Developmental Plasticity in Plants

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As sessile organisms, plants are unable to seek out environmental conditions optimal for their growth and development but instead must complete their life cycles in the environment in which they are growing. However, plants are remarkably plastic, such that a single genotype is able to give rise to a wide range of phenotypes. Developmental plasticity has profound implications for plant evolution and ecology and can make important contributions to improving yield stability in agriculture. In this review, we discuss the genetic control mechanisms that underlie plasticity and their implications for plant evolution, using the control of flowering time in Arabidopsis as an example. Furthermore, we consider how rapid improvements in quantitative genetic resources provide opportunities to analyze the molecular mechanisms that regulate developmental plasticity more directly and completely.

Higher plants are complex multicellular organisms that are autotrophic. They depend on the collection of relatively simple molecules from the environment—water, carbon dioxide, and mineral nutrients—and use energy from sunlight to power the conversion of these simple molecules into the complex macromolecules from which they are built. To collect sufficient quantities of these basic resources, plants must have very large surface areas in contact with both the above- and below-ground environments. As a result, they are sessile, unable to seek out environmental conditions optimal for their successful growth and development. Instead, wherever a seed germinates, the emerging plant must complete its life cycle at that location, whatever the environment. Furthermore, the environment is likely to change dynamically during the lifetime of the plant. Given these constraints, it is not surprising that plant development is remarkably plastic, such that a single plant genotype is able to give rise to a wide range of phenotypes depending on the prevailing environmental conditions (Bradshaw 1965; Sultan 2000).

This plasticity in development is possible because of the continuous nature of plant development (Palmer et al. 2012). During embryogenesis, the basic body axes of the plant are established, including the main apical–basal axis with the shoot apical meristem (SAM) at one end and the root apical meristem at the other. After germination, this axis is elaborated by the development of root and shoot systems, built through the action of the primary embryonic meristems, as well as the establishment of new secondary meristems, giving rise to lateral roots and shoots and higher-order branches. Each root and shoot can branch to different degrees, elongate to different degrees, and, particularly the shoot, produce a range of different specialized structures of varying types, to various degrees. For example, shoots can produce leaves adapted to shade or bright sunlight and, at some point, switch from making leaves to making floral organs. By integrating environmental information into the regulation of these growth and developmental processes, plant form can be modulated according to the environment in which the plant is growing. In this way, the final morphology of the plant depends on the environment, defining it as plastic.

The extraordinary phenotypic plasticity of plants has received attention in two interlinked contexts. First, developmental plasticity has profound implications for plant evolution. If a single genotype can result in multiple phenotypes, this weakens the link between selection at the level of the phenotype and changes in allele frequency at the level of the genotype. This suggests an explanation for why there are so many fewer plant species than animal species on the planet. However, it also raises interesting questions about the constraints on plasticity and the limits to the selective advantages that it brings, begging the question, under what circumstances is plasticity selected against? Second, it is becoming increasingly possible to investigate the genetic control mechanisms underlying plasticity. A priority in this context is the question of the relationship between plasticity in a trait and the mean value of that trait. As mentioned above, these two areas are interlinked, because understanding the genetic architecture of plasticity will contribute to understanding its evolution. In this review, we discuss progress toward this goal and the opportunities presented by spectacular improvements in quantitative genetic resources, enabled by cheap and easy DNA sequencing.

THE EVOLUTIONARY SIGNIFICANCE OF PLASTICITY

Nonadaptive Plasticity

As described above, the ability of plants to modulate their development in response to the environment is usually considered to be adaptive, contributing positively to reproductive success. However, before considering the implications for adaptive developmental plasticity, it is
important to point out that phenotypic variation between plants grown in different environments may not affect fitness at all, or it may have a negative impact on fitness. Such variation may, for example, reflect variation in a trait as a result of mutation or selection of other traits that are linked (Alpert and Simms 2002), or it may reflect poorly buffered development with negative effects on individual fitness as the inevitable result of stress or resource limitation (Sultan 2000; Steinger et al. 2003; Weiner 2004; Van Kleunen and Fischer 2005).

Although adaptive plasticity is generally considered to be of most evolutionary and ecological interest, all these types of plasticity influence a plant’s chances of survival and reproduction and will thus affect processes such as colonization. Nonadaptive plasticity reduces the likelihood of persistence in a new environment that is stressful and thus increases the strength of selection. Grether (2005) suggests that when a species encounters novel environmental conditions that trigger phenotypic changes with reduced fitness, selection will favor genetic changes that counteract these changes to restore the phenotype to its ancestral state. When this form of adaptive evolution, termed “genetic compensation,” has occurred, genetic differences among populations have evolved so that the mean trait values of populations in different environments may appear to be more similar when measured in their native habitat than when grown under common environmental conditions. Thus, genetic compensation reduces phenotypic variation but promotes genetic divergence (Ghalambor et al. 2007; Fitzpatrick 2012).

Alternatively, stressful environments may increase phenotypic variation via the expression of cryptic genetic variation. Under typical environmental conditions, the observed phenotypic diversity is only a fraction of the diversity that is actually available, but under stressful conditions, cryptic genetic variation may be revealed. By chance, this stress-induced, nonadaptive plasticity could produce a phenotype that supports survival and reproduction, thereby allowing further adaptation to the new environmental conditions (Rutherford 2000; Gibson and Dworkin 2004; Ghalambor et al. 2007).

Adaptive Plasticity

The ability to modulate development according to the environment can clearly be adaptive by maintaining reproductive success in a changing environment. This is true both for a single individual growing in fluctuating environment and for populations growing in heterogeneous environments. These properties mean that phenotypic plasticity has important roles in the evolutionary diversification of plants (Via and Lande 1985; Schlüter 1986; Scheiner 1993) and in the distribution and persistence of plant species, for example, in response to global climate change (Reusch and Wood 2007; Chevin et al. 2010; Nicotra et al. 2010; Palmer et al. 2012). Populations with high plasticity in selectively important traits can rapidly adapt phenotypically to changes in environmental conditions without the need for genetic adaptation (Valladares et al. 2007; Crispo 2008). Similarly, successful invasive species often show higher plasticity than the native species with which they are competing, allowing them to adjust rapidly to their new geographical area (Sultan 2003; Lande 2009).

The ability of a single genotype to compete successfully in diverse environments can result in reduced genetic variation at the population level, which can inhibit trait evolution (Schlichting 1986; Sultan 2000). If developmental plasticity results in adaptive phenotypes that match the optimum for relevant traits in a range of environments, there is no opportunity for directional selection and thus no genetic differentiation among populations across the environmental range (Ghalambor et al. 2007). Alternatively, if plasticity allows a population to persist in a changed or new environment, but does not deliver the optimal phenotype, the phenotype may become genetically fixed by genetic assimilation (Pigliucci et al. 2006) and improved further by natural selection. This phenomenon, also known as the “Baldwin effect” (Simpson 1953), can accelerate phenotypic adaptation to new growth conditions by allowing an entire population to persist by adaptive plasticity, with successful phenotypes honed by natural selection (Crispo 2008; Lande 2009).

Costs and Limits

Given its obvious benefits, the observation that there is substantial genetic variation in plasticity among natural plant populations suggests that it must also have costs (Pigliucci and Schlüter 1995; Linhart and Grant 1996; Van Kleunen and Fischer 2005). According to DeWitt et al. (1998), costs are evident when, under similar environmental conditions, the fitness of a plastic organism is lower compared with the fitness of a nonplastic organism with the same mean trait value. Several costs for plasticity have been identified (DeWitt et al. 1998; Auld et al. 2010). The most straightforward are those associated with sensing the environmental conditions and transducing these signals into developmental responses. More difficult to assess are costs that could result from the additional genetic complexity that may be required for plasticity. This could result in genetic costs from linkage between loci that promote plasticity and loci conferring low fitness. There could also be pleiotropic effects of loci affecting plasticity on other traits and epistatic interactions between plasticity-related loci and other loci. All of these could reduce the fitness of plastic organisms in comparison with those that have more fixed developmental programs.

In addition to these costs, plasticity clearly has limits that can also reduce fitness. This can be the case where there are developmental range limits such that the developmental outcome of a plastic response cannot produce trait values as near to the optimum as a more constrained, fixed developmental program. For example, in extreme environments, the optimum for a trait may be very far from the population mean and not attainable by an organism that can adopt a range of trait values around the mean.
Because of this, phenotypic plasticity is often associated with lifestyle. Generalists living in moderate environments, with strong competition, often display a high degree of plasticity, whereas specialists that are restricted to harsher environmental conditions display a low degree of plasticity (Sultan 2003).

As well as developmental range limits, plasticity can be limited by information reliability. This occurs when the cues used to assess the environment are insufficiently reliable or specific to provide an accurate assessment or when the environmental conditions are so variable that the information available at any one time gives little indication of future conditions. This latter case is closely related to so-called lag-time limits. Here, the amount of time between sensing and responding to environmental cues results in a temporary mismatch between environment and phenotype (Van Kleunen and Fischer 2005; Valladares et al. 2007). This is particularly problematic if the environment changes at a rate faster than the lag time, such that these mismatches persist. Furthermore, unlike biochemical or physiological plasticity, where changes are generally reversible, developmental modifications are often irreversible (Pigliucci et al. 2006) and therefore represent a more significant risk. If the environment reverts, it may not be possible to adopt the previous developmental state, or it may preclude the adoption of further alternative states, should the environment change again. Thus, the potential for phenotypic plasticity is limited by the history of plant responses of the plant (Weinig and Delph 2001).

When Is Adaptive Plasticity a Selective Advantage?

Because of these limits and costs to plasticity, it is clear that selection will act on plasticity as well as on mean trait value, and thus the plasticity of a trait should itself be considered a quantitative trait (Bradshaw 1965; Alpert and Simms 2002; Nicotra et al. 2010). Because of selection, the degree or pattern of plasticity varies among species, populations, and genotypes. Furthermore, plasticity can vary within a single genotype, depending for example on the environmental conditions or developmental stage of the individual (Schlichting 1986; Sultan 2000; Alpert and Simms 2002), so that plasticity is itself a plastic trait (Sultan 2000; Huber et al. 2012).

Natural selection will favor locally adapted plants when the environment is constant, whereas phenotypic plasticity may be more beneficial in spatially or temporally heterogeneous environments (Givnish 2002). Whether natural selection actually favors plasticity depends on the scale and amplitude of spatial heterogeneity and the duration of the environmental variation. Plasticity may be most advantageous if the environment varies at a scale between the size of the responding part of the plant (e.g., the leaf in the case of sun vs. shade leaves or even the whole plant for root vs. shoot balance) and the dispersal range of the offspring (Alpert and Simms 2002). Recently, Baythavong (2011) has been able to support this hypothesis with empirical data by characterizing the plant communities associated with serpentine and nonserpentine soils. Serpentine soils, derived from ultramafic rocks, exhibit low levels of essential macronutrients, low calcium–magnesium ratios, and toxic concentrations of several elements (Branco and Ree 2010). Sites containing patches of serpentine soil represent a significantly finer grained habitat than sites without serpentine patches. Plasticity in plant populations growing on a site containing patches of serpentine soil was found to be higher than in populations growing on sites with uniformly nonserpentine soils (Baythavong 2011).

Plasticity may also be most beneficial if the environmental variation matches the developmental range of the plastic genotype and if the amount of time needed to express the plastic response is shorter than the duration of the environmental state (Padilla and Adolph 1996; Alpert and Simms 2002). Because the effectiveness of the plastic response greatly depends on its timing, these environmental variations are even more likely to select for plasticity if they are predictable and thus preceded by environmental cues that accurately indicate future conditions (Alpert and Simms 2002).

Another important consideration is the complexity of the environment. A heterogeneous environment often varies with respect to multiple factors, such as light, temperature, humidity, and nutrient status, and the spatial and temporal scales may differ for each of these factors (Valladares et al. 2007). All of these different environmental cues must be detected and integrated to maximize individual plant fitness. This involves prioritization of phenotypic variation that addresses the most limiting environmental factor and resolution of conflicting inputs into the same output.

Despite the long history of both theoretical and empirical research on plasticity, the mechanisms that regulate developmental plasticity are still poorly understood. Understanding how environmental cues are integrated with endogenous developmental programs to produce a functional phenotype will provide more insight into the ecological and evolutionary implications of phenotypic plasticity and its evolution.

THE MOLECULAR MECHANISMS OF DEVELOPMENTAL PLASTICITY

As described above, in 1965 Bradshaw noted that there is considerable variation in plasticity, both within species and among closely related species, and proposed that the plasticity of a trait is under genetic control. The discussion over the nature of this genetic control has been somewhat theoretical and based around the idea that traits are determined by distinct sets of structural genes that contribute directly to the trait and regulatory genes that control the activity of the structural genes. The two main models proposed to account for the genetic control of plasticity involve different roles for structural versus regulatory genes (Schlichting and Pigliucci 1993; Via 1993).

In the first model, allelic sensitivity, structural genes are constitutively expressed, but the level of expression is...
important and the molecular analysis of trait plasticity in an ecological context. In this model, direct selection on the trait value leads to a strongly correlated response of trait plasticity, and plasticity is achieved as a result of the combined environmental sensitivity of many structural genes, with no specific “plasticity genes” (Via 1993). In contrast, in the second model (regulatory control), the expression of structural genes is mediated by regulatory genes, for example, genes encoding transcription factors that control the level of expression of multiple structural genes. If the activity of these genes is under environmental control, such genes could act as “plasticity genes,” exerting environmentally dependent control over the expression of structural genes, enabling the amount and pattern of the plastic response to evolve independently of the mean trait value (Schlichting 1986; Schlichting and Pigliucci 1993, 1995). Under this model, different regulatory genes can be expressed in different environments, enabling independent responses to different environmental inputs. This could contribute to the evolution of plasticity, because genes with environment-specific expression may experience relaxed selection and thus a higher relative mutation rate compared with constitutively expressed genes (Van Dyken and Wade 2010; Wunder 2012). This type of regulatory architecture could also support the evolution of control mechanisms that anticipate environmental change, for example, through circadian gating (Schlichting and Pigliucci 1995; Givnish 2002).

Allelic sensitivity and regulatory control are obviously not mutually exclusive. Furthermore, they start with the notion that there are separable structural and regulatory genes, which is clearly not the case. Perhaps the most useful aspect of these models is in their predictions about “plasticity genes.” In the allelic sensitivity model, there are no “plasticity genes,” and variation in mean trait values and plasticity variation go hand in hand. In the regulatory control model, there are distinct plasticity genes, and changes in which can affect plasticity without affecting mean trait values. Exploration of these issues requires the molecular analysis of trait plasticity in an ecological context. This type of analysis is underway for several important and/or tractable traits, the most advanced of which is the environmental control of flowering, principally studied in Arabidopsis. The genetic architecture of flowering time plasticity and its implications for its adaptive significance are discussed below.

**PLASTICITY IN ARABIDOPSIS FLOWERING TIME**

The optimal time to flower for an annual plant such as Arabidopsis is a complex calculation. For Arabidopsis, flowering is a one-off irreversible decision that leads to the production of seed and the end of the life cycle. During the vegetative phase, shoot and root proliferation allows nutrient and photosynthate accumulation. The transition to flowering curtails this phase of resource accumulation, capping fecundity. The optimal time to spend in the vegetative phase to maximize fitness depends on the environment. This includes relatively predictable aspects of the environment caused by the season, such as day length and average temperature, and less predictable aspects, such as nutrient availability, salinity, or unseasonable temperature fluctuations. Monitoring reliable seasonal indicators such as day length provides information about whether the conditions for growth are likely to be favorable and get better or worse in the future. Under circumstances where a prolonged period of favorable conditions is anticipated, remaining vegetative, thereby accumulating resources, is likely to improve fitness. In contrast, indicators of less predictable, unfavorable conditions may promote early flowering, ensuring that at least some seeds are produced, or they may slow growth and delay flowering, freeing resources for protection from stressful conditions.

Initial efforts to understand flowering time focused on the mechanisms that integrate seasonal cues into the flowering program. Current understanding of these mechanisms has been summarized in excellent reviews (for example, see Mouradov et al. 2002; Putterill et al. 2004; Ausin et al. 2005; Corbesier and Coupland 2006; Michaels 2009; Amasino 2010; Jariilo and Pineiro 2011; Srikanth and Schmid 2011). Regulatory frameworks established through the study of the seasonal regulation of flowering are now being elaborated via more recent analyses of how less predictable environmental variables affect the system.

**Seasonal Regulation of Flowering**

Molecular genetic analysis of flowering time began with classical forward genetic approaches to identify genes involved in the major seasonal regulatory inputs into flowering, namely, day length and winter cold. These studies identified a core set of regulators that act as integrators for multiple inputs. The main floral integrators are SOC1 (SUPPRESSOR OF OVEREXPRESSION OF CONSTANS 1), FT (FLOWERING LOCUS T), and FD, and they in turn regulate the expression of floral meristem identity genes including AP1 (APETALA1) and LFY (LEAFY) (Kobayashi et al. 1999; Abe et al. 2005; Wigge et al. 2005; Yoo et al. 2005) that drive the floral program. Arabidopsis is a facultative long-day plant; its flowering is accelerated in long days (16/8 h light/dark) and delayed in short-day conditions (8/16 h light/dark). Day length is detected by a so-called external coincidence mechanism involving the interaction of the circadian clock and light detection. In leaves, the clock protein GI (GIGANTEA) promotes the expression of CO (CONSTANS) such that it peaks during the night in short-day conditions, but this occurs at dusk when plants are grown under long days (Suarez-Lopez et al. 2001). In darkness, the CO protein is rapidly degraded and thus never accumulates under short days. In long days, at the end of the day, light and CO coincide. The light is detected by the photoreceptors cry1 (cryptochrome 1) and cry2 and phyA (phytochrome A), resulting in stabilization of...
CO (Valverde et al. 2004). CO induces the expression of FT (Yanovsky and Kay 2002). FT encodes a small, soluble protein that moves through the phloem toward the SAM, where it interacts with the bZIP transcription factor FD to activate the floral meristem identity genes (Abe et al. 2005; Wigge et al. 2005).

In the meristem, independent of CO, expression of the floral integrators FT, SOC1, and FD is regulated by FLC (FLOWERING LOCUS C), a member of the MADS box transcription factor family. FLC acts as a suppressor of flowering, and its expression is down-regulated during prolonged exposure to cold (vernalization) (Michaels and Amasino 1999). Arabidopsis accessions that live at more northerly latitudes have high levels of FLC expression driven by the FRI (FRIGIDA) gene (Clarke and Dean 1994). As a result, these accessions require a period of prolonged cold before they flower, defining the winter annual life history. In the short growing season available to these plants, natural selection apparently favors germination in the autumn, overwintering as a slow-growing plant, and rapid establishment of faster growth as soon as the spring arrives, and flowering when the photoperiod reaches the required length. Down-regulation of FLC by prolonged cold is maintained after vernalization by chromatin modification at FLC by a Polycomb repression complex (Doyle and Amasino 2009; Jiang et al. 2009). Interestingly, this repression is mitotically stable and functions as a memory of prior cold exposure, permitting rapid flowering at the appropriate photoperiod. FLC is also the main target of the so-called autonomous pathway that allows flowering eventually even in the absence of environmental conditions that accelerate flowering. Different autonomous pathway genes repress FLC expression using different mechanisms. FCA and FY control FLC gene expression by regulating polyadenylation (Manzano et al. 2009). FLD (FLOWERING LOCUS D) and FVE are involved in histone deacetylation and thus may repress FLC expression by changing chromatin structure (He et al. 2003; Ausin et al. 2004; Kim et al. 2004).

Recently, an additional endogenous pathway for flowering has been described involving two microRNAs, miR156 and miR172. In Arabidopsis, the expression of these miRNAs is independent of day length but is developmentally controlled. As development proceeds and the plant matures, miR156 levels decline, releasing the transcriptional repression of genes of the SPL (SQUAMOSA PROMOTER BINDING-LIKE) family (Wang et al. 2009). SPL transcription factors are positive regulators of floral integrators and may increase levels of miR172 (Wu et al. 2009; Yamaguchi et al. 2009). In turn, miR172 promotes flowering by down-regulating repressors of FT (Aukerman and Sakai 2003; Mathieu et al. 2009).

The plant growth regulator GA (gibberellin) also has an important role in the activation of flowering, which is especially evident under short-day conditions when the photoperiodic pathway is not activated. Under short days, GAs activate the expression of SOC1 and the floral meristem identity gene LFY (Blazquez et al. 1998; Moon et al. 2003). The activation of LFY by GA is possibly also mediated through miR159, which directs cleavage of mRNA encoding GAMYB-related transcription factors that regulate the activity of LFY (Gocal et al. 2001; Achard et al. 2004). Under long days, GA is required in the vascular tissue, where it increases the expression of FT and its closest homolog TSW (TWIN SISTER OF FT). In the meristem, GA activates the transcription of SPL genes but is not required to activate SOC1 as reported under short-day conditions (Hisamatsu and King 2008; Porri et al. 2012).

**Regulation of Flowering Time in Response to Nutrient Availability**

The inorganic nutrient nitrate (NO₃⁻) is a major source of nitrogen for plants. Low NO₃⁻ does not statistically significantly reduce the number of days to flowering (Kant et al. 2011; Marin et al. 2011), but when flowering time is scored on the basis of leaf number, plants grown on low NO₃⁻ flower earlier, suggesting that differences in flowering time are masked by differences in growth rate (Marin et al. 2011). Consistent with this view, in Arabidopsis, several studies have shown that the number of days to flowering is not significantly affected by mineral nutrition (Zhang and Lechowicz 1994; Pigliucci and Schlidtung 1995). However, when plants germinated and grown on high nutrient levels for several weeks were transferred to low nutrient levels, flowering was promoted. This effect of nutrient deficiency was more pronounced in short days than in long days (Kolar and Senkova 2008).

In an alternative approach to reduce growth differences between treatments, Marin et al. (2011) studied the effect of varying concentrations of NO₃⁻ on flowering while providing constant N nutrition through glutamine supply. Decreased NO₃⁻ resulted in earlier flowering but only in short-day conditions, consistent with previous results (Kolar and Senkova 2008). Mutants and transgenic lines affecting the autonomous, GA, photoperiod, or vernalization pathways were still able to respond to nitrate. Strikingly, the triple mutant fca-1 co-2 ga1-3 did not flower under either short- or long-day conditions but did flower when grown under low-NO₃⁻ conditions. Other abiotic stresses such as high light or high temperature could not induce flowering in this triple mutant. Furthermore, low NO₃⁻ also accelerated flowering in floral integrator mutants such as ft and fdl and floral identity mutants such as lfy. These findings suggest that the response to nitrate is not part of a general stress response but is regulated by a separate pathway, parallel to the photoperiod, GA, and autonomous pathways, that enters downstream from known floral integrators (Fig. 1). Interestingly, transgenic lines that constitutively overexpressed FLC did not respond to nitrate, but plants grown on low NO₃⁻ have reduced levels of FLC mRNA (Kant et al. 2011; Marin et al. 2011). Possibly, low nitrate represses FLC but only when it is expressed from its own promoter (Fig. 1).
Figure 1. Schematic model of the pathways that control flowering time in Arabidopsis. Genetic studies have identified regulatory pathways (in bold) that integrate the seasonal environmental cues of day length and winter into the endogenous flowering program. Less predictable environmental cues, such as nutrient availability, salinity, or unseasonal temperature fluctuation, also affect flowering time. Most of these responses are regulated through the well-described pathways, modifying the activity of genes involved in the photoperiodic, autonomous, and GA floral pathways. Floral integrators, such as FT, SOC1, and FD, act as hubs that integrate multiple inputs to regulate flowering by modulating the expression of floral meristem identity genes, such as AP1 and LFY. Some environmental cues, however, such as low nitrate ($\text{NO}_3^-$), may regulate flowering time independently of these pathways.

### Regulation of Flowering Time in Response to Ambient Temperature

In Arabidopsis, exposure to high temperature results in extension of the plant axes and early flowering (Gray et al. 1998). These phenotypes are similar to those produced in response to shading (Franklin 2008). Interestingly, phyB, which has a prominent role in the perception of shading, is also involved in temperature-sensitive flowering. When grown under a constant temperature of 22°C, the phyB mutant displays an early flowering phenotype compared with wild type. Accordingly, FT expression is greatly enhanced. However, the early flowering phenotype of the phyB mutant is abolished when grown at the lower temperature of 16°C, and FT transcript levels are reduced to wild-type levels at this temperature. Additional loss of PHYE next to PHYB accelerates flowering at 16°C and triggers an elevation in FT transcription. These data support a role for FT in the ambient-temperature-dependent induction of flowering mediated through the bHLH transcriptional regulator PIF4 (PHYTOCHROME INTERACTING FACTOR 4) (Koini et al. 2009). Mutants with lesions in PIF4 showed a loss of thermal induction of flowering and abolished FT expression at high ambient temperature in short days, whereas overexpression of PIF4 caused early flowering. Kumar et al. (2012) showed that PIF4 directly binds to the FT promoter near the transcriptional start site and that temperature-dependent regulation of FT by PIF4 proteins is controlled at the level of chromatin accessibility. The protein activity of PIF4 is controlled by DELLA proteins; these are repressors of GA-responsive growth. DELLA proteins are able to bind to the DNA-recognition domain of PIF4 (de Lucas et al. 2008), preventing PIF4 from binding to the promoter of FT. The phytohormone GA induces degradation of DELLA proteins via the ubiquitin/proteasome pathway. Accordingly, GA application increased the...
expression of FT (Hisamatsu and King 2008). These findings suggest that PIF4 acts as an integrator between the ambient-temperature-sensing pathway and the GA floral induction pathway (Kumar et al. 2012) (Fig. 1).

Regulation of Flowering Time in Response to Salinity

In contrast to the previously discussed environmental cues, salt stress delays flowering in Arabidopsis. Flowering in response to salt is regulated in a dose-dependent manner. Moderate salt stress promotes vegetative growth, but delays flowering, whereas high concentrations of salt repress both vegetative and reproductive growth. Salt-treated wild-type plants contain reduced levels of bioactive GAs and substantially reduced expression of LFY (Achard et al. 2006). Exogenous GA application can partially rescue both the late-flowering phenotype and LFY expression levels of salt-treated plants, suggesting that the regulation of flowering time by salt is partly mediated by the GA pathway. Mutants with lesions in FLC or FCA, however, are still responsive to salinity, suggesting that the autonomous pathway is not involved in salt-induced repression of floral transition (Fig. 1).

CO and FT mRNA levels are reduced by salt stress in a dose-dependent manner and the co-2 mutant is not responsive to moderate salt stress (Fig. 1) (Li et al. 2007). An additional mechanism to regulate FT expression in response to salinity involves the transcription factor NTM1-LIKE8 (NTL8), a member of the NAM, ATAF1/2, CUC2 (NAC) transcription factor family. The expression of NTL8 is induced by high salt. The NTL8 protein is primarily localized at the plasma membrane, but if released, NTL8 translocates to the nucleus to regulate the expression of target genes. Overexpressing nuclear NTL8 delays flowering and is associated with the reduced expression of FT. In the ntl8-1 knockout mutant, however, the salt-induced repression of FT is diminished (Kim et al. 2007). In contrast to FT, its gene family member BFT (BROTHER OF FT AND TFL1) is induced under high salinity, whereas other environmental factors, such as cold and drought, only slightly affect its expression. The salt induction of BFT expression is largely mediated by the stress-related phytohormone ABA (abscisic acid). Overexpression of BFT delayed flowering under both normal growth conditions and high salinity, whereas the bft-2 mutant was insensitive to high salinity. BFT regulates the meristem identity genes, such as LFY and AP1, but only AP1 appears to be involved in BFT regulation of flowering under high salinity (Ryu et al. 2011).

Central Regulation of Flowering Time and Flowering Time Plasticity

As is clear from the evidence presented above, different environmental cues accelerate or delay the transition to flowering. The genes FLC and FT, which are the central players in the well-described seasonal flowering time pathways, are also key components in the regulation of flowering in response to less predictable environmental cues. Indeed, most of these environmentally induced responses are regulated through the photoperiodic, autonomous, or GA floral pathways (Fig. 1). Nevertheless, some environmental cues may regulate flowering time independently of these pathways, bypassing known floral integrators and identify genes. Another recurring feature is that these environmental inputs frequently influence flowering by multiple signaling routes, resulting in both redundancy and complexity in the mechanisms that control flowering.

These features have interesting hybrid properties between the allelic sensitivity and regulatory control models. On the one hand, there are clearly “plasticity genes.” For example, FRI functions to set the level of FLC, thereby determining the extent to which a plant requires vernalization to flower. If vernalized, FRI and fri plants flower at the same time, but without vernalization, their flowering times are dramatically different. This defines FRI as a “plasticity gene.” Furthermore, there is clearly a regulatory hierarchy allowing multiple different environmental inputs to be integrated by an overlapping set of common targets to regulate a single output, characteristic of the regulatory control model. However, the redundancy in the system, with one environmental signal targeting multiple genes in the regulatory network, is reminiscent of the allelic sensitivity idea, and some environmental inputs, such as nitrate, have substantial effects on flowering time through as yet unknown mechanisms that apparently feed in downstream from known regulators, closer to the “structural genes.”

Natural Variation in Flowering Time Plasticity

All of these studies, although building a detailed network for the control of the floral transition, were not really intended to specifically address the question of plasticity. The mutant screens that were the mainstay of the identification of the genes described involved selection for mean trait value in a single environment. Many different environments have been used in different screens, but nonetheless, this approach does not screen directly for plasticity per se. This requires individuals with identical genotypes to be grown in two or more environments. Therefore, the use of classical mutant screens to identify candidate genes involved in phenotypic plasticity is challenging. In a typical mutant screen, single M2 plants are selected based on phenotype, but to identify plasticity mutants, M3 families would have to be collected and scored in two relevant environmental conditions. Moreover, plasticity is difficult to study using null alleles, which are the most common type recovered from mutant screens. Null alleles simultaneously remove the function of the gene and any ability to modulate it, making it difficult to separate mean trait value from plasticity in the trait. These problems can be overcome through the use of natural quantitative variation mapped in recombinant inbred lines using traditional linkage mapping or genome-wide association (GWA) mapping to identify quantitative trait loci (QTL). Both molecular tools are based on the statistical association
of phenotypic data with genotypic data (Ikram and Char- don 2010).

Traditional linkage mapping in recombinant inbred lines (RILs) is well suited to the identification of genomic regions governing plasticity. RILs are almost completely homozygous, allowing replicates to be grown and analyzed across several environments. Mapping resolution and genetic diversity in the mapping population is determined by the number of founder lines and the generations of intermating and selfing used to make the RILs (Bergelson and Roux 2010). For example, Kover et al. (2009b) developed a set of RILs derived from a multiparent advanced generation intercross MAGIC of 19 accesses of Arabidopsis. The mapping resolution for the MAGIC lines can be as high as 300 kb, so these lines can be used for fine-mapping QTL including phenotypic plasticity.

In natural accessions, recombination events have accumulated over many generations, so the mapping resolution in GWA studies can be higher than in traditional mapping. Self-fertilizing species such as Arabidopsis are well suited for GWA studies. Once the accessions have been densely genotyped, they can be used to analyze different traits among different environments (Nordborg and Weigel 2008). These approaches have now been quite extensively used to map variation in flowering time, including studies carried out on the same lines in different conditions (see, e.g., Schwartz et al. 2009; Brachi et al. 2010; Li et al. 2010; Strange et al. 2011). Most of the subsequent molecular analysis has focused on known loci, already identified in classical mutant screens. Although this has revealed important data on the molecular basis for natural variation in flowering time, it has been less relevant from the point of view of analysis of phenotypic plasticity. This may change as the focus shifts to previously uncharacterised loci or studies of plastic responses to nonseasonal inputs. Similarly, analysis of traits other than flowering time that can be modulated more continuously is now under way (see, e.g., Filiault and Maloof 2012). For these inputs and traits, there is likely to be less straightforward selection on plasticity, so the genetic architecture of plasticity in these traits may be more revealing.

A further use of these lines that can, for example, help to distinguish between allelic sensitivity and regulatory control models is through selection experiments (Scheiner 2002). These are challenging in higher plants owing to their relatively long generation times. Nevertheless, several artificial selection studies have been conducted with Nicotiana rustica and Arabidopsis to evaluate the independence between mean trait value and trait plasticity. These have variously targeted the mean trait values, the plasticity of the trait, or a combination of the two for selection (Brumpton et al. 1977; Jinks et al. 1977; Callahan and Pigliucci 2005; Springate et al. 2011). The findings of these studies suggest that phenotypic plasticity is at least partially under separate genetic control from the trait value, supporting the regulatory control hypothesis. For example, in an artificial selection experiment carried out by Kover et al. (2009a), outbred lines of Arabidopsis were selected for early flowering under either winter-annual or spring-annual simulated conditions. At the end of the experiment, the selected lines always flowered earlier than the control lines, independent of the growth conditions. However, when grown in the conditions in which they were selected, the lines always flowered earlier than the lines selected in the alternative environment. These findings suggest that some of the selected genes affect flowering time under both growing conditions, whereas others are involved specifically in the environmental control of flowering and therefore could be described as “plasticity genes.” Molecular analysis suggests that one of these is FRI (Scarcelli and Kover 2009).

Conclusion and Prospects

Developmental plasticity has shaped the evolutionary history of plants so that understanding the genetic architecture of plasticity is a prerequisite for understanding plant evolution and ecology. Rapid progress has been achieved through advances in plant developmental genetics, with opportunities for the future provided by post-genomic quantitative genetics. These advances are timely because understanding plasticity is also paramount for progress in crop science. Phenotypic plasticity is an important trait for the yield stability of crops. Crops have typically been bred for uniform growth and higher yields under local conditions and often have been selected for reduced genotype-by-environment interactions (Palmer et al. 2012). Thus far, directional selection has not focused on phenotypic plasticity (Nicotra et al. 2010), despite the fact that yield stability and quality in crops such as cereals and rapeseed is often compromised by fluctuating weather conditions (Peltonen-Sainio et al. 2011). Therefore, selective breeding aimed at the plasticity of traits indirectly and directly associated with yield may allow the development of crops with improved yield stability in an increasingly unpredictable environment.

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