Far-red light promotes *Botrytis cinerea* disease development in tomato leaves via jasmonate-dependent modulation of soluble sugars

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**Abstract**
Plants experience a decrease in the red:far-red light ratio (R:FR) when grown at high planting density. In addition to eliciting the shade avoidance response, low R:FR also enhances plant susceptibility to pathogens via modulation of defense hormone-mediated responses. However, other mechanisms, also affected by low R:FR, have not been considered as potential components in FR-induced susceptibility. Here, we identify FR-induced accumulation of leaf soluble sugars as a novel component of FR-induced susceptibility. We observed that phytochrome inactivation by FR or phytochrome B mutation was associated with elevated leaf glucose and fructose levels and enhanced disease severity caused by *Botrytis cinerea*. By experimentally manipulating internal leaf sugar levels, we found that the FR-induced susceptibility in tomato was partly sugar-dependent. Further analysis revealed that the observed sugar accumulation in supplemental FR occurred in a jasmonic acid (JA)-dependent manner, and the JA biosynthesis mutant *def1* also displayed elevated soluble sugar levels, which was rescued by exogenous methyl jasmonate (MeJA) application. We propose that the reduced JA responsiveness under low R:FR promotes disease symptoms not only via dampened induction of defense responses, but also via increased levels of soluble sugars that supports pathogen growth in tomato leaves.

**KEYWORDS**
Botrytis, defense, jasmonate, phytochromes, soluble sugars, tomato

**INTRODUCTION**
As photoautotrophic organisms, plants harness light energy into sugars through photosynthesis. Red (R) and blue (B) light are absorbed by plant tissue while far-red (FR) light is reflected by or transmitted towards neighbouring vegetation. At high planting density, plants experience a drop in the R:FR ratio due to the depletion of R and the omnipresence of FR radiation in the canopy. Changes in the R:FR ratio in the environment are sensed by specialized photoreceptors called phytochromes, where phyB plays a dominant role and can be found in two photoconvertible forms, the active (Pfr) and inactive (Pr) form (Franklin, 2008). The perception of low R:FR conditions subsequently leads to photoconversion of phyB into the inactive Pr form that is restricted to the cytoplasm. This phenomenon prevents phyB interaction with the bHLH-type phytochrome interacting factors (PIF) transcription factors and their consequent inactivation and degradation by
et al., 2018). The resulting accumulation of PIFs initiates downstream growth responses, largely via increased auxin synthesis and response in Arabidopsis known as shade avoidance (Hornitschek et al., 2012; Li et al., 2012; Pantazopoulou et al., 2017).

The perception of low R:FR or the genetic inactivation of phyB in plants is known to affect plant immunity towards distinct pathogens (Ballaré, 2014; Ballaré & Pierik, 2017; Pieterse, Pierik, & Van Wees, 2014). This so-called FR-induced susceptibility is an example of the well-known growth-defense trade-off, and has been studied in a number of plants species and pathosystems. Phytochrome inactivation upon low R:FR promotes lesion development induced by the necrotrophic fungus Botrytis cinerea and enhances Mamestra brassicae and thrips feeding performance in Arabidopsis and tomato (Cortés, Weldegergis, Boccalandro, Dicke, & Ballaré, 2016; Izaguirre, Mazza, Blondini, Baldwin, & Ballaré, 2006; Ji et al., 2019). Compromised plant defense towards B. cinerea in low R:FR conditions goes via a decrease in secondary metabolite production and defense gene activation in Arabidopsis (Cargnel, Demkura, & Ballaré, 2014; De Wit et al., 2013).

Interestingly, even though plant growth and immunity are known to influence each other, these are not passive trade-offs, but involve molecular control. During FR enrichment, the gibberellin (GA) pathway is strongly activated to promote shoot elongation via degradation of growth-inhibiting DELLA proteins. However, DELLA proteins not only inhibit growth, but also promote jasmonic acid (JA) signalling by physically interacting with JAZ proteins, negative defense regulators (Cerrudo et al., 2012; Hou et al., 2010; Leone, Keller, Cerrudo, & Ballaré, 2014). DELLA removal in low R:FR therefore releases JAZ proteins that in turn sequester MYC transcription factors, thereby preventing the activation of JA-associated defense gene expression (Chico et al., 2014; Leone et al., 2014; Pieterse et al., 2014). In addition, low R:FR conditions were also shown to reduce the pool of bioactive JAs available in Arabidopsis plant tissue upon exogenous MeJA treatment and was associated with increased susceptibility to B. cinerea (Fernández-Milmanda et al., 2020). Although the DELLA-JAZ interaction or the decrease in JA response explains part of the FR-induced susceptibility, there likely are several other layers of regulation involved in balancing shade avoidance and defense against pathogens and pests.

In addition to modulating plant defense against pathogens, phytochrome signalling has also been described to influence carbohydrate levels in plants. In Arabidopsis, plants lacking phytochromes showed an increase in soluble sugars and starch during daytime compared to wild-type plants indicating the involvement of phytochrome in the control of primary metabolism (Yang, Seaton, Krahmer, & Halliday, 2016). In addition, low R:FR treatment promotes carbohydrate levels in various species, including Brassica rapa (De Wit et al., 2018), Allium cepa (Lercari, 1982), Cucumis sativus (Xiong, Patil, Moe, & Torre, 2011) and Nicotiana tabacum (Kasperbauer, Tso, & Sorokin, 1970).

Carbohydrates generated by plants are also the primary nutrient source for plant pathogens. It is known that plants strictly regulate sugar fluxes to restrict pathogens from accessing those resources. However, plant pathogens have developed the capacity to hijack the sugar transport machinery of their host to redirect carbohydrates towards the infection site (Chen et al., 2010; Doidy et al., 2012; Lapin & Van den Ackerveken, 2013). For instance, pathogenic bacteria and fungi were shown to promote the expression of different sets of SWEET sugar transporters in Arabidopsis upon infection showing a pathogen-specific sugar transport manipulation (Chen et al., 2010). In grapevine, B. cinerea has been shown to upregulate VvSWEET4 and further analysis on Arabidopsis showed that sweet4 mutants are less vulnerable to the fungus (Chong et al., 2014). Although these studies mostly focus on the impact of a pathogen infection on the sugar allocation of the host, the impact of enhanced soluble sugar levels (e.g., via phytochrome inactivation) on disease severity has not received much attention.

High-sugar concentrations in plant tissue, although beneficial for the pathogen, are often associated with stronger plant defense potential. Studies have shown the involvement of sugars in the induction of secondary metabolites such as phenylpropanoids and/or defense gene induction such as MAPK or PR genes (reviewed in Moghaddam & Van Den Ende, 2012). In addition, a negative correlation between the defense hormone JA and soluble sugar levels was observed in Nicotiana attenuata (Machado, Arce, Ferrieri, Baldwin, & Erb, 2015). JA biosynthesis mutant plants displayed elevated monosaccharide levels in leaf tissue as well as increased Manduca sexta larval mass upon infection compared to wild-type plants (Machado et al., 2015). Also, a recent study reported that the disease severity induced by B. cinerea is negatively correlated to the relative fructose content in tomato stem (Lecompte et al., 2017).

Nevertheless, JA and sugar signalling pathways are not always negatively correlated. Studies in Arabidopsis have shown that glucose and JA can act synergistically to increase glucosinolate (GS) levels which have been shown to promote plant resistance towards B. cinerea in Arabidopsis (Buxdorf, Yaffe, Barda, & Levy, 2013; Guo et al., 2013; Kliebenstein, Rowe, & Denby, 2005). Overall, even though it becomes clear that plant soluble sugar status can influence plant immunity, it is still unclear whether elevated soluble sugar levels would promote or inhibit disease development.

Here, we propose that phyB inactivation could enhance disease severity via soluble sugar accumulation in plant tissue. We show that tomato plants exposed to WL + FR or lacking phyB have increased levels of glucose and fructose in leaves compared to WL-treated plants. By experimentally modulating soluble sugar levels in tomato leaves, we associate high soluble sugar levels to enhanced symptom development induced by B. cinerea. Furthermore, a tomato mutant deficient in JA biosynthesis exhibited high soluble sugar levels in leaf tissue accompanied with elevated disease severity. Consistent with these findings, soluble sugar levels as well as plant susceptibility was diminished by exogenous MeJA both in the JA biosynthesis mutant and wild-type plants showing the occurrence of JA-mediated soluble sugar modulation in tomato leaves. As JA responsiveness is decreased in tomato phytochrome mutants (Cortés et al., 2016), we propose that the elevated soluble sugar levels found in WL + FR-treated plants can be mediated by reduced JA signalling in low R:FR light. These results
indicate that the FR-induced susceptibility in tomato leaves is partly controlled by phyB signalling as it regulates JA-mediated soluble sugar modulation in turn affecting disease severity induced by pathogens.

2 | MATERIALS AND METHODS

2.1 | Plant growth and light treatments

Seeds from tomato (Solanum lycopersicum) cv. Moneymaker (LA2706) and phyB1phyB2 double mutant (LA4364, cv. Moneymaker), were obtained from the C. M. Rick Tomato Genetics Resource Center and propagated by the Horticulture and Product Physiology group from Wageningen University as part of the LED it Be 50% consortium. Tomato cv. Castlemart and JA biosynthesis mutant defenseless 1 (def1) (cv. Castlemart), kindly provided by Dr. Maria J. Pozo (Dept Soil Microbiology and Symbiotic Systems, Granada, Spain), were sown in 9 cm pots with regular potting soil (Primasta soil, the Netherlands). Plants were grown for 4 weeks after sowing in climate chambers (MD1400; Snijders, the Netherlands) in long-day photoperiod (8-hr dark/16-hr light) at 150 μmol m⁻² s⁻¹ photosynthetically active radiation (PAR) using white light (WL, R:FR ratio = 5.5) generated by Philips GreenPower LED research modules (Signify B.V.). The calculated phytochrome photostationary state (PSS) value was 0.8 equivalent to the proportion of active phytochrome (Pfr) within the total of active and inactive (Pr) phytochrome present (Sager, Smith, Edwards, & Cyr, 1988). For additional FR treatments, 3-week-old plants were exposed for 5 days, starting at Zeitgeber time (ZT) = 3 on the first day, to WL supplemented with uniform far-red (FR) Philips GreenPower LED research modules referred to as WL + FR (R:FR = 0.14, PSS value = 0.5). Local FR radiation was performed by using custom FR LEDs (724–732 nm) attached onto a flexible arm (Figure S1). The top leaflet of either the third or fourth oldest leaf of 4-week-old WL-exposed plants was illuminated with FR. WL-exposed plants which did not receive local FR illumination were taken as control.

2.2 | Pathogen growth conditions and bioassays

2.2.1 | Botrytis cinerea

B. cinerea strain Bc 05.10 (Van Kan, Van’t Klooster, Wagemakers, & Dees, 1997) and a pepper isolate (Denby, Kumar, & Kliebenstein, 2004; Grant, Chini, Basu, & Loake, 2003) were maintained on half strength Potato dextrose agar medium (PDA ½, BD Difco™) and grown for approximately 10 days at room temperature under natural daylight conditions. The spore suspensions were prepared according to Van Wees, Van Pelt, Bakker, and Pieterse (2013) and diluted to a final concentration of 1.5 x 10⁵ spores ml⁻¹ in half strength potato dextrose broth (PDB ½, BD Difco™) prior to inoculation. Bioassays were performed on detached tomato leaflets previously pretreated with WL or WL + FR for 5 days. The leaflets were placed in square Petri dishes onto Whatman® filter soaked with 6 ml of tap water to avoid dehydration. The adaxial side of the leaflets was drop-inoculated three to six times with 5 μl of spore suspension. Plates were sealed with PARAFILM® M and incubated for 3 days in their respective light pretreatment conditions (WL or WL + FR). Pictures were taken 3 days post inoculation (dpi) and lesion areas were assessed by going along the lesion edges using the polygon selection tool of the ImageJ software.

In vitro mycelium diameter measurements were performed by depositing a 5 μl droplet of a 1.5 x 10⁵ spores ml⁻¹ solution onto 1.5% granulated agar (BD Difco™) plates supplemented with homogenates of WL- or WL + FR-treated leaves (1 gFW ml⁻¹), or onto PDA ½ supplemented with increasing concentrations of D-glucose (Duchefa Biochemie; 10, 20 and 50 g L⁻¹). Inoculated plates were incubated for 3 days in WL at room temperature. The newly grown mycelium diameter was measured by using a digital caliper.

2.2.2 | Phytophthora infestans

P. infestans strain P188069 (Kamoun et al., 1997) was maintained on Rye sucrose agar medium (RSA, Caten & Jinks, 1968) for 10 days in darkness at room temperature. Cultured plates were flooded with 10 ml of sterile MilliQ water and incubated at 4°C for 3 hr. Zoospores were collected, counted and adjusted to a concentration of 5 x 10⁶ zoospores ml⁻¹ prior to the bioassays. The abaxial side of detached tomato leaflets was drop-inoculated several times with 10 μl of zoospores suspension and leaflets were incubated for 3 days in their respective light pretreatment conditions (WL or WL + FR). The lesion areas were measured after 3 dpi with the ImageJ software. Statistical analysis was performed with student t-test, p-value <.05.

In vitro mycelium diameter measurements were performed by depositing a plug of mycelium (Ø 0.6 cm) in the centre of ½ strength V8 agar plates (1.5% granulated agar, BD Difco™) supplemented with increasing concentrations of D-glucose (Duchefa Biochemie; 0, 10, 20 and 50 g L⁻¹). Inoculated plates were incubated for 10 days in darkness. The newly grown mycelium diameter was measured by using a digital caliper.

2.2.3 | Pseudomonas syringae

P. syringae pathovar tomato strain DC3000 (Whalen, Innes, Bent, & Staskawicz, 1991) was grown in liquid King’s medium B (KB; King, Ward, & Raney, 1954) and incubated overnight in an orbital shaker at 220 rpm at 28°C. The next day, the bacterial culture was centrifuged at 7,800 rpm (7,170 G) for 5 min. The bacterial pellet was washed three times with sterile MilliQ water and resuspended in 10 mM MgSO₄4. The inoculum was adjusted to OD₆₆₀ = 0.025 (OD₆₆₀ = 1 x 10⁹ cells ml⁻¹) and supplemented with 0.02% Silwet L-77 (Lehle seeds). The two first lateral leaflets of the third leaf of WL- and WL + FR-pretreated intact tomato plants were dipped in the bacterial inoculum for 3 s and whole plants were subsequently placed in their respective light treatment for 3 days at 95% RH. Two leaf
transferred in square Petri dishes and inoculated with B. cinerea spores as described above. Colony forming units (CFU) were counted and converted in CFU per cm² of leaf surface area.

For glucose-mediated bacterial growth measurements were performed by growing bacteria in liquid KB until OD₆₆₀ reached 0.1. For replicate, 4 ml of bacterial suspension was transferred into a new tube and supplemented with 1 ml of water or with glucose solutions at different concentrations (10, 20 and 50 g L⁻¹). The OD₆₆₀ was measured every hour for 3 hr.

2.3 | In planta modulation of soluble sugar levels

Tomato leaflets of the third leaf were detached under water and placed in square Petri dishes containing Whatman® filter paper (bottom 1/3 cut out) soaked with 2.5 ml of tap water to avoid dehydration. Plates were placed on a metallic rack at a 30° angle. The bottom of the tilted plates was filled with 13 ml of 0.5 M glucose solution for glucose supplementation experiments or 100 μM 3-(3,4-dichlorophenyl)-1,1-dimethylurea (DCMU, Sigma®) for sugar starvation experiments. Tap water was used as a control in mock conditions. Only the petiole of the leaflet was immersed in the solution. Plates were kept at a 30° angle and incubated in WL or FR conditions for 24 hr prior to sugar quantifications or bioassays with B. cinerea. Prior to inoculation, the leaflets were transferred into square Petri dishes and drop-inoculated following the procedure described above.

2.4 | Plant hormone treatments

Three-week-old tomato plants were placed under either WL or WL + FR for 5 days and were sprayed daily with a 50 μM methyl jasmonate (MeJA, Sigma-Aldrich®) or mock solution (0.1% EtOH) supplemented with 0.1% Tween20 (Duchefa Biochemie). Hormone treatments were performed at 10 a.m. every day (ZT = 3). For sugar quantifications, leaf discs (ø 0.6 cm) originating from each infected leaflet were punched for 5 days and were sprayed daily with a 50 g L⁻¹ glucose solution or 100 μM glucose solution (used as standards) were mixed with 185 μl of reaction mix containing 170 μl MilliQ, 10 μl of Buffer (Solution 1) and NADP⁺/ATP (Solution 2) in a transparent flat-bottomed 96-well plate. The absorbance at 340 nm (A₃₄₀nm) was measured at the start of the experiment as well as after a 15 min incubation at 37°C (R₀ and R₁). All wells were supplemented with 2 μl of yeast invertase (100 mg ml⁻¹, Sigma-Aldrich®) diluted in buffer (Solution 1) were added to each well and incubated for 25 min at 37°C before the final A₃₄₀nm measurement (R₃). The glucose standard curve equation (y = ax + b) was used for soluble sugar quantifications. Formulas used to determine soluble sugar concentrations:

\[
\text{Glucose} \left( \frac{\mu\text{mol}}{\text{mgFW}} \right) = \frac{(R₂ - R₁) - b}{a} + 135
\]

\[
\text{Fructose} \left( \frac{\mu\text{mol}}{\text{mgFW}} \right) = \frac{(R₃ - R₁) - b}{a} + 135
\]

\[
\text{Sucrose} \left( \frac{\mu\text{mol}}{\text{mgFW}} \right) = \frac{(R₄ - R₁) - b}{a} + 135
\]

3 | RESULTS AND DISCUSSION

3.1 | Phytochrome inactivation enhances soluble sugar levels and disease severity of B. cinerea in tomato

Previous studies have shown that Arabidopsis phytochrome mutants accumulate more soluble sugars during the day than the wild-type does (Yang et al., 2016). First, we studied whether, as in Arabidopsis, a phyB mutation would affect soluble sugar levels in tomato leaves and could in turn lead to increased disease severity towards the necrotrophic fungus B. cinerea. Since tomato has two PHYB genes, we grew the phyB1phyB2 double mutant under white light LEDs (WL) for 3 weeks. We recorded stem elongation for 5 days prior to quantifying soluble sugar levels in leaves and inoculation with B. cinerea spores (Figure 1). Tomato phyB1phyB2 plants exhibited a constitutive shade
avoids a strong stem elongation phenotype (Figure 1a) as well as elevated glucose and fructose levels in leaves compared to wild-type Moneymaker plants (Figure 1b). This was accompanied by an increased foliar susceptibility to B. cinerea in phyB1phyB2 compared to wild-type Moneymaker plants (Figure 1c). Possibly, the increase in soluble sugars in phyB1phyB2 mutants might benefit B. cinerea growth in planta, thereby enhancing disease development on tomato leaves.

Next, we studied whether alike the phyB1phyB2 double mutant (Figure 1b), inactivating phytochrome by supplementing the WL background with FR LED light (WL + FR) would increase soluble sugar levels in leaves. Over a course of 5 days of light treatment, we observed that soluble sugar levels, mainly glucose and fructose, were significantly increased in WL + FR-exposed leaves compared to the WL control starting at 3 days of treatment (Figure 2a–d). Altogether, our data demonstrate that the increase in soluble sugar levels occurring phytochrome mutants previously shown in Arabidopsis (Yang et al., 2016) is also observed in tomato (Figure 1b and Figure 2a). It seems paradoxical that WL + FR conditions would at the same time promote energy and carbon-requiring shoot elongation and soluble sugar accumulation in leaf tissue. However, since we measured sugars in tomato leaflets specifically, we cannot rule out the possibility that soluble sugars would accumulate in leaves at the expense of other plant organs. We hypothesize that WL + FR would cause a gradual sugar accumulation in leaf tissue beyond levels that would be fully consumed in the dark period leading to leftover carbohydrates in the leaves from the third day of treatment onwards (Figure 2a). Indeed, we could only observe an increase in susceptibility when leaves had been exposed to WL + FR for at least 4 days (Figure 2e), consistent with the proposition that FR-induced sugar accumulation in tomato leaves could promote B. cinerea development in plant tissue.

3.2 | Supplemental FR promotes symptom development induced by pathogens with distinct lifestyles

The increase in soluble sugar levels observed upon WL + FR treatment in tomato plants led us to speculate that not just B. cinerea, but also other pathogens with different lifestyles might develop better on such leaf material since they all principally feed on sugar resources from their host. We performed bioassays on WL- and WL + FR-treated plants using B. cinerea (B.c. 05.10) as well as a pepper isolate of B. cinerea to test the specificity of the FR-induced susceptibility of tomato leaves in response to different B. cinerea isolates (Figure 3a,b). We also tested the bacterial hemibiotroph Pseudomonas syringae pv. tomato DC3000 (Figure 3c) already described to induce more severe symptoms on Arabidopsis plants exposed to FR-enriched light conditions (De Wit et al., 2013). Lastly, we tested the oomycete Phytophthora infestans (88069) to study the effect of supplemental FR on tomato susceptibility towards a biotrophic pathogen (Figure 3d). Interestingly, WL + FR-exposed tomato plants displayed significantly more severe symptoms than the WL controls to all pathogens tested (Figure 3a–d). Also, all pathogens showed increased growth rate when exposed to increasing concentrations of glucose in vitro (Figure 3e–h). These data indicate that the FR-induced susceptibility in tomato is not specific to B. cinerea as supplemental FR increased symptom development caused by an array of pathogens.

We hypothesize that this phenomenon might partly be caused by the increase in soluble sugar levels upon phytochrome inactivation by WL + FR perception in tomato leaves. As an independent indication for this hypothesis, we used the B. cinerea-tomato pathosystem to study the effect of leaf chemical content in WL + FR-treated leaves, compared to WL-treated leaves, on B. cinerea mycelial growth in vitro. To this end, we mixed ground WL or WL + FR-treated leaf material to
a minimal water-agar medium and recorded *B. cinerea* mycelial growth after 3 days (Figure S2). As expected, the mycelium did not develop well in the water-agar control, but the addition of ground leaf material boosted the mycelial development of the fungus in vitro and to a higher extent on WL + FR-treated plant material than WL, even though the quantitative effect was rather small at this homogenate concentration (1 gFW ml\(^{-1}\)).

Additional FR was already described to increase Arabidopsis susceptibility towards *B. cinerea* and *P. syringae* via a dampening of the jasmonic acid (JA) and salicylic acid (SA) defense pathways, respectively (Cerrudo et al., 2012; De Wit et al., 2013). In addition, our results suggest that elevated soluble sugar levels found in tomato leaves in response to WL + FR might also promote lesion development via an accelerated growth of these pathogens in tomato leaves.

### 3.3 Modulation of leaf soluble sugar levels affects *B. cinerea* disease severity in tomato

Next, we checked whether a modulation of internal sugar levels in tomato leaves could affect *B. cinerea* lesion development. To this end, we use a detached leaflet assay allowing for passive solution uptake by the petiole prior to inoculation with the fungus (Figure 4a). After 4 days in either WL or WL + FR, tomato leaflets were detached and supplemented with either a 0.5 M glucose, 100 μM of the photosynthesis inhibitor DCMU or water as a control for 24 hr prior to sugar quantification (Figure 4b,d) or bioassays with *B. cinerea* (Figure 4c,e). Although WL + FR-treated plants already displayed an increase in soluble sugars compared to WL-treated plants in mock conditions, the glucose solution uptake by the leaflets increased the glucose levels in planta after 24 hr as well as the lesion area induced by *B. cinerea* (Figure 4c,e). Consistent with this, a DCMU treatment gave opposite effects and reduced soluble sugar levels in the leaflets in WL + FR light, which was accompanied by an increased resistance to *B. cinerea* compared to control conditions (Figure 4e). These results suggest that internal sugar levels are one of the components determining disease severity in tomato leaves upon infection with *B. cinerea*. Interestingly, the lesion areas were similar between mock-treated WL + FR-exposed plants and glucose-treated WL plants even though leaf sugar levels were significantly different between both treatments (Figure 4, c). As the bioassays shown in Figure 4 were all performed in WL conditions, the possibility of WL + FR affecting directly the pathogen growth can be ruled out. Nevertheless, we cannot exclude the effect of light quality on direct plant defense via, for example, hormone-mediated defense pathways or on the production of defense metabolites which could affect plant immunity upon WL + FR exposure (Cargnel et al., 2014).
In order to investigate whether a targeted FR enrichment (Local FR, hereafter) could have a direct influence on sugar levels and disease severity in a local and remote manner, we designed FR lighting setups allowing for spatially restricted supplemental FR illumination (Figure S1). We illuminated the top leaflet of the third oldest leaf (L3) for 5 days (Figure 5a) and observed a strong petiole elongation of the illuminated leaf compared to plants that did not receive any supplemental FR (Figure 5d) and quantified soluble sugars and disease symptoms in this local L4 (L4), as well as in the now older, distal L3 (L3). Upon local FR illumination on L4, we observed that both L3 and L4 itself exhibited higher soluble sugar content as well as increased disease symptoms compared to WL (Figure 5c). Next, we swapped the treatment of the leaves: we locally exposed the top leaflet of L4 to supplemental FR (Figure 5d) and quantified soluble sugars and disease symptoms in this local L4 (L4), as well as in the now older, distal L3 (L3). Upon local FR illumination on L4, we observed that both L3 and L4 itself exhibited higher soluble sugar content as well as increased disease symptoms compared to WL (Figure 5c). Next, we swapped the treatment of the leaves: we locally exposed the top leaflet of L4 to supplemental FR (Figure 5d) and quantified soluble sugars and disease symptoms in this local L4 (L4), as well as in the now older, distal L3 (L3). Upon local FR illumination on L4, we observed that both L3 and L4 itself exhibited higher soluble sugar content as well as increased disease symptoms compared to WL (Figure 5c). Next, we swapped the treatment of the leaves: we locally exposed the top leaflet of L4 to supplemental FR (Figure 5d) and quantified soluble sugars and disease symptoms in this local L4 (L4), as well as in the now older, distal L3 (L3). Upon local FR illumination on L4, we observed that both L3 and L4 itself exhibited higher soluble sugar content as well as increased disease symptoms compared to WL (Figure 5c).
the younger FR-illuminated leaf to the older non-FR-illuminated leaf could have occurred. However, we cannot exclude the presence of a mobile signal, other than sugars themselves, that would be transported from the illuminated leaf towards an older leaf and trigger sugar accumulation and enhanced pathogen growth in that location. Future studies, using for example radiolabeled CO\textsubscript{2} applied to the different leaf tips may help distinguish between these hypotheses.

3.5 The activation of JA signalling inhibits soluble sugar accumulation in tomato leaflets

Upon low R:FR perception in Arabidopsis or genetic inactivation of phyB in tomato, JA responsiveness is reduced (Cerrudo et al., 2017; Cortés et al., 2016). In addition, JA levels in plant tissue are negatively correlated to the soluble sugar levels in turn modulating disease progression in \textit{Nicotiana attenuata} plants (Machado et al., 2015). JA biosynthesis or perception mutants in tobacco displayed an increase in glucose and fructose levels but no changes in sucrose levels in leaf tissue compared to the wild-type (Machado et al., 2015) similar to what we observed for the \textit{phyB1phyB2} double mutant and WL + FR-treated tomato plants (Figure 1b and Figure 2a, respectively). To follow this up, we investigated whether the elevated soluble sugar levels observed in WL + FR-treated tomato leaves could be mediated via a JA-dependent mechanism. We exposed 3-week-old tomato plants either to WL or WL + FR and sprayed the plants daily with either a 50 μM methyl jasmonate (MeJA) or a mock solution for 5 days prior to quantifying the soluble sugar content in the third oldest leaf (Figure 6a). Under WL + FR conditions the leaflets contained a higher total sugar content compared to WL conditions in the mock treatment. However, exogenous MeJA treatment dramatically reduced the levels of all soluble sugars measured (Figure 6a). The soluble sugar depletion induced by exogenous MeJA treatment was also associated with increased foliar resistance in WL-treated plants but not in WL + FR-treated plants possibly because of a reduced JA responsiveness in the latter (Cortés et al., 2016; Figure 6d). In accordance, the JA biosynthesis mutant \textit{defenseless1} (def1) displayed constitutively elevated glucose levels as well as increased symptom severity.
upon *B. cinerea* infection, compared to its wild-type background Castlemart (Figure 6b,e).

These observations confirm a connection between JA and soluble sugar levels where JA-deficient plants accumulate more soluble sugars in leaf tissue. Moreover, again we notice a correlation between elevated soluble sugar levels and enhanced pathogen infection. These data suggest that JA may play a role in soluble sugar homeostasis in tomato leaves upon WL + FR perception. Next, we attempted to rescue the def1 mutation by providing these plants with exogenous MeJA (50 μM) every day for 5 days prior to quantifying soluble sugar levels or inoculating with *B. cinerea* (Figure 6c,f). As expected, we observed a clear decrease in soluble sugar levels in def1 mutant leaves when treated with exogenous MeJA compared to the mock (Figure 6c). Also, the exogenous addition of MeJA on def1 mutants could partially rescue plant resistance to *B. cinerea* (Figure 6f) showing the inhibitory effect of JA signalling on soluble sugars in regulating symptom development in tomato leaves. Interestingly, the susceptibility of def1 plants was significantly increased by the perception of supplemental FR even though the sugar levels remained unchanged (Figure 6b,e). This observation indicates that more factors than just changes in soluble sugar levels and/or JA-mediated defenses are affected by WL + FR. This, in turn could modulate either direct plant immune responses or pathogen growth rate within plant tissue leading to more severe disease symptoms on tomato leaves.

Based on these results, we hypothesize that the decreased JA responsiveness induced by WL + FR causes a soluble sugar accumulation which in turn promotes *B. cinerea* development in planta compared to WL conditions. Consistently, additional FR light had no effect on soluble sugar levels in the JA-deficient def1 mutant (Figure 6b,c). The latter experiment, however, should be interpreted cautiously since the wild-type background in which the def1 mutant occurs (cv Castlemart) had only slight changes in soluble sugar
concentrations between WL- and WL + FR-treated plants (Figure 6b; \( p = .056 \)). The negative correlation between JA and soluble sugar levels was already described by Machado et al. (2015). In addition, the JA biosynthesis mutant displayed an increased feeding performance by the chewing insect *Manduca sexta*. However, these caterpillars were shown to perform worse on high-sugar diets. This shows the opposite of what we describe here for WL + FR conditions, where elevated sugar levels occur together with elevated symptom development in tomato leaves by an array of pathogens (Figure 3). However, as the pathosystem and experimental conditions used by Machado et al. (2015) are hugely different from our study system it is not possible to predict whether the sugar-resistance relations in these studies contrast each other. Nevertheless, both studies describe a negative relation between JA and soluble sugar levels.

Although we found a negative effect of JA on soluble sugar levels, the disease phenotypes observed do not always perfectly fit the soluble sugar levels (Figure 6). This observation can be explained by the effect of WL + FR on other aspects of plant immunity such as defense hormone signalling or secondary metabolite biosynthesis. These plant immunity routes are regulated by JA, and have been previously shown to be regulated by supplemental far-red, both at the level of JA response (Chico et al., 2014; Leone et al., 2014; Pieterse et al., 2014) and JA synthesis (Fernández-Milmanda et al., 2020). As mentioned previously, glucose and JA signalling act synergistically to promote glucosinolate (GS) biosynthesis in response to *B. cinerea* in Arabidopsis (Buxdorf et al., 2013; Guo et al., 2013; Kliebenstein et al., 2005). Even though GS are specific to Brassicaceae, it would be interesting to determine which secondary metabolites are associated with tomato

**FIGURE 6**  Jasmonic acid signalling inhibits sugar accumulation in tomato associated with increased lesion development. (a–c) Leaf total soluble sugar levels including glucose (pink), fructose (yellow) and sucrose (blue) fractions as well as (d–f) lesion area induced by *B. cinerea* on detached leaflets at 3 dpi in (a and d) Moneymaker plants exposed for 5 days to WL or WL + FR conditions and treated daily with 50 μM MeJA or a mock solution, (b and e) in tomato cv. Castlemart plants and the JA biosynthesis mutants def1 (defenseless 1) exposed to WL or WL + FR conditions for 5 days or (c and f) in the def1 mutant treated for 5 days in WL and WL + FR and treated daily with a 50 μM MeJA or a mock solution. Different letters represent significant differences according to ANOVA, Tukey’s post-hoc test (\( p < .05 \)). Data represent mean ± SEM, \( n = 3-8 \) plants per treatment.
resistance to B. cinerea and how they would be influenced by either WL + FR perception or glucose supplementation in future studies.

4 | CONCLUSION AND PERSPECTIVES

Our results show that a decrease in JA responsiveness previously observed in tomato (Cortés et al., 2016) coupled with elevated soluble sugar levels promotes pathogen growth in plant tissue leading to increased lesion development. We demonstrate that supplemental FR, either applied to the whole plant or on a targeted leaflet, interferes with soluble sugar homeostasis in tomato leaves in a local and remote fashion thereby affecting pathogen growth capacity in plant tissue. We also show evidence about the role of JA in the regulation of soluble sugar levels upon WL + FR exposure. Although the soluble sugar status of the plants alone does not always predict the susceptibility phenotype upon infection by B. cinerea, we show that the modulation of soluble sugar levels in tomato leaves is mostly based on a JA-dependent mechanism and is overall associated with severity of disease symptoms.

In the context of global climate change and the simultaneously increasing global food demand, novel agricultural approaches are needed to increase crop yield under challenging environmental conditions. One option to do so is to grow plants closer together, whilst maintaining yield and resilience. However, we show that neighbour detection via a reduction in the R:FR ratio elevates foliar susceptibility to pathogenic microorganisms via a rapid increase in soluble sugar levels. Nevertheless, in greenhouse systems, applying supplemental light using LEDs is relatively straightforward. Assuming that supplemental red LEDs would be applied as inter-lighting to counteract the naturally occurring drop in R:FR due to the absorption of red light by the crop stand it might be possible to maintain plant resistance even at high density. In a dense plant canopy, the elevation of soluble sugars in WL + FR could possibly be controlled by increasing the R:FR ratio in the middle of the canopy in order to reduce the sugar-mediated susceptibility that occurs systemically in older, but not younger leaves than the FR-exposed ones (Figure 5). Higher in the canopy, there is still ample direct light from top lamps and outside to keep R:FR relatively high. Overall, increasing plant resistance at high density by using LED lighting seems possible and could partly solve the FR-induced susceptibility in tomato towards pathogens. Also, as sugar supplies are beneficial to most pathogens, we believe that balancing soluble sugar levels in planta by smart LED lighting plans would rescue tomato susceptibility to other pathogens than B. cinerea as well. Future studies on timing and spatial distribution of light quality all over a mature tomato stand are needed to understand if these positive effects of FR-enriched light can be achieved without having the negative effects on tomato pathogen resistance.

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CONFLICT OF INTEREST

The authors declare no conflict of interest.

AUTHOR CONTRIBUTIONS

Sarah Courbier and Ronald Pierik designed the study and experiments with additional input from Kaisa Kajala, Sanne Grevink, Emma Sluijs, Pierre-Olivier Bonhomme and Sarah Courbier carried out the experiments. Sarah Courbier drafted the article and designed the figures. Ronald Pierik and Sarah Courbier discussed results. Ronald Pierik, Sarah Courbier and Saskia C. M. Van Wees wrote the final manuscript, Kaisa Kajala commented on the final draft.

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