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Non-reversion of Impatiens in the absence of meristem commitment

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as in red-flowered Impatiens, meristem commitment a supply of a leaf-derived signal is needed to maintain
is absent, but that purple-flowered plants maintain flowering in the *Impatiens* meristem (Pouteau *et al.*, 1997 flowering in LD conditions because of a more perman-
of genes involved in the flowering process (Ma, 1994). ent supply of signal from their leaves than occurs in red-flowered plants. A working hypothesis is proposed Members of the former group function in the transition
to explain how a signal from the leaves can retain a from vegetative to reproductive growth whilst organ to explain how a signal from the leaves can retain a controlling role during flower development. identity genes are thought to control flower development

floricaula. tem identity gene, floricaula (flo) of Antirrhinum majus

floral state appears, in many plants, to be a prerequisite 1995). In *Impatiens*, however, the transcription of the for floral development. At this point the meristem *Impatiens flo* homologue, *Imp-flo* was strikingly dif for floral development. At this point the meristem *Impatiens flo* homologue, *Imp-flo* was strikingly different, becomes autonomous for flowering and the signals with the *flo* transcript found in the vegetative as well a becomes autonomous for flowering and the signals produced by the leaves are no longer significant to the floral and reverting meristems (Pouteau et al., 1997). development (McDaniel, 1992). In red-flowered plants of This pattern of vegetative transcription has also been Impatiens balsamina, a point of commitment does not observed in Nicotiana and pea (Kelly et al., 1995; Hofer exist and development of the terminal flower proceeds as $et al., 1997)$ and suggests that transcription of flo is

Abstract dictated by the inductive status of the leaves, which is **Purple-flowered plants of** *Impatiens balsamina* **main-**

tained floral development on transfer from inductive

Lyndon, 1986, 1990; Pouteau et al., 1995, 1997). Thus,

short days (SD) to long days (LD), a treatment in which

in the manner outlined in the ABC model (Coen and Key words: Floral commitment, Impatiens, floral reversion, Meyerowitz, 1991). The transcription of the floral merishas been linked with the point of commitment to flower (Bradley *et al.*, 1996) and overexpression of its homo-
logue, *leafy*, in *Arabidopsis thaliana* has been shown to Commitment (or determination) of the meristem to a be sufficient to initiate flowering (Weigel and Nilsson, floral state appears, in many plants, to be a prerequisite 1995). In *Impatiens*, however, the transcription of th

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flowered line of *Impatiens* is described. In common with red-flowered plants, purple-flowered plants were found to
lack meristem commitment but, in contrast to red-
flowered plants, were able to maintain their floral state
flowered on a daily basis for 15 d. on transfer to LD. Plants were thus non-reverting but
uncommitted, with non-reversion being reliant on the Removal): Leaves unfolded during the first 5 SD were left on
presence of leaves that remain induced on transfer of presence of leaves that remain induced on transfer of plants to LD. The results for purple-flowered Impatiens after the first 5 SD were removed on a daily basis for 15 d. support the idea that a leaf-derived signal can directly These treatments are summarized in Fig. 1. The number of control the developmental phases of the floral meristem, leaves removed is given in Table 1.5 SD+LD, 5 SD-LLR and possibly through direct interaction with organ identity 5 SD-SLR mature plants were dissected on days 55–56. possibly through direct interaction with organ identity

Results Materials and methods

Seed was collected from purple-flowered plants originally In our original work on *Impatiens balsamina*, the process derived from the same batch of mixed flower colour seeds of the of flowering and reversion was described in a population
 Impatiens balsamina cv. Dwarf Bush Flowered as the red-

flowered plants from mixed seed that incl period of at least 4 weeks cold storage, seeds were imbibed on showed the most uniform reversion whilst purple-flowered
moist filter paper in Petri dishes for 72 h in 16 h photoperiods plants did not revert on transfer to moist filter paper in Petri dishes for 72 h in 16 h photoperiods at 23 °C. Seeds with a radicle length of 3–4 mm after this time
were selected and sown at a depth of 1 cm in F1 compost
(Levington) with 35 seeds per tray. After sowing, plants were
placed in growth cabinets under control temperature was maintained at 21 °C. Each tray received The aim of this work was to study non-reversion in 200 cm³ of tap water and was rotated (both the tray itself and purple-flowered plants in order to illuminate furt 200 cm^3 of tap water and was rotated (both the tray itself and

around the cabinet) daily.

Plants were initially grown in LD, given by 8 h of light

provided by tungsten bulbs and fluorescent tubes at 263–

280 μ mol m⁻² s⁻¹ followed by 16 h of light from tungsten bulbs

only a only, at 3–4 μ mol m⁻² s⁻¹ measured at the top of the plants on day 0, After around 8–10 LD, when a sufficient number of plants had a first true leaf between 7 and 11 mm in length, as
uniform a population as possible was selected and used for
experimentation. The plants had initiated 8.9 ± 0.79 and
 9.8 ± 0.98 leaves and primordia for re treatments began on this day; $8 h$ of tungsten and fluorescent

using M2 compost (Levington). In one purple-flowered experi-
next of leaf serrations, $\langle 50\%$ petal pigmentation and no
ment of those used to calculate the combined data presented in petiole—see Battey and Lyndon. 1984),

insufficient to cause flowering or act as a marker for
commitment in these species.
In this paper, the floral development of a purple-
ID and no more leaves were removed.
LD and no more leaves were removed.

and 5 SD+SD-LR plants were dissected when the terminal genes.
flower was fully opened (from day 40 onwards).

Flowering in red-flowered and purple-flowered plants Plant growth

purple-flowered plants corresponded approximately to light as above followed by 16 h of darkness.

On day 0, and at all transfer and fixation times, at least five

plants were selected at random and dissected to determine the

number of leaves and primordia present. On day 1 ment of those used to calculate the combined data presented in petiole—see Battey and Lyndon, 1984), petals, petals
Table 2, plants experienced darkness during the day on day 17 with some staminate characteristics, stamens variable. However, purple-flowered plants produced Leaf removal
Leaf removal experiments were carried out to establish the some plants. In some plants petal and stamen development Leaf removal experiments were carried out to establish the plants. In some plants, petal and stamen development
basis for non-reversion of purple-flowered *Impatiens* (see Results was reiterated and stamen initiation was s (i) 5 SD+LD: no leaf removal. to red-flowered plants which remain completely vegetative

Fig. 1. Diagram of experimental treatments carried out in the leaf removal experiment. Treatments are described in detail in the Materials and methods and the Results sections of the text. The diagrams indicate which leaves were removed; for the numbers of leaves removed refer to Table 1.

aTotal removed in these treatments may have included some of the outer bracts.

Table 2. Development of red and purple-flowered plants grown under continuous SD

The starting node of various phases of floral development at the terminal meristem is indicated. On commencing flowering the meristem stops initiating axillary structures and internodes are no longer elongated. Combined data from four experiments. $n=31$ and $n=24$ for purple and red-flowered plants, respectively, unless otherwise stated. $n =$ Number of plants.

aCarpels were not always visible at time of dissection. Approximately 25% purple plants showed some petal-stamen reiteration.

in LD, purple-flowered plants exposed to continuous LD lished over a greater number of nodes than in SD plants. flowered, though much later than SD plants. The terminal Stamens and carpels were not always visible at the time flower of these plants opened around day 70 and the first of dissection on days 69–70. bract was initiated between nodes 30–40 (as opposed to nodes 11–15 in SD). Floral development was character-
ized by a prolonged phase of initiation of leaves with Non-reversion of purple-flowered plants modified venation and/or shape prior to the first bract Purple-flowered plants were given a treatment of 5 SD,

and development of fully pigmented petals was accomp- followed by LD until dissection. In red-flowered plants

Fig. 3. Floral diagrams depicting development at the terminal meristem of purple-flowered *Impatiens* in SD and 5 SD treatments. Diagrams are representations constructed using data collected from the dissection of mature plants. For clarity one whorl of each class of organ identity is shown except in C, D and E where continued petal initiation and indeterminacy is illustrated by many whorls. (Normally there are 5–6 organs per whorl.) The number at the bottom right of each floral diagram indicates the proportion of plants in the treatment which developed in the manner indicated by the diagram. Br, bract; Pe, petal; St, stamen; Ca, carpel. Numbers given after these initials correspond to the range of organ number of this type found in the plants developing in this manner. Intermediate organs are leaves with some petal features. Bract number in 5 SD-LLR and 5 SD+SD-LR plants may be an underestimate as a result of removal of some bracts during leaf removal in these treatments. Where petal initiation was still in progress at the time of dissection, the total given is to the last pigmented petal (young petals at the centre were often green and unexpanded at the time of dissection). Where no carpels are shown, petal or stamen or a repeating pattern of petal-stamen-petal was continuing at the time of dissection.

to leaf production after floral development has begun; petal number in SD plants was 30.3 ± 7.9 and in 5 SD flowering is halted and leaf production resumed after the $+LD$ plants, in which petal initiation was often production of a pseudoflower of organs containing some continuing at the time of dissection, 46.6 ± 11.0). areas of petal pigment (Battey and Lyndon, 1984; Pouteau et al., 1997) (Fig. 2B; compare with Fig. 2A). In contrast
to red-flowered plants, purple-flowered plants did not
revert on transfer to LD, but instead continued flower In red-flowered and purple-flowered plants, a floral revert on transfer to LD, but instead continued flower In red-flowered and purple-flowered plants, a floral stimu-
development (Fig. 2D: compare with Fig. 2C). Similar lus is produced in the leaf in SD (Pouteau *et al.*, 1 development (Fig. 2D; compare with Fig. 2C). Similar results were obtained in a 2 SD+LD and 8 SD+LD unpublished results). SD leaves of red-flowered plants treatment (data not shown), both known reversion treat-
ments in red-flowered plants (Pouteau *et al.*, 1997, and this, because of the absence of meristem commitment, ments in red-flowered plants (Pouteau et al., 1997, and unpublished data). In $\overline{5}$ SD+LD purple-flowered plants causes reversion (Pouteau *et al.*, 1997). The non-reversion internodes were lost and flowers had, at first sight, a of purple-flowered plants suggested that either meristem similar phenotype to SD plants. On closer inspection, the commitment occurred in these plants or the induced state effect of the LD component of the treatment was evident of the leaves could persist in LD. In order to determine through the lack of visible stamens at the centre of most the likely maintenance mechanism, a leaf removal experiplants (Fig. 2D). The number of floral organs in this ment was carried out. If leaves remained in control of

this treatment results in reversion of the terminal meristem treatment was greater than in SD flowering plants (mean

Fig. 2. Development of the terminal meristem of *Impatiens*. Red-flowered plants: (A) terminal flower of SD plant; (B) reversion to leaf production occurring after 5 SD+LD. Purple-flowered plants: (C) terminal flower of SD plant and (D) 5 SD+LD plant. (E) Reversion of the terminal meristem in 5 SD-SLR treatment. (F) Terminal flower of 5 SD-LLR plant and (G) 5 SD+SD-LR plant. (H) Petal-leaf and petal-bract organs as observed in a number of plants from 5 SD+LD, 5SD-LLR treatments.

SD leaves or as a result of a quantitative build-up of true for only half of the 5 SD-LLR plants at the time of stimulus in LD leaves, their removal would be expected dissection. These results suggest that $5 SD + SD$ -LR plants to affect flowering. In contrast, flowering would be produced floral stimulus at a higher level than those unaffected by leaf removal if the meristem was committed. maintained in LD (5 SD-LLR), but undergoing the same All plants were given 5 SD and transferred to LD as leaf removal treatments, and also suggest that the level above. At the time of transfer, in some of the plants one of floral stimulus produced by leaves unfolded in SD is cotyledon and the leaves that had unfolded during the higher in SD than in LD. 5 SD period (SD leaves) were removed (5 SD-SLR). In other plants, SD leaves were left intact on the plant but
those leaves unfolding in LD (LD leaves) were removed **Discussion** for a period of 15 d after transfer (5 SD-LLR) (plants When leaves that had unfolded in SD were removed from were inspected daily and any unfolding leaves were purple-flowered plants on transfer into LD after 5 SD removed). No leaves were removed from a further set of (5 SD-SLR treatment), reversion occurred at the terminal plants (5 SD+LD controls) (Fig. 1; Table 1). meristem. This is evidence that meristem commitment

When SD leaves were removed (5 SD-SLR treatment), 7 lack of reversion in 5 $SD + LD$ plants demonstrates that out of 9 plants reverted (Fig. 3D). At the time of dissec- there is a commitment of the whole plant to the floral tion (when SD controls had developed fully-opened ter- state. This result supports the idea, implied by redminal flowers) these 7 plants had little or no pigmentation flowered plants, that meristem commitment is not a of any organ, no petals except small, green unexpanded prerequisite for flowering (Battey and Lyndon, 1990). petals in 2 plants and in all plants internodes were lost Expression of the meristem identity gene floricaula above then reinstated creating a pseudoflower of between 5 and a threshold level correlates with meristem commitment in 18 intermediate (leaves with petal features) organs Antirrhinum (Bradley et al., 1996). However, in red- (Fig. 2E). Finally, from the centre of the pseudoflower flowered plants of Impatiens the flo homologue is leafy organs, often with hairy abaxial surfaces and ser-
expressed in vegetative, floral and 5 $SD+3$ LD meristems rated edges were emerging. The other 2 plants in this (Pouteau *et al.*, 1997). Similarly in meristems of purpletreatment flowered, but produced approximately twice as flowered plants sampled after 8 LD, 8 SD or 5 SD+ many petals as SD plants and no stamens were visible at 3 LD transcription of *Imp-flo* could be seen in bands at the time of dissection (Fig. 3D). the base of the primordia (data not shown). This expres-

maintain floral development (Fig. 2F) and were develop- be that the absence of meristem commitment in both mentally similar to those in the 5 $SD+LD$ treatment these lines of *Impatiens* is a result of a failure to achieve (described above) though only half of the 5 SD-LLR a threshold level of flo . plants lacked visible stamens (Fig. 3E). Within the In the absence of meristem commitment, the leaves of 5 SD-LLR and 5 SD+LD treatments, a few plants had Impatiens retain a dominant control over the developmosaic organs consisting of a leaf-like tip with a petaloid mental state of the terminal meristem and persistence of base (Fig. 2H). the induced state of leaves is an adequate substitute for

flowered plants were able to maintain flowering on trans- flowered plants the ability of leaves to maintain an fer to LD through the retention of an induced state in induced state in LD is absent, giving rise to reversion the SD leaves. The LD influence did not cause reversion, (Pouteau *et al.*, 1997) the requirement for these leaves but did cause flowering to deviate from that of SD plants (SD leaves) to be intact for flowering of the purplesuggesting that the leaves unfolded in SD produced a flowered line to occur implies that they are capable of lower level of stimulus on transfer of the plants to LD. maintaining an induced state in LD. In the work described To test this, plants were grown in SD and leaves unfolding here, an enhanced induced state was apparent in those during the first 5 SD of the treatment were left intact, leaves unfolding during SD. However, even in continuous but those unfolding subsequently were removed on a LD purple-flowered plants eventually achieved an anomdaily basis for 15 d. Plants remained in SD for the whole alous type of flowering so presumably LD leaves achieved of the experiment ($5 \text{ SD} + \text{SD-LR}$, see Fig. 1 and Table 1). a low level of induction. This implies that the reverting The results are shown in Figs 3B and 2G (compare with 5 SD-SLR plants would be expected eventually to flower 5 SD-LLR, Fig. 3E). 5 SD+SD-LR plants produced under the control of the LD leaves. The results of this smaller numbers of bracts and petals than 5 SD-LLR experiment also imply that the stimulus-producing ability plants, reflecting a more rapid transition through of leaves induced in SD diminished to a lower level on the steps of flower development. In addition, all transfer to LD. Whilst the retention of the induced state

flowering, either by persistence of the inductive ability of 5 SD+SD-LR plants had visible stamens whilst this was

The results of this experiment are summarized in Fig. 3. does not occur in purple-flowered plants. However, the In contrast all the 5 SD-LLR plants were able to sion pattern was identical to red-flowered plants. It may

This leaf removal experiment suggested that purple- apical autonomy in floral development. Whilst in red-

species (Bernier et al., 1981), the nature of determination floral fate in *Impatiens* does not conform to spatial models to this state and the specific role its persistence plays in in which the flower is structured by the rigid confines of flowering is largely unknown. domains and compartments (see Bowman *et al.*, 1989).

In Antirrhinum and Arabidopsis leaf induction appears to lead to the expression of an irreversible floral pathway involving meristem and organ identity genes $(Ma, 1994)$. **Conclusion** On this basis a logical explanation for reversion might be

If this model of flowering in *Impatiens* is correct then the

that, in the absence of meristem commitment a signal

the flowering normally charged in number flow

Impatiens during flowering and reversion suggest that the flowering process is not only stimulus-dependent, but also **Acknowledgements**
driven by the attainment of critical threshold levels of
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timulus: levels which a stimulus; levels which are necessary in order to initiate
and maintain expression of specific elements of the floral
process. Thus our working hypothesis is that there is a
quantitative (grant number AT45/559). We would li lus and the progress of floral morphogenesis. Using this (University of Edinburgh) for their encouragement and hypothesis, for example, the prolonged petal initiation of enthusiasm. 5 SD+LD plants (Fig. 3C) is interpreted as an arrest of the floral process arising from insufficient stimulus for activation of stamen initiation when plants are transferred **References** from SD to LD. The swift transition to stamen initiation Battey NH. 1985. Growth and development at the shoot apex in SD plants (Fig. 3A) reflects the stronger stimulus-
producing ability of leaves in SD than in LD. This PhD thesis, University of Edinburgh. producing ability of leaves in SD than in LD. This PhD thesis, University of Edinburgh.
 Battey NH, Lyndon RF. 1984. Changes in apical growth and hypothesis also explains why the extent of flower develop hypothesis also explains why the extent of flower develop-
ment found in red-flowered plants increases with the
number of SD (Battey and Lyndon, 1984; Pouteau *et al.*,
Battey NH, Lyndon RF, 1986. Apical growth and modifi 1997). A similar situation, also implying involvement of of the development of primordia during re-flowering of the level of floral stimulus is the whorl-by-whorl deter-
mination of flowers of Silane coali-rese in vitro (Donnicon 58, 333–41. mination of flowers of *Silene coeli-rosa in vitro* (Donnison
 et al., 1991; Donnison and Francis 1993, 1994). In
 Impatiens, further evidence that genes influencing organ
 Battey NH, Lyndon RF. 1988. Determination stimulus is the non-appearance of stamens in most

5 SD+LD and some of the 5 SD-LLR purple-flowered

plants at the time of dissection, and the 'superflowering'

described by Simon (1973) which occurred when plants

were ex

of the leaves has been documented for a number of developmental plasticity suggests that determination of

that, in the absence of meristem commitment a signal
derived in stable flowering normally observed in purple-flowered
deviced from induced leaves must persist in order to allow
the continued expression of this pathway. Rem

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