by studies in a variety of systems (for review, see Thomas and Vince-Prue 1997). For example, in Xanthium [cocklebur], exposing a single leaf to one short-day cycle is sufficient to induce flowering, even when plants are subsequently returned to noninductive long days (Naylor 1941). Naylor, as well as others, noticed that initiation of flowers is much more robust after noninduced leaves are removed, suggesting that stimulating export of photosynthetic assimilates through the phloem stream causes also increased export of florigen from the induced leaf. It was later demonstrated that florigen indeed moves with about the same velocity as photosynthetic assimilates, further supporting the notion that the phloem portion of the plant vasculature is the conduit for florigen (e.g., see King et al. 1968).

But the most stringent tests of the florigen hypothesis were those that employed grafting of induced leaves onto noninduced plants. Georg Melchers (1937), who, like Büning, spent much of his career in the Southern German town of Tübingen, showed that leaves of both short-day- and long-day-requiring strains of Hyoscyamus [henbane] produce a signal that can cause a biennial Hyoscyamus strain to flower in the first year [biennial plants normally have an obligate requirement for passing through winter before they will flower]. A final example from this era is provided by the work of Jan Zeevaart (1958), who later collaborated with Melchers’ student Anton Lang [e.g., see Lang and Melchers 1948]. Zeevaart (1958) showed that an induced leaf of Perilla [shiso, an herb widely used in many East and Southeast Asian cuisines] retains its ability to cause flowering of a recipient plant, even if it had been moved from inducing to noninducing conditions 3 mo ago. This is, however, not universal. The induced leaves of Ipomoea [Pharbitis] nil (morning glory) produce florigen only for a few days in the absence of a continuous stimulus (Imamura 1967).

Genetics to the rescue

Despite the initial rapid advances in understanding central features of photoperiod measurements and signaling, further attempts at isolating florigen or revealing its molecular nature failed. This led some to propose that florigen was not a single substance, but rather a mixture of different molecules. Some went so far to suggest that it might even involve an electric signal [Karege et al. 1982].

Most of the essential components are now known, largely thanks to pioneering genetic analyses done with Arabidopsis thaliana, a facultative long-day plant (Napp-Zinn 1969). Popular laboratory strains flower within a few weeks under long days, but much later in short days. While variation in flowering behavior of naturally occurring A. thaliana strains had already been noticed by Friedrich Laibach (1951), George Rédei (1962) was the first to isolate flowering mutants in a defined background. Loss-of-function mutations at three loci, CONSTANS (CO), GIGANTEA (GI), and LUMINIDEPENDENS (LD), cause plants to flower later than the standard strain under inductive long days. Rédei had noticed that the co line flowers at about the same time in long and short days, which was the reason he christened the mutant constans. Rédei was, however, more interested in the fact that the late-flowering strains appeared to be more vigorous [reflected by the name gigantea for one of the other mutants], and he did not further exploit them to dissect the physiology of flowering. This was left to Maarten Koornneef et al. (1991), who performed the first large-scale genetic screens for mutants with abnormal flowering behavior. Koornneef used less colorful names than Rédei, and named all of them “f” [for flowering] followed by one or two additional letters (fca, fd, fe, fha, fpa, ft, fve, fwa, and fy). Because some journals insist that one spell out gene names, we later invented the name FLOWERING LOCUS T for the FT gene [Karailsky et al. 1999]. In contrast to Rédei, Koornneef used these mutants to focus on the control of flowering...
time. He observed that several flower much later than the wild type in long days, but that they are about as late as wild type under noninductive short days. He therefore postulated that the CO, FT, and GI genes, and possibly also FHA, are specifically involved in the perception of the photoperiod signal [Koornneef et al. 1991], which we today know from molecular analyses to be correct.

Koornneef isolated additional mutants impaired in several different processes, ranging from light and hormone response to floral organ formation, and his successes inspired many others to conduct their own mutant hunts. In addition to morphological traits, several scientists began to exploit reporter genes, and one of these screens led to the isolation of mutants defective in the circadian clock [Millar et al. 1995]. Many clock genes have in the meantime been cloned, with additional factors identified by reverse genetics, and their functional analysis, coupled with whole-genome studies, has allowed the development of sophisticated models of circadian regulation in A. thaliana [Locke et al. 2006; Zeilinger et al. 2006]. For a detailed discussion of the circadian clock, the reader is referred to recent reviews [Gardner et al. 2006; McClung 2006].

**Molecular basis of the external coincidence model**

It is primarily through the analysis of two genes that we understand today the molecular basis of the external coincidence model in plants: CO, which encodes a B-box zinc finger protein [Putterill et al. 1995], and FT, which encodes a small transcription cofactor [Fig. 2; Kardailsky et al. 1999; Kobayashi et al. 1999].

Although the exact biochemical function of CO protein is not entirely understood, CO is an important regulator of FT mRNA expression [Kobayashi et al. 1999; Samach et al. 2000; Suárez-López et al. 2001; Yanovsky and Kay 2002; An et al. 2004]. Genome-wide analyses with loss-of-function mutants have confirmed FT as the major early target of CO [Wigge et al. 2005]. CO on its own does not bind DNA, but it likely acts as part of a CCAAT-box-binding complex involving HAP proteins [Ben-Naim et al. 2006; Wenkel et al. 2006; Cai et al. 2007]. CO is probably recruited to the FT promoter, based on experiments with an overexpressed version of CO that can be post-translationally controlled. However, because several other potential direct targets identified in this study [Samach et al. 2000] could not be confirmed in analyses of co mutants [Wigge et al. 2005], additional experiments are needed to demonstrate unambiguously that FT is directly regulated by CO. FT is known to be a direct target of FLOWERING LOCUS C (FLC) and SHORT VEGETATIVE PHASE (SVP), transcriptional repressors that mediate the effects of exposure to winter-like conditions or to more modest changes in ambient temperature [Hellwell et al. 2006; Searle et al. 2006; Lee et al. 2007]. In addition, chromatin modification appears to play an important role in preventing inappropriate FT expression [Piñeiro et al. 2003; Takada and Goto 2003; Germann et al. 2006; Turck et al. 2007].

When the Weigel and Araki groups [Kardailsky et al. 1999, Kobayashi et al. 1999] cloned the CO gene, both overlooked that FT mRNA abundance has a circadian cycle. Since CO mRNA had been reported to accumulate at the shoot apex, where flowers are produced [Simon et al. 1996], the focus at the time was on processes thought to be much downstream from the photoperiodic signal. It was therefore a major advance when it was discovered that FT mRNA expression is not only much higher in long days, but also has a circadian pattern, with peak levels during the second half of a 24-h cycle [Harmer et al. 2000; Suárez-López et al. 2001]. There is very little FT expression in short days, but upon transfer from short to inductive long days, FT is immediately induced in a CO-dependent manner, and returning plants to short days causes FT expression to subside within a day, indicating a very immediate effect of day length on FT mRNA accumulation [Imaizumi et al. 2003; Corbesier et al. 2007].

In contrast to FT, CO mRNA is easily detected in both long and short days. Its abundance cycles regardless of day length, with a prominent peak in the night following either a long or a short day, and an earlier shoulder in the afternoon only in long days [Fig. 2; Suárez-López et al. 2001]. The observation that CO mRNA expression does

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**Figure 2.** Photoperiod-dependent activation of FT mRNA expression in the leaf of the long-day plant A. thaliana. CDF1 mRNA is expressed highly at the beginning of the day. FKF1 protein is only active in the light and causes destabilization of CDF1 protein, so that in long days, CDF1 protein is at very low levels during the second half of the light phase. Since CDF1 represses CO mRNA expression, this in turn allows for an earlier increase of CO mRNA levels in long days. CO protein is only stable in the light, hence it accumulates only in long days. It in turn activates FT mRNA expression.
not automatically lead to accumulation of FT mRNA suggested that the CO/FT module might be at the heart of a molecular external coincidence mechanism. A first line of indirect evidence in support of this hypothesis was provided by Yanovsky and Kay (2002), who studied the toc1 (timing of CAB expression 1) mutant, in which the period of the endogenous clock is shortened to ~21 h compared with ~24.5 h of the wild type. While many other clock mutations alter both the phase and level of CO mRNA expression (Suárez-López et al. 2001), toc1 affects only the phase angle of CO, such that CO mRNA peaks earlier in short days and therefore in the light, while high levels of CO mRNA are restricted to the dark in short-day grown wild-type plants (Yanovsky and Kay 2002). When the day–night cycle is shortened to 21 h and thus matched to the endogenous period in toc1 mutants, the CO pattern no longer differs from that in wild type and the short-day early flowering phenotype of toc1 is suppressed (Strayer et al. 2000; Yanovsky and Kay 2002).

Additional, more sophisticated manipulations of length and structure of the light/dark cycle, similar to what had been used in the early days of studying the circadian clock (Nanda and Hammer 1958; Pittendrigh 1960), confirmed that bringing peak levels of CO mRNA into the light phase promotes flowering of wild-type plants under short photoperiods (Rodén et al. 2002).

That FT is only induced when high levels of CO mRNA are present in the light pointed to post-transcriptional regulation of CO activity. In an elegant series of experiments, George Coupland and colleagues (Valverde et al. 2004) demonstrated that far red and blue light, which are respectively perceived by phytochrome A and cryptochrome 2 (the latter encoded by the flowering time gene FHA; Guo et al. 1998), regulate the stability of CO protein through inhibition of proteasome-dependent CO degradation. Thus, the light-sensitive flowering regulator that Bünning (1936) proposed to be produced with a circadian rhythm does indeed exist: It is the CO protein. In the dark, CO is likely recruited to the proteasome by SUPPRESSOR OF PHYA-105 (SPA) proteins, which in turn interact with the E3 ubiquitin ligase CONSTITUTIVE PHOTOMORPHOGENIC 1 [COP1] (Laubinger et al. 2004, 2006).

Interestingly, light of another quality, red light, which is perceived by phytochrome B, promotes, rather than inhibits, degradation of CO, and thereby delays flowering (Mockler et al. 2003; Valverde et al. 2004). The mechanism remains unclear since misexpression experiments have suggested a nonautonomous effect of phytochrome B on CO (Endo et al. 2005). Finally, the PHYTOCHROME AND FLOWERING TIME 1 [PFT1] protein has been proposed to specifically mediate the effects of phytochrome B on FT independently of CO (Cerdán and Chory 2003). Since PFT1 has recently been identified as a subunit of Mediator, a multiprotein complex that has a general role in transcriptional regulation (Bäckström et al. 2007), this hypothesis may need to be revisited.

The function of CO and FT is likely to be universal, since their orthologs in rice, Heading-date 1 (Hd1, also known as Photoperiod Sensitivity 1, Se1) and Hd3a, play very similar roles. Not only is rice, a grass, a distant relative of A. thaliana among flowering plants, but it is also a short-day plant. The groups of Masahiro Yano, Ko Shimamoto, and Takeshi Izawa (Yano et al. 2000; Izawa et al. 2002; Kojima et al. 2002; Hayama et al. 2003) showed that, similar to the situation in A. thaliana, Hd1/Se1 and Hd3a promote flowering (or heading as it is called in rice) in response to inductive photoperiods, in this case short days. In addition to activating Hd3a under inductive conditions, Hd1/Se1 represses Hd3a under noninductive conditions. Plants with compromised Hd1/Se1 activity thus flower later than wild type in short days (under inductive conditions), but earlier in long days (under noninductive conditions) (Fig. 3; Yano et al. 2000). The situation might be similar in A. thaliana, as some investigators, though not all, found co mutants to flower a bit earlier than wild type in short days (Ređei 1962; Koornneef et al. 1991; Reeves and Coupland 2001).

Intriguingly, the repressive function of Hd1/Se1 requires the red-light receptor phytochrome B. Loss of phytochrome B activity therefore leads to early flowering, similar to the effects of phytochrome B mutations in A. thaliana (Reed et al. 1993; Ishikawa et al. 2005), albeit for a different reason: While phytochrome B antagonizes the activation of FT by CO in A. thaliana (Cerdán and Chory 2003; Halliday et al. 2003; Valverde et al. 2004; Endo et al. 2005), it stimulates the repression of Hd3a by FT.
Hd1/Se1 in rice (Fig. 3; Ishikawa et al. 2005). A further major difference between the two species emerged from the discovery of Early heading date 1 (Ehd1), which encodes an apparent transcription factor with a GARP DNA-binding domain. Ehd1, which is expressed independently of Hd1/Se1, is a positive regulator of Hd3k and other FT-like genes in rice, but has no obvious counterpart in the A. thaliana genome (Doi et al. 2004).

As alluded to earlier, CO expression profiles are similar but not identical under long and short days, with substantial levels of CO mRNA detected earlier and in the light when photoperiods are long (Suárez-López et al. 2001). The initial shoulder that precedes the later peak of CO mRNA in long days requires activity of FLAVIN-BINDING, KELCH REPEAT, F-BOX 1 [FKF1], which itself is transcriptionally regulated by the circadian clock (Fig. 2). FKF1 mRNA has a rather narrow peak of expression, which is in the second half of a long day, but toward the end of a short day. The FKF1 protein, a blue-light receptor, shows a broader peak, resulting in FKF1 being present during much of the second half of the light phase in long days, but only during the dark phase in short days (Imaizumi et al. 2003). Regulation of FKF1 activity by light provides a mechanism for how FKF1 promotes the daytime shoulder of CO mRNA in long days. Light-activated FKF1 affects CO mRNA accumulation by destabilizing CYCLING DOF FACTOR 1 [CDF1], a direct repressor of CO transcription [Imaizumi et al. 2005]. This occurs most likely through an SCF complex that includes FKF1 as specificity component (Han et al. 2004; Takahashi et al. 2004). Whether CDF1 and its homologs are the only factors mediating FKF1 activity still needs to be demonstrated, since so far only partial CDF loss-of-function lines with rather weak phenotypes have been examined [Imaizumi et al. 2005]. Among other factors that have less precisely defined roles but are also important for shaping the CO mRNA profiles is RED AND FAR-RED INSENSITIVE 2 [Rffi2], a RING-domain zinc finger protein [Chen and Ni 2006]. In summary, photoperiod controls both the mRNA expression pattern of CO and the stability of CO protein, even though the Bünning model would have required only one level of regulation.

**Molecular basis of florigen**

Now that we have discussed the molecular essence of the external coincidence model, we turn to the second big question in flowering biology: the molecular nature of florigen. Grafting experiments demonstrated that CO acts upstream of the mobile signal, making the CO target FT an excellent candidate for florigen [An et al. 2004; Ayre and Turgeon 2004]. The FT promoter is active in the companion cells of the phloem, and in situ hybridization in a mutant with derepressed FT expression confirmed that the transcript accumulates in the phloem (Takada and Goto 2003; An et al. 2004). FT expression in phloem companion cells is essential for its function, as recently shown by tissue-specific inactivation of FT mRNA using artificial microRNAs [Mathieu et al. 2007]. Consultation of an expression atlas based on microarray experiments confirmed that FT mRNA is more abundant in leaves, particularly in cotyledons and the distal portion of older leaves, than at the shoot apex (Wigge et al. 2005).

Different from FT, tissue-specific overexpression experiments indicated that CO is ineffective when present only at the shoot apex, while specific overexpression in the phloem of leaves has similar effects as ubiquitous overexpression [An et al. 2004; Ayre and Turgeon 2004; Corbesier et al. 2007]. In stark contrast, tissue-specific overexpression of FT causes early flowering, regardless of whether this occurs in the phloem of leaves and stem or at the shoot apex [An et al. 2004], which was the first result that immediately suggested FT exerting its role at the shoot apex.

The initial sequence analysis of FT had not been very revealing, as FT was found to encode a small globular protein with structural similarity to a variety of enzyme inhibitors in fungi and animals, including Raf kinase inhibitor protein [Kardailsky et al. 1999; Kobayashi et al. 1999; Ahn et al. 2006]. FT might play a general role as a molecular scaffold [Pnueli et al. 2001], but the best understood function is as a transcriptional cofactor, interacting directly with the bZIP protein FD. In contrast to FT, FD mRNA is found only at the shoot apex, with a moderate increase in expression levels upon photoperiodic induction that appears to be FT independent [Abe et al. 2005; Wigge et al. 2005]. At the shoot apex, FD likely recruits FT to the promoter of APETALA1 [AP1], an important regulator of flower fate, although this conclusion is based on experiments with FD-overexpressing plants, and the interaction found was fairly weak. The role of FT as a transcriptional cofactor is, however, supported by the effects of fusing it to the heterologous VP16 activation domain [Wigge et al. 2005].

The finding that photoperiod-activated CO regulates FT transcription in the phloem of leaves, but FT protein acts at the shoot apex, suggested that FT mRNA or protein is translocated from the leaf to the shoot apex, and that one of them is the mobile florigen signal. The companion cells, in which FT is transcribed, directly contact the sieve elements, which form the conducting tissue of the phloem. It has been known for several years that small globular proteins such as free green fluorescent protein [GFP] not only readily enter sieve elements from the companion cells [most likely through intercellular channels called plasmodesmata], but also that they are transported across large distances through the phloem and can be unloaded from the phloem into surrounding tissue [Imlau et al. 1999; Stadler et al. 2005]. At 27 kDa, GFP is considerably larger than FT, a globular protein of 20 kDa [Ahn et al. 2006], suggesting that FT moves between cells at least as well as free GFP. Taken together, the simplest scenario is that FT protein itself constitutes the long-distance signal [Abe et al. 2005; Wigge et al. 2005].

Unfortunately, there was some initial confusion because of erroneous reports of long-distance trafficking of FT mRNA [discussed in Zeevaart 2006]. In a study that
met with initial skepticism because of this earlier report, and that needs to be appreciated more widely, Eliezer Lifschitz et al. (2006) were the first to address directly the question of FT mRNA versus protein as florigenic substance, employing grafting techniques. The SINGLE FLOWER TRUSS (SFT) gene of tomato is the FT ortholog, and even though tomato is day neutral, sft mutants flower very late. grafts from SFT-overexpressing plants (albeit not from wild-type plants) consistently and strongly rescued the flowering defect of sft mutants. Since no SFT mRNA could be detected in the recipient, these authors concluded that either SFT protein itself is the mobile signal or that SFT elicits a secondary mobile signal (Lifschitz et al. 2006). In tomato, the FD ortholog SPBG is expressed widely (Pnueli et al. 2001), and such a secondary signal is a distinct possibility. In A. thaliana, however, this seems less likely, since FT appears to act directly at the shoot apex. Importantly, Lifschitz et al. (2006) also demonstrated the universality of the systemic signal, by showing that a graft of tomato overexpressing SFT could accelerate flowering in the classic Maryland Mammoth variety of tobacco, thereby reproducing, using modern tools, the findings of Melchers (1937) with henbane.

A string of recent papers has explored the molecular basis of the florigen signal in additional detail (Corbesier et al. 2007; Jaeger and Wigge 2007; Lin et al. 2007; Mathieu et al. 2007; Tamaki et al. 2007). Two of these studies (Jaeger and Wigge 2007; Tanaki et al. 2007) even claim to have demonstrated movement of FT protein from leaves to the shoot apex, even though direct experimental evidence has not been produced. Nevertheless, together with the previous findings (Abe et al. 2005; Wigge et al. 2005; Lifschitz et al. 2006), the arguments for FT protein (or its orthologs) being the long-distance signal are now quite strong.

Several investigators made use of lines that overexpressed FT fusion proteins in phloem companion cells. Coupland and colleagues (Corbesier et al. 2007) showed that an FT-GFP fusion protein in A. thaliana could move across a graft junction from the shoot to the root, in line with known properties of other GFP fusion proteins. However, in contrast to most GFP fusions, and similar to free GFP alone, FT-GFP can be unloaded from the phloem near the shoot apex (Stadler et al. 2005; Corbesier et al. 2007). Although even the overexpressed FT-GFP could not be detected in the entire domain where the FT partner FD is expressed, the effects on flowering suggested that protein below the detection threshold was present throughout the shoot apical meristem (Corbesier et al. 2007). Export from the phloem was also demonstrated by Jaeger and Wigge (2007), using a fusion of FT to a small Myc tag, importantly, export from the phloem was prevented when the FT-Myc protein was directed to the nucleus with a heterologous nuclear localization signal. This finding is consistent with a general feature of many plant proteins, for which trafficking to adjacent cells is only prevented by efficient subcellular targeting, as proposed by Wu et al. (2003) and recently demonstrated for the SHORTROOT transcription factor (Cui et al. 2007). Markus Schmid and colleagues (Mathieu et al. 2007) also trapped FT in phloem companion cells by fusing it to a nuclearly localized triple-GFP protein. Exploiting transgenically expressed tobacco etch virus (TEV) protease and a TEV protease recognition site engineered into this fusion, they could then release FT from this much larger precursor, which in turn caused early flowering. Similar conclusions regarding the export of an Hd3a-GFP fusion from the phloem were reached by Shimamoto’s group (Tamaki et al. 2007) using rice. In this case, the fusion appeared to travel considerably further than in A. thaliana; however, as the authors duly note, free GFP shows a similar movement behavior, making it difficult to interpret the significance of their findings.

Change-of-function experiments, in which genes are misexpressed, have the disadvantages that expression levels are difficult to control and that one can never be quite sure that the promoter used is indeed completely silent in a specific tissue. Furthermore, all the experiments of “complementing” co and ft mutants with CO and FT linked to a variety of different promoters actually led to much earlier flowering than seen in wild type. As already discussed, grafting experiments provide a more stringent assay for long-distance movement (An et al. 2004; Ayre and Turgeon 2004; Lifschitz et al. 2006; Corbesier et al. 2007). This route was also taken by the laboratory of Bill Lucas (Jorgensen et al. 1998), who has a long-standing interest in long-distance signaling, particularly by RNA molecules. The Lucas group (Lin et al. 2007) exploited the graft compatibility of two cucurbit species with different photoperiodic requirements for flowering. Their approach was particularly elegant, because sequence differences between the two species allowed differential detection of host and donor FT proteins after grafting. Using mass spectrometry, Lin et al. (2007) found FT protein from the donor in the phloem sap of the host, while this was not the case for FT mRNA. Importantly, similar to classic grafting experiments, induction of flowering in this system did not require overexpression in the donor. FT protein has also been detected by mass spectrometry in phloem sap of Brassica napus (rapeseed), the caveat being that these experiments were performed on inflorescence stems; that is, long after the plants had been stably induced to flower (Giavalisco et al. 2006).

In conclusion, while it so far has not been possible to directly identify FT protein that has been produced in leaves in the very cells at the shoot apex that also express its partner FD [an admittedly technically very challenging experiment], little doubt remains that FT (along with paralogs such as TSF; Michaels et al. 2005; Yamaguchi et al. 2005; Mathieu et al. 2007) is a major component of florigen. It is difficult to exclude, and indeed is perhaps likely, that other molecules appear in supporting roles. For example, several microRNAs (miRNAs) function in flowering control [e.g., see Aukerman and Sakai 2003; Schmid et al. 2003; Chen 2004; Schwab et al. 2005]. While there is not yet any experimental evidence for long-range action of these miRNAs,
some of them have been detected in phloem sap as well [Yoo et al. 2004].

**Molecular events at the shoot apex**

In *A. thaliana*, *FT* and *FD* are characterized by complementary expression patterns. *FT* expression is activated in a day-length-dependent manner, but does not provide regional specificity for flower initiation, since it is activated primarily in leaves. Positional information for flower formation requires *FD*, which is expressed at the shoot apex, albeit largely independently of inductive photoperiods. The outcome is that through the interaction of *FT* and *FD* proteins, flowers are formed at the right time, in the right place [Abe et al. 2005; Wigge et al. 2005].

How, then, does the *FT/FD* complex control the formation of flowers, which start out as groups of undifferentiated cells at the flanks of the shoot apical meristem? The only direct target of this complex proposed so far is *AP1* (Wigge et al. 2005). As pointed out by Searle et al. [2006], expression of *AP1* is a relatively late event during floral induction, preceded by the activation of several other positive floral regulators, such as *FRUITFULL* ([FUL], *SUPPRESSOR OF OVEREXPRESSION OF CO 1* [SOC1, also known as AGAMOUS-LIKE 20 (AGL20)], its paralog *AGL42*, and *AGL24*, all of which like *AP1* encode MADS domain proteins. Apart from delayed activation of these genes at the shoot apex of *ft* or *fd* mutants, several are affected by misexpression of *FT* or *FD* [Yu et al. 2002; Schmid et al. 2003; Abe et al. 2005; Moon et al. 2005; Teper-Bammolker and Samach 2005; Wigge et al. 2005; Searle et al. 2006]. Many of these proteins interact with each other, as well as with *SVP*, a MADS domain protein that represses flowering [de Folter et al. 2005].

The effects of *FT/FD* on both early and late flowering activators at the shoot apex suggest a feed-forward mechanism, in which activation of early factors such as *SOC1* enhances the direct activation of later factors such as *AP1*. A feed-forward mechanism, with *FT/FD* affecting expression of late targets such as *AP1* both directly and indirectly through *SOC1*, would explain the relatively strong effect of *soc1* mutations in a *ft* background [Moon et al. 2005; Yoo et al. 2005]. Intriguingly, negative feedback of *AP1* on *SOC1*, *AGL24*, and *SVP* expression contributes to the formation of normal flower primordia, even though these effects are relatively modest [Yu et al. 2004; Liu et al. 2007].

A similar theme of dual levels of interconnected regulation is apparent in the roles of the floral repressor *FLC*, which mediates the effects of winter-like conditions. *FLC* counteracts activation of *FT* by *CO* in the leaf and, in addition, directly represses expression of the *FT* cofactor *FD* at the shoot apex [Searle et al. 2006]. Finally, apart from being a target of *FT* at the shoot apex, as first shown by Schmid et al. [2003], *SOC1* is also expressed in leaves, and it is negatively regulated by *FLC* at both sites [Borner et al. 2000; Lee et al. 2000; Samach et al. 2000; Hepworth et al. 2002; Schmid et al. 2003; Michaels et al. 2005; Helliwell et al. 2006; Searle et al. 2006]. In leaves, *SOC1* is regulated independently of *CO* by the clock [Blázquez et al. 2002; Fujiiwara et al. 2005; Wigge et al. 2005]. Whether or how *SOC1* in leaves might affect flowering is, however, unknown, since *FT* expression in leaves appears to be unaffected by loss of *SOC1* activity [Lee et al. 2000; Moon et al. 2005].

Even in the absence of *FT* and *FD* activity, all the early apex regulators such as *SOC1* are eventually activated, revealing a photoperiod-independent component of regulation. A good candidate for such a pathway is one that relies on gibberellin hormones, which not only regulate, for example, *SOC1* [Borner et al. 2000; Blázquez et al. 2002; Moon et al. 2003], but also the flower meristem identity gene *LEAFY* (*LFY*) [Blázquez et al. 1998]. In contrast to *FT*, *FD*, *SOC1*, and so on, *LFY* activity is essential for the development of normal flowers, acting partially redundantly in flower formation with the *FT/FD/ AP1* axis [Weigel et al. 1992; Ruiz-Garcia et al. 1997; Abe et al. 2005; Wigge et al. 2005]. While early studies suggested that *LFY* functions primarily in parallel with *FT*, it is now clear that *LFY* is both a target of the *FT* and the gibberellin pathways [Nilsson et al. 1998; Blázquez and Weigel 2000; Schmid et al. 2003; Eriksson et al. 2006]. Furthermore, contrary to common wisdom, gibberellins have a clear effect on flowering in long days, although it is not known whether they also function downstream from the photoperiod pathway in *A. thaliana*, as suggested for other species [Pharis et al. 1987; Blázquez et al. 1998; Griffiths et al. 2006; King et al. 2006; Willige et al. 2007]. The regulatory interactions described in this and earlier sections are summarized in Figure 4.

**Research into photoperiodic flowering—time to move on?**

While core aspects of how photoperiod is perceived and how the signal is relayed from the leaves to the shoot apex are known, by no means do we completely understand the flowering process. Many open questions remain. For example, early apex regulators such as *FD*, *SOC1*, or *FUL* show a different pattern of expression from their targets *LFY* and *AP1*. The latter are expressed specifically in flower primordia, while the early regulators are found also, or mainly, in the shoot apical meristem, with their exact spatial relationships not known in much detail. Application of sophisticated live-imaging techniques, as they have been used for the study of meristem regulators [e.g., see Heisler et al. 2005], is almost certain to provide important new insights.

Another powerful approach is the manipulation of the spatial distribution of gene activity, but this has so far been mostly achieved by tissue-specific overexpression, which almost always leads to gain-of-function phenotypes and thus confounds the interpretation of results. A more sensitive assay is the ablation of normal gene function in specific tissues or cells. A new and very effective tool for this purpose is available in the form of artificial miRNAs [amiRNAs] [Alvarez et al. 2006; Schwab et al. 2006], and these have already been employed to demon-
strate that FT and TSF not only can function, when over-expressed, in phloem companion cells [An et al. 2004; Abe et al. 2005], but that they also, and perhaps more importantly, are required in these cells [Mathieu et al. 2007]. Extension of this approach would be rather useful to formally demonstrate that clock components upstream of CO, as well as CO itself, are required in phloem companion cells. It could also be used to revisit the question of CO function outside the leaves. CO mRNA has been detected by in situ hybridization at the shoot apex (Simon et al. 1996), and both reporter fusion experiments [An et al. 2004] and microarray data (Schmid et al. 2003, 2005) confirm that CO mRNA levels at the shoot apex are quite substantial.

A related question is why CO mRNA peaks during the night under both long and short photoperiods, since it appears to be the earlier FKF1-dependent induction of CO that is primarily responsible for the effect of long photoperiods on flowering [Suárez-López et al. 2001; Imaizumi et al. 2003]. It is noticeable that all CO regulators show a circadian expression pattern. Thus, the late peak of CO expression might simply be a consequence of upstream factors that are expressed in a circadian fashion, which may further point to built-in redundancy in CO regulation. It might be possible to address this issue by combining amiRNAs with circadian promoters, to ablate CO mRNA expression during specific times of the 24-h cycle. The late peak of CO expression might also represent an atavism important in species that discriminate more finely between different daylengths than A. thaliana does. In this context, studies of plants that show intraspecific variation in photoperiod response are particularly promising [Böhlenius et al. 2006].

A further problem that begs more extensive study is the mechanism that underlies maintenance of the florally induced state. When A. thaliana plants are returned from inductive long days to noninductive short days, FT expression in leaves ceases within a day, even though the transition to flowering has been irreversibly triggered [Corbesier et al. 2007]. Could the answer lie in the surprising expression pattern of FT outside the rosette leaves? After the transition to flowering, FT is most highly expressed in stem leaves on the inflorescence, in sepals, which are the green, leaf-like organs surrounding flowers, and in developing fruits and seeds [Kobayashi et al. 1999]. It is noticeable that the FT repressor FLC is also highly expressed in developing seeds, but at a later stage (Schmid et al. 2005).

Finally, the CO/FT module is being redeployed by plants in other photoperiod-dependent processes, such as
in trees. The peak of CO expression is later during the day–night cycle for northern aspen trees than for southern trees. Thus, when exposed to 20-h days, the CO peak already falls into the dark for the northern trees, FT is no longer activated, and growth is not supported anymore. Conversely, the CO peak for the southern trees is still in the light, FT remains active, and dormancy continues to be repressed [after data from Böhlenius et al. 2006].

Figure 5. Latitudinal variation in CO-dependent FT expression in trees. The peak of CO expression is later during the day–night cycle for northern aspen trees than for southern trees. Thus, when exposed to 20-h days, the CO peak already falls into the dark for the northern trees, FT is no longer activated, and growth is not supported anymore. Conversely, the CO peak for the southern trees is still in the light, FT remains active, and dormancy continues to be repressed [after data from Böhlenius et al. 2006].

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