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### Authors

Bjornson, Marta  
Balcke, Gerd Ulrich  
Xiao, Yanmei  
[et al.](#)

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## Integrated omics analyses of retrograde signaling mutant delineate interrelated stress response strata

Marta Bjornson<sup>1,2</sup>, Gerd Ulrich Balcke<sup>3</sup>, Yanmei Xiao<sup>1</sup>, Amancio de Souza<sup>1,±</sup>, Jin-Zheng Wang<sup>1,±</sup>, Dina Zhabinskaya<sup>4</sup>, Ilias Tagkopoulos<sup>5</sup>, Alain Tissier<sup>3</sup>, and Katayoon Dehesh<sup>1,±,\*</sup>

<sup>1</sup>Dept. of Plant Biology, University of California, Davis, CA 95616

<sup>2</sup>Dept. of Plant Sciences, University of California, Davis, CA 95616

<sup>3</sup>Dept. of Physics, University of California, Davis, CA 95616

<sup>4</sup>Dept. of Computer Science, University of California, Davis, CA 95616

<sup>5</sup>Dept. of Cell and Metabolic Biology, Leibniz-Institute of Plant Biochemistry, Halle, Germany

### Abstract

To maintain homeostasis in the face of intrinsic and extrinsic insults, cells have evolved elaborate quality control networks to resolve damage at multiple levels. Interorganellar communication is a key requirement for this maintenance, however the underlying mechanisms of this communication have remained an enigma. Here we integrate the outcome of transcriptomic, proteomic, and metabolomics analyses of genotypes including *ceh1*, a mutant with constitutively elevated levels of both the stress-specific plastidial retrograde signaling metabolite methylerythritol cyclodiphosphate (MEcPP) and the defense hormone salicylic acid (SA), as well as the high MEcPP but SA deficient genotype *ceh1/eds16*, along with corresponding controls. Integration of multi-omic analyses enabled us to delineate the function of MEcPP from SA, and expose the compartmentalized role of this retrograde signaling metabolite in induction of distinct but interdependent signaling cascades instrumental in adaptive responses. Specifically, here we identify strata of MEcPP-sensitive stress response cascades, among which we focus on selected pathways including organelle-specific regulation of jasmonate biosynthesis; simultaneous induction of synthesis and breakdown of SA; and MEcPP-mediated alteration of cellular redox status in particular glutathione redox balance.

Collectively, these integrated multi-omic analyses provided a vehicle to gain an in-depth knowledge of genome-metabolism interactions, and to further probe the extent of these interactions and delineate their functional contributions. Through this approach we were able to pinpoint stress-mediated transcriptional and metabolic signatures and identify the downstream processes modulated by the independent or overlapping functions of MEcPP and SA in adaptive responses.

\*Corresponding author: Katayoon Dehesh, University of California, Riverside, Tel: 951-827-6370, Fax: 951-827-5155, kdehesh@ucr.edu.

±Current address: Institute for Integrative Genome Biology, and Dept. of Botany and Plant Sciences, University of California, Riverside, CA 92521

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## Keywords

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## Introduction

To maintain cellular homeostasis in the face of frequent changes in their internal and external environment, plants must delicately balance and fine-tune adaptive responses at the tissue, cellular, and organellar levels. To achieve this homeostasis, tight regulation of a variety of cellular processes, ranging from changes in nuclear gene expression to alterations in mRNA or protein stability is required. Of relevance to these adaptive processes, inter-organellar communication via signals such as the general-stress-induced plastidial retrograde metabolite 2-*C*-methyl-D-erythritol-2,4-cyclodiphosphate (MEcPP) have a key function in coordinating stress responses (Xiao *et al.*, 2012).

MEcPP is a bifunctional chemical entity that serves both as a metabolic intermediate of the methyl erythritol phosphate (MEP) pathway responsible for production of isoprenoid precursors, and as a signaling metabolite. Constitutively expressing *HYDROPEROXIDE LYASE 1* (*ceh1*) mutant plants were found to accumulate high levels of MEcPP, which in turn results in a constitutive production of the otherwise defense-associated hormone salicylic acid (SA) (Xiao *et al.*, 2012). Generation of a double mutant containing high MEcPP and negligible levels of SA via introducing the SA defective mutant *eds16* (mutant in *ISOCHORISMATE SYNTHASE1*, also known as *sid2*) into the *ceh1* background unraveled distinct signaling roles of MEcPP (Xiao *et al.*, 2012). These specific roles include induction of constitutive expression of one branch of jasmonic acid (JA) responsive genes (Lemos *et al.*, 2016), a constitutive general stress response (Benn *et al.*, 2016) (GSR), and induction of gene expression of a branch of the unfolded protein response (UPR) in the endoplasmic reticulum (Walley *et al.*, 2015). Collectively, these findings identify MEcPP as an interorganellar communication signal, with an as-yet-unknown mechanism of action.

As an initial step to untangle the role and mechanism of MEcPP action, we characterized the *ceh1* mutant using microarrays (Walley *et al.*, 2015), followed by expanding these data through RNA-seq on *ceh1*, *ceh1/eds16*, *eds16*, and parent lines. Subsequently, we performed proteomic analyses on these genotypes to address the well-established regulatory function of translation in stress responses (Mustroph *et al.*, 2009; Zanetti *et al.*, 2005; Juntawong *et al.*, 2014; Walley *et al.*, 2015). We finally extended the analyses to metabolomics to gain insight into the metabolic status of these lines. The integration of transcriptomics, proteomics, and metabolomics data resulted in the most comprehensive analyses of the function of plastidial retrograde signaling, while circumventing reported deficiencies of the individual approaches (Wanichthanarak *et al.*, 2015; Rajasundaram and Selbig, 2016; Larsen *et al.*, 2015; Rajasundaram *et al.*, 2014; Hyun *et al.*, 2014; Zeng *et al.*, 2013; Hirai *et al.*, 2004; Wilson *et al.*, 2015; Vijayakumar *et al.*, 2016).

Collectively, the integration of three omics outputs delineates the role of MEcPP from SA in regulation of adaptive responses, and reveals a MEcPP-mediated function in controlling SA production and decay, organelle-specific regulation of JA biosynthesis, as well as enhanced oxidation in the pool of glutathione.

## Results and Discussion

### Gene Set Enrichment Analysis of Transcriptome and Proteome Data

To determine the nature of metabolic disruptions in the *ceh1* mutant, we undertook gene set enrichment analysis (GSEA) to identify metabolic pathways/processes with altered transcription/translation. To evaluate the differences in both transcription and translation, and to differentiate which of these are due directly to MEcPP vs. the elevated SA in *ceh1*, we conducted GSEA analyses on both proteomics and transcriptomics data, comparing each of *ceh1*, *ceh1/eds16*, and *eds16* to parental controls. Transcriptomics data detected more genes (7884, 551 of which were differentially regulated in at least one genotype) than proteomics (4192 proteins of which 35 were differentially regulated in at least one genotype) with a greater number differentially regulated, indicative of the greater sensitivity of this technique. These analyses revealed a number of robustly altered pathways (Table S1, gene lists of relevant pathways in Table S2). Few pathways were significantly modified in *eds16* relative to parent, so for simplicity induced and suppressed pathways at the RNA and protein levels in only *ceh1* and *ceh1/eds16* are shown in Figure 1.

Among the upregulated pathways are several which are similarly altered in *ceh1* and *ceh1/eds16*, indicating that they are changed primarily by MEcPP, with little or no input of SA. Of particular note is “Protein processing in the endoplasmic reticulum,” which is upregulated in both *ceh1* and *ceh1/eds16* at both the RNA and protein level. In addition to this previously reported role of MEcPP in induction of the unfolded protein response in the ER of *ceh1* (Walley *et al.*, 2015), using GSEA analyses on transcriptomics data enabled us to identify “Protein export” and “Phagosome” as differentially modified pathways in *ceh1* and “Proteasome” in both *ceh1* and *ceh1/eds16*, thereby extending the portfolio of disrupted protein folding and stability in response to MEcPP and associated SA accumulation. Other upregulated pathways identified only in transcriptomics data include “Plant-pathogen interaction” and “Glutathione metabolism.” This may suggest translational or post-translational suppression of these pathways, or merely be the consequence of an intrinsically different detection limit between the two ‘omics approaches – indeed more than half of the proteins associated with these pathways were not detected in any genotype (Walley *et al.*, 2015).

Among suppressed pathways are “Photosynthesis” and “Carbon fixation in photosynthetic organisms,” downregulated in both *ceh1* and *ceh1/eds16* at the RNA level, indicating a role of MEcPP in repressing these pathways independently of SA. In general, these disruptions in the *ceh1* mutant may be considered more likely to be due to specific signaling by MEcPP than perturbation of the MEP pathway, which theoretically is believed to be redundant with the cytosolic mevalonate pathway. In plants, a large proportion of the produced isoprenoids participate in photosynthetic processes and only a minor of the isoprenoid flux contribute to the synthesis of hormones (Vranová *et al.*, 2012; Vranová *et al.*, 2013). The light activation

of the MEP-pathways genes as opposed to light-inhibition of the MVA pathway genes, concomitant with increased synthesis of isoprenoid-derived metabolites such as phytyl chains and carotenoids after illumination is a prime example of distinct function of the two isoprenoid biosynthesis pathways (Vranová *et al.*, 2013; Ghassemian *et al.*, 2006; Rodríguez-Concepción, 2006). As such, the disruption of photosynthesis-related pathways in *ceh1* may in part be due to perturbation of the MEP pathway and by extension chlorophyll biosynthesis, as represented by the downregulation of “Porphyrin and chlorophyll metabolism” in both *ceh1* and *ceh1/eds16* (Figure 1). In contrast to RNA levels, protein levels of “Photosynthesis” and “Carbon fixation in photosynthetic organisms” in *ceh1/eds16* as compared to *ceh1* are unaffected and enhanced, respectively. This is in line with published results that SA signaling affects protein translation or stability of these pathways (Rivas-San Vicente and Plasencia, 2011).

Alpha-linolenic acid metabolism, which comprises mostly JA biosynthesis, is among those pathways notably affected by both MEcPP and SA. This pathway is upregulated in both *ceh1* and *ceh1/eds16* at the protein level, but only *ceh1/eds16* at the RNA level. These data support previous observations that while SA exerts an inhibitory function, MEcPP induces jasmonate biosynthesis and responses (Lemos *et al.*, 2016). Finally, a number of pathways upregulated in *ceh1/eds16* specifically at the protein level are, conversely, downregulated in *ceh1* at the protein and RNA level, highlighted in red in Figure 1. These pathways are largely associated with central metabolism, and are likely modulated by SA levels, known to inhibit both photosynthesis and respiration via both transcriptional and post-translational mechanisms (Rivas-San Vicente and Plasencia, 2011).

Independent analyses of transcriptomic and proteomic data allowed identification of lowly-expressed pathways utilizing RNA-seq while leveraging proteomic data to identify specific protein level modulation and increase confidence in transcriptomic data (Table S1 and S2). In summary we identified several classes of differentially regulated pathways: those affected a) solely by MEcPP, such as protein stability; b) primarily by MEcPP with partial SA contribution, including stress signaling pathways such as jasmonate biosynthesis, pathogen response and glutathione metabolism; and c) predominantly by SA with some contribution of MEcPP, primarily related to central metabolism.

### Selected metabolic disruptions caused by HDS mutation

The substantial alteration of metabolic pathways identified by transcriptomic and proteomic analyses prompted us to extend our studies to metabolomics, by first assaying a panel of 90 small molecules primarily associated with central carbon metabolism in all the genotypes, of which 82 accumulated differentially among the four tested genotypes (Table S3). A subset of these results is shown in Figure 2, with the remaining metabolites' information in Figure S1. As a confirmatory analysis, we initially examined intermediates of the MEP pathway, finding the expected accumulation of MEcPP upstream of HDS in both *ceh1* and *ceh1/eds16*. In contrast with previously published results (Gonzalez-Cabanelas *et al.*, 2015), we found similar over-accumulation of earlier pathway intermediates, up to and including MEP (Figure 2A). Of note, dimethylallyl pyrophosphate (DMAPP) is not reduced in lines with mutated HDS, reflective of function of the redundant mevalonate pathway in producing this

metabolite. We did not detect significant changes in mevalonate pathway genes/proteins, but this does not rule out post-translational regulation of this pathway.

Additional notable disruptions to central metabolism were in the Calvin-Benson-Bassham (CBB) cycle and the tricarboxylic acid (TCA) cycle, with an overall buildup of CBB intermediates and depletion of TCA cycle intermediates in *ceh1* and *ceh1/eds16* (Figure 2B), consistent with a potential block in the CBB cycle, and reduced sugar availability (Rojas-González *et al.*, 2015). More information about this block may be inferred by the fact that while sedoheptulose 1,7-bisphosphate accumulates in *ceh1* and *ceh1/eds16*, sedoheptulose 7-phosphate is depleted relative to WT levels. The gene encoding sedoheptulose 1,7 bisphosphatase, which mediates this conversion, is not only downregulated in *ceh1* and *ceh1/eds16* at the transcript and protein level (Table S4, Walley *et al.*, 2015), it is also post-translationally regulated by chloroplast conditions including pH, Mg<sup>2+</sup>, feedback inhibition, and redox status (Raines *et al.*, 1999). Thus the evident lack of sedoheptulose 1,7 bisphosphatase activity may suggest differences in the chloroplastic environment of MEcPP-overaccumulating lines. Intriguingly, the stronger modulation of a number of CBB intermediates in *ceh1*, as compared to *ceh1/eds16*, clearly differentiate input of SA from MEcPP in the process (Figure 2B), corroborating the observation that pathway “Carbon fixation in photosynthetic organisms” is induced in *ceh1/eds16* and conversely suppressed in *ceh1* (Figure 1). However, in the TCA cycle, most metabolites are similarly disrupted in *ceh1/eds16* as in *ceh1* alone, suggesting some effect on these pathways either of the MEcPP signal or the disruption of the MEP pathway, but independently of SA accumulation.

Another noted trend is the redox status of the various reducing currencies of the cell (Figure 2C). While NAD and NADP are present at lower levels, their reduced forms NADH and NADPH are present at greater levels in *ceh1* and *ceh1/eds16*. In contrast, the existing glutathione pool is more oxidized, with more GSSG and less GSH in *ceh1* and *ceh1/eds16*. Collectively, these data suggest altered redox potential in response to accumulation of MEcPP.

Finally, a number of assayed amino acids are differentially accumulated in *ceh1* and *ceh1/eds16* (Figure 2D). Specifically, the levels of over half of the amino acids show a strong SA-dependent trend (Figure 2D), present at greater levels in *ceh1* than in *ceh1/eds16*. This corroborates the previous finding that SA increases free amino acid concentrations in algae (Kovacik *et al.*, 2010), perhaps via effects on protein synthesis and/or breakdown. MEcPP also influences amino acid levels independently of SA; a subset of amino acids, including tryptophan and methionine, are reduced in both *ceh1* and *ceh1/eds16*. Of these tryptophan is of note since the tryptophan breakdown pathway “Tryptophan metabolism” is upregulated in *ceh1/eds16* at the protein level, reflecting dampening effects of SA (Figure 1). Additionally, constitutively high SA in *ceh1* may further contribute to sequestration/reduction of the pool of chorismate, a precursor shared by SA and tryptophan biosynthetic pathways. These processes combined present an example of how transcriptomic/proteomic data may inform metabolomics data analyses. Finally a subset of assayed amino acids, including glycine and serine, is present at greater levels in both *ceh1* and *ceh1/eds16*, but with stronger enrichment in *ceh1* than *ceh1/eds16*. Notably, these amino acids play an important role in the rescue of Rubisco oxygenase activity, photorespiration. Photorespiration is not represented directly by

a unique KEGG pathway, but is a part of the pathways “Glycine, serine, and threonine metabolism,” and “Glyoxylate and decarboxylate metabolism,” both of which are downregulated in *ceh1* by the combined input from SA and MEcPP (Figure 1).

Collectively, the integration of transcriptomics, proteomics and metabolomics analyses enabled us to confirm expected results, as in MEcPP accumulation, to suggest underlying mechanisms, as in tryptophan depletion, and to identify areas for further investigation, as in glutathione redox balance and potential impact on photorespiration.

### MEcPP regulation of salicylic acid biosynthesis and metabolism

High SA levels in the *ceh1* mutant (Xiao *et al.*, 2012), prompted us to construct a “SA biosynthesis and metabolism” pathway using Pathvisio (van Iersel *et al.*, 2008; Kutmon *et al.*, 2015). This was used to visualize *ceh1* and *ceh1/eds16* transcript, protein, and metabolite levels relative to control (Figure 3), and the data from *eds16* are shown as a separate Pathvisio output (Figure S2).

The data confirmed the previously reported enhanced levels of *ICS1* expression in *ceh1* (Xiao *et al.*, 2012). To better understand mechanism(s) involved in this regulation, we assayed transcript and protein levels of known regulators of *ICS1*. Direct transcriptional regulation of this gene has only recently begun to be elucidated (Chen *et al.*, 2009; Zhang *et al.*, 2010; Wang *et al.*, 2015; Zheng *et al.*, 2012; Van Verk *et al.*, 2011; Zheng *et al.*, 2015), and the interactions or hierarchy of *ICS1*-binding transcription factors remains to be discovered. In *ceh1*, the majority of published positive regulators of *ICS1*, namely *CBP60g*, *SARD1*, *TCP8* and *9*, *NTL9*, *CHE* and *WRKY28* are themselves upregulated, suggesting some upstream mechanisms of MEcPP-mediated induction of *ICS1*. Intriguingly, the 3 NAC domain transcription factors (ANAC019, 055 and 072) known to suppress accumulation of SA and mediate coronatine (JA-Ile mimic)-induced re-opening of stomata are also upregulated (Zheng *et al.*, 2012). This suggests priority of one or more of the *ICS1* inducers over these TFs, with implications for robustness of SA production.

Upregulation of *ICS1* in *ceh1* has been shown to result in increased levels of total SA, as measured via gas chromatography/mass spectrometry (Xiao *et al.*, 2012). Fine-tuned analyses using liquid chromatography/mass spectrometry determined that the majority of the total SA previously measured in *ceh1* is in fact conjugated SA, namely, SA glucoside (SAG, Figure 3 and Figure S2). This is consistent with previous reports showing that total SA is largely present as SAG (Vlot *et al.*, 2009), a more soluble derivative of SA facilitating vacuolar storage (Hennig *et al.*, 2002; Klessig and Malamy, 1994; Seo *et al.*, 1995). This led us to examine other known SA derivatives and their respective modifying enzymes in MEcPP-overaccumulating backgrounds, including transport form methyl SA (MeSA) (Park *et al.*, 2007), SAG and salicylate glucose ester (SGE) as SA storage forms (Song *et al.*, 2008; Dean and Delaney, 2008), the less well understood potential SA enhancers SA-SO<sub>3</sub> (Baek *et al.*, 2010) and SA-Asp (Zhang *et al.*, 2007), as well as the SA breakdown products dihydroxy benzoic acids (DHBA) (Bartsch *et al.*, 2010; Zhang *et al.*, 2013; Li *et al.*, 2014). Most of these metabolites were below detection level in our analyses, though undetectable levels of MeSA are most likely due to the volatile nature of this compound resulting in its loss during sample preparation.

The most abundant SA derivatives were the abovementioned SAG and DHBA glucosides (DHBAG, Table S3). Like SA, and consistent with previous reports (Bartsch *et al.*, 2010), DHBA was detected solely as a sugar ester. In Arabidopsis, DHBA conjugates with both glucose and xylose have been detected, with the xylose activity dominant; as such DHBAG is detected primarily in the absence of the relevant xylosyltransferase, UGT89A2 (Li *et al.*, 2014). UGT89A2 is downregulated in the *ceh1* mutant (Table S5), and interestingly, Col-0, the parent genotype for *ceh1*, typically accumulates DHBA xylosides, rather than DHBAG. The mechanism and functional consequences of this shift are not yet clear, but suggest some differential function of these two glycosides in SA breakdown or signaling. The glucosyltransferase mediating DHBAG formation is not yet known, but several candidate UDP-glucosyltransferases with activity on DHBA *in vitro* (Lim *et al.*, 2002) are upregulated in *ceh1* (Table S5).

In spite of only negligible total levels of SA in *ceh1/eds16*, the large majority of the SA conjugation and breakdown enzymes differentially regulated in *ceh1* are similarly regulated in the double mutant. This suggests that the differential regulation of these enzymes in *ceh1* is not simply the established consequence of accumulation of SA (Seo *et al.*, 1995), but rather an MEcPP-mediated specific response. The rationale and mechanism behind MEcPP inducing both SA production and breakdown/conjugation remains to be elucidated, but may be related to perturbation of other stress signals in the *ceh1* mutant, such as jasmonate signaling and redox balance.

### Misregulation of chloroplast-localized steps of jasmonate biosynthesis

The KEGG pathway “Alpha-linolenic acid metabolism,” largely composed of jasmonate biosynthesis, is among upregulated pathways in both *ceh1* and *ceh1/eds16*. Biosynthesis of jasmonic acid (JA), an oxylipin stress hormone, begins in the chloroplast with deesterification of fatty acids, followed by their oxidation and subsequent cyclization. The cyclic intermediate 12-oxophytodienoic acid (OPDA) is then transported from the chloroplast to the peroxisome to generate JA. JA itself is then transported from the peroxisome to the cytosol, where it may be conjugated to isoleucine to generate the active signaling molecule JA-Ile, or to a methyl-group to generate methyl JA (MeJA) a long-distance transportable derivative (Figure 4). OPDA, the jasmonate intermediate, has been shown to act as an independent signaling molecule, with partially overlapping responses with JA (Savchenko *et al.*, 2014). To understand regulation of jasmonate biosynthesis in *ceh1* with and without SA accumulation, we overlaid transcript, protein, and metabolite levels of jasmonate synthesis to gain insight into the network connecting MEcPP to jasmonate biosynthesis and signaling pathways, in the presence and absence of high SA levels (Figure 4). Most genes of this pathway were not differentially expressed in *eds16* (Figure S3), and thus for simplicity only differences of *ceh1* and *ceh1/eds16* relative to parent are presented in Figure 4.

We previously reported that OPDA, but not JA itself, is present at greater levels in *ceh1* and *ceh1/eds16* than parent plants (Lemos *et al.*, 2016). Here, LC-MS based metabolomics analyses presented no increase in free OPDA in *ceh1* and *ceh1/eds16*, but rather an increase in arabidopsides, in which OPDA and dinor-OPDA are esterified to glycerol backbones



(Stelmach *et al.*, 2001; Hisamatsu *et al.*, 2003; Hisamatsu *et al.*, 2005). Consistent with previous results, these arabidopsides accounted for the majority (~90%) of detected OPDA (Table S3)(Nilsson *et al.*, 2012). Arabidopsides have been found thus far only in the genus *Arabidopsis* (Mosblech *et al.*, 2009) and *Cirsium arvense* (Hartley *et al.*, 2015), where they have been shown to increase in response to wounding (Kourtchenko *et al.*, 2007) and in the hypersensitive response (Andersson *et al.*, 2006). They are suggested to act as a pool of OPDA to facilitate prolonged or rapid jasmonate response, as well as direct signaling and chemical roles (Mosblech *et al.*, 2009; Andersson *et al.*, 2006; Stelmach *et al.*, 2001). Arabidopsides have faced scrutiny for potentially being artifacts of hormone extraction techniques; although conditions known to produce these artifacts were not part of extraction (Nilsson *et al.*, 2012), the previously detected OPDA increase in *ceh1* (by GC-MS) may be partially due to contamination from arabidopside-localized OPDA, de-esterified from the glycerol backbone during sample preparation for gas chromatography.

Strong upregulation of jasmonate biosynthesis at both protein and RNA levels in *ceh1* and *ceh1/eds1* is observed in the majority of enzymes catalyzing reactions up to OPDA biosynthesis in the chloroplast (e.g., LOX and AOC family members). Inversely, of enzymes catalyzing the conversion of OPDA to JA, only the first step, OPR3, is strongly upregulated; all other steps are only minutely altered. This differential regulation, while consistent with production of arabidopsides, also presents a bottleneck that may contribute to the previously published elevated levels of OPDA. Of note, the proteins of JA biosynthesis that are differentially regulated are predominantly chloroplast localized, as opposed to the peroxisome-located steps of JA biosynthesis that are slightly, if at all altered, thus suggestive of a compartment-specific role of MEcPP in altering jasmonate biosynthesis, and thereby determining the relative levels of jasmonates species to tailor plant adaptive responses.

Finally, in addition to differential regulation of jasmonate biosynthesis, OPDA localization may also be altered. In order to proceed to JA biosynthesis, OPDA must exit the chloroplast (through a mechanism(s) not yet fully understood) and be imported into the peroxisome byperoxisomal ATP-binding cassette (ABC) transporter 1 (PXA1, (Theodoulou *et al.*, 2005), whose transcription is downregulated in the *ceh1* mutant, thus potentially contributing to OPDA accumulation. This mechanism is not inconsistent with OPDA sequestration into membranes as arabidopsides, but the effect of MEcPP must be interpreted with caution; not only is CGI58 (Park *et al.*, 2013) a PXA1-interacting protein also involved in OPDA transport upregulated in MEcPP-overaccumulating lines (Table S4), but also the downregulation of *PXA1* is significant only in *ceh1*, in the presence of elevated SA. This suggests one underlying regulatory mechanism of antagonistic cross talk between SA and JA.

Higher levels of jasmonate in *ceh1/eds16as* compared to that of the *ceh1* is a clear display of SA's functional footprint in each of the observed pathways including reduced PXA1 and enhanced OPDA biosynthesis enzyme and arabidopside accumulation as *ceh1/eds16*, allowing greater jasmonate accumulation. Finally, in accordance with previous results (Lemos *et al.*, 2016), jasmonate-responsive genes are more highly upregulated in *ceh1/eds16* than *ceh1* (Figure 4). Antagonism between SA and JA is well established, but nuances are still being revealed. Although initial reports suggested that SA inhibits JA biosynthesis

(Pena-Cortés *et al.*, 1993; Harms *et al.*, 1998), recent studies have suggested instead that SA does not affect JA biosynthesis but rather JA signaling (Van der Does *et al.*, 2013; Spoel *et al.*, 2003; Ndamukong *et al.*, 2007), and that SA/JA interactions may not be solely antagonistic, but even synergistic depending on timing and relative concentrations (Mur *et al.*, 2006). Surprisingly, our results suggest that in the context of MEcPP-induced constitutive SA and OPDA production, SA does have a dampening effect on jasmonate biosynthesis, refining our view of, and offering mechanistic insight into interplay between two key defense pathways.

### Altered Glutathione redox coupling and impact in *ceh1*

Interplay between SA and JA is partially mediated by redox homeostasis, key in SA signaling and an important component of SA responses. Several other lines of evidence suggest altered redox homeostasis in *ceh1*: the pool of glutathione in *ceh1* and *ceh1/eds16* is largely oxidized in comparison to the parent, while other reducing equivalents show the opposite trend (Figure 5A, Supplemental Table S6). Moreover, “Glutathione metabolism” was identified as an upregulated pathway in *ceh1* and *ceh1/eds16* at the RNA level. Accordingly, we investigated glutathione metabolism under elevated MEcPP.

We first utilized the KEGG “Glutathione metabolism” pathway, identifying sub-pathways for glutathione biosynthesis, oxidation/reduction, and conjugation as schematically displayed (Figure 5A), with complete glutathione-associated genes available for all assayed lines (Figure S4). The altered reduction status of the glutathione pool in MEcPP-overaccumulating genotypes is presented by comparison of oxidized glutathione (GSSG, increased levels in *ceh1/eds16* and more strongly increased in *ceh1* relative to parent) to reduced glutathione (GSH, slightly increased in *ceh1* but decreased in *ceh1/eds16* relative to parent). The GSH/GSSG ratio, although not a perfect proxy for the reduction potential of this redox couple or its compartment-specific effect on cellular redox potential (Schafer and Buettner, 2001; Diaz-Vivancos *et al.*, 2015), can suggest important changes in glutathione-mediated redox status, and the GSH/GSSG ratio in MEcPP-overaccumulating lines is only approximately 1/4 to 1/3 that of parent lines (Supplemental Table S6). These results were repeated using independent tissue and LC-MS conditions with a modified extraction, yielding similar fold changes in glutathione redox status (Method S1, Figure S5, Table S6). To investigate underlying mechanisms we first questioned whether diminished GSH biosynthesis may contribute to greater glutathione oxidation. The major regulators of glutathione synthesis are considered to be post-translationally regulated GSH1 enzyme activity and cysteine availability (Noctor *et al.*, 2012; Queval *et al.*, 2009). While GSH1 enzyme activity was not examined here and cysteine levels are not readily detected by the utilized LC-MS conditions, neither *ceh1* nor *ceh1/eds16* display altered levels of GSH1 protein, and in fact *ceh1/eds16* even has enhanced levels of the  $\gamma$ -glutamylcysteine intermediate (Figure 5A), supporting the notion that glutathione biosynthesis is not a bottle neck.

A significant difference however is observed in glutathione redox regulation. Glutathione serves many key signaling and detoxifying roles in plant cells, one of which is regenerating reduced ascorbic acid (AscA) from dehydroascorbate (dhAscA), followed by subsequent

regeneration of GSH from NADPH (Figure 5A). These reactions, catalyzed by the dehydroascorbate reductase (DHAR) and glutathione reductase (GR) gene families, provide a major reactive oxygen species (ROS) detoxification mechanism for the cell and as such these reducing currencies are generally tightly balanced (Mittler *et al.*, 2004). A possible explanation for the imbalance in *ceh1* is transcriptional regulation of *GR1* and *2*, the two differentially compartmentalized plant glutathione reductases which are both downregulated in *ceh1/eds16*, while only *GR2*, the chloroplastic isoform is downregulated in *ceh1* (Yu *et al.*, 2013; Tzafrir *et al.*, 2004; Marty *et al.*, 2009). Interestingly, a recent report has shown that the stress-induced upregulation of *GR1* and *2* is dependent on signals originating in functional chloroplasts (Garnik *et al.*, 2016), which may be disrupted *ceh1* and *ceh1/eds16*. The root cause of the observed altered GSH/GSSG ratio in MEcPP-accumulating lines is not yet clear. However, one could assume ROS as a causative agent, simply because of the evolutionarily conserved role of this signaling molecule in response to unfavorable cues (Mittler *et al.*, 2011).

Aside from direct roles as a reducing agent, glutathione serves in glutathionylation of various substrates, controlling protein redox status and function, and metabolite solubility and degradation. These functions also appear to be disrupted in *ceh1*. Among the families of glutathione-S-transferases (GSTs) and glutaredoxins (GRXs) responsible for these processes, the plant-specific tau (GSTU) and phi (GSTF) families possessed a large number of upregulated genes (Figure 5B, Figure S4). The mechanism and physiological implications of upregulation of GSTF and GSTU in *ceh1* is not immediately obvious; indeed the regulation and function of the majority of GSTs and GRXs, and specifically that of the GSTU and GSTF families, is elusive, partly due to their large and partially redundant families (Sappl *et al.*, 2009). A subset of GSTs has been demonstrated to be upregulated by SA, bacterial attack, and hydrogen peroxide (Sappl *et al.*, 2009), but there is no clear overlap between these stress-induced GSTs and GSTs induced in *ceh1*. Efforts directed at identification of GSTU and GSTF substrates have indicated roles for these family members in glutathionylating small molecules, thereby rendering them more soluble, e.g. for xenobiotic detoxification (Noctor *et al.*, 2012; Noctor *et al.*, 2011; Krajewski *et al.*, 2013). However, the individual substrate specificity *in planta*, and the relevance thereof, is not clear. Thus, potential effects of GSTU and GSTF upregulation in *ceh1* remain to be discovered. One effect may include depletion of the reduced glutathione pool, contributing to general glutathione oxidation. Glutathionylation of small molecules often leads to degradation, of both the molecule and the conjugated amino acids. Thus, upregulation of the GSTU and GSTF families may both be a response to diminished availability of reduced glutathione, and contribute to this problem by diverting reduced glutathione to eventual degradation.

### Exogenous reduced glutathione does not rescue *ceh1* phenotypes

The *ceh1* mutant is dwarfed relative to parent plants and this phenotype is not consistently rescued in *ceh1/eds16*, suggesting that although constitutive SA signaling can result in dwarfism, this is not the sole reason for compromised growth in *ceh1*. In contrast, double mutant lines of *ceh1* and the key general stress response (GSR) regulator calmodulin-binding transcriptional activator CAMTA3 (Benn *et al.*, 2014) (*ceh1/camta3*), with greatly diminished GSR, do grow to a larger size than *ceh1*, though still smaller than parent plants

(Benn *et al.*, 2016). The difference between phenotypes of *ceh1/eds16* vs. *ceh1/camta3* may lie in the specific vs. general nature of the signaling pathways – both UPR and SA responses are partially reduced in *ceh1/camta3*. Like the GSR, redox balance, and particularly glutathione, has numerous connections with stress transduction pathways disrupted in *ceh1*: redox signaling in fact induces the GSR (Bjornson *et al.*, 2014), the ER stress response both induces and is induced by redox signals (Fedoroff, 2006; Ozgur *et al.*, 2014), SA has oxidizing and reducing interactions with the redox status of the cell and with glutathione specifically, depending on timing and concentration (Herrera-Vasquez *et al.*, 2015), and redox balance of glutathione influences SA/JA cross-inhibition (Ndamukong *et al.*, 2007). Thus, redox balance and particularly glutathione redox status may represent a common pathway influencing the multitude of stress signaling pathways disrupted in *ceh1*.

To address this possibility and based on the report that application of exogenous GSH can lead to a larger and more reduced glutathione pool (Cheng *et al.*, 2015; Tausz *et al.*, 2004); we grew *ceh1* on exogenous GSH to assay for rescue of plant size (Figure 6). Because of the effects of redox on the GSR, we also assayed for the GSR in these conditions using the *4xRapid Stress Response Element:LUCIFERASE (4xRSRE:LUC)* reporter line (Walley *et al.*, 2007). The previously reported constitutive expression of *4xRSRE:LUC* in *ceh1* and *ceh1/eds16* (Benn *et al.*, 2016), was not altered in plants grown on GSH-supplemented media as measured by LUC activity driven by *RSRE* (Figure 6A). In fact, glutathione supplementation only led to significant changes in *eds16* and *ceh1/eds16*, where it actually increased luminescence. This surprising result may not be the effect of increased *4xRSRE:LUC* transcription, but rather LUC enzyme activity; indeed, LUC is more stable and active in more reduced environments (Czupryna and Tsourkas, 2011). The change in LUC activity specifically in SA deficient lines is well aligned with the interplay between this hormone and redox homeostasis, although the underlying mechanism is still unknown (Herrera-Vasquez *et al.*, 2015). Additionally, in the presence of GSH, only *ceh1/eds16* and *eds16* experience a marginal but significant increase in plant size (Figure 6B). This would also be consistent with exogenously supplied glutathione compensating for a redox imbalance in the absence of SA signaling.

The lack of an increase in *ceh1* size under glutathione supplementation suggests that one or more stress signaling pathway(s) is still active in reduced glutathione-supplemented *ceh1*, with the extension that MEcPP activates SA, JA, and GSR signaling independently of its effect on redox balance of glutathione. We cannot rule out the possibility that disruption of the MEP pathway leads to some size effect in *ceh1* independently of MEcPP signaling, but the partial rescue of plant size in *ceh1/camta3* mutants argues against this being the sole cause of the reduced size of *ceh1*. Rather, accumulation of MEcPP may result in activation of each pathway individually, or via some as-yet-undiscovered coordinating signaling node. Indeed, it has been shown that Ca<sup>2+</sup> signaling is required for exogenously applied MEcPP GSR induction (Benn *et al.*, 2016), suggesting the existence of at least one intermediary for MEcPP modulation of cellular status.

## Conclusions

Over-accumulation of MEcPP in the *ceh1* mutant leads to numerous biochemical and physiological stress response phenotypes. The interrelation and interface among stress transduction pathways prevents the drawing of irrefutable conclusions by analyses of each phenotype in isolation. Through integration of multi-omics outputs and pathway-oriented analyses, we have identified mechanisms leading to misregulation of *ICS1* and reduced conversion of OPDA to JA - a clear indication of a compartmentalized stress response cascade - as well as other perturbations, including regulation of SA breakdown and imbalances in central metabolism, amino acid levels, and redox status. Integrated analysis of these complementary omics datasets has allowed us to generate a more nuanced picture of retrograde mediated stress signaling strata, interrelation, and compartmentalization, and to identify targets of investigation in the uncharted territories of interorganellar stress signaling.

## Experimental Procedures

### Plant growth conditions

For all “-omics” experiments, *Arabidopsis thaliana* plants of the genotypes *pHPL:LUC* (parent), *ceh1*, *eds16*, and *ceh1/eds16* were grown under 16h light / 8h dark, and at 23-25°C. Three weeks old seedlings were harvested and flash-frozen in liquid nitrogen.

For *4xRSRE:LUC* visualization, seedlings were grown on ½ strength MS media, without or with 500µM glutathione. Seedlings were imaged for *4xRSRE:LUC* activity as described previously (Walley *et al.*, 2007; Bjornson *et al.*, 2014; Benn *et al.*, 2014).

### RNA-seq

Libraries for RNA-seq were prepared according to the HTR method (Kumar *et al.*, 2012), and sequenced on an Illumina HiSeq200 in 50bp single end read mode. Reads were mapped to the Arabidopsis genome using TopHat2 (Kim *et al.*, 2013) on the public usegalaxy.org instance of Galaxy (Afgan *et al.*, 2016) (counts by gene and statistical analyses available in Table S4). RNAseq data is available through the Gene Expression Omnibus GSE96666.

### Proteomics

Proteomics sample and data preparation has been described (Walley *et al.*, 2015).

### Metabolomics

Metabolites extraction and profiling from five biological replicates per genotype was as previously reported (Balcke *et al.*, 2011; Treutler *et al.*, 2016), with additional alterations described in Supporting Information Method S1. MS conditions and MRM transitions for the targeted approach are presented in Supporting Information Table S7. MS<sup>1</sup> and MS<sup>2</sup> raw spectra were deconvoluted using MS-DIAL (Tsugawa *et al.*, 2015). Families of altered metabolites were identified using the web application MetFamily (Treutler *et al.*, 2016), and levels of selected metabolites were quantified using the software Multiquant 3.0 (Sciex).

## Gene Set Enrichment Analysis

Gene set enrichment analysis (GSEA) was carried out using the R (Team, 2016) package Generally Applicable Gene set Enrichment analysis (GAGE), available through Bioconductor (Luo *et al.*, 2009). For gene sets we used the KEGG database of *Arabidopsis thaliana* metabolic pathways (Ogata *et al.*, 1999; Kanehisa *et al.*, 2016).

Analysis was carried out for RNA and protein. Briefly, for RNA counts were normalized using DESeq2 (Love *et al.*, 2014), and the normalized counts were used as input for GAGE.

For proteins, a similar procedure was followed, using iTRAQ intensities (Walley *et al.*, 2015) and paired biological replicates. Due to the smaller dynamic range of protein measurements, uncorrected p values were occasionally considered when providing support to RNA-seq results, as noted in Table S1.

## Heatmap

Assayed metabolites were first sorted according to KEGG pathway/molecular category, then heat maps were created using the ggplot2 (Wickham, 2009) package in R. For consistency, limits of log<sub>2</sub> fold change -2 (blue) to 2 (yellow) were used throughout the heatmap and pathway visualizations, although numerical values are available in Tables S3, S4, and S6.

## Pathway visualization

With the exception of SA biosynthesis and breakdown, pathway visualizations are based on KEGG (Ogata *et al.*, 1999; Kanehisa *et al.*, 2016) pathway templates. Relevant steps and Arabidopsis family members were highlighted and expanded in custom-made Pathvisio (Kutmon *et al.*, 2015; van Iersel *et al.*, 2008) pathways, with metabolite data converted to fold change and corrected p values calculated in R.

## Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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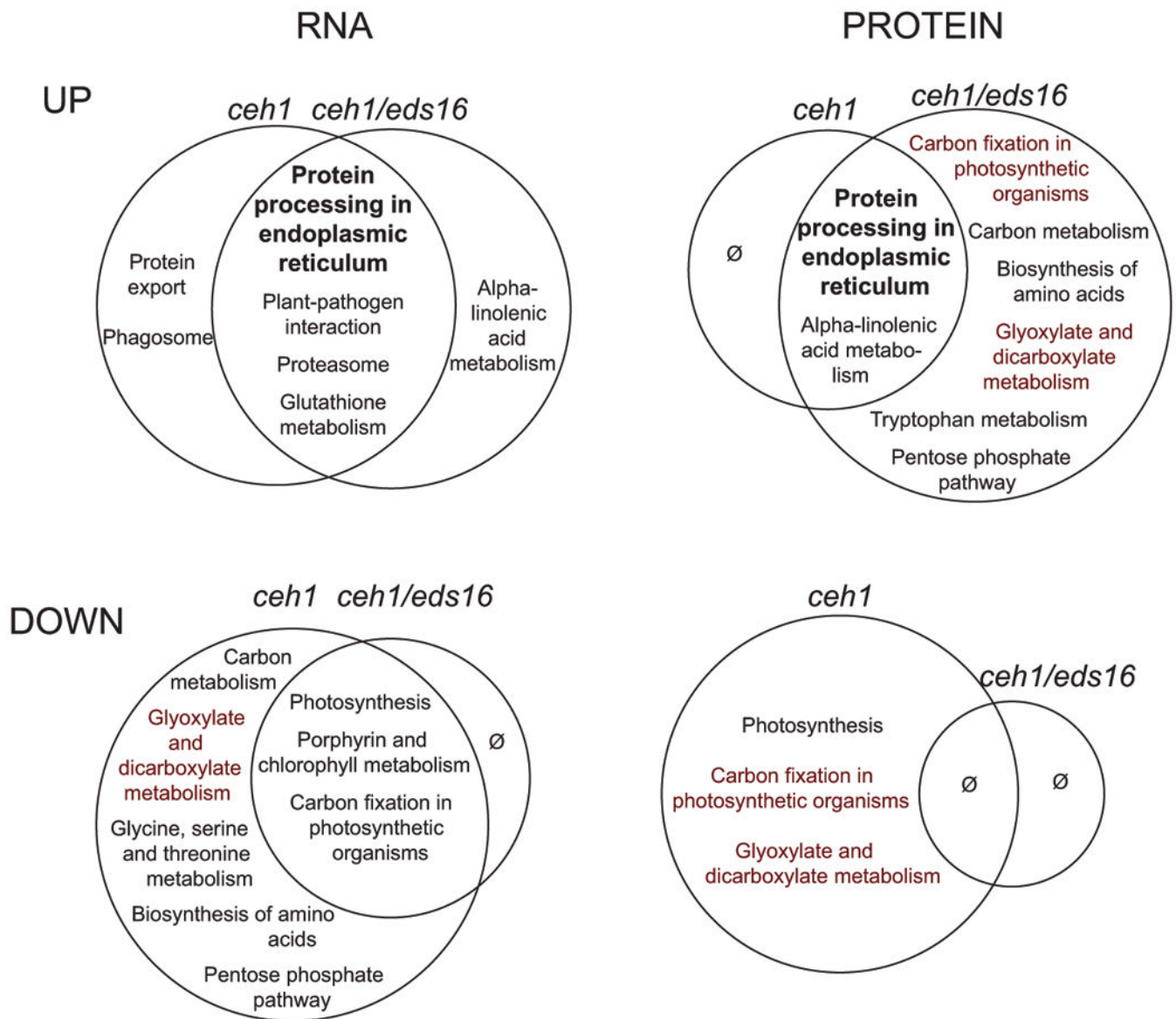
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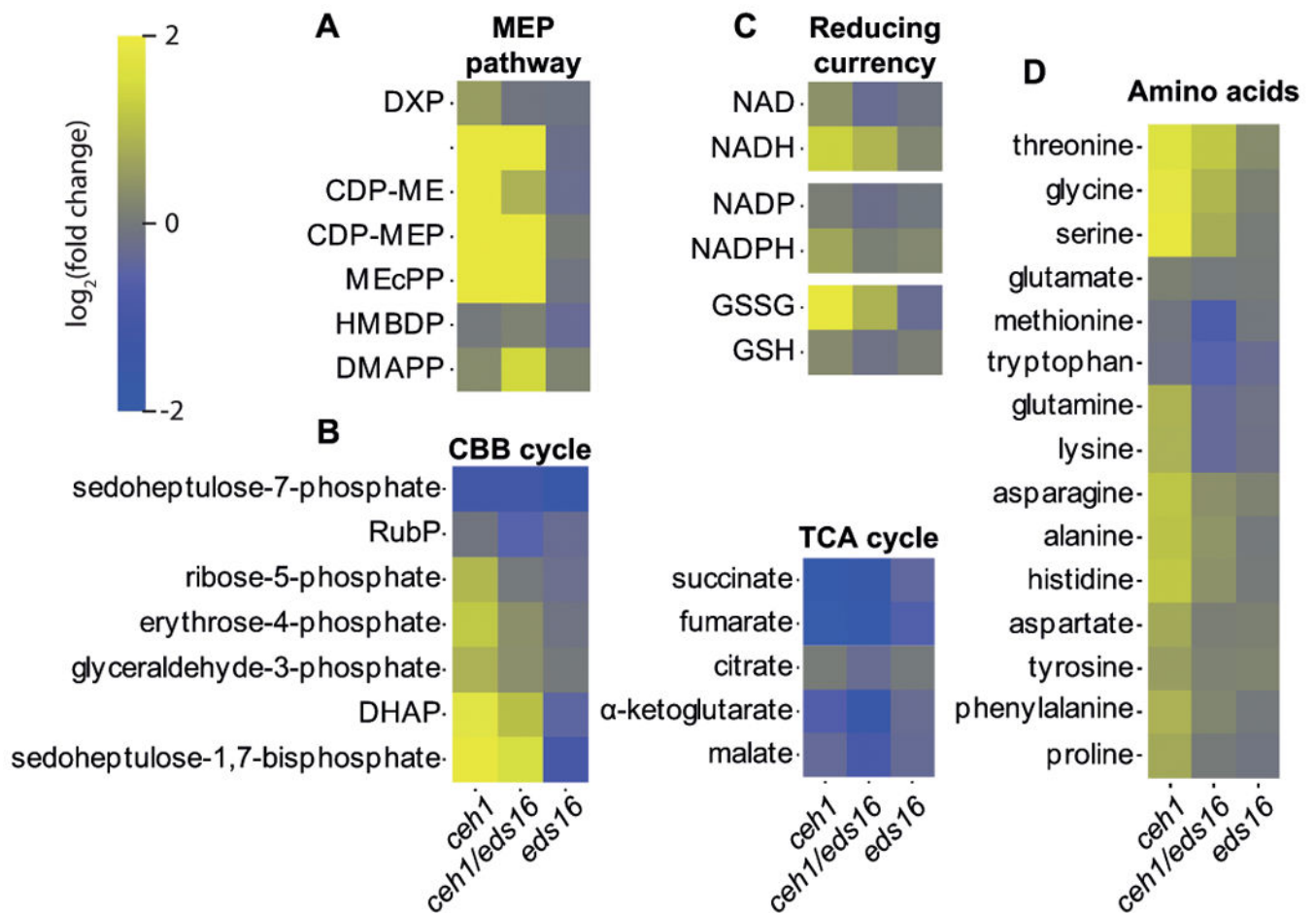
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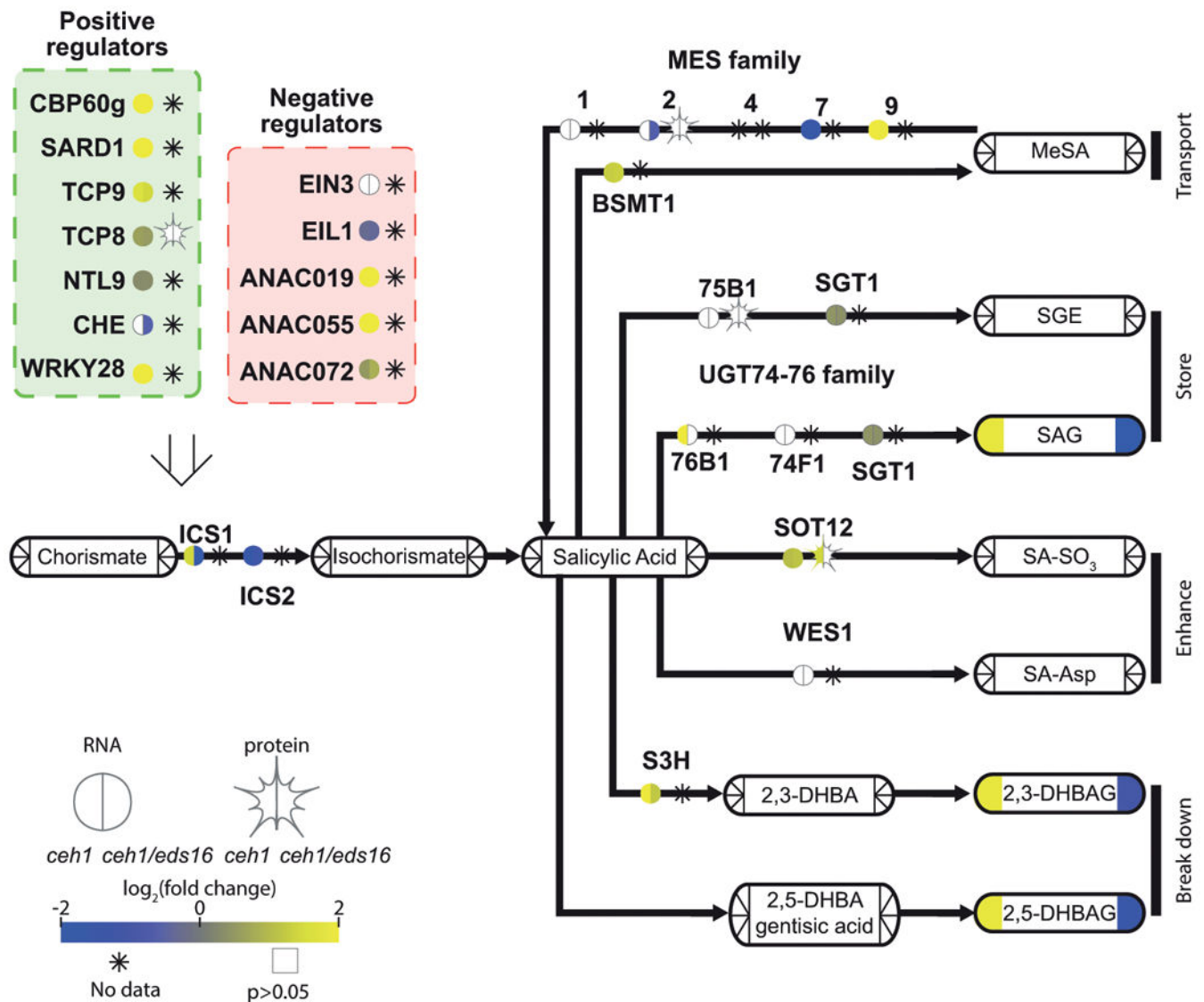
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**Figure 1. Differentially regulated pathways in *ceh1* and *ceh1/eds16* relative to parent line** Depicted modulated pathways that are up (upper row) or down (lower row) regulated in *ceh1* (left side of Venn diagrams) and *ceh1/eds16* (right side of Venn diagrams) were identified. For RNAseq data (left column) pathways with an FDR-corrected p value of less than 0.05 are shown. For proteomics data, due to lower dynamic range, uncorrected p values were used as a less reliable indicator of significance, particularly when in concordance with transcriptomic data. Pathways in bold are those similarly regulated at RNA and protein level in both displayed genotypes, while pathways in red are inversely regulated between *ceh1* and *ceh1/eds16*. ∅ indicates no significantly differentially regulated pathways.



**Figure 2. Selected metabolite sets accumulating differentially among *ceh1*, *ceh1/eds16*, and *eds16*** (A) MEP pathway intermediates, (B) central metabolism related Calvin Benson Bassham (CBB) and tricarboxylic acid (TCA) cycle metabolites, (C) Reducing currencies and (D) amino acids differentially accumulate in the displayed genotypes relative to parent lines. Shading represents  $\log_2$ -fold change for each metabolite (rows) relative to wild type for each genotype (columns), from -2 (darkest) to 2 (lightest). Abbreviations: DXP: Deoxy-xylulose phosphate, MEP: methyl-erythritol phosphate, CDP-ME: diphosphocytidyl-methylerythritol, CDP-MEP: diphosphocytidyl-methylerythritol phosphate, MEcPP: methyl-erythritol cyclodiphosphate, HMBDP: hydroxymethylbutenyl pyrophosphate, DMAPP: dimethylallyl pyrophosphate, RubP: ribulose 1,5 bisphosphate, DHAP: dihydroxyacetone phosphate, NAD: nicotinamide adenine dinucleotide, NADP: nicotinamide adenine dinucleotide phosphate.



**Figure 3. Salicylic acid biosynthesis and degradation is regulated by MEcPP**

A pathway schematic for SA biosynthesis and breakdown, with transcription factors/enzymes shown by bare text with a circle, representing RNA, and star, representing protein levels. Metabolites are shown by rounded rectangles. For metabolites, RNA, and protein, the log<sub>2</sub>-fold change for *ceh1* vs parent (left) and *ceh1/eds16* vs. parent (right) is represented by shading, from -2 (darkest) to 2 (lightest). Non-detectable products in each section are shown by an asterisk, while empty symbols with grey borders represent non-significant fold changes relative to parent. Arrows represent chemical conversions, catalyzed by the enzymes shown on the line. The open arrow from transcription factors to ICS1 represents direct transcriptional regulation of this enzyme by these transcription factors. Abbreviations: CBP60g: calmodulin-binding protein 60g, SARD1: systemic acquired resistance deficient 1, TCP8/9: teosinte branched 1, cycloidea, PCF domain containing 8/9, NTL9: NTM1-like 9, CHE: CCA1 hiking expedition, EIN3: ethylene-insensitive 3, EIL1: EIN3-like1, ANAC019,055,072: Arabidopsis Nam/ATAF1,CUC2-domain containing TF 19/55/72,

ICS1/2: isochorismate synthase  $\frac{1}{2}$ , MES: methyl esterase, BSMT: SA methyl transferase, MeSA: methyl-salicylic acid, UGT: UDP-glucose transferase, SGT: SA glucose transferase, SGE: salicylate glucose ester, SAG: salicylic acid glucoside, SOT12: sulphotransferase 12, SA-SO<sub>3</sub>: sulphosalicylic acid, WES1: =GH3.5 WESO1, S3H: salicylate-3-hydroxylase, DHBA: dihydroxybenzoic acid, DHBAG: DHBA glucoside

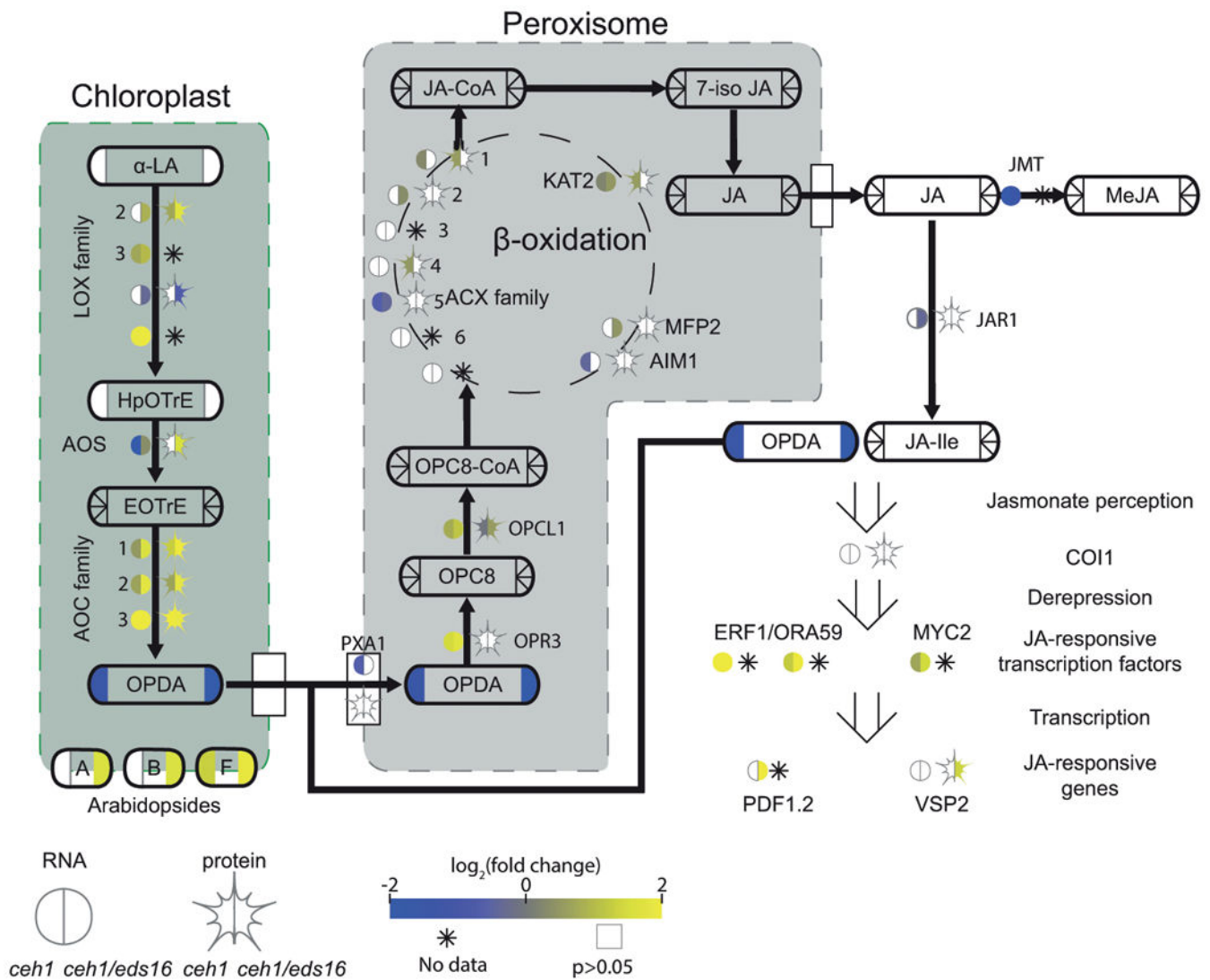
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**Figure 4. MECPP and SA alter jasmonate biosynthesis and signaling**

A pathway schematic for JA biosynthesis and breakdown, with transcription factors/enzymes shown by bare text with a circle, representing RNA, and star, representing protein levels. Metabolites are shown by rounded rectangles. For metabolites, RNA, and protein, the  $\log_2$ -fold change for *ceh1* vs parent (left) and *ceh1/eds16* vs. parent (right) is represented by shading, from -2 (darkest) to 2 (lightest). Non-detectable products in each section are shown by an asterisk, while empty symbols with grey borders represent non-significant fold changes relative to parent. Arrows represent chemical conversions, catalyzed by the enzymes shown on the line, or metabolite transport. Peroxisome-localized steps in conversion of 12-OPDA to JA are shown in the grey box, while simplified chloroplast-localized steps leading to 12-OPDA synthesis are in the green. Open arrows represent metabolite/protein, protein/protein, and protein/DNA (transcription regulation) interactions. Abbreviations:  $\alpha$ -LA:  $\alpha$ -linolenic acid, LOX: lipoxygenase, HpOTrE: hydroperoxyoctadecatrienoic acid, AOS: allene oxide synthase, EOTrE: epoxyoctadecatrienoic acid, AOC: allene oxide cyclase, OPDA: oxophytodienoic acid, PXA1: peroxisomal ABC transporter 1, OPR3: OPDA reductase 1,

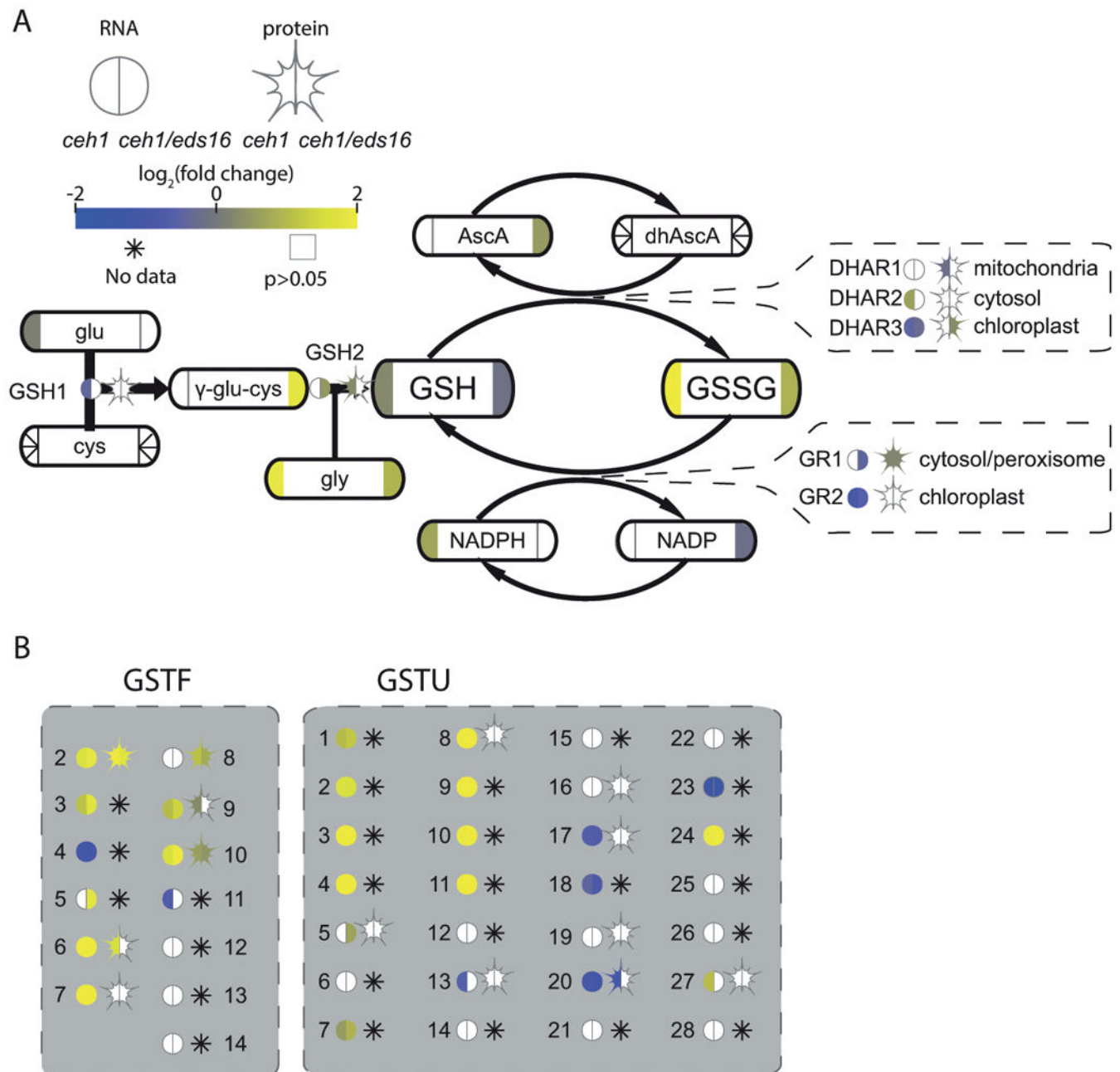
OPC8: oxopentenylcyclopentenyl octanoate, OPCL1: OPC8 coA ligase1, ACX: acyl-coA oxidase, KAT2: ketoacyl-coA thiolase 2, MFP2: multifunctional protein 2, AIM1: abnormal inflorescence meristem 1, MeJA: methyl-JA, COI1: coronatine insensitive 1, ERF1: ethylene response factor 1, ORA59: octadecanoid-responsive Arabidopsis AP2/ERF 59, MYC2: (=JIN1) MYC-domain containing transcription factor, PDF1.2: plant defending 1.2, VSP2: vegetative storage protein 2.

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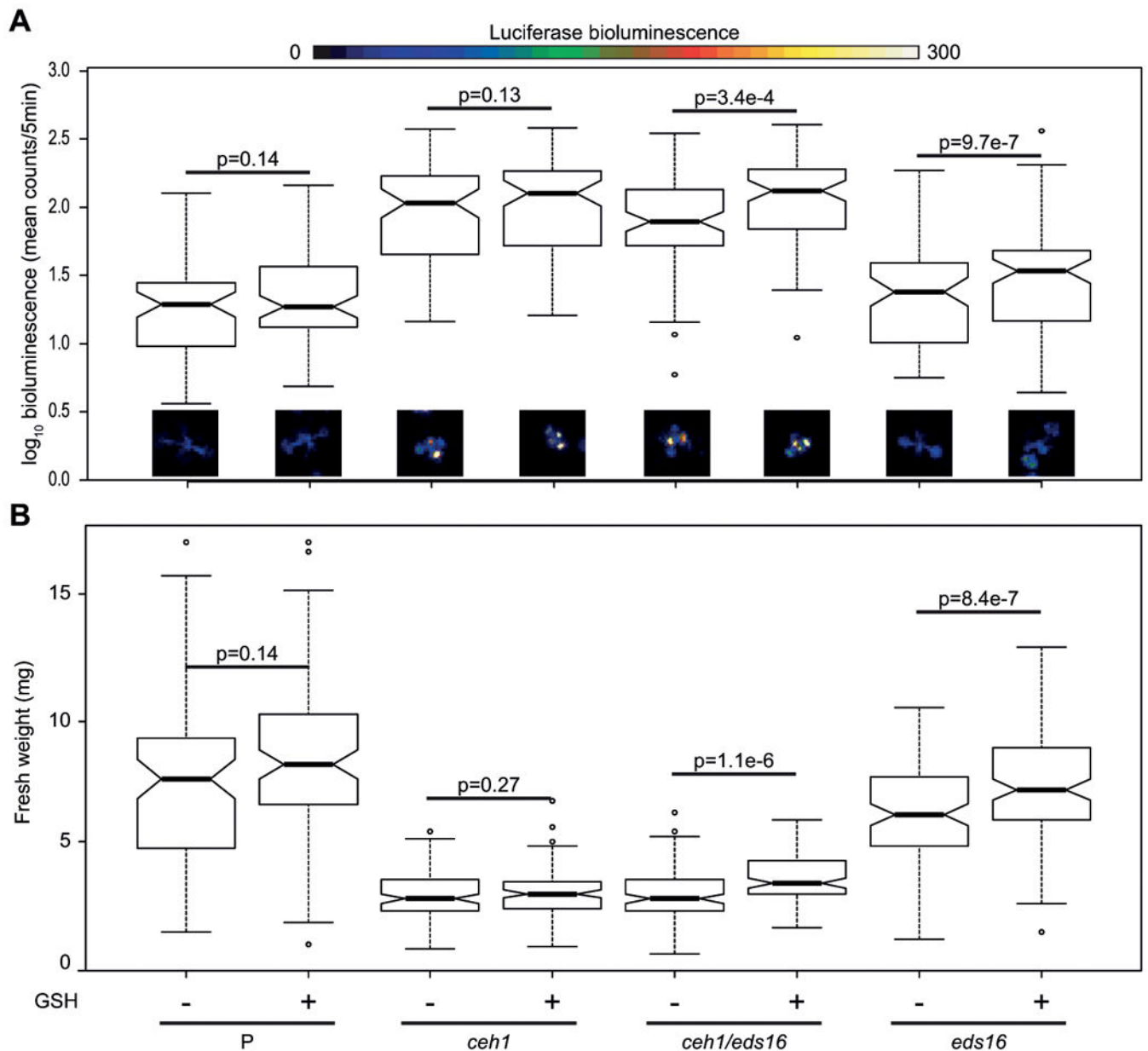
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**Figure 5. MEcPP influences glutathione redox balance and function**

(A) A partial simplified pathway schematic for glutathione biosynthesis and function, with enzymes shown by bare text with a circle, representing RNA, and star, representing protein levels. Metabolites are shown by rounded rectangles. For metabolites, RNA, and protein, the  $\log_2$ -fold change for *ceh1* vs parent (left) and *ceh1/eds16* vs. parent (right) is represented by shading, from -2 (darkest) to 2 (lightest). Non-detectable products in each section are shown by an asterisk, while empty symbols with grey borders represent non-significant fold changes relative to parent. Arrows represent chemical conversions, catalyzed by the enzymes shown on the line. Callouts for the DHAR and GR enzyme families also show subcellular

localization of these enzymes. **(B)** Glutathione S-transferases of the phi (GSTF) and tau (GSTU) families are shown in grey boxes. Abbreviations: glu: glutamate, cys: cysteine, GSH1: (=γ-ECS1, RML1, PAD2) glutathione synthetase 1, gly: glycine, GSH2: glutathione synthetase 2, GSH: reduced glutathione, GSSG: oxidized glutathione, AscA: ascorbic acid, dhAscA: dehydroxyascorbic acid, DHAR: dehydroxyascorbic acid reductase, GR: glutathione reductase, GSTF: glutathione S-transferase phi family, GSTU: glutathione S-transferase tau family.



**Figure 6. Exogenous glutathione does not rescue GSR or size phenotypes of *ceh1***

Parent (P), *ceh1*, *ceh1/eds16*, and *eds16* seedlings were grown in media with or without reduced glutathione supplementation. The GSR, as measured via 4xRSRE:LUC activity (A) was examined in each line, represented by notched boxplots of quantified luminescence ( $\log_{10}$  transformed to achieve normality) and by false-color images, with a scale bar above showing conversion from luminescence to color. Notched box plots show possible outliers (circles), the interquartile range (box), 95% confidence interval (notch), and median. Non-overlapping notches is considered strong evidence for statistically different populations. Following imaging, seedlings were harvested and weighed (B). For each genotype, p values for comparisons between (+) and (-) glutathione were calculated using a one-way ANOVA,

blocking by experimental replicate in the statistical program R,  $n > 75$  seedlings per genotype/treatment combination over three experiments.

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