

# Interplay between Jasmonic Acid, Phosphate Signaling and the Regulation of Glycerolipid Homeostasis in Arabidopsis

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## Title

Interplay between Jasmonic Acid, Phosphate Signaling and the Regulation of Glycerolipid Homeostasis in Arabidopsis

#### Heading

Jasmonic acid control of glycerolipid homeostasis

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#### Abstract.

Jasmonic acid (JA) biosynthesis and signaling are activated in Arabidopsis cultivated in phosphate (Pi) deprived conditions. This activation occurs mainly in photosynthetic tissues and is less important in roots. In leaves, the enhanced biosynthesis of JA coincides with membrane glycerolipid remodeling triggered by the lack of Pi. We addressed the possible role of JA on the dynamics and magnitude of glycerolipid remodeling in response to Pi-deprivation and resupply. Based on combined analyses of gene expression, JA biosynthesis and glycerolipid remodeling in wild type Arabidopsis and in the coi1-16 mutant, JA signaling seems important in the determination of the basal levels of phosphatidylcholine (PC), phosphatidic acid (PA), monogalactosyldiacylglycerol (MGDG) and digalactosyldiacylglycerol (DGDG). JA impact on MGDG steady state level and fluctuations seem contradictory. In the coi1-16 mutant, the steady state level of MGDG is higher, possibly due to a higher level of PA in the mutant, activating MGD1, and to an increased expression of MGD3. These results support a possible impact of JA in limiting the overall content of this lipid. Concerning lipid variations, upon Pi-deprivation, JA seems rather associated with a specific MGDG increase. Following Pi-resupply, whereas the expression of glycerolipid remodeling genes returns to basal level, JA biosynthesis and signaling genes are still upregulated, likely due to a JA-induced positive feedback remaining active. Distinct impacts on enzymes synthesizing MGDG, *i.e.* downregulating MGD3, possibly activating MGD1 expression and limiting the activation of MGD1 via PA, might allow JA playing a role in a sophisticated fine tuning of galactolipid variations.

## Keywords

glycerolipid remodeling, jasmonic acid, MGD1, MGD3, phosphate signaling.

#### Introduction.

Plants have to cope with frequent variations of nutrients in soils. Phosphorus, in the form of soluble inorganic phosphate (Pi), is one of the most limiting elements (Rellan-Alvarez et al. 2016). The response to Pi starvation has been extensively studied in *Arabidopsis thaliana*, a plant assimilating Pi mainly via PHT1 transporters (Mitsukawa et al. 1997; Muchhal et al. 1997; Nussaume et al. 2011; Okumura et al. 1998). A lack of Pi triggers changes at multiple organization levels and time scales, ranging from early biochemical and metabolic tuning of Pi allocation, to genome scale transcription reprogramming, leading to an integrated metabolic, physiological and developmental response (Chiou and Lin 2011; Plaxton and Tran 2011; Yang and Finnegan 2010; Zhang et al. 2014). Primary long term phenotypic changes include: the secretion of phosphatases in the soil to scavenge Pi from organic sources, an enhanced expression of Pi transporters, a reduced growth of primary root and shoot, an enhanced growth of lateral roots and root hairs, an accumulation of anthocyanins and an intense remodeling of membrane glycerolipids, saving Pi from phospholipids (Peret et al. 2011).

The major early effector in low Pi signaling is the transcription factor PHR1 (Phosphate starvation response 1) (Bustos et al. 2010; Misson et al. 2005; Misson et al. 2004; Rubio et al. 2001; Thibaud et al. 2010). Concerning the control of Pi incorporation, PHR1 triggers the expression of miR399 in the shoot of *Arabidopsis*, and this microRNA moves to the root where it represses the expression of *PHO2*, encoding a ubiquitin ligase involved in the degradation of PHO1 (Lin et al. 2008; Liu et al. 2014b; Pant et al. 2008) and of several members of Phosphate transporters PHT1 family (Huang et al. 2013). The PHO1 Pi transporter then accumulates in the root xylem parenchyma, and operates in the transfer of Pi up to the shoot (Hamburger et al. 2002; Liu et al. 2012).

A later remodeling of lipids, marked by a replacement of phospholipids by Pi-free plastid glycolipids, is also under the control of PHR1 (Bustos et al. 2010; Pant et al. 2015). In brief, a phospholipid breakdown is triggered in non-plastidial membranes, mostly at the level of phosphatidylcholine (PC), involving the NPC4 and NPC5 phospholipases C (Gaude et al. 2008; Nakamura et al. 2005) and the PLD<sup>1</sup> and PLD<sup>2</sup> phospholipases D (Cruz-Ramirez et al. 2006). The mechanisms involved in the decline of phosphatidylglycerol (PG) in the plastid is unknown. Concomitantly, the syntheses of phosphorus-free sulfoquinovosyldiacylglycerol (SQDG), monogalactosyldiacylglycerol (MGDG) and digalactosyldiacylglycerol (DGDG) increase in plastids (Awai et al. 2001; Boudiere et al. 2014; Dormann and Benning 2002; Essigmann et al. 1998; Hartel and Benning 2000; Jouhet et al. 2004; Jouhet et al. 2003; Jouhet et al. 2007; Yu et al. 2002). In the thylakoids, a net PG-to-SQDG replacement occurs (Essigmann et al. 1998), whereas DGDG is exported from the plastid and relocates to the plasma membrane (Andersson et al. 2003) and tonoplast (Andersson et al. 2005), where a PC-to-DGDG

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replacement occurs. DGDG also relocates to mitochondria (Jouhet et al. 2004; Michaud et al. 2016). Genes involved in this lipid remodeling, i.e. the SQDG synthase (*SQD2*), two MGDG synthase isoforms (*MGD2* and *MGD3*) and DGDG synthases (*DGD1* and *DGD2*), are therefore markers of the low Pi response and of the remodeling of lipids.

It was recently shown that Pi deficiency also induced the biosynthesis of jasmonic acid (JA) and its derivative JA-Isoleucine (JA-Ile), therefore activating the JA signaling pathway (Khan et al. 2016). The biosynthesis of JA relies on the supply of alpha-linolenic acid (C18:3) derived from MGDG and DGDG, converted into oxo-phytodienoic acyls (OPDA) via a plastid lipoxygenase/allene oxide synthase (LOX/AOS) pathway (Andreou et al. 2009). OPDA is exported to the peroxisome and then serves as precursor for the synthesis of JA (Andreou et al. 2009; Wasternack and Hause 2013). Based on qRT-PCR studies, a lack of Pi induced the expression of gene markers of JA biosynthesis (lipoxygenase 2, LOX2) and signaling (jasmonate zim-domain protein 10, JAZ10) (Khan et al. 2016). A puzzling question is that, although multiple studies had been performed to analyze the transcriptomic response of roots and leaves of Arabidopsis to low Pi (Bustos et al. 2010; Hammond et al. 2003; Misson et al. 2005; Morcuende et al. 2007; Muller et al. 2007; Woo et al. 2012), only one reported a change in the expression of genes involved in JA biosynthesis and signaling (Morcuende et al. 2007). It was also shown that JAZ10 was initially not induced in the phr1-1 mutant, but reached the wild type induction level after 10 days on Pi-deficient medium, thus indicating an early (but partial) control by PHR1. The pho1-7 mutation (Khan et al. 2016) was introgressed into Arabidopsis lines impaired in JA biosynthesis (the *aos* mutant defective for AOS (Park et al. 2002)) and JA signaling (the *coi1-34* mutant affecting the level of the COI1, an F-box protein interacting with JA-IIe (Acosta et al. 2013; Yan et al. 2013; Yan et al. 2009)). In the obtained pho1-7 aos and pho1-7 coi1-34 double mutants, the expression of the low-Pi responsive gene MGD3 was significantly reduced compared to the pho1-7 parent. These experiments suggest that JA could influence MGD3 expression, either by a transcriptional activation of MGD3 that would be additive with the upregulation in response to low Pi, or by amplifying the magnitude of MGD3 upregulation by low-Pi. Other studies support a link between galactolipid and JA levels. Consistently with the low Pi-response, Methyl-JA was shown to upregulate MGD1, in a context of increased DGDG production (Taki et al. 2005). By contrast, a mutant with 90% reduction of DGDG level due to the impairement of DGD1 also shows a JA overproduction (Lin et al. 2016). Altogether, these studies show that different lipid remodeling patterns, with opposite variations of galactolipid levels, coincide with an increase in JA, suggesting that a sophisticated system of regulation by this oxylipin might operate.

In the present study, the transcriptomic changes induced by Pi deprivation and resupply were reexamined, refining the kinetics of gene expression responses. Previous reports showing that JA biosynthesis and signaling were activated upon Pi starvation were confirmed (Khan et al. 2016), but

showed that the expression of corresponding genes following Pi-resupply was not necessarily tuned down back to basal levels. We also confirmed that the activation of JA biosynthesis was different in roots and shoots (Khan et al. 2016), correlating with different patterns of upregulation of JA biosynthesis and signaling genes. We then attempted to refine our understanding of the interplay between low-Pi response, JA signaling and the control of glycerolipid homeostasis.

**Results and discussion.** 

# Transcriptional response to phosphate variations reveals different patterns of genes involved in jasmonic acid metabolism and signaling in different organs.

*Arabidopsis* seedlings were grown on modified media supplemented with either 500 μM KH<sub>2</sub>PO4 (High Pi or "HPi") or 200 μM KCl (Low Pi or "LPi"). For the analysis of Pi deprivation and resupply, seeds were cultivated in LPi conditions for 7 days and then transferred for 30 min, 1 h or 3 h either to LPi (LPi-LPi; "Pi-deprived") or HPi (LPi-HPi; "Pi-resupply"). Plants cultivated in HPi conditions for 7 days before transfer onto fresh HPi medium (HPi-HPi; "Pi-supplemented") acted as controls (Fig. 1). Roots and shoots of *Arabidopsis* grown under these Pi regimen were carefully cut and collected using a razor blade, RNA was extracted and used to generate RNA-seq libraries. We thus obtained gene expression profiles in roots (Supplementary Table S1) and shoots (Supplementary Table S2) of *Arabidopsis* for each Pi regimen.

Genes that were differentially expressed in Pi-supplemented and Pi-deprived conditions were determined. To that purpose, compiled RNA-seq data was obtained from 0.5, 1 and 3 hours following transfers from HPi to HPi (Pi-supplemented) and LPi to LPi (Pi-deprived), respectively, and used to compare transcript abundance in shoots and roots. Using the non-parametric NOISeq method (Zheng and Moriyama 2013), only genes with a stable expression for each Pi concentration could be considered, based on the NOISeq p-value cutoff set at 0.05. Differentially expressed genes were then selected based on a Log-fold-change ratio threshold, i.e. |Log2FC| > 1.

In roots, 494 genes were down-regulated and 879 were up-regulated in low-Pi condition (Supplementary Table S3). In the list of upregulated genes, we sought enriched GO terms and functional annotations using the DAVID method (<u>http://david.abcc.ncifcrf.gov</u>) (Huang et al. 2007). Consistently with known effects of Pi deprivation, terms corresponding to phosphatase (P-value =  $1.8.10^{-8}$ ), phosphate ion transport (P-value =  $3.8.10^{-10}$ ), glycerophospholipid catabolic process (P-value =  $3.3.10^{-3}$ ), galactolipid biosynthetic process (P-value =  $1.5.10^{-5}$ ), phosphatidylcholine 1-acylhydrolase activity (P-value =  $8.1.10^{-2}$ ), cell wall biogenesis/degradation (P-value =  $2.6.10^{-1}$ ) or transcription regulation (P-value =  $5.4.10^{-1}$ ) were enriched. The term corresponding to cytochrome P450 proteins was also enriched (P-value =  $8.1.10^{-4}$ ) although we could not determine the precise pathway(s) in which these CYP proteins could be involved. In details, glycerolipids marker genes known to be upregulated in low Pi (Misson et al. 2005; Morcuende et al. 2007) were confirmed, i.e. *MGD2* (AT5G20410; Log2FC = 4.1), *MGD3* (AT2G11810; Log2FC = 6.6), *DGD1* (AT3G11670; Log2FC = 1.4), *DGD2* (AT4G00550; Log2FC = 1.8), *SQD2* (At5g01220; Log2FC = 4.3), *PLD*, (AT3G05630; Log2FC = 4.5) and *NPC4* (AT3G03530; Log2FC = 4.5) (Supplementary Table S3).

In shoots, 505 genes were down-regulated and 1,215 genes were up-regulated (Supplementary Table S4). We also sought enriched GO terms and functional annotations using the DAVID method in upregulated genes. As above, terms corresponding to acid phosphatase (P-value =  $3.9.10^{-10}$ ), protein phosphatase 2C (P-value =  $5.5.10^{-1}$ ), phosphate ion transport (P-value =  $2.10^{-6}$ ), cellular phosphate ion homeostasis (P-value =  $7.1.10^{-7}$ ), glycerophospholipid catabolic process (P-value =  $6.2.10^{-3}$ ), galactolipid biosynthetic process (P-value =  $1.1.10^{-3}$ ), P-type ATPase (P-value =  $2.8.10^{-1}$ ), cell wall modification (P-value =  $8.6.10^{-1}$ ) or transcription regulation (P-value =  $5.2.10^{-1}$ ) were enriched, confirming past analyses. By contrast with roots, additional terms corresponding to oxylipin biosynthetic process (P-value =  $9.7.10^{-2}$ ), together with cytochrome P450 (P-value =  $3.2.10^{-2}$ ) and secondary metabolites biosynthesis, transport, and catabolism (P-value =  $7.2.10^{-2}$ ) were also identified. Glycerolipids marker genes upregulated in low Pi (Misson et al. 2005; Morcuende et al. 2007) were also confirmed, i.e. *MGD2* (Log2FC = 4.6), *MGD3* (Log2FC = 8.6), *DGD2* (Log2FC = 1.3), *SQD2* (Log2FC = 4.7), *PLD* $\zeta$ 2 (Log2FC = 5.4) and *NPC4* (Log2FC = 5.7) (Supplementary Table S4). Transcriptional changes in shoots appeared therefore to involve oxylipins and, overall the magnitude of upregulation of marker genes was higher compared with roots.

Overall, this comparison of Arabidopsis grown in Pi-supplemented vs. Pi-deprived conditions validates the study dataset. The enrichment of GO terms corresponding to oxylipins in the comparison performed with green tissues supports therefore the possible activation of Jasmonic acid biosynthesis pathway reported earlier (Khan et al. 2016), at least in shoots.

We then focused our analysis on the expression of genes involved for JA biosynthesis (AOS, AOC1, AOC2, AOC3, LOX2, LOX4, OPR3, JAR1, ACS1, ACS2, ACX1, AIM1, MFP2, KAT2, KAT5 and JMT), JA signaling (FT AP2/ERF, JAZ1 to JAZ10, COI1), and as a control, glycerolipid remodeling (NPC4, NPC5, PLD<sup>2</sup>, PLD<sup>2</sup>, MGD1, MGD2, MGD3, DGD1, DGD2). Differential expression was measured in roots and shoots, 0.5 h after medium transfer, and highlights the same marker genes as those detected after the non-parametric comparison using all gene expression levels measured 0.5, 1 and 3 hours following medium transfer (Table 1). Concerning JA synthesis and signaling, AOS (in shoots), AOC1 (in both roots and shoots), AOC2 and AOC3 (in shoots), LOX2 (in roots and to some extent in shoots), LOX4 (about two-fold increase in both roots and shoots), JAZ5 (in roots) and JAZ7 (in shoots) appeared upregulated in low Pi condition. Some modulated responses could therefore be observed between roots and shoots, but in both cases, genes involved in JA biosynthesis and signaling were significantly upregulated in response to Pi deprivation. Interestingly about half of the selected genes coding for JA biosynthesis or signaling components contained in their promoter region, one or more Pi-deprivation responsive elements (P1BS box 1, a known PHR1 binding element (Rubio et al. 2001)) (Table 1).

This comparison supports the fact that subsets of genes involved in JA biosynthesis might be coregulated with genes involved in glycerolipid remodeling in response to Pi availability. We wondered whether these genes could be tuned down back to control level upon addition of Pi. The design of our experiment allowed investigating whether genes could be finely tuned following such resupply. To determine if these genes returned to basal expression levels after resupply differentially expressed genes and GO enrichment following Pi-resupply were determined, comparing the expression levels after 0.5, 1 and 3 hours in shoots. To that purpose, a partition of differentially expressed genes was performed using a K-mean method, with the number of partitions set to 10 (Dolch et al. 2017; Liu et al. 2014a). Each cluster consisted of genes with similar expression patterns (i.e. expression curves after 0.5, 1 and 3 hours) following Pi-resupply, with representative nearest mean curves, serving as prototypes for the clusters (shown in Fig. 2). Three clusters comprise genes upregulated following resupply, i.e. Cluster 2 consisting of genes with a strong and regular expression increase  $(Log_{2FC} > 2)$ , Cluster 4 with genes mainly upregulated 3 h following Pi-resupply and Cluster 5 with genes highly upregulated 1 h following Pi-resupply. Three clusters comprise genes downregulated following Piresupply, i.e. Cluster 7 consisting of genes with moderate but significant expression decline, Cluster 8 with genes mainly downregulated after 1 h resupply, and Cluster 10, a very large cluster containing genes with a regular expression decrease. This later cluster comprises genes with moderate to strong variation levels (Log2FC < -2). Based on GO term enrichment using two independent methods, GOseq (P-value <  $5.10^{-2}$ ) and DAVID (P-value  $\leq 1.10^{-2}$ ), we analyzed these clusters to detect possible biological processes (BP) or molecular functions (MF) with a correlated dynamics of gene expression following Pi-resupply (Fig. 2). GO terms enriched in Cluster 10 highlight genes involved in the maintenance and repair of photosynthetic machinery, chloroplast division, cell mitosis and leaf morphogenesis, illustrating that the developmental reorientation in shoots does not occur within a short period following Pi resupply, but should rather require a longer time to return back to their level in control conditions.

GO terms corresponding to BP Jasmonic acid mediated signaling pathway, BP Regulation of jasmonic acid mediated signaling or BP Response to jasmonic acid were found enriched in Clusters 2, 4 and 5, *i.e.* showing therefore an upregulation upon Pi-resupply, whereas GO terms corresponding to BP Phosphate ion homeostasis, BP Phosphorus metabolic process, BP Lipid storage, BP Sulfolipid biosynthetic process, BP Galactolipid metabolic process, MF 1,2-diacylglycerol 3-beta-galactosyltransferase activity (i.e. synthesis of MGDG) were mainly found enriched in Cluster 8, showing a restoration of control expression level for these Pi-responsive genes (Fig. 2).

This analysis highlights that although genes involved in glycerolipid remodeling and JA metabolism or signaling are upregulated when plants are deprived of Pi, they appear to have uncoupled patterns of

expression following Pi-resupply. We addressed therefore the following questions: could JA signaling be involved in the dynamics and/or magnitude of the glycerolipid remodeling triggered by Pideprivation? Could JA signaling be involved in a reverse glycerolipid remodeling, returning back to initial state following Pi-resuply?

# Jasmonic acid biosynthesis and signaling and glycerolipid remodeling are coupled following Pi deprivation and uncoupled following Pi resupply

The expression of some representative genes was examined via qRT-PCR (Fig. 3, white bars) in the leaves of wild type Col-0 Arabidopsis grown under different Pi regimen.

- When comparing Pi-supplemented and Pi-deprived conditions, genes involved in JA biosynthesis were upregulated (*LOX2*, *AOC1*, *AOC2*) or unchanged (*AOS*). The selected reporter gene for JA signaling, *JAZ10*, was consistently upregulated in Pi-deprived condition. As expected, all representative genes involved in glycerolipid reprogramming were upregulated (*NPC4*, *PLD*, *Z*, *MGD2*, *MGD3*, *DGD1*, *DGD2*, *SQD2*). The expression of *MGD1* increased slightly.
- Upon Pi-resupply, genes involved in glycerolipid reprogramming were tuned down. Only MGD1 expression was enhanced. In contrast, all genes involved in JA biosynthesis and response were upregulated by the refeeding with Pi.

Previous reports have clearly demonstrated the production of JA and JA-IIe in *Arabidopsis* deprived of Pi, with different levels in roots and shoots (Khan et al. 2016). We sought whether the regulation of JA biosynthesis genes in the various Pi-regimen analyzed here, was also correlated with changes in JA levels in planta. To that purpose, we used an *Arabidopsis* transgenic line expressing the  $\beta$ -glucuronidase (GUS) reporter gene, fused with *JAZ1* under the cauliflower mosaic virus 35S promoter, p35S::JAZ1-GUS (Thines et al. 2007). Based on the interaction of JA-IIe with COI1, a component of the Skp1/Cullin/F-box SCF<sup>COI1</sup> ubiquitin E3 ligase complex (Feys et al. 1994; Xie et al. 1998), JAZ1 is degraded. In this reporter system, an *in vivo* production of JA ad JA-IIe is therefore detected by a loss of GUS staining. The p35S::JAZ1-GUS transgenic line was cultivated under various phosphate regimen and the histochemical analysis was consistent with a low JA content in both roots and shoots in Pi-supplemented condition (Fig. 4, A), a production of JA mainly localized in the stem and leaves in Pi-deprived condition (Fig. 4, B) and no apparent decrease in JA content following Pi-resupply in leaves (Fig. 4, C). *In planta*, the level of JA appears therefore consistent with the variations of JA biosynthesis and response genes (Fig. 3, white bars).

We also sought whether the very rapid down-regulation of genes involved in glycerolipid remodeling (within hours following Pi-resupply) had an effect on a restoration of the control lipidomic profile. Using wild type samples collected in parallel, we extracted and analyzed glycerolipids 3 hours following a resupply in Pi (Fig. 5, Col 0). We compared the profiles with that obtained in Pi-supplemented, used as a control. In photosynthetic tissues, plants grown in Pi-deprived medium exhibited the expected increase in SQDG, concomitant with a decrease in PG, consistent with the well-established SQDG-to-PG replacement in thylakoid membranes. In shoots, MGDG level increased slightly, DGDG level nearly doubled, whereas that of phospholipids and particularly PC decreased, consistently with the DGDG-to-PC exchange triggered by Pi deprivation. In roots, although lipid contents where much lower and more technically challenging to quantify, variations of SQDG, PG and MGDG were not observed in these nongreen tissues, decrease in phospholipids were not statistically significant, whereas an increase of DGDG was clearly induced in Pi-deprived condition. In both roots and shoots, within a day following Piresupply, no significant change could be observed. Contrary to the down tuning of genes involved in glycerolipid remodeling (Fig. 3), these analyses did not support any glycerolipid remodeling back to control level rapidly after a resupply in Pi.

Taken together, these analyses show that JA biosynthesis and signaling genes are upregulated in Pideprived condition compared to Pi-supplemented condition, however, following Pi-resupply, expression of these genes is even more activated. Jasmonic acid presence in Pi-deprived condition and following a resupply in Pi is confirmed *in planta*. It is likely that JA and JA-derivatives accumulated upon Pi deprivation might maintain an upregulation of JA biosynthesis genes even though Pi signaling is attenuated, following the well-known positive feedback of the JA pathway, namely JA-induced JA biosynthesis (Sasaki et al. 2001). Only MGD1 expression followed the trend of JA biosynthesis and signaling genes, suggesting that this gene might be related to this interplay between JA signaling and responses to Pi variations.

# Comparison of the gene expression reprogramming and the lipid remodeling in the WT and coil genetic backgrounds.

To confirm the link between JA signaling and glycerolipid remodeling induced by Pi variations, we considered a genetic background impaired in JA signaling. In previous works, the expression of the low-Pi responsive *MGD3* gene had been analyzed in the *pho1-7 aos* and *pho1-7 coi1-34* double mutants: its expression was significantly reduced compared to the *pho1-7* parent, suggesting that JA might control lipid remodeling. However, the increase of SQDG, MGDG and DGDG relatively to phospholipids was unchanged in *pho1-7 aos* and *pho1-7 coi1-34* double mutants compared to *pho1-7* (Khan et al. 2016). Here, we used a *coi1-16* mutant, impaired at the level of the COI1 dependent JA perception, subjected to various Pi regimen, to consider all possible Pi-responsive pathways. The *coi1-16* background is also known to contain a mutation in the *PEN2* gene (Westphal et al. 2008), a glycosyl hydrolase involved in the synthesis of antifungal glucosinolates (Bednarek et al. 2009). The *coi1-16* 

mutant was previously used to unravel the interplay between JA and the response to potassium variations, with no effect which could be attributed to the *pen2* mutation (Armengaud et al. 2010). No biotic stress was exerted in the present study and only genes related to the *COI1* role and lipidome remodeling were examined. We compared gene expression and glycerolipid profiles in WT (Col-0) plants and *coi1-16* lines (Fig. 3, black bars and Fig. 5).

In the *coi1-16* mutant, the expression of most genes involved in JA biosynthesis or signaling was consistently reduced compared to the wild type, and was unaltered by variations in Pi, besides a very low upregulation observed for *JAZ1* or *JAZ10*. Interestingly, the expression of *AOC1* follows a pattern of a Pi-responsive gene in the *coi1-16* background (Fig. 3, black bars) although no P1BS box could be detected in its promoter region (Table 1). This suggests that expression of *AOC1* is controlled in response to Pi-availability by an unknown pathway and that JA masks this control by enhancing *AOC1* expression.

Genes involved in Pi-dependent glycerolipid remodeling are differentially expressed in the coi1-16 background (Fig. 3, black bars). PLD $\zeta_2$  and MGD2 show little changes. The overall expression of NPC4, MGD3 and DGD1 and to some extent of DGD2 and SQD2 is increased compared to WT, in particular in Pi-deprived condition, suggesting a negative control by JA. *MGD1* seems to show a slightly opposite response to Pi variations in coi1-16 compared to Col-0, with a downregulated expression in Pi-deprived condition, unchanged after Pi-resupply (Fig. 3). However, this slight decrease observed in multiple analyses was not statistically significant based on a Dunett's test. MGD3 expression is significantly activated in low-Pi in both WT and coi1-16, with a magnitude of upregulation moderated by JA. By contrast with previous analyses in the pho1-7 coi1-34 double mutants and pho1-7 parent (Khan et al. 2016), data shown in Fig. 3 suggest that the basal level of MGD3 expression is increased in the coi1-16 mutant when compared to Col-0. Although we have no clear explanation, this apparent discrepancy might be due to the *pho1-7* mutation, impairing phosphate assimilation constitutively. In this previous work, the decreased expression in MGD3 was apparently contradictory with the observed increase in galactolipids (Khan et al. 2016); here the level of expression of MGD3 (together with an increased level of PA, a known activator of MGD1, see below) is consistent with the higher level of MGDG in the coi1-16 mutant. Altogether, these results suggest that JA exerts a sophisticated tuning of the expression of genes involved in glycerolipid homeostasis, regulating genes controlling phospholipid homeostasis in the endomembrane system (NPC4) and galactolipid metabolism in chloroplasts (MGD3, DGD1, and to some extent MGD1, DGD2 and SQD2).

We sought whether these results were consistent with previous analyses of transcriptome change in Arabidopsis subjected to JA. In a recent comprehensive study, a whole-genome transcriptional

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expression analysis was performed based on RNA extracted from leaves over 14 consecutive time points within 16 h following application of methyl JA (Hickman et al. 2017). Expression patterns of differentially expressed genes were clustered in 27 groups, 1 to 14 exhibiting an increased expression whereas 15 to 27, a downregulation. In this dataset, *MGD1* belonged to cluster 8 (consistent with the tendency observed in Fig. 3) and *MGD3* to cluster 16 (supporting the higher expression level we observed in the *coi1-16* mutant). This confirms the results reported on Figure 3 with a balanced control of MGDG synthesizing genes by JA, via opposite effects on MGD1 and MGD3.

When analyzing glycerolipids in the coi1-16 mutant exposed to various Pi regimen, the most striking differences are observed in leaves at the levels of MGDG, DGDG, PC and phosphatidic acid (PA) (Fig. 5, A). Firstly, the basal level of PC appears lower in *coi1-16* and responds to Pi-deprivation like the WT. PC is the sole lipid showing a strong modification at its fatty acid composition level (Fig. 6). In this analysis, diacyls are expressed as the sum of carbon contained in the two fatty acid and the number of double bonds they harbor. PC-34 corresponds mainly to PC containing one fatty acid with 16 carbons and one fatty acid with 18 carbons and PC-36 corresponds to PC containing two 18-carbon fatty acids. In the coi1-16, PC-34-2 relative proportion increases, whereas that of PC-36-3 and PC-36-4 decrease, suggesting that molecular species enriched in fatty acids with 18 carbons are responsible for the global decrease in PC. Secondly, PA level is significantly higher in the coi1-16 background. For this lipid intermediate, no change in fatty acid content is detected and no variation is observed in response to Pi. These data do not provide any evidence for the origin of this pool of PA, should it derive from a subpool or PC or another source. The subcellular distribution of this PA pool and its generation machinery being unknown, we could not speculate on the mechanism of its production in response to JA. It is known that PA can reach the chloroplast and activate galactolipid synthesis (Benning 2009; Botella et al. 2016; Dubots et al. 2012). Thirdly, and consistently with a high PA level, MGDG content appears higher than in the wild type, and remains unchanged regardless of Pi variations. DGDG level is twice as high in *coi1-16* as that in Col-0, and is even more increased in response to Pi-deprivation. The fatty acyl profiles in MGDG and DGDG showed little changes, except a slight increase in 18:3/18:3 in DGDG in roots (from 56.4% of total DGDG fatty acyls in Col-0 to 64.5% in coi16-1), reflecting an important contribution of MGD1 in most conditions in the synthesis of the bulk of galactolipids and a visible impact of the action of MGD3 in root non-photosynthetic plastids. It has been previously reported that even when MGD3 was overexpressed, basal level of MGDG could show no change (Murakawa et al. 2014). Here, the increase in MGDG in the *coi16-1* background appears therefore most importantly due to MGD1 activation by PA.

Altogether these results highlight that more mechanisms are involved, rather than just a regulation by gene transcription, in the complex modulation of lipid homeostasis by JA. Interestingly, all mechanisms dissected here converge toward a control of galactolipid level.

#### Conclusion

We confirmed previous reports showing that JA biosynthesis and signaling are activated in Arabidopsis photosynthetic tissues when plants are cultivated in Pi-deprived conditions. Given the well-known role of JA in response to insects, this discovery was initially evaluated as a possible link between Pi deficiency and an enhanced herbivory resistance. JA could however act on other cellular responses, as shown here. In leaves, activation of JA coincides with a glycerolipid remodeling triggered by the lack of Pi. Concerning a possible action of JA on glycerolipid remodeling, our analysis of the coi1-16 mutant is consistent with a control of the basal levels of MGDG, which is higher in this mutant possibly via an increased production of PA (an activator of MGDG synthesis) and by an increased expression of MGD3. This supports a role of JA in lowering MGDG production, at least by inhibiting MGD3 expression. By contrast, upon Pi-deprivation, a role of JA in MGDG increase seems to occur, possibly by a moderate stimulation of the expression of MGD1, which needs to be confirmed in the future. When adding Pi to deprived plants, whereas glycerolipid remodeling genes are tuned back to normal, JA biosynthesis and signaling genes are even more activated. Our study shows that in this apparently reverse condition, JA also plays a role, and is therefore uncoupled from the low-Pi response. As a preliminary investigation of mechanisms involved, the effect of JA on two MGDG synthesis enzymes, *i.e.* a control of the activation of MGD1 (via an increase in PA) and a repression of MGD3 expression, might contribute to a fine tuning of MGDG variations in either Pi-deprivation or Pi-resupply. The existence of this balanced control of MGDG synthesis is supported by the complete loss of MGDG tuning in response to Pi variations in the coi1-16 and is consistent with previous whole gene expression analyses. JA interplay with plant response to Pi variations is therefore complex, highlighting a fine tuning of the magnitude of gene expression levels in response to Pi. In the future, important questions need to be addressed, including the understanding of the lag between the rapid restoration of glycerolipid remodeling gene expression following Pi-resupply and the actual restoration of the glycerolipid profile; the elucidation of the mechanisms involved in the control of glycerolipid homeostasis by JA; and the evaluation of a possible involvement of JA in other glycerolipid remodeling processes occurring in response to other abiotic and biotic stresses and in relation with other hormonal controls.

## Material and Methods.

#### Plant material and growth conditions.

All Arabidopsis thaliana lines used in this study were in the Col-0 ecotype. The coi1-16 mutant and p35S::JAZ1-GUS (Thines et al. 2007) transgenic lines were provided by Dr Anthony Champion (IRD Montpellier). For all experiments plants were grown on modified Murashige and Skoog (MS) medium (Arnaud et al. 2014) supplemented with 2  $\mu$ M FeCl<sub>2</sub> and either 500  $\mu$ M KH<sub>2</sub>PO4 (High Pi or "HPi") or 200 µM KCl (Low Pi or "LPi"). For the analysis of Pi starvation and resupply, seeds were cultivated in LPi conditions for 7 days and then transferred for 30 min, 1 hour or 3 hours either on LPi (LPi-LPi; "Pideprived") or HPi (LPi-HPi; "Pi-resupply"). The untreated control for these experiments was performed by growing plants on HPi conditions for 7 days before transferring them onto fresh HPi medium conditions during the appropriate control time (HPi-HPi; "Pi-supplemented"). Cultivations were performed on solid or liquid medium, as indicated (Kanno et al. 2016). For cultivation on agar plates, Low Pi Plant Agar (SIGMA A1296) was added to the supplemented MS medium at 0.8%. Seedlings were stratified in darkness (4°C, 24 hours), and grown in a 16-h-light/8-h-dark photoperiod at 21-24°C, in white light in vertical plates. For experiments using GUS reporters, seedlings were grown in liquid culture (24 wells falcon sterile plates, 3 mL MS medium per well, 10-15 seeds per well). Transfers were performed by moving directly the seedlings into the new medium. For lipids analyses and transcript level measurements, seedlings were germinated on stripes of Sefar Nitex 100 µM (30 - 40 seeds per stripe) placed into square plates of solid agar medium, and grown vertically. Transfers were performed by moving the Nitex stripes on new medium.

#### RNA extraction and RT-qPCR analyses

At least 30 individual seedlings were harvested per condition and stored in liquid nitrogen. A minimum of three biological replicates were collected per genotype/treatment. RNA was then isolated using the RNeasy Plant Mini Kit (Qiagen), following the manufacturer's instructions. Possible traces of DNA were eliminated with the RNase-Free DNase Set (Qiagen). 1 µg of RNA was used to obtain complementary DNA (cDNA) with the QuantiTect Reverse Transcription Kit (Quiagen). The quantitative polymerase chain reactions (qPCR) were carried out using SsoAdvanced<sup>™</sup> Universal SYBR® Green Supermix (Biorad) and Biorad CFX96 Real-time PCR Detection System, according to the manufacturer's instructions. Three technical replicates were analyzed for each biological replicate. The TUB2 gene (AT1G75785) was used as reference. Primer pairs (forward, fwd and reverse, rev) used for amplification were the following: MGD1 (AT4G31780) fwd, TGGTTCGAGTCTCCGTAGGT and rev, CAAAGCTCTCCGGAACCACT; MGD2 (AT5G20410) fwd, ACAAGAAATTGGCATCTGCAT and rev, CATCAGAGGATGCACGCTAA; DGD1 (AT3G11670) fwd, ACGGTGAAGATGCAGTCGAGT and rev, CCACAAACTTCCCCATGGCT; DGD2 (AT4G00550) fwd, ACGGTGAAGATGCAGTCGAGTCGAG and rev, TTTCCCATCGCCAAGGCTTCTG; SQD2 (At5G01220) fwd, TACCTGAAGCTCGGATTGCT and rev, TGTGAGAGTTCATCGCCTTG; PLDZ2

(AT3G05630) fwd, TCACGACAAGCAAGAACAGGTTAG and rev, AGTGCAGAGGAAGAGAGCACCATC; NPC4 (AT3G03530) fwd, TCCAAACCCGGGTCATCCTA and rev, GTTCATAACCGCGGAGGACA; AOC1 (AT3G25760) fwd, ACTCCTACTCGAGCTCTCTCTCAG and rev, GTTCTTGAACTTTGCTTGGTCTGG; AOC2 (AT3G25770) fwd, ACTGGAGCCTAGCGGAGTTA and rev, ACACAGCGATACGAGAAACAT; AOS (AT5G42650) fwd, GGTGGCGAGGTTGTTTGTGATTG and rev, TTCCTAACGGCGACGTACCAAC; LOX2 (AT2G18790) fwd, CAAGGATGCTGGCCTCTTAC and rev, TCGTCTCGTAACCATGAAAATC; LOX4 (AT1G72520) fwd, GGAAGACCACATCATCGGTCAAC and rev, AAACGGTTCGTCTCTAACGCTTG; JAZ1 (AT1G19180) fwd, AGCTTCACTTCACCGGTTCTTGGA and rev, TCTTGTCTTGAAGCAACGTCGTCA; JAZ10 (AT5G13220) fwd, TCGCAAGGAGAAAGTCACTGCAAC and rev, CGATTTAGCAACGACGAAGAAGAC; TUB2 (AT5G62690) fwd, GAGCCTTACAAGG.

#### **RNA-seq analyses**

Roots and shoots of Arabidopsis grown under various Pi regimen were carefully cut and collected using a razor blade and RNA was extracted as described above. RNA quality and integrity were determined using the Nanodrop 1000 Spectrophotometer and Agilent Bioanalyser. Only high-quality RNA samples (Abs<sub>260/280 nm</sub> ratios of 2.0–2.1) were used for RNA-seq library generation with the Illumina Truseq Stranded Total RNA sample prep kit. RNA-seq libraries were multiplexed and loaded per lane into the Illumina HiSeq flow cell v3. All sequencing protocols were carried out as per the manufacter's instructions using the Illumina HiSeq 1000 and HiSeq control software. RNA-seq reads were analysed using the Cufflinks package (Trapnell et al. 2012), version 2.1.1 and mapped onto the Arabidopsis TAIR10 genome (Kim et al. 2015) with the TAIR10 transcriptome annotation, using the DESeq2 method (Love et al. 2014; Varet et al. 2016). Clustering was achieved based on expression profiles as described previously (Dolch et al. 2017). In brief, partition of differentially expressed genes was performed using a K-mean method, with a number of partitions set to 6 and a clustering based on a Euclidian distance (Liu et al. 2014a). For each group we sought whether gene ontology (GO) terms could be enriched, either by the DAVID method (http://david.abcc.ncifcrf.gov) (Huang et al. 2007), using the corresponding Refseq gene IDs and with a default p-value threshold of 0.1, or using the GOseq R package (Young et al. 2010) with an identical p-value threshold. Based on GO enriched terms, a focused analysis of acyl-lipid and oxylipin pathways was performed. Illumina reads of all samples have been submitted to the Sequence Read Archive at the National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov/sra) under accession number SRP133280.

#### Lipid extraction and analyses.

Roots and shoots of Arabidopsis grown under various Pi regimen were carefully cut and collected using a razor blade and a minimum of 100 µg fresh weight were stored in liquid nitrogen (3 biological replicates). Samples were lyophilized and lipids were extracted by the Folch method (Folch et al. 1957). Total glycerolipids were quantified from their fatty acids: in an aliquot fraction of extracted lipids, a known quantity of 15:0 was added and the fatty acids were converted into methyl esters (FAME) by a 1 hour incubation in 3 mL 2.5%  $H_2SO_4$  in pure methanol at 100°C (Jouhet et al. 2003). The reaction was stopped by addition of 3 mL water and 3 mL hexane. The hexane phase was analyzed by a gas chromatography-flame ionization detector (GC-FID) (Perkin Elmer) on a BPX70 (SGE) column. FAME were identified by comparison of their retention times with those of standards (Sigma) and quantified by the surface peak method using 15:0 for calibration. For quantification of lipid classes by highperformance liquid chromatography coupled to tandem mass spectrometry (LC-MS/MS), fractions of extracted lipids corresponding to 25 nmol were suspended in 100  $\mu$ L of chloroform/methanol 2:1, (v/v) containing 125 pmol of internal standards and analyzed as previously described (Jouhet et al. 2017). Internal standards were either obtained from Avanti Polar Lipids Inc. (for diacylglycerol, DAG 18:0-22:6; phosphatidylcholine, РС 18:0-18:0; phosphatidylethanolamine, ΡE 18:0-18:0; phosphatidylinositol, PI 18:0-18:0; phosphatidylserine, PS 18:0-18:0; phosphatidylglycerol, PG 18:0-18:0; phosphatidic acid, PA 18:0-18:0 and diphophatidylglycerol, DPG 14:0-14:0-14:0-14:0), synthesized by D. Lafont (Amara et al. 2010; Amara et al. 2009) (for galactolipids, MGDG 18:0-18:0 and DGDG 16:0-16:0) or purified from spinach thylakoid (Deme et al. 2014) and hydrogenated (Buseman et al;2006) (for sulfoquinovosyldiacylglycerol, SQDG 16:0-18:0). The HPLC separation method was adapted from (Rainteau et al. 2012). Lipid classes were separated using an Agilent 1200 HPLC system using a 150 mm×3 mm (length × internal diameter) 5 μm diol column (Macherey-Nagel), at 40°C. The mobile phases consisted of hexane/isopropanol/water/ammonium acetate 1 M, pH 5.3 [625/350/24/1, (v/v/v/v)] (A) and isopropanol/water/ammonium acetate 1M, pH 5.3 [850/149/1, (v/v/v)] (B). The injection volume was 20  $\mu$ L. After 5 min, the percentage of B was increased linearly from 0% to 100% in 30 min and stayed at 100% for 15 min. This elution sequence was followed by a return to 100% A in 5 min and an equilibration for 20 min with 100% A before the next injection, leading to a total runtime of 70 min. The flow rate of the mobile phase was 200 µL/min. The distinct glycerolipid classes were eluted successively as a function of the polar head group. Mass spectrometric analysis was done on a 6460 triple quadrupole mass spectrometer (Agilent) equipped with a Jet stream electrospray ion source under following settings: drying gas heater, 260°C; drying gas flow 13 L.min<sup>-1</sup>; sheath gas heater, 300°C; sheath gas flow; 11 L.min<sup>-1</sup>; nebulizer pressure, 25 psi; capillary voltage, ± 5000 V; nozzle voltage, ± 1000. Nitrogen was used as collision gas. The quadrupoles Q1 and Q3 were operated at widest and unit resolution respectively. PC analysis was carried out in positive ion mode by scanning for precursors of m/z 184 at collision energy (CE) of 34 eV. SQDG analysis was carried out

in negative ion mode by scanning for precursors of m/z -225 at a CE of -56eV. PE, PI, PS, PG, PA, MGDG and DGDG measurements were performed in positive ion mode by scanning for neutral losses of 141 Da, 277 Da, 185 Da, 189 Da, 115 Da, 179 Da and 341 Da at CEs of 20 eV, 12 eV, 20 eV, 16 eV, 16 eV, 8 eV and 8 eV, respectively. Quantification was done by multiple reaction monitoring (MRM) with 30 ms dwell time. DAG and TAG species were identified and quantified by MRM as singly charged ions [M+NH4]+ at a CE of 16 and 22 eV respectively with 30 ms dwell time. DPG species were quantified by MRM as singly charged ions [M-H]<sup>-</sup> at a CE of -45 eV with 50 ms dwell time. Mass spectra were processed by MassHunter Workstation software (Agilent) for identification and quantification of lipids. Lipid amounts (pmol) were corrected for response differences between internal standards and endogenous lipids and by comparison with a quality control (QC). QC extract corresponds to an Arabidopsis lipid extract previously qualified and quantified by thin layer chromatography and gas chromatography coupled to ion flame detection (Jouhet et al. 2017).

#### GUS activity analyzes.

An *Arabidopsis* transgenic line expressing the β-glucuronidase (GUS) reporter gene, fused with *JAZ1* under the cauliflower mosaic virus 35S promoter, p35S::JAZ1-GUS (Thines et al. 2007) was cultivated under various phosphate regimen. Seedlings were prefixed, immediately after collection in ice-cold 90% acetone for 20 min on ice, then rinsed with cold water for 5 min, vacuum infiltrated for 10 min on ice with staining solution (50 mM sodium phosphate buffer pH 7.0, 0.2% Triton-X-100, 10 mM potassium ferrocyanide, 10 mM potassium ferricyanide, 1 mM X-Gluc) and incubated at 37°C in the dark for the indicated time. Samples were then cleared by progressive dehydration through ethanol series up to 100% and progressively re-hydrated prior to observation. Imaging was then performed using an Olympus SZX12 binocular microscope (with WHS-10X magnification system) equipped with a Nikon DXM1200C digital camera.

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**Table 1. Differential expression of genes involved in jasmonic acid biosynthesis and signaling in Pi-starved versus Pi-replete conditions.** In the "Pi-supplemented" conditions, plants were grown for 7 days on HPi medium and then transferred to a fresh HPi medium. In the "Pi-deprived" condition, plants were grown for 7 days on LPi medium and then transferred to a fresh LPi medium. After 7 days, plants were transferred on an identical medium and then collected for RNA extraction. Based on whole genome RNA seq data, the differential expression of selected genes was determined in both shoots and roots and expressed in Log2FC. Genes involved in gene remodeling triggered by Pi deprivations, known to be upregulated, are given as internal control.

Short description     P1BS box I     Shoots Log2FC     Roots       JA biosynthesis	Log2FC	
JA biosynthesis		
AOS     AT5G42650     allene oxide synthase     0     2.03587     -0.3	00299	
AOC1     AT3G25760     allene oxide cyclase 1     0     2.84213     2.5	<u>6893</u>	
AOC2 AT3G25770 allene oxide cyclase 2 1 <u>1.04551</u> 0.55	50625	
AOC3 AT3G25780 allene oxide cyclase 3 0 <u>1.04178</u> -0.3	11482	
LOX2 AT3G45140 lipoxygenase 2 1 0.940651 <u>1.5</u>	<u>3607</u>	
LOX4 AT1G72520 lipoxygenase 4 2 0.958026 0.77	73372	
OPR3     AT2G06050     oxophytodienoate-reductase 3     1     0.757211     0.3	0015	
JAR1 AT2G46370 Auxin-responsive GH3 family protein 0 0.740317 0.08	41236	
ACS1 AT4G05160 AMP-dependent synthetase and ligase family protein 0 0.642895 0.14	15448	
ACS2 AT5G63380 AMP-dependent synthetase and ligase family protein 0 0.420248 -0.3	30739	
ACX1 AT4G16760 acyl-CoA oxidase 1 0 0.409011 0.42	13446	
AIM1 AT4G29010 enoyl-CoA hydratase/isomerase family 1 0.408754 0.22	21613	
MFP2     AT3G06860     multifunctional protein 2     0     0.246093     0.36	6468	
KAT2     AT2G33150     peroxisomal 3-ketoacyl-CoA thiolase 3     3     0.200535     0.20	)1582	
KAT5     AT5G48880     peroxisomal 3-keto-acyl-CoA thiolase 2     1     0.190344     0.44	10682	
JMT AT1G19640 jasmonic acid carboxyl methyltransferase 0 -0.465981 -0.1	71726	
JA signaling		
FT AP2/ERF AT3G50260 cooperatively regulated by ethylene and jasmonate 1 1 1.46763 -0.1	24066	
JAZ1 AT1G19180 jasmonate-zim-domain protein 1 2 0.818662 -0.2	02647	
JAZ2 AT1G74950 jasmonate-zim-domain protein 2 0 0.323852 -0.02	55306	
JAZ3 AT3G17860 jasmonate-zim-domain protein 3 2 0.726571 0.12	23497	
JAZ4 AT1G48500 jasmonate-zim-domain protein 4 1 0.45042 0.19	96468	
JAZ5 AT1G17380 jasmonate-zim-domain protein 5 0 <u>1.00279</u> -0.5	15517	
JAZ6 AT1G72450 jasmonate-zim-domain protein 6 1 0.353224 -0.1	81203	
JAZ7 AT2G34600 jasmonate-zim-domain protein 7 / 0 0.314164 <u>1.2</u>	<u>6975</u>	
JAZ8 AT1G30135 jasmonate-zim-domain protein 8 / _ 0 0.428381 0.46	55424	
JAZ9 AT1G70700 jasmonate-zim-domain protein 9 2 -0.264564 -0.5	40653	
JAZ10 AT5G13220 jasmonate-zim-domain protein 10 0 -0.160352 0.3	39158	
COI1 AT2G39940 RNI-like superfamily protein / _ 0 -0.19787 -0.1	68118	
Glycerolipid remodelling		
NPC4     AT3G03530     non-specific phospholipase C4     7     3     6.29128     4.7	4817	
NPC5 AT3G03540 non-specific phospholipase C5 Nd 1 Nd	٧d	
PLDÇ1 AT3G16785 phospholipase D zeta 1 0 0 0.204936 0.01	01217	
PLDÇ2 AT3G05630 phospholipase D zeta 2	4813	
MGD1 AT4G31780 monogalactosyldiacylglycerol synthase 1 2 0.38767 0.69	95703	
MGD2 AT5G20410 monogalactosyldiacylglycerol synthase 2 4 4.49902 4.3	<u>9337</u>	
MGD3 AT2G11810 monogalactosyldiacylglycerol synthase 3 0 8.67635 6.4	8075	
DGD1 AT3G11670 digalactosyldiacylglycerol synthase 1 1 0.718678 1.4	<u>5735</u>	
DGD2AT4G00550digalactosyldiacylglycerol synthase 201.8451.9	9015	
SQD2AT5G01220sulfoquinovosyldiacylglycerol synthase44.455934.4	<u>5593</u>	

#### **Figure Legends**

**Figure 1. Experimental design.** Three phosphate growing conditions were compared. In the "Pi-supplemented" conditions, plants were grown for 7 days on HPi medium and then transferred to a fresh HPi medium. In the "Pi-deprived" condition, plants were grown for 7 days on LPi medium and then transferred to a fresh LPi medium. In the "Pi-resupply" condition, plants were grown for 7 days on LPi medium and then transferred to a fresh HPi. Plants were then collected for various analyses, *i.e.* 0.5, 1 and 3 hours after transfer. In some analyses, plants were collected 24 hours after transfer.

Figure 2. K-mean clustering of gene expression profiles in Pi-deprived Arabidopsis following a resupply with phosphate. Plants were grown for 7 days on LPi medium and then transferred to a fresh HPi, corresponding to the "Replenished" condition. Plants were then collected 0.5, 1 and 3 hours after transfer. RNA was extracted and gene expression determined as described in the Methods section. A partition of differentially expressed genes was performed using a K-mean method, with a number of partitions set to 10 and a clustering based on a Euclidian distance (Liu et al. 2014a). Each cluster consists of genes with similar expression profiles following Pi-resupply, with representative nearest mean curves shown, serving as prototypes. Three clusters comprise genes upregulated following Pi resupply (Clusters 2, 4 and 5), whereas three clusters comprise genes which expression is downregulated (Clusters 7, 8, 10). Based on gene ontology (GO) term enrichment using two independent methods, GOseq (P-value <  $5.10^{-2}$ ) and DAVID (P-value  $\leq 1.10^{-2}$ ), genes involved in enriched molecular function (MF) and biological processes (BP) related to phosphate incorporation and homeostasis, glycerolipid remodeling and jasmonic acid biosynthesis and signaling are indicated.

Figure 3. Expression of genes involved in JA biosynthesis and signaling and in lipid remodeling in the WT Col0 strain and in the *coi1-16* mutant of *Arabidopsis thaliana*. Plants were cultivated as described in Fig. 1. Leaves were carefully collected 3 hours after medium transfer, RNA were extracted and gene expression levels were evaluated by RT-qPCR. Data were normalized as described in Methods, using the expression of TUB2 as a reference. Experiments correspond to biological triplicates. Error bars show standard deviations. White bars, expression levels in Col0; solid bars, expression levels in *coi1-16*. Two-way ANOVA Dunett's test: (\*) P-value <  $10^{-1}$ ; (\*\*) P-value <  $10^{-2}$ ; (\*\*\*) P-value <  $10^{-3}$ ; (\*\*\*) P-value <  $10^{-4}$ .

**Figure 4**. **Histochemical detection of the GUS activity in** *Arabidopsis* **355::JAZ1::GUS transgenic plants, submitted to different phosphate growing conditions. A**, **Pi-supplemented condition.** Plants were grown for 7 days on HPi medium and then transferred to a fresh HPi medium for 24 hours. Scale bar: 1mm. **B**, **Pi-deprived condition.** Plants were grown for 7 days on LPi medium and then transferred to a fresh LPi medium for 24 hours. Scale bar: 500 μm. **C**, **Pi-resupply condition.** Plants were grown for 7 days on LPi medium for 7 days on LPi medium and then transferred to a fresh HPi medium and then transferred to a fresh HPi medium for 24 hours. Scale bar: 500 μm. **C**, **Pi-resupply condition.** Plants were grown for 7 days on LPi medium and then transferred to a fresh HPi medium for 24 hours.

**Figure 5. Glycerolipid profiles in** *Arabidopsis thaliana* **Col-0 and coi-16 mutant treated in various phosphate regimen. A. Glycerolipid profiles in shoots. B. Glycerolipid profiles in roots.** Plants were grown as shown in Fig. 1, *i.e.* 7 days on HPi medium and then transferred to a fresh HPi medium for 3 hours in Pi-supplemented condition (Pi-supplem.); 7 days on LPi medium and then transferred to a fresh LPi medium for 3 hours in Pi-deprived

condition and eventually 7 days on LPi medium and then transferred to a fresh HPi medium for 3 hours in Piresupply condition. Roots and shoots were carefully collected, and glycerolipids were extracted as described in Methods. Each lipid class is expressed in nmol per mg of dry weight (DW). Green bars, glycerolipid profiles in Col-0; brown bars, glycerolipid profiles in coi1-16. Data correspond to three independent biological replicates. Error bars correspond to standard deviation. DAG, diacylglycerol; DGDG, digalactosyldiacylglycerol; DPG, diphosphatidylglycerol; MGDG, monogalactosyldiacylglycerol; PA, phosphatidic acid; PC, phosphatidylcholine; PE, phosphatidylethanolamine; PG, phosphatidylglycerol; PI, phosphatidylinositol; SQDG, sufoquinovosyldiacylglycerol, TAG, triacylglycerol. Significant differences between Col-0 and coi1-16 are indicated by a star (p-value < 0.05; two-tailed t-test).

**Figure 6. Leaf phosphatidylcholine diacyl profiles in** *Arabidopsis thaliana* **Col-0** and **coi-16** mutant treated in **various phosphate regimen.** Plants were grown as shown in Fig. 1, *i.e.* 7 days on HPi medium and then transferred to a fresh HPi medium for 3 hours in Pi-supplemented condition (Pi-supplem.); 7 days on LPi medium and then transferred to a fresh LPi medium for 3 hours in Pi-deprived condition and eventually 7 days on LPi medium and then transferred to a fresh HPi medium for 3 hours in Pi-deprived condition. Shoots were carefully collected, and glycerolipids were extracted as described in Methods. Phosphatidylcholine (PC) diacyl moieties were analyzed as described in Methods. Data correspond to three independent biological replicates. Error bars correspond to standard deviation. Diacyls are expressed as the sum of carbon contained in the two fatty acids and the number of double bonds they harbor. For instance, PC containing a 16:0 and an 18:2 fatty acids is expressed as PC-34-2; PC containing two 18:2 fatty acids is expressed as PC-36-4. Green bars, PC profiles in Col-0; brown bars, PC profiles in coi1-16. Data correspond to three independent biological replicates. Significant differences between Col-0 and coi1-16 are indicated by a star (p-value < 0.05; two-tailed t-test).

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Plant & Cell Physiology Page 28 of 32 2729 genes; GO enrichment analysis 3482 genes; GO enrichment analysis 1255 genes; GO enrichment analysis using Goseq (P-value < 5.10<sup>-2</sup>) using Goseq (P-value < 5.10<sup>-2</sup>) using Goseq (P-value < 5.10<sup>-2</sup>) GO term GO term GO term BP Phosphate ion transmembrane BP Phosphate ion transmembrane BP Phosphate ion transmembrane transport transport transport BP Galactolipid biosynthetic process BP Galactolipid biosynthetic process BP Galactolipid biosynthetic process BP Endoplasmic reticulum to BP Fatty acid oxidation BP CDP-diacylglycerol biosynthetic chloroplast transport process BP Jasmonic acid biosynthetic process BP Jasmonic acid mediated signaling BP Response to jasmonic acid BP Regulation of jasmonic acid pathway MF Jasmonoyl-isoleucinemediated signaling BP Regulation of jasmonic acid hydroxylase activity BP Response to jasmonic acid mediated signaling **BP** Root epidermal cell BP Response to jasmonic acid differenciation **Cluster 2 Cluster 4 Cluster 5** 20 20 20 20 20 LogFC normalized counts (1h vs. 0.5h and 3h vs. 0.5h) C -20 -20 -20 -20 -20 0.5h 1h 0.5h 0.5h 0.5h 1h 0.5h 3h 1h 3h 1h 3h 3h 1h 3h **Cluster 7 Cluster 8** Cluster 10 20 20 20 20 20 0 C C  $\sim$ -20 -20 -20 -20 -20 0.5h 0.5h 1h 3h 1h 3h 0.5h 1h 3h 0.5h 1h 3h 0.5h 1h 3h 440 genes; GO enrichment analysis 5744 genes; GO enrichment analysis using Goseq (P-value < 5.10<sup>-2</sup>) using Goseq (P-value < 5.10<sup>-2</sup>) GO term GO term BP Phosphate ion homeostasis **BP** Photosynthesis **BP** Phosphorus metabolic process BP ATP synthesis coupled proton transport **BP** Lipid storage BP photosystem I assembly BP Sulfolipid biosynthetic process BP photosystem II assembly BP Galactolipid metabolic process BP photosystem II stabilization MF 1,2-diacylglycerol 3-betagalactosyltransferase activity BP photosystem II repair BP Jasmonic acid stimulus BP mitotic nuclear division BP Jasmonic acid mediated signaling **BP** chloroplast fission pathway

BP leaf morphogenesis

JA biosynthesis and signaling Glycerolipid remodeling



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Phosphatidylcholine diacyl profile