INTRODUCTION

In addition to their role as providers of carbon and energy, sugars fulfill a signaling role in coordination with hormonal signaling pathways (Rolland et al., 2006) controlling various plant physiological processes, probably also including innate immunity (Bolouri Moghaddam and Van den Ende, 2012). Distinct glucose, sucrose, and fructose signaling pathways can be discerned (Cho and Yoo, 2011; Li et al., 2011). These signaling pathways may be strongly influenced by the activities of sucrose splitting enzymes (vacuolar, cell wall and neutral invertases, sucrose synthase or SuSy; Koch, 2004) since they have strong impact on sucrose to hexose ratios, which might be an important parameter in plant responses, especially under stress (Xiang et al., 2011). It can be speculated that cellular sucrose to hexose ratios translate into certain levels of fructose-6-phosphate (F6P), an emerging regulatory molecule in plant growth and stress responses (Lann et al., 2006; Vandersteene et al., 2012; Wingler et al., 2012). T6P levels are likely controlled by the balance between its synthesis from UDPGlc and glucose 6-phosphate (G6P) by trehalose-6-phosphate synthase (TPS) and its breakdown by trehalose-6-phosphate phosphatase (TPP; Lann et al., 2006).

Next to growth and stress responses, it can be expected that sugar signaling is of great importance in flowering time control. This major developmental transition directly affects yield (DELLAs, PHYTOCHROME INTERACTING FACTORS, invertases, and trehalose-6-phosphate) might fulfill central roles in the transition to flowering as well. The emerging concept of “sweet immunity,” modulated by the clock, might at least partly rely on a sucrose-specific signaling pathway that needs further exploration.

Keywords: circadian clock, DELLA, flowering time, invertase, PIF , sugar signaling, sweet immunity, T6P

Sugars do not only act as source of energy, but they also act as signals in plants. This mini review summarizes the emerging links between sucrose-mediated signaling and the cellular networks involved in flowering time control and defense. Cross-talks with gibberellin and jasmonate signaling pathways are highlighted. The circadian clock fulfills a crucial role at the heart of cellular networks and the bilateral relation between sugar signaling and the clock is discussed. It is proposed that important factors controlling plant growth (DELLAs, PHYTOCHROME INTERACTING FACTORS, invertases, and trehalose-6-phosphate) might fulfill central roles in the transition to flowering as well. The emerging concept of “sweet immunity,” modulated by the clock, might at least partly rely on a sucrose-specific signaling pathway that needs further exploration.

CIRCADIAN REGULATION OF FLOWERING

In Arabidopsis, flowering can be autonomous or induced by gibberelins (GAs), as internal signals, or by the photoperiod and vernalization as external signals. FLOWERING LOCUS T (FT), SUPPRESSOR OF OVEREXPRESS OF CONSTANS 1 (SOC1), SQUAMOSA PROMOTER BINDING PROTEIN-LIKE (SPL) and LEAFY (LFY) transcription factors are among the best characterized floral pathway integrators, next to others (Matoukas et al., 2012; Yamaguchi and Abe, 2012). Both FT and SOC1 are inhibited by FLOWERING LOCUS C (FLC) in the autonomous and vernalization dependent pathways, while FT and SOC1 are activated by the photoperiodic protein CONSTANS (CO; Lee and Lee, 2010). During the day-night cycle, rhythmic expression of the core circadian clock components, CCA1, LHY, and TOC1 control the expression of GL, an activator of CO (Murphy et al., 2011).

The induction of flowering through the transport of phloem-mobile signals (FT and GA) to the apex is well-documented (Corbesier et al., 2007; Tamaki et al., 2007; Turnhull, 2011; Matoukas et al., 2012; Yu et al., 2012). It is known since long that phloem-mobile sucrose may represent an additional critical factor in controlling the transition to flowering (Corbesier et al., 1998; Roldan et al., 1999; Ohno et al., 2001). This would represent another function for sucrose next to its known roles in many other plant regulatory and signaling mechanisms including growth, development, and stress-related responses (Wind et al., 2010).
PLACING FLOWERING INTO THE BIGGER PICTURE: CENTRAL ROLES FOR DELLAS AND miRNAs

DELLA proteins are crucial players in GA signaling pathways involved in plant growth control (Harberd et al., 2009; Figure 1). GA inhibits DELLA protein levels, which are inhibitors of PHYTOCHROME INTERACTING FACTORS (PIFs), acting as growth enhancers (Nouze et al., 2011; Stewart et al., 2011; Figure 1). Recently, miR156 and miR172 were found as important factors controlling plant developmental transitions (Yamaguchi and Abe, 2012). It was found that miR156 acts as a negative regulator of SPL gene expression. SPLs stimulate LFY and MAD-box genes (Borner et al., 2000; Vekemans et al., 2012) and the production of miR172, which in turn stimulates reproductive competency and flowering through its inhibitory action on APETALA2 (AP2), TARGET OF EAT1 (TOE), SCHLAMMUTZEN (SNZ), and SCHNARCHZAPEN (SNZ), inhibitors of FT (Zhu and Hellwigg, 2010; Yamaguchi and Abe, 2012; Figure 1). miR172 is also under control of the clock by GI (Jung et al., 2007; Figure 1). The missing mechanistic link between GA signaling and flowering was recently established, by defining a role for DELLA as a general flowering inhibitor. DELLA inhibits SPL gene expression and miR172 production (Galvao et al., 2012; Yu et al., 2012; Figure 1). Therefore, DELLA proteins are now considered both as growth and flowering inhibitors. Accordingly, transgenic plants overexpressing DELLA proteins or plants expressing mutant DELLA repressors show dwarfism and delayed flowering (Dill et al., 2004; Hamama et al., 2012). What is more, at lower GA levels, some DELLA proteins were found to act as strong activators of the jasmonate (JA) signaling pathway (Wild et al., 2012), a major pathway controlling plant defense responses (Yang et al., 2012; Figure 1). It can be concluded that DELLAs occupy a central and crucial position in plant growth, development and flowering as well as in stress responses (Figure 1).

HOW DO SUGAR SIGNALS INTERACT WITH FLOWERING NETWORKS?

The relation between sugar metabolism/signaling and floral transition received extensive attention lately (Turnbull, 2011; King, 2012). The study of Heyer et al. (2004) already provided clear evidence that flowering time control is strongly influenced by modifying sugar balances in the apex. They placed yeast invertase under the control of a meristem-specific promoter and compared apoplastic and cytosolic localized invertase versions. Intriguingly, transition to flowering was hastened by the expression of the invertase in the cell wall, while a flowering delay was observed when the invertase was localized in the cytosol. This indicated that invertases with a different localization might fulfill a crucial role in transition to flowering. It was recently proposed that high sucrose levels are associated with high T6P levels (Winger et al., 2012), but it should be noted that this correlation depends on the activity of sucrose splitting enzymes, such as invertases (Figure 1). Although the molecular mechanism for the control of transition to flowering by sugars remains to be further investigated, a possible scenario is that T6P acts as a positive mediator of some PIF isoforms (Figure 1). Indeed, it was reported that PIF5 overexpression leads to early flowering, both under long day and short day conditions (Nouze et al., 2011), strongly suggesting that PIFs might be linked to floral transition. Moreover, in hypocotyl elongation studies it was found that sucrose stimulates several PIF isoforms, even in the dark (Liu et al., 2011; Stewart et al., 2011; Lilley et al., 2012; Sairanen et al., 2012). Therefore, similar to DELLA proteins which were only recently recognized as important players in flowering time control, it can be expected that some PIF isoforms may be involved as well. However, it remains to be demonstrated whether such underlying PIF-mediated mechanisms account for the stimulation of FT gene expression by sucrose (King et al., 2008; Figure 1). Also, it would be interesting to investigate the mechanisms involved in the sucrose-mediated upregulation of the LFY gene (Matsoukas et al., 2012; Figure 1). Another emerging link between flowering and sucrose metabolism/transport was reported by Sos et al. (2011). These authors demonstrated that the indeterminate domain 8 (IDD8) transcription factor plays a role in FT-dependent flowering induction, via modulation of the SULF4 activity. Further, Conervi et al. (2012) reported that the starch to sucrose transition is important during autonomous flowering. In conclusion, sucrose seems to interact in many ways with the flowering network, and further studies are needed to fully understand these connections at the molecular level.

HOW DOES SUGAR SIGNALING INTERACT WITH THE CLOCK?

On the one hand, it was recently reported that the clock’s core central oscillator genes GI, TOC1, and CCA1 are stimulated by sucrose (Knight et al., 2008; Dalchau et al., 2011), suggesting that the clock is entrained by metabolic signals such as sugars, possibly independent from phytochrome-mediated light perception. On the other hand, it has been demonstrated that the enzymatic activity and expression of a vacuolar invertase gene in petioles of sugar beet follows a circadian rhythm (González et al., 2005). Furthermore, it is well-known that vacuolar invertases are stimulated by GA (González and Cejudo, 2007; Choubane et al., 2012). A function of vacuolar invertases as stimulators of stomatal opening has recently been suggested (Antunes et al., 2012; Nt, 2012), in addition to their well-described role in cellular elongation processes (Wang et al., 2010). These observations fit well with the overall idea that invertases fulfill a central (Figure 1) and crucial role coordinating carbon dioxide uptake, photosynthesis, and plant growth through GA- and sugar-mediated signaling pathways, with a clear connection to the flowering time control network (Figure 1), as explained above. Intriguingly, expression of FT in guard cells also promoted stomatal opening (Kinoshiba et al., 2011). This suggests a role for FT in stimulating carbon dioxide uptake and fixation, to produce the necessary carbon skeletons that are required for the flowering process.

Noteworthy, the cell wall invertase LINS of tomato, induced by JA signaling and considered as a pivotal enzyme for the integration of various signals, is also regulated by a diurnal rhythm (Proels and Roitsch, 2009). Intriguingly, the rhythms of the
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FIGURE 1 | Emerging sugar signaling connections to cellular networks involved in plant growth, defense, and floral transition. Simplified schematic presentation of a selection of crucial players in plant growth, flowering transition, and defense responses, and their interactions. Metabolites are not boxed; proteins are in colored boxes. Arrows (→) signify stimulation, while an inhibitory interaction is presented by the ◐ symbol. Red arrows refer to the putative effect of the Suc/INV/T6P module. Full arrows indicate established relationships. Dotted arrows indicate established relationships that are in need for further exploration (unraveling of mechanistic details). Dashed arrows represent rather speculative connections that remain to be confirmed. Straightforward symbols are used for light and the clock. Stomatal opening is also schematically presented. APA2, APETALA2; CO, CONSTANS; DELLA, DELLA protein; FT, Flowering locus T; GA, gibberellin; INV, invertase (or any other sucrose splitting enzyme); JA, jasmonate; LFY, LEAFY; miR156, micro RNA 156; miR172, micro RNA 172; NO, nitric oxide; PIF, Phytochrome Interacting factor; SPL, Squamosa Promoter Binding Protein-like; Suc, sucrose; TOE, TARGET OF EAT1; SMZ, SCHLAFMÜTZE; SNZ, SCHNARCHZAPEN. For more details on floral transition networks, readers are referred to Matsoukas et al. (2012) and Yu et al. (2012), where apex and leaf-located processes are discerned.

two above-mentioned vacuolar and cell wall invertases were not synchronized, perhaps reflecting differential diurnal patterns in growth dynamics. These and other observations strongly suggest that there is an intimate interplay and reciprocal relationship between sugar metabolism/signaling and the plant circadian clock. So, besides light as the most important stimulus influencing the clock’s components through phytochromes and cryptochromes, endogenous sugar signals, hormones, and stresses also entrain the clock (Arana et al., 2011; Facella et al., 2012; Goodspeed et al., 2012;
Seung et al., 2012). Vice-versa, the clock is involved in regulating the biosynthesis of GA (Blázquez et al., 2002) and JA (Shin et al., 2012), suggesting that the clock shows putative bilateral relationships with these hormones as well (Figure 1).

**SUGAR SIGNALING IN RHYTHMIC IMMUNITY**

 Sugars as signaling molecules are well-known activators of various pattern-recognition receptor genes (Johnson and Ryan, 1990; Herbers et al., 1996a,b). There is mounting evidence that, in addition to plant cell wall or fungal-derived oligosaccharides, also sugars such as sucrose could be involved in plant priming and innate immunity responses (Gomez-Ariza et al., 2007; Birch et al., 2009; Bolouri Moghaddam and Van den Ende, 2012; Sonnewald et al., 2012). One of the best studied pathways in plant defense responses is the sucrose-specific signaling pathway that leads to the production of anthocyanins (Teng et al., 2005; Soliniec et al., 2006). Despite huge research efforts, a number of (transcription) factors involved in this pathway remain to be identified. Possibly T6P is involved (Wingerl et al., 2012), but the underlying mechanisms need further investigation. During sucrose-mediated signaling, Ca2+ seems to be involved as well, probably by stimulating sucrose uptake into the cell (Shin et al., 2013). Both light and most plant hormones influence this pathway (reviewed in Das et al., 2012). Nitric oxide (NO) and pathogen-derived elicitors are also able to trigger the production of anthocyanins (Tosi et al., 2011; Cai et al., 2012; Figure 1).

Strikingly, transgenic plants expressing a mammalian NO synthase become disease resistant to a broad array of plant pathogens (Chun et al., 2011) highlighting the importance of NO signaling. Perhaps NO is an integral part of the sucrose-specific pathway leading to anthocyanin production, and this is an interesting area of further investigation. Furthermore, NO was found to regulate DELLA contents and PIF expression (Lozano-Juste and Leon, 2011). The effect of sugar signaling on plant immunity may, at least partly, depend on the expression and activation of kinases such as the mitogen-activated protein kinases (MAPKs). It has been recently demonstrated that sucrose can rapidly activate Ca2+ and CaMK, a unique sucrose-specific MAPK from *Cephalotachylum fuchsiaeum* (Li et al., 2012), but it is unknown whether other plants (such as *Arabidopsis*) also have such sucrose-specific MAPKs. It has been shown that many plant factors involved in plant immune responses are regulated by the clock (Farias et al., 2010; Bhandari et al., 2012; Wang et al., 2011). Plants probably evolved this type of regulation to maximize levels of defense compounds (toxins, defense hormones) and/or sweet immunostimulators at those moments of the day when the encounter with the pathogen/herbivore is more likely to occur. For instance, JA levels show a diurnal oscillation that is synchronized with insect feeding behavior (Goodspeed et al., 2012). Similarly, a clock-controlled variation in resistance to the virulent bacterial pathogen *Pseudomonas syringae* pv. *tomato* DC3000 (Pst DC3000) was discovered in *Arabidopsis* (Bhandari et al., 2011).

Interestingly, flowering time control and defense signaling pathways in plants seem to have points of convergence too (Liu et al., 2012). The rice spotted leaf 11 mutant shows an enhanced resistance to *Magnaporthe grisea* and *Xanthomonas oryzae* pv. *oryzae* (Yin et al., 2000; Liu et al., 2012; Marini et al., 2012) and the spotted leaf 11 gene expression is induced by both incompatible and compatible rice-blast interactions (Zeng et al., 2004). Moreover, the spotted leaf 11 protein is involved in flowering time regulation in rice (Liu et al., 2012). This dual role in control of flowering time and defense has also been demonstrated for the *Arabidopsis* ortholog Plant U-box 13, but the molecular mechanisms involved and the possible links with sugar signaling events remain unclear.

**CONCLUSION**

Many aspects of plant growth, development, floral transition, and defense responses are regulated by circadian rhythms as well as by sugar signaling events. This mini review focused on the emerging links between sugar signaling, the clock, floral transition, and immune responses. Overall, GA and JA signaling pathways greatly determine plant growth versus defense responses, with DELLA and PIF proteins as central players. The recent finding that some DELLA proteins are also key players in floral transition urges further research on the possible involvement of PIFs in floral transition processes, since some data suggest that PIF expression may be under direct control by sugar signals, perhaps mediated by T6P. Invertases may be important to control T6P levels, taking a central position in these networks. Furthermore, putative new roles are emerging for invertases (e.g., stomatal opening).

It is also proposed that efficient defense responses might not only rely on hormones and on cell wall or pathogen-derived saccharides, but perhaps also on sucrose, through a sucrose-specific signaling pathway, perhaps (partly) resembling (or overlapping) with the sucrose-mediated pathway controlling anthocyanin biosynthesis in *Arabidopsis*. However, the putative new role of CA7 acting in this pathway remains to be identified, as well as the (transcription) factors involved in the upper part of the pathway, and this remains a challenging task.

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