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Stem transcriptome during shade avoidance

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Stem transcriptome reveals mechanisms to reduce the energetic cost of shade-avoidance responses in tomato

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Whilst the most conspicuous response to low red to far-red ratios (R:FR) of shade light perceived by phytochrome is the promotion of stem growth, additional, less obvious effects may be discovered by studying changes in the stem transcriptome. Here we report rapid and reversible stem transcriptome responses to R:FR in tomato (*Solanum lycopersicon*). As expected, low R:FR promoted the expression of growth-related genes, including those involved in the metabolism of cell-wall carbohydrates and in auxin responses. In addition, genes involved in flavonoid synthesis, isoprenoid metabolism and photosynthesis (dark reactions) were overrepresented in clusters showing reduced expression in the stem of low R:FR-treated plants. Consistent with these responses, low R:FR decreased the levels of flavonoids (anthocyanin, quercitin, kaempferol) and selected isoprenoid derivates (chlorophyll, carotenoids) in the stem, and severely reduced the photosynthetic capacity of this organ. However, lignin contents were unaffected. Low R:FR reduced the stem levels of jasmonate, which is a known inducer of flavonoid synthesis. The rate of stem respiration was also reduced in low R:FR-treated plants, indicating that by downsizing the stem photosynthetic apparatus and the levels of photoprotective pigments under low R:FR, tomato plants reduce the energetic cost of shade-avoidance responses.
Due to the optical properties of green leaves, canopy shade is characterized by low red / far-red ratios (R:FR) of the light. One of the most conspicuous responses of plants exposed to low, compared to high R:FR, is the promotion of stem growth (Smith, 1982; Casal and Smith, 1989; Ballaré, 1999; Morelli and Ruberti, 2002; Franklin and Whitelam, 2005; Casal, 2012). A taller stem exposes the leaves to higher and better light strata within the canopy. Conversely, leaf growth may increase, decrease or remain unchanged when the plants receive low R:FR.

In sparse canopies, the horizontal leaves are exposed to the high R:FR of full sunlight while the vertical stems can receive low R:FR, caused by selective reflection of FR by the green foliage. This provides an early warning signal of the presence of neighbours (Ballaré et al., 1987). At later stages of canopy development the leaves become mutually shaded and exposed to low R:FR. The low R:FR reaching only the stem causes a promotion of stem growth that is rapidly reversed to the pre-stimulation values upon return to high R:FR. However, when the low R:FR also reaches the leaves, the promotion is more persistent and only returns to the pre-stimulation values 24 h after the plants are exposed to high R:FR (Casal and Smith, 1988). Therefore, there is a positive correlation between the threat indicated by the shade signals and the persistence of the growth response.

In Arabidopsis seedlings grown under continuous white light for 7 d (at which point they were producing their first true leaves), transfer to low R:FR significantly promotes the expression of cell wall- cell elongation-, cell division- and auxin-associated genes (Devlin et al., 2003). In Arabidopsis plants grown for 19 d under continuous white light before transfer to darkness, a pulse of far-red light given at the end of white light promoted the expression of auxin- and brassinosteroid-responsive genes both in the leaf petioles and blades, despite the contrasting growth responses of these parts of the leaf to far-red light (Kozuka et al., 2010). Conversely, several xyloglucan endotransglucosylase-hydrolase (XTH) genes showed increased expression in the petiole but not in the blade, which correlates with the growth response (Kozuka et al., 2010).

Previous studies have described organ-specific transcriptome responses observed when dark-grown seedlings of Arabidopsis (Ma et al., 2005; López-Juez et al., 2008) or soybean (Li et al., 2011) are exposed to light for the first time. Despite the large stem-growth response to R:FR, our knowledge of stem transcriptome responses to R:FR is very limited. Here we used light-grown tomato (Solanum lycopersicon) seedlings to investigate the stem transcriptome responses to R:FR. By comparing the stem (first internode) and the first pair of true leaves we identified shared and organ-
specific responses. We relate the differential stem and leaf transcriptome responses to organ-selective hormonal and physiological responses.

RESULTS

Shared and specific stem and leaf transcriptome responses to low R:FR

Plants of tomato were grown for 24 d under continuous white light. When the first internode (the stem between the cotyledonary node and the first leaf pair node) was 6.4 ±0.4 mm (mean, SE), half of the plants were exposed to supplementary FR reaching the leaves and the stem from both sides, to lower the R:FR of the horizontally propagating light from 4.6 (white light alone) to 0.05 (white light plus FR). Four days later (see photograph of representative plants in Fig. 1A), the first internode and the first pair of leaves were harvested separately. Total RNA was extracted from these samples, processed and hybridized to tomato Affymetrix microarrays. Microarray data were filtered by presence criteria and subjected to factorial ANOVA with R:FR (high or low) and organ (stem or leaves) as main factors. The genes showing significant effects of treatments ($p < 0.05$, $q < 0.003$) were divided in three groups according to the significance of the effects of R:FR and its interaction with organ (Table S1). A first group included genes showing significant effects of R:FR ($p < 0.05$, $q < 0.04$) and no R:FR by organ interaction. Four clusters of this gene group showed promotion by low R:FR (A1, A2, A5, A9) and five showed reduced expression in response to low R:FR (A3, A4, A6, A7, A8) (Fig. 1B, Table S1). Most genes of this first group also showed significant effects of organ, which were additive to the R:FR effects. A second group included genes showing differential responses to R:FR in the stem and leaves (significant interaction, $p < 0.05$, $q < 0.09$). Two clusters of this gene group showed promotion of expression by low R:FR in the stem (I1, I5), three clusters showed promotion of expression by low R:FR in the leaves (I2, I8, I11), two clusters showed inhibition of expression by low R:FR in the stem (I3, I9) and three clusters showed inhibition of expression by low R:FR in the leaves (I6, I7, I10) (Fig. 1B, Table S1). Few genes showed opposite responses (quantitatively modest) in stem and leaves (cluster I4). Some genes showed reduced expression in response to R:FR in both organs but the effects were significantly higher in the stem (cluster I9). A third, residual group of genes showed no interaction, no significant effects of R:FR and significant effects of organ, and were not further analysed (Table S1).

The total number of genes showing statistically significant responses to low R:FR in the same direction in both organs was larger than the number of genes
responding significantly more in the leaves or the stem (Fig. 1D). However, the latter picture was inverted when the analysis was restricted to those genes with statistically significant responses with a magnitude of at least two fold. In other words, organ-independent changes are more numerous but the largest changes in gene expression are organ specific (Fig. 1D).

**Rapid and reversible stem transcriptome responses to low R:FR**

A second microarray experiment was designed to investigate the early kinetics of stem transcriptome responses to low R:FR. Plants were grown under continuous white light, exposed to supplementary FR for either 1 or 4 h and harvested 0, 1 or 3 h after returning to high R:FR (seven conditions, including the control never exposed to low R:FR). To minimise the impact of circadian rhythms the plants were grown under continuous white light and harvested simultaneously. Microarray data were filtered by presence criteria and subjected to ANOVA (Table S2). The genes showing significant effects of treatments ($p < 0.01$, $q < 0.10$) were used for cluster analysis. A total of 424 genes (clusters S1-S5) showed increased and 410 genes (clusters S6-S9) showed reduced expression in response to low R:FR (Fig. 1C, Table S2). These clusters differed in the kinetics of response, from those that responded rapidly to the transfer from high to low R:FR and showed a rapid reversal of the response after transfer to high R:FR (clusters S2, S8 and S6, the latter with transient overcompensation after the return to high R:FR) to those showing a gradual response to low R:FR and to subsequent high R:FR (clusters S7 and S9) (Fig. 1C). Many genes were significantly affected by both 4 h and by 4 d of low R:FR compared to high R:FR controls. These genes showed a strong correlation in the extent of relative response at both time points (Fig. 1E), supporting the robustness of the reported changes in gene expression.

**Overrepresented GO terms**

We searched for overrepresented GO terms within each cluster (Fig. 1). The overview of both experiments revealed three major areas of transcriptome response in the stem: a) Cell-wall carbohydrate process related genes are overrepresented in Cluster I5, b) Flavonoid process related genes are overrepresented in clusters I3, and S7, and c) Photosynthesis related genes and photosynthetic pigment related genes (e.g. carotenoid metabolism genes within the GO terms “isoprenoid biosynthetic process” and “steroid metabolic process”) are overrepresented in clusters S7, S8 and S9. In subsequent sections we investigate these functions in further detail. Transport-related
genes were overrepresented in cluster I4 but the responses were small in magnitude. The leaves showed photosynthetic- and translation-related genes overrepresented in cluster I6 and flavonol-related genes overrepresented in cluster I8 (Fig. 1). The clusters describing organ-specific responses contain most of the genes with a significant effect of R:FR of at least two-fold and yielded most of the overrepresented GO terms.

**Low R:FR differentially affects stem and leaf growth**

The growth rate of the stem showed a rapid promotion by low R:FR. When the plants were returned back to high R:FR, the rate of stem growth also returned rapidly to the pre-stimulation values (Fig. 2A). However, following a short (1 h) exposure to low R:FR, the stem growth rate showed an overcorrection upon the return to high R:FR leading to a complete compensation of the promotion 3 h later. After 4 d of treatment, the stem showed the characteristic response to low R:FR while the leaf area and dry weight were unaffected (Fig. 2B). In accordance with the latter pattern, cell-wall carbohydrate process genes are overrepresented in cluster I5, showing promotion by low R:FR in the stem and not in the leaves (Fig. 1B). Cell wall-associated genes are also promoted by low R:FR in Arabidopsis (Devlin et al., 2003). Cluster I5 includes several XTH genes and XTH genes had previously been associated to the promotion of hypocotyl and petiole growth responses to low R:FR in Arabidopsis (Kozuka et al., 2010; Sasidharan et al., 2010). Some of these XTH genes showed rapid responses to R:FR (Fig. 2C). Auxin-signalling genes are promoted by low R:FR in Arabidopsis (Devlin et al., 2003; Sessa et al., 2005; Kozuka et al., 2010) and this was also the case in the stem of tomato (Fig. 2C). These results collectively establish a base of coincidence with previous transcriptome reports in Arabidopsis, which contrasts with the tomato-stem-specific patterns reported in subsequent sections.

Some observations require clarification before continuing with the main stream analysis of this work. Previous results had shown that mustard (Casal and Smith, 1988) and tomato (Casal et al., 1995) plants exposed to low R:FR for more than 3 h have elevated rates of stem growth after transfer to high R:FR. This was confirmed under the current conditions, as upon transfer to high R:FR, the stem of plants previously exposed to 4 h low R:FR grew more during the subsequent 20 h (1.7 ±0.2 mm, mean ±SE, P <0.01, n= 42) than the plants previously exposed to only 1 h low R:FR (1.1 ±0.1 mm) or high R:FR controls (1.0 ±0.1 mm). However, this long-term promotion was not observed immediately after the 4 h of low R:FR (Fig. 2A).
Low R:FR differentially affects stem and leaf flavonoids

Figure 3 presents the expression patterns of all the genes of the flavonoid biosynthesis pathway (starting with p-coumaril CoA and chalcone synthase) and the upstream phenylpropanoid pathway that showed significant responses to low R:FR in one or both microarray experiments. The genes involved in steps upstream of p-coumaril CoA or in the branch towards lignin showed contrasting responses to R:FR and lignin levels were unaffected by R:FR (Fig. 3). Conversely, the genes involved in the early steps of flavonoid biosynthesis, between p-coumaril CoA and dihydrokaempferol showed a rapid reduction in expression in the stem in response to low R:FR, which in some cases remained 4 d later (Fig. 3). Three genes involved in subsequent steps between dihydroquercetin and anthocyanin showed a severe reduction of expression in the stem after 4 d of treatment. Consistent with the patterns of expression of upstream genes in the flavonoid pathway, anthocyanin levels were high only in the stem of high R:FR controls and very low in the stem of low R:FR-treated plants and the leaves of both groups (Fig. 3). Kaempferol and quercetin, which derive from dihydrokaempferol, also showed reduced levels in the stem of low R:FR-treated plants.

The flavonoid synthesis genes that show reduced expression in the stem of tomato plants exposed to low R:FR did not respond to low R:FR in whole Arabidopsis seedlings (based on the analysis of publicity available data (Sessa et al., 2005)). In Arabidopsis, mutations at the PHYB or PHYD genes involved in the perception of high R:FR, affect the accumulation of anthocyanin in the stem (Aukerman et al., 1997). Therefore, a putative response of anthocyanin synthesis genes in the stem of Arabidopsis could become diluted by the comparatively large proportion of biomass corresponding to cotyledons and expanding leaves.

Flavonol levels increased in the leaves (mainly quercetin and rutin) in response to low R:FR (Fig. 3). In the case of rutin, this leaf response could be in part due to the promotion of the most highly expressed UDP-glucose:flavonoid 3-O-glucosyltransferase (UFGT) genes in the leaves. Noteworthy, the At2g36790 gene, which encodes a UDP-glucose:flavonol-3-O-glycoside-7-O-glucosyltransferase, showed a 3-fold promotion of expression by low R:FR in the Arabidopsis leaf-rich samples (high R:FR: 32 ±0, 4 d Low R:FR: 117 ±8) (Sessa et al., 2005). In other words, this flavonol biosynthesis gene increases its expression in response to low R:FR in the leaves of tomato (Fig. 3) and in leaf-rich Arabidopsis samples.
Low R:FR differentially affects photosynthesis and respiration in the leaf and stem

Photosynthesis-related genes are overrepresented in clusters S7 and S9, showing a rapid reduction in stem expression in response to low R:FR (Fig. 1C). These genes are linked mainly to dark reactions (Calvin cycle) (Fig. 4D). In addition, genes related to isoprenoid biosynthesis and steroid metabolism, which represent steps upstream chlorophyll and carotenoid biosynthesis, are respectively overrepresented in clusters S8 and S9 (Fig. 1C). These observations prompted us to measure the rates of net carbon dioxide exchange against irradiance in the stem and in the leaves of tomato plants exposed to low R:FR for 4 d. In the controls exposed to high R:FR, the stem showed a significant photosynthetic capacity, equivalent to approximately 1/3 of the leaf capacity at saturating irradiance (Fig. 4B). Noteworthy, in low R:FR-treated stems net carbon dioxide exchange reached the compensation point at 100 μmol . m⁻² . s⁻¹ and remained close to zero at higher irradiances (Fig. 4B). Furthermore, the rate of stem mitochondrial respiration (i.e. exchange rates at irradiance =0) was approximately half in the plants treated with low R:FR. The internal concentration of carbon dioxide was lower in low R:FR-treated stems, indicating that the limitation was non-stomatal (Fig. 4C). The levels of chlorophyll (Fig. 4E) and carotenoids (Fig. 4F) were also severely reduced in the stem of low R:FR-treated plants, which is consistent with a strong downszie of the photosynthetic apparatus.

The leaves also showed some reduction in maximum photosynthesis (rates of net carbon dioxide exchange at saturating irradiance) (Fig. 4A), which are consistent with previous observations in the phyB mutant of Arabidopsis (Boccalandro et al., 2009). However, the magnitude of the leaf response is small when compared to that of the stem. Photosynthesis-related genes, in particular genes linked to light reactions, are overrepresented in cluster I6, showing reduced expression in the leaves in response to 4 d of low R:FR (Fig. 1B).

Low R:FR differentially affects leaf and stem jasmonic acid levels

Jasmonic acid (JA), abscisic acid (ABA), ethylene (ACC) and auxin (IAA) have been reported to stimulate the synthesis of flavonoids (Feys et al., 1994; Jeong et al., 2004; Robson et al., 2010; Lewis et al., 2011). To investigate the signals potentially involved in the organ-specific patterns of expression of flavonoid biosynthesis genes and flavonoid compound levels we measured the contents of these hormones in the stem and leaves of plants exposed 4 d to low R:FR, compared to the high R:FR controls.
The levels of JA were significantly reduced in the stem of low R:FR-treated plants (Fig. 5), showing a pattern consistent with that of anthocyanin levels and the expression of anthocyanin-synthesis genes. Noteworthy, the expression of the 12-Oxophytodienoate-10,11-Reductase gene involved in JA synthesis (Schaller et al., 1998; He et al., 2002) showed a rapid decrease in the stem of low R:FR-treated plants (Table S2). The levels of ABA increased in the leaves of low R:FR-treated plants, those of ACC showed a similar promotion by low R:FR in the stem and leaves and IAA contents were not significantly affected by R:FR (Fig. 5).

Light-dependent induction of anthocyanin by JA

To investigate the physiological role of the stem-specific reduction in JA content we selectively added JA to the stem of plants exposed to either high or low R:FR. Exogenous JA increased anthocyanin levels in the high R:FR controls (Fig. 6A), suggesting that the higher levels of endogenous JA under this light condition (compared to low R:FR) would be a requisite for the observed high anthocyanin levels. Exogenous JA had no effects in low R:FR-treated plants (Fig. 6B), suggesting that the reduction of anthocyanin contents by low R:FR would be the result of reduced sensitivity to JA.

Previous results have provided genetic evidence for a negative role of JA signalling in shade avoidance (Robson et al., 2010). Addition of JA to the stem promoted the growth of this organ in tomato plants grown under high R:FR but it did not promote stem growth at low R:FR (Fig. 6B). Therefore, JA also reduced shade avoidance in our system (note reduced difference between high and low R:FR in JA-treated stems).

DISCUSSION

When plants are shaded by neighbours, the photoassimilates that fuel growth become limited by light availability. In response to the low R:FR signals of dense vegetation the growth of the stem is promoted to place the leaves at a higher position, with better chances to capture light for photosynthesis (Smith, 1982; Casal and Smith, 1989; Ballaré, 1999; Morelli and Ruberti, 2002; Franklin and Whitelam, 2005; Casal, 2012). This shade-avoidance response involves enhanced growth when the available photoassimilates may already be limiting. Thanks to the high sensitivity to small reductions in R:FR caused by far-red light reflected from neighbours, in growing
canopies the stem growth response can anticipate actual shade (Ballaré et al., 1987) but even so, shade-avoidance reactions continue after actual shade is established and photoassimilates become scant. The analysis of the tomato stem transcriptome and follow-up experiments reported here reveal mechanisms that reduce the energetic cost of shade avoidance responses.

Genes involved in early steps of the flavonoid-synthesis pathway and downstream genes leading to anthocyanin synthesis showed comparatively high levels of expression in the stem of plants exposed to high R:FR, a strong (and often rapid) reduction in response to low R:FR, and lower expression levels in the leaves (Fig. 3). A similar pattern was observed for the end product, as anthocyanin levels were high only in the stem of high R:FR controls. The biosynthesis of flavonols diverges from that of anthocyanin beyond dihydrokaempferol. The levels of the flavonols kaempferol and quercetin were also reduced in the stem in response to low R:FR. Flavonol levels were high in the leaves, where they were not reduced by low R:FR (Fig. 3) in accordance with previous results in *Brassica napus* (Gerhardt et al., 2008) and *Betula pendula* (Tegelberg et al., 2004). Actually, in tomato leaves we observed increased quercetin and rutin levels in response to low R:FR, which correlated with the promotion of expression by low R:FR of some flavonol-related genes in these organs (Fig. 3). Low R:FR also reduced the expression of early genes in the isoprenoid and steroid pathways in the stem (Fig. 1C). Some of the downstream products of the latter genes, such as chlorophyll and carotenoids, were also severely reduced in the stem (Fig. 4E, F) while ABA, which is another downstream product, was unaffected in this organ (Fig. 5), indicating the occurrence of additional control points.

In order to investigate hormonal changes potentially involved in the observed flavonoid responses we measured the levels of IAA, ABA, JA and ACC in the stem and leaves as these hormones had been implicated in the induction of flavonoids. In young Arabidopsis seedlings low R:FR increased the abundance of IAA (Tao et al., 2008; Hornitschek et al., 2012) but this is not the case in the petiole and lamina of Arabidopsis leaves (Kozuka et al., 2010) or in the stem and leaves of tomato (Figure 5). We did observe the promotion of auxin-related genes (Fig. 2C) typically reported for Arabidopsis. One of the possibilities is that in Arabidopsis leaves and tomato organs low R:FR cause IAA tissue redistribution rather than changes in organ IAA levels. Diversion of auxin towards outer stem cell layers has also been proposed for Arabidopsis hypocotyls (Keuskamp et al., 2010). Auxin transport could be favoured by the reduced flavonoid levels observed under low R:FR (Brown et al., 2001). Low R:FR increased ACC levels in tomato as reported for sorghum (Finlayson et al., 1998; Finlayson et al., 1999) and tobacco (Pierik et al., 2004) and increased ABA levels in the
leaves as reported for the phyB mutant of Arabidopsis (González et al., 2012). However the patterns of these hormone changes do not reflect the organ dependency of flavonoid responses. Only JA decreased in the stem and not in the leaves in parallel with the flavonoid patterns. Previous reports had also shown no JA changes in the leaves of Arabidopsis exposed to different R:FR (Moreno et al., 2009). Adding JA selectively to the stem increased anthocyanin levels in the stem of seedlings exposed to high R:FR but not of those exposed to low R:FR (Fig. 6). These results suggest that anthocyanin levels are high under high R:FR due to increased JA levels and increased sensitivity to JA.

In Arabidopsis, the coi1, jar1, jin1, and jai mutants show enhanced promotion of hypocotyl elongation by low R:FR, indicating that JA biosynthesis and signalling genes reduce shade-avoidance responses (Robson et al., 2010). Application of JA to the stem of tomato yielded a result consistent with the latter observation as it reduced the response to low compared to high R:FR (Fig. 6A). Although in contrast to the case of Arabidopsis, this reduced response in tomato was caused by enhanced growth under high R:FR and not reduced growth under low R:FR. The observed reduction of JA levels in the stem of tomato exposed to low R:FR could therefore be part of a positive feed-back loop of shade-avoidance responses in this organ.

In tomato, non-leaf green organs, including the stem, are potentially quite active photosynthetically (Hetherington et al., 1998). However, in some cases the tomato stem shows no net carbon dioxide exchange because its photosynthetic capacity can be enough to re-fix the respired carbon dioxide (Xu et al., 1997). Here we show that in plants grown under high R:FR the stem certainly has significant photosynthetic capacity, equivalent to approximately one third that of the leaves. However, under low R:FR this capacity is substantially reduced and net rates of carbon dioxide exchange are close to zero at irradiances above the compensation point (Fig. 4B). There would be no benefit in maintaining the maximum potential of the photosynthetic apparatus when shade limits the expression of this potential. The reduction of photosynthesis has a non-stomatal origin (Fig. 4C) and could be caused by limitations in the Calvin Cycle (Fig. 4D). In addition, the rates of mitochondrial respiration are also strongly reduced by low R:FR.

Taken together, the results indicate that in response to low R:FR the stem photosynthetic apparatus is downsized. In the stem, photosynthetic and photoprotective pigment abundances decline, the photosynthetic capacity becomes minimized and a positive relationship between R:FR and JA levels appear as an important component of the control of anthocyanin content. In turn, these changes reduce the energetic cost of producing and maintaining a longer stem and this is
manifested in lower respiration rates. The low R:FR-induced savings are very selective as they mainly involve the stem and not the leaves and, within the stem, they involve processes linked to light (which is becoming scant) and not processes related to the stem physical support function, as lignin concentrations are not reduced.

MATERIALS AND METHODS

Plant Material and growth conditions

Seeds of tomato (Solanum lycopersicon, La Germinadora, Buenos Aires, Argentina) were sown on wet paper at 22 ºC under continuous white light (PAR= 51 µmol m⁻² s⁻¹). Seven-day-old seedling were transplanted to 70 cm³ pots containing equal amounts of perlite (Perlome; Buenos Aires), peat most (Finca Don Calvino, Buenos Aires) and vermiculite (Intersum, Buenos Aires) and watered as needed with a solution containing 1 g L⁻¹ of Hakaphos R (Compos, Buenos Aires). Plants were grown under continuous light at 130 µmol m⁻² s⁻¹ of PAR (Li-188B sensor; Li-COR) provided by fluorescent tubes and the temperature was 22 ºC. The R:FR (Skye meter SKR 100, remote probe SHR1100; Skye Instruments) was 4.6.

Light treatments

Twenty four-day-old plants were separated in two groups. The control group continued to grow under the high R:FR (= 4.6) conditions described above. The low R:FR (= 0.05) group received continuous supplementary far-red light (7.8 µmol m⁻² s⁻¹) provided from both sides by incandescent lamps in combination with a water filter (10 cm width), a red filter (no. 026, Lee Filters), and three blue acrylic filter (2.5 mm thick, Paolini 2031, Buenos Aires).

Stem and Leaf growth.

The length of the first internodes (the stem between the cotyledonary node and the first true leaf node) was measured with a ruler to the nearest millimetre. Dry weight was determined after drying at 70 ºC during 1 d. Leaf area was measured with a LI-3100 meter (LI-COR, Lincoln, Nebraska). For detailed kinetics, the plants were photographed with a Canon Power Shot A520 camera and the images of the different time points were aligned using Photoshop 7.0 to record internode length increments.
Microarray Experiments

In a first experiment, the first pair of leaves and the first internode of both low R:FR and control plants (three biological replicates) were harvested in liquid nitrogen, and total RNA was extracted with RNAEasy Plant Mini Kit (Qiagen). In a second experiment, only the first internode was harvested at different times after transfer from high to low and from low to high R:FR (two biological replicates per time point). cDNA and cRNA synthesis and hybridization to Affymetrix Tomato Gene Chips were performed following Affymetrix instructions. Expression data were normalized to the total sum of expression values of each microarray (Clarke and Zhu, 2006), restricted by presence criteria (presence flags for all the replicates of at least one condition), and used either for factorial ANOVA with R:FR and organ as factors or for one-way ANOVA (seven treatments). P values provided by ANOVA were used to calculated \( q \)-values (Storey and Tibshirani, 2003). The cut-off values for the effects of treatments, R:FR, organ and interaction in the factorial ANOVA of the first experiment and for the effects of treatments in the second experiment were set at \( P < 0.05 \) or \( q < 0.10 \), whichever was reached first. These values are provided in Tables S1 and S2. To identify the clusters we used the standard setting of the dChip Software (Li and Wong, 2003).

Lignin

Relative lignin levels were determined using the thioglycollic acid method (Bruce and West, 1989). Samples containing 200 mg of plant material were stored at -20 °C and subsequently homogenized. Absolute methanol (25 mL) was added to the samples, which were vacuum filtered and rinsed with further methanol on filter paper. The solid material was dried at 60 °C for at least 24 h, combined with 5 ml 2 M HCl and 0.5 ml 80% (v/v) thioglycollic acid (Riedel-deHean, Germany) and placed in boiling water for 4 h. After centrifugation (30000 g, 10 min, 4 °C) the pellets were washed with distilled water, re-suspended in 5 ml 0.5 M NaOH and incubated at 25 °C for 18 h. After further centrifugation, 1 mL of concentrated HCl was added to the supernatants and incubated for 24 h at 4°C. Finally, the samples were centrifuged (1000 g, 10 min, ca 25°C) and the pellet re-suspended in 3 ml of 0.5 M NaOH, where absorbance was determined spectrophotometrically at 280 nm.
**Flavonol levels**

Samples were ground with liquid nitrogen and flavonoids extracted in methanol:hexane (1:0.5 v/v per 0.1 g fresh weight) and stirred overnight at 25 °C with rotation at 200 rpm, followed by centrifugation at 13,000 rpm for 10 min. The methanol fraction was dried in a rotary evaporator and the sample resuspended in 400 µL of methanol (protocol based on (Torres et al., 2005). Resuspended extracts (5µL) were injected into an Agilent 1100 HPLC system fitted with an Eclipse XDB-C18 column. Gradient elution was conducted at a flow of 1 mL / min, at 40°C. Mobile phase solvents (HPLC grade) were water:acetic acid (99.90:0.10, v/v, A) and acetonitrile:acetic acid (99.90: 0.10, v/v, B). The initial concentration was 15 % B and linear-gradient elution was carried out to 36.5 % B at 30 min. Detection was achieved with the Agilent 1100N G1365 MWD module at 270 nm, 280 nm, 254 nm and 220 nm.

**Chlorophyll, carotenoid and Anthocyanin levels**

The samples were harvested in 1ml Acetone (pure solvent) and incubated in darkness at -20 °C for at least 3 d. Absorbance was measured at 661.6, 644.8, and 470 nm to calculate chlorophyll and carotenoids levels (Lichtenthaler and Buschmann, 2001). For anthocyanin levels, plant material was extracted in 1 ml 1% (w/v) HCl methanol. Measurements of A530 were corrected for chlorophyll absorption (657 nm) (Mancinelli et al., 1991).

**Leaf Photosynthesis**

A portable gas-exchange system (LI-COR 6400; LI-COR) was used to obtain curves of CO₂ exchange against PAR provided by the red and blue diode gas –exchange system (6400-02B LED light source; LI-COR) in fully expanded first leaves and first internode stems. The area included in the 6 cm² chamber was recorded for each sample. Measurements started at 2000 µmol m⁻² s⁻¹ of PAR in the references chamber, and decreased stepwise to 1500, 1000, 750, 500, 400, 300, 200, 100, 70, 35 and 0 µmol m⁻² s⁻¹.

**Analysis of hormone abundances**

Freeze dried stem and leaf tissue was finely chopped with a razor blade and the masses were measured (2.74 to 3.2 mg dry weight). A mixture of stable isotope
labelled hormones including 0.76 ng $^{13}$C$_2$-JA, 1 ng $^{13}$C$_6$-IAA, 10 ng $^2$H$_6$-ABA and 10 ng $^2$H$_4$-ACC was added to each sample replicate in a 1.7 mL centrifuge tube. 500 μL of methanol warmed to 55 °C was added, the tissue was ground with a micropestle, and re-extracted with 500 μL of methanol and then with 500 μL of 80% ethanol. The extractions were centrifuged and the cleared supernatants were pooled after each extraction. The pooled extracts were dried and the residue was re-suspended in 800 μL of chloroform and partitioned against 1 mL of H$_2$O adjusted to pH 9.0 with NH$_4$OH. The aqueous fraction was recovered, adjusted to pH 5.0 with acetic acid and partitioned against 1 mL of ethyl acetate. The organic fraction (JA, IAA, ABA) was dried and then methylated with ethereal diazomethane. The aqueous fraction containing ACC was dried, re-dissolved in 100 μL of 0.1 N acetic acid and applied to a 100 μL Dowex 50 column pre-washed with 500 μL 0.1 N acetic acid. The column was washed with 500 μL 0.1 N acetic acid. 80 μL of 2 N NH$_4$OH was applied to the column and eluted to waste, and the free ACC was collected in the next 180 μL of 2 N NH$_4$OH and then dried. The ACC was derivatized with 80 μL of 3 mM phthalic anhydride at 90 °C for 40 min, dried, and then methylated with ethereal diazomethane. Samples were analyzed on an Agilent 7890A/5975C XL GC-MS equipped with a 0.25 mm x 30 M DB-5MS column (0.25m film) using pulsed splitless injection. He was used as the carrier gas at 0.75 mL/ min. The inlet was maintained at 250 °C and the oven was ramped from 45oC (2.25 min initial hold) to 250 °C at 40 °C per minute, held at 250 °C for 3 minutes, and then ramped to 290 °C at 40 °C per minute. The ion source temperature was maintained at 230 °C and the quadrupole was heated to 150 °C. The ion source was operated in electron impact mode and both scan and selected ion data were acquired. Two ion pairs were monitored for each hormone, and the larger fragments were used for quantification (JA- 193,195,224,226; IAA- 130,136,189,195; ABA- 162,166,190,194; ACC- 213, 216, 245, 249 m/z).

**Supplemental Data**

The following materials are available in the online version of this article.

**Supplemental Table S1.** Stem and leaf transcriptome responses to R:FR.
**Supplemental Table S2.** Rapid changes in stem transcriptome in response to R:FR.
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**FIGURE LEGENDS**

**Figure 1.** Shared and specific responses of the stem and leaves of tomato to low R:FR. **A.** Tomato plant grown under low R:FR for 4 d compared to the high R:FR control. **B.** Clusters of genes with stem and/or leaf expression affected by 4 d low compared to high R:FR. Clusters A1-A9 show additive effects of R:FR and organ and clusters I1- I11 show interaction between R:FR and organ. **C.** Clusters of genes with early stem expression responses to low R:FR (S1-S9). In B and C, data are means and SE of all the genes represented in the cluster (the number of genes is indicated in brackets), clusters with less than 30 genes are not included (see Table S1) and overrepresented ontology terms are indicated. **D.** Number of genes with expression promoted or inhibited by 4 d low, compared to high R:FR in the stem and or leaves (the number of genes with a fold change of at least two is given in brackets). **E.** Correlation of the log10 low R:FR / high R:FR expression ratio of the genes significantly affected after 4 h and 4 d by low R:FR.

**Figure 2.** Promotion of stem growth by low R:FR. **A.** Stem length increment as affected by 1 or 4 h of low R:FR and the subsequent transfer back to high R:FR. The growth rate is indicated close to the corresponding regression line. **B.** Size (length or area) and dry weight of the stem and first pair of leaves in seedlings exposed to 4 d of low R:FR, compared to the high R:FR controls. **C.** Low R:FR promotes the expression of auxin response genes (*IAA7, IAA4, Les. 3702, Les. 4097*) and cell-wall genes (*XTH9, CELL WALL INVERTASE*). In A and B, data are means ±SE of 7 plant replicates and significant effects of low compared to high R:FR are indicated, *t* test, ***: P <0.001. Data shown in B are from Tables S1 and S2.

**Figure 3.** Reduced levels of flavonoids in the stem of low R:FR-treated plants. The chart shows the steps of lignin and flavonoid (anthocyanin, flavonol) biosynthesis catalyzed by enzymes encoded by genes showing significant expression responses to R:FR. The expression in the stem and leaf of plants treated for 4 d with low R:FR compared to the high R:FR control and/or in the stem of seedlings exposed 1 or 4 h to low R:FR and subsequently transferred back to high R:FR is shown when significant differences were observed (Data from Tables S1 and S2). The levels of lignin, kaempferol, rutin, quercetin and anthocyanin in plants treated for 4 d with low R:FR compared to the high R:FR control is indicated (data are means ±SE of 4-12 plant replicates and significant effects of low compared to high R:FR are indicated, *t* test, *: P <0.05, **: P <0.01). UGG: UDP-glucose:glucosyltransferase, UFGT: Flavonoid 3-
glucosyl transferase, C4H: Trans-cinnamate 4-monooxygenase, CHl: Chalcone isomerase-like protein, CHS: Chalcone synthase F3H: Flavanone 3 beta-hydroxylase, IFR: Isoflavone reductase, FLS: Flavonol synthase, ANS: Anthocyanin synthase, DFR: Dihydroflavonol 4-reductase, RT: Rhamnosyltransferase, HQT: Hydroxycinnamoyl CoA quinate transferase. The chart is based on BRENDA data base (http://www.brenda-enzymes.info) (Scheer et al., 2011).

**Figure 4.** Low R:FR severely reduces the stem photosynthetic capacity. **A.** Net carbon dioxide exchange in the leaves of plants treated for 4 d with low R:FR compared to their high R:FR control. **B.** Idem in the stem of these plants. **C.** Internal carbon dioxide concentration for the stems shown in B. **D.** Low R:FR reduces the expression of Calvin Cycle protein genes. **E.** Chlorophyll content in the stem and leaf of plants treated for 4 d with low R:FR compared to the high R:FR control. **F.** Idem for carotenoid content. In A-C, E and F data are means ±SE of 6 (A), 3 (B, C) or 7 (E, F) plant replicates and significant effects of low compared to high R:FR are indicated, $t$ test, *: P <0.05, **: P <0.01, ***: P <0.001. In A, maximum net carbon dioxide exchange is significantly different if the rates are corrected by using rates at 300 μmol.m⁻².s⁻¹ as covariate. Data shown in D are from Table S2.

**Figure 5.** Low R:FR alters the hormonal balance. Jasmonic acid (JA), abscisic acid (ABA), ethylene (ACC) and auxin (IAA) in the stem and leaves of plants treated for 4 d with low R:FR compared to their high R:FR control. Data are means ±SE of five plant replicates and significant effects of low compared to high R:FR are indicated, $t$ test, *: P <0.05, ***: P <0.001.

**Figure 6.** Exogenous JA applied to the stem promotes stem anthocyanin (A) and stem growth (B) in high but not in low R:FR-treated plants. The stem was painted with a brush soaked in a water solution containing 450 μM JA and 0.015 % v/v of Silwet, twice during the first day of treatment and daily during the subsequent 3 d of treatment. Controls were painted similarly but without JA. Data are means ±SE of 3 (A) or 7 (B) plant replicates and significant effects of JA are indicated, $t$ test, *: P <0.05, **: P <0.01, ***: P <0.001.
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