ORIGINAL RESEARCH



Identification of new potential downstream transcriptional targets of the strigolactone pathway including glucosinolate biosynthesis

Alicia M. Hellens^{1,2} | Tinashe G. Chabikwa^{1,3†} | Franziska Fichtner^{1,2,4} | Philip B. Brewer^{1,2,5} | Christine A. Beveridge^{1,2} |

Correspondence

Alicia M. Hellens and Christine A. Beveridge, School of Biological Sciences, University of Queensland, St. Lucia, QLD 4072, Australia. Email: a.hellens@uq.net.au and c.beveridge@ uq.edu.au

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Abstract

Strigolactones regulate shoot branching and many aspects of plant growth, development, and allelopathy. Strigolactones are often discussed alongside auxin because they work together to inhibit shoot branching. However, the roles and mechanisms of strigolactones and how they act independently of auxin are still elusive. Additionally, there is still much in general to be discovered about the network of molecular regulators and their interactions in response to strigolactones. Here, we conducted an experiment in Arabidopsis with physiological treatments and strigolactone mutants to determine transcriptional pathways associated with strigolactones. The three physiological treatments included shoot tip removal with and without auxin treatment and treatment of intact plants with the auxin transport inhibitor, N-1-naphthylphthalamic acid (NPA). We identified the glucosinolate biosynthesis pathway as being upregulated across strigolactone mutants indicating strigolactoneglucosinolate crosstalk. Additionally, strigolactone application cannot restore the highly branched phenotype observed in glucosinolate biosynthesis mutants, placing glucosinolate biosynthesis downstream of strigolactone biosynthesis. Oxidative stress genes were enriched across the experiment suggesting that this process is mediated through multiple hormones. Here, we also provide evidence supporting non-auxin-mediated, negative feedback on strigolactone biosynthesis. Increases in strigolactone biosynthesis gene expression seen in strigolactone mutants could not be fully restored by auxin. By contrast, auxin could fully restore auxin-responsive gene expression increases, but not sugar signaling-related gene expression. Our data also point to alternative roles of the strigolactone biosynthesis genes and potential new signaling functions of strigolactone precursors.

In this study, we identify a strigolactone-specific regulation of glucosinolate biosynthesis genes indicating that the two are linked and may work together in regulating

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¹School of Biological Sciences, University of Queensland, St. Lucia, Queensland, Australia

²ARC Centre for Plant Success in Nature and Agriculture, The University of Queensland, St Lucia, Queensland, Australia

³QIMR Berghofer Medical Research Institute, Brisbane, Queensland, Australia

⁴Institute for Plant Biochemistry, Heinrich Heine University, Düsseldorf, Germany

⁵School of Agriculture, Food and Wine, The University of Adelaide, Glen Osmond, South Australia, Australia

[†]The author died prior to the submission of this paper.

stress and shoot branching responses in Arabidopsis. Additionally, we provide evidence for non-auxin-mediated feedback on strigolactone biosynthesis and discuss this in the context of sugar signaling.

KEYWORDS

Arabidopsis thaliana, auxin, flavonoids, glucosinolates, strigolactones, transcriptomic

1 | INTRODUCTION

Strigolactones (SLs) are a group of recently discovered plant hormones that play a role in many aspects of plant growth and development (reviewed in Brewer et al., 2013). SLs were first associated with, and subsequently named after, the parasitic weed of genus Strigg or witchweed (Cook et al., 1972). Exogenous SLs exuded from plant roots in poor nutrient conditions induce Striga germination and provide beneficial symbiotic interactions with arbuscular mycorrhizal (AM) fungi (Bouwmeester et al., 2007). In 2008, SLs were identified as plant hormones that have a role in signaling, particularly in the inhibition of shoot branching or axillary bud outgrowth (Gomez-Roldan et al., 2008; Umehara et al., 2008). SLs have since been intricately studied and are clearly an important aspect of plant growth and development because they act in networks with most other plant hormones and are impacted by nutrient and signaling pathways (recent reviews: Barbier et al., 2019; Bürger & Chory, 2020; Faizan et al., 2020; Kelly et al., 2023; Saeed et al., 2017).

In the past decade or so, much of the SL biosynthetic and signaling pathway has been elucidated (reviewed in Yoneyama & Brewer, 2021). In Arabidopsis, bioactive SLs are synthesized from β-carotene through consecutive reactions catalyzed by β-carotene DWARF27 (D27): CAROTENOID DIOXYGENASE7 and 8 (CCD7 and CCD8), encoded by MORE AXILLARY GROWTH3 and 4 (MAX3 and MAX4) genes, respectively; MAX1, which encodes a cytochrome P450 CYP711A1; a carlactonoic acid methyltransferase (CLAMT); and LATERAL BRANCHING OXIDO-REDUCTASE (LBO) (Abe et al., 2014; Alder et al., 2012; Brewer et al., 2016; Mashiguchi et al., 2022; Seto et al., 2014). More recently, SL has been demonstrated to be catabolized by carboxylesterase enzyme, CXE15 (Xu et al., 2021). SL perception and signaling involve the F-box protein MORE AXILLARY GROWTH2 (MAX2) and the α/β -fold hydrolase DWARF14 (D14), which form a complex that binds SUPPRESSOR OF MAX2 1-LIKE target proteins SMXL6, SMXL7, and SMXL8 (also known as DWARF53 [D53]) for degradation to release downstream genes from inhibition and allowing SL responsiveness (Arite et al., 2009; Chevalier et al., 2014; Ishikawa et al., 2005; Soundappan et al., 2015; Stirnberg et al., 2007; Wang et al., 2015). MAX2 is also involved in karrikin signaling (Nelson et al., 2011).

Genes that show similar expression patterns, or are co-expressed, have the potential to encode proteins with a related function, or which participate in similar biological processes (Eisen et al., 1998; Marcotte et al., 1999). Some aspects of the SL pathway are co-regulated, such as SL biosynthesis genes MAX3 and MAX4. This knowledge allowed for the discovery of other SL biosynthesis

components *LBO* and the Arabidopsis ortholog of rice *D27*, as they were both found to be upregulated in other SL mutants in a similar manner to the previously confirmed SL biosynthesis genes *MAX3* and *MAX4* (Brewer et al., 2016; Waters, Brewer, et al., 2012). After identifying the co-regulated expression patterns, mutants for *lbo* and *d27* were identified and shown to have enhanced branching and ultimately to act on the SL pathway. Given the success of this approach in identifying new SL biosynthesis genes (Brewer et al., 2016; Waters, Brewer, et al., 2012), we postulate that additional aspects of the SL regulatory network can be deciphered by interpreting co-regulated genes.

Increased SL biosynthesis gene expression in SL-deficient mutants is indicative of a negative feedback loop that has long been associated with the plant hormone auxin (Hayward et al., 2009; Ligerot et al., 2017). Auxin is one of the most well-researched plant hormones. Auxin promotes SL biosynthesis gene expression (Yoneyama & Brewer, 2021) and, in turn, SL inhibits auxin biosynthesis, transport, and canalization (Crawford et al., 2010; Ligerot et al., 2017; Shinohara et al., 2013; Zhang et al., 2020). Apically derived auxin upregulates the SL biosynthesis genes MAX3, MAX4, LBO, and D27 (Bennett et al., 2006; Brewer et al., 2016; Hayward et al., 2009; Shen et al., 2012; Waters, Brewer, et al., 2012). MAX2- and D14-dependent SL perception, in turn, inhibits auxin transport (Bennett et al., 2006; Crawford et al., 2010; Ligerot et al., 2017). In the SL-deficient mutants, max3 and max4, MAX2-mediated inhibition of auxin transport does not occur because of lack of SL signal perception and transduction. High endogenous levels of auxin in SL mutants (Beveridge, 2000) indicate that auxin may contribute to negative feedback, resulting in increased SL biosynthesis gene expression in SL-deficient mutants (Brewer et al., 2009; Foo et al., 2005; Hayward et al., 2009; Waters, Brewer, et al., 2012). However, the magnitude of upregulation of MAX3 and MAX4 gene expression in response to exogenous auxin is much less than the increase observed in SL mutants (Bennett et al., 2006; Hayward et al., 2009). This indicates that some other factors may be involved. Additionally, auxin is not the only contributor to SL-mediated responses, and non-auxin signals also regulate aspects of plant development (Bainbridge et al., 2005; Barbier et al., 2019; Faizan et al., 2020). We therefore hypothesize that there is also a non-auxin-mediated feedback on SL biosynthesis.

Sucrose has recently re-emerged as a major player in apical dominance (Barbier, Lunn, & Beveridge, 2015; Mason et al., 2014) and may act via a number of signaling pathways. Trehalose 6-phosphate (T6P) is a sucrose-specific signaling molecule, with T6P levels being strongly correlated with sucrose levels in plants (Fichtner & Lunn, 2021). T6P levels strongly correlate with bud outgrowth in pea (Fichtner

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et al., 2017). Additionally, transgenic Arabidopsis plants with increased levels of T6P show increased branching (Fichtner et al., 2021), implicating T6P in the regulation of lateral bud outgrowth. Sucrose can be broken down into glucose and fructose. The enzyme HEXOKINASE (HXK) is the first enzyme in glycolysis that adds a phosphate to glucose to form glucose 6-phosphate. In Arabidopsis, the HXK1-deficient mutant, *gin2*, has reduced rosette branching and also shows increased *MAX2* expression (Barbier et al., 2021). Interestingly, sucrose antagonizes the SL-induced repression of shoot branching (or tillering) in rice, partially by inhibiting *OsMAX2* gene expression and by antagonizing the SL-mediated degradation of D53 (Bertheloot et al., 2020; Patil et al., 2021).

To determine novel processes co-regulated with SLs, and to investigate auxin-related feedback, we conducted a large-scale, genome-wide gene expression study. This study was previously used to discover LBO, which is co-expressed with MAX3 and MAX4 (Brewer et al., 2016). We revisited this prior data set to extrapolate more information. Recent progress in transcriptome analysis has investigated the response to SL, and SL mutants separately (Kumar et al., 2019: Wang et al., 2020). Our study compared SL-modifying physiological treatments with SL mutants. Mutant genotypes combined with physiological treatments in wild type (WT) allowed us to determine processes that are differentially regulated and co-regulated by SL independent of auxin. We identified differentially expressed and co-expressed genes in SL biosynthesis (max1, max3) and signaling (max2), mutants, as well as in WT Arabidopsis plants in response to decapitation with, and without the addition of auxin, indole-3-acetic acid (IAA), and with auxin transport inhibitor, N-1-naphthylphthalamic acid (NPA). Decapitation removes an auxin source but also removes a sink and therefore increases available sucrose (Kebrom, 2017). All mutants and treatments were compared with intact WT plants. By manipulating the physiological network in this way, we were able to classify different groups of co-expressed genes, providing a solid basis to discover new research directions for understanding the networks involved in SL and auxin regulation.

2 | RESULTS

2.1 | Overview and functional enrichment

Fully elongated hypocotyls were used as the tissue sample in this study of signals regulating SL biosynthesis genes, because grafting experiments have demonstrated that WT hypocotyls can inhibit branching in SL biosynthesis mutant plants (Bainbridge et al., 2005; Beveridge, 2000; Hayward et al., 2009). Gene expression was profiled from the hypocotyls of 2-week-old Arabidopsis WT (1) intact (WT_Intact), (2) decapitated (WT_Decap), (3) decapitated and treated apically with auxin (WT_Decap_IAA) and plants (4) treated with the auxin transport inhibitor NPA (WT_NPA), as well as of mutants in SL biosynthesis and signaling genes: (5) max1, (6) max2, (7) and max3. Cell expansion and growth have ceased in hypocotyls by the treatment time of 2 weeks, making this tissue even more attractive to look at gene regulation caused by genotype and treatment

(Gendreau et al., 1997). The mutants and treatments all affect the SL pathway in some way. *max1* and *max3* are deficient in SLs, whereas *max2* cannot perceive SLs. Decapitated and NPA-treated samples inhibit auxin delivery to the hypocotyl, thus affecting SL biosynthesis gene expression (Brewer et al., 2009; Hayward et al., 2009). Decapitation + IAA treatment replaces the loss of auxin in decapitated samples and so helps determine processes that are not solely auxin regulated.

The physiological treatments had a relatively strong effect on the transcriptome. As expected, principal component analysis (PCA) analysis of the normalized intensity values showed clustering of the replicates of each treatment/mutant. The first component (57.07%) discriminated between the plants according to physiological treatments (Figures 1a and S1). The second component (11.7%) separated SL mutants from WT Intact and WT Decap IAA. As might be expected, all SL mutants were clustered together and WT Decap is close to WT_NPA on the graph, indicating overlapping transcriptional responses (Figures 1a and S1). Results of differentially expressed gene (DEG) analysis relative to the WT Intact are shown in Figure 1b and Table S1. Genes were deemed "differentially expressed" by absolute LogFC > 2 (|LogFC≥2|) compared with WT expression values and using an adjusted P-value (adj P-value) cut-off of .05 to assign statistical significance. The highest number of DEGs was observed in the WT_Decap (1097) and WT_NPA (1111) treatments, which indicates dynamic transcriptional responses for these treatments. Applying auxin after decapitation decreased the number of DEGs compared with WT_Decap (292). Many DEGs are shared among the three max mutants, and differences between them are discussed below. The combined DEGs of the three max mutants (246 combined) were fewer than any of the individual physiological treatments (Figure 1b).

Affymetrix probe IDs of the DEGs were used for functional enrichment of biological processes based on gene ontology (GO) (Ashburner et al., 2000) using the Database for Annotation, Visualization and Integrated Discovery (DAVID) v6.8 (Huang et al., 2009). Functional enrichment analysis showed that many biological processes were differentially regulated among treatments/mutants forming a distinction that mimics the clustering pattern observed by PCA analysis (Figure 1a). There was commonality in the enriched biological processes (GO terms) from DEGs of the three max mutants compared with WT (Figure 1c). WT_Decap and WT NPA also share many common GO terms, apart from the response to auxin group, which was identified for all WT treatments, but not the mutants. The WT_Decap_IAA GO terms were largely unique compared with other WT samples. Some details of these differences are described below together with a co-expression analysis.

2.2 | Co-expression analysis reveals three distinct clusters pertaining to SL biosynthesis and signaling

Gene regulatory networks are used to characterize the correlation patterns among genes. Groups of genes that are correlated in their

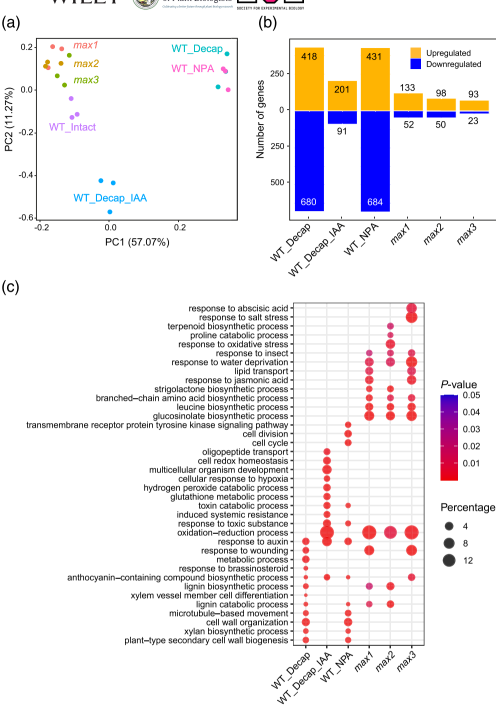


FIGURE 1 Whole-genome differential gene expression analysis. (a) Clusters of samples based on similarity determined by principal component analysis (PCA) show max mutants clustering together and close to wild-type intact (WT Intact) and separate from two other clusters that contain decapitated (WT Decap) and N-1-naphthylphthalamic acid (NPA)-treated (WT NPA) plants separate from decapitation + indole-3-acetic acid (IAA) (WT_Decap_IAA)-treated plants. (b) Total number of differentially expressed genes (DEGs) when compared with WT Intact. Genes with a LogFC of two and an adjusted P-value of <.05 were considered differentially expressed. WT Decap- and WT NPA-treated samples have the largest number of DEGs. The three strigolactone mutants show a much smaller transcriptional response. (c) Gene ontology (GO) categories enriched from DEGs identified in (b). GO terms determined using Database for Annotation, Visualization and Integrated Discovery (DAVID) online database for GO enrichment analysis (Dennis et al., 2003) with a P-value of <.05 are considered significantly enriched. Percentage indicates the number of genes differentially expresses in that GO term as a percentage of total genes in that GO term for Arabidopsis.

pattern of expression cluster to form modules that can often be linked to biological functions, such as hormonal biosynthesis. Weighted gene co-expression network analysis (WGCNA) was used to construct gene networks and detect modules of genes (Langfelder & Horvath, 2008). We obtained a network with scale-free topology composed of 26 modules of gene expression correlated with mutants and/or physiological treatments (Figures 2 and S2). WGCNA assigned to each module a unique color label that was used as a specific module identifier in the analyses. The modules were composed, on average, of 437 genes (median gene number per module, 159). The "gray" (or improper) module contains 346 probe sets that were not allocated in any of the 26 modules.

We identified three modules of interest relevant to the SL pathway and which indicated separation of co-expression among genes of the SL pathway. These modules each contained a MAX gene described henceforth as module type genes: MAX3 in brown (617 genes), MAX1 in blue (1921 genes), and MAX2 in turquoise (4238 genes) (Figure 2 and Table S2). Genes in the MAX3 module broadly showed an increase in expression in the SL mutants compared with WT. The MAX1 and MAX2 modules differed compared with the MAX3, showing a relatively modest change in expression in SL mutants, with a clear response to NPA and to decapitation, which was restored with auxin. The three SL-type genes and their neighbors (strongly correlated genes; P < .05) were extracted into

max2

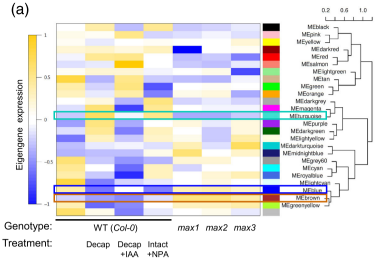
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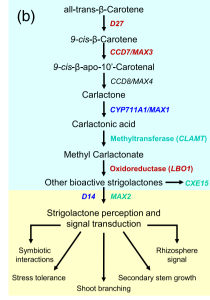


FIGURE 2 Weighted gene co-expression network analysis separates strigolactone (SL) genes into different co-expression groups. (a) Heat map and dendrogram of 26 identified groups of genes based on the pattern of gene co-expression across the entire dataset. Three groups were identified to contain SL genes of interest: MAX1 (blue), MAX2 (turquoise), and MAX3 (brown) (Table S2). (b) Diagram of SL biological pathway. The color of the text of the genes indicates the co-expression group to which they belong.

groups, and the data were used to create a GO term network using the ClueGO extension in cytoscape for each module (Figure S3 and

The MAX3-like group was extracted from the brown module (224 out of the 617 genes). It consists of SL biosynthesis genes MAX3, MAX4, D27, and LBO (Figures 2 and 3) and is also enriched in genes involved in glucosinolate (GSL) biosynthesis pathway (Figure S3) as discussed later. The MAX3-like group is categorized by a pattern of differential expression in the max mutants.

The MAX1-like group was extracted from the blue module (736 out of the 1921 genes) and includes MAX1 and the SL signaling gene D14 but also includes genes involved in T6P synthesis, TREHALOSE 6-PHOSPHATE SYNTHASE 1 (TPS1), and dephosphorylation, like TREHALOSE 6-PHOSPHATE PHOSPHATASE E (TPPE) (Figure 3b). This group is characterized by having genes with differential expression in decapitated and NPA-treated samples. Although MAX1 and D14 are upregulated in these samples, TPS1 and TPPE are downregulated; both correlated and anti-correlated genes are included (Figure 3b).

The MAX2-like group, which was extracted from the turquoise module (1341 out of the 4238 genes), contains MAX2 and is enriched in genes involved in auxin and light signaling (Figures 3b and S3). The recently discovered SL catabolism gene CARBOXYLES-TERASE 15 (CXE15) and the carlactonic methyltransferase (CLAMT) both belong in the turquoise co-expression module with MAX2 (Mashiguchi et al., 2022; Wakabayashi et al., 2021; Xu et al., 2021) (Figure 3b and Table S2). However, neither genes were strictly co-expressed with MAX2 and so are not included in the MAX2-like group.

Treatments with WT plants highlight 2.3 auxin-responsive genes and anthocyanin pathway genes

As expected, DEGs in common from WT_Decap, WT_Decap_IAA, WT NPA plants are enriched in the auxin-activated signaling pathway category (Figures 1a and 4a). Genes involved in anthocyanin-containing compound biosynthesis were enriched in WT_Decap, WT_Decap_IAA, and WT_NPA plants (Figures 1c and 4). Anthocyanin biosynthesis genes such as UDP-GLUCOSE:FLAVO-NOID 3-O-GLUCOSYLTRANSFERASE (UF3GT), MYB75/PRODUCTION OF ANTHOCYANIN PIGMENT 1 (PAP1), and DIHYDROFLAVONOL 4-REDUCTASE (DFR) are largely repressed in these treated WT plants compared with untreated WT (Figure 4b). Among the genes that showed common expression among all the treated WT plants (decapitation with or without IAA and intact with NPA), compared with intact WT, were 75 genes also differentially expressed between WT and SL mutants. Some of these were in the anthocyanin group although not suppressed in the mutants to the extent in WT treated plants (Figure 4b). Previous studies have shown effects of sugars and auxin on anthocyanin content (Ozeki & Komamine, 1986; Teng et al., 2005). Additionally, MAX2 was suggested to impact anthocyanin content, as max2 mutants have downregulated anthocyanin biosynthesis gene expression (Richmond et al., 2022). This potential regulation of anthocyanins has not been widely repeated and is yet to be tested by direct measurement of anthocyanin levels, or related to the high endogenous auxin or auxin signaling in SL mutants. Therefore, we measured anthocyanin content in whole rosettes of WT (Col-0), max2,

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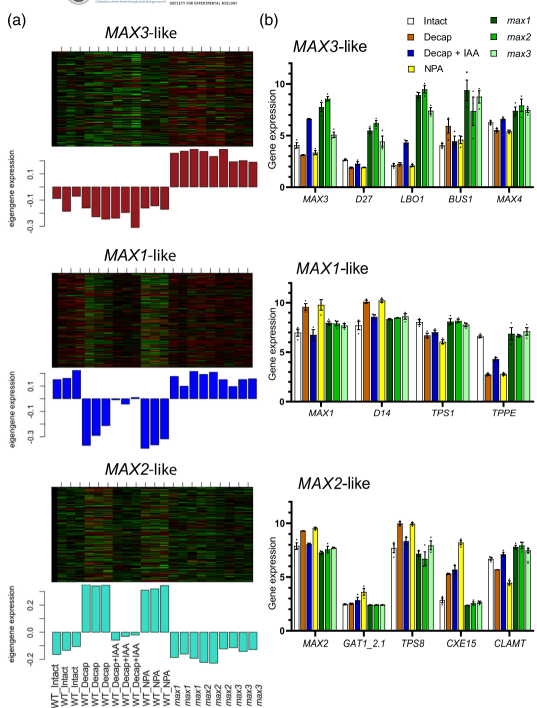


FIGURE 3 Functional analysis of strigolactone (SL)-like co-expression groups. (a) Eigengene expression in individual replicates for MAX3-like, MAX1-like, and MAX2-like co-expression groups identified in Figure 2. (b) Absolute gene expression as raw robust multichip average values of representative genes from each MAX-like co-expression group. Data presented as mean \pm SEM (n=3).

d14, and in the auxin-overproducing line 35S:YUCCA1 (Zhao et al., 2001). High auxin content plants, 35S:YUCCA1, did in fact have the highest accumulation of anthocyanins, and max2 had significantly lower anthocyanin accumulation than both WT and 35S: YUCCA1 plants (Figure 4c).

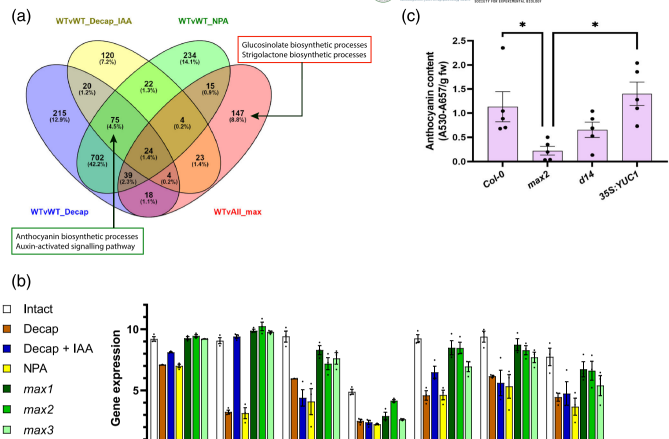
2.4 | SLs may be strongly associated with GSLs in Arabidopsis

We sought to identify and characterize DEGs common among the max mutants used in this study. Thirty-seven genes were common

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PAP1

UDPGT

DFR

DEGs among all *max* mutants (Figure 5a). These genes are significantly enriched for GSL biosynthetic processes (*P*-value: 7.50e⁻⁶) (Figures 5b and S4 and Table S3). Further investigation determined that all these GSL genes belonged to the process of, specifically, aliphatic GSL (aGSL) biosynthesis. Some aGSLs are derived from branched chain amino acids, and the biosynthesis of these branched chain amino acids was also a significantly enriched GO term for all the *max* mutants (Figure 1c). Over 40% of all aGSL biosynthesis genes are upregulated in the SL mutants (Figures 5c and S4). For example, GSL biosynthesis genes MYB29, CYTOCHROME P450 79F1/BUSHY 1 (CYP79F1/BUS1) (henceforth referred to as BUS1), ISOPROPYLMALATE ISOMERASE 1 and 2 (IPMI1 and IPMI2), and BILE ACID TRANSPORTER 5 (BAT5) are upregulated in all three *max* mutants (Figure 5b) and appear to be highly expressed specifically in the leaf axils of plants

CIP7

BHLH32

(Figure S5) (Tantikanjana et al., 2001; Winter et al., 2007). Interestingly, the *bus1* mutant has a bushy growth phenotype (Hansen et al., 2001; Reintanz et al., 2001; Tantikanjana et al., 2001). SL analog, GR24, can be used to complement the hyperbranching phenotype of the SL deficient biosynthesis mutants *max1*, *max3*, and *max4* but not in SL signaling mutants, for example, *max2* (Gomez-Roldan et al., 2008; Umehara et al., 2008). To determine whether *BUS1* is involved in SL biosynthesis, GR24 was applied to rosette buds of WT (Col-0), *max4*, *max2*, and *bus1* mutants. Consistent with published results, GR24 complements the branching phenotype of the *max4* mutant but not the *max2* mutant (Figure 5d). Similarly, the enhanced branching phenotype of *bus1* cannot be complemented by GR24 application suggesting that GSL's connection to branching inhibition might be downstream of SL signaling (Figure 5d).

LDOX

UF3GT

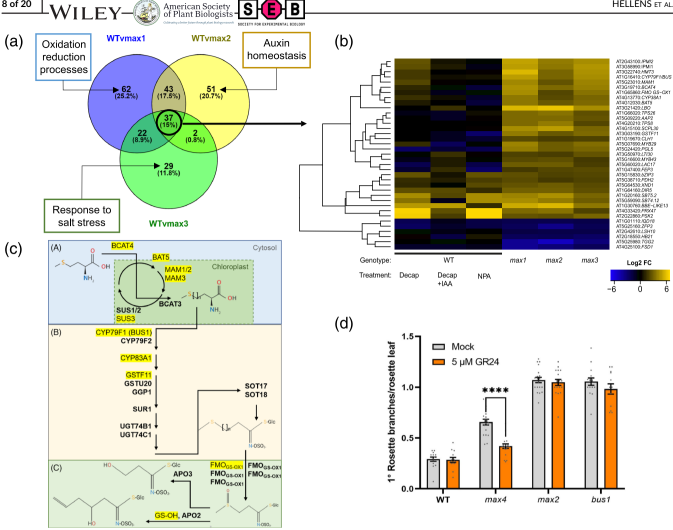


FIGURE 5 Strigolactone (SL) mutants highly upregulate glucosinolate biosynthesis. (a) Overlap between differentially expressed genes (DEGs) for max1, max2, and max3 mutants. Top gene ontology (GO) term for each individual mutant is highlighted (full list in Table S3). Percentages in each bubble represent the percentage of all genes in the entire Venn diagram in that bubble. (b) Expression for the 37 genes commonly identified as differentially expressed in max mutants. This gene list shows enrichment in glucosinolate metabolism. (c) Diagram of aliphatic glucosinolate biosynthesis pathway showing genes (highlighted in yellow) that are upregulated in max mutants. The aliphatic glucosinolate biosynthesis pathway is split into three components: (A) chain elongation, part of which occurs in the chloroplast; (B) biosynthesis of core glucosinolate structure; and (C) secondary modifications. For more detailed explanation, see Sønderby et al., 2010), (d) Exogenous SL (GR24) does not restore high branching phenotype of the bus 1 mutant to WT. Data are presented as mean \pm SEM (n=13-20) two-way ANOVA P-value of <.0001.

2.5 SL negative feedback is only partly regulated by auxin

Consistent with previous reports, we find that SL biosynthesis genes D27, MAX4, MAX3, and LBO are upregulated in max1, max2, and max3 SL mutant hypocotyls (Bennett et al., 2006; Brewer et al., 2016; Hayward et al., 2009; Shen et al., 2012; Waters, Brewer, et al., 2012) (Figure 6a). This negative feedback loop has been suggested to be auxin mediated (Hayward et al., 2009; Ligerot et al., 2017). However, if auxin was solely responsible, exogenous auxin would be expected to result in the same increase in SL biosynthesis gene expression as observed in SL biosynthesis mutants. Using a transcriptomic approach, we can confirm auxin uptake and signaling by examining transcriptional responses of

auxin-related genes. Here, we used a ratio of the fold change gene expression compared with WT_Intact across all three max mutants used in this study to the fold change gene expression in WT_Decap_IAA compared with WT_intact plants (Figure 6b). A ratio of 1 (the x-axis) would indicate that IAA/auxin application to decapitated plants has the same effect on SL biosynthesis as an SL biosynthesis gene mutation, a ratio below 1 indicates high auxin regulation (bars below the x-axis), and a ratio above 1 indicates that it is something different in the SL mutants causing gene expression changes (bars above the x-axis) (Figure 6b). Genes covering the major classes of auxin-responsive genes (PIN, IAA, GH3, and SAUR) were selected to compare against. For all 16 auxin-responsive genes, the ratio of gene expression is equal to or less than one, implying that their expression can mostly be explained by the

FIGURE 6 Feedback regulation of strigolactone (SL) biosynthesis genes in SL mutants is predominantly auxin-independent. (a) Auxin-related genes in SL mutant hypocotyls show similar expression to WT. Significant (*P* = .1) auxin-related genes were derived from gene ontology (GO) terms from differentially expressed genes (DEGs) from WT_Decap_IAA and WT_NPA treatments. Cluster 1 contains auxin-related genes upregulated in WT_Decap and WT_NPA consistent with inhibition of polar auxin transport. Cluster 2 contains genes typically upregulated by auxin. Cluster 3 contains SL biosynthesis genes (except MAX1). Cluster 4 contains genes downregulated by WT_Decap and WT_NPA treatments. (b) The ratio of fold change in MAX mutants and fold change in WT_Decap_IAA treated show selected SL biosynthesis (in orange), and sugar responsive genes (in blue) are highly expressed in SL biosynthesis mutants compared with auxin-treated samples. Auxin-responsive genes (in red) all show a low (≤1) ratio of MAX: WT_Decap_IAA fold change expression indicating predominantly auxin regulation of expression.

addition of auxin (red bars in Figure 6b). However, the ratio of gene expression for SL biosynthesis genes D27, MAX4, MAX3, and LBO are all above 1 (orange bars in Figure 6b). A high ratio

indicates that auxin alone is not responsible for the increased SL biosynthesis and provides circumstantial evidence for non-auxin feedback on SL biosynthesis gene expression.

Several sugar-responsive genes were highlighted as being co-regulated with the SL biosynthesis gene, MAX1 (Figure 3b). When the fold change gene expression ratio between SL mutant and IAA-treated plants is applied to these genes, they also show a ratio above 1, like the SL biosynthesis genes (blue bars in Figure 6b). This indicates that these sugar-responsive genes are also not solely controlled by auxin. Using available data from several microarray experiments through Genevestigator (Hruz et al., 2008), we confirmed that SL genes are more responsive to treatments with sugar perturbations than those with auxin perturbations (Figure S6). MAX1 is not included in this analysis because unlike other SL biosynthesis genes (D27, MAX4. MAX3, and LBO). MAX1 is not upregulated in any SL mutant. However, MAX1 is upregulated by decapitation and NPA treatments. (Figure 6a). The SL receptor genes MAX2 and D14 are both also upregulated in WT Decap and WT NPA (Figure 6a). This indicates that transcriptional regulation of MAX1 in response to auxin is more similar to MAX2 and D14 than to D27, MAX4, MAX3, and LBO.

3 | DISCUSSION

In this study, we used whole genome differential gene expression and WGCNA to study transcriptional changes and patterns associated with disruptions to the SL pathway in Arabidopsis to identify potential new interactions between SL and other pathways. WT Decap and WT_NPA treatments showed the largest transcriptional responses and were quite similar (Figure 1a). This is consistent with the shoot tip being a source of auxin and affecting multiple aspects of development. WT_Decap_IAA showed fewer DEGs than WT_Decap alone, consistent with auxin being able to restore some, but not all of the transcriptional response caused by decapitation (Cline, 1996; Crawford et al., 2010). This comparison of the treatments in WT enables a grouping of genes involved in the auxin-independent decapitation responses including wounding and loss of the main shoot sink tissue, the shoot tip. In contrast, relative to these large responses, mutants in SL signaling have comparatively few transcriptional differences compared with WT Intact. It is often noted that very few genes are transcriptionally regulated by the loss of SLs compared with other plant hormones (Lantzouni et al., 2017; Mashiguchi et al., 2009; Smith & Li, 2014; Wang et al., 2020), which is somewhat surprising given the number of phenotypic differences in SL mutants compared with WT plants (Brewer et al., 2013).

Differential gene expression across this study has highlighted enrichment of genes involved in secondary metabolite biosynthesis, particularly anthocyanins and GSLs, indicating that plant defense mechanisms are being modulated by SL and physiological perturbations in Arabidopsis. Transcriptional responses unique to and common to all of the *max* mutants included altered expression of GSL biosynthesis genes, and it is therefore likely that a connection exists between SLs and GSLs (Figure 5). Previous research showed that Arabidopsis mutants deficient in key steps of the GSL biosynthesis had increased shoot branching, indicating a potential role for GSLs in the regulation of shoot architecture besides their key role in defense

(Bak & Feyereisen, 2001; Hansen et al., 2001; Reintanz et al., 2001). Anthocyanin biosynthesis genes are downregulated across the data set, particularly by physiological treatments (Figure 4b). It is likely that SL and auxin are together enhancing anthocyanin biosynthesis (Ha et al., 2014; Patil et al., 2021). Additionally, we revealed some evidence for non-auxin-mediated negative feedback on SL biosynthesis.

3.1 Interaction between SLs and GSL metabolism

GSLs are secondary metabolites characteristic of the Brassicaceae family. GSLs can be divided into three groups based on their amino acid precursor, aGSLs are derived from Ala, Leu, Ile, Val, and Met (Figure 5c uses methionine as an example), benzenic GSLs are derived from Phe or Tyr; and indolic GSLs are derived from Trp (Sønderby et al., 2010). The major role for GSLs in plants lies in their toxicity to insects, thus preventing herbivory (Chew, 1988; Halkier & Gershenzon, 2006). However, with over 100 known GSLs with more than 40 different genes involved in GSL biosynthesis, it is likely that GSLs perform a wide range of roles in plants (Francisco et al., 2016; Sønderby et al., 2010). Indeed, a number of GSL mutants exhibit significant growth defects, including enhanced shoot branching and adventitious rooting; thus, it is likely that cross-talk plant hormones and GSL biosynthesis (Burow et al., 2015; Burow & Halkier, 2017; Francisco et al., 2016; Robert-Seilaniantz et al., 2011).

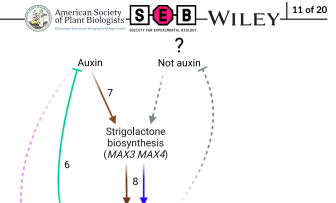
All three of the SL mutants showed significant upregulation of genes in the aGSL biosynthesis pathways, indicating that SLs might be inhibiting GSLs (Figures 4, 5c, and S4) (Sønderby et al., 2010). Of genes commonly upregulated by max mutants (Figures 5B and S4), the transcription factor MYB29 regulates the expression of aGSL biosynthesis genes in Arabidopsis, including BAT5, which belongs to the bile acid transporter family (Araki et al., 2013; Zhang et al., 2017; Zuluaga et al., 2019). IPMI1 and IPMI2, which catalyze plastidic chain elongation reactions, are also targets of MYB29 (Augustine & Bisht, 2017; Gigolashvili et al., 2009; Sønderby et al., 2010). BUS1 and CYP79F2 perform partially redundant functions in aGSL synthesis in Arabidopsis (Chen et al., 2003; Hansen et al., 2001). bus1 single and bus1/cyp79f2 double-knockout plants are GSL deficient (Reintanz et al., 2001; Tantikanjana et al., 2004). Interestingly, like max mutants, bus1 Arabidopsis mutants exhibit an increased branching phenotype (Reintanz et al., 2001) (Figure 5d). Given that exogenous SL is unable to recover the mutant bus1 phenotype, it is possible that aGSLs are regulated downstream of MAX2 (Figure 5d). bus1 plants are aGSL deficient; however, the max mutants (also showing a bushy phenotype) show enhanced aGSL biosynthesis further suggesting that aGSL inhibition of shoot branching is not due to inhibiting SL biosynthesis and may be activated as part of a feedback process to suppress branching under SL deficiency. It is likely that SLs and GSLs are working together to some extent as shoots of bus1 and SL mutants have higher levels of endogenous cytokinins (Tantikanjana et al., 2001; Young et al., 2014). The increase in CK levels in bus1 is far higher than

ever reported for SL mutants and this is consistent with the observation that CK pathways were upregulated in WT_Decap and WT_NPA but not in SL mutant hypocotyls (Table S3).

Auxin signaling has been found to affect aGSL levels in Arabidopsis. Triple-knockout *iaa5/iaa6/iaa19* plants have reduced aGSL levels and impaired drought tolerance. Application of GSLs to these mutants restores the stomatal regulation and drought tolerance (Salehin et al., 2019). *IAA5*, *IAA6*, and *IAA19* are well-known auxin-responsive genes (Weijers & Wagner, 2016) and were strongly promoted by auxin (Figure 6a). *IAA5* and *IAA19* expression was decreased in WT_Decap and WT_NPA treatments and increased in WT_Decap_IAA compared with WT (Figure 6a). These treatments had, comparatively, little or no effect on aGSL biosynthesis transcript levels (Figure S4C,D) compared with the high level of promotion in SL mutants. This indicates that high level of aGSL gene expression in SL mutants is not simply due to auxin signaling.

HOMEOBOX PROTEIN 21 (HB21) is among the 37 common DEGs between the three SL mutants (Figure 5b) and is downregulated in the max mutants. HB21 is a direct target of BRANCHED1 (BRC1), the central integrator of shoot branching signals (Wang et al., 2020), and acts redundantly with HB40 and HB53 to inhibit branching (González-Grandío et al., 2017; Whipple et al., 2011). BRC1 is downregulated in axillary buds of SL mutants (Braun et al., 2012; Dun et al., 2012). Because of BRC1 unfortunately not being a probe on the Affymetrix chip, we could not test BRC1 expression directly. However, as BRC1 is a well-established, bud-specific marker gene, we do not anticipate BRC1 expression in the hypocotyl (Figure S8) (Aguilar-Martínez et al., 2007; Braun et al., 2012; Klepikova et al., 2016). Therefore, HB21 downregulation in hypocotyls of max mutants might be independent of BRC1. Interestingly, HB21 is also known to regulate GSL accumulation, and hb21 mutant Arabidopsis plants have diminished aGSL levels (Li et al., 2014, 2018), further linking SL signaling, aGSL biosynthesis, and branching in Arabidopsis.

Another established aspect of SL signaling is the promotion of associations with AM fungi (Bouwmeester et al., 2007). However, many Brassica species do not form associations with AM fungi, including Arabidopsis (Kruckelmann, 1975). GSLs are a characteristic of brassicas and so may be part of the explanation that these plants cannot form symbiotic associations. GSLs have been linked to impacting microbe diversity in the rhizosphere (Bressan et al., 2009; Burow & Halkier, 2017). Additionally, Arabidopsis GSL biosynthesis genes are upregulated in response to AM fungi colonization in a neighboring host (Fernández et al., 2019). It may be that GSL biosynthesis is upregulated in this situation to inhibit AM fungi colonization although the spores are present. The SL involvement in AM fungi associations is likely a conserved response (Fernández et al., 2019; Lanfranco et al., 2018), but perhaps SLs can somewhat inhibit the GSL-mediated inhibition of AM fungi (Figure 7). GSL measurements in SL mutants in Arabidopsis as well as monitoring AM fungi colonization in non-GSLproducing Arabidopsis mutants and regular AM host species in response to different SL and GSL treatments would be needed to better understand this interaction.



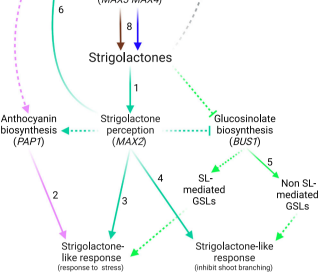


FIGURE 7 Summary of strigolactone (SL) signaling. Model of SL regulation of secondary metabolites and non-auxin-mediated feedback on SL biosynthesis. Dashed lines indicate ideas reported in this manuscript. Numbers are references to previously published interactions. (1) Hamiaux et al. (2012). (2) Landi et al. (2015). (3) Pandey et al. (2016). (4) Stirnberg et al. (2002). (5) Sønderby et al. (2010). (6) Ligerot et al. (2017). (7) Hayward et al. (2009). (8) Gomez-Roldan et al. (2008).

3.2 | Interactions between SLs and anthocyanins

Anthocyanins are an important class of flavonoids that represent a large group of ubiquitous plant secondary metabolites (Liu et al., 2018). They are glycosylated polyphenolic compounds involved in stress responses, especially high light and nutrient stress and so are fundamental in plants and their adaptation to the environment (Liu et al., 2018; Tanaka et al., 2008). Anthocyanins also largely play a protective role in plants (Agati et al., 2012; Gould et al., 2002; Landi et al., 2015; Zheng et al., 2020). Anthocyanin biosynthesis genes are downregulated compared with WT_Intact, across the data set, but especially in the WT_Decap, WT_Decap_IAA, and WT_NPA plants (Figure 4b). Auxin has been shown to regulate anthocyanin biosynthesis synergistically with cytokinins (Ji et al., 2015; Liu et al., 2014). Cytokinins promote anthocyanin biosynthesis and have been shown to induce DFR and UF3GT expression (Deikman & Hammer, 1995; Wang et al., 2019). Although co-treatment of auxin and cytokinins significantly enhances cytokinin-induced anthocyanin production, high auxin concentration (>9-µM NAA) strongly inhibits anthocyanin biosynthesis, even in the presence of cytokinin (Ji et al., 2015; Liu

et al., 2014). As well as hormones, sucrose induces anthocyanin production. Sucrose upregulates *PAP1* and *DFR*, which catalyze the first committed step in anthocyanin biosynthesis (Das et al., 2012; Teng et al., 2005). It is possible that altered auxin, cytokinins, or sugar levels in the stem after decapitation enhance anthocyanin biosynthesis.

SLs have also been shown to regulate anthocyanin accumulation in plants (Ferrero et al., 2018; Ito et al., 2015). This could suggest that SLs regulate anthocyanin accumulation downstream of auxin or sugar signaling, supported by max2 rosettes having decreased anthocyanin accumulation, as MAX2 is implicated in both auxin and sugar signaling pathways (Figure 4c) (Barbier, Péron, et al., 2015; Ha et al., 2014; Ligerot et al., 2017; Patil et al., 2021). Recent findings have revealed that SLs play a role in anthocyanin accumulation through upregulation of anthocyanin biosynthesis genes such as PAP1 (Wang et al., 2020). Wang et al. (2020) described max3 mutants as showing decreased anthocyanin content compared with WT, which our study consolidates, as the max3 mutant shows decreased expression of anthocyanin biosynthesis genes, including PAP1 (Figures 4b and 7). MAX2, although being essential to SL reception, is also essential for karrikin signaling by interacting with α/β hydrolase KARRIKIN INSENSITIVE 2 (KAI2) instead of D14 (Nelson et al., 2011; Waters, Nelson, et al., 2012). Karrikins can enhance anthocyanin accumulation, and this accumulation is dependent of KAI2. However, unlike max2 mutants, kai2 mutants do not show reduced anthocyanin accumulation compared with WT plants (Bursch et al., 2021). These findings agree with recent suggestions that MAX2 is a master regulator of flavonoid biosynthesis with max2 showing downregulated flavonoid biosynthesis and metabolism in roots, which is not seen in max4 or d14 mutants (Richmond et al., 2022) (Figure 4c).

3.3 | Transcriptional differences among SL biosynthesis mutants

The physiological treatments provide a good framework to identify genes that respond to major signaling changes within the plant relating to shoot branching and apical dominance. SL mutants enable us to narrow down which parts of the framework are specifically involved in regulation of or by SLs.

DEG and WGCNA analyses highlighted differences among SL mutants (Figure 5a). Although an independent function of MAX2 is expected considering its role in the perception (as opposed to biosynthesis) of SL, and its involvement in karrikin signaling, MAX3 and MAX1 act linearly in the SL biosynthesis pathway (Abe et al., 2014; Alder et al., 2012; Nelson et al., 2011; Stirnberg et al., 2007). MAX1 encodes a cytochrome P450 enzyme that converts carlactone to carlactonic acid, whereas MAX3 encodes a carotenoid cleavage dioxygenase that converts 9-cis- β -carotene to 9-cis- β -apo-10'-carotenal (Figure 2b). Consequently, the max1 mutant accumulates carlactone, whereas max3 mutants do not and instead accumulate 9-cis- β -carotene. max1 mutants showed a greater number of transcriptional changes than max2 and max3 mutants (Figure 1b). In accordance, another recent microarray analysis of SL mutants also reported a

difference in transcriptional responses in the SL mutants, highlighting substantial transcriptional changes in Arabidopsis *max1* mutant leaves compared with other SL mutants (Kumar et al., 2019). A higher transcriptional response in *max1* compared with *max3* might suggest that the build-up of carlactone is biologically relevant. Carlactone added to *max* mutants can inhibit rosette branching in Arabidopsis, even partially in the *max2* mutant background, which is deficient in SL sensing (Abe et al., 2014). Therefore, part of the carlactone effect on shoot branching seems to be not mediated by D14-MAX2 degradation of SMXLs opening a new avenue of enquiry (Abe et al., 2014; Kumar et al., 2019).

3.4 | Feedback regulation of SL biosynthesis

SL mutants have increased expression of SL biosynthesis genes (D27, MAX3. MAX4. and LBO), indicative of a negative feedback loop (Figure 6a) (Hayward et al., 2009). The perception of SL by MAX2 may contribute to negative feedback on SL biosynthesis genes via inhibiting auxin transport (Ligerot et al., 2017). However, auxin may not be solely responsible for negative feedback on SL biosynthesis in SL mutants. Application of auxin neither rescues the enhanced branching phenotype nor SL biosynthesis gene expression back to WT levels in SL mutants, nor does it restore depleted SL contents to roots of decapitated plants (Bennett et al., 2006; Hayward et al., 2009; Yoneyama et al., 2015) (Figures 6a and 7). To further examine the extent of non-auxin feedback regulation of SL biosynthesis gene expression, we plotted the ratio of gene expression in SL mutants to that of the same genes in auxin-treated plants (Figure 6b). If auxin was solely responsible for controlling SL biosynthesis, a ratio of 1, or below, is expected because the high gene expression change in SL mutants would be balanced with the high gene expression of auxin response after auxin treatment. As expected, all auxin-responsive genes (in red; Figure 6b) fall below the threshold of 1. In contrast, all SL biosynthesis genes (in orange) show a very high ratio above 1. The inability of auxin to restore SL biosynthesis gene levels to intact WT levels in max mutants suggests some non-auxin-related negative feedback on SL biosynthesis genes (Figure 6b).

Sugars or sugar signaling is a candidate for a non-auxin signal regulating SL biosynthesis gene expression and needs to be tested experimentally. Consistent with this, sugar perturbations generally have a larger effect on the transcriptional regulation of SL biosynthesis genes *MAX1* and *D27* than do auxin perturbations, but this is not the case for *MAX3* (Figures S6 and S7). Given that *MAX3* is not differentially expressed in treatments that affect sugar levels, sugar itself is not a great candidate for a common non-auxin signal regulating SL biosynthesis gene expression.

Sucrose is known to have an inhibitory effect on SL perception (Bertheloot et al., 2020; Fichtner et al., 2021; Patil et al., 2021). Sugar signaling-related genes were strongly correlated with SL genes MAX1 (i.e., TPS1 and TPPE) and MAX2 (i.e., TPS8) (Figure 3b) raising the hypothesis that sugar signaling could be affecting expression of these genes. MAX2 expression is suppressed by sugars (Patil et al., 2021),

research.

whereas MAX1 expression is enhanced (Figure S7). It is important to notice here that regulation of MAX1 gene expression is different from the other SL biosynthesis genes. In particular, MAX1 is not as highly expressed in SL mutants than other SL biosynthesis genes (Figure 3b). These similarities in expression among MAX1, MAX2, and also D14 suggest that MAX1 may be regulated by sugars similar to the SL signaling genes (Barbier, Péron, et al., 2015; Bertheloot et al., 2020; Patil et al., 2021). The interactions between auxin, sugars,

and SL levels and signaling is therefore an interesting topic for further

3.5 | Conclusion

The aim of this research was to identify novel components in the SL regulatory network and to provide a tool for additional network analysis for the regulation of the SL pathway. GSLs and anthocyanins are both highlighted as potentially new targets of SL signaling. Additionally, we present evidence for non-auxin-mediated feedback on SL biosynthesis and propose that components of sugar signaling could be partially involved in this process (Figure 7).

4 | EXPERIMENTAL PROCEDURES

4.1 | Plant materials and growth conditions

For the microarray experiment, Arabidopsis seeds were stratified for 72 h at 4°C and then sown on University of California potting mix type C and vermiculite (3:2 v/v). The plants were grown for 2 weeks (until they had five to six leaves expanded) in a temperature- and humidity-controlled growth chamber under 24°C with 16-h light/21°C with 8-h dark photoperiod providing 180 μ mol m $^{-2}$ s $^{-1}$. For anthocyanin measurements (Figure 4c) and SL treatments (Figure 5c), Arabidopsis seeds were stratified for 72 h at 4°C and then sown on UQ23 potting mix (70% composted pine bark 0–5 mm, 30% coco-peat). Plants were grown for 7 weeks (SL treatments) in temperature- and humidity-controlled growth chambers (16 h:8 h, light:dark, 150 \pm 20 μ mol m $^{-2}$ s $^{-1}$, 22°C:20°C, day:night).

4.2 | Treatments for microarray

Two-week-old WT Arabidopsis plants (cv. Colombia-0 or Col-0) were subjected to three different treatments: decapitation (WT_Decap), decapitation + IAA (WT_Decap_IAA), and NPA (WT_NPA) as described in Brewer et al. (2016). Treatments were compared with intact WT and three SL mutants *max1-1*, *max2-1*, and *max3-1* (Booker et al., 2004; Stirnberg et al., 2002). Decapitation involved the removal of the vegetative shoot apex and three to four expanding leaves with scalpel and fine forceps, followed immediately by application of



lanolin (<10% EtOH) to the cut surface. For the decapitation + auxin treatment, lanolin was supplemented with 3 mg mL $^{-1}$ of IAA. For NPA treatments, 3 mg ml $^{-1}$ of NPA was applied in a lanolin ring (<10% EtOH) around the top of the hypocotyl. Hypocotyls were collected as three biological replicates of 120–150 plants per treatment or mutant and flash frozen in liquid nitrogen and stored at -80° C. Treatments for each replicate were performed simultaneously, and each biological replicate was harvested over a period of 60–80 min at 24 h after treatment.

4.3 | RNA isolation

Total RNA was isolated using NucleoSpin RNA plant kits (Machery-Nagel, Düren, Germany) and further purified and concentrated using the RNeasy MinElute Cleanup Kit (Qiagen, Hilden, Germany) according to manufacturer's protocol. RNA was quantified using a NanoDrop 1000 Spectrophotometer (Thermo Scientific, Wilmington, Delaware, USA).

4.4 | Microarray hybridization

Hybridization, washing, and scanning were performed at the Clive & Vera Ramaciotti Centre for Gene Function Analysis (University of NSW, Sydney), using Arabidopsis ATH1 genome oligonucleotide chips. High-quality biotin-labeled complementary RNA (cRNA) was prepared from 4-µg total RNA per sample according to the standard Affymetrix protocol, hybridized to whole-genome Arabidopsis ATH1 GeneChips and scanned with an Affymetrix GCS3000 Scanner (Affymetrix, Santa Clara, CA) according to the manufacturer's recommended protocol (GeneChip Expression Analysis Technical Manual).

4.5 | Microarray data pre-processing

The scanned probe array images (.DAT files) were converted into .CEL files using the GeneChip Operating Software (Affymetrix). Affymetrix CEL-data files were preprocessed using the statistical language and environment R version 3.2.2 (Ihaka & Gentleman, 1996). Probe intensities were normalized for background correction using the robust multiarray average method corrected for GC content of the oligo, an algorithm known as GCRMA (Harr & Schlötterer, 2006; Irizarry et al., 2003). GCRMA background correction was used in conjunction with the only perfect match (PM) correction algorithm (Seo & Hoffman, 2006). Gene expression values were calculated from the PM probes using the median polish summary method (Harr & Schlötterer, 2006). Raw gene expression values are supplied in Table S4. The above-mentioned methods were executed using the "affy" (Gautier et al., 2004) and "simpleaffy" (Wilson & Miller, 2005) packages.

4.6 | Microarray data statistical analysis

The limma package (Ritchie et al., 2015; Smyth, 2004) was used to perform moderated *t*-statistics between the treatments to identify DEGs. For each probe, fold change and corresponding *P*-values measuring the statistical significance of DEGs were calculated using empirical Bayesian statistics, which moderates the standard errors within each probe (Ritchie et al., 2015). Benjamini–Hochberg's method was used to correct *P*-values for multiple testing (Benjamini & Hochberg, 1995). We used an adjusted *P*-value (adj *P*-value) cut-off of .05 and absolute LogFC of >2 (|LogFC≥2|) to assign statistical significance. Gene functional annotations obtained from The Arabidopsis Information Resource (TAIR) database (Huala et al., 2001) were used to annotate the genes and match gene IDs to their corresponding probe set numbers provided by the GeneChip manufacturer (Affymetrix).

4.7 | PCA

PCA is a popular multivariate statistical tool that is used to summarize a large number of variables to a smaller number of derived variables that may be readily visualized in two- or three-dimensional space. PCA was applied to summarize and visualize the entire gene expression matrix (all treatment and genes) using the R package "ggfortify" (Tang et al., 2016).

4.8 | Functional enrichment analysis

GO enrichment analysis infers biological meaning from systems biology experiments (Hill et al., 2008). Array IDs of DEGs per comparison against WT_Intact as queries in the DAVID online database for GO enrichment analysis (Dennis et al., 2003). Cytoscape software with the ClueGO package was used to identify and visualize significant (P < .05) biological processes represented in each group.

4.9 | Graphs and heatmap rendering

GraphPad Prism 9 or R programming tools for plotting data, "ggplot2" (Wickham, 2016) and "gplots" (Warnes et al., 2009), were used to render graphs and heatmaps.

4.10 | WGCNA

A signed co-expression network was constructed with the WGCNA package in R (Langfelder & Horvath, 2008). WGCNA identifies modules composed of genes that are connected in terms of the topological overlap mapping metric (TOM), a neighborhood proximity measurement that quantifies the degree of sharedness among network neighbors. We used signed network and soft-thresholding power ($\beta=3$)

based on the scale-free model fitting index $R^2 > .9$ (Figure S1A) to obtain an adjacency matrix. The adjacency matrix was transformed into a TOM with the topological overlap (TO)-based dissimilarity (1-TOM). This step resulted in a clustering tree/dendrogram (Figure S1B) whose branches were identified for cutting, depending on their shape. The dynamic tree-cutting algorithm cut the hierarchal clustering tree into modules, that is, clusters of highly co-expressed genes. To obtain moderately large and distinct modules, the default minimum module size was set to 30 genes, and a minimum height for merging modules was .25. Modules were summarized by a hierarchical clustering dendrogram, and module structure was visualized by a heatmap and topological overlap matrix plot. WGCNA assigned to each module a unique color label, and the relationships between each module and treatment/mutants was analyzed to identify the highly correlated modules. Finally, we selected clusters of genes highly correlated with SL biosynthesis and signaling genes and exported them to Cytoscape for network visualization (Shannon et al., 2003).

4.11 | Anthocyanin measurements

For anthocyanin measurements, plants were grown in temperature-and humidity-controlled growth chambers (16 h:8 h, light:dark, 140 \pm 10 μ mol m $^{-2}$ s $^{-1}$, 22°C:18°C, day:night). Plants were harvested 5 days before bolting, snap frozen in liquid nitrogen, and ground to a fine powder. To 30–60 mg of powder, 600 μ L of 1% HCl in methanol was added and the mixture left overnight at 4°C. The following day, 600 μ L of chloroform and 200 μ L of water were added. Samples were centrifuged for 10 min at 16,000 x g. Anthocyanin content is presented as (A $^{530\,\text{nm}}$ –A $^{657\,\text{nm}}$) per gram of fresh weight.

4.12 | SL treatments

Five microliters of 0- or 5- μ M rac-GR24 solution in acetone and Tween20 was applied to rosette buds on day 21. Ten microliters was applied on days 24, 32, 35, and 38. On days 27 and 29, 10 μ L of 0- or 10- μ M GR24 was applied. Branching was scored on day 56, and number of primary rosette branches was normalized by number of leaves.

5 | ACCESSION NUMBERS

AT2G43100, AT5G07690, AT5G09220, AT1G20160, AT2G18550, AT2G22860, AT5G16600, AT5G24420, AT5G15830, AT1G65860, AT1G01110, AT4G25100, AT1G16410, AT1G19670, AT5G23010, AT3G19710, AT3G50970, AT4G15100, AT4G12030, AT2G42610, AT5G59090, AT1G66020, AT4G33420, AT5G38710, AT1G64160, AT5G60020, AT3G22740, AT3G58990, AT1G30760, AT5G25980, AT3G03190, AT4G13770, AT5G64530, AT5G25160, AT1G47400, AT3G21420, AT4G20210, AT1G04240, AT1G04250, AT1G04550, AT1G15580, AT1G17140, AT1G19850, AT1G23080, AT1G28010, AT1G29510, AT1G34670, AT1G48660, AT1G56650, AT1G57560,

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AT1G70940, AT1G72430, AT1G73590, AT1G75500, AT1G76520, AT1G77110, AT1G78100, AT2G01940, AT2G16580, AT2G21050, AT2G23170, AT2G26170, AT2G33310, AT2G38120, AT2G44990, AT3G14370, AT3G15540, AT3G28345, AT3G53480, AT3G59900, AT3G62100, AT3G63300, AT4G00880, AT4G14560, AT4G17980, AT4G22620, AT4G27260, AT4G28640, AT4G30290, AT4G32880, AT4G34760, AT4G36110, AT5G13360, AT5G19530, AT5G20820, AT5G39610, AT5G43700, AT5G47370, AT5G54490, AT5G54510, AT5G55250, AT5G62380, AT5G65670, AT5G67480, AT1G03055,

AUTHOR CONTRIBUTIONS

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Alicia M. Hellens, Franziska Fichtner, Philip B. Brewer, and Christine A. Beveridge designed the research; Alicia M. Hellens, Franziska Fichtner, and Philip B. Brewer performed the research; Alicia M. Hellens, Tinashe G. Chabikwa, Franziska Fichtner, Philip B. Brewer, and Christine A. Beveridge analyzed the data; Alicia M. Hellens and Tinashe G. Chabikwa wrote the paper. All authors except Tinashe G. Chabikwa edited the manuscript and approved the final version.

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CONFLICT OF INTEREST STATEMENT

The Authors did not report any conflict of interest.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are openly available in the National Center for Biotechnology Information Gene expression omnibus (NCBI GEO) at https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE226903 using reference number [GSE226903].

ORCID

Alicia M. Hellens https://orcid.org/0000-0001-9715-8012

Franziska Fichtner https://orcid.org/0000-0002-4508-5437

Philip B. Brewer https://orcid.org/0000-0003-4871-9260

Christine A. Beveridge https://orcid.org/0000-0003-0878-3110

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