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An ORP3-independent pathway uses 4,5-didehydro-jasmonate for jasmonate synthesis

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23 Abstract

24 Biosynthesis of the phytohormone jasmonoyl-isoleucine (JA-Ile) requires reduction of the JA 25 precursor 12-oxo-phytodienoic acid (OPDA) by OPDA-reductase-3 (OPR3). Previous 26 analyses of opr3-1 Arabidopsis mutant suggested an OPDA signaling role, independent of 27 JA-Ile and its receptor COI1; this hypothesis was challenged, as *opr3-1* is a conditional allele 28 not completely impaired in JA-Ile biosynthesis. To clarify the role of OPR3 and OPDA in 29 JA-independent defenses, we isolated and characterized a loss-of-function opr3-3 allele. 30 Strikingly, opr3-3 plants remained resistant to necrotrophic pathogens and insect feeding, and 31 activated COI1-dependent JA-mediated gene expression. Analysis of OPDA derivatives 32 identified 4,5-didehydro-JA in wounded wild-type and opr3-3 plants. OPR2 was found to 33 reduce 4,5-didehydro-JA to JA, explaining the accumulation of JA-Ile and activation of JA-34 Ile-responses in opr3-3 mutants. Our results demonstrate that in absence of OPR3 OPDA 35 enters the β -oxidation to produce 4,5-ddh-JA as a direct precursor of JA and JA-Ile, which 36 identifies an OPR3-independent pathway for JA biosynthesis.

38 Introduction

39 Jasmonoyl-isoleucine (JA-Ile; 1), a lipid-derived phytohormone essential for plant survival 40 in nature, regulates plant responses to many stresses including defense against insects, nematodes, and necrotrophic fungal and bacterial pathogens¹⁻⁵. JA-Ile also regulates growth 41 and developmental processes such as pollen viability, stamen development and senescence^{5,6}. 42 43 The bioactive form of the hormone, (+)-7-iso-JA-Ile (2), acts as "molecular glue" to induce formation of the COI1-JAZ co-receptor complex⁷⁻¹⁰. JA-IIe-mediated COI1-JAZ interaction 44 45 triggers ubiquitination of JAZ repressors and their degradation by the proteasome, which in turn activates several transcription factors that regulate specific physiological responses^{1,7,10}. 46

47 The octadecanoid pathway is responsible for biosynthesis of JA (3) and its derivatives (Supplementary Results, Supplementary Fig. 1)^{5,6,11}. Jasmonate biosynthesis begins with 48 release of α -linolenic acid (18:3) from plastidial membrane lipids by lipases and its 49 50 oxygenation by 13-lipoxygenase⁵. The coupled dehydration-cyclization reaction promoted 51 by allene oxide synthase (AOS) and allene oxide cyclase (AOC) then generates the first 52 cyclopentenone oxylipin, 12-oxo-10,15(Z)-phytodienoic acid (OPDA; 4), in the chloroplast (Supplementary Fig. 1)^{5,11}. OPDA is transported into the peroxisome and reduced to 8-(3-53 oxo-2-(pent-2-envl)cyclopentyl)octanoic acid (OPC-8) by OPDA reductase 3 (OPR3)¹¹. 54 55 Arabidopsis has at least two additional OPR enzymes, OPR1 and OPR2. Although these 56 enzymes can reduce unnatural stereoisomers of OPDA in vitro, experimental evidence of their involvement in OPDA reduction in vivo has not been provided yet^{12,13}. Peroxisomal 57 58 β-oxidation rounds OPC-8 undergoes three to generate 6-(3-oxo-2-(pent-2-59 envl)cyclopentyl)hexanoic acid (OPC-6), 4-(3-oxo-2-(pent-2-envl)cyclopentyl)butanoic acid (OPC-4) and finally JA, as an equilibrium of (+)-7-iso-JA and (-)-JA stereoisomers^{5,14,15}. 60 61 2,3-dinor-12-oxo-10,15(Z)-phytodienoic acid (dnOPDA; 5) is a 16-carbon analog of OPDA 62 synthesized from 7,10,13-hexadecatrienoic acid (16:3) by a parallel hexadecanoid pathway (Supplementary Fig. 1)^{5,6,16}. dnOPDA is thought to follow the same pathway as OPDA,
giving rise to OPC-6, OPC-4 and JA (Supplementary Fig. 1)^{5,6,16}. Finally, the cytoplasmic
JA-amido synthetase JAR1 conjugates JA with isoleucine (Ile), to generate bioactive JA-Ile
[(+)-7-*iso*-JA-Ile], which is in equilibrium with its inactive epimer (-)-JA-Ile^{8,17}.

67 JA biosynthesis or signaling mutants are sterile and more susceptible than wild-type (WT) 68 plants to insects and necrotrophic pathogens. Available opr3 alleles are also sterile but, in 69 contrast to all other mutants in the pathway, the opr3-1 allele showed near-WT resistance to insects (i.e., Bradysia impatiens) and pathogens (i.e., Alternaria brassicicola)^{18,19}. This 70 mutant accumulated OPDA, but only minute amounts of JA were detected¹⁹. A putative role 71 72 was therefore proposed for OPDA or OPDA derivatives in activation of defenses, independent of JA or other cyclopentanones¹⁹. Further analysis of the opr3-1 allele showed it 73 74 to be a conditional mutant in which the T-DNA insertion in an OPR3 intron could be spliced 75 out in specific conditions. Upon Botrytis cinerea infection, this line produced OPR3 transcripts, accumulated JA, and showed partial resistance²⁰. Although this finding provides 76 77 a plausible explanation for pathogen resistance in opr3-1, the debate remains, since several 78 studies reported JA- or COI1-independent roles for OPDA using tools with certain limitations such as *opr3-1*, RNAi silencing *OPR3*, or weak alleles of $coil^{21-30}$; additional *opr3* alleles 79 80 described so far do not help in addressing the OPDA role, as they might not be loss-of-81 function alleles 31 .

To further analyze the role of OPR3 and OPDA in defense, we isolated a true loss-offunction *opr3-3* mutant and characterized JA-mediated plant responses. *opr3-3* plants were resistant to insect feeding and necrotrophic pathogen infection, and activated JA-related gene expression in response to these stresses similarly to the original *opr3-1* allele. These responses were fully COI1-dependent and therefore cannot be attributed to the proposed OPDA defense-signaling function. Our data indicate that, in the absence of OPR3, OPDA 88 could give rise to a COI1 ligand. Quantification of OPDA derivatives showed that opr3-3 89 accumulated low JA and JA-Ile levels, which explained the phenotypic observations. In 90 contrast, β-oxidation intermediates such as OPC-6 and OPC-4 were not detected, although we 91 identified 4,5-didehydro-JA (4,5-ddh-JA; 6) as an alternative jasmonate that accumulated in 92 both WT and opr3-3 plants. We also showed that 4,5-ddh-JA is not conjugated to Ile to form 93 4,5-ddh-JA-Ile (7), but is reduced to JA by OPR2. Our results define an alternative pathway 94 for JA biosynthesis and demonstrate that OPDA can enter the β -oxidation pathway in the 95 absence of OPR3.

97 **Results**

98 Identification of a knockout (KO) allele of opr3

99 To clarify the controversy generated by the conditional opr3-1 mutant, we sought a 100 complete loss-of-function opr3 allele in T-DNA collections. Of the putative mutants selected, only the SK24765 line was sterile (Col-0 background, Saskatoon collection³²), a well-101 102 described feature of *opr3* mutants^{18,33}. Exogenous JA restored fertility to SK24765 plants 103 (Fig. 1a). We confirmed T-DNA insertion at position G1982 within the fifth exon in all 104 sterile plants tested, and termed the SK24765 line opr3-3 (Supplementary Fig. 2a). As 105 anticipated, the opr3-3 mutation is recessive and heterozygous opr3-3 plants segregate 1:3 106 sterile:fertile plants.

To determine whether *opr3-3* was a complete loss-of-function allele, we analyzed *OPR3* expression by qPCR in various conditions including wounding, insect feeding and fungal infections. *opr3-3* did not accumulate any *OPR3* transcripts in any of these conditions (Fig. 1b). Consistent with previous reports²⁰, however, OPR3 expression was detected in *opr3-1* plants infected by *B. cinerea* and *A. brassicicola* (Supplementary Fig. 2b). These results confirm that the *opr3-1* allele is conditional and indicated that *opr3-3* allele is a complete KO.

114 *opr3-3* activates defense responses

To assess the proposed JA-independent role for OPDA in plant defense, we compared the response of *opr3-3*, WT and *coi1-30* mutants to an insect (*Spodoptera littoralis*), fungal pathogens (*B. cinerea*, *A. brassicicola*) and mechanical wounding. As predicted, *coi1-30* mutants were more susceptible to all pathogens and insect than WT plants, due to the impaired activation of JA-dependent defenses (Fig. 2a and Supplementary Fig. 3). In contrast, *opr3-3* showed near-WT resistance to *B. cinerea* and *A. brassicicola*, not markedly different from that of the leaky *opr3-1* allele; the *opr3-3* response to *S. littoralis* was
intermediate between WT and *coi1-30* (Fig. 2a and Supplementary Fig. 3).

123 Expression analyses showed that all stress conditions induced several JA marker genes in 124 opr3-3 mutants (Fig. 2b,c). This induction was lower than in WT plants, but still much 125 higher than the fully impaired induction in *coi1-30* mutants. This finding was unanticipated, 126 as this KO allele was not thought to produce JA, and thus should not activate the JA pathway. 127 To evaluate whether the defense responses in opr3-3 depended on the JA-Ile receptor 128 COI1, we generated an opr3-3 coi1-30 double mutant and challenged it with fungi 129 (B. cinerea, A. brassicicola), an insect (S. littoralis), and wounding. Infection symptoms, 130 spore production, larval weight and induction of defense gene expression for the opr3-3 131 coil-30 mutant were indistinguishable from those of coil-30 (Fig. 2 and Supplementary 132 Fig. 3). The data suggest that the complete loss-of-function mutant opr3-3 can activate JA-133 mediated gene expression and defense responses in a COI1-dependent manner.

134 Analysis of JAs in *opr3-3* plants

To understand how *opr3-3* activates JA-mediated responses in a COI1-dependent manner, we measured levels of OPDA-derivatives by liquid chromatography-mass spectrometry (LC-MS) after wounding. Consistent with the KO nature of the *opr3-3* mutation, OPC-4 accumulated in wounded WT but not *opr3-3* plants (Supplementary Fig. 4a). This indicated that lack of a functional OPR3 impairs OPDA reduction, and thus the production of the β-oxidation intermediate OPC-4. The conditional *opr3-1* mutant was very similar to *opr3-3* (Supplementary Fig. 4b).

142 In contrast to current models of the JA biosynthesis pathway, wounding induced 143 accumulation of low JA and JA-Ile levels in *opr3-3*, similar to the levels reported for the 144 conditional *opr3-1* allele (Fig. 3a-c and Supplementary Fig. 4c,d)¹⁹. These data suggested the 145 existence of an alternative OPR3-independent pathway for JA biosynthesis.

146 **4,5-didehydro-JA** accumulates in *opr3* and WT

In WT plants, OPDA is reduced by OPR3 in the peroxisome and is funneled into the β -oxidation pathway to produce JA. Based on the wide range of molecules that undergo peroxisomal β -oxidation, this process seems to have limited specificity ^{14,15}. We thus hypothesized that in the absence of OPR3, OPDA might enter the β -oxidation pathway without prior reduction of the cyclopentenone ring to the corresponding cyclopentanone; in this case, it would produce 4,5-didehydrojasmonate (4,5-ddh-JA) as the non-reduced cyclopentenone-analog of JA³⁴.

154 To test this hypothesis, we chemically synthesized 4,5-ddh-JA and its Ile conjugate 4,5-155 ddh-JA-Ile (as quantification standards; see Methods and Supplementary Fig. 5a) and 156 analyzed their accumulation in WT and *opr3-3* plants after wounding. WT plants transiently 157 accumulated low 4,5-ddh-JA levels in response to wounding (Fig. 3d), which indicated that 158 the direct entry of OPDA into β -oxidation is a natural alternative in WT plants. opr3-3 159 accumulated approximately 20-fold more 4,5-ddh-JA than WT plants after wounding (Fig. 160 3d). The conditional opr3-1 mutant also accumulated 4,5-ddh-JA at levels similar to opr3-3 161 (Supplementary Fig. 6). These data show that 4.5-ddh-JA is a natural oxylipin that 162 accumulates in WT plants in response to wounding, and that this accumulation is promoted in 163 opr3 alleles. They also suggest that 4,5-ddh-JA is responsible for defense activation in opr3 164 alleles. Finally, 4,5-ddh-JA-Ile was not detected in WT or opr3 plants, which implies that 165 4,5-ddh-JA might not be a JAR1 substrate (Supplementary Fig. 5b).

166 4,5-didehydro-JA triggers COI1-dependent responses

To assess whether 4,5-ddh-JA acts as a signaling molecule, we analyzed its root-growth inhibitory activity, a typical COI1-mediated response. WT Col-0 plants grown in the presence of 4,5-ddh-JA showed inhibited root growth comparable to that of the JA control treatment (Fig. 4a,b). The *opr3-3* response to 4,5-ddh-JA was similar to that of the WT, which indicates that this molecule acts downstream of OPR3. Nonetheless, *jar1-1* and *coi1-1*were insensitive to 4,5-ddh-JA, indicating that conjugation to Ile by JAR1 (directly or
through conversion to JA) and perception by COI1 are necessary for 4,5-ddh-JA activity
(Fig. 4a,b).

175 We also studied the effect of 4,5-ddh-JA on hormone-induced degradation of JAZ 176 repressors, another typical COI1-mediated response. Both JA and 4.5-ddh-JA rapidly induced JAZ1 protein degradation in Arabidopsis 35S:JAZ1-GUS transgenic plants (Fig. 4c). 177 178 In addition, 4,5-ddh-JA caused JAZ1-GUS degradation in opr3-3, which indicates that OPR3 179 is not necessary for 4,5-ddh-JA activity. However, neither JA nor 4,5-ddh-JA triggered 180 JAZ1-GUS degradation in *jar1-1* and *coi1-1*, which corroborated the finding that JAR1 is 181 needed to generate the bioactive form of the signaling molecule, which is perceived by COI1. 182 These results show that 4,5-ddh-JA is a signalling molecule that requires JAR1 and COI1 to 183 induce JA-mediated responses.

184 **4,5-didehydro-JA** is the source of JA in *opr3-3*

185 Given that 4,5-ddh-JA was bioactive, but its Ile conjugate could not be detected after 186 wounding, we reasoned that 4,5-ddh-JA might be reduced in the cell to produce JA. To test 187 this hypothesis, we measured JA accumulation after exogenous 4,5-ddh-JA treatment. Both 188 WT and opr3-3 plants accumulated similarly high JA levels after 4,5-ddh-JA application 189 (Supplementary Fig. 7). To further confirm that 4,5-ddh-JA can be converted to JA, and to 190 detect potential intermediates in this alternative pathway, we fed wounded plants with 191 deuterated α -linolenic acid ([²H₅]18:3) and analyzed accumulation of deuterated 18:3 derivatives. We detected labeled OPDA ($[^{2}H_{5}]OPDA$) and dnOPDA ($[^{2}H_{5}]dnOPDA$) in WT 192 and *opr3-3* plants (Fig. 5). $[{}^{2}H_{5}]$ hexadecatrienoic acid ($[{}^{2}H_{5}]$ 16:3) was not detected, which 193 shows that $[^{2}H_{5}]$ dnOPDA is not produced by the parallel hexadecanoid pathway but it is 194 derived from $[^{2}H_{5}]$ OPDA by a single round of β -oxidation in WT and *opr3-3* plants (Fig. 5b). 195

196 The β-oxidation intermediates OPC-6 and OPC-4 were detected as the deuterated derivatives ^{[²H₅]OPC-6 and ^{[²H₅]OPC-4 only in WT but not in *opr3-3* plants (Fig. 5b), which suggests}} 197 198 that the canonical β-oxidation of typical OPDA derivatives (OPC) is lost in the absence of 199 OPR3. Both WT and, to a greater extent, opr3-3 plants accumulated deuterated tetranor-OPDA ($[{}^{2}H_{5}]$ tnOPDA; 8) and 4,5-ddh-JA ($[{}^{2}H_{5}]$ 4,5-ddh-JA), which implies that dnOPDA 200 201 can be converted into 4.5-ddh-JA in two further rounds of β -oxidation. Despite the absence 202 of OPC-6 and OPC-4, we detected deuterated JA ($[^{2}H_{5}]JA$) and JA-Ile ($[^{2}H_{5}]JA$ -Ile) in WT 203 and opr3-3 plants, which indicates that JA can be synthesized by a 4,5-ddh-JA-mediated 204 biosynthetic pathway alternative to the canonical OPC-mediated β -oxidation pathway (Fig. 205 5b). In summary, these data show that an OPR3-independent JA-biosynthetic pathway 206 occurs naturally in WT plants and that the flux through this pathway is increased in absence 207 of OPR3.

208 4,5-didehydro-JA is reduced to JA by OPR2

209 We sought enzymes responsible for conversion of 4,5-ddh-JA to JA in the absence of OPR3. Cytosolic OPR1 and OPR2 are the enzymes most similar to peroxisomal OPR3³⁵. 210 211 Expression of both OPR1 and OPR2 is induced after wounding, although to a lesser extent than OPR3 (Supplementary Fig. 8a; Genevestigator)^{19,36}. To test the hypothesis that OPR1 212 213 and OPR2 reduce 4,5-ddh-JA to JA in opr3-3 plants, loss-of-function opr1-1 and opr2-1 214 mutants were selected and crossed with opr3-3 to generate the double mutants opr1-1 opr3-3 215 and opr2-1 opr3-3 (Supplementary Fig. 8b-d); triple mutants could not be obtained by 216 crossing, as OPR1 and OPR2 are contiguous genes. The single opr1-1 and opr2-1 mutants 217 accumulated JA and 4,5-ddh-JA levels similar to those of WT plants after wounding (Fig. 218 6a,b). Wound-induced levels of 4,5-ddh-JA were as high in opr1-1 opr3-3 and opr2-1 opr3-3 219 as in opr3-3, or even slightly higher in opr2-1 opr3-3 (Figure 6b). In contrast, wound-220 induced JA accumulation was reduced in opr1-1 opr3-3 compared to opr3-3 and almost

undetectable in *opr2-1 opr3-3*, which suggested that OPR2 is the main enzyme responsible
for JA reduction from 4,5-ddh-JA (Fig. 6a).

As predicted, JA-Ile accumulation mirrored that of JA. *opr1-1* and *opr2-1* accumulated JA-Ile at levels similar to WT plants. These levels were almost unaffected in *opr1-1 opr3-3* compared to *opr3-3* and undetectable in *opr2-1 opr3-3* (Supplementary Fig. 8e).

To confirm that OPR2 is the main enzymatic activity in 4,5-ddh-JA reduction to JA, we measured JA accumulation after exogenous treatment with 4,5-ddh-JA. *opr3-3* and *opr1-1 opr3-3* plants accumulated similarly high JA and JA-IIe levels after 4,5-ddh-JA application, whereas, the *opr2-1 opr3-3* double mutant showed significantly lower JA and JA-IIe levels (Fig. 6c). These findings show that OPR2 is primarily responsible for JA reduction from 4,5ddh-JA.

232 To directly test whether OPR1 and OPR2 are capable of reducing 4,5-ddh-JA to yield 233 JA, the two enzymes were expressed as N-terminally His-tagged fusion proteins in E. coli 234 and purified from bacterial extracts. Recombinant OPR2 reduced 4,5-ddh-JA at the expense of NADPH with a catalytic efficiency (K_{cat}/K_M) of 3750 M^{-1} *s⁻¹ (K_{cat} of 0.819 s⁻¹ and V_{max} of 235 0.819 M^{-1} *s⁻¹), while no reduction was observed for OPR1 (Fig. 6d). The apparent K_M of 236 237 OPR2 for 4,5-ddh-JA, determined as an indirect measure for substrate affinity, was 7-fold higher (218 μ M) than the K_M of OPR3 for its substrate OPDA (35 μ M)¹². Therefore, the 238 239 affinity of OPR2 for 4,5-ddh-JA is lower but in the same order of magnitude to that of OPR3 240 for OPDA.

To assess the role of OPR1 and OPR2 in pathogen responses, the *opr* double mutants were infected with *B. cinerea* and *A. brassicicola*. As anticipated, *opr3-3* showed near-WT resistance to the fungal infection, very similar to *opr1-1 opr3-3*, whereas *opr2-1 opr3-3* were more susceptible than any of the genetic backgrounds tested (Fig. 6e and Supplementary Fig. 8f-h). Consequently, induction of JA-regulated defence genes was similar in WT and *opr3-3*

- plants, reduced in *opr1-1 opr3-3* and considerably lower in *opr2-1 opr3-3* (Fig. 6g and
 Supplementary Fig. 8i).
- 248 Wound induction of JA-marker gene expression (JAZ1, JAZ5, AOS and MYC2) was
- reduced in the double mutants compared to the single *opr3-3* (Fig. 6h and Supplementary Fig.
- 250 8j). Induction was also lower in *opr2-1 opr3-3* than *opr1-1 opr3-3*, further suggesting OPR2
- predominance over OPR1.
- In all, the genetic, biochemical and physiological data show that OPR2 mediates 4,5-ddh-
- 253 JA transformation into JA after wounding.

255 Discussion

256 Here we identify an OPR3-independent pathway for JA biosynthesis that involves direct 257 entry of OPDA into the β -oxidation pathway to produce dnOPDA, tnOPDA and 4,5-ddh-JA, 258 which is then reduced to JA by OPR2. To clarify the controversy regarding the activity of JA 259 precursors in the absence of OPR3 and the role of COI1 in their function, we obtained and 260 characterized a full loss-of-function opr3-3 allele unable to express any OPR3 transcript in 261 basal or stress conditions. This knockout mutant was unexpectedly able to activate JA-262 dependent defenses that were fully dependent on COI1. Consistent with the full KO nature 263 of opr3-3, the typical OPDA β-oxidation derivatives and JA precursors OPC-6 and OPC-4 264 were not detected in this mutant; even so, opr3-3 accumulated small amounts of JA and JA-265 Ile in stress situations, which suggested an OPR3- (and OPC)-independent pathway for JA 266 biosynthesis. LC-MS quantification of deuterated derivatives after feeding plants deuterated 267 α -linolenic acid/18:3 showed accumulation of labeled OPDA, dnOPDA, tnOPDA, 4,5-ddh-268 JA, JA, and JA-Ile in opr3-3, but neither OPC-6 nor OPC-4. These results demonstrate that 269 in the absence of OPR3, OPDA can enter the β -oxidation pathway to produce non-reduced 270 OPDA derivatives (dnOPDA, tnOPDA and 4,5-ddh-JA). Labeled dnOPDA, tnOPDA and 271 4,5-ddh-JA were also identified in WT plants, which indicates that this alternative JA-272 biosynthetic pathway occurs naturally and is potentiated in the absence of OPR3 in opr3 273 mutants.

It was traditionally considered that hexadecatrienoic acid is the only dnOPDA source ¹⁶.
Our results indicate that OPDA is an alternative source for dnOPDA production through a
single β-oxidation cycle, a conversion that would require OPDA conjugation to CoA. In
support of this hypothesis, the OPC8-CoA ligase OPCL1 also accepts OPDA as substrate ^{37,38}.
Moreover, an analysis of Arabidopsis acyl-CoA synthase substrate specificity identified some
enzymes with high affinity for OPDA and dnOPDA, and the *At5g63380* gene was proposed

to encode an OPDA:CoA ligase ³⁷, although its relevance in JA synthesis remains to be demonstrated. OPDA conversion to OPC-8 by OPR3 was thus thought to occur also at the level of CoA conjugates ^{37,39}; OPDA-CoA is a high-affinity substrate of the β -oxidation enzyme acyl-CoA oxidase *in vitro*, and direct OPDA entry into β -oxidation has been hypothesized ⁴⁰.

285 Our genetic analysis of KO mutants complemented the LC-MS quantification and 286 supported 4,5-ddh-JA as the precursor of JA and JA-Ile in opr3-3. Previous studies showed 287 that the OPDA reductases OPR1 and OPR2 can reduce OPDA and other oxylipins in vitro, but with less efficiency than OPR3 and different stereospecificity ^{13,14,24,41,42}. This low 288 289 efficiency, together with the lack of peroxisomal location of OPR1 and OPR2 that contrasts 290 with peroxisomal OPR3, ruled out a role for OPR1 and OPR2 in the JA biosynthetic pathway 291 *in vivo*¹³. Consistent with this idea, we did not detect OPC-6 or OPC-4 in *opr3-3*, which 292 confirmed that OPR1 and OPR2 cannot (even partially) replace OPR3 in OPDA reduction. 293 opr2 nonetheless show reduced JA accumulation when crossed with opr3, which indicates 294 that OPR2 is necessary for 4,5-ddh-JA reduction to JA, likely in the cytoplasm. Confirming 295 the genetic data, recombinant OPR2 was found to catalyze the NADPH-dependent reduction 296 of 4.5-ddh-JA to yield JA. The need for JAR1 for 4.5-ddh-JA activity in vivo and the lack of 297 the 4,5-ddh-JA-Ile conjugate also support the concept that 4,5-ddh-JA must be reduced to JA 298 and subsequently conjugated to Ile for activity.

Despite the low JA levels detected in the original analysis of opr3-1 (4% of that in WT)¹⁹ and in more recent studies ⁴³, opr3-1 has long been assumed to be an appropriate tool for uncoupling OPDA and JA synthesis and for dissecting OPDA-specific responses ^{5,18,19}. This allele has been widely used to show JA-independent roles of OPDA, which in many cases also seemed to be independent of the JA-Ile receptor COI1 ^{19,22-29}. *opr3-1* is nonetheless a conditional allele able to express *OPR3* and to produce notable amounts of JA after fungal infection (up to 30% of JA levels in infected WT plants) ²⁰. The extensive use of this conditional allele has generated much confusion in the field; therefore, JA-independent functions of OPDA should be revised. This matter was aggravated by the simultaneous use of weak *coi1* alleles, which also suggested COI1-independent functions of OPDA, and by exogenous OPDA treatments that might not represent endogenous functions ²²⁻²⁹.

310 Our results using opr3-3 coi1-30 double mutants demonstrate that defense responses in 311 opr3-3, including the activation of defense gene expression, are fully COI1-dependent. 312 Although we cannot rule out induction of COI1-independent effects by exogenous OPDA treatment ^{19,22,24,25,29}, OPDA effects that depend on COI1 in vivo are unlikely to be mediated 313 314 by any JA-independent function of OPDA. OPDA-mediated effects that are independent of 315 COI1, on the other hand, may still be attributed to the activity of OPDA per se. The OPDA 316 molecule carries a highly reactive α_{β} -unsaturated carbonyl group that defines the reactive electrophile species (RES) 44,45. RES activity induced by exogenous OPDA treatment might 317 318 thus explain the reported OPDA responses through its binding to cyclophilin 20-3, which in turn regulates cellular redox homeostasis²⁵. 319

OPR3 orthologues are not found in lower plants such as Bryophytes, but genes with notable similarity to *OPR2* have been identified ⁴⁶⁻⁴⁹. Despite of the lack of OPR3, JA and JA-IIe have been detected in several liverworts and mosses, which suggests that the OPR3independent JA biosynthetic pathway reported here is conserved in lower plants ⁵⁰. This alternative *OPR2*-dependent pathway might thus be the original and only way to synthesize JA in ancestral land plants, still present in some extant Bryophytes, whereas the current OPR3-dependence would be a later acquisition in evolution, found only in vascular plants.

Here we identify an OPR3-independent pathway for JA-Ile synthesis that occurs naturally in WT plants and is potentiated in *opr3* mutants. This pathway involves direct peroxisomal β-oxidation of OPDA to dnOPDA, tnOPDA and 4,5-ddh-JA that, after leaving the

- 330 peroxisome, is reduced to JA in the cytosol by OPR2. That this alternative pathway has a
- 331 crucial role in certain biological processes or in response to certain environmental stimuli is
- an attractive hypothesis that awaits further study.

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506 Figure legends

507 **Figure 1**. *opr3-3* is a complete loss-of-function allele

(a) Inflorescences of 7-week-old plants are shown. *opr3-3* mutants are sterile and JA
treatment (bracket) rescues male fertility. Scale bars, 1 cm.

- 510 (b) Gene expression analysis of *OPR3* in wild-type (Col-0) and *opr3-3* plants in response to
- 511 Botrytis cinerea, Alternaria brassicicola, Spodoptera littoralis, or wounding. Statistically
- 512 significant OPR3 expression differences of control vs. challenged plants (Student's t-test; **
- 513 p <0.01 and *** p <0.001). ACT8 was used as housekeeping control gene. Each biological
- 514 sample consisted of tissue pooled from 5-10 plants (n = 5). Means \pm SD of 4 technical
- 515 replicates. Each experiment was repeated at least twice with similar results.

516 Figure 2. opr3-3 mutants activate defense responses

517 (a) Box-plots of spore quantification or larval weight of Col-0 plants and mutant infected

518 with *B.cinerea* (n = 15; 3 dai), *A.brassicicola* (n = 15; 9 dai), or challenged with *S.littoralis*

519 larvae (n = 70; 7 dai). Horizontal lines are medians, boxes show the interquartile range and 520 error bars show the full data range. Letters above columns indicate significant differences 521 evaluated by one-way ANOVA/Tukey HSD post hoc test (p <0.01). Experiments were

522 repeated four times with similar results (or twice in the case of Spodoptera).

523 (b and c) Expression of JA-regulated genes in challenged plants (n = 10). *PDF1.2* and 524 CYP79B3, or JAZ7, LOX3 and AOS were measured by real-time PCR in untreated plants 525 (control, C) and in plants challenged for 3 days with B. cinerea (Bc) or 9 days with A. 526 brassicicola (Ab), or 48 h with S. littoralis larvae (Sl) or 30 min after wounding. Data in (b) 527 and (c) are shown as mean \pm SD (SE in *S.littoralis*) of three technical replicates expressed as 528 relative fold change normalized to ACT8. Experiments were repeated three times with 529 similar results. Statistically significant expression differences compared to Col-0 or opr3-3 are highlighted (Student's t-test; * p <0.05; ** p <0.01; *** p <0.001). 530

531 Figure 3. JA accumulation in Col-0 and *opr3-3* plants

- 532 (a) Structure of JA-Ile, JA and 4,5,-ddh-JA.
- 533 (b) Time-course accumulation of JA (nmoles/fresh weight (g)) in Col-0 (grey bars) and opr3-
- 534 *3* (black bars) after wounding. Four-week-old plants (n = 10) were wounded, and damaged
- 535 leaves collected after the indicated time. Data shown as mean \pm SD of four biological
- 536 replicates. Experiments were repeated three times with similar results.
- 537 (c) Levels of 4,5-ddh-JA (pmoles/fresh weight (g)) of Col-0 (grey bars) and opr3-3 (black
- bars) plants after wounding as in (b) (n = 10). Data shown as mean \pm SD of four biological
- replicates. Experiments were repeated three times with similar results.
- 540 (d) Accumulation of JA-Ile (pmoles/fresh weight (g)) in Col-0 (grey bars) and *opr3-3* (black
- bars) plants after wounding as in (b) (n = 10). Data shown as mean \pm SD of four biological
- 542 replicates. Experiments were repeated three times with similar results.

543 Figure 4. 4,5-didehydro-JA triggers JA-regulated COI1-dependent responses

- 544 (a and b) Arabidopsis Col-0 and mutant seedlings grown for 10 days on control medium (-)
- 545 or medium supplemented with 50 µM 4,5-ddh-JA (a). Bar, 1 cm. Quantification of root
- 546 length of Col-0 and mutant seedlings (n = 20) in control medium (-) or medium
- 547 supplemented with 50 μ M JA or 4,5-ddh-JA (b). Data shown as mean \pm SD. The experiment
- 548 was repeated three times with similar results. Letters above columns indicate significant
- 549 differences (one-way ANOVA/post-hoc Tukey HSD Test, p <0.01).
- 550 (c) GUS-staining visualization of JAZ1 stability in roots of 7-day-old transgenic Arabidopsis
- 551 35S:JAZ1:GUS on Col-0 or mutant backgrounds. Seedlings were treated with 5 µM JA or
- $25 \,\mu\text{M}$ 4,5-ddh-JA (1 h). The experiment was repeated three times with similar results. Scale
- 553 bars, 1 mm.

554 Figure 5. OPDA conversion into 4,5-ddh-JA, JA, and JA-Ile

Accumulation (10^6 arbitrary unit/fresh weight (g)) of radiolabeled derivatives of 18:3 in Colo and *opr3-3* plants. Five-week-old plants (n = 10) were wounded and fed with deuterated 18:3 ([2H_5]18:3) and accumulation of deuterated 18:3 derivatives was analyzed after 30 min. Results shown as means ± SD of four biological replicates. OPDA/dnOPDA biosynthesis takes place in the chloroplasts (top compartment), whereas OPDA/dnOPDA reduction and ®oxidation occur in the peroxisome (lower compartment). JA is transformed into JA-Ile in the cytosol.

- 562 Figure 6. OPR2 converts 4,5-ddh-JA into JA
- 563 (a and b) Time-course accumulation of JA or 4,5-ddh-JA in five-week-old Col-0 and opr
- mutants (n = 10) after wounding. Data are shown as mean \pm SD of four biological replicates.
- 565 (c) Accumulation of JA and JA-Ile after exogenous 4,5-ddh-JA treatment in opr3-3, opr1-
- 566 *lopr3-3* and *opr2-lopr3-3* mutants (n = 10). Mean \pm SD of four biological replicates after 567 subtraction of basal levels.
- 568 (d) NADPH-dependent reduction of 4,5-ddh-JA by OPR2. The reaction rate (μ M*s-1) of 569 recombinant OPR1 and OPR2 (1 μ M) was assayed under steady-state conditions with 570 increasing substrate concentrations (10 to 1000 μ M of (±)-4,5-ddh-JA). Mean ± SD of three 571 technical replicates.
- 572 (e) Box-plots of fungal spore quantification of Col-0 and mutant plants (n = 15) infected with 573 *B. cinerea* (7 dai). Letters above columns indicate significant differences (one-way 574 ANOVA/post-hoc Tukey HSD Test, p < 0.01).

575 (f) Expression of JA-regulated *PDF1.2* after *B. cinerea* infection or *JAZ5* after wounding in 576 Col-0 and mutant plants (n = 5). Gene expression was measured by RT-qPCR in untreated 577 plants (control, C) and in plants after infection with *B. cinerea* (7 dai) or 1 h after wounding 578 (n = 5). Data shown as mean \pm SD of three technical replicates expressed as relative fold 579 change normalized to *ACT8*.

- 580 Letters above columns in (a), (c), (g), and (h) indicate significant differences compared to
- 581 expression in *opr3-3* plants (Student's t-test, p <0.05).
- 582 All experiments were repeated at least twice with similar results.

584

585 **Online Methods**

586 Plant material and growth conditions

587 Arabidopsis thaliana Col-0 is the genetic background of wild-type and mutant lines used in this study, with the exception of *opr3-1*, which is in the Ws background ¹⁸. Plants were 588 grown in Johnson's medium at 21°C in a 16-h light/8-h dark cycle, as described ^{7,8}. The KO 589 590 lines opr1-1 (SALK 145353), opr2-1 (SALK 116381) and opr3-3 (SK24765) were obtained from the NASC and the Saskatoon collection ³². Homozygous lines were selected by PCR 591 592 using the T-DNA-specific and gene-specific primer combination LB GW1/OPR3 F1 for 593 opr3-3, LBb1.3/OPR1-F1 for opr1-1 and LBb1.3/OPR2-F1 for opr2-1. Double mutants were 594 generated by crossing of single mutant, and double-homozygous plants were identified by 595 PCR analyses. After bolting, flower buds of sterile mutant plants were treated with a 0.5 mM 596 MeJA solution (Sigma) dissolved in 0.1% Tween 20 (Calbiochem), daily for 2 weeks.

597 **Root measurements**

For root-growth inhibition assays, root length of 20 to 30 seedlings was measured 10 days after germination, alone or in the presence of 50 μ M jasmonic acid (JA; Sigma) or 50 μ M 4,5ddh-JA. Pictures were taken with a Nikon D1-x digital camera and root length was measured using ImageJ software. Data were analyzed by one-way ANOVA/Tukey HSD post hoc test (p <0.01). Three independent biological replicates (20-30 seedlings each) were measured for each sample. Data are shown as mean ± SD.

604 Fungal infection analyses

Seeds were grown directly in soil as described in ⁵¹. *coi1-30* and *opr3-3 coi1-30* mutants were selected in plates with Johnson's medium containing 0.5 μ M coronatine (Sigma), and transferred to soil after 7 days. At least 15 leaves of five-week-old plants (3 leaves/plant) were inoculated with *B. cinerea* suspension of 5 × 10⁶ spores/ml PDB (Difco) as described in 51 . Images of disease symptoms were taken 6 to 9 days after inoculation. Spores were quantified in a hemocytometer under a light microscope (Leica DMR UV/VIS). Five inoculated leaves of five different plants were pooled for each biological sample, and three to seven independent biological replicates were measured for each treatment. Data were analyzed by one-way ANOVA/Tukey HSD post hoc test (p <0.01). This experiment was repeated three times with similar results. Data are shown as mean ± SEM.

615 *A. brassicicola* infection assays were performed as described for *B. cinerea*, inoculating 616 each leaf with 20 μ l of a suspension of 10⁶ *A. brassicicola* spores/ml PDB. Data (analyzed by 617 one-way ANOVA/Tukey HSD post hoc test, p <0.01) are shown as mean \pm SEM. Images of 618 disease symptoms were acquired and spores quantified as for *B. cinerea*.

619 Insect bioassays

620 Plants were grown for three weeks in a growth chamber (short-day 10/14h photoperiod, 621 20°C, 65% relative humidity). coi1-30 and opr3-3 coi1-30 mutant were selected in the same 622 conditions as for fungal infection assays. Five-week-old plants were placed in transparent 623 plastic boxes and forty newly hatched Spodoptera littoralis larvae were placed on 70 plants 624 for seven days of feeding, when larvae were collected and weighed. Data were analyzed on 625 log-transformed values by one-way ANOVA/Tukey HSD post hoc test (p <0.01). The 626 experiment was repeated three times independently, with similar results. Data are shown as 627 mean \pm SEM.

628 Quantitative RT-PCR

Quantitative RT-PCR was performed using biological samples of tissue pooled from 5-10
plants. RNA was extracted and purified using Trizol reagent (Invitrogen) followed by the
High Pure RNA isolation kit (Roche), including DNase digestion to remove genomic DNA
contamination. cDNA was synthesized from 1 μg total RNA with the high-capacity cDNA
reverse transcription kit (Applied Biosystems). For gene amplification, 4 μl from a 1:10

634 cDNA dilution was added to 7.5 µL of Power SYBR Green (Applied Biosystems) and gene-635 specific primers (Supplementary Table 1). Quantitative PCR was performed in 96-well 636 optical plates in a 7500 or HT 7900 Real Time PCR system (Applied Biosystems) using 637 standard thermocycler conditions (an initial hold at 50°C for 120 s, 95°C for 10 min, 638 followed by a two-step SYBRPCR program of 95°C for 15 s and 60°C for 60 s for 40 cycles). 639 Relative expression values are the mean \pm SEM of three to four technical replicates relative 640 to the basal wild-type control using ACT8 as housekeeping gene. Data were analyzed by 641 unpaired Student's t-test. The experiment was repeated three times independently, with 642 similar results.

643 **Phytohormone analysis**

644 Phytohormone measurements were performed using biological samples of tissue pooled 645 from 5-10 plants and at least three independent biological replicates were measured for each 646 treatment. This experiment was repeated twice or three times with similar results. Data 647 (analyzed by unpaired Student's t-test.) are shown as mean \pm SD. (-)-Jasmonic acid (JA), cis-648 12-oxo-phytodienoic acid (OPDA) and N-(-)-jasmonoyl isoleucine (JA-Ile) were purchased 649 from OlChemim Ltd, dinor-12-oxo-phytodienoic acid (dnOPDA) from Cayman Chemical Company, OPC-4 and OPC-6 described in⁸, 4,5-ddh-JA and 4,5-ddh-JA-Ile were synthesized 650 651 (see below). The deuterium-labeled internal standards ²H₂-N-(-)-jasmonoyl isoleucine $([^{2}H_{5}]JA-Ile)$ and $^{2}H_{5}$ -cis-12-oxo-phytodienoic acid $([^{2}H_{5}]OPDA)$ were obtained from 652 OlChemim Ltd., ²H₅-jasmonic acid ([²H₅]JA) from CDN Isotopes and ²H₅-dinor-12-oxo-653 654 phytodienoic acid ($[^{2}H_{5}]$ dnOPDA) from Cayman Chemical Co.

Endogenous JA, JA-Ile, OPDA, dnOPDA, OPC-4, OPC-6, tnOPDA, 5-ddh-JA and 4,5ddh-JA-Ile and the corresponding ${}^{2}H_{5}$ -phytohormones in plants were analyzed using high performance liquid chromatography-electrospray-high-resolution accurate mass spectrometry (HPLC-ESI-HRMS). The hormones were extracted and purified as follows: 0.25 g frozen

plant tissue (ground to a powder in a mortar with liquid N₂) was homogenized with 2.5 ml 659 precooled (-20°C) methanol:water:HCOOH (90:9:1, v/v/v with 2.5 mM Na-660 diethyldithiocarbamate) and 25 µl of a stock solution of 1000 ng ml⁻¹ deuterium-labeled 661 internal standards [²H₅]JA and [²H₅]dnOPDA, 200 ng ml⁻¹ [²H₅]JA-Ile and 400 ng ml⁻¹ 662 663 $[^{2}H_{5}]$ OPDA in methanol. Samples were extracted by shaking in a Multi Reax shaker 664 (Heidolph Instruments) (60 min, 2,000 rpm, room temperature). After extraction, solids were 665 separated by centrifugation (10 min, 20,000 G, room temperature) in a Sigma 4-16K 666 Centrifuge (Sigma Laborzentrifugen), and re-extracted with an additional 1.25 ml extraction 667 mixture, followed by shaking (20 min) and centrifugation. Pooled supernatants (2 ml) were 668 separated and evaporated at 40°C in a RapidVap Evaporator (Labconco Co). The residue 669 was redissolved in 500 μ l methanol/0.133% acetic acid (40:60, v/v) and centrifuged (10 min, 670 20,000 RCF, room temperature) before injection into the HPLC-ESI-HRMS system.

671 Hormones were quantified using a Dionex Ultimate 3000 UHPLC device coupled to a 672 Q Exactive Focus Mass Spectrometer (Thermo Fisher Scientific) equipped with an HESI(II) 673 source, a quadrupole mass filter, a C-trap, a HCD collision cell and an Orbitrap mass 674 analyzer, using a reverse-phase column (Synergi 4 mm Hydro-RP 80A, 150 x 2 mm; 675 Phenomenex). A linear gradient of methanol (A), water (B) and 2% acetic acid in water (C) 676 was used: 38% A for 3 min, 38% to 96% A in 12 min, 96% A for 2 min and 96% to 38% A 677 in 1 min, followed by stabilization for 4 min. The percentage of C remained constant at 4%. Flow rate was 0.30 ml min⁻¹, injection volume 40 μ l, and column and sample temperatures 678 679 were 35 and 15°C, respectively. Ionization source working parameters were optimized (see 680 Supplementary Table 2).

For phytohormone detection and quantification, we used a full MS experiment with MS/MS confirmation in the negative-ion mode, using multilevel calibration curves with the internal standards. MS^1 extracted from the full MS spectrum was used for quantitative

analysis, and MS^2 for confirmation of target identity. For full MS, a m/z scan range from 62 684 685 to 550 was selected, resolution set at 70.000 full width at half maximum (FWHM), automatic gain control (AGC) target at 1e⁶ and maximum injection time (IT) at 250 ms. A mass 686 687 tolerance of 5 ppm was accepted. The MS/MS confirmation parameters were resolution of 17,500 FWHM, isolation window of 3.0 m/z, AGC target of 2e⁵, maximum IT of 60 ms, loop 688 689 count of 1 and minimum AGC target of 3e³. Instrument control and data processing were 690 carried out with TraceFinder 3.3 EFS software. Accurate masses of phytohormones and 691 internal standard are reported in Supplementary Table 3. Their principal fragments for these molecules are shown in Supplementary Table 3, with the exception of $[^{2}H_{5}]4.5$ -ddh-JA. 692 $[^{2}H_{5}]$ ddh-JA-Ile, $[^{2}H_{5}]$ OPC-4, $[^{2}H_{5}]$ OPC-6 and $[^{2}H_{5}]$ tnOPDA. 693

694 JAZ1-GUS degradation assays

The 35S:JAZ1-GUS in wild-type and *coi1-30* background were described ^{10,52}. The 695 696 35S:JAZ1-GUS marker line was introgressed into opr3-3 and jar1-1 backgrounds by crossing, 697 and double homozygous lines were used for further analyses. 35S:JAZ1-GUS seedlings were 698 grown vertically on MS plates and 6-day-old seedlings were treated for 1 h with jasmonate solution as described ⁵³. To visualize GUS activity, samples were placed in staining solution 699 and incubated (overnight, 37°C) as described in ⁵³. Tissue was then soaked several times in 700 701 75% ethanol and kept in 5% glycerol for photography with a Nikon D1-x camera. The 702 analysis was performed using 5-15 plants per sample. This experiment was repeated at least 703 three times with similar results.

704 Analysis of OPR1 and OPR2 activity

The open reading frames of Arabidopsis OPR1 and OPR2 were previously cloned into the expression vectors pQE-30 and pQE-31 (Qiagen) and kindly provided by Florian Schaller ^{12,54}. For protein expression, an over-night culture of the expression constructs in *E. coli* BL21-CodonPlus (DE3)-RIL (Agilent Technologies) was used to inoculate 400 mL LB

medium to an OD₆₀₀ of 0.01. The culture was grown at 37°C to OD₆₀₀ of 0.8, when protein 709 710 expression was induced by addition of 1 mM isopropyl-1-thio- β -D-galactopyranoside. After 711 4 h at 30°C, cells were harvested by centrifugation and lysed in BugBuster (5 mL per gram 712 packed cell weight; Merck Life Science) containing 1 mM PMSF. The lysates were cleared 713 by centrifugation (20000 xg, 20 min, 4° C), recombinant OPRs were purified by metal chelate 714 affinity chromatography on Ni-nitrilotriacetate (Ni-NTA) agarose (Qiagen) following the 715 supplier's protocols, and dialyzed against 25 mM Tris/HCl pH 7.5. The concentration of 716 OPR1 and OPR2 was determined spectrophotometrically at 445 nm using a molar extinction coefficient of $\epsilon_{445} = 11600 \text{ M}^{-1} \text{cm}^{-1}$. 717

718 Activity of OPR1 or OPR2 was measured spectrophotometrically by recording 719 NADPH consumption at 340 nm in disposable UV micro cuvettes. Activity assays were 720 performed under steady state conditions at 25°C in 0.2 mL 50 mM Na₂HPO₄/NaH₂PO₄ buffer 721 pH 7.0 containing 0.1 % (v/v) Triton X-100 and 20 mM glucose/1 U/ μ L glucose oxidase as 722 an oxygen consuming system. Reaction mixtures contained recombinant OPR1 or OPR2 at a 723 concentration of 1 µM, 200 µM NADPH, and the racemate (±)-4,5-didehydro-jasmonic acid 724 at a range of concentrations from 10 μ M to 1 mM. This racemate includes natural and 725 unnatural *trans* isomers. Apparent kinetic constants were derived by fitting the data to the 726 Michaelis-Menten equation by the non-linear least squares method using the Enzyme 727 Kinetics module 1.3 of the Sigmaplot version 10.0 (Systat Software GmbH). Assays were 728 performed using three technical replicates and data are shown as mean \pm SD. This experiment 729 was repeated twice with similar results.

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731 Online Methods References

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745		
746	Data A	vailability

- All data generated or analysed during this study are included in this published article (and its
- 748 supplementary information files).
- 749
- 750 Supplementary Results
- 751 Supplementary Fig. 1. Biosynthesis and intracellular flux of jasmonates in Arabidopsis
- 752 Supplementary Fig. 2. opr3 mutants in Arabidopsis
- 753 Supplementary Fig. 3. Responses of *opr3* mutants to fungal infection
- 754 Supplementary Fig. 4. Accumulation of JA derivatives in wild-type plants and opr3
- 755 mutants
- 756 Supplementary Fig. 5. Structures of 4,5-ddh-JA and 4,5-ddh-JA-Ile
- 757 Supplementary Fig. 6. Accumulation of 4,5-ddh-JA in wild-type plants and opr3 plants
- 758 Supplementary Fig. 7. Analysis of *OPR* expression and characterization of *opr* mutants
- 759
- 760 Supplementary Table 1. List of primers used
- 761 Supplementary Table 2. Ionization source working parameters
- 762 Supplementary Table 3. Masses of phytohormones and internal standard and their
- 763 principal fragments

Competing financial interests statement

- The corresponding author declares on behalf of all co-authors that there are no competing
- financial interests.



Figure 1. opr3-3 is a complete loss-of-function allele



Figure 2. opr3-3 mutants activate defense responses



Figure 3. JA accumulation in Col-0 and opr3-3 plants



Figure 4. 4,5-didehydro-JA triggers JA-regulated COI1-dependent responses



Figure 5. OPDA convention into 4,5-ddh-JA, JA, and JA-Ile



Figure 6. OPR1 and mainly OPR2 convert 4,5-ddh-JA into JA

Supplementary Information

Identification of an OPR3-independent pathway for jasmonate biosynthesis

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Supplementary Fig. 1. Biosynthesis and intracellular flux of jasmonates in Arabidopsis Scheme of the biosynthetic pathway of JA-Ile generated from plastidial membrane lipids. Pathway intermediates (described in the text) are 18:3 (α-linolenic acid), 16:3 (hexadecatrienoic acid), OPDA (12oxo-phytodienoic acid), dnOPDA(dinor-oxo-phytodienoic acid), OPC-8 (8-(3-oxo-2-(pent-2-enyl) cyclopentyl)octanoic acid), OPC-6 (6-(3-oxo-2-(pent-2-enyl)cyclopentyl)hexanoic acid), OPC-4 (4-(3oxo-2-(pent-2-enyl)cyclopentyl)butanoic acid), JA (jasmonic acid) and JA-Ile (jasmonoyl-Isoleucine). Black arrows define the octadecanoid pathway, blue arrows indicate the parallel hexadecanoid pathway. Biosynthetic enzymes (described in the text) are shown in green: 13-LOX (13-lipoxygenase), AOS (allene oxide synthase), AOC (allene oxide cyclase), OPR3 (OPDA reductase 3) and JAR1 (jasmonic acid-amido synthetase).



Supplementary Fig. 2. opr3 mutants in Arabidopsis

(a) Scheme of the *OPR3/AT2G06050* locus, including the T-DNA insertion that defines the *opr3-1* allele ^{18,33} in Ws accession and *opr3-3* (SK24765) allele in Col-0 accession.

(b) Expression of *OPR3* after fungal infection and wounding of Col-0 and mutant plants (n = 10) measured by real-time PCR in untreated plants (control, C) and plants challenged for 3 days with *B. cinerea* (*Bc*) or 9 days with *A. brassicicola* (*Ab*). Values are the mean \pm SD of three technical replicates expressed as relative fold change normalized to *ACT8*. Experiments were repeated twice with similar results. Statistically significant expression compared to untreated plants is highlighted (Student's t-test; ** p <0.01; *** p <0.001).



Supplementary Fig. 3. Responses of opr3 mutants to fungal infection

(a) Expression of JA-regulated genes after fungal infection, insect challenge and wounding of Col-0 and mutant plants (n = 10). *ChiB*, *Thio2.1*, *JAZ10* and *JAZ5* were measured by real-time PCR in untreated plants (control, C) and plants challenged for 3 days with *B. cinerea* (*Bc*) or 9 days with *A. brassicicola* (*Ab*), or 48 h with *S. littoralis* larvae (*Sl*) or 30 min after wounding (W) (n = 10). Values are the mean \pm SD of three technical replicates expressed as relative fold change normalized to *ACT8*. Experiments were repeated twice with similar results. Statistically significant wound-induced expression compared to untreated plants are highlighted (Student's t-test; * p <0.05; ** p <0.01; *** p <0.001).

(b and c) Infection of Arabidopsis plants (n = 15) with *B. cinerea*. Quantification of the spores of *B. cinerea* grown on different mutant lines was calculated 72 h post-inoculation. Data shown as box-plots of mean of 5 biological replicates of 5 leaves each. Letters above columns indicate significant differences (one-way ANOVA/post-hoc Tukey HSD Test, p <0.05). The experiment was repeated three times with similar results. Representative leaves of plants infected with *B. cinerea* are shown in (c). Scale bars, 1 cm.

(d and e) Infection of Arabidopsis plants with *A. brassicicola*. Fungal spores, grown on different mutant lines were counted 9 days post-inoculation, are shown as box-plots of mean of 5 biological replicates of 5 leaves each. Letters above columns indicate significant differences (one-way ANOVA/post-hoc Tukey HSD Test, p < 0.05). The experiment was repeated three times with similar results. Representative leaves of plants infected with *A. brassicicola* are shown in (e). Scale bars, 1 cm.

(f) Expression of JA-regulated genes after fungal infection and wounding of Col-0 and mutant plants (n = 10). *PDF1.2*, *JAZ7* and *AOS* were measured by real-time PCR in untreated plants (control, C) and plants challenged for 3 days with *B. cinerea* (*Bc*) or 9 days with *A. brassicicola* (*Ab*) or 30 min after wounding (W) (n = 10). Values are the mean \pm SD of three technical replicates expressed as relative fold change normalized to *ACT8*. Experiments were repeated twice with similar results. Statistically significant wound-induced expression compared to untreated plants is highlighted (Student's t-test; ** p <0.01; *** p <0.001).



Supplementary Fig. 4. Accumulation of JA derivatives in wild-type plants and *opr3* mutants Accumulation of OPC-4 (pmoles/fresh weight (g)) (a-b), JA (pmoles/fresh weight (g)) (c) and JA-Ile (pmoles/fresh weight (g)) (d) in wounded wild-type (grey bars) and *opr3* mutants (black bars) plants. Four-week-old plants (n = 10) were wounded and damaged leaves collected at the time indicated. Data for *opr3-3* compared to Col-0 are shown in (a), whereas *opr3-1* compared to Ws is shown in (b) (c) and (d). Data shown as mean \pm SD of four biological replicates. The experiment was repeated three times with similar results.





(+)-4,5-ddh-JA (9)

(-)-4,5-ddh-JA (10)





(+)-4,5-ddh-JA-(S)-lle (**11**)

(-)-4,5-ddh-JA-(S)-lle (12)



Supplementary Fig. 5. Structures of 4,5-ddh-JA and of 4,5-ddh-JA-Ile

(a) Chemical structures of 4,5-ddh-JA, used as an enantiomeric mixture of the (-)- and (+)-forms, (-)-4,5-ddh-JA-Ile and (+)-4,5-ddh-JA-Ile.

(b) Chromatogram of pure (+)-4,5-ddh-JA-Ile (top panel) and of WT and *opr3-3* plants.



Supplementary Fig. 6. Accumulation of 4,5-ddh-JA in wild-type plants and *opr3* plants (a and b) Accumulation of 4,5-ddh-JA (pmoles/fresh weight (g)) in wounded wild-type (grey bars) and *opr3* mutants (black bars) plants. Four-week-old plants (n = 10) were wounded and damaged leaves collected at the time indicated. Data for *opr3-3* compared to Col-0 are shown in (a), whereas *opr3-1* compared to Ws are shown in (b). Data shown as mean \pm SD of four biological replicates. The experiment was repeated three times with similar results.



Supplementary Fig. 7. JA accumulation after exogenous 4,5-ddh-JA treatment in Col-0 and *opr3-3* plants. Data shown as mean \pm SD of four biological replicates after subtraction of basal JA levels. JA accumulation was not statistically different (Student's t-test p <0.01) in Col-0 compared to *opr3-3* plants (n = 10). The experiment was repeated twice with similar results.



Supplementary Fig. 8. Analysis of OPR expression and characterization of opr mutants

(a) *OPR1*, *OPR2* and *OPR3* expression in Col-0 and *opr3-3* mutant plants (n = 10) 30 min after wounding (W, dark grey bars). Unwounded plants were included as controls (C, light grey bars). Values are the mean \pm SD of three technical replicates expressed as relative fold change normalized to *ACT8*. The experiment was repeated three times with similar results.

(b and c) Genomic scheme of the loss-of-function opr1-1 (b) (SALK_145353) and opr2-1 (c) (SALK_116381) mutant lines and the exogenous T-DNA insertions.

(d) Expression analysis by RT-PCR of WT plants and T-DNA insertion lines of *OPR1* and *OPR2* using *EF1* as control transcripts. The experiment was repeated three times with similar results.

(e) Accumulation of JA-Ile (pmoles/fresh weight (g)) in wounded Col-0, *opr* single and double mutant plants (n =10). Five-week-old plants were wounded and damaged leaves collected at time 0 (white bars), 30 (light grey), 60 (dark grey) or 180 min (black) after wounding. Data shown as mean \pm SD of four biological replicates. The experiment was repeated three times with similar results.

(f) Box-plots of fungal spore quantification of Col-0 and mutant plants (n = 15) infected with *A*. *brassicicola* (9 dai). Letters above columns indicate significant differences (one-way ANOVA/post-hoc Tukey HSD Test, p < 0.01).

(g-h) Representative leaves of Col-0 and mutant plants infected with *B. cinerea* and *A. brassicicola* are shown in (g) and (h) respectively. Scale bars, 1 cm.

(i) Expression of JA-regulated *ChiB* and *PDF1.2* (in thousands) after fungal infection of Col-0 and mutant plants (n = 10) measured by real-time PCR in untreated plants (control, C) and plants challenged for 7 days with *B. cinerea* (*Bc*) or 9 days with *A. brassicicola* (*Ab*). Values are the mean \pm SD of three technical replicates expressed as relative fold change normalized to *ACT8*. Experiments were repeated twice with similar results. Letters above columns indicate significant differences compared to expression in *opr3-3* plants (Student's t-test, P < 0.05).

(j) Expression of early JA-regulated genes after wounding in Col-0 and mutant plants (n = 10). *AOS*, *JAZ1* and *MYC2* expression were measured by real-time PCR in untreated plants (control, C, grey bars) and in plants 1 h after wounding (W, dark grey bars). Values are the mean \pm SD of three technical replicates expressed as relative fold change normalized to *ACT8*. The experiment was repeated at least twice with similar results. Letters above columns indicate significant differences compared to expression in *opr3-3* plants (Student's t-test, P < 0.05).

Supplementary Table 1 List of primers used

Name	Sequence
OPR1 qPCR-F	5'-ATCCAGGAGCATTAGGGCT-3'
OPR1 qPCR-R	5'-CGCTTTCCTCATCGGCAT-3'
OPR2 qPCR-F	5'-TCCAGAAGCATTAGGGCTGT-3'
OPR2 qPCR-R	5'-TGATGTTGAAAGCACATATAAAAGC-3'
OPR3-3 qPCR-F	5'-GCATGGAAGCAAGTTGTGGAAGCA-3'
OPR3-3 qPCR-R	5'-CATGCGCCCCGTGGATCTCAAT-3'
OPR3-3 qPCR-F2	5'-ATCTCTCTCATCGAGTGGTT-3'
OPR3-3 qPCR-R2	5'-CCTCCATTAGGTTGATACACTG-3'
PDF1.2 qPCR-F	5'-CACCCTTATCTTCGCTGCTC-3'
PDF1.2 qPCR-R	5'-GTTGCATGATCCATGTTTGG-3'
CYP79B3 qPCR-F	5'-CTTTGCTTACCGCTGATGAA-3'
CYP79B3 qPCR-R	5'-GCGTTTGA TGGGTTGTCTG-3'
AOS qPCR-F	5'-GCGACGAGAGATCCGAAGA-3'
AOS qPCR-R	5'-CTCGCCACCAAAACAACAAA -3'
LOX3 qPCR-F	5'-CACTGCAATTCACAAGCAACC-3'
LOX3 qPCR-R	5'-CAAAGGAGGAATCGGAGAAGC-3'
JAZ1 qPCR-F	5'-CACGTCTGTGAGAAGCTAGGC-3'
JAZ1 qPCR-R	5'-TTCTGAGTTCGTCGGTAGCC-3'
JAZ5 qPCR-F	5'-AAAGATGTTGCTGACCTCAGTG-3'
JAZ5 qPCR-R	5'-CCCTCCGAAGAATATGGTCA-3'
JAZ7 qPCR-F	5'-TTCGGATCCTCCAACAATCCCA-3'
JAZ7 qPCR-R	5'-TCAAGACAATTGGATTATTATGTTACAGT-3'
MYC2 qPCR-F	5'-GTGCGGGATTAGCTGGTAAA-3'
MYC2 qPCR-R	5'-ATGCATCCCAAACACTCCTC-3'
ACT8 qPCR-F	5'-CCAGTGGTCGTACAACCGGTA-3'
ACT8 qPCR-R	5'-TAGTTCTTTTCGATGGAGGAGCTG-3'
Name	Sequence
LB_GW1	5'-GCTTTCGCCTATAAATACGACGGATCGT-3'
SLBb1.3	5'-ATTTTGCCGATTTCGGAAC-3'
OPR1_F1	5'-AACACACTACATTACATTATTGATAACA-3'
OPR2_F1	5'-GAAACACATTACATTACTGATAACACGA-3'
OPR3_F1	5'-GCATGGAAGCAAGTTGTGGAAGCA-3'

Supplementary Table 2

Ionization source working parameters

Instrumental parameters	Value
Sheath gas flow rate	44 au
Auxiliary gas flow rate	11 au
Sweep gas flow rate	1 au
Spray voltage	3.5 kV
Capillary temperature	340 °C
S-lens RF level	50
Auxiliary gas heater temperature	300 °C

Supplementary Table 3

Masses of phytohormones and internal standard and their principal fragments

Analyte	[M-H] ⁻¹ Phytohormone
JA	209.11832
Ja-Ile	322.20238
OPDA	291.19657
dn-OPDA	263.16527
4,5-ddh-JA	207.10267
4,5-ddh-JA-Ile	320.18673
OPC-4	237.14962
OPC-6	265.18092
² H ₅ -JA	214.1497
² H ₅ -JA-Ile	327.23377
² H ₅ -OPDA	296.22795
² H ₅ -dnOPDA	268.19665
² H ₅ -tnOPDA	235.13397
² H ₅ -4,5-ddh-JA	212.13405
² H ₅ -ddh-JA-Ile	325.21812
² H ₅ -OPC-4	242.181
² H ₅ -OPC-6	270.2123
² H ₅ -JA	214.1497
² H ₅ -JA-Ile	324.21494
² H ₅ -OPDA	296.22795
² H ₅ -dnOPDA	268.19665
	·
Analyte	[M-H] ⁻¹ Fragment
JA	59.01297
T T1	120.00725

JA	59.01297
Ja-Ile	130.08735
OPDA	165.12843
dn-OPDA	165.12843
4,5-ddh-JA	163.11282
4,5-ddh-JA-Ile	130.08735
OPC-4	125.09715
OPC-6	96.95968
² H ₅ -JA	61.02555
² H ₅ -OPDA	170.15994
² H ₅ -dnOPDA	170.15994
² H ₅ -JA	61.02555
² H ₅ -JA-Ile	131.0937
² H ₅ -OPDA	170.15994
² H ₅ -dnOPDA	170.15994

Supplementary Note 1

Chemical synthesis

Analytical and chromatographical methods. Gas chromatography-mass spectrometry (GC-MS) was carried out using an Agilent mass selective detector model 5977E connected to an Agilent model 7820A gas chromatograph. A capillary column of 5% phenylmethylsiloxane (12 m, 0.33 μ m film thickness) with helium as the carrier gas was used. The temperature was raised from 80°C to 320°C at a rate of 10°C/min. Reversed-phase HPLC (RP-HPLC) was carried out using a column of 250 x 10 mm Nucleosil 100-7 C₁₈ eluted with methanol-water-acetic acid (55:45:0.02, v/v/v) at a flow rate of 4 mL/min, whereas straight-phase HPLC (SP-HPLC) was performed with a column of Nucleosil 50-7 (250 x 10 mm) using the solvent systems indicated at a flow rate of 4 mL/min. NMR spectra were recorded on CDCl₃ solutions using Bruker 500 or 600 MHz instruments.

Chemicals. Methyl (±)- jasmonate, L-isoleucine and all other chemicals used were purchased from Sigma-Aldrich.

Methyl (\pm)-4,5-didehydrojasmonate (13). The title compound was prepared by modification of a previously described protocol ⁵⁴. Thus, methyl (\pm)-jasmonate (1 mmol, 224 mg) was added to dry *N*,*N*-dimethylformamide (3 mL) containing diethyl allyl phosphate (2 mmol, 388 mg), Na₂CO₃ (2.4 mmol, 254 mg) and palladium(II) acetate (0.12 mmol, 27 mg). The solution was purged with argon and stirred at 80°C for 24 h. Water was added and the mixture extracted with diethyl ether. After drying over MgSO₄ the solvents were evaporated leaving a residue of 250 mg. Purification was performed on a silica gel column (5 g) which was eluted with diethyl ether/hexane (1:9, v/v), 15 fractions of 15 mL. Unreacted methyl jasmonate was

recovered in fractions 5-7 and methyl 4,5-ddh-JA was present in fractions 9-14. The latter were combined and evaporated, leaving 76 mg of the title compound (yield, 34%) accompanied by 4 mg of an unknown allyl adduct of methyl 3,7-ddh-JA and traces of methyl 7,8-ddh-JA and methyl 3,7-ddh-JA. Preparative SP-HPLC using a solvent system of 2-propanol-hexane (1:99, v/v) afforded the pure title compound as a colorless oil. The mass spectrum showed prominent ions at m/z 222 (30%, M⁺), 193 (15, M⁺ - C₂H₅), 167 (13), 154 (70, rearrangement with loss of the C-8 to C-12 side chain), 133 (30), 107 (25), and 95 (100). The UV spectrum (EtOH) showed λ_{max} 217 nm and the ¹H NMR spectrum showed signals at δ 0.98 (3H, t, J = 7.5 Hz), 2.05-2.14 (3H, m), 2.30-2.37 (1H, m), 2.49 (1H, dd, J = 15.8, 8.3 Hz), 2.50-2.58 (1H, m), 2.60 (1H, dd, J = 15.8, 6.7 Hz), 3.00-3.06 (1H, m), 3.72 (3H, s), 5.28 (1H, dtt, J = 10.8, 7.5, 1.7 Hz), 5.49 (1H, dtt, J = 10.8, 7.4, 1.7 Hz), 6.20 (1H, dd, J = 5.8, 2.0 Hz), and 7.64 (1 H, dd, 5.8, 2.4 Hz). This spectrum was in full agreement with that previously published ⁵⁴.

Methyl (+)-(*3S*,*7R*)-*4*,*5*-*didehydrojasmonate* (**14**). The (+)-rotatory form of the side chain *trans* isomer of methyl 4,5-didehydrojasmonate is the 3S,7R enantiomer ⁵⁵ and is the form which is stereochemically related to natural methyl (-)-jasmonate (3R,7R). It should be noted that introduction of the ring double bond reverses the sign of optical rotation and that the change in configurational assignment at C-3 is a consequence of the Cahn-Ingold-Prelog rules. In the present work, the title compound was prepared by palladium-catalyzed dehydrogenation of the methyl ester of (-)-JA (22 mg), which was available since a previous study ⁵⁶. The material obtained (9 mg) showed λ_{max} (EtOH) 217 nm and the mass spectrum was identical to that given above for the (±) form. (±)-4,5-Didehydrojasmonic acid (6). Methyl (±)-4,5-didehydrojasmonate (100 mg, 0.45 mmol) was added to a solution of 24 mg of LiOH (1 mmol) in 6 mL of water and 24 mL of tetrahydrofuran. The solution was stirred at 23°C for 18 h. The product obtained following extraction with diethyl ether was subjected to preparative SP-HPLC using a solvent system of 2-propanol-hexane-acetic acid (4:96:0.02, v/v/v). This provided the pure title compound as a colorless oil (59 mg) showing λ_{max} 217 nm. The mass spectrum of a methyl-esterified sample was identical to that of methyl 4,5-ddh-JA, and the mass spectrum of the trimethylsilyl (Me₃Si) ester derivative showed prominent ions at m/z 280 (22%, M⁺), 251 (7, M⁺ - C₂H₅), 212 (18, rearrangement with loss of the C-8 to C-12 side chain), 148 (21), 117 (15, O=C=O⁺SiMe₃) and 73 (100, Me₃Si⁺).

(+)-(3S,7R)-4,5-Didehydrojasmonic acid (9). Methyl (+)-4,5-didehydrojasmonate (9 mg, 0.04 mmol) was treated with LiOH (2.4 mg, 0.1 mmol) using the above-described protocol. The pure title compound was obtained following SP-HPLC as described above. Its properties including the UV spectrum and mass spectrum were identical to those recorded for the racemic compound.

Coupling of (\pm) -4,5-didehydrojasmonic acid (6) to (S)-isoleucine. (\pm) -4,5-Didehydrojasmonic acid (90 mg, 433 µmol) was dissolved in 18 mL of redistilled ethyl acetate containing 9.8 mg (97 µmol) triethylamine. *O*-(Benzotriazol-1-yl)-*N*,*N*,*N'*,*N'*-tetramethyluronium tetrafluoroborate (TBTU, 161 mg, 501 µmol) was added and the solution stirred at 23°C for 30 min. (S)-Isoleucine (171 mg, 1303 µmol) suspended in 9 mL of dry *N*,*N*-dimethylformamide was added and the mixture stirred at 23°C for 18 h. Extraction with ethyl acetate at pH 2 afforded a product that was purified on a silica gel column (5 g). Elution with ethyl acetate/chloroform/acetic acid (4:6:0.05, v/v/v) afforded a diasteromeric mixture of comparable amounts of the (*S*)isoleucine conjugates of (+)- and (-)-4,5-ddh-JA. These were resolved by preparative RP-HPLC which afforded an earlier eluting and a later eluting conjugate (retention volumes, 86 and 104 mL, respectively). The methyl esters of these diastereomers showed a small separation on GC-MS analysis where, under the conditions used, the earlier-eluting and later-eluting conjugates had retention times of 16.68 and 16.74 min, respectively.

In view of previous data recorded for the (+)- and (-) (*S*)-isoleucine conjugates of (\pm)-JA ⁸, the results suggested that the earlier and later eluting 4,5-ddh-JA conjugates obtained as described above were the (-)- and (+)-isomers, respectively. In order to firmly establish this point a separate experiment was conducted in which (+)-4,5-ddh-JA (5 mg) was coupled to (*S*)-isoleucine. Analysis of the product using RP-HPLC showed a single peak which matched that ascribed to the (+) conjugate obtained from (\pm)-4,5-ddh-JA, thus unequivocally showing that the diastereomer that eluted faster on RP-HPLC was the (-)- or (3R,7S) isomer, whereas the slower-eluting diastereomer was the (+)- or (3S,7R) isomer.

N-[(+)-(3S,7R)-4,5-didehydrojasmonoyl]-(S)-isoleucine (11). The above-mentioned later eluting (S)-isoleucine conjugate (16 mg) was a colorless semisolid whose UV spectrum (EtOH) showed λ_{max} 217 nm (ϵ 10,800). An aliquot was dissolved in 50 µL of methanol and treated with diazomethane and taken to dryness after about 5 s (longer times of treatment led to the formation of byproducts). Analysis by GC-MS showed a single peak and the mass spectrum showed prominent ions at m/z 335 (27%, M^+), 306 (16, M^+ - C₂H₅), 276 (22, M^+ - CH₃COO), 190 (11, M^+ - [NH-CH(OOCCH₃)-C₄H₉ + H]), 146 (44, [NH-CH(OOCCH₃)-C₄H₉ + 2 H]), 128 (34, [CH(OOCCH₃)-C₄H₉ - H]), and 86 (100, 146 - CH₃COOH). The structure (Suppl. Figure 5a) was fully supported by the ¹H and ¹³C NMR spectra (Supplementary Note Table 1).

N-[(-)-(3R,7S)-4,5-didehydrojasmonoyl]-(S)-isoleucine (12). The earlier eluting (S)isoleucine conjugate (15 mg) showed λ_{max} 217. An aliquot was treated with diazomethane (5-10 s) and gave a single peak on GC-MS. The mass spectrum was virtually identical to that of the (+)-conjugate. The structure (Supplementary Fig. 5a) was fully supported by the ¹H and ¹³C NMR spectra (Supplementary Note Table 2). of N-[(+)-(3S,7R)-4,5-didehydrojasmonoyl]-(S)-Alkali-promoted cyclization *isoleucine (11).* A sample of *N*-[(+)-4,5-didehydrojasmonoyl]-(*S*)-isoleucine (34 mg) was dissolved in methanol (4.5 mL) and 2 M NaOH in water (0.5 mL) was added. The solution was kept at 23°C for 1 h and then acidified and extracted with ethyl acetate. The product was purified by RP-HPLC, which showed a main peak at a retention volume of 100 mL. This material was subjected to SP-HPLC using a solvent system of ethanol-hexane-acetic acid (5:95:0.1, v/v/v). Two peaks appeared, *i.e.* cyclization product (CP)-1 (15) (5 mg; 60 mL effluent) and CP-2 (16) (17 mg; 76 mL effluent) (Supplementary Note Fig. 1). The UV spectra of these two materials lacked the absorption band present in the parent compound, thus indicating the disappearance of the conjugated ring double bond.

Cyclization product-2 (**16**). GC-MS analysis of the methyl ester of CP-2 showed prominent ions at m/z 335 (10%, M⁺), 306 (5, M⁺ - C₂H₅), 279 (30, M⁺ - C₄H₈), 276 (100, M⁺ - COOCH₃), 250 (26, 306 - C₄H₈), 247 (24, 306 - COOCH₃) and 208 (21). This spectrum combined with the lack of the ring double bond as indicated by UV spectroscopy suggested a bicyclic structure in which the nitrogen atom of the isoleucine residue had attacked C-4 of the didehydrojasmonoyl residue (Supplementary Note Fig. 1). An alternative reaction involving Michael attack by the α carbon anion of the isoleucine moieity on C-4 could be ruled out by NMR spectrometry (Supplementary Note Table 3).

Cyclization product.1 (15). On GC-MS, the methyl ester of CP-1 eluted somewhat later compared to CP-2 (retention times, 16.8 and 16.1 min, respectively). Its mass spectrum was similar to that of the methyl ester of CP-2 and showed prominent ions at m/z 335 (8%, M⁺), 306 (2, M⁺ - C₂H₅), 279 (22, M⁺ - C₄H₈), 276 (100, M⁺ - COOCH₃), 250 (5, 306 - C₄H₈), 247 (12, 306 - COOCH₃) and 208 (15). Treatment of CP-1 with NaOH produced an equilibrium mixture of CP-2 and CP-1 in proportions about 3:1, thus indicating that the 2(*Z*)-pentenyl side chain is *cis*-oriented with respect to the amide ring in CP-1 and *trans*-oriented in CP-2 (Supplementary Note Fig. 1).

Supplementary Note Table 1

Pos.	δ ¹³ C [ppm]	¹³ C mult.	δ ¹ H [ppm] mult. (J[Hz]]
1	170.7	S	-
2	40.5	t	2.543 dd (14.5;6.5) / 2.360 dd (14.5;8.6)
3	43.9	d	3.097 m
4	166.2	d	7.700 dd (5.8;2.4)
5	133.6	d	6.183 dd (5.8,2.0)
6	210.7	s	-
7	51.0	d	2.08
8	27.8	t	2.518 m / 2.314 br dt (14.4;7.3)
9	124.5	d	5.276 dtt (10.8;7.5;1.6)
10	134.6	d	5.475 dtt (10.8;7.3;1.5)
11	20.6	t	2.066 dqd (7.4;7.4;1.6)
12	14.2	q	0.959 t (7.6)
1'	56.4	d	4.630 dd (8.4;4.8)
2'	37.4	d	2.066 dqd (7.4;7.4;1.6)
3'	25.1	t	1.496 dqd (13.6;7.5;4.6) / 1.213 m
4'	11.6	q	0.954 t (7.5)
5	15.5	q	0.969 d (6.8)
6'	174.8	S	-
NH	-	-	6.027 d (8.4)

NMR data of (+)-4,5-didehydro-JA-Ile (solvent: CDCl₃)

¹H chemical shifts with only two decimal places are chemical shifts of HSQC correlation peaks

Supplementary Note Table 2

Pos.	δ ¹³ C [ppm]	¹³ C mult.	δ ¹ H [ppm] mult. (J[Hz]]
1	170.6	S	-
2	40.4	t	2.518 dd (14.6;6.9) / 2.395 dd (14.6;8.2)
3	43.8	d	3.097 m
4	166.2	d	7.661 dd (5.7;2.4)
5	133.6	d	6.182 dd (5.7,2.0)
6	210.8	S	-
7	51.0	d	2.101 ddd (7.8;4.7;2.4)
8	27.8	t	2.507 m / 2.312 br dt (14.5;7.8)
9	124.4	d	5.249 dtt (10.8;7.5;1.5)
10	134.6	d	5.464 dtt (10.8;7.3;1.4)
11	20.6	t	2.059 dqd (7.4;7.4;1.5)
12	14.2	q	0.953 ^a t (7.6)
1'	56.4	d	4.653 dd (8.5;4.7)
2'	37.5	d	1.971 m
3'	25.1	t	1.494 dqd (13.6;7.5;4.5) / 1.215
			ddq(13.6;9.3;7.5)
4'	11.6	q	0.952 ^a t (7.6)
5'	15.5	q	0.970 d (6.8)
6'	175.1	S	-
NH	-	-	6.066 d (8.6)

NMR data of (-)-4,5-didehydro-JA-Ile (solvent: CDCl₃)

^a may be interchanged

Supplementary Note Table 3

Pos.	δ ¹³ C [ppm]	¹³ C mult.	δ ¹ H [ppm] mult. (J[Hz]]
1	174.9	s	-
2	37.0	t	2.71 / 2.390 m
3	38.7	d	2.68
4	58.0	d	4.301 td (7.2; 3.4)
5	43.4	t	2.71 / 2.541 dd (19.3; 7.2)
6	216.3	s	-
7	53.1	d	2.184 m
8	27.0	t	2.450 m / 2.274 dtd-like (14.4;8.1;1.2)
9	124.0	d	5.256 dtt (10.8;7.6;1.6)
10	134.9	d	5.520 dtt (10.8;7.4;1.4)
11	20.6	t	2.046 dqd (7.6;7.6;1.6)
12	14.1	q	0.956 (7.6)
Me	52.0	q	3.715 (s)
1´	60.4	d	4.318 d (10.6)
2´	35.2	d	2.03
3´	25.1	t	1.373 dqd (13.7;7.4;3.1) / 1.054 ddq
			(13.7;9.7;7.4)
4´	10.8	q	0.871 t (7.6)
5´	16.3	q	0.950 d (6.6)
6´	170.4	S	-

NMR data of CP-2 methyl ester (solvent: CDCl₃)

¹H chemical shifts with only two decimal places are chemical shifts of HSQC correlation peaks



Supplementary Note Figure 1.

(+)-4,5-didehydrojasmonoyl]-(*S*)-isoleucine (11), CP-2 (16) and CP-1 (15)

Alkali-promoted conversions of N-[(+)-4,5-didehydrojasmonoyl]-(S)-isoleucineshowing attack by the isoleucine nitrogen at C-4 of the didehydrojasmonoyl residue.The primary product CP-2 is in base-catalyzed equilibrium with CP-1, in which the2(Z)-pentenyl side chain of the jasmonoyl residue is cis-oriented with respect to theamide ring.

Supplementary Note References

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