Activity of the purified plant ABC transporter NtPDR1 is stimulated by diterpenes and sesquiterpenes involved in constitutive and induced defenses

Baptiste Pierman §1, Frédéric Toussaint §1, Aurélie Bertin §9, Daniel Lévy §9, Nicolas Smargiasso §9, Edwin De Pauw §9, and Marc Boutry §1,2

From the §1Institut des Sciences de la Vie, Université catholique de Louvain, B-1348 Louvain-la-Neuve, Belgium, the §9Laboratoire Physico Chimie Curie, Institut Curie, Paris Sciences et Lettres Research University, CNRS UMR168, and Sorbonne Universités, Université Pierre et Marie Curie Paris 06, 75005 Paris, France, and §3Mass Spectrometry Laboratory, Molecular Systems Research Unit, University of Liège, B-4000 Liège, Belgium

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Within the plant ATP-binding cassette transporter family, pleiotropic drug resistance (PDR) transporters play essential functions, such as in hormone transport or defense against biotic and abiotic stresses. NtPDR1 from Nicotiana tabacum has been shown to be involved in the constitutive defense against pathogens through the secretion of toxic cyclic diterpenes, such as the antimicrobial substrates cembrene and sclareol from the leaf hairs (trichomes). However, direct evidence of an interaction between NtPDR1 and terpenes is lacking. Here, we stably expressed NtPDR1 in N. tabacum BY-2 suspension cells. NtPDR1 was purified as an active monomer glycosylated at a single site in the third external loop. NtPDR1 reconstitution in proteoliposomes stimulated its basal ATPase activity from 21 to 38 nmol of Pi mg⁻¹ min⁻¹, and ATPase activity was further stimulated by the NtPDR1 substrates cembrene and sclareol, providing direct evidence of an interaction between NtPDR1 and its two substrates. Interestingly, NtPDR1 was also stimulated by capsidiol, a sesquiterpene produced by N. tabacum upon pathogen attack. We also monitored the transcriptional activity from the NtPDR1 promoter in situ with a reporter gene and found that, although NtPDR1 expression was limited to trichomes under normal conditions, addition of methyl jasmonate, a biotic stress hormone, induced expression in all leaf tissues. This finding indicated that NtPDR1 is involved not only in constitutive but also in induced plant defenses. In conclusion, we provide direct evidence of an interaction between the NtPDR1 transporter and its substrates and that NtPDR1 transports compounds involved in both constitutive (diterpenes) and induced (sesquiterpenes) plant defenses.

The ATP-binding cassette (ABC)³ superfamily of proteins is ubiquitous in all kingdoms of life. The family is mainly composed of primary active transporters, which constitute two nucleotide-binding domains (NBDs) and two transmembrane domains (TMDs) (1, 2). While the TMDs bind and translocate substrates across membranes, the NBDs hydrolyze ATP and provide the energy for active transport. The NBDs contain several well-conserved motifs such as Walker A and B; the D, H, and Q loops; and the ABC signature motif (3, 4). The family is particularly large in plants with more than 100 members in both Arabidopsis thaliana and Oryza sativa (5–7). In comparison, the human genome encodes for 49 ABC transporters (8). This large number of ABC proteins in plants can be related to their terrestrial and sessile life style, which makes them particularly exposed to (a)biotic stresses (9–11).

Members of the full-size ABCG subfamily (also called pleiotropic drug resistance (PDR)), which is only found in plants and fungi (12), play various essential roles in plants (for reviews, see Refs. 7, 9, and 12–15). Known functions of full-size ABCGs include hormone transport (16–20), heavy metal detoxification (21, 22), transport of cuticle precursors (23), plant defense against pathogens (24–29), transport of monoterpene indole alkaloids (30, 31), and transport of β-caryophyllene (32). In contrast, very little information is available concerning the enzymatic properties of plant PDRs or, more generally, plant ABC transporters. To our knowledge, only one plant PDR transporter has been purified and characterized. The full size ABCG NpPDR5 from Nicotiana plumbaginifolia was expressed in Nicotiana tabacum Bright Yellow-2 (BY-2) suspension cells. The purified transporter displayed basal ATPase activity in the absence of substrates, which are still unknown (33).

NtPDR1 from N. tabacum was first identified in N. tabacum BY-2 cells (34) as a gene activated by the INF1 elicitin, a molecule triggering the biotic stress response (35). In BY-2 cells, ³The abbreviations used are: ABC, ATP-binding cassette; NBD, nucleotide-binding domain; TMD, transmembrane domain; DDM, dodecyl-β-D-maltoside; PDR, pleiotropic drug resistance; MNG, decyl maltose neopentyl glycol; TEV, tobacco etch virus; Nt, N. tabacum; Np, N. plumbaginifolia; BY-2, Bright Yellow-2; Nb, N. benthamiana; Ni-NTA, nickel-nitrilotriacetic acid; GUS, β-glucuronidase.

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1 Both authors contributed equally to this work.

2 To whom correspondence should be addressed. Tel.: 3210473621; Fax 3210473872; E-mail: marc.boutry@uclouvain.be.
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both the NtPDR1 transcript and protein levels increased upon incubation with methyl jasmonate, a stress-response hormone (25, 35). This induction can be related to plant defense upon infection by pathogens (36). In the plant, NtPDR1 is constitutively expressed in the root epidermis as well as in the long glandular trichomes. These trichomes synthesize and secrete large amounts of antimicrobial cyclic diterpenes that constitute a first and constitutive line of plant defense against pathogens (37). Although these diterpenes freely diffuse through membranes, active transport is necessary to keep the concentration of those toxic compounds as low as possible within the plant cells that synthesize them. Indeed, BY-2 cell lines overexpressing NtPDR1 displayed a stronger tolerance to cyclic diterpenes, and transport assays pointed to the diterpenes sclareol and cembrene as putative substrates (25). NbABCG1, the NtPDR1 ortholog, and NbABCG2, a closely related gene, in the species Nicotiana benthamiana have been shown to be involved in induced defense against the pathogen Phytophthora infestans (28). Plants silenced for both NbABCG1 and NbABCG2 were shown to be more sensitive to the cytotoxic effect of the sesquiterpene capsidiol. Moreover, the secreted/intracellular ratio of capsidiol was reduced, suggesting that NbABCG1/2 might transport capsidiol. Preinvasive defense was also affected in the silenced lines, and one hypothesis is that NbABCG1/2 may also transport diterpenes involved in this type of defense (28). However, direct evidence of the interaction between either NtPDR1 or NbABCG1/2 and sesqui- and/or diterpenes is still lacking to clarify the roles of the transporters.

In this study, we describe the overexpression of NtPDR1 in N. tabacum BY-2 cells, its purification, and its reconstitution into liposomes. We demonstrate that NtPDR1, and not NpPDR5, a transporter involved in plant defense against insects used as a control, had its ATPase activity stimulated by the sesqui- and/or diterpenes is still lacking to clarify the roles of the transporters.

Results
Expression of NtPDR1 in BY-2 suspension cells

The coding sequence of NtPDR1 (25) was fused to an N-terminal purification tag with the following scheme: His10-StrepII tag-tobacco etch virus protease cleavage site (supplemental Fig. S1). The recombinant gene was inserted in an expression cassette under the control of a strong constitutive promoter (38) and transferred to the N. tabacum BY-2 cell (34) genome using Agrobacterium tumefaciens-mediated transformation. The gene was inserted together with the nptII kanamycin resistance gene for selection and mcherry, a gene encoding a fluorescent protein, which was used for the selection of highly expressing clones. Transformed cell lines were selected on kanamycin, and 60 lines were kept according to their mCherry fluorescence. Then, the lines were screened for NtPDR1 expression through Western blotting using HisProbe (Thermo Scientific), and the constitutively expressed plasma membrane H+ -ATPase was used as a loading control. The clones displayed expression of the NtPDR1 at different levels (see Fig. 1A for an example) due to the position effect because the integration in the plant genome occurs randomly. It has been shown that every cell from a primary BY-2 transgenic line does not express the transgene in a homogenous manner. This heterogeneity might result from a double transformation event or appear later as a result of gene silencing. Nevertheless, it is possible to increase the homogeneity via subcloning (39). Because of this additional step performed on the elite line (T23), we obtained two subclone lines (SC7 and SC11) displaying significantly greater NtPDR1 expression (Fig. 1B). Scaling up of NtPDR1 production was performed in a 4-liter stirred-tank bioreactor. A fresh weight of about 1 kg of cells was obtained that typically yielded between 2 and 2.5 g (protein) of microsomal fraction.

Solubilization and purification

We screened several detergents for solubilization (supplemental Fig. S2), and n-dodecyl β-D-maltoside (DDM) was found to be the most efficient with a solubilization yield above 90% according to gel quantification. We then followed the purification procedure recently used for NpPDR5 (33) with minor modifications as described under “Experimental procedures.” The purification process included two affinity chromatography steps on Ni-NTA resin, the removal of the His10-StrepII tag by the addition of TEV protease, and a final incubation on Ni-NTA (Fig. 2). The resulting flow-through, which contained NtPDR1, was concentrated by ultrafiltration. A single band corresponding to NtPDR1 was obtained (Fig. 2) without visible contamination or degradation fragments upon SDS-PAGE and Coomassie Blue staining. The purification from 1 g of microsomal fraction yielded about 0.4 mg of purified NtPDR1.

Oligomeric state of NtPDR1

Gel filtration of the purified NtPDR1 was performed to assess the monodispersity and oligomeric state of NtPDR1 (Fig. 3A).
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Figure 2. Purification of NtPDR1. NtPDR1 was solubilized from 800 mg of a microsomal fraction and purified as described under “Experimental procedures.” Twenty microliters of solubilized material (SN), flow-through of the first Ni-NTA column (FT1), elutions of the first and second Ni-NTA columns (E1 and E2), elution of the desalting column (P10), the flow-through and elution of the third column (FT3 and E3), and 5 μl of concentrated flow-through 3 were analyzed by SDS-PAGE and Coomassie Blue staining. The arrow shows at which step the purification tag was cleaved using TEV protease. The band corresponding to NtPDR1.

Seventy percent of NtPDR1 was eluted in a peak at an apparent size of about 240 kDa, which is consistent with the size of a monomeric NtPDR1 (162 kDa) surrounded by the detergent micelle. Smaller peaks of greater size probably correspond to oligomers or aggregates. We further analyzed the main peak of NtPDR1 by negative-stain electron microscopy (Fig. 3B). The raw images of negatively stained proteins showed that solubilized NtPDR1 was monodisperse. To enhance the signal over noise ratio, 2D classification was carried out using more than 5,300 particles (Fig. 3B). The classes displayed in Fig. 3B most likely represent different views and thereby orientations of the sample adsorbed onto an EM grid. Our unbiased reference-free analysis revealed that NtPDR1 was monomeric. Compared with reported analyses of other ABCs transporters, the particles showed recognizable features of ABCs including U or V shapes with, at one end, greater densities likely corresponding to NBDs.

Glycosylation of NtPDR1

Five N-glycosylation sites are predicted for NtPDR1 (Asn-5, -641, -738, -1231, and -1386) of the native predicted enzyme (supplemental Fig. S3). Purified NtPDR1 was proteolyzed either by trypsin or by multienzymatic limited digestions to increase the sequence coverage. The resulting peptides were then analyzed by LC-MS/MS. Peptides covering positions 5, 1231, and 1386 were all detected without any glycosylation. No free or glycosylated peptide covering position 641 was identified. This site is predicted to be localized very close to a membrane-spanning domain (supplemental Fig. S3) and is probably not accessible to the glycosylation machinery or to proteases. In contrast, Asn-738 is localized in an external loop and was found to be glycosylated. The glycosylation profile (Fig. 4) corresponds to a typical plant profile with GnGnXF as the most abundant glycan (supplemental Fig. S4). Deglycosylation was performed using peptide N-glycosidase A to confirm site 738 occupancy. After this step, the deglycosylated peptide (displaying 1 mass unit difference due to the Asn to Asp modification) was observed for site 738, whereas no Asn to Asp modification was observed for other sites, confirming previous results (data not shown). We also tested the glycosylation profile of NpPDR5 purified in a very similar manner as NtPDR1 (33). Two glycosylation sites are predicted, but only free peptides were found covering those positions (data not shown).

Purified NtPDR1 is active

ABC proteins are known to exhibit basal ATPase activity. NtPDR1 was purified in an active form because it was able to hydrolyze ATP (Fig. 5A). In the presence of DDM and asolectin, a mixture of phospholipids, NtPDR1 displays a $K_m$ of 1.14 ± 0.30 mM Mg-ATP and a $V_{max}$ of 21.24 ± 2.39 nmol of P$_i$·min$^{-1}$·mg$^{-1}$. In addition, NtPDR1 basal ATPase activity was inhibited by the well-known ABC inhibitors vanadate, miconazole, and, to a lesser extent, oligomycin A (Fig. 5B).

Reconstitution into liposomes

To assess the activity of NtPDR1 in a lipid environment, purified NtPDR1 was reconstituted in asolectin liposomes. The reconstitution efficiency was assessed by flotation centrifugation on a sucrose gradient. More than 90% of the NtPDR1 signal was found in the top fractions of the gradient (supplemental Fig. S5A), demonstrating that the transporter was reconstituted in liposomes. A protease assay using trypsin demonstrated that the proteoliposomes were well-sealed (supplemental Fig. S5B). Indeed, the antibody against NtPDR1 was obtained using a peptide corresponding to a region present in the second NBD. Upon reconstitution, part of the transporter is expected to be oriented in the membrane with its NBDs inside the liposome. In that case, only partial digestion by trypsin should occur because few cleavage sites are available outside of the vesicles. Indeed, different bands corresponding to partial digestion of NtPDR1 were visible when the proteoliposomes were incubated with trypsin. All these bands were not present when the liposomes were destabilized by Triton X-100. Altogether, these results show that NtPDR1 was well-reconstituted in sealed vesicles. The same approach was used to reconstitute NpPDR5, and similar results were obtained (supplemental Fig. S6). These NpPDR5 proteoliposomes served as a control in ATPase activity tests.

Reconstitution of NtPDR1 increases its basal ATPase activity

Basal ATPase activity of reconstituted NtPDR1 was first assessed in the absence of putative substrates (Fig. 6A). NtPDR1 displayed a $K_m$ of 1.00 ± 0.14 mM ATP, which is in the range of the typical ATP concentration of plant cells (40). The apparent $V_{max}$ was 37.6 ± 2.0 nmol of P$_i$·min$^{-1}$·mg$^{-1}$, a 77% increase compared with the value observed for the solubilized enzyme. It has to be noted that this $V_{max}$ is underestimated because only the proteins with their NBDs at the external side of the membranes have access to ATP. The reconstituted NtPDR1 ATPase activity was almost completely abolished by vanadate (Fig. 6B).

NtPDR1 is specifically activated by defense terpenoids

In vivo toxicity and transport assays (25) supported the hypothesis that NtPDR1 is involved in the transport of cyclic
diterpenes such as sclareol and cembrene but could not discriminate between direct and indirect effects. It is well-known that the ATPase activity of ABC transporters can be stimulated by their substrates. However, this is not the case for all ABC transporters. For example, yeast PDR proteins are not stimulated by their substrates (41). Here, we investigated the potential stimulation of NtPDR1 upon the addition of different terpenoids. NtPDR1 ATPase activity was stimulated by the cyclic diterpenes sclareol (55%) and cembrene (89%) with $K_m$ values of 3.4 $\pm$ 1.4 and 21.3 $\pm$ 6 $\mu$m, respectively (Fig. 7, A and B). Because its N. benthamiana homologs NbPDR1 and NbPDR2 have been proposed to transport capsidiol, the effect of this cyclic sesquiterpene on ATPase activity was also tested on NtPDR1. Capsidiol stimulated NtPDR1 (120%) with a $K_m$ of 41.7 $\pm$ 10.2 $\mu$m (Fig. 7C). In contrast, neither the monoterpene eucalyptol nor the linear diterpene geranylgeraniol stimulated NtPDR1 activity (Fig. 7D). In addition, none of the NtPDR1-stimulating compounds increased the NpPDR5 basal ATPase activity (Fig. 7E), indicating that the stimulation of NtPDR1 activity was specific.

NtPDR1 expression is induced in leaf tissues by methyl jasmonate

In leaf tissues, NtPDR1 is constitutively expressed in the head cells of long glandular trichomes where the defense diterpenes are synthesized, secreted, and take part in the constitutive plant defense (25). The stimulation of NtPDR1 by capsidiol raises the hypothesis that this transporter is also involved in induced defense in leaf tissues because capsidiol was shown to be produced upon pathogen attack (42). We thus tested whether the $\beta$-glucuronidase (GUS) reporter gene controlled by the NtPDR1 transcription promoter was activated in the presence of methyl jasmonate, a typical hormone involved in plant defense. Although GUS activity was confined to trichomes in
control conditions, leaf incubation with methyl jasmonate induced GUS expression in the whole leaf tissue (Fig. 8).

**Discussion**

Despite their essential roles, plant ABC transporters have been poorly characterized at the enzymatic level. One explanation for this lack of data resides in the difficulty of achieving heterologous expression in organisms such as bacteria or yeast (9). *N. tabacum* BY-2 cells have been used to functionally characterize different ABC transporters such as *Arabidopsis* ABCB4 (43, 44) and ABCG40 (19), MtABCG10 from *Medicago truncatula* (45), and PhABCG1 from *Petunia hybrida* (46). For the latter as well as NpPDR5 (33), purification of the protein was also carried out. BY-2 cells offer among others the advantages of proper membrane composition and glycosylation. The lipid composition of the membranes has an impact on protein folding, structure, and function (47, 48). In plants, the plasma membrane is particularly rich in sterols and polar sphingolipids (49, 50), which might be crucial for membrane protein stability. It would be interesting to test these lipids during reconstitution in liposomes.

Size-exclusion chromatography and single-particle analysis by electron microscopy of NtPDR1 showed that solubilized NtPDR1 is a monomer. The length of ∼9 nm is similar to lengths of other ABCG transporters including the plant

![Figure 5. Basal ATPase activity of purified NtPDR1.](image)

- **A**, the ATPase activity of purified NtPDR1 was measured in the presence of the indicated Mg-ATP concentration. Free Mg$^{2+}$ was kept at 1 mM. $K_m$ and $V_{max}$ are 1.14 ± 0.30 mM Mg-ATP and 21.24 ± 2.39 nmol-min$^{-1}$-mg$^{-1}$, respectively. $B$, the specific activity was assessed in the presence and absence of 50 μM vanadate, 10 μM miconazole, or 100 μM oligomycin A. Error bars represent S.D.

- **B**, the specific activity was assessed in the presence and absence of 50 μM vanadate. Error bars represent S.D.

![Figure 6. Basal ATPase activity of reconstituted NtPDR1.](image)

- **A**, the ATPase activity of reconstituted NtPDR1 (4 μg) was measured in the presence of the indicated Mg-ATP concentration. Free Mg$^{2+}$ was kept at 1 mM. $K_m$ and $V_{max}$ are 1.04 ± 0.14 mM Mg-ATP and 37.64 ± 2.01 nmol-min$^{-1}$-mg$^{-1}$, respectively. $B$, the specific activity was assessed in the presence and absence of 50 μM vanadate. Error bars represent S.D.
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Figure 7. ATPase activity of reconstituted NtPDR1 (A–D) or NpPDR5 (E) activity in the presence of different terpenoids. A, sclareol. $K_m = 3.36 \pm 1.36 \mu M$, and the maximal stimulation is $54 \pm 4\%$. B, cembrene. $K_m = 21.27 \pm 6.02 \mu M$, and the maximal stimulation is $89 \pm 7\%$. C, capsidiol. $K_m = 41.47 \pm 10.19 \mu M$, and the maximal stimulation is $125 \pm 12\%$. D, geranylgeraniol and eucalyptol at $50 \mu M$. E, relative activity of reconstituted NpPDR5 in the presence or absence of $50 \mu M$ cembrene, sclareol, or capsidiol. Error bars represent S.D.

Figure 8. Stimulation of NtPDR1:GUS expression by methyl jasmonate. Leaf discs of N. tabacum plants expressing the GUS reporter gene under the control of the NtPDR1 promoter (25) were incubated with water or with $250 \mu M$ methyl jasmonate, and the in situ GUS activity was monitored 24 h later. A, control leaf disk incubated with water. The left panel shows a top view of a leaf disk. GUS expression is limited to trichomes. The right panel shows a trichome in detail. B, leaf disk incubated with $250 \mu M$ methyl jasmonate. Expression was observed in the whole leaf tissue. Arrows indicate examples of stained trichomes head cells.
NpPDR5 (33) as well as the mammalian ABCG2 (Protein Data Bank code 5NJ3) and ABCG5/G8 (Protein Data Bank code 5DO7). This is shorter than the 12–13 nm reported for several ABCs of other subfamilies such as ABCA1 (Protein Data Bank code 5XJY), ABCB1 (Protein Data Bank code 5KOY), and ABCC1 (Protein Data Bank code 5U9J). This is related to shorter intracytoplasmic helices in ABCG transporters, suggesting a similar topology for NtPDR1.

Glycosylation is a common co- and post-translational modification of proteins moving through the secretory pathway and plays a crucial role in protein folding, trafficking, stability, and activity (51). Because glycosylation is not strictly conserved between plants and other eukaryotes (52), glycosylation defects might explain the difficulty in overexpressing some plant ABC transporters in these organisms. Interestingly, glycosylation has also been shown to affect the transport activity of the human ABCC4 (MRP4) transporter (53). The authors demonstrated that the absence of glycosylation was not deleterious for ABCC4 localization but affected its transport activity in a substrate-specific manner. Because our analysis demonstrated that NtPDR1 is indeed glycosylated, we believe it was safer to use a homologous expression system to ensure correct glycosylation. The single glycosylation site is localized within a ~60-residue external loop in the first TMD. Curiously, the corresponding predicted site in the second TMD was not found glycosylated. Glycosylation asymmetry was also found in the external loops of the human heterodimer ABCG5/ABCG8 for which two and one glycosylation sites were identified, respectively (54). In contrast, no glycosylation was found for NpPDR5.

The cyclic diterpenes cembrene and sclareol have been identified as putative substrates of NtPDR1. However, direct evidence of an interaction between these compounds and the protein was lacking. The same is true for any other plant ABC transporter. We thus cannot exclude that transport that was observed at the cell level (25) was an indirect consequence of NtPDR1 expression. To solve this question, we first tried to monitor the transport of a radiolabeled analog of sclareol (55) that was shown to be transported in N. tabacum BY-2 cells overexpressing NtPDR1 (25). To do so, we followed the accumulation of the sclareol analog inside proteoliposomes, but we could not obtain reliable transport data. This illustrates the difficulty in showing direct transport of hydrophobic substrates by in vitro assays. For instance, close to half of the human ABC transporters are thought to facilitate lipids or lipid-related compounds. In many cases, the substrates have been indirectly identified by monitoring accumulation of compounds in humans with specific diseases or in knock-out mice (56). The lack of evident transport is probably due to the high hydrophobicity of the substrates, which readily diffuse and accumulate inside the membrane. We indeed observed a fast and huge background of the diterpenes associated with the liposomes in the absence of ATP. Even in the case of active transport toward the liposome lumen, the compounds will likely re-enter the membrane by passive diffusion. In planta, lipid transfer proteins have been shown to be involved in the transport of hydrophobic diterpenes and sesquiterpenes from the membrane vicinity to lipid droplets to avoid re-entering in the plasma membrane (57, 58).

For some ABC transporters, an alternative to transport assays is the stimulation of the ATPase activity by the substrate (e.g. Refs. 59 and 60). Indeed, the addition of the diterpenes cembrene and sclareol as well as the sesquiterpene capsidiol did stimulate the ATPase activity with $K_m$ values in the micromolar range, indicating direct interaction between NtPDR1 and these substrates. None of these compounds were found to have an impact on NpPDR5 basal activity, indicating that the interaction with NtPDR1 was specific. Neither the monoterpene eucalyptol nor the linear diterpene geranylgeraniol stimulated the NtPDR1 ATPase activity. This is consistent with in vivo data (25).

The observation that the diterpenes sclareol and cembrene as well as the sesquiterpene capsidiol stimulate NtPDR1 ATPase activity is very interesting because these two types of compounds are involved in different steps of plant defense. Diterpenes are constitutively synthesized in, and secreted from, the head cells of glandular trichomes (61) where NtPDR1 is constitutively expressed (25). Capsidiol and other sesquiterpenes are synthesized as a response to pathogen attack in tissues next to the site of pathogen invasion (62, 63). This is in agreement with the induced expression of the GUS reporter under the control of the NtPDR1 promoter in whole leaf tissues upon addition of the defense hormone methyl jasmonate, whereas expression was confined to the trichome secretory cells in the absence of this hormone. NtPDR1 is thus involved in transporting compounds involved in both constitutive (diterpenes) and induced (sesquiterpenes) defenses.

An important question that remains to be addressed is the subcellular localization of NtPDR1. This transporter has been localized to the plasma membrane in BY-2 cells, and this is in agreement with its transport activity (25) as well with the fact that diterpenes are secreted from the trichome head cells (64). However, one hypothesis might be that NtPDR1 is localized to internal vesicles in which substrates are transported before being released out of the cell upon fusion of the vesicles with the plasma membrane. The advantage would be to considerably increase the secretion capacity, which would be in agreement with the massive secretion of diterpenes from tobacco trichomes. Indeed, diterpenes together with sucrose esters also secreted from the trichomes might represent more than 10% of the dry weight of the whole tobacco leaf (64). Future work should therefore focus on the subcellular localization of NtPDR1 in trichomes but also in other leaf tissues upon pathogen attack.

In conclusion, this work provides direct evidence of the interaction between a plant ABC transporter and its substrates. We demonstrated in vitro that the activity of purified NtPDR1 is specifically stimulated by both cyclic di- and sesquiterpenes involved in constitutive and induced plant defenses. We also provide another example of an ABC protein for which N. tabacum BY-2 cells are a convenient host for protein expression, purification, reconstitution, and biochemical characterization. Our pipeline could be used for other plant ABC proteins for which very little is known at the biochemical level (65).
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Experimental procedures

Plant materials

*N. tabacum* L. cv. BY-2 suspension cells (34) were cultured in MS medium (0.44% (w/v) Murashige and Skoog salts (MP Biomedicals), 0.02% (w/v) KH$_2$PO$_4$, 5.10−3% (w/v) myoinositol, 5·10−3% (w/v) thiamine, and 2·10−5% (w/v) 2,4-dichlorophenoxyacetic acid, pH 5.8 (KOH)) supplemented with 100 μg/ml kanamycin. The cultures were grown in the dark at 25 °C and shaken at 90 rpm. The lines were subcultured (diluted 10 times in fresh medium) every week. They were shaken at 90 rpm. The lines were subcultured (diluted 10 times in fresh medium) every week.

Genetic engineering and BY-2 cell transformation

The isolation of the microsomal fraction was performed as described (33). Briefly, the cells from a 6-day-old bioreactor were harvested by filtration through Miracloth (Calbiochem) and washed with ice-cold homogenization buffer (50 mM Tris-HCl (pH 8.0), 2 mM EDTA, 0.25 M sorbitol, 20 mM DTT, and protease inhibitor mixture (1 mM PMSF and 1·10−4% (w/v) each of leupeptin, pepstatin, aprotinin, and chymostatin)). The cells were resuspended in 1 volume of homogenization buffer and ground twice with glass beads (0.8–1-mm diameter; VWR) in a DYNO-MILL Type KDL. The homogenate was centrifuged at 3,700 × g (Beckman-Coulter Avanti J-E centrifuge and JLA-9.1000 rotor) for 5 min, and the supernatant was centrifuged at 6,000 × g for 5 min (same rotor) to remove the cell debris and the heavy subcellular fraction. The supernatant was centrifuged at 100,000 × g for 40 min to pellet the membranes (microsomal fraction). The pellet was suspended in suspension buffer (330 mM sucrose, 3 mM KH$_2$PO$_4$ (pH 7.8), and 3 mM KCl). Protein concentration was assessed using the Bradford method (69) with bovine serum albumin as the standard.

Solubilization assays

The solubilization assay was adapted from Toussaint et al. (33). Microsomal proteins were diluted at 2.5 mg/ml in the solubilization buffer (20 mM Tris-HCl (pH 7.5), 50 mM NaCl, 20 mM imidazole, 10% (w/v) glycerol, and protease inhibitor mixture) supplemented with 1% detergent (DDM, decyl maltose neopentyl glycol (MNG), octyl β-glucoside, Triton X-100, tetradecyl octaethylene glycol ether (C$_{14}$E$_{8}$), CHAPS, lauryldimethyl amine oxide, or SDS). All samples were incubated for 30 min under agitation on a wheel at 4 °C. After centrifugation for 15 min at 100,000 × g and 4 °C (Beckman Optima MAX ultracentrifuge and TLA-55 rotor), the supernatant was kept, and the pellet was resuspended in the same volume of solubilization buffer by sonication (ultrasonic processor Sonics Vibra-Cell 75022; amplitude, 30). The supernatants and pellets were analyzed by SDS-PAGE and Western blotting.

Purification of NtPDR1 and NpPDR5

The purification was performed as described (33). Briefly, the proteins were solubilized in DDM and purified twice by nickel-affinity chromatography. Then, the His-StrepII tag was removed, the proteins were incubated a third time with Ni-NTA, and purified NtPDR1 or NpPDR5 were recovered in the flow-through and concentrated by centrifugal ultrafiltration. One difference between NtPDR1 and NpPDR5 purification was that the imidazole concentration used for the incubation on the third Ni-NTA column was 10 mM (NtPDR1) or 15 mM (NpPDR5) due to the greater tendency of NtPDR5 to bind to Ni-NTA with its endogenous histidine residues.

Size-exclusion chromatography

Purified proteins were loaded on a Superdex 200 10/300 GL size-exclusion chromatography column (GE Healthcare) linked to the ÅKTA Explorer system. The column was pre-equilibrated with 20 mM Tris-HCl (pH 7.5), 50 mM NaCl, and 10%
(w/v) glycerol supplemented with 0.03% MNG. The elution was carried out at 0.45 ml/min, and the absorbance at 280 nm was recorded.

**Reconstitution of NtPDR1 and NpPDR5**

The reconstitution protocol was adapted from Geertsm et al. (70). Fifty mg of asolectin from soybean (Sigma, reference number 11145) were dissolved in chloroform and dried under N\(_2\). The lipidic film was dissolved in diethyl ether and dried again under N\(_2\) to remove traces of chloroform. The lipid film was suspended in 1 ml of reconstitution buffer (20 mM Tris-HCl (pH 7.5) and 50 mM NaCl) and sonicated on ice for 15 min (ultrasonic processor Sonics Vibra-Cell 75022; amplitude, 30; pulses of 6 s). The suspension was submitted to five cycles of flash freezing in liquid nitrogen and slow thawing at room temperature, and the liposomes were flash frozen and stored at −80 °C. When needed, 1 ml of liposomes at 50 mg/ml was slowly thawed at room temperature and extruded 11 times through a 400-nm polycarbonate filter (Whatman) using a Mini Extruder (Avanti Polar Lipids). The suspension was diluted to 4 mg/ml asolectin with reconstitution buffer (total volume, 12.5 ml) and destabilized by 10 successive additions of 3.75 mg of Triton X-100 (final concentration, 3 mg/ml) with 1 min of gentle agitation in between. Purified NtPDR1 or NpPDR5 was added to 3 ml of destabilized liposomes at a protein/lipid ratio of 1:100 (w/w) and agitated for 15 min at 4 °C. To remove detergents, the suspension was agitated for 16 h at 4 °C with successive additions of 40 mg/ml SM-2 Bio-Beads (Bio-Rad) after 0, 30, 90, and 840 min. The suspension was filtered on a Poly-Prep column (Bio-Rad), and the beads were washed with 2 ml of reconstitution buffer. The proteoliposomes were collected by centrifugation for 20 min at 267,000 \(\times\) g (Beckman Optima MAX ultracentrifuge and MLA-80 rotor). The proteoliposomes were suspended in reconstitution buffer to 10 mg/ml lipids and used for further analysis.

**Protease assay**

Thirty microliters of reconstitution buffer supplemented or not with 1 \(\mu\)g of trypsin (Sigma), 1.6% Triton X-100 (Sigma), and 20 \(\mu\)g of trypsin inhibitor (Sigma) were incubated for 10 min on ice. Fifteen microliters of proteoliposomes (1.5 \(\mu\)g of proteins) were added, and the solution was incubated for 20 min at 37 °C. The reaction was stopped by the addition of 20 \(\mu\)g of trypsin inhibitor and incubation for 10 min on ice, and then the samples were analyzed by immunoblotting.

**Flotation assay**

The sucrose flotation assay was adapted from Pernstich et al. (71). Proteoliposomes (5 \(\mu\)g of reconstituted proteins) in 150 \(\mu\)l of reconstitution buffer supplemented with 50% (w/v) sucrose were layered at the bottom of a 2-ml centrifugation tube (Beckman Optima MAX ultracentrifuge and TLS-55 rotor) and overlaid with 1.75 ml of reconstitution buffer supplemented with 40% (w/v) sucrose and 200 \(\mu\)l of reconstitution buffer. The tube was centrifuged for 12 h at 200,000 \(\times\) g. Aliquots of 300 \(\mu\)l were recovered, and 30 \(\mu\)l of each aliquot were analyzed by immunoblotting. Proteoliposomes (0.5 \(\mu\)g of reconstituted proteins) were used as the input control.

**ATPase activity**

For solubilized NtPDR1, the ATPase assay was performed as described (33) in 50 mM Tris-HCl (pH 7.5), 50 mM NaCl, 60 \(\mu\)g/ml pyruvate kinase (Roche Applied Science), 32 \(\mu\)g/ml lactate dehydrogenase (Roche Applied Science), 0.1% (w/v) asolectin and DDM, 20 mM KNO\(_3\), 0.2 mM (NH\(_4\))\(_2\)MoO\(_4\), 10 mM NaN\(_3\), 4 mM phosphoenol pyruvate, 0.4 mM NADH, 4 mM ATP, and 5 mM MgCl\(_2\) except when stated otherwise. The ATP hydrolytic activity of reconstituted NtPDR1 or NpPDR5 was assessed according to Galán et al. (60). The reaction mixture (100 \(\mu\)l) contained 20 mM Tris-HCl (pH 7.5), 50 mM NaCl, 60 \(\mu\)g/ml pyruvate kinase, 32 \(\mu\)g/ml lactate dehydrogenase, 4 mM phosphoenol pyruvate, 0.4 mM NADH, 20 mM KNO\(_3\), 0.2 mM (NH\(_4\))\(_2\)MoO\(_4\), 10 mM NaN\(_3\), 4 mM ATP, and 5 mM MgCl\(_2\). The reaction was started by the addition of 40 \(\mu\)l of proteoliposomes (4 \(\mu\)g of proteins), and the absorbance at 340 nm was measured at 37 °C using a microplate reader (SPECTROstar Nano, BMG Labtech). All activity data represent the mean ± S.D. of at least three replicates.

**Electron microscopy and single-particle analysis**

The purified proteins were diluted into a detergent-containing buffer (20 mM Tris (pH 7.5), 50 mM NaCl, and 0.03% MNG) at a concentration of 0.02 mg/ml. The sample was adsorbed on a carbon-coated glow-discharged EM grid (300 mesh; Electron Microscopy Science) for 30 s before being stained with a 2% uranyl formate solution. The data were collected using a LaB6 G2 Tecnai (FEI, Eindhoven, The Netherlands) electron microscope operated at an acceleration voltage of 200 kV. The images were acquired using a 4,000 \(\times\) 4,000 CMOS F416 camera (Tietz Video and Image Processing Systems, Germany) in an automated manner using the EMTools software suite (Tietz Video and Image Processing Systems) with defocus varying from −1 to −3 \(\mu\)m. The images were collected at a magnification of 50,000, a dose of 10 electrons/Å\(^2\), and a pixel size of 2.13 Å.

From 28 images, 5,301 boxes of 135 \(\times\) 135 pixels were manually windowed out using the boxer EM software suite (72). The picked particles were normalized against the background and appended into a single SPIDER file (73). A first round of reference-free alignment and classification was performed using SPIDER to obtain 40 unbiased classes. Those classes were then used as references for a multireference alignment followed by a classification that was iterated four times until no change was observed. Eventually 129 classes were obtained.

**Glycosylation profiling**

Purified NtPDR1 was reduced, alkylated, and purified as described (74). The protein was then resuspended in proteolysis buffer supplemented with 0.1% (w/w) Rapigest (Waters). Two proteolysis methods were performed: trypsin digestion as described previously (74) and multienzymatic (using a trypsin, chymotrypsin, and Glu-C mixture) limited proteolytic digestions (for 2 h at 37 °C). An aliquot of the resulting peptides was then subjected to deglycosylation at 37 °C for 24 h using peptide N-glycosidase A (0.2 milliunit/100 \(\mu\)g; Roche Applied Science) in a citrate/phosphate buffer adjusted to pH 5 and supplemented with 0.005% (w/w) Rapigest.
Peptides were then analyzed by LC-MS/MS. Briefly, peptides were separated by 1D reverse-phase chromatography (M-Class, HSS T3 C18 column, Waters) using a 70-min gradient (5–40% acetonitrile/water and 0.1% formic acid) at 600 nl/min. Eluted peptides were then analyzed by nanoelectrospray ionization-MS on a quadrupole Orbitrap hybrid mass spectrometer (Q Exactive Plus, Thermo Fisher Scientific) operated in positive mode. Survey scans were recorded at 70,000 resolving power (full width at half-maximum) from 400 to 1,750 m/z. MS/MS spectra were recorded using a “data-dependent analysis” scheme in which the 12 most abundant ions observed in MS mode were selected for fragmentation experimentation (normalized collision energy set at 28).

MS/MS spectra were searched for typical oxonium ions (m/z 204.09 and 366.14) indicating the presence of glycopeptides. For all candidates, the presence of the Y1 ion (fragment containing peptide + N-acetylhexosamine) was checked to confirm the glycosylation site. The composition of the assigned spectra was determined using Glycomod software (available at [http://web.expasy.org/glycomod/])4 and manual interpretation of MS/MS spectra. Relative abundances were then determined by integrating MS1 signals using Skyline 3.0.1. Alternatively, MS/MS spectra were subjected to a database search using the Mascot 2.2 search engine and the TrEMBL database restricted by integrating MS/MS spectra. Relative abundances were then determined of MS/MS spectra. Relative abundances were then determined by integrating MS1 signals using Skyline 3.0.1. Alternatively, MS/MS spectra were subjected to a database search using the Mascot 2.2 search engine and the TrEMBL database restricted by integrating MS/MS spectra. Relative abundances were then determined of MS/MS spectra. Relative abundances were then determined by integrating MS1 signals using Skyline 3.0.1. Alternatively, MS/MS spectra were subjected to a database search using the Mascot 2.2 search engine and the TrEMBL database restricted by integrating MS/MS spectra. Relative abundances were then determined of MS/MS spectra. Relative abundances were then determined by integrating MS1 signals using Skyline 3.0.1. Alternatively, MS/MS spectra were subjected to a database search using the Mascot 2.2 search engine and the TrEMBL database restricted by integrating MS/MS spectra. Relative abundances were then determined of MS/MS spectra. Relative abundances were then determined by integrating MS1 signals using Skyline 3.0.1. Alternatively, MS/MS spectra were subjected to a database search using the Mascot 2.2 search engine and the TrEMBL database restricted by integrating MS/MS spectra. Relative abundances were then determined of MS/MS spectra. Relative abundances were then determined by integrating MS1 signals using Skyline 3.0.1. Alternatively, MS/MS spectra were subjected to a database search using the Mascot 2.2 search engine and the TrEMBL database restricted by integrating MS/MS spectra. Relative abundances were then determined of MS/MS spectra.

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GUS reporter activity

Leaf discs of N. tabacum plants expressing the GUS reporter gene under the control of the NtPDR1 promoter (25) were submerged in water or 250 μM methyl jasmonate for 24 h, and GUS reporter activity was measured as described (75).

Author contributions—M. B. conceived and coordinated the study. B. P. and F. T. performed most of the experiments. A. B. and D. L. performed the electron microscopy analysis. N. S. and E. D. P. performed the glycosylation analysis. All authors reviewed the results and wrote the paper.

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