Human macrophage migration inhibitory factor (MIF) is an atypical chemokine implicated in intercellular signaling and innate immunity. MIF orthologs (MIF/D-DT-like proteins, MDLs) are present throughout the plant kingdom, but remain experimentally unexplored in these organisms. Here, we provide an in planta characterization and functional analysis of the three-member gene/protein MDL family in Arabidopsis thaliana. Subcellular localization experiments indicated a nucleo-cytoplasmic distribution of MDL1 and MDL2, while MDL3 is localized to peroxisomes. Protein–protein interaction assays revealed the in vivo formation of MDL1, MDL2, and MDL3 homo-oligomers, as well as the formation of MDL1-MDL2 hetero-oligomers. Functionally, Arabidopsis mdl mutants exhibited a delayed transition from vegetative to reproductive growth (flowering) under long-day conditions, but not in a short-day environment. In addition, mdl mutants were more resistant to colonization by the bacterial pathogen Pseudomonas syringae pv. maculicola. The latter phenotype was compromised by the additional mutation of SALICYLIC ACID INDUCTION DEFICIENT 2 (SID2), a gene implicated in the defense-induced biosynthesis of the key signaling molecule salicylic acid. However, the enhanced antibacterial immunity was not associated with any constitutive or pathogen-induced alterations in the levels of characteristic phytohormones or defense-associated metabolites. Interestingly, bacterial infection triggered relocalization and accumulation of MDL1 and MDL2 at the peripheral lobes of leaf epidermal cells. Collectively, our data indicate redundant functionality and a complex interplay between the three chemokine-like Arabidopsis MDL proteins in the regulation of both developmental and immune-related processes. These insights expand the comparative cross-kingdom analysis of MIF/MDL signaling in human and plant systems.

Human macrophage migration inhibitory factor (MIF) is a small (114 amino acids, 12.345 kDa) multifunctional protein that is best known for its role as an atypical cytokine/chemokine to regulate innate immunity. In fact, MIF was the first human cytokine to be discovered over five decades ago (1–3). Dysregulation of MIF has been associated with acute and chronic inflammatory diseases such as septic shock, rheumatoid arthritis, and atherosclerosis, as well as autoimmune conditions and cancer (4–7). The expression of MIF in the human organism is not limited to immune cells but is broadly detectable in various tissues and cell types. Unlike other cytokines/chemokines, MIF lacks a canonical N-terminal signal peptide and is released from preformed intracellular stores into the extracellular environment via stimulus-regulated unconventional secretion (1, 8). MIF’s cytokine/chemokine activities are mediated through interaction with its cognate receptor CD74 and/or by noncognate engagement of one of its three CXC chemokine receptors in a cell-, tissue-, or disease-specific manner (1, 3, 9, 10). CD74 is a single-pass transmembrane protein, also known as the human HLA class II histocompatibility antigen γ chain (9, 11). Upon MIF binding, signaling from cell-surface-expressed CD74 prominently regulates cell proliferation, apoptosis, and inflammatory gene expression. The chemokine receptors of MIF, CXCR2, CXCR4, and CXCR7, belonging to the class of heptahelical membrane proteins, function as G-protein-coupled receptors (GPCRs) and are the bona fide receptors for classical chemokines such as CXCL8, CXCL12, and CXCL11, respectively. MIF engages...
these receptors via a structural mimicry mechanism to promote leukocyte recruitment responses that can play an important pathogenic role in cardiovascular and inflammatory diseases (6, 7, 10).

In addition to its cytokine/chemokine functions in the extracellular space (“MIF the cytokine”), MIF has a number of suggested intracellular functions mediated by protein–protein interactions or enzymatic activity. This is in line with its high degree of evolutionary conservation, and it has been speculated that intracellular, enzymatic activities of MIF are evolutionarily ancient (“MIF the enzyme”) (3). Reported enzymatic activities comprise a tautomerase activity linked to an N-terminal proline-containing catalytic pocket (3, 12), an oxidoreductase activity that is dependent on a central CXXC motif (3, 13), a nuclear endonuclease activity (3, 14), as well as a chaperone-like role (3, 15). However, the precise molecular mechanisms of these activities and their potential interplay are incompletely understood and their physiological significance remains elusive. Many vertebrates including humans possess a paralog of MIF, termed D-dopachrome tautomerase (D-DT) or MIF-2. In humans, D-DT/MIF-2 shares approximately 35% amino acid identity and a high degree of architectural similarity with MIF, and the protein recapitulates some of MIF’s pathogenic activities. However, the physiological and pathogenic functions of D-DT are less well characterized than those of MIF (16, 17).

MIF-like proteins are conserved in most eukaryotes, including plants, with signs of neofunctionalization in some taxa (18, 19). Previously, we performed a comprehensive in silico analysis of plant MIF/D-DT-like (MDL) proteins, focusing on the dicotyledonous reference plant species Arabidopsis thaliana (20). We found that seed plants typically express two (gymnosperms) or three (angiosperms) different paralogs, which in Arabidopsis have been named MDL1 (AT5G51660), MDL2 (AT5G01650), and MDL3 (AT3G01660). Publicly accessible microarray data indicate that MDL1 and MDL2 are essentially expressed constitutively in aerial plant organs with little responsiveness to abiotic or biotic stress cues, while expression of MDL3 appears to be stress-inducible. The latter gene exhibits strongly enhanced transcript accumulation in leaves upon various abiotic (cold treatment, osmotic and oxidative stress, wounding, UV-B exposure) and biotic (microbial elicitors, various pathogens) stress factors. MDL3 shows in addition coexpression with a number of prominent genes involved in plant immunity (20). Structure prediction and preliminary experimental data suggest that all three Arabidopsis MDL proteins resemble the secondary and tertiary structure of human MIF (20, 21).

In contrast to human and murine MIF, very little is known about the function of plant MDLs. Analysis of recombinant epitope-tagged Arabidopsis MDLs revealed an unexpected lack of tautomerase activity, which is possibly conditioned by an amino acid polymorphism in their catalytic clefts. Surprisingly, the three MDLs can bind to the human MIF receptors CD74 and CXCR4 (which are absent from plants), activate signaling activities downstream of these in human immune cells, and substitute for human MIF in leukocyte recruitment. These findings disclose cross-kingdom mimicry of human MIF by these plant orthologs and reflect their (partial) functional conservation (21, 22). Reminiscent of the situation of human immune cells and some parasitic pathogens (19, 23), plant-feeding aphids secrete an MIF ortholog into host cells to suppress plant immune responses, which is necessary for their survival and nourishment (24).

Here, we functionally characterized the three Arabidopsis MDL proteins in planta by a comprehensive set of biochemical, cell biological, and genetic experiments. We validated the respective gene models, determined subcellular protein localizations, and studied their interactions, revealing the formation of MDL homo- and, in part, hetero-oligomers. Using a set of mdl mutants, we further uncovered roles for MDLs in the control of flowering time and bacterial pathogenesis.

Results

Arabidopsis MDL1 and MDL2 are subject to alternative splicing

The TAIR (The Arabidopsis Information Resource; https://www.arabidopsis.org/) database catalogs two different gene models for MDL1 (designated MDL1.1 and MDL1.2), four different gene models for MDL2 (MDL2.1, MDL2.2, MDL2.3, and MDL2.4), leading to three different predicted protein variants), and a single gene model for MDL3 (Fig. S1A, (20)). Protein structure prediction suggests that these transcript variants give rise to MDL forms with different C-terminal tail regions (Fig. S1B). To confirm experimentally the existence of the respective MDL transcript versions and to explore a putative tissue specificity of their expression, we first developed sets of splice variant-specific oligonucleotide primer pairs for reverse transcriptase–polymerase chain reaction (RT-PCR) (Fig. S1C). We validated the specificity of these primer pairs by using plasmid DNA harboring cloned versions of the various predicted MDL transcripts (except MDL2.4) as a template (Fig. S1D). Semiquantitative RT-PCR analysis of RNA samples from various Arabidopsis organs indicates that (i) all five tested MDL1 and MDL2 transcript variants exist (the used primer pairs did not discriminate between MDL2.2 and MDL2.3; Fig. S1C), (ii) MDL1.1 and MDL2.1 appear to be the predominant transcript versions, and (iii) that there is no pronounced organ specificity in the accumulation of any of the various splice variants (Fig. S1E).

Subcellular localization of Arabidopsis MDL proteins

We previously performed in silico analysis of the three MDL proteins regarding the presence of subcellular targeting signals. This revealed a putative nuclear localization signal for MDL1 and a predicted C-terminal peroxisomal targeting sequence (PTS1) for MDL3 (20). Proteomic studies further reported MDL1 and MDL2 as being stromal chloroplast proteins (25, 26) and MDL3 as being a peroxisomal protein (27, 28). Accordingly, the Arabidopsis Cell electronic Fluorescent Pictograph (eFP) Browser at BAR ePlant (https://bar.utoronto.ca/eplant/) presents MDL1 and MDL2 preferably localizing to chloroplasts and MDL3 to peroxisomes (Fig. S2).
Figure 1. Subcellular localization of MDL proteins. A, transient expression of N-terminally (upper part) and C-terminally (lower part) mCherry-tagged MDL fusion proteins in Arabidopsis mesophyll protoplasts. Imaging was performed by CLSM. From left to right: mCherry signal displayed in magenta; chlorophyll autofluorescence shown in green; overlay of mCherry and autofluorescence signals. B, transient expression of N-terminally mCherry-tagged MDL fusion proteins in Arabidopsis mesophyll protoplasts derived from a transgenic GFP-PTS1 reporter line. Imaging was performed by CLSM. From left to right: mCherry signal displayed in magenta; GFP signal shown in green; overlay of mCherry and GFP signals. C, subcellular localization of N-terminally mCherry-tagged MDL fusion proteins in leaf epidermis and mesophyll cells, stably and constitutively expressed in three-week-old seedlings of respective transgenic lines. Imaging was performed by CLSM. Left: mCherry signal displayed in magenta; right: overlay of mCherry (magenta) and chlorophyll autofluorescence (green) signals. D, subcellular localization of N-terminally mCherry-tagged MDL fusion proteins in root tip cells, stably and constitutively expressed in
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To explore the subcellular localization of the MDLs experimentally, we generated N- and C-terminally mCherry-tagged variants and expressed these transiently in Arabidopsis leaf mesophyll protoplasts under the control of the constitutive cauliflower mosaic virus 35S promoter. We recorded mCherry fluorescence and chlorophyll autofluorescence in transformed protoplasts by confocal laser scanning microscopy (CLSM). This revealed mainly cytoplasmic and frequently nuclear localization of MDL1 (both MDL1.1 and MDL1.2) and MDL2 (both MDL2.1 and MDL2.2), irrespective of the site of tagging (N- or C-terminal). We further noted a punctate mCherry fluorescence pattern for the N-terminally tagged MDL3 fusion protein and nucleocytoplasmic localization for the C-terminally tagged MDL3 variant. No mCherry-derived fluorescence was observed in chloroplasts, which could be unequivocally identified by their characteristic chlorophyll-mediated autofluorescence, for any of the MDL fusion proteins (Fig. 1A). Colocalization of mCherry-MDL3 with a canonical peroxisomal marker (GFP-PTS1; (29)) upon transient expression in leaf mesophyll protoplasts derived from a transgenic GFP-PTS1 marker line supports the presence of a C-terminal PTS1 targeting signal, MDL3 interacted with the PTS1 receptor PEX5. Its second interacting protein, AT5G64160, lacks any recognizable protein domains but was already found previously as an interactor of MDL3 (31).

In the following, we focused on the putative homo- and hetero-oligomerization of the MDLs by exploring all possible pairwise interactions between the three proteins using three different in planta experimental approaches. As a first approach, we used the split luciferase assay, which is based on the complementation of N- and C-terminal luciferase fragments translationally fused to the proteins of interest. Successful establishment of luciferase activity by fragment complementation can be assessed quantitatively by luminometric measurements. We transiently coexpressed the three MDLs N-terminally tagged with cLuc (C-terminal luciferase fragment) and nLuc (N-terminal luciferase fragment) in Nicotiana benthamiana and scored the resulting luminescence. This revealed noticeable luciferase activities upon coexpression of cLuc-MLD1 in combination with nLuc-MDL1 and nLuc-MDL2, cLuc-MDL2 in combination with nLuc-MDL1 and nLuc-MDL2, as well as cLuc-MDL3 in combination with nLuc-MDL2 and nLuc-MDL3. Accordingly, the respective pooled luminescence intensity values of multiple experimental replicates differed in a statistically significant manner from empty vector controls, which was not the case for the other tested pairwise combinations (Fig. 2, B and C).

As a second approach, we transiently coexpressed epitope-tagged (FLAG and mCherry) versions of various combinations of the three MDLs in N. benthamiana and performed co-immunoprecipitation (co-IP) assays using epitope-directed antibodies. This revealed pull-down of mCherry-MDL1 together with FLAG-MDL1 and FLAG-MDL2 of mCherry-MDL2 together with FLAG-MDL1, as well as of mCherry-MDL3 together with FLAG-MDL3 (Fig. 2D).

Finally, as a third approach, we took advantage of the transgenic Arabidopsis Col-0 lines constitutively expressing one of the three mCherry-tagged MDLs (Fig. 1C) and a monoclonal antibody directed against MDL2 (designated ATM 20C8; Fig. S3A) to perform co-IP experiments under seminative conditions in Arabidopsis leaf extract. This procedure yielded pull-down of mCherry-MDL1 together with MDL2 as well as mCherry-MDL2 (here MDL2.2) together with MDL2 (Fig. 2E). In summary, data of four independent protein interaction assays (Y2H, split luciferase, co-IP in

five-day-old seedlings of respective transgenic lines. Imaging was performed by CLSM. From left to right: mCherry signal displayed in magenta; DAPI signal shown in blue; overlay of mCherry (magenta) and DAPI (blue) signals. Images shown in (A–C) are maximum projections of z-stacks, micrographs shown in (D) represent one focal plane. Scale bars represent 25 μm.
Figure 2. Arabidopsis MDL proteins form homo- and in part hetero-oligomeric complexes. A, Y2H assay. Candidate interactions of a previous high-throughput screen were validated by pairwise one-on-one matings using DNA-binding domain (DB) DB-MDL1, -MDL2, and -MDL3 or empty vector (EV) as bait for prey activation domain (AD)-tagged candidate interactors, including EV. Interactions were assayed by growth on selective plates using the HIS3 reporter. The experiment was performed three times with similar results.

B and C, split-luciferase assay. cLUC-MDL1, -MDL2, and -MDL3 were transiently co-expressed with nLUC-MDL1, -MDL2, and -MDL3 or nLUC-EV (empty vector) in N. benthamiana. At 1 day post inoculation (dpi) (for cLUC-MDL1 combinations) or 2 dpi (for cLUC-MDL2 and cLUC-MDL3 combinations), plants were sprayed with 1 mM luciferin for the detection of luminescence. B, Quantification of data from the split-luciferase complementation assay. The luminescence of four independent experiments was quantified and is shown as a boxplot. Statistical significance between the interactions of the three MDLs and the EV control was determined with a two-way multipaired ANOVA test.
indicative of the split luciferase and co-IP experiments in *N. benthamiana*

Table 1

| Protein 2 | MDL1 | MDL2 | MDL3 |
|-----------|------|------|------|
| MDL1      | Y2H: yes | Y2H: yes | Y2H: no |
| co-IP (Nb): yes | co-IP (Nb): yes | co-IP (Nb): no |
| split luc: yes | split luc: yes | split luc: no |
| MDL2      | Y2H: yes | Y2H: no | Y2H: no |
| co-IP (Nb): yes | co-IP (Nb): yes | co-IP (Nb): no |
| split luc: yes | split luc: yes | split luc: yes |
| MDL3      | Y2H: no | Y2H: no | Y2H: no |
| co-IP (Nb): no | co-IP (Nb): no | co-IP (Nb): yes |
| split luc: no | split luc: no | split luc: yes |

*Y2H*, yeast two-hybrid assay; co-IP (Nb), co-immunoprecipitation in *N. benthamiana*; split luc, split luciferase assay; co-IP (At), co-immunoprecipitation in *A. thaliana*.

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**Table 1**

**Summary of MDL protein–protein interaction data**

**Protein 1**

**Protein 2**

MDL1

Y2H: yes

co-IP (Nb): yes

split luc: yes

Y2H: yes

co-IP (Nb): yes

split luc: yes

Y2H: no

co-IP (Nb): no

split luc: no

MDL2

Y2H: yes

co-IP (Nb): yes

split luc: yes

Y2H: no

co-IP (Nb): no

split luc: yes

MDL3

Y2H: no

co-IP (Nb): no

split luc: no

Y2H: no

co-IP (Nb): yes

split luc: yes

**Arabidopsis** *mdl* mutants exhibit a delayed flowering phenotype

During the crossing procedure and propagation of the *mdl* mutants, we noticed that some of these appeared to initiate flowering later than isogenic Col-0 wild-type plants. To explore this potential mutant phenotype systematically and to discriminate a putative general delay in plant development from a delayed switch from vegetative to reproductive growth, we performed experiments under controlled conditions. In these experiments, we included, apart from the Col-0 wild-type, mutants short vegetative phase-41 (*svp*-41; (32)) and constans-10 (co-10 (33)), which are known for early and late flowering, respectively, as additional controls. We grew the two *mdl* triple mutants and the respective single mutants both under short- (8 h light; SD) and long-day (16 h light; LD) conditions and scored the total number of leaves (rosette plus cauline leaves) upon bolting as an indirect readout of flowering time. Under LD conditions, Col-0 wild-type plants had a median number of 15 leaves when transitioning to flowering, similar to previous reports (32). Control mutants *svp*-41 and *co*-10 showed the expected early and late flowering phenotypes in LD conditions, with a median of 8 and 53 leaves, respectively. Both *mdl* triple mutants differed in leaf numbers in a statistically significant manner from Col-0 wild-type, with medians of 23 (*mdl*1-1 *mdl*2-2 *mdl*3-3) and 21 (*mdl*1-2 *mdl*2-1 *mdl*3-2) leaves, respectively. The values of the six corresponding *mdl* single mutants were in a similar range as for the two triple mutants, though in tendency somewhat lower, but still significantly different from Col-0 wild-type according to statistical analysis (Fig. 3, A and B). Under SD conditions, on the other hand, the various *mdl* single and triple mutants showed a similar number of leaves at the transition from vegetative to reproductive growth as Col-0 plants and *co*-10 mutant plants (median of 65–81 leaves; no statistically significant difference). Only *svp*-11 mutants still flowered earlier (median of 21 leaves; Fig. 3C), as expected. In summary, the *mdl* mutants exhibit a consistent photoperiod-dependent delayed flowering phenotype, which is most pronounced in the two triple mutants and weaker but still recognizable in the six single mutant lines.

(***< 0.001, ns = not significant**). Raw data and exact statistical values for this graph can be found in in the supplemental source data file (Supporting file 1). C. Representative photographs of *N. benthamiana* leaves and the emitted luminescence (from blue = low to red = high). D. co-immunoprecipitation of epitope-tagged MDLs. FLAG-MDL1, -MDL2, and -MDL3 and mCherry-MDL1, -MDL2, and -MDL3 were transiently coexpressed in *N. benthamiana*. At 2 dpi, proteins were extracted and immunoprecipitation was performed for the mCherry-MDL proteins using α-RFP agarose (capturing mCherry). mCherry-MDL proteins were detected with an α-RFP antibody and FLAG-MDL proteins with a monoclonal α-FLAG antibody. Ponceau staining was used as a loading control. Co-immunoprecipitation was repeated twice with similar results. E. *in planta* protein–protein interaction of MDL2. Three-week-old transgenic *Arabidopsis* seedlings stably expressing mCherry-MDL fusion proteins (transgenic lines; “TL”) were used for the immunoprecipitation of mCherry-MDL proteins using α-RFP agarose. mCherry-MDLs were detected with an α-RFP antibody and MDL2 was detected with the monoclonal α-MDL2 antibody ATM 20C8. Ponceau staining was used as a loading control. Co-immunoprecipitation was repeated twice with similar results.

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Arabidopsis mdl mutants possess unaltered infection phenotypes to adapted and nonadapted powdery mildew fungi

Given that MDL3 is coexpressed with several key components of plant defense (20), we suspected that at least mdl3 mutants might play a role in the plant response to pathogens. We thus challenged mdl mutants with various plant pathogens to reveal whether they show any altered infection phenotype. We first inoculated the two sets of mdl mutants (single and triple mutants) with conidiospores of the fungus

| AGI identifier | Gene name | Mutant | T-DNA line | Site of insertion |
|----------------|-----------|--------|------------|------------------|
| AT5G57170      | MDL1      | mdl1-1 | SAIL_772_G01| Exon 2           |
|                |           | mdl1-2 | GK_786_B08 | Intron 2         |
| AT5G01650      | MDL2      | mdl2-1 | SALK_024488C| Intron 1         |
|                |           | mdl2-2 | SALK_078883| Intron 2         |
| AT3G51660      | MDL3      | mdl3-2 | SM_3_31346 | 3' untranslated region |
|                |           | mdl3-3 | GK_750_H06 | Intron 1         |

![Figure 3](image_url)
Plant chemokine-like MDL proteins

Golovinomyces orontii, which is an adapted powdery mildew pathogen of Arabidopsis (34). The resulting macroscopically visible infection phenotype (white powder-like fungal conidiation) was comparable for Col-0 and the tested mdl mutants at 7 days post inoculation (dpi) (Fig. S6A). Similarly, the microscopically assessed host cell entry rates of G. orontii conidiospores at 48 hours post inoculation (hpi) were similar for Col-0 wild-type plants and the two mdl triple mutants (median of 88–91%; no statistically significant difference; Fig. S6B). By contrast, the highly resistant mlo2 mlo6 mlo12 control mutants (35, 36) allowed very low levels of host cell entry (median of 0–2%; Fig. S6B) and lacked the characteristic signs of fungal sporulation (Fig. S6A), as expected.

Since MDL3 is coexpressed with genes that play a key role in resistance to nonadapted pathogens such as PEN2 (37) and PEN3 (38) and because MDL3, like PEN2, localizes to peroxisomes (Fig. 1B; (37)), we also tested the outcome of interactions between mdl mutants and the nonadapted barley powdery mildew pathogen, Blumeria graminis fsp. hordei (Bgh). Col-0 wild-type plants had a median Bgh host cell entry rate of 10%, while the pen2-1 mutant, which is defective in nonhost resistance, allowed 34% (median) penetration by Bgh. All six tested mdl single mutants showed similar host cell entry rates as Col-0 (median of 8–13%, no statistically significant difference to Col-0, Fig. S6C).

Arabidopsis mdl mutants exhibit enhanced basal resistance to Pseudomonas syringae pv. maculicola

Next, we explored the infection phenotype of the mdl mutants upon challenge with the bacterial pathogen P. syringae pv. maculicola strain ES4326 (Psm), which is virulent on Arabidopsis. For this purpose, we took advantage of a strain that carries a chromosomal integration of the Photorhabdus luminescens luciferase gene (here designated as Psm lux). This strain is well established and allows to assess bacterial propagation by measuring the luminescence emitted from infected leaves (39, 40). We infiltrated bacterial suspensions in rosette leaves of five-week-old Col-0 wild-type and mdl mutant plants and scored the macroscopic infection phenotype and resulting luminescence at 3 dpi. We observed the characteristic Psm-induced leaf chlorosis in Col-0 plants, while we noticed reduced disease symptoms (chlorosis and water-soaked lesions) in the case of the mdl triple mutants (Fig. 4A). This correlated well with a strongly (approximately tenfold) reduced luminescence (i.e., bacterial propagation) in the mdl triple mutants as compared with Col-0. The bacteria-derived luminescence of the six corresponding mdl single mutants was intermediate, i.e., lower than in the case of Col-0 but higher than in the respective mdl triple mutants (Figs. 4B and S7).

In addition to the enhanced basal resistance phenotype of mdl mutants with Psm in local leaves, MDL3 shows characteristics of genes potentially involved in systemic acquired resistance (SAR), an induced form of resistance in systemic tissue after local infections with (hemi-)biotrophic pathogens (41). These features comprise (i) increased transcript levels of MDL3 (but not MDL1 or MDL2) in systemic leaf tissue of Arabidopsis Col-0 in the course of biologically induced SAR, and (ii) induction of MDL3 transcript accumulation by piperoc acid (a precursor of the key SAR signaling molecule N-hydroxy-piperic acid) application in a FLAVIN-DEPENDENT MONOOXYGENASE1 (FMO1)-dependent manner ((42, 43); Fig. S8A). We therefore also explored a potential deficiency of mdl3 mutants to develop SAR. To this end, we infiltrated primary (local) leaves of Col-0 and mdl3 mutant plants with either buffer (MgCl2, mock control) or Psm, which is a potent inducer of SAR (44). Two days later, we challenged systemic leaves with Psm lux and determined bacterial luminescence in these leaves at 3 dpi. This experiment revealed a strong reduction in bacterial titers in the systemic leaves of all plants that were inoculated with Psm bacteria in local leaves. Since there was no detectable difference between Col-0 wild-type plants and the two tested mdl3 mutants, we conclude that mdl3 mutants retain the capacity to induce SAR (Fig. S8B). The reduced bacterial luminescence in the mock-treated mdl3 mutant plants in comparison to Col-0 wild-type nevertheless further corroborated the enhanced resistance of mdl3 mutants to Psm (cf. Figs. 4B and S7).

Enhanced basal resistance of Arabidopsis mdl mutants to Psm is compromised by additional mutation of the salicylic acid biosynthesis gene SID2

Salicylic acid (SA) is a key phytohormone that orchestrates the plant immune response to various pathogens, including bacteria (45). We thus hypothesized that SA might be involved in the enhanced disease resistance of the mdl mutants. To test this proposition experimentally, we took advantage of genetic analysis by crossing the mdl1-1 and mdl3-2 single mutants as well as the mdl1-1 mdl2-2 mdl3-3 triple mutant with the sid2-1 mutant, which is defective in the gene encoding isochorismate synthase 1, a key enzyme in salicylic acid biosynthesis (46, 47). We challenged homozygous mdl1-1 sid2-1 and mdl3-2 sid2-1 double mutant as well as mdl1-1 mdl2-2 mdl3-3 sid2-1 quadruple mutant plants and respective controls (Col-0, sid2-1) with Psm lux by leaf infiltration and recorded the resulting bacterial luminescence at 3 dpi. The outcome of this assay recapitulated in tendency the enhanced resistance phenotype of mdl1-1, mdl3-2, and mdl1-1 mdl2-2 mdl3-3 (Fig. 5, A and B, cf. Fig. 4) and revealed the expected supersusceptibility phenotype of the sid2-1 mutant (46). Consistently, all genotypes harboring the sid2-1 mutation in addition to the mdl alleles showed the typical Psm disease symptoms (Fig. 5A) and permitted bacterial proliferation at levels similar to sid2-1 plants (Fig. 5B; mdl1-1 sid2-1 and mdl3-2 sid2-1 not shown). These data indicate that the sid2-1 mutation is genetically epistatic to the mdl mutations. They further suggest that the enhanced resistance phenotype of mdl mutants might depend on the defense-based accumulation of the signaling molecule SA.

Given that according to genetic analysis SA contributes to the enhanced bacterial resistance of mdl mutants, we wondered whether the accumulation of SA and/or other relevant signaling and defense-associated molecules would be altered in these mutants following pathogen challenge. We thus determined the
levels of total SA, jasmonic acid-isoleucine (JA-Ile), N-hydroxy-pipolic acid (NHP), abscisic acid (ABA), camalexin, indole-3-carboxylic acid (ICA), and some others in the rosette leaves of Col-0 control plants, two mdl1 single mutants, and both mdl triple mutants. The respective leaf samples were collected in mock-treated and Psm-inoculated plants at 32 hpi. We found a strong increase of total SA, NHP, ABA, camalexin, and ICA as well as a marked decrease of JA-Ile upon bacterial inoculation, but did not notice any statistically significant difference between the genotypes tested. Likewise, we did not observe any constitutive accumulation or depletion of any of these molecules in the mock-treated mdl mutants as compared with Col-0 wild-type plants (Figs. 5C and S9). Taken together, the enhanced disease resistance phenotype of mdl mutant plants does not correlate with any detectable differences in the constitutively present and inducible levels of the signaling and defense metabolites analyzed.

**Bacterial infection induces changes in Arabidopsis MDL transcript and protein levels and triggers subcellular relocation of MDL proteins**

With the aim to characterize the role of the MDL genes/proteins during bacterial pathogenesis further, we next studied transcript and protein accumulation in time-course experiments upon Psm challenge. In accordance with publicly accessible microarray data (http://bar.utoronto.ca/efp/cgi-bin/efpWeb.cgi), semiquantitative RT-PCR revealed opposite trends for MDL1 and MDL2 on the one hand and MDL3 on the other hand, with reduced transcript levels of MDL1 (from 12 hpi onwards) and MDL2 (at 48 hpi) and an increase in MDL3 transcript levels (from 12 hpi onward; Fig. S10). Immunoblot analysis with the monoclonal α-MDL2 antibody, which does not cross-react with bacterial proteins (Fig. S11A), indicated a putative moderate increase (approximately twofold) in MDL2 protein abundance at later time points during Psm infection (at 24 hpi and 48 hpi; Fig. S11B).

To find out whether bacterial colonization would result in a change of subcellular protein localization, we analyzed leaf samples of seedlings of the above-mentioned transgenic lines stably expressing mCherry-MDL1.1, mCherry-MDL1.2 mCherry-MDL2.2, and mCherry-MDL3 (Fig. 1, C and D) upon spray challenge with Psm by CLSM. We observed the seemingly erratic formation of prominent cytoplasmic MDL1.1 and MDL1.2 (and to a lesser degree also MDL2.2) protein aggregates at the peripheral lobes of leaf epidermal cells following bacterial infection (Fig. 6A). Plasmolysis with 1 M MgCl₂...
Figure 5. Enhanced disease resistance of the Arabidopsis mdl mutants to Psm depends on SA biosynthesis but is associated with unaltered SA levels. A and B, three mature leaves of five-week-old Arabidopsis plants were infiltrated with Psm lux (OD$_{600}$ 0.0005). A, representative macroscopic infection phenotypes at 3 dpi of Col-0 wild-type, sid2-1, mdl1-1 mdl2-2 mdl3-3 triple, and mdl1-1 mdl2-2 mdl3-3 sid2-1 quadruple mutant plants. Crosses in magenta mark the inoculated leaves. B, boxplots represent bacterial titers of Psm lux at 0 dpi and 3 dpi in leaves of mdl1-1 and mdl3-2 single as well as mdl1-1 mdl2-2 mdl3-3 triple mutants in comparison to Col-0 wild-type, the sid2-1 single mutant, mdl1-1 sid2-1 and mdl3-2 sid2-1 double mutants as well as the mdl1-1 mdl2-2 mdl3-3 sid2-1 quadruple mutant. Titers were recorded via bacterial bioluminescence and are given as RLU cm$^{-2}$. The number of evaluated plants per genotype and time point is given as $n$ above the x-axis with each plant value representing the mean of three leaves. Statistical significance between the plant genotypes was determined for the 0 dpi and 3 dpi datasets, each in comparison to the respective Col-0 control, with a two-way multipaired ANOVA test (****$p<0.0001$, ***$p<0.001$, **$p<0.01$, ns = not significant). Shown are data from one representative experiment of two independent biological replicates with similar results. Raw data and exact statistical values for this graph can be found in the supplemental source data file (Supporting file 1). C, local levels of total SA at 32 hpi in leaves of two independent mdl1 single and two independent mdl1 mdl2 mdl3 triple mutants and Col-0 wild-type following infiltration with Psm (OD$_{600}$ 0.005) or buffer (10 mM MgCl$_2$, mock control). Boxplots represent data from two independent biological replicates.
resulted in retraction of the fluorescent signal from the cell walls, which suggests that these aggregates were indeed formed in the cytoplasm and are not present in the extracellular (apoplastic) space (Fig. 6B). This is consistent with the finding that in immunoblot analysis the vast majority of the MDL2 pool appears to be intracellular following Psm challenge, with only a weak or no signal in apoplastic washing fluids (Fig. S12). In summary, these experiments provide evidence for transcriptional and posttranscriptional MDL dynamics upon bacterial infection.

Discussion

MIF and MIF-like proteins such as D-DT have been studied primarily in humans and animals (1, 3, 16) and in some of their unicellular pathogens, e.g., members of the genera Plasmodium and Leishmania (19, 48). It has been shown that these protists deliver their own MIF variants into their hosts to interfere with immune signaling by impeding host MIF functionality (49–51). Similarly, the release of aphid MIFs into a plant host has been demonstrated to modulate plant immunity (24). Apart from the latter report, a comprehensive in silico analysis (20),

Figure 6. Subcellular relocalization of stably expressed mCherry-tagged MDL proteins in response to Psm infection. A, three-week-old transgenic Arabidopsis seedlings stably expressing N-terminally mCherry-tagged MDL fusion proteins were spray-inoculated with Psm (OD_{600} 0.2) or mock-treated (10 mM MgCl₂). Displayed are subcellular localizations of mCherry-MDL fusion protein in leaves (epidermal and mesophyll cells) determined by CLSM at 2 dpi. B, seedling leaves from (A) after treatment with 1 mM MgCl₂ for the induction of plasmolysis. All images shown are maximum projections of z-stacks. Scale bars represent 25 μm.
recombinant protein work (21, 52), and phylogenetic studies (18), plant MIF proteins have not been subject to functional investigation, in particular not in their native organismal context. Despite the sequence-relatedness to their human counterparts, MDL proteins thus belong to the still large group of Arabidopsis proteins with yet unknown biochemical and biological function (53). Here we provide a basic experimental characterization of the Arabidopsis MDL gene/protein family, revealing a contribution of the three MDLs in this plant species to the modulation of bacterial infection and the control of flowering time.

We investigated the subcellular localization of the three Arabidopsis MDLs by expressing fluorophore-tagged protein variants both transiently in Arabidopsis leaf mesophyll protoplasts (Fig. 1, A and B) and stably in transgenic lines (Fig. 1, C and D). Both approaches yielded consistent results—nucleo-cytoplasmic localization of MDL1 and MDL2 irrespective of the site of protein tagging (N- or C-terminal) and peroxisomal localization of N-terminally tagged MDL3. Peroxisomal localization of MDL3 in our studies is in line with previous observations (27, 28, 54) and the presence of a canonical C-terminal peroxisomal targeting sequence (PTS1). Accordingly, masking of the C-terminal PTS1 sequence by a C-terminal fluorophore-tag results in nucleo-cytoplasmic instead of peroxisomal localization (Fig. 1A). The low molecular mass below the nuclear size exclusion limit of approximately 60 kDa may allow passive diffusion of cytosolic MDL1 and MDL2 into the nucleus. Human MIF has been found to be actively transported into the nucleus by the protein apoptosis-inducing factor in neurons undergoing hypoxic or excitotoxic stress (14), raising the possibility that comparable processes might be operative in plants as well. Overall, we assume that the detection of MDL1 and MDL2 in former chloroplast proteomic studies (25, 26) might be due to cytoplasmic contaminations in the chloroplast preparations or the “stickiness” of MDL proteins to these organelles.

Four different types of protein–protein interaction assays indicate homo- and hetero-oligomerization of the three Arabidopsis MDL proteins (Table 1). The few discrepancies between the outcomes with individual assays could be due to (1) the type, position, and length of the tags used to analyze the interactions, (2) the lack of expression of particular constructs in a given system, (3) unfavorable conditions for the interaction in the nucleus (Y2H), or (4) the lack of required post-translational modifications in the yeast system (Y2H). Particularly MDL1 and MDL2 seem to have the capacity to form homo-oligomers on the one hand as well as MDL1–MDL2 hetero-oligomers on the other hand (Fig. 2 and Table 1). Human MIF crystallizes as a noncovalent homo-trimer. An MIF trimer is required for the tautomerase activity of MIF since the catalytic cavity is jointly formed by residues from different subunits, and it is the trimer that is believed to bind to the MIF receptor CD74 and to represent the physiologically relevant form (55–57). The subunit–subunit interfaces within the human MIF trimer have been characterized by an intertwining loop structure. This interaction comprises prominent contributions from residues 38–44 and 96–102 of β-strands 2 and 5, respectively, as well as of the central β-sheet of one subunit, and from residues 48–50 and 107–109, respectively, from two adjacent subunits, which extend the 4-stranded central β-sheet into a 6-stranded sheet structure (55). However, cross-linking experiments performed under physiological concentrations of human MIF in solution have suggested that MIF oligomeric species may dynamically change between monomeric, dimeric, and trimeric forms. In addition, MIF engagement of its chemokine receptors CXCR2 and CXCR4 has been suggested to occur at low nanomolar concentrations at which the monomer would be the favored species (6, 10). Although an in silico comparison insinuates also a trimeric structure for the MDLs (20, 21), their specific oligomeric species in planta remains unclear.

Besides MDL–MDL interactions, no other proteins were identified as interaction partners of MDL1 or MDL2 in an Y2H screen. This might indicate that these two proteins do not require further partners. However, it must be taken into account that only about 48% of Arabidopsis genes are represented by the screened ORFeome and that the employed Y2H implementation has a documented assay sensitivity of ~35% (31). In the case of MDL3, the peroxisomal import receptor PEX5 was identified as an interactor, which is consistent with the presence of a peroxisomal PTS1 import signal and subcellular localization of MDL3 in peroxisomes. The additional two proteins found in the Y2H screen (AT2G42060 and AT5G64160) will need to be validated by in planta approaches in future studies. Finally, targeted interaction studies might be performed with known interactors of human MIF that are also present in plants, such as the COP9 signalosome subunit CSN5/JAB1 or the redox proteins peroxiredoxin-1 and superoxide dismutase-1 (3).

We obtained homozygous T-DNA knockout mutants for genes MDL1 and MDL2 and respective mdl1 mdl2 double knockout lines (Table 2 and Table S1), which are all transcript null mutants (Fig. S4). At least for the mdl2 mutants also the respective protein is undetectable (Fig. S3B). These findings indicate that MDL1 and MDL2 are dispensable in Arabidopsis, both individually and in combination, for growth and reproduction. By contrast, the available mdl3 mutants with T-DNA insertions in an intron and in the 3' untranslated region retained residual transcript levels (Fig. S4). We thus currently do not know whether MDL3 is an essential gene in Arabidopsis that results in lethality when fully inactivated.

We discovered that the mdl mutants possess a delayed flowering phenotype in LD conditions, which was most pronounced in the two independent mdl1 mdl2 mdl3 triple mutant lines (Fig. 3). This phenotype is apparent under LD conditions but not in SD settings. It will require extensive additional studies (e.g., genetic epistasis analysis with well-characterized flowering time mutants) to place the MDL proteins more precisely in the complex network of pathways controlling flowering time in various environments (58).

We found that mdl mutants exhibit unaltered infection phenotypes in response to adapted and nonadapted powdery mildew fungi (Fig. S6) but enhanced disease resistance to the bacterial pathogen Psm (Fig. 4). Enhanced disease resistance in plant...
mutants is frequently the consequence of autoimmunity, i.e., a constitutive activation of plant defense prior to pathogen challenge, which is typically correlated with retarded plant growth (dwarfism) and/or the occurrence of spontaneous cell death in leaf tissue (“lesion mimic mutants”; (59)). At the molecular level, such mutants also often show constitutive expression of defense marker genes and/or the accumulation of defense signaling molecules in the absence of any pathogen (60–62). However, mdl mutants have a normal growth habit (Fig. S5) and lack recognizable lesions and detectable changes in the level of defense-associated phytohormones and metabolites in unchallenged plants. The tested molecules also do not hyperaccumulate upon bacterial infection (Figs. 5C and S9). We thus conclude that the increased disease resistance of mdl mutants to Psm is unlikely to be due to constitutive defense or the enhanced activation of common defense pathways. The observed supersusceptibility of mdl sid2 mutants (Fig. 5B) might thus not indicate SA dependence of the mdl resistance phenotype but hint to the SA-dependent defense sector acting in parallel to the MDL pathway(s). We, therefore, hypothesize that the MDL proteins could rather serve as host susceptibility factors for P. syringae bacteria that might be targeted by conventional or unconventional bacterial effector proteins. Precedence for such effector-targeted susceptibility factors exists in various plant pathosystems (63–67). These susceptibility factors might be in turn guarded by cytoplasmic plant immune sensors, and the absence of their guarders could result in their inappropriate activation and thus enhanced disease resistance (67). As an alternative scenario we cannot rule out that MDL proteins act as negative regulators of noncanonical plant defense responses.

In the human system, recombinant MIF protein has been described to enhance biofilm formation of Pseudomonas aeruginosa, an opportunistic bacterial pathogen with a broad host range that is distantly related to Psm (68). Biofilm formation could promote bacterial infection and thus explain the proposed role of MDLs as host susceptibility factors. Although also the phytopathogen P. syringae has the capacity to form biofilms (69), this is, however, an unlikely scenario for the Arabidopsis–Psm interaction since Psm bacteria remain localized extracellularly during plant colonization, and no evidence was found for the secretion of MDL proteins during bacterial pathogenesis (Figs. 6B and S12).

Bacterial challenge triggers subcellular relocation of transgenically expressed and fluorophore-tagged MDL1 and MDL2, resulting in its aggregation at the peripheral lobes of epidermal cells (Fig. 6A). Recruitment of plasma membrane-localized defense-related proteins to sites of attempted entry has been reported for powdery mildew fungi (70, 71). This response can be recapitulated by the exogenous application of conserved pathogen elicitors such as the flg22 epitope or chitin (72). It depends on cytoskeleton function and intracellular protein trafficking (73). We are, however, not aware of any reports describing similar protein redistribution upon bacterial infection, especially considering the nucleo-cytoplasmic (nonmembrane) localization of MDL1 and MDL2. Unlike the situation for human MIF (3, 8), our experimental data do not support pathogen-triggered secretion of MDL proteins to the extracellular space (Figs. 6B and S11). The physiological relevance of the change in subcellular localization thus remains to be explored.

It is noteworthy that both mutant phenotypes discovered in this study—i.e., enhanced resistance to Psm and delayed flowering—are detectable in all mdl single mutants (including the mdl3 mutants that still retain residual transcript levels; Fig. S4) and exacerbated in the respective mdl triple mutants. This finding points to the notion that all three MDL proteins may cofunction in the same cellular pathway where they might act in partial redundancy. Such a scenario could be reconciled most easily with all three proteins cooperating within the same protein complex, with a protein assembly lacking one paralog being less active. However, while the three MDL genes are coexpressed in rosette leaves, they show nonoverlapping expression patterns in other plant organs (20). Moreover, the distinctive subcellular localization of MDL1 and MDL2 (nucleo-cytoplasmic; Fig. 1) on the one hand and MDL3 (peroxisomal; Fig. 1) on the other hand, as well as the lack of convincing evidence for physical interaction between MDL1/MDL2 and MDL3 (Fig. 2 and Table 1), argue against this hypothesis. Nonetheless, at least in rosette leaves, the MDL proteins might act as nonessential components in consecutive steps of a given cellular pathway.

Some plant mutants that exhibit both an altered flowering time and immune phenotype have been described. An example is the elevated level of resistance to the fungal pathogen Fusarium oxysporum seen in late-flowering mutants and the significant correlation of late flowering and enhanced disease resistance in natural Arabidopsis accessions. The respective study identified the photoperiodic pathway flowering time regulator GIGANTEA as a susceptibility factor for F. oxysporum infection (74). By contrast, mutants defective in the transcriptional regulator gene MED18 show late flowering in combination with enhanced susceptibility to fungal (Botrytis cinerea and Alternaria brassicicola) infection (75). In addition, in particular mutants with deficiencies in the biosynthesis or signaling pathways of defense-associated phytohormones (SA, JA-Ile, and ethylene) exhibit both deregulated plant immunity and altered flowering time. Examples comprise SA mutants such as sid2 and eds5 (76), the JA-Ile receptor mutant coi1 (77), and ethylene-insensitive mutants such as etr1, ein2, and ein3 (78). The MDL proteins are novel players that likewise modulate both the control of plant immunity and flowering time. The mechanistic details of how these three proteins engage in the cross talk between these two processes remain to be explored. The elucidation of these specifics may feed back to the human system, where intracellular MIF functions remain poorly understood. This research may thus also bring about new insights into pathomechanisms underlying human diseases, thereby closing the loop of cross-kingdom MIF studies.

**Experimental procedures**

**Plant material and growth conditions**

All Arabidopsis lines used in this study (Table S1) are in the genetic background of accession Columbia-0 (Col-0). T-DNA
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individually transplanted into bigger pots, either singly in small day (LD) conditions. Two weeks after seeding, seedlings were and vermiculite, and placed in controlled short- (SD) or long-day (LD) conditions. Two weeks after seeding, seedlings were obtained by intermutant crossings. All mutant lines were finally confirmed through DNA sequencing of the respective PCR amplicons.

**Arabidopsis** seeds were stratified, sown in an 8:1:1 mixture of soil (ED73, Balster Einheitserdewerk GmbH, Fröndenberg, Germany; sterilized by 10 min microwaving before use), sand, and vermiculite, and placed in controlled short- (SD) or long-day (LD) conditions. Two weeks after seeding, seedlings were individually transplanted into bigger pots, either singly in small pots of 6 cm diameter or as 1–4 plants per 9 × 9 × 8 cm pots depending on the purpose. Plants for flowering experiments, crossings, and seed production were grown in a 16 h light/8 h dark LD cycle at 20 °C, a relative humidity of 60–70%, and with a photon flux density of 80–100 μmol m⁻² s⁻¹. Plants for all other experiments were generally grown in SD cycles of 10 h light/14 h dark or of 8 h light/16 h dark in the case of flowering experiments, 20 °C, 60–70% relative humidity, and 80–100 μmol m⁻² s⁻¹ photon irradiance.

*N. benthamiana* plants were grown in SoMi513 soil (HAWITA, Vechta, Germany) in 9 × 9 × 8 cm pots in a 10 h light/14 h dark SD cycle at 20 °C, 80–90% relative humidity, and with a light intensity of 80–100 μmol m⁻² s⁻¹. Plants infiltrated with *A. tumefaciens* were transferred to an LD cycle of 16 h light at 23 °C/8 h dark at 20 °C, 60–65% relative humidity, and a light intensity of 105–120 μmol s⁻¹ m⁻².

**Cloning**

Coding sequences of **MDL1.1**, **MDL1.2**, **MDL2.1**, **MDL2.2**, and **MDL3** were cloned via GATEWAY Cloning Technology (Invitrogen, Carlsbad, CA, USA) into the respective entry (pDONR207) and destination vectors. The destination vectors used in this study are summarized in Table S2.

The destination vectors p35S::mCherry-GWY and p35S::GWYmCherry were generated by introducing the amplified mCherry PCR product into pExHisHA via either restriction sites EcoRI and SmaI or SmaI and XbaI, respectively. The GWY cassette was introduced into the recombinant p35S::mCherry vector by using the restriction site SmaI. The destination vectors pAMPAT-cLUC-GWY and pAMPAT-nLUC-GWY were generated by introducing either cLUC-GWY or nLUC-GWY from pCAMBIA-cLUC-GWY or pCAMBIA-nLUC-GWY (80), respectively, into pAMPAT-MCS-Sacl via the restriction sites HindIII and SacI. The GWY cassette was introduced into the transgenic plants using the NucleoSpin RNA Plus kit (Macherey-Nagel, Düren, Germany) following the manufacturer’s instructions. The resulting cDNA was diluted 1:10 before application in PCR. RT-PCR was performed as a standard application with GoTag G2 DNA Polymerase using 2.5 μl of diluted cDNA per 50 μl reaction. Gene AT4G26410 was used as a reference gene (81). The respective primer pairs used for RT-PCR are listed in Table S3.

**Gene expression analysis by reverse transcriptase–polymerase chain reaction (RT-PCR)**

Total RNA was extracted from rosette leaves of **Arabidopsis** plants using the NucleoSpin RNA Plus kit (Macherey-Nagel, Düren, Germany) following the instructor’s manual. One microgram of total RNA was treated with DNase I (Sigma-Aldrich, Merck KGaA, Darmstadt, Germany) prior to the reverse transcription into cDNA using the High-Capacity RNA-to-cDNA kit (Applied Biosystems, Invitrogen, Carlsbad, CA, USA) following the manufacturer’s instructions. The resulting cDNA was diluted 1:10 before application in PCR. RT-PCR was performed as a standard application with GoTag G2 DNA Polymerase using 2.5 μl of diluted cDNA per 50 μl reaction. Gene AT4G26410 was used as a reference gene (81). The respective primer pairs used for RT-PCR are listed in Table S3.

**Transient gene expression in Arabidopsis protoplasts**

Transfection of **Arabidopsis** mesophyll protoplasts was performed by the tape-sandwich method (82). Leaves of five-week-old **Arabidopsis** plants were fixed with the upper epidermal surface on autoclave tape. Subsequently, the lower epidermal surface was ripped off with the help of Tesa tape. Leaves were placed peeled in an enzyme solution containing cellulose OnozukaTM R-10 and macerozymeTM R-10 (both Yakult, Tokyo, Japan) and incubated gently shaking at room temperature until cell walls were digested and mesophyll protoplasts distributed in the solution. The protoplast-containing solution was collected by centrifugation at 100 × g for 3 min at 4 °C, washed twice with 15 ml prechilled W5 (154 mM NaCl, 125 mM CaCl₂, 5 mM KCl, 5 mM glucose, and 2 mM MES, pH 5.7), and incubated on ice for 30 min. The protoplasts were collected by centrifugation at 100 × g for 3 min and subsequently resuspended in 4 ml MMG solution (0.4 M mannitol, 15 mM MgCl₂, and 4 mM MES, pH 5.7). For the transfection of protoplasts, 20 μl containing 7 μg of plasmid DNA was added to 200 μl of the prepared protoplasts. Subsequently, 220 μl PEG solution (40% (w/v) PEG 4000, 0.1 M CaCl₂, 0.2 M mannitol) was added, gently mixed, and incubated at room temperature for 10 min. Thereafter, the transfected protoplasts were washed thrice with 0.9 ml W5 solution by centrifugation at 100 × g for 3 min. After the last washing step, the protoplasts were resuspended in 0.9 ml W5 and incubated in the dark overnight at room temperature.

**Confocal laser scanning microscopy (CLSM)**

For localization of fluorophore-tagged proteins, leaves of transgenic **Arabidopsis** seedlings were placed in a droplet of water on a glass slide and shielded with a coverslip. Analyses were performed with a Leica TCS SP8 lightning confocal microscope and the LAS-X software package (Leica Microsystems, Wetzlar, Germany). Pictures and z-stacks were recorded with a 20x water or 63x water immersion objective (Leica). For DAPI staining, seedlings were incubated in 1:500 diluted DAPI (AppliChem GmbH, Darmstadt, Germany) solution (DAPI stock solution 5 mg/ml in dimethylformamide...
(Sigma-Aldrich)) for 2 h. Fluorescence signals were recorded in sequential scan mode with the following specifications: GFP excitation at 496 nm (argon laser) and emission at 499–523 nm; mCherry excitation at 561 nm (DPSS 561 diode) and emission at 589–622 nm; chlorophyll autofluorescence excitation at 476 nm and emission at 681–762 nm; DAPI excitation at 405 nm (Diode 405) and emission at 420–470 nm.

For the analysis of subcellular localization of fluorophore-tagged proteins after Psm inoculations, three-week-old SD-grown transgenic Arabidopsis seedlings were inoculated with Psm at an optical density OD at 600 nm (OD₆₀₀) of 0.2 or, for control, treated with 10 mM MgCl₂. Treatments were performed by spraying the solutions, each supplemented with 0.04 % v/v Silwet L-77 (Momentive Performance Materials GmbH, Leverkusen, Germany), twice from all sides. One day before and for 1 day after the treatment, the seedlings were covered with a lid to induce opening of the stomata in high humidity. Treated seedlings were analyzed by CLSM at 4, 24, and 48 hpi using the specifications given above.

**Generation of transgenic Arabidopsis plants**

Binary vectors with the open reading frames (ORFs) of MDL1.1, MDL1.2, MDL2.1, MDL2.2, and MDL3, N-terminally fused to mCherry, were introduced into Arabidopsis Col-0 wild-type plants through A. tumefaciens-mediated transformation by the floral dip method (83). A. tumefaciens strain GV3101 with the helper plasmid pMP90:RK was used for the transformation. T1 seedlings of T0 seedlings recovered from the transformed plants were selected by spraying with BASTA solution (BASF, Ludwigshafen am Rhein, Germany). Sterilized T1 seeds resulting from BASTA-resistant T1 plants were individually seeded on MS plates containing BASTA. The developing T2 generation was screened for seedlings with an approximately 3:1 survival rate. T2 seeds, resulting from T2 plants with a 3:1 survival rate, were reasessed on BASTA-containing plates with MS medium to identify homozygous individuals (all seedlings survive). In addition, seedlings were checked for proper mCherry fluorescence. Homozygous individuals were multiplied and used in targeted applications.

**Flowering time experiments**

For the determination of the flowering time, Arabidopsis plants were stratified for uniform germination and grown under SD (8 h light/16 h dark) or LD (16 h light/8 h dark) cycles in controlled conditions. Uniform growth of the plants was followed by weekly counting the rosette leaves. The development of primary shoots was assessed by checking the plants every other day. Flowering time (in days) was scored when primary shoots reached 0.5 mm and was additionally assessed as the total number of rosette and cauline leaves of the main stem at the stage of full flowering.

**Powdery mildew infection assays with G. orontii and B. graminis f.sp. hordei**

Inoculation assays with G. orontii were performed with four-week-old Arabidopsis plants grown in SD conditions by leaf-to-leaf transfer of conidiospores using heavily infected leaves of eds1-2 [Col-0] mutant plants (84), which were the cultivation host for G. orontii. At 2 days post inoculation (dpi), leaf samples were collected from the infected target plants and placed in 80% (v/v) ethanol for the removal of leaf pigments. In the following, fungal structures were stained in a Coomassie solution (45% v/v methanol, 10% v/v acetic acid, 0.05% w/v Coomassie blue R-250; Carl Roth, Karlsruhe, Germany) for their visualization with a Zeiss Axio phot microscope (Carl Zeiss AG, Jena, Germany). Fungal host cell entry rates were quantified by the percentage of germinated fungal spores developing haustoria and secondary hyphae over germinated spores with restricted growth by plant formed appressoria. At least 100 germinated spores per leaf (excluding stomatal and vessel regions) and four leaves per plant were evaluated to calculate a medium penetration success per plant. In addition, extra infected plants were evaluated at 7 dpi for their macroscopically visible disease symptoms.

Inoculation assays with B. graminis f.sp. hordei (Bgh) were performed as described before (85). In brief, four- to five-week-old Arabidopsis plants were inoculated by dusting off Bgh conidia from heavily infected barley plants. Subsequently, the inoculated Arabidopsis plants were transferred back to SD conditions. At 2 dpi, infected leaves were sampled and placed in 80% (v/v) ethanol for the removal of leaf pigments. In the following, fungal structures were stained in an aniline blue solution (150 mM K₂HPO₄, 0.01% w/v aniline blue; Sigma-Aldrich, Munich, Germany) for 1 day to visualize callose via fluorescence with an Axio phot microscope (Carl Zeiss AG, Jena, Germany), using UV excitation of 327–427 nm and a DAPI/aniline blue filter of 417–477 nm emission. Fungal host cell entry rates of Bgh were quantified by the percentage of inhibited penetration attempts (stained papilla) over successful penetration (haustorium inside the epidermal cell). At least 100 interaction sites were evaluated per leaf of three plants per genotype and experiment.

**P. syringae infection assays**

Bacterial infection assays were performed with P. syringae pv. maculicola ES4326 (Psm) and a Psm strain constitutively expressing the P. linitialis luxCDABE operon (Psmlux; (39). Psm and Psmlux were grown on King’s B plates supplemented with 50 μg ml⁻¹ rifampicin for Psm and Psmlux and 25 μg ml⁻¹ kanamycin for Psmlux for 2 days at 28 °C. Overnight (log phase) liquid cultures in King’s B medium were prepared from the plates, washed thrice in 10 mM MgCl₂, and diluted to the desired optical density in 10 mM MgCl₂.

For basal bacterial growth assays, three mature leaves of five-week-old SD-grown Arabidopsis plants of uniform and healthy appearance were infiltrated from the abaxial side with Psmlux at an OD₆₀₀ of 0.0005 using a needleless 1 ml syringe. The bacterial growth in leaves was determined indirectly by recording the bacterial bioluminescence in leaf discs (r = 3 mm, adaxial side up) individually placed in 96-well microtiter plates (flat bottom) filled with 250 μl 10 mM MgCl₂. The
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average bioluminescence was recorded for 10 s per well with a 10 s with the “Centro XS” LB 960 Microplate Luminometer” and the corresponding “MicroWin 2000 software” (Berthold Technologies, https://www.berthold.com/). Three leaf discs per plant were recorded and the average auto-luminescence (background of untreated leaves) was subtracted to determine the average bioluminescence per plant. Increasing luminescence values (given in relative light units, RLU) correlate linearly with increasing bacterial titers and reflect bacterial growth determined by the standard plate assay (39, 40).

For analyses of metabolites and protein accumulation after Psm inoculation, infiltrations were performed as described above except for using Psm at an OD600 of 0.005 for inoculations and 10 mM MgCl2 as mock infiltrations. Leaf samples for metabolite analysis (six leaves per sample and three samples from six plants per genotype) were collected at 32 hpi. Leaf samples for protein analysis (two leaves from two plants per sample and three samples from six plants per genotype) were collected at 0, 2, 4, 6, 12, 24, and 48 hpi. In both cases, the samples were instantly shock-frozen in liquid nitrogen and stored at −80 °C for the following respective analysis.

SAR assays were carried out with five-week-old SD-grown Arabidopsis plants of healthy and uniform appearance. Three lower mature leaves were either infiltrated with Psm at an OD600 of 0.005 for inoculations or with 10 mM MgCl2 as a mock treatment. Two days later, three upper (systemic) mature leaves of the same plants were challenged with infiltrations of Psm lux at an OD600 of 0.0005. The bacterial growth in the challenged leaves was determined 3 dpi via bioluminescence as described above.

Transient expression in N. benthamiana

A. tumefaciens (strain AGL1) harboring appropriate constructs and the GV2260 p19 helper strain were used for transient expression in N. benthamiana as described before (85) with minor modifications. In brief, bacterial cultures were resuspended in infiltration media (10 mM MES, pH 5.6, 10 mM MgCl2, 200 μM acetosyringone) to an OD600 of 0.5 and incubated at room temperature for 2 h. For coinfiltration, equal volumes of each A. tumefaciens transformant were mixed and infiltrated with a needleless syringe from the abaxial side into fully expanded leaves of four to six-week-old N. benthamiana plants.

Protein isolation and immunodetection

For protein accumulation experiments, four leaves of different five-week-old Arabidopsis plants were homogenized in 200 μl protein extraction buffer (50 mM Tris pH 8.0, 1 mM EDTA, 1 mM DTT, 1 mM PMSF) and centrifuged for 20 min at 15,000 × g at 4 °C to remove cell debris. The protein concentration was determined using the Bradford protein assay (86). Unless otherwise stated, 5 μg total protein was used for the separation of proteins by sodium dodecylsulfate–polyacrylamide gel electrophoresis (SDS-PAGE). Each experiment was repeated at least once with similar results. Protein samples were denatured in 1xNuPAGE LDS Sample buffer (Thermo Fisher Scientific, Darmstadt, Germany) by boiling at 95 °C for 5 min prior to gel loading. Thereafter, samples were subjected to SDS-PAGE, transferred to a nitrocellulose membrane (0.2 μm, Carl Roth, Karlsruhe, Germany), and used for immunodetection according to the manufacturer’s instructions. Primary antibodies were purchased from Chromotek (6G6, α-RFP, Planegg-Martinsried, Germany), Cell Signalling (D6W5B, α-FLAG, Frankfurt, Germany) or generated (20C8, α-MDL2, see below). Secondary antibodies were purchased from Cell Signalling (7074, α-rabbit IgG HRP, Frankfurt, Germany), Merck (A9037, α-rat IgG HRP, Darmstadt, Germany), and Thermo Fisher Scientific (32,430, α-mouse IgG HRP, Darmstadt, Germany). Chemiluminescence detection of antigen–antibody complexes was performed with SuperSignal West Femto Western substrate (Thermo Fisher Scientific, Darmstadt, Germany). As a loading control, nitrocellulose membranes were stained in Ponceau S (AppliChem GmbH, Darmstadt, Germany) solution. For protein accumulation after Psm inoculation, infiltrations were performed as described above, except for using Psm at an OD600 of 0.005 for inoculations and 10 mM MgCl2 as mock infiltrations.

Generation of monoclonal α-MDL2 antibodies

For the production of MDL2-specific monoclonal antibodies, highly purified recombinant MDL2 protein was prepared using Rosetta (DE3)-competent Escherichia coli cells, which expressed a pET21-derived gene construct to yield a C-terminally 6xHis-tagged MDL2 (MDL2-6xHis) protein product. Bacterial expression and protein purification were carried out essentially as described previously (21, 52). Wistar rats were immunized with 60 μg purified full-length MDL2-6xHis, 5 nmol CpG (TIB MOLBIOL, Berlin, Germany) and an equal volume of Incomplete Freund’s adjuvant (IFA; Sigma, St Louis, USA)). After a 6-week interval, a boost injection without IFA was given 3 days before the fusion of rat spleen cells with IFA was given 3 days before the fusion of rat spleen cells with P3X63Ag8.653 myeloma cells using standard procedures (87, 88). Hybridoma supernatants were screened in a bead-based flow cytometry assay (iQue, Intellicyte; Sartorius, Göttingen, Germany). Briefly, MDL2-6xHis protein was captured to activated 3D Carboxy beads (PolyAN, Berlin, Germany) and incubated for 90 min with hybridoma supernatant and Atto 488-coupled isotype-specific monoclonal mouse-anti-rat IgG secondary antibodies. Antibody binding was analyzed using ForeCyt software (Sartorius, Göttingen, Germany). Positive supernatants were further assayed for their potential by western blot. Hybridoma cells from selected supernatants were subcloned at least twice by limiting dilution to obtain stable monoclonal cell lines. The experiments of this study were performed with the purified antibody clone ATM 20C8 (IgG2b/k).

Yeast two-hybrid (Y2H) assays

Y2H screens were performed as described before (89). Briefly, ORFs encoding MDL1.1, MDL2.1, or MDL3 were...
transferred by Gateway cloning from pDONR207 vectors into the GAL4 DNA-binding domain (DB)-encoding Y2H vector pDEST-pPC97, and subsequently transformed into the yeast strain Y8930. These constructs were screened by yeast mating against a collection of 12,000 Arabidopsis ORFs fused to the Gal4 activation domain (AD) in the yeast strain Y8800 (30). The screening was done as a binary mini-pool approach, i.e., each DB-ORF was screened against pools of 188 AD-ORFs. Interactions were assayed by growth on selective plates using the HIS3 reporter, adding 1 mM 3-amino-1,2,4-triazole (3-AT) to suppress background growth. This primary screen was carried out once and identities of candidate interactors were verified by Sanger sequencing. All candidate interactions were verified by pairwise one-on-one mating in three independent experiments. Only pairs scoring positive in all three assays were considered as bona fide interaction partners.

**Co-immunoprecipitation**

For co-IP, 500 μl of homogenized plant material (of two-week-old Arabidopsis seedlings or 2 dpi of transiently transformed N. benthamiana) was mixed with 1 ml extraction buffer (50 mM Tris-HCl, pH 8.0, 1 mM EDTA, 1 mM DTT, 1 mM phenylmethanesulfonyl fluoride (AppliChem GmbH, Darmstadt, Germany)) and centrifuged at 15,000 × g at 4 °C to remove cell debris. The protein concentration was determined using the Bradford protein assay and 500 μg of soluble proteins was incubated with 10 μl α-RFP-Trap Agarose slurry (ChromoTek GmbH, Planegg-Martinsried, Germany) for 1 h at 4 °C under constant agitation. Thereafter, beads were washed six times in 200 μl protein extraction buffer. After the last washing step, the protein extraction buffer was removed completely and the beads were resuspended in 50 μl SDS sample buffer. Briefly, before separation by SDS-PAGE, proteins were eluted by boiling the samples for 10 min at 95 °C.

**Split luciferase complementation assays**

For split luciferase complementation assays, transiently transformed leaves of N. benthamiana were sprayed with 1 mM D-luciferin (PerkinElmer, Rodgau, Germany) solved in water supplemented with 0.01% (v/v) Tween-20 at 2 days after infiltration. Leaves were kept in the dark for 10 min before luminescence was detected with a ChemiDoc XRS+ imagine system. Intensities/mm² of different combinations were evaluated using the Image Lab software (BioRad, Feldkirchen, Germany). For each combination of interaction partners, four independent experiments consisting of two different plants and two leaves per plant were evaluated.

**Preparation and analysis of apoplastic wash fluid (AWF)**

For the analysis of AWFs, five-week-old SD-grown Arabidopsis plants were sprayed with Psm or mock-treated (10 mM MgCl₂) as described above or left untreated. Four plants of each treatment and genotype were sampled at 24 and 48 hpi. To extract AWF, three leaves each of four to five Arabidopsis plants were cut and submerged in a beaker with infiltration buffer (20 mM MES, 2 mM CaCl₂, 0.1 M KCl, pH 6.0). To infiltrate, the beaker was placed in a desiccator, and a vacuum was applied for 5 min. After infiltration, the leaves were carefully padded dry and placed, the petiole facing up, in 1.5 ml tubes. The tubes were then centrifuged at 200 × g for 20 min at 4 °C. The AWF was collected from the bottom of the tubes and fluid from the same treatment was pooled. For leaf samples, two leaves from different plants were harvested and snap-frozen in liquid nitrogen. Protein was extracted from leaf samples as described above, with the addition of 0.1 % Tween to the extraction buffer. For immunoblotting, 10 μl of AWF or 7.5 μg of leaf protein was mixed with Laemmli buffer and denatured at 95 °C for 10 min. The samples were separated by SDS-PAGE, subsequently blotted onto a nitrocellulose membrane, and used for immunodetection as described above. In addition to α-MDL2, α-PsbA/D1 (AS05 084, PsbA/D1 protein of PS II, Agrisera, Vännäs, Sweden) was used as a control for cytosolic contamination.

**Analysis of phytohormones and defense-associated metabolites**

Phytohormones were extracted with methyl-tert-butyl ether (MTBE), reversed phase-separated using an ACQUITY ultra performance liquid chromatography (UPLC) system (Waters Corp., Milford, MA, USA), and analyzed by nanoelectrospray ionization (nanoESI) (TriVersa Nanomate; Advion Biosciences, Ithaca, NY, USA) coupled with an AB Sciex 4000 QTRAP tandem mass spectrometer (AB Sciex, Framingham, MA, USA) employed in scheduled multiple reaction monitoring mode (90). The reversed phase separation of constituents was achieved by UPLC using an ACQUITY UPLC HSS T3 column (100 mm × 1 mm, 1.8 μm; Waters Corp., Milford, MA, USA). Solvents A and B were water and acetonitrile/water (90:10, v/v), respectively, both containing 0.3 mmol/l ammonium formate (adjusted to pH 3.5 with formic acid).

For absolute quantification of SA, SA glycoside (SAG), JA-Ile, NHP, ABA, camalexin, and ICA, the following internal standards were added to the plant material before extraction: 10 ng D₄-SA (C/D/N Isotopes Inc, Pointe-Claire, Canada), 50 ng D₁₃-C₆-SAG (kindly provided by Petr Karlovsky, Göttin- gen, Germany), 10 ng D₃-JA-Leu (kindly provided by Otto Miersch, Halle/Saale, Germany), 50 ng D₅-Pip (Merck KGaA, Darmstadt, Germany), 50 ng D₉-NHP (91), 10 ng D₆-ABA (C/D/N Isotopes Inc, Pointe-Claire, Canada), and 20 ng D₇-IAA (Eurisotop, Freising, Germany). Mass transitions and optimized parameters for the detection of these phytohormones and defense-associated metabolites by mass spectrometry are shown in Table S4. After data processing, the absolute amounts of SA and SAG were added to express the total SA levels.

**Statistical analysis**

In this study, “biological replicates” refers to data from fully independent experiments, whereas “technical replicates” refers to data from samples of the same experiment or plant. Statistical analyses and graph generation were performed using the GraphPad Prism software (GraphPad Prism Software Inc, San Diego, CA, USA).
San Diego, CA, USA). In boxplot graphs, the center lines show the medians and upper and lower box limits indicate the 25th and 75th percentiles, respectively. Outliers are indicated by dots. Statistical analyses were performed using either a one-way analysis of variance (ANOVA) test or a two-way ANOVA test, in either case with Tukey’s method for multiple comparisons (multipaired ANOVA) as indicated in the figure legends. Raw data and details of the statistical analysis for all figures can be found in the supplemental source data file (Supporting file 1).

Data availability

All data and materials are available upon request. Access to the monoclonal α-MDL2 antibody and the transgenic MDL lines requires a material transfer agreement (MTA).

Supporting information—This article contains supporting information (21, 32, 33, 37, 42, 43, 46, 52, 84, 35, 36).

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Abbreviations—The abbreviations used are: ABA, abscisic acid; ANOVA, analysis of variance; AWP, apoplastic wash fluid; CLSM, confocal laser scanning microscopy; co-IP, co-immunoprecipitation; DAPI, 4’,6-diamidino-2-phenylindole; D-DT, D-dopachrome tautomerase; dpi, days post inoculation; hpi, hours post inoculation; ICA, indole-3-carboxylic acid; IA-Ile, jasmonic acid-isoleucine; LD, long-day; MDL, MDL/D-DT-like protein; MIF, macrophage migration inhibitory factor; NHP, N-hydroxy-pipelic acid; OD600, optical density at 600 nm; ORF, open reading frame; RLU, relative light units; RT-PCR, reverse transcription-polymerase chain reaction; SA, salicylic acid; SAR, systemic acquired resistance; SD, short-day; SDS-PAGE, sodium dodecylsulfate–polyacrylamide gel electrophoresis; Y2H, yeast two-hybrid.

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