



OXIDATIVE SIGNAL-INDUCIBLE1 induces immunity by coordinating N-hydroxypipecolic acid, salicylic acid, and camalexin synthesis

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Summary

- Expression of OXIDATIVE SIGNAL-INDUCIBLE1 (OXI1) is induced by a number of stress conditions and regulates the interaction of plants with pathogenic and beneficial microbes.
- In this work, we generated Arabidopsis *OXI1* knockout and genomic *OXI1* overexpression lines and show by transcriptome, proteome, and metabolome analysis that *OXI1* triggers *ALD1*, *SARD4*, and *FMO1* expressions to promote the biosynthesis of pipecolic acid (Pip) and N-hydroxypipecolic acid (NHP).
- OXI1 contributes to enhanced immunity by induced SA biosynthesis via CBP60g-induced expression of SID2 and camalexin accumulation via WRKY33-targeted transcription of PAD3. OXI1 regulates genes involved in reactive oxygen species (ROS) generation such as RbohD and RbohF. OXI1 knock out plants show enhanced expression of nuclear and chloroplast genes of photosynthesis and enhanced growth under ambient conditions, while OXI1 overexpressing plants accumulate NHP, SA, camalexin, and ROS and show a gain-of-function (GOF) cell death phenotype and enhanced pathogen resistance. The OXI1 GOF phenotypes are completely suppressed when compromising N-hydroxypipecolic acid (NHP) synthesis in the fmo1 or ald1 background, showing that OXI1 regulation of immunity is mediated via the NHP pathway.
- Overall, these results show that OXI1 plays a key role in basal and effector-triggered plant immunity by regulating defense and programmed cell death via biosynthesis of salicylic acid, N-hydroxypipecolic acid, and camalexin.

Introduction

As sessile organisms, plants have developed various strategies to survive different stresses. Their ability to adapt to changing environments largely relies on sensing and signaling processes, which are based on receptors and signaling networks. Plant plasma membrane-localized pattern-recognition receptors (PRRs) perceive microbe-derived molecular patterns, leading to pattern-triggered immunity (PTI) (Alhoraibi *et al.*, 2019). Pattern-triggered immunity is the first layer of plant immunity and is essential for plants to resist diverse pathogens (DeFalco & Zipfel, 2021). Among the best-known PRRs is FLAGELLIN SENSITIVE2 (FLS2), which perceives a conserved 22-amino acid epitope (flg22) of the N-

To overcome PTI, pathogens deliver effector proteins into host cells, resulting in effector-triggered susceptibility (ETS). Resistance to pathogens can occur when specific intracellular receptors, called resistance (R) proteins, recognize race-specific avirulence (avr) effectors. Under these conditions, the R proteins

terminus of bacterial flagellin (Gómez-Gómez & Boller, 2000). Upon ligand binding, FLS2 interacts with BRASSINOSTEROID INSENSITIVE1-ASSOCIATED KINASE1 (BAK1) to form an active receptor complex (Chinchilla *et al.*, 2007). Within minutes, PRRs trigger downstream responses, including a transient influx of calcium ions from the apoplast, activation of calcium-dependent protein kinases, the production of extracellular reactive oxygen species (ROS), and the activation of mitogen-activated protein kinases (MAPKs) (Ligterink *et al.*, 1997; Jeworutzki *et al.*, 2010; Ranf *et al.*, 2011; Bigeard *et al.*, 2015).

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activate strong defense reactions, termed effector-triggered immunity (ETI) (Jones & Dangl, 2006). Effector-triggered immunity usually triggers a localized cell death at the infection site, called the hypersensitive response (HR), which restricts pathogen growth (Caplan *et al.*, 2008).

Plant defense comprises PTI and ETI, and a successful defense against pathogens relies on mutual potentiation of immunity by both PTI and ETI components (Pruitt et al., 2021). Patterntriggered immunity and ETI are associated with the induction of systemic acquired resistance (SAR) in uninfected distal parts of the plant, which provides broad-spectrum resistance against subsequent pathogen challenges (Gruner et al., 2013). The establishment of SAR in response to an inducing pathogen inoculation requires the accumulation and coordinated action of the immune-regulatory metabolites salicylic acid (SA) and Nhydroxypipecolic acid (NHP) (Nawrath & Métraux, 1999; Hartmann et al., 2018; Hartmann & Zeier, 2019; Zeier, 2021). In Arabidopsis, the ISOCHORISMATE SYNTHASE1 (ICS1)catalyzed conversion of the shikimate pathway intermediate chorismate to isochorismate is a key biochemical event for the pathogen-induced biosynthesis of SA (Wildermuth et al., 2001). ENHANCED DISEASE SUSCEPTIBILITY5 exports isochorismate from the plastid to the cytosol, where the cytosolic amidotransferase avrPphB SUSCEPTIBLE3 (PBS3) conjugates glutamate to isochorismate to produce isochorismate-9glutamate, which spontaneously decomposes into SA (Rekhter et al., 2019; Torrens-Spence et al., 2019).

N-hydroxypipecolic acid is synthesized through N-hydroxylation of the nonprotein amino acid pipecolic acid (Pip) by flavin-dependent monooxygenase (FMO1) (Chen *et al.*, 2018; Hartmann *et al.*, 2018). Pipecolic acid is derived from L-Lys by consecutive transamination and reduction steps that involve the aminotransferase AGD2-LIKE DEFENSE RESPONSE PROTEIN1 (ALD1) and the reductase SAR-DEFICIENT4 (SARD4), respectively (Návarová *et al.*, 2012; Ding *et al.*, 2016; Hartmann *et al.*, 2017).

The OXIDATIVE SIGNAL-INDUCIBLE1 (OXII) protein kinase has emerged as a potential player linking the activation of MPK3/6 to disease resistance (Rentel et al., 2004; Petersen et al., 2009). OXIDATIVE SIGNAL-INDUCIBLE1 (OXI1) also plays a role in coordinating the beneficial interaction with the fungus Piriformospora indica (Camehl et al., 2011). Interestingly, OXI1 is regulated transcriptionally and at the protein kinase level by hydrogen peroxide (H₂O₂) (Rentel et al., 2004). Reactive oxygen species of plants are either by-products of photosynthesis or are induced during abiotic stresses and pathogen attack. Among the different forms of ROS, plants produce singlet oxygen (¹O₂), superoxide radicals (O₂⁻), or H₂O₂ (Apel & Hirt, 2004). Whereas superoxide can spontaneously or enzymatically convert to H₂O₂ and hydroxyl radicals in the presence of metals, the production of these ROS can also increase under stress conditions when the absorbed light energy supersedes the amount of useful energy passing through the photosynthetic electron transport chain. Under these conditions, OXI1 regulates high light-induced programmed cell death via singlet oxygen signaling (Beaugelin et al., 2019).

Major advances in understanding ROS-producing enzymes, ROS sensing as well as compartmentalization and systemic ROS signaling have recently been made (Waszczak et al., 2018; Castro et al., 2021). In this study, we show that OXII plays an important role in regulating the trade-off between plant growth and defense. Using transcriptomics, proteomics and metabolomics analyses, we provide evidence that OXII is a negative regulator of photosynthesis, but positively regulates biosynthesis of key defense factors, ROS, and camalexin. Importantly, we show that OXII regulates various steps in the biosynthesis of NHP and SA, the two key regulators of SAR. Accordingly, OXII knockout and overexpressor plants show opposite behavior in plant growth and both basal and ETI. Taken together, our study shows that OXII coordinates plant defense responses and growth.

Materials and Methods

Plant material and growth conditions

Arabidopsis thaliana Columbia (Col-0) was used as wild-type (WT) in this study. The 2.2 kb *OXII* promoter and the genomic sequence of *OXII* with the 5' UTR and 3' UTR were amplified by polymerase chain reaction (PCR) and cloned into pCambia 3300 carrying BASTA resistance (Forzani *et al.*, 2011). The HA tag was cloned at the Sal I site found at the ATG site of *OXII*. The construct was used to transform Col-0 *oxi1-2* (Gabi_355H08) using the floral dip method and analyzed by qRT-PCR and immunoblotting with HA antibody. For *OXII* expression analysis, an *OXII::GUS* line was used (Forzani *et al.*, 2011).

Arabidopsis plants were grown individually in Jiffy-7 pellet in a controlled cultivation chamber with a 12 h : 12 h, light : dark at 110 $\mu mol~m^{-2}~s^{-1}light$ intensity and a relative humidity of 60%. Day and night temperatures were set to 22°C and 20°C, respectively. Nonflowering 5-wk-old plants were used for bacterial disease assay and ROS burst assay. For microarray analysis of seedlings, surface-sterilized seeds were vernalized at 4°C in the dark for 3 d and then germinated on 1/2 strength Murashige and Skoog (1/2MS, M6899; Sigma) plates, containing 0.5% sucrose (S5016; Sigma), 1% agar (A1296; Sigma), and 0.5% MES (M8250; Sigma), grown at 23°C, 60% humidity, and 125 $\mu mol~m^{-2}~s^{-1}$ light intensity with a 16 h : 8 h, light : dark photoperiod. The plants were transferred to a new 60 mm \times 60 mm Petri dish with 10 ml 1/2MS medium overnight before treatment.

Immunoblotting

Immunoblotting was carried out as described (Frei dit Frey et al., 2014). Briefly, 10 µg of total protein extracts from roots was denatured with SDS loading buffer and then separated by 8% SDS/PAGE before transfer to polyvinylidene difluoride membranes (Millipore) by electroblotting. Membranes were probed with Ponceau S (P3504; Sigma) and anti-HA IgG (Covance Carnegie Center, Princeton, NJ, USA). Membranes were developed by enhanced chemiluminescence, as recommended by the manufacturer (GeneImage; Amersham Biosciences).

Pathogen infection assays

Spray and injection infections with Pst DC3000 and Pst DC3000 AvrRpm1 (kindly supplied by J. Parker; MPI, Cologne, Germany) and determination of bacterial titers were performed on 4- to 5wk-old plants as previously described (Berriri et al., 2012). Spray inoculations were performed with bacterial suspensions at 108 CFU ml⁻¹ in 10 mM MgCl2 with 0.04% Silwet L-77. Infiltration experiments were performed with bacterial suspensions at 10⁵ CFU ml⁻¹ in 10 mM MgCl₂. At least four plants of each genotype were used per experiment, and the experiments were repeated at least three times. Bacterial numbers in mutant or transgenic lines were compared with Col-0 using a two-tailed Student's t-test. Hyaloperonospora arabidopsidis isolate Emwa1 (kindly supplied by J. Parker, MPI) was inoculated onto 2-wk-old seedlings at 5×10^4 spores ml⁻¹ as previously described (Berriri *et al.*, 2012). To assess plant cell death and *H. arabidopsidis* infection structures, trypan blue staining of the leaves was performed 6-d postinoculation (dpi). Experiments were repeated three times with similar results. Salicylic acid contents were determined after bacterial spray inoculation as previously described (Berriri et al., 2012). For Pseudomonas syringae pv maculicola strain ES4326 (Psm) carrying the Photorhabdus luminescens luxCDABE (lux) operon (Psm lux) (Zeier et al., 2004; Fan et al., 2008), bacteria were grown in King's B medium supplemented with the appropriate antibiotics at 28°C under constant shaking. Overnight log-phase cultures were washed four times with 10 mM MgCl₂ solution and diluted to different final OD_{600} levels for leaf inoculation. The diluted bacterial solutions were pressure-infiltrated from the abaxial side of the leaves using a needleless syringe, covering the whole surface of the respective leaf. To assess bacterial growth in naïve 5-wk-old plants (basal resistance), three fully grown leaves per plant (Berriri et al., 2012) were infiltrated with a suspension of Psm lux at $OD_{600} = 0.001$ between 10:00 h and 11:00 h. At 60 h, the bacterial growth in the infiltrated leaves was determined by measuring the bacterial bioluminescence in leaf disks as described (Hartmann et al., 2018).

Histochemical staining

For studying *OXI1* expression, an *OXI1*::GUS line was used (Forzani *et al.*, 2011). Plant tissues were fixed in 90% acetone for 30 min at 4°C, washed three times with 50 mM sodium phosphate buffer (pH 7.0), and subsequently stained for up to 16 h in 50 mM sodium phosphate buffer (pH 7.0), 2 mM K3Fe (CN6), 2 mM K4Fe(CN6) containing 1 mM 5-bromo-4-chloro-3-indolyl-D-glucuronide (Duchefa, Haarlem, the Netherlands). Tissues were cleared in ethanol and visualized with a stereo microscope (MZ16FA; Leica).

To determine H_2O_2 accumulation, 5-wk-old plants grown on jiffy pellets in plant growth chambers (Percival Scientific, Perry, IA, USA) under 16 h : 8 h, light : dark photoperiod at 22°C, 60% humidity, were used. The leaves were gently vacuum-infiltrated with 1 mg ml $^{-1}$ DAB in water (D8001; Sigma-Aldrich), and incubated in the dark for 4–5 h at 22°C on a shaker at 80–100 rpm. Following incubation, the samples were destained in bleaching solution (ethanol : acetic acid : glycerol,

3:1:1). Destained leaves were mounted on microscopic glass slides in 30% glycerol and pictured using Canon EOS 6D.

For cell death staining, detached leaves were treated with trypan blue (1 mg ml⁻¹) according to Beaugelin *et al.* (2019). The trypan blue solution was prepared by dissolving 40 mg trypan blue in 10 ml lactic acid (85%, w/w), 10 ml phenol (buffer equilibrated, pH 7.5–8.0), 10 ml glycerol, and 10 ml distilled water. Leaves were incubated in the trypan blue solution for 40 min, then immediately rinsed with ethanol and washed overnight in ethanol.

GC-MS metabolomics

The metabolomics study was performed on samples from two independent experiments, each including four biological replicates (Methods S1; Table S1). Semiautomated integration of chromatograms was performed with QUANLYNX software (Waters, Milford, MA, USA) after conversion of the raw data into NetCDF format. A model ion was chosen manually to perform a correct integration for 42 peaks, with the following criteria: Gaussian shape, high relative abundance in the mass spectrum, and the absence of saturation in any sample (Table S1). Metabolites were annotated with confidence levels according to the Chemical Analysis Working Group (Sumner et al., 2007), on the basis of both spectral and retention index matches with public database (Kopka et al., 2005). Peak surfaces were then normalized according to the internal standard peak surface (ribitol) and sample fresh weight (FW) (Table S2). Raw metabolomics dataset (accession number MSV000090447) can be downloaded from the publicly available MassIVE repository at the UCSD Center for Computational Mass Spectrometry website: https://massive. ucsd.edu/.

Quantification of defense metabolites

The levels of camalexin, SA, SAG, SGE, Pip, and NHP were determined by gas chromatography/mass spectrometry (GC-MS)-based analyses of plant extracts as detailed in Hartmann et al. (2018) and Stahl et al. (2019). Briefly, three leaves per plant were inoculated with a suspension of Psm at $OD_{600} = 0.005$, and control plants were mock-infiltrated with a 10 mM MgCl₂ solution. Leaves from one plant were harvested at 10-, 20-, and 48-h postinfection (hpi) after treatment, pooled for one biological replicate, and frozen in liquid nitrogen. Fifty milligrams of pulverized, frozen tissue samples was extracted twice with 1 ml of MeOH: H₂O (80: 20, v/v). For internal standardization, 1 µg of D₄-SA, D₉-Pip, D₉-NHP, indole-3-propionic acid (IPA), and salicin were added. Six hundred micoliters of the extract was evaporated to dryness, and free hydroxyl groups of the analytes were converted into their trimethylsilyl derivatives by adding 20 µl of pyridine, 20 µl of N-methyl-Ntrimethylsilyltrifluoroacetamide (MSTFA) containing trimethylchlorosilane (v/v) and 60 µl of hexane and heating the mixture to 70°C for 30 min. The cooled samples were transferred to a GC vial and diluted fivefold with hexane. Two microliters of the sample mixture was separated on a gas chromatograph (GC 7890A; Agilent Technologies, Santa Clara, CA, USA) equipped with a Phenomenex ZB-35 (30 m \times 0.25 mm \times 0.25 μ m)

capillary column using the following temperature program: 70° C for 2 min, with 10° C min⁻¹ to 320° C, and 320° C for 5 min. Mass spectra were recorded with a 5975C mass spectrometric detector (Agilent Technologies) between m/z 50 and m/z 750 in the electron ionization (EI) mode. Metabolites were quantified by integrating peaks of selected ion chromatograms using MSD CHEMSTATION software v.E.02.01.1177 (Agilent Technologies): SA (m/z 267) relative to D₄-SA (m/z 271), Pip (m/z 156) relative to D₉-Pip (m/z 165), NHP (m/z 172) relative to D₉-NHP (m/z 181), camalexin (m/z 272) relative to IPA (m/z 202), and SAG (m/z 267) and SGE (m/z 193), both relative to salicin (m/z 268).

RNA extraction and qRT-PCR analysis

Total RNA was extracted from 4-wk-old plants grown on jiffy pellets in plant growth chambers (Percival Scientific) under 16 h: 8 h, light: dark photoperiod at 22°C, 60% humidity, using NUCLEOSPIN Plant RNA (Macherey Nagel, Düren, Germany) kit following the manufacturer's protocol. First-strand cDNA was synthesized from 1.5 µg of total RNA using Super-Script III First-Strand Synthesis SUPERMIX kit (Invitrogen). The diluted cDNA (10 times) was used to perform quantitative RT-PCR (qRT-PCR) using SsoAdvanced Universal SYBR Green Supermix (Bio-Rad) on StepOnePlus Real-Time PCR System (Applied Biosystems, Waltham, MA, USA). The data were analyzed using Bio-Rad Cfx Manager software. ACTIN was used as a housekeeping gene for normalization of gene expression levels. Analyses were performed in triplicate and was repeated thrice with independent RNA samples. Primers used in this study are listed in Table \$3.

Transcriptome analysis

Four-week-old plants were collected for RNA analysis. Microarray analysis was carried out using the CATMAV.6.2 array based on Roche-Nimble Gene technology as described (Frei dit Frey et al., 2014). The arrays contained 219 684 primers representing all the Arabidopsis thaliana genes: 37 309 probes corresponding to TAIRV.8 annotation (including 476 probes of mitochondrial and chloroplast genes); 1796 probes corresponding to EUGENE software predictions and 5328 probes corresponding to repeat elements; 1322 probes for miRNA/MIR, 329 probes for other RNAs (rRNA, tRNA, snRNA, and soRNA); and several controls. Three independent biological replicates of the microarray analysis were performed (Table S4). For each biological repetition, 5-wk-old plants grown under 16 h: 8 h, light: dark photoperiod at 22°C, 60% humidity, were collected, and RNA samples were obtained by pooling more than 50 plants. Total RNA was extracted using Qiagen RNAeasy according to the supplier's instructions. For each comparison, one technical replicate with fluorochrome reversal was performed for each biological replicate (six hybridizations per comparison, RIN score cutoff = 7). The labeling of cRNAs with Cy3-dUTP or Cy5dUTP (Perkin-Elmer-NEN Life Science Products, Waltham, MA, USA) and the hybridization to the slides were performed as previously described (Lurin et al., 2004). Two-micron scanning was performed with InnoScan900 scanner (InnopsysR, Carbonne,

France), and raw data were extracted using MAPIXR software (InnopsysR).

Transcriptome analysis was performed as described (Frei dit Frey et al., 2014). Briefly, for each array, the raw data comprised the logarithm of median feature pixel intensity at wavelengths 635 nm (red) and 532 nm (green). For each array, a global intensity-dependent normalization, using LOESS (Locally reweighted Scatter Plot Smoothing), was performed to correct the dye bias. The differential analysis was based on the log ratios averaging over the duplicate probes and over the technical replicates. Hence, the numbers of available data for each gene equal the number of biological replicates and are used to calculate the moderated t-test. Under the null hypothesis, no evidence that the specific variances vary between probes is highlighted by limma (Ritchie et al., 2015), and consequently the moderated t-statistic is assumed to follow a standard normal distribution. To control the false discovery rate, we calculated adjusted P values using the optimized FDR approach (Storey & Tibshirani, 2003). We considered as being differentially expressed the probes with an adjusted P value \leq 0.05. Analysis was made with the R software. The function SqueezeVar of the library LIMMA was used to smooth the specific variances by computing empirical Bayes posterior means. The library KERFDR was used to calculate the adjusted P-values. The analysis to find overrepresented categories in the gene sets was obtained with AMIGO (Carbon et al., 2008), which is based on a hypergeometric test.

Results

OXI1 overexpression results in a gain-of-function phenotype induces ROS and leaf cell death in late development

Most of the functional analysis of OXII was made so far in the Arabidopsis Wassilewskija (Ws) background, hindering a genetic analysis with a large set of available mutants such as in the Columbia accession. Moreover, constitutive OXI1 overexpression function was studied using the strong constitutive 35S cauliflower mosaic virus promoter (Petersen et al., 2009). For these reasons, we generated a set of pOXI1::HA-OXI1 transgenic lines in the Col-0 oxi1-2 background (Beaugelin et al., 2019). Some OXII lines were slightly smaller than WT and started to show a gainof-function (GOF) leaf cell death phenotype after 4 wk of development (Figs 1a, S1). Western blot analysis of these lines showed elevated expression levels (OE1 and OE2), whereas other lines (C1 and C2) resembled WT plants and showed low expression levels (Fig. 1b). C1 and C2 were therefore considered as complementation lines. The cell death phenotype in the OXI1-OE lines consistently started with the darkening of the center of the rosettes, which then spread to the petioles and the leaves. The pattern of cell death development parallels the expression pattern of OXI1 as evidenced in pOXI1::GUS analysis, which showed strong expression along the vascular system (Fig. 1c) and was also reflected by the localized accumulation of ROS (Fig. 1d) and cell death (Fig. 1e) as evidenced by diamine benzidine (DAB) and trypan blue staining, respectively.

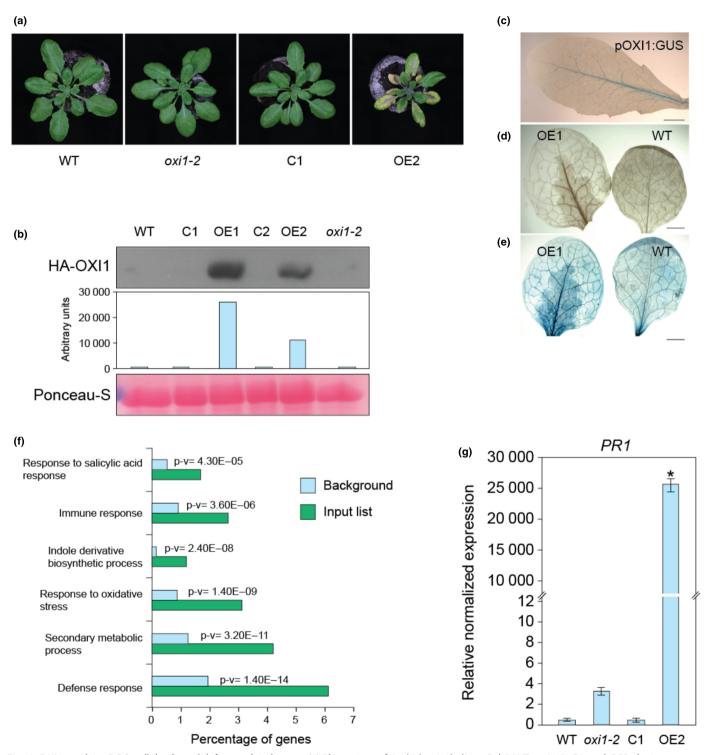


Fig. 1 OX/17 regulates ROS, cell death, and defense-related genes. (a) Phenotypes of Arabidopsis thaliana Col-0 WT, oxi1-2, C1, and OE2 plants grown for 4 wk on soil. (b) Western blot of 4-wk-old Col-0 WT, C1, OE1, C2, OE2, and oxi1-2 plants with HA antibody. The bottom panel represents small ribulose bisphosphate carboxylase (Rbcs) staining by Ponceau S. (c) GUS staining of 4-wk-old leaf of pOX/11::GUS plant. Bar, 1 mm. (d) Diaminobenzidine staining for H_2O_2 in leaves of 4-wk-old OE1 and Col-0 WT plants. Bar, 1 mm. (e) Trypan blue staining of leaves of 4-wk-old OE1 and Col-0 WT plants. Bar, 1 mm. (f) GO analysis of significantly upregulated genes ($log_2(FC)$ (P < 0.05)) of 4-wk-old OE2 when compared with Col-0 WT plants. Gene ontology term analysis was performed using the AGRIGO database (for a complete list of differentially regulated genes, see Table S4). (g) qRT-PCR of PR1 gene expression levels of Col-0 WT, oxi1-2, C1, and OE2 in three independent experiments after normalization to actin (n = 6). Values represent the mean \pm SE, * indicates significant difference from WT, $P \le 0.05$, as determined by Student's t-test.

Transcriptome analysis of OXI1 lines

To obtain insight into the underlying molecular processes giving rise to the cell death GOF phenotype in the OXII overexpression lines, we performed a global transcriptome analysis of three OXII-OE lines. Compared with Col-0, these lines showed 1055 upregulated genes (fold change ≥ 2) and 1004 downregulated genes (fold change ≤ 2) (Table S4). A gene ontology (GO) analysis of the upregulated genes revealed a significant overrepresentation of genes involved in metabolic and immune-related processes (Fig. 1f). For metabolic processes, mostly secondary metabolic pathways, such as amino acid- and indole-derivative biosynthetic processes, were induced (Fig. 1f; Table S4). Moreover, ion homeostasis, defense, SA, and JA pathways were significantly upregulated for the immune-related processes (Fig. 1f; Table S4), and PR1 gene expression levels were strongly enhanced upon overexpression of OXII (Fig. 1g).

Proteome analysis of OXI1 lines

In order to understand the changes induced by OXII at the protein level, we carried out whole proteome analysis of WT, oxi1-2, C1, and OE2 lines by LC-MS/MS of 4-wk-old plants. Proteomes of all the lines from four independent biological replicates were analyzed (Table \$5). Proteins were considered to be differentially expressed with a \log_2 ratio of > ± 1 and FDR < 0.05. Eighty-eight proteins were upregulated and 12 downregulated in WT compared with OE2 as visualized in the volcano plot (Fig. 2a). Enrichment analysis was made using AGRIGO followed by removal of redundant GO terms using REVIGO. The comparison of OE2 with WT showed 88 upregulated proteins involved in defense response, innate immune response, and SAR (Fig. 2a). The 12 downregulated proteins showed no enrichment in GO terms. These data are consistent with the role of OXI1 as a positive regulator of defense and cell death. Next, we compared oxi1-2 with OE2 (Fig. 2b). One hundred and forty proteins were enriched in the GO terms defense response, programmed cell death, and SAR (Fig. 2b). The 37 upregulated proteins in oxi1-2 did not show any significant GO terms. Finally, on comparing oxi1-2 with C1, the 23 downregulated proteins in oxi1-2 did not show any significant GO terms, but the 122 upregulated proteins in oxi1-2 belonged to the GO terms photosynthesis, photosynthesis light reaction, and oxidation-reduction process (Fig. 2c).

To test whether *OXI1* regulates the photosynthesis proteome at the level of transcription, we performed qRT-PCR on a number of photosynthesis genes. As shown in Fig. 2d, in comparison with WT and the *C1* complementation line, these photosynthesis-related genes show enhanced expression in *oxi1-2* but all of them were strongly repressed in *OE2*. In summary, these results are consistent with a negative role of *OXI1* in photosynthesis and metabolism.

OXI1 induces salicylic acid biosynthesis and SA responses

The transcriptome analysis suggested that *OXI1* regulates SA biosynthesis. Isochorismate synthase ICS1 also known as SID2 (SA-

INDUCTION-DEFICIENT2) is the key enzyme for SA biosynthesis (Wildermuth et al., 2001), transcript levels of which strongly accumulated in OE2 (Fig. 3a,b). SA-INDUCTION-DEFICIENT2 expression is redundantly regulated by the transcription factors Cbp60g and SARD1 (Zhang et al., 2010), which were both found to be upregulated in OXII overexpression lines (Fig. 3a, b). To quantify the levels of free SA, SA-β-glucoside (SAG), and SA glucose ester (SGE) upon pathogen challenge, we infected 4-wk-old Arabidopsis Col-0 WT, oxi1-2, C1, and OE2 plants with Psm. In contrast to oxi1-2, which showed enhanced susceptibility, OE2 showed enhanced resistance toward *Psm* when compared with WT or C1 plants (Fig. 4a). Constitutive accumulation of all of SA, SAG, and SGE were found in OE2 already in the absence of pathogen challenge (Figs 3c, S2). In addition, the levels of free and total SA (i.e. the sum of SA, SAG, and SGE) increased to similar levels in WT, oxi1-2, and the C1 complementation line in response to Psm inoculation, at 10, 20, and 48 hpi. Interestingly, OE2 was also primed for enhanced SA production in the earlier phases of the Psm interaction, which manifested itself in strongly enhanced levels of SAG, SGE, and total SA at 10 and 20 hpi (Figs 3c, S2). While WT, oxi1-2, and C1 showed a systemic increase in SA, SAG, and SGE at 48-h post-Psm inoculation, the high constitutive levels of the three SA forms were not further increased in the systemic leaves of pathogen-challenged OE2 (Figs 3c, S2). Overall, these results indicate that OXII overexpression already poised plants for maximal SA synthesis before pathogen attack.

Salicylic acid signaling is mediated by the transcriptional regulator NPR1 which can shuttle from the cytoplasm to the nucleus in a redox-dependent manner and is dependent on the thioredoxins TRX-h3 and h5 (Tada et al., 2008). Nuclear NPR1 then associates with TGA transcription factors to induce SA-dependent genes (Mou et al., 2003; Zhang et al., 2003). Interestingly, OXI1-OE lines show strongly enhanced levels of TRX-h3 and TRX-h5 transcripts, suggesting that SA signaling might also be enhanced in these plants (Fig. 3a,b). A major part of OXI1-regulated genes are SA-responsive genes. The list (Table S4) includes multiple WRKY transcription factors (WRKY18, 40, and 60), PR genes (PR1, PR2, PR3, PR4, and PR5), and the metabolic enzymes PBS3, BAP1, and alpha-dioxygenase DOX1, which is involved in the protection against oxidative stress and cell death (De León et al., 2002).

OXI1 enhances WRKY33 transcript levels and camalexin biosynthesis

The transcriptome analysis indicated that the transcription factor WRKY33 and its direct downstream target PAD3 are strongly upregulated in OXI1 overexpression plants (Fig. 4b). Since CYP79B2 and PAD3 encode key enzymes for camalexin synthesis (Fig. 4c), we quantified camalexin levels in WT, oxi1-2, C1, and OE2 (Fig. 4d). Camalexin levels were strongly induced in all plant lines at 20 hpi with Psm. Moreover, in mock-treated OE2 plants, camalexin levels were elevated in the absence of pathogen inoculation. In addition, the pathogen-induced accumulation of camalexin was more pronounced in OE2 than in the other lines under investigation at 20 hpi (Fig. 4d). Interestingly, we also

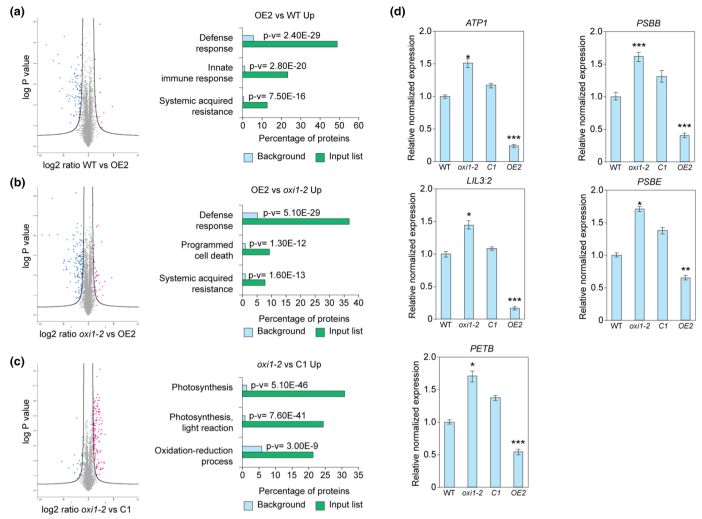


Fig. 2 Global proteome profile shows a role of OX/1 in photosynthesis and defense responses. (a) Volcano plot depicting the differentially expressed proteins in Arabidopsis thaliana Col-0 WT vs OE2. Gene ontology enrichment of the downregulated proteins. (b) Volcano plot depicting the differentially expressed proteins in oxi1-2 vs OE2. Gene ontology enrichment of the downregulated proteins. (c) Volcano plot depicting the differentially expressed proteins in oxi1-2 vs C1. GO enrichment of the upregulated proteins. (a–c) The fold enrichment for the GO terms was calculated based on the frequency of proteins in the deregulated group compared with their frequency in the total proteome (log₂ FC) (P < 0.05) using the AgriGo database (for complete protein lists see Table S5). (d) Gene expression of photosynthesis-related genes ATP1, LIL1, PETB, PSBB, and PSBE in 4-wk-old Col-0 WT, oxi1-2, C1, and OE2 lines. qRT-PCR mean values in relation to WT of three independent experiments after normalization to actin \pm SE (n = 6), * indicates significant difference from WT, $P \le 0.05$, **, $P \le 0.01$ as determined by Student's t-test.

observed strongly increased *CYP82C2* transcript levels in *OE2* plants (Fig. 4b). In contrast to *PAD3*, *CYP82C2* converts IAOx into 4-OH-ICN (Fig. 4c), another indolic compound with a key function in pathogen resistance (Rajniak *et al.*, 2015). In summary, these data show that *OXI1* activates the defense-related indolic pathway in Arabidopsis and strongly promotes the accumulation of the phytoalexin camalexin.

OXI1 triggers reactive oxygen species biosynthesis and homeostasis

Free radicals can be produced in Arabidopsis by several pathways, including from nitrous acid, which can be generated by at least two pathways upon pathogen attack. NO can be synthesized via

the nitrate reductases NIA1 and NIA2 or via an unknown mechanism by the GTPase NO-associated protein 1 (AtNOA1) (Lozano-Juste & León, 2010). In OXII overexpressor lines, we found enhanced expression of NIA2 (Table S4), suggesting that the lines might produce enhanced levels of NO. Reactive oxygen species and NO can both act as signaling molecules of free radicals and are both involved in pathogen defense. Upon pathogen infection of leaves, plants induce superoxide production via several means, including the NADPH oxidases RbohD and RbohF (Torres Dangl, 2005). OXIDATIVE SIGNAL-INDUCIBLE I GOF lines showed enhanced transcript levels of RbohD and RbohF (Fig. 5a,b) and accumulation of H₂O₂ (Fig. 5c). Moreover, a significant set of genes with roles in ROS homeostasis are identified in OXII-OE lines, including ascorbate

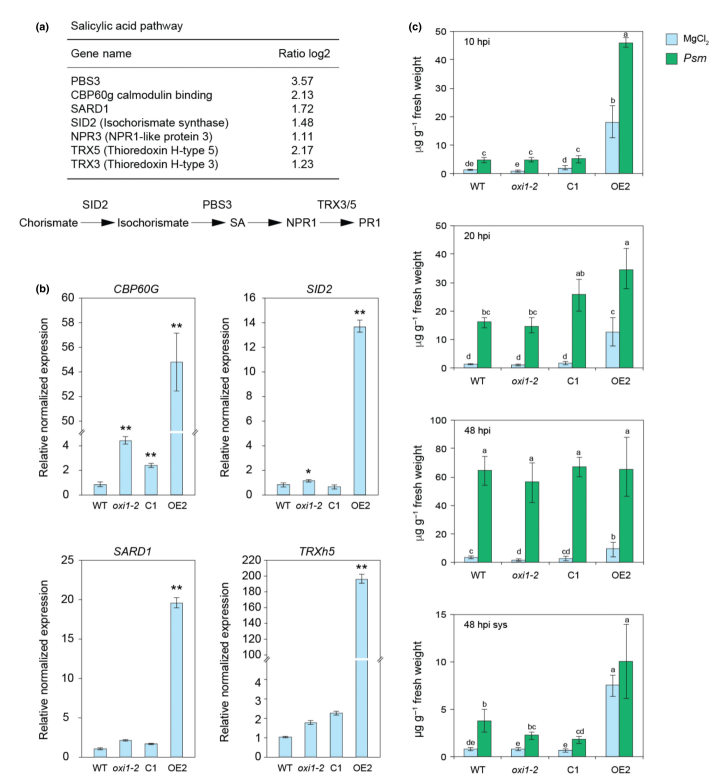
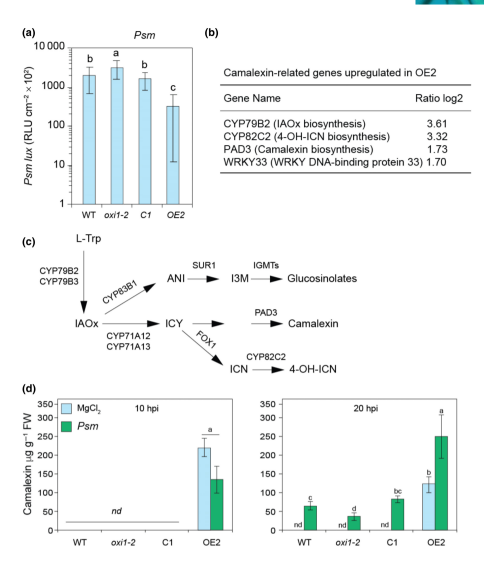


Fig. 3 OXI1 regulates SA biosynthesis, signaling, and basal immunity. (a) Selected upregulated genes related to SA biosynthesis in 4-wk-old OE2 compared with Arabidopsis thaliana Col-0 WT plants. (b) Cbp60g, SID2, SARD1, and TRXh5 gene expression levels in 4-wk-old Col-0 WT, oxi1-2, C1 complementation, and OE2 overexpression lines. qRT-PCR mean values in relation to Col-0 WT of three independent experiments after normalization to actin \pm SE (n=6), *, $P \le 0.05$; **, $P \le 0.05$; indicate significant differences from WT, as determined by Student's t-test. (c) Total levels of SA (sum of free SA, SAG, and SGE) in Psm-inoculated or mock-treated Arabidopsis leaves at 10, 20, and 48 hpi (hours postinfection), and in distal, systemic (sys) leaves at 48 hpi. Values \pm SE are given in $\mu g g^{-1}$ fresh weight (n=4) of three independent experiments. Different letters above the bars denote statistically significant differences (P < 0.05), analysis of variance and Post hoc Tukey's HSD test).

Fig. 4 OXI1 controls basal immunity and camalexin biosynthesis. (a) Growth of Psm lux in 4-wk-old Arabidopsis thaliana Col-0 WT, oxi1-2, C1, and OE2 plants. Bacterial numbers were assessed at 60 hpi (hours post infection) with the bioluminescent Psm lux strain and expressed as relative light units (rlu) per cm² leaf area. Data represent the mean \pm SD of at least 20 leaf replicates (n = 6-7). Different letters denote significant differences (P < 0.005, analysis of variance and post hoc Tukey's HSD test). (b) Selected upregulated genes in 4-wk-old OE2 plants related to (c) glucosinolate, camalexin, and 4-OH-ICN biosynthesis. Arrows denote enzymatic steps. Tryptophan (L-Trp); IAOx, indole-3-acetaldoxime; 4OH-ICN; ICY, indole-3-cyanohydrin; ANI, aci-nitro indole; 4-OH-ICN, 4-hvdroxvindole-3carbonylnitrile; I3M, indol-3-ylmethyl glucosinolate; ICN, indole-3-carbonylnitrile. (d) Camalexin levels in 4-wk-old Col-0 WT, oxi1-2, C1, and OE2 lines upon infection with *Psm* at 10 and 20 hpi. Values \pm SE are given in $\mu g g^{-1}$ fresh weight (n = 4) of three independent experiments. Different letters above the bars denote statistically significant differences (P < 0.05, analysis of variance and post hoc Tukey's HSD test). nd, nondetectable.



oxidase 1, the Cu/Zn superoxide dismutase *CSD1* (Fig. 5a), the oxidoreductase *TXR5*, or a number of peroxidases such as *PER32* or *PER50* (Table S4).

OXI1 induces polyamine biosynthesis gene expression

Polyamines play a role in plant growth, development, and in abiotic and biotic stress responses, including pathogen defense (Takahashi & Kakehi, 2010). In plants, putrescine is derived from arginine through several biosynthetic steps involving arginine decarboxylase (ADC), agmatine iminohydrolase (AIH), and N-carbamoylputrescine amidohydrolase (CPA) (Takahashi & Kakehi, 2010). In *OXII* overexpressor plants, ADC gene expression levels were upregulated (Fig. 5d) and arginine and putrescine also accumulated (Figs 6, S3). Spermidine is synthesized from putrescine by spermidine synthase (SPDS) which requires decarboxylated S-adenosyl methionine (dc-SAM) as aminopropyl donor. Decarboxylated S-adenosyl methionine itself is derived from methionine by a two-step synthesis via S-adenosyl methionine synthase (SAMS) and S-adenosyl methionine decarboxylase

(SAMDC). Interestingly, both *OXI1* GOF lines did not show accumulation of spermidine in the metabolite profile, and SPDS, SAMS, and SAMDC transcript levels were downregulated (Table S4). Polyamines can also contribute to apoplastic H₂O₂ production by polyamine oxidases (PAO) (Moschou *et al.*, 2012). Therefore, it is also possible that some ROS in the *OXI1* GOF lines might be derived from polyamine oxidation reactions, although no upregulation of *PAO* gene expression levels was observed in any of the two lines.

OXI1 promotes basal resistance against virulent biotrophic pathogens

To verify the function of *OXII* in basal resistance in the Col-0 accession, we spray-infected 4-wk-old Col-0 WT, *oxi1-2*, *CI*, and *OE2* with *Pst* DC3000 and determined bacterial proliferation levels at 3 dpi. In contrast to the *oxi1-2* mutant plants, which showed enhanced susceptibility, *OE2* showed enhanced resistance toward *Pst* DC3000 (Fig. 5e). In summary, the observed phenotypes correlate well with the function of *OXII*

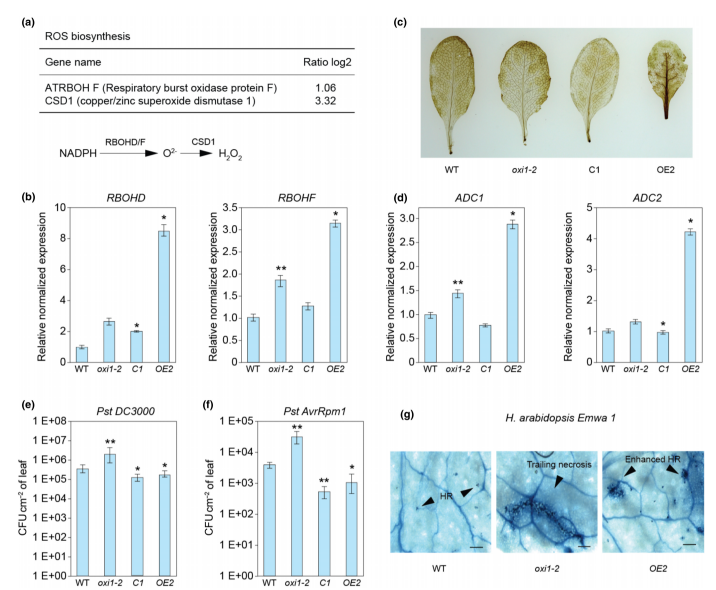


Fig. 5 *OXI1* triggers ROS production and enhances basal and effector-triggered immunity. (a) Selected upregulated genes related to reactive oxygen species biosynthesis in *OE2* compared to *Arabidopsis thaliana* Col-0 wild-type (WT) plants. (b) *RBOHD* and *RBOHF* gene expression levels in 4-wk-old Col-0 WT, *oxi1-2*, *C1*, and *OE2* lines. qRT-PCR mean values in relation to WT of three independent experiments after normalization to actin \pm SE (n = 6), *, significant difference from WT, $P \le 0.05$, **, $P \le 0.01$ as determined by Student's *t*-test. (c) Diaminobenzidine staining of H₂O₂ in 4-wk-old Col-0 WT, *oxi1-2*, *OE2*, and *C1* lines. (d) *ADC1* and *ADC2* expression in Col-0 WT, *oxi1-2*, *C1*, and *OE2* lines. qRT-PCR mean values in relation to Col-0 WT of three independent experiments after normalization to actin \pm SE (n = 6), *, significant difference from WT, $P \le 0.05$, as determined by Student's *t*-test. (e) Growth of *Pst* DC3000 and (f) *Pst* DC3000 A*vrRpm1* in 4-wk-old Col-0 WT, *oxi1-2*, *C1*, and *OE2* plants. Bacterial titers of *Pst* and *Pst* A*vrRpm1* were determined at 3 dpi by spray inoculation of plants. Bars represent the average of at least four independent replicate experiments, and error bars show SD (n = 6). *, P < 0.05 and **, P < 0.01 by Student's *t*-test. (g) Growth of *H. arabidopsidis* Emwa1 in 4-wk-old Col-0 WT, *oxi1-2*, and *OE2* plants. Cell death structures were stained in leaves with lactophenol trypan blue 6 d after *H. arabidopsidis* Emwa1 inoculation. Bar, 100 μm.

as positive regulator of defense genes and antimicrobial substances.

OXI1 positively affects effector-triggered immunity

We also investigated the role of *OXI1* in ETI responses in the Col-0 background. We spray-infected plants with the avirulent *Pst AvrRpm1* strain that induces ETI due to the presence of the CC-NB-LRR resistance gene *RPM1*. As shown in Fig. 5f, *oxi1-2* was compromised in resistance against the avirulent *Pst AvrRpm1*

strain that is recognized by the CC-NB-LRR R protein RPM1. By contrast, CI and OE2 showed enhanced resistance against Pst AvrRpm1 (Fig. 5f). Surprisingly, CI showed enhanced resistance against Pst DC3000 and Pst AvrRpm1, suggesting that even small enhancement of OXII expression might be sufficient to trigger a primed defense state.

To test the role of *OXI1* susceptibility to the oomycete *Hyaloperonosopora parasitica*, in Col-0 plants, 2-wk-old WT, *oxi1-2*, and *OE2* were sprayed with *H. arabidopsis Emwa1* (Hpa Emwa1) (Fig. 5g). While a trailing necrosis phenotype in *oxi1-2*

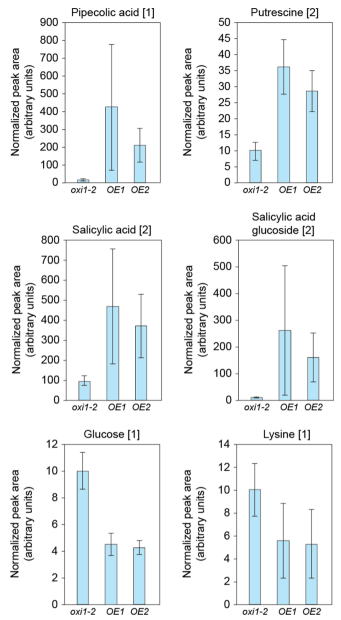


Fig. 6 *OXI1* regulation of metabolism. Global metabolite analysis was performed on 4-wk-old *Arabidopsis thaliana* Col-0 WT, *oxi1-2*, *OE1*, and *OE2* lines. Bar plots of pipecolic acid, putrescine, salicylic acid, salicylic acid glucoside, glucose and lysine (mean \pm SD) representing normalized GC–MS peak areas in *oxi1-2*, *OE1*, and *OE2* compared with Col-0 WT (Mann–Whitney *U*-test, P < 0.05).

plants indicated enhanced susceptibility, HR at the infection sites of Col-0 plants correlated with their resistant phenotype, whereas *OE2* showed enhanced cell death spreading around infection sites. Overall, our results show that *OXI1* plays a role in ETI mediated by CC-NB-LRR resistance genes.

Metabolome analysis of *OXI1* reveals its function in regulating secondary metabolism

To further investigate the function of *OXI1*, we performed a global metabolomics analysis of *OE1*, *OE2*, and *oxi1-2* plants. In

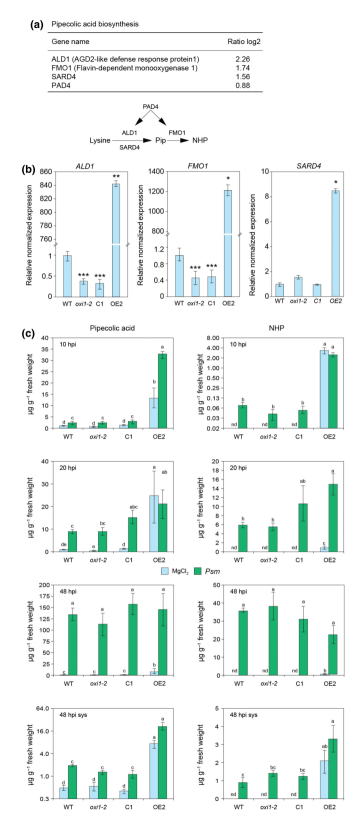
comparison with the other lines including Col-0 WT plants, oxi1-2 mutant plants showed only small differences in a number of metabolites. By contrast, both OXII-OE lines showed very strong signatures, with several metabolites that accumulated compared with WT or decreased in oxi1-2 mutant plants. Free SA or its glucoside were among the 10 most differentially accumulated metabolites in the OXI1-OE lines (Figs 6, S3). Interestingly, the most differently accumulated metabolite was pipecolic acid (Figs 6, S3), whose accumulation is required for local and systemic defense responses in Arabidopsis (Návarová et al., 2012). We also found major differences in tyrosine, tryptophan, and the branched chain amino acids leucine, isoleucine, and valine, all of which are marker metabolites during a localized pathogen infection (Zeier, 2013). On the contrary, OXII-OE lines showed lower levels of glucose and fructose and the amino acids lysine, proline, glycine, and glutamine when compared to WT plants (Figs 6, S3).

OXI1 regulates pipecolic acid and N-hydroxypipecolic acid biosynthesis

Accumulation of Pip is necessary for local and systemic defense responses in Arabidopsis. Pipecolic acid is derived from the amino acid lysine, whose levels were depleted in OXI1 overexpression lines. The transaminase ALD1 converts L-Lys to 2,3dehydropipecolic acid which is then reduced to Pip by SARD4 (Ding et al., 2016; Hartmann et al., 2017). Finally, Pip can be hydroxylated to NHP by FMO1 (Chen et al., 2018; Hartmann et al., 2018). N-hydroxypipecolic acid is the actual immuneactive and SAR-inducing signal of this pathway (Hartmann & Zeier, 2019; Zeier, 2021). Upon pathogen attack, ALDI, SARD4, and FMO1 are strongly upregulated and mutants in any of these genes show enhanced susceptibility to biotrophic pathogens (Song et al., 2004; Mishina & Zeier, 2006; Hartmann et al., 2017). In accordance with the transcriptome and metabolite data, ALD1, SARD4, and FMO1 transcripts (Fig. 7a,b), and Pip and NHP metabolite levels accumulated already in mocktreated OE2 plants without pathogen attack (Fig. 7c). Upon inoculation with Psm, Pip and NHP strongly accumulated in WT, oxi1-2, C1, and OE2 plants from 10 hpi onward (Fig. 7c). Besides their constitutively enhanced Pip and NHP levels, OE2 plants showed a primed biosynthesis of Pip and NHP, both at the sites of *Psm* infection (at 10 hpi for Pip, at 20 hpi for NHP) and in distal leaves (48 hpi) (Fig. 7c). These results indicate that OXI1 promotes the biosynthesis of Pip and NHP via the transcriptional upregulation of ALD1, SARD4, and FMO1.

To test whether the enhanced resistance phenotype of *OXI1*-overexpressing plants is mediated by NHP, we crossed the *OXI1-OE* line *OE2* with *ald1* or *fmo1*, respectively. Remarkably, the strong pathogen resistance phenotype of *OE2* was completely suppressed in homozygous *OE2 ald1* and *OE2 fmo1* lines (Fig. 8a), indicating that the NHP pathway is essential for *OXI1* function in immunity.

We next verified that the enhanced NHP levels in *OE2* plants were fully dependent on *ALD1* and *FMO1* (Fig. 8b). As expected, the constitutive accumulation of Pip was completely



suppressed in *OE2 ald1* plants (Fig. 8b). Moreover, *OE2 fmo1* lines showed strongly attenuated constitutive accumulation of Pip (Fig. 8b). This indicates that *OXI1*-promoted NHP biosynthesis positively regulates the biosynthesis of its precursor Pip,

Fig. 7 *OX11* induces pipecolic acid and NHP biosynthesis. (a) Selected upregulated genes related to pipecolic acid (Pip) and N-hydroxypipecolic acid (NHP) biosynthesis in *OE2* compared with *Arabidopsis thaliana* Col-0 wild-type (WT) plants. (b) *ALD1*, *FMO1*, and *SARD4* gene expression levels in 4-wk-old Col-0 WT, oxi1-2, C1, and OE2 lines. qRT-PCR mean values in relation to WT in three independent experiments after normalization to actin \pm SE (n = 6), *, P ≤ 0.05; **, P ≤ 0.01; ***, P ≤ 0.001 indicate significant differences from WT, as determined by Student's t-test. (c) Pipecolic acid levels (left) and NHP levels (right) in Psm-inoculated or mock-treated Arabidopsis leaves at 10, 20 and 48 hpi (hours postinfection), and in distal, systemic (sys) leaves at 48 hpi. Values \pm SE are given in μ g g⁻¹ fresh weight (n = 4) of three independent experiments. Different letters above the bars denote statistically significant differences (P < 0.05, analysis of variance and post hoc Tukey's HSD test). nd, nondetectable.

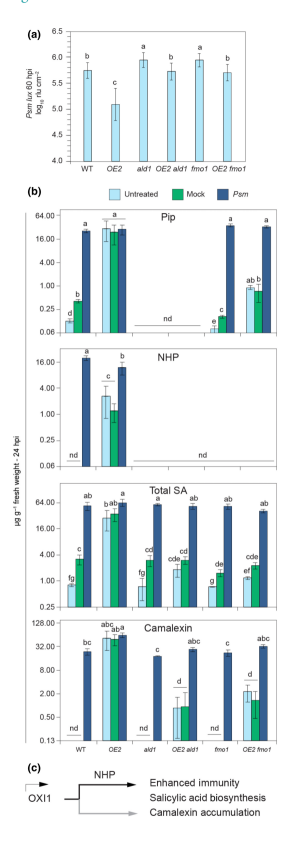
and thus confirms the existence of a positive feedback loop in the induction of the NHP biosynthetic pathway (Hartmann & Zeier, 2019). Moreover, these data confirm that Pip and NHP accumulation is promoted by *OXII*-dependent regulation of *ALD1* and *FMO1*.

Since *OXII-OE* lines show constitutive accumulation of SA, we also asked whether the SA accumulation promoted by *OXII* occurs independently of the NHP pathway. As seen in Fig. 8(b), total SA levels are partially suppressed in *OE2 ald1* and *OE2 fmo1* lines under nontreated and mock-treated conditions. We also determined the levels of free and conjugated forms of SA, but these results (Fig. S4) did not reveal a different regulation than those seen for total SA (Fig. 8b). These data suggest that *OXII-*induced activation of the NHP pathway directly controls SA accumulation. Upon pathogen challenge, however, an NHP-independent mechanism seems to drive SA accumulation, as all lines accumulated SA to similar levels under these conditions (Fig. 8b,c).

Finally, we also verified whether the constitutive camalexin accumulation in *OXI1* GOF lines is linked to the NHP pathway. Similar to the situation of SA, camalexin levels were partially suppressed in *OE2 ald1* and *OE2 fmo1* lines under nontreated and mock-treated conditions (Fig. 8b). These data suggest that *OXI1* regulation of the NHP pathway directly impinges on camalexin accumulation. Upon pathogen challenge, however, all lines accumulated camalexin to similar levels (Fig. 8b), suggesting that an NHP-independent mechanism drives camalexin accumulation under pathogen conditions.

Discussion

OXIDATIVE SIGNAL-INDUCIBLE1 protein kinase is activated by treatment of plants with cellulose or H₂O₂ and links the activation of MPK3/MPK6 to disease resistance (Rentel et al., 2004; Petersen et al., 2009) as well as to the beneficial interaction with the endosymbiotic fungus Piriformospora indica (Camehl et al., 2011). Interestingly, the OXII gene is also strongly induced at the transcriptional level by various abiotic and biotic conditions that are usually associated with the occurrence of accumulated ROS levels, such as high light or heavy metals (Smeets et al., 2013; Beaugelin et al., 2019). To gain more insight into the function of OXII in plants, we generated a



number of lines that express the genomic *OXI1* locus in the *oxi1-2* knockout background of Col-0. Aside from identifying weak expression lines that complement the *oxi1-2* phenotype, *OXI1* overexpression lines showed cell death phenotypes in plants after

Fig. 8 Role of the NHP pathway in OX/1-induced basal resistance to virulent Pseudomonas syringae. (a) Naïve Arabidopsis of the indicated lines were inoculated with bioluminescent P. svringae pv maculicola ES4326 (Psm) expressing the luxCDABE operon from Photorhabdus luminescens (Psm lux) by syringe infiltration of three leaves each with a bacterial suspension of OD_{600} (optical density at 600 nm) = 0.001. As a measure of plant susceptibility, the numbers of bacteria were determined at 60 hpi (hours postinfection) in inoculated leaves by luminescence quantification and expressed as relative light units (rlu) per cm² leaf area (Gruner et al., 2013). Bars indicate the mean \pm SD of at least 15 leaf replicates $(n \ge 21)$. Different letters denote significant differences (P < 0.05, analysis of variance and post hoc Tukey's HSD test). (b) Accumulation of defenserelated metabolites in different Arabidopsis after bacterial inoculation. Levels (in $\mu g g^{-1}$ fresh weight (FW)) of Pip, NHP, total SA (sum of free (unbound) SA, SA-β-glucoside (SAG) and SA glucose ester (SGE)), and camalexin in leaves inoculated with compatible P. syringae pv maculicola (Psm) at 24 hpi. The leaf metabolite levels of plants infiltrated with 10 mM MgCl₂ (mock) or of untreated plants (nt) harvested at the same time is also given. Bars represent means \pm SD of four biological replicates (n=4). One replicate sample consisted of four leaves from one plant. Different letters denote significant differences (P < 0.05, Kruskal–Wallis H-test). nd, not detected. (c) Model of OXI1 mediated immunity. OXI1 expression results in activation of the NHP pathway, thereby inducing SA and camalexin accumulation and pathogen resistance. A minor branch of OXI1enhanced levels of SA, camalexin and immunity, is regulated by an NHPindependent branch.

4-5 wk of growth. These results are in contrast to those of Petersen et al. (2009) who generated OXII-overexpressing lines under the 35S CaMV promoter in the Ws-2 background. Petersen et al. (2009) only reported one line and found no significant difference in the behavior of the oxi1-1 knockout line in the Ws-2 background. By contrast, we observed that high OXI1 levels are correlated with a GOF cell death phenotype in untreated plants and in basal and ETI. Although Petersen et al. (2009) detected OXI1 transcripts in the Ws-2 line, OXI1 protein was not measured. We previously also generated stable transgenic 35 S::HA-OXI1 cDNA lines and in several of these lines we observed gene expression by qRT-PCR, but none of the lines had detectable OXI1 protein levels, indicating gene silencing and that constitutive expression of OXII is not tolerated. We therefore generated HA-tagged pOXI1::HA-OXI1 lines expressing the entire genomic OXII locus. As shown in this work, coherent transcriptome, proteome, and metabolome analysis reveals a causal correlation of OXI1 protein levels with the GOF cell death and pathogen resistance phenotype and establishes OXI1 as a positive regulator of programmed cell death and immunity.

Our metabolite analysis also revealed that *OXI1* promotes the accumulation of Pip and NHP, which are lysine-derived metabolites that have been shown to be essential components of SAR (Hartmann & Zeier, 2019). Pipecolic acid accumulates upon pathogen infection in both infected (local) and uninfected (distal) leaves (Hartmann *et al.*, 2017, 2018) and is generated by the chloroplast-localized lysine aminotransferase ALD1 and the dehydropipecolate reductase SARD4 (Ding *et al.*, 2016; Hartmann *et al.*, 2017). The importance of Pip in immunity is demonstrated in *ald1* mutants, which are compromised in pathogen-induced defense in both local and distal leaves (Song *et al.*, 2004; Návarová *et al.*, 2012). Interestingly, *OXI1-OE* plants show highly induced transcript levels of the key Pip

biosynthesis enzymes ALD1 and SARD4, suggesting that OXI1 regulates Pip biosynthesis.

L-Pip is further hydroxylated to the SAR-activating metabolite NHP by FMO1. N-hydroxypipecolic acid biosynthesis by FMO1 is induced at the transcriptional level by pathogens (Hartmann et al., 2018), and our results suggested that OXII triggers the accumulation of Pip and NHP via transcriptional upregulation of ALD1 and FMO1. To test this hypothesis, we crossed OE2 with ald1 and/or fmo1 to generate OE2 ald1 and OE2 fmo1 and compared Pip and NHP levels in these lines with those of WT, ald1, and fmo1. The constitutive and inducible accumulation of NHP in OXI1-OE lines was completely suppressed in OE2 ald1 and OE2 fmo1 under nontreated and mock-treated conditions, while Pip accumulation was fully blocked in OE2 ald1 (Fig. 8b). This confirms that Pip and NHP accumulation under OXI1-overepressing conditions is mediated by ALD1 and FMO1, respectively. Interestingly, the OXI1-triggered accumulation of Pip is markedly attenuated in OE2 fmo1 (Fig. 8b), indicating that Pip accumulation is promoted by NHP. This supports our previous hypothesis on the existence of a positive feedback loop in the NHP biosynthetic pathway (Hartmann & Zeier, 2019; Yildiz et al., 2021; Zeier, 2021). Importantly, the enhanced pathogen resistance of OXII overexpression lines was completely suppressed in homozygous OE2 ald1 and OE2 fmo1 lines (Fig. 8a). These results show that NHP functions as a main mediator of the here-described function of OXI1 in plant immunity.

N-hydroxypipecolic acid induces the expression of a set of major plant immune genes to enhance defense readiness, amplifies resistance responses, acts synergistically with SA in the induction of SAR, promotes hypersensitive cell death, and primes plants for enhanced immune mobilization in cases of future pathogen challenge (Hartmann & Zeier, 2019). Pipecolic acid has also been shown to induce SA accumulation and the transcriptional induction of NHP biosynthetic genes in an FMO1dependent manner, suggesting that elevated NHP levels trigger SA biosynthesis and amplify its own production (Bernsdorff et al., 2016). In this context, hormone, metabolite, and chemical analysis indicated the strong accumulation of SA and its glucoseconjugated forms SAG and SGE upon OXII expression at a time when the plants developed a cell death phenotype. Importantly, we could correlate these events with the expression of several genes involved in SA biosynthesis, such as CBP60g and SARD1, two key transcriptional regulators of SID2 (Zhang et al., 2010), encoding the isochorismate synthase enzyme responsible for SA biosynthesis (Wildermuth et al., 2001). In agreement with upregulation of CBP60g and SARD1, SID2 was equally upregulated and plants showed enhanced SA levels already in the absence of pathogen challenge.

The expression of the isochorismate synthase enzyme SID2 is strongly upregulated in *OXI1* overexpression lines (Fig. 3a,b) and the plants accumulate high levels of salicylic acid (Figs 3c, S2) and show enhanced cell death during development. Programmed cell death in plants is dependent on the accumulation of SA. To test whether the cell death phenotype of *OXI*1 GOF lines is dependent on SA, we crossed *OE2* with *sid2* mutants, which are

compromised in SA biosynthesis. *OE2 sid2* plants partially suppressed the dwarf and cell death phenotype of the *OXI1-OE* line (Beaugelin *et al.*, 2019), indicating that *OXI1* regulates cell death by controlling the expression of the SA biosynthetic pathway.

Our data suggest that activation of OXI1 results in NHP and SA production. Our results show that the *OXI1*-driven activation of SA biosynthesis is dependent on NHP (Fig. 8). These data support the existence of an NHP- and SA-dependent selfamplification machinery operating during the establishment of plant SAR (Zeier, 2021). Elevated NHP levels are causative for a primed state in plants (Yildiz et al., 2021), and this is further confirmed by the potentiated NHP, SA, and camalexin accumulation in OXII-OE plants following Psm attack. OXIDATIVE SIGNAL-INDUCIBLE1 belongs to the most highly and fastest induced Arabidopsis genes and proteins that react to a variety of stress conditions including pathogen attack (Rentel et al., 2004). Our work here shows that OXII plays a central role in the production of NHP, SA, and camalexin together contributing to enhanced innate immunity (Fig. 8c). However, recent work suggests that NHP is not upstream of SA, but acts in parallel to mutually amplify each other (Sun et al., 2020). Interestingly, the analysis of OE2 ald1 and OE2 fmo1 lines suggests that OXI1 also regulates an NHP-independent smaller signaling branch to contribute to SA accumulation (Fig. 8c). To further clarify the complex fine regulation, it will be interesting to investigate whether OE2 sid2 plants are not affected in the upregulation of NHP

In addition to NHP and SA, our data show that OXI1 is also a key regulator of ROS accumulation and signaling in Arabidopsis. The interaction of ROS with SA in immunity has been well-documented (Herrera-Vásquez et al., 2015), but little knowledge exists vet on the link between ROS and NHP. One component that could be a common denominator linking ROS to SA is glutathione, as demonstrated for NPR1, which thereby acts as a cellular redox sensor (Mou et al., 2003). Our transcriptome and metabolome analysis also indicated that OXI1 activates the defense-related indolic pathway and strongly promotes camalexin accumulation. Glutathione might not only be linking ROS to SA but also to camalexin biosynthesis as γ-glutamyl cyclotransferase mutants of the defense-related indole pathway not only inhibit camalexin synthesis but also NHP and SA biosynthesis (Wang et al., 2022). However, a number of other reports (Návarová et al., 2012; Bernsdorff et al., 2016; Hartmann et al., 2018; Yildiz et al., 2021) demonstrate that enhanced NHP levels prime for enhanced camalexin biosynthesis are consistent with the metabolic phenotypes of the OXII-OE2, the OXI1-OE2 fmo1, and the OXI1-OE2 ald1 lines (Fig. 8b). Overall, our results indicate that OXI1 is at the heart of the regulation of a number of key factors of plant immunity and that further experiments are warranted to uncover the full complexity of OXII functioning in defense.

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Author contributions

CF generated and analyzed transgenic lines. MH performed pathogen assays. RL performed global metabolome analysis. MH, LA, SR and JZ analyzed selected metabolites. AAR and NR performed transcript analysis. AH, SCS, NR and HN performed the proteome analysis. JZ and HH designed the work and wrote the paper. AAR and MH contributed equally to this work.

Competing interests

None declared.

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Data availability

The proteome data that support the findings of this study are openly available in PRIDE under the following project number PXD023853. Microarray data were deposited at CATdb (http://urgv.evry.inra.fr/cgi-bin/projects/CATdb/consult_expce.pl?experiment_id=337) (Project AU2010-01_OXI1) and GEO (Project GSE35941) according to the 'Minimum Information About a Microarray Experiment' standards. Global proteome sample preparation, data acquisition, and analysis by LC–MS/MS (Methods S2).

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Supporting Information

Additional Supporting Information may be found online in the Supporting Information section at the end of the article.

- **Fig. S1** Characterization of OXIDATIVE SIGNAL-INDUCIBLE1 (*OXII*) lines.
- **Fig. S2** OXIDATIVE SIGNAL-INDUCIBLE1 (*OXII*) regulation of SA biosynthesis.
- **Fig. S3** OXIDATIVE SIGNAL-INDUCIBLE1 (*OXII*) regulation of metabolism.

Fig. S4 SA derivatives in different OXIDATIVE SIGNAL-INDUCIBLE1 (OXII) Arabidopsis lines after mock- or *Pseudomonas syringae* pv *maculicola* inoculation.

Methods S1 GC-MS metabolomics.

Methods S2 Global proteome sample preparation, data acquisition, and analysis by LC–MS/MS.

Table S1 List of the 42 variables detected in Arabidopsis rosettes and identification parameters.

Table S2 GC–MS metabolomic raw data set: relative content of 42 metabolites in Arabidopsis leaves extracts (arbitrary unit: normalized peak area of integrated EICs).

Table S3 Nucleotide sequences for qRT-PCR analysis.

Table S4 Upregulated genes in Arabidopsis OXIDATIVE SIGNAL-INDUCIBLE1 (*OXII*) lines.

Table S5 Proteome analysis of Arabidopsis OXIDATIVE SIGNAL-INDUCIBLE1 (*OXII*) lines.

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