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**Research Article**

**A novel plant enzyme with dual activity: an atypical  
Nudix hydrolase and a dipeptidyl peptidase III**

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## Abstract

In a search for plant homologues of dipeptidyl peptidase III (DPP III) family, we found a predicted protein from the moss *Physcomitrella patens* (UniProt entry: A9TLP4), which shared 61% sequence identity with the *Arabidopsis thaliana* uncharacterized protein, designated Nudix hydrolase 3. Both proteins contained all conserved regions of the DPP III family, but instead of the characteristic hexapeptide HEXXGH zinc-binding motif, they possessed a pentapeptide HEXXH, and at the N-terminus, a Nudix box, a hallmark of Nudix hydrolases, known to act upon a variety of nucleoside diphosphate derivatives. To investigate their biochemical properties, we expressed heterologously and purified *Physcomitrella* (PpND) and *Arabidopsis* (AtND) protein. Both hydrolyzed, with comparable catalytic efficiency, the isopentenyl diphosphate (IPP), a universal precursor for the biosynthesis of isoprenoid compounds. In addition, PpND dephosphorylated four purine nucleotides (ADP, dGDP, dGTP, and 8-oxo-dATP) with strong preference for oxidized dATP. Furthermore, PpND and AtND showed DPP III activity against dipeptidyl-2-arylamide substrates, which they cleaved with different specificity. This is the first report of a dual activity enzyme, highly conserved in land plants, which catalyses the hydrolysis of a peptide bond and of a phosphate bond, acting both as a dipeptidyl peptidase III and an atypical Nudix hydrolase.

**Keywords:** *Arabidopsis thaliana*; enzyme kinetics; metalloprotease; plant biochemistry; *Physcomitrella patens*; substrate specificity.

## Introduction

The peptidase family M49 (dipeptidyl peptidase III family) was first distinguished among metallopeptidases based on the unique zinc binding motif, the hexapeptide HEXXGH (Fukasawa *et al.*, 1998; Chen and Barrett, 2004), and by the presence of additional four conserved regions (consensus sequences) in the primary structures of its members (Abramić *et al.*, 2004). All five evolutionary conserved regions are situated in the C-terminal half of the dipeptidyl peptidase III (DPP III) molecule (Abramić *et al.*, 2004). Determination of the first three dimensional structure, from the yeast DPP III, revealed a two-domain protein with a novel fold (Baral *et al.*, 2008), as an additional feature of the family M49.

The biochemical properties of this zinc-peptidase isolated from various (mostly mammalian) organisms, have been extensively characterized, as have been structure-activity studies of heterologously (in *Escherichia coli*) expressed enzymes from human, rat, bacterial and yeast origin (Fukasawa *et al.*, 1998; Fukasawa *et al.*, 1999; Abramić *et al.*, 2000; Chen and Barrett 2004; Salopek-Sondi *et al.*, 2008; Špoljarić *et al.*, 2009; Jajčanin-Jozić *et al.*, 2010; Karačić *et al.*, 2012; Vukelić *et al.*, 2012; Abramić *et al.*, 2015). In contrast to biochemistry, molecular enzymology, and structural biology (Baral *et al.*, 2008; Bezerra *et al.*, 2012) knowledge of peptidases belonging to the DPP III family, their physiological roles have been much less elucidated. As DPP III is a cytosolic enzyme, broadly distributed in cells and tissues, it is assumed to participate in the final stages of intracellular protein catabolism (Abramić *et al.*, 1988). In addition, several studies indicated a role for DPP III in the mammalian pain-modulatory system (Chiba *et al.*, 2003; Baršun *et al.*, 2007), and its participation in the endogenous defense against oxidative stress is well supported (Liu *et al.*, 2007). It has been shown that DPP III is an activator in the Keap1-Nrf2 signaling pathway (Hast *et al.*, 2013), which is the major regulator of cytoprotective responses to oxidative and electrophilic stress (Kansanen *et al.*, 2013).

In search for plant homologues of the M49 family, we have found a predicted protein of 770 amino acids from the moss *Physcomitrella patens* (UniProtKB code: A9TLP4), and an *Arabidopsis thaliana* uncharacterized protein of 772 amino acids, designated Nudix hydrolase 3 in UniProtKB (entry code Q8L831; gene: NUDT3). A multiple sequence alignment revealed that *Physcomitrella* protein A9TLP4, and *A. thaliana* protein Q8L831 contain all conserved sequence regions of the M49 family with one exception: instead of the

characteristic hexapeptide zinc-binding motif HEXXGH, they possess the pentapeptide HEXXH. In addition, near their N-termini both plant proteins contained a variant of the «Nudix box» motif (Figure 1), a characteristics of the Nudix hydrolase superfamily.

The Nudix superfamily is widespread among eukaryotes, bacteria, archaea, and viruses and consists mainly of pyrophosphohydrolases that act upon substrates of general structure nucleoside diphosphates linked to some moiety X to yield nucleoside monophosphate plus phosphate-X (Bessman *et al.*, 1996; McLennan 2006; Xu *et al.*, 2006). These enzymes are characterized by a conserved sequence motif GX<sub>3</sub>EX<sub>7</sub>REUXEEXGU (Nudix box), where U represents a bulky hydrophobic amino acid such as Ile, Leu, or Val and X is any amino acid. Their substrates include (deoxy)ribonucleoside diphosphates and intact and oxidatively damaged nucleoside triphosphates, nucleotide sugars, dinucleotide coenzymes, dinucleoside polyphosphates, and capped RNA (Mildvan *et al.*, 2005). The Nudix box forms a loop- $\alpha$  helix-loop structural motif which functions as catalytic site (Mildvan *et al.*, 2005). Additional sequences associated with this motif influence the substrate specificity of the enzyme (Yoshimura and Shigeoka 2015).

A large number of Nudix hydrolases from various organisms such as bacteria, yeast, algae, nematodes, plants and vertebrates have been identified and characterized (Dunn *et al.*, 1999; McLennan 2006; Ogawa *et al.*, 2008; Yoshimura and Shigeoka 2015). The prototype member of the Nudix superfamily is the MutT pyrophosphohydrolase from *E. coli*, a monomeric protein of 129 amino acids. Its three dimensional structure has been determined, and its catalytic mechanism studied in detail (Mildvan *et al.*, 2005). It has been proposed that the biological role of MutT is to remove mutagenic nucleotides, since its best substrate *in vitro* is 8-oxo-dGTP (Mildvan *et al.*, 2005).

Not only Nudix hydrolases, but also the isopentenyl diphosphate isomerases (IDIs) share the Nudix motif (Yoshimura and Shigeoka 2015). IDI is an metalloenzyme that catalyzes interconversion of isopentenyl diphosphate and dimethylallyl diphosphate (Bonanno *et al.*, 2001). Phylogenetic analysis revealed two distinct families of IDI, IDI1 and IDI2 (Berthelot et al 2012). IDI1s, found from bacteria to human, are members of the Nudix superfamily (Berthelot *et al.*, 2012). *E. coli* IDI1 is a single  $\alpha/\beta$  domain protein that resembles *E. coli* MutT (Bonanno *et al.*, 2001).

The specific functions of many Nudix hydrolases have not been elucidated, due to their broad substrate specificities. However, increasing evidence is emerging showing that Nudix enzymes may play protective, regulatory, and signaling roles in metabolism (Mildvan *et al.*, 2005; Yoshimura and Shigeoka 2015).

This is the first report of proteins with both Nudix and DPP III domains. Intrigued by potential for roles in plant stress response and cytoprotection, we undertook the heterologous expression and biochemical characterization of the *P. patens* A9TLP4 and *A. thaliana* Q8L831 proteins. We hypothesized that both proteins have dipeptidyl peptidase III activity, due to the presence of evolutionary conserved regions of the M49 family in their primary structure (Figure 1). In addition, we investigated whether they could act as a Nudix enzyme as well.

## Results

### Search for plant homologues of DPP III family

Plant dipeptidyl peptidases III have been recognized only recently as data from plant genomes sequencing projects have become publicly available. Among the first such hits by BLAST similarity search, we found the homologue of DPP III family in the moss *Physcomitrella patens* subsp. *patens*, which is used as a model organism for studies on plant physiology, development and evolution (Prigge and Bezanilla 2010, Müller *et al.*, 2015). Mosses share fundamental physiological processes with higher plants, although they diverged early in evolution. Sequencing of the *P. patens* genome enabled comparisons with other species, which revealed genomic changes accompanying the evolutionary movement to land, including acquisition of genes for tolerating terrestrial stresses (temperature variation and water availability) (Rensing *et al.*, 2008).

The *Physcomitrella patens* homologue - a predicted protein A9TLP4 - showed 30.1% identity (*E*-value: 4e-22) with the *Bacteroides thetaiotaomicron* DPP III (UniProt entry Q8A6N1), and 22.3% identity (*E*-value: 4.7e-7) with the human orthologue of M49 family (UniProt Q9NY33).

We tentatively named the *P. patens* uncharacterized protein A9TLP4, because it contained the Nudix box and DPP III (M49) family consensus sequences, as «*P. patens* Nudix-DPP III»

(PpND) and searched online databases of plant genomes for its homologues. Using Conserved Domain Architecture Retrieval Tool (Geer *et al.*, 2002), we obtained 76 non-redundant sequences containing both Nudix and M49 domain. This type of domain architecture was found only in land plants (not shown).

Among the PpND homologues found, an *Arabidopsis thaliana* protein (UniProt: Q8L831) annotated as «Nudix hydrolase 3», product of *NUDT3* gene, attracted our interest, since its enzyme activity as a Nudix hydrolase was not experimentally confirmed. It had 61.6% identity to PpND (BLAST *E*-value: 0.0). Ogawa *et al.*, (2005), who extensively investigated cytosolic Nudix hydrolases in *A. thaliana*, suggested that the product of AtNUDT3 is not likely a typical Nudix hydrolase enzyme, because its predicted amino acid sequence shows an extremely low similarity to those of other AtNUDTs, and its molecular mass (86.9 kDa) is much higher than those of the enzymes in other organisms.

### **Heterologous production, purification and biochemical characterization of the recombinant *P. patens* PpND and the recombinant *A. thaliana* Nudix hydrolase 3 (AtND)**

The cDNA cloning, expression and purification of the recombinant PpND and AtND was performed as described in the materials and methods section.

The purified His-tagged proteins were analyzed by SDS-PAGE (Figure 2A) and both exhibited a single protein band, whose molecular weight was estimated to be about 88 000, similar to the theoretical molecular mass. Native PAGE analysis of PpND and AtND also showed one protein band (Figure 2B), indicating that each protein exists in solution as a monomer. The isoelectric point as determined by isoelectric focusing was about pH 5.1 for PpND and about pH 5.5 for AtND. Circular dichroism (CD) spectra were obtained for both proteins indicating similar secondary structures (Supplementary Figure 1).

Peptidase activity of purified PpND and AtND was investigated by assaying the hydrolysis of chromogenic substrates of dipeptidyl peptidases, aminopeptidases and endopeptidase. Among them, both plant proteins cleaved only dipeptidyl-2-naphthylamides (Table 1). The best substrate of PpND was Arg-Arg-2-naphthylamide (Arg<sub>2</sub>-2NA), which is known as the preferred synthetic substrate of dipeptidyl peptidase III type of enzyme (Ellis and Nuenke 1967). In addition, five other dipeptidyl-2-naphthylamides were hydrolyzed by both plant enzymes: Ala-Arg-, Pro-Arg-, Phe-Arg-, Ala-Ala- and Gly-Arg-2NA (Table 1).

The biochemical properties of plant enzymes were further characterized using their hydrolytic action on Arg<sub>2</sub>-2NA. The optimal pH for the hydrolysis of this substrate was determined at pH 7.4, for PpND, and at pH 7.9 for AtND. Only Co<sup>2+</sup> ions (among examined Co<sup>2+</sup>, Mg<sup>2+</sup>, Mn<sup>2+</sup> and Zn<sup>2+</sup> ions) activated the PpND and AtND significantly. Both enzymes showed patterns of inhibition typical for DPPs III: strong inhibition with chelating agents (EDTA, o-phenanthroline) and the sulfhydryl reagent *p*-hydroxymercuri-benzoate.

### **Kinetic analyses of PpND and AtND with peptidase substrates**

Screening for peptidase activity indicated the difference in specificity between the moss and *Arabidopsis* enzyme (Table 1). To further investigate the catalytic properties of PpND and AtND hydrolysis of dipeptidyl-2-naphthylamides, which were selected as a good substrates according to the initial results (Table 1), kinetic analyses were performed as described in Materials and methods. The two best substrates of the PpND were Arg<sub>2</sub>-2NA and Ala-Arg-2NA. The kinetic parameters for Arg<sub>2</sub>-2NA hydrolysis catalyzed by *P. patens* enzyme were determined to be:  $K_m = 10.63 \pm 1.72 \mu\text{M}$ ,  $k_{\text{cat}} = 0.822 \pm 0.180 \text{ min}^{-1}$ ,  $k_{\text{cat}}/K_m = 1289 \text{ M}^{-1} \text{ s}^{-1}$ , and for hydrolysis of Ala-Arg-2NA:  $K_m = 17.21 \pm 0.75 \mu\text{M}$ ,  $k_{\text{cat}} = 0.633 \pm 0.157 \text{ min}^{-1}$ ,  $k_{\text{cat}}/K_m = 613 \text{ M}^{-1} \text{ s}^{-1}$ , revealing a slight preference for diarginyl derivative (two-fold higher  $k_{\text{cat}}/K_m$ ). To investigate the AtND specificity, kinetic parameters were determined for four dipeptidyl-2-naphthylamides, selected as a good substrates. As shown in Table 2, AtND preferred Ala-Arg-2NA. The catalytic efficiency (the  $k_{\text{cat}}/K_m$  value) for this substrate was 19-fold and 9-fold higher compared to that for Arg<sub>2</sub>-2NA and Pro-Arg-2NA.

### **Investigation of the predicted isopentenyl diphosphate isomerase (IDI) activity**

In addition to the Nudix hydrolases, Nudix superfamily encompasses other non-hydrolytic enzymes. including the isopentenyl diphosphate (IPP) isomerases (Yoshimura and Shigeoka 2015). Interestingly, Gunawardana *et al.*, (2009) predicted that AtNUDT3 (AtND) is a member of isopentenyl diphosphate isomerase (IDI) family, based on a comprehensive bioinformatic analysis of 25 Nudix hydrolases encoded in the *Arabidopsis* genome. When we aligned the primary structure of PpND and AtND to proteins with confirmed IDI activity (human IDI1, *E. coli* IDI, *Halobacterium* IDI) (Supplementary Figure 2), we noticed that most of the residues necessary for IDI function, inferred from structural studies on human and *E. coli* IDI (Wouters *et al.*, 2003; Zhang *et al.*, 2007), are conserved in plant proteins. However, in PpND and AtND, the signature active-site motif of IDIs, NTCCSH, is replaced



by ISSAGH, in which the catalytic (second) cysteine is substituted by an alanine residue (Ala76 in PpND sequence). Interestingly, IDI activity has been reported for an enzyme from the archaea *Halobacterium* sr. NRC1 (UniProt: Q9HDP40), although the catalytic Cys is replaced by Ala (Supplementary Figure 2) (Hoshino and Egushi 2007). Therefore, both PpND and AtND were tested for IDI activity by a color complementation assay (Wang *et al.*, 2009), as described in Materials and methods. Colonies expressing PpND and AtND were yellow, indicating the absence of isomerase activity (Supplementary Figure 3).

### **Nudix hydrolase activity**

The presence of a Nudix box motif at the N-termini of these two proteins motivated further investigation of their potential Nudix hydrolase activity. For that purpose, we used a continuous fluorescence assay (Xu *et al.* 2013), that employs fluorescently tagged *E. coli* phosphate binding protein (PBP) as a phosphate sensor. We screened PpND and AtND with substrates of Nudix hydrolases, with the common structure nucleoside–diphosphate-X. In addition, we examined isopentenyl diphosphate (IPP) as a substrate for hydrolysis by PpND or AtND. Although these plant enzymes did not have IDI activity, based on their structural similarity (Supplementary Figure 2) we hypothesized that they could bind IPP, in order to hydrolyze it.

### **Substrate screening**

Figure 3 illustrates the screening results of both enzymes against the library of 73 substrates (full list of substrates is given in Supplementary Table 1), along with their  $v_i/[E_0]$  values ( $s^{-1}$ ). White bars indicate the cases in which the activity against a substrate was measured as part of group mixture. Grey bars indicate the screened activity for a substrate that was assayed individually against the enzyme. More details are given in Supplementary Table 2.

The screening results against individual substrates initially found that PpND exhibited the highest activity against IPP ( $v_i/[E_0]$  value:  $0.06 s^{-1}$ ), 8-oxo-dATP ( $0.02 s^{-1}$ ), ADP ( $0.03 s^{-1}$ ), and dGTP ( $0.03 s^{-1}$ ). Notable activity for AtND was only found against IPP ( $0.04 s^{-1}$ ).

### **Michaelis-Menten kinetics**

For all substrates that were found to have notable activity in the screening, kinetic parameters were measured carefully. Supplementary Figure 4 shows the results of PpND and AtND

against these substrates. A summary of the  $k_{\text{cat}}/K_m$  ratios with their standard errors is given in Table 3.

The most reactive substrate of PpND is IPP, with a  $k_{\text{cat}}/K_m = (1.9 \pm 0.16) \times 10^4 \text{ M}^{-1} \text{ s}^{-1}$ , followed by 8-oxo-dATP with a  $k_{\text{cat}}/K_m = (1.1 \pm 0.64) \times 10^4 \text{ M}^{-1} \text{ s}^{-1}$ . Despite showing high activity readings in the screening process, ADP was found to have activity closer to that of dGTP and the less reactive dGDP, on the order of magnitude of  $10^2 \text{ M}^{-1} \text{ s}^{-1}$ . For IPP, the comparison of  $v_i/[E_0]$  values with and without inorganic pyrophosphatase (PPase) found no appreciable change in activity. AtND was recorded to have a  $k_{\text{cat}}/K_m$  of  $(1.8 \pm 0.13) \times 10^4 \text{ M}^{-1} \text{ s}^{-1}$  against IPP. The addition of PPase also did not have an affect in the recorded activity measured.

Additionally, because IPP was a new substrate for this screen panel, IPP was also screened against MutT enzyme, to determine whether it could lead to any artifactual enzyme activity. No such activity was seen (not shown).

### **Identification of isopentenyl monophosphate (IP) as a product of Nudix hydrolase activity of PpND towards IPP**

The activity of PpND toward IPP was additionally examined by LC-MS/MS as described in Materials and methods. Reactions with two concentrations of PpND (60 nM and 600 nM) were performed at 37 °C for 10 min and their products were analyzed. Negative ESI tandem mass detection was run in multiple reaction monitoring (MRM) mode. The values of  $m/z$  of IPP and IP standard parent ions were 244.70 and 164.90, respectively. Collision-induced dissociation of parental ions yielded characteristic fragments of IPP ( $m/z=226.7, 158.5, 79.0$ ) and IP ( $m/z= 148.6, 96.9, 79.0$ ). As shown in Supplementary Figure 5, IP was identified as the product of PpND enzyme reaction toward IPP as a substrate. In addition, the progress of enzyme reaction (Supplementary Figure 5, B and C) was shown in comparison to the control (Supplementary Figure 5A). Based on calculation, the ratio of characteristic ion  $m/z=79.00$ , generated from IPP and those from IP was 4.5:1 (IPP:IP) in enzyme reaction with 60 nM PpND (Supplementary Figure 5B) and 1:4 (IPP:IP) in reaction with 10 times higher concentration of PpND (Supplementary Figure 5C).

## Structure – activity relationships

We observed a significantly lower  $k_{cat}$  value for hydrolysis of the diarginyl-2NA substrate catalyzed by the PpND and AtND, in comparison with human and bacterial orthologues of the DPP III family (Jajčanin-Jozić and Abramić 2013; Vukelić *et al.*, 2012). This was not surprising, since the plant enzymes contain a pentapeptide HECCH, instead of the characteristic active-site hexapeptide HEXXGH. The importance of this hexapeptide motif for full activity of DPP III had been shown previously for the rat orthologue, by mutational analysis (Fukasawa *et al.*, 1999). Substitution of E592 in *Physcomitrella* DPP III, corresponding to the catalytic glutamate in the canonical DPP III motif, HEXXXH, resulted in a complete loss of activity towards Arg<sub>2</sub>-2NA, while the IPP hydrolase activity remained (Figure 4, PpND\_E592A). Two insertion variants were produced to investigate if introduction of the consensus active-site motif into the *P. patens* protein would enhance peptidase activity. A variant with the hexapeptide HECCGH, created by inserting a glycine into the pentapeptide H<sup>591</sup>ECCH<sup>595</sup> motif of wild-type PpND, showed a 29-fold decreased hydrolytic activity towards Arg<sub>2</sub>-2NA compared to the wild-type enzyme. A second mutant with the hexapeptide HECLGH, designed to mimic the active-site motif of the *Bacteroides thetaiotaomicron* DPP III (Vukelić *et al.*, 2012), again resulted in a much less active (57-fold) enzyme. CD measurements showed that the reduced activity is not due to changes in the secondary structure (not shown). Separate expression of the isolated DPP III domain resulted in substantial loss in activity, while expression of the Nudix domain failed to yield soluble Nudix domain protein.

As the first glutamate of the Nudix box motif is essential for catalysis in different Nudix hydrolases (Gunawardana *et al.*, 2009), E92 of the PpND RE<sup>92</sup>LQEE motif was replaced by alanine. The variant PpND\_E92A showed no IPP hydrolase activity, but retained DPP III activity towards Arg<sub>2</sub>-2NA (Figure 4). As A76 from the PpND motif ISSA<sup>76</sup>GH aligns with the catalytic cysteine of the active-site motif of IDIs (Supplementary Figure 2), variant PpND\_A76C was generated to investigate whether a potential isomerase activity could be found. As with wild-type PpND, this was not detected in the color complementation assay (not shown). Finally, the double mutant PpND\_A76C\_E92A was prepared, based on the possibility that IPP isomerase activity might only be detected when IPP hydrolase activity is not present due to the design of the *in vivo* assay. However, this double mutant (PpND\_A76C\_E92A) also did not have IPP isomerase activity (not shown).

## Discussion

Proteolytic enzymes (peptidases) are involved in many aspects of plant physiology, growth and development (Schaller 2004). Considering their numerous and diverse biological functions, it is not surprising that a large number of peptidases has been recognized in plant genomes. According to the *MEROPS* database (<http://merops.sanger.ac.uk/>) (Rawlings *et al.*, 2014), the count of known and putative peptidases in plants *P. patens* and *A. thaliana* is 343 and 745, respectively. However, only a limited number of proteolytic enzymes from these two plants have been characterized by biochemical approaches, none of them being reported as a dipeptidyl peptidase.

In this study, recombinant A9TLP4 protein originating from the moss *Physcomitrella patens* and recombinant NUDT3 protein from the flowering plant *Arabidopsis thaliana* were obtained by heterologous expression in *E. coli*, and their enzymatic properties were examined. Using bioinformatic approaches we have recognized these two proteins as atypical members of the metallopeptidase family M49 (DPP III family), which instead of the hexapeptide HEXXGH active-site zinc-binding motif contain the pentapeptide HECCH. In addition they harbour a variant Nudix box motif, a characteristics of Nudix hydrolases, near their N-termini. Therefore, we called them «Nudix-DPP III» (ND), PpND and AtND.

Experimental characterization of purified proteins revealed biochemical properties (hydrolysis of DPP III specific substrate, Arg<sub>2</sub>-2NA and activation by Co<sup>2+</sup> ions) which classify them as members of the DPP III family. The catalytic efficiency ( $k_{cat}/K_m$ ) of both plant enzymes for Arg<sub>2</sub>-2NA was similar to the yeast DPP III, that has 12-fold higher  $k_{cat}/K_m$  for Arg<sub>2</sub>-2NA hydrolysis (Jajčanin-Jozić *et al.*, 2010), but much lower (3 orders of magnitude), compared to the values determined for human and bacterial DPPs III (Vukelić *et al.*, 2012; Jajčanin-Jozić and Abramić 2013 ). At present, the structural basis for this low efficiency of the plant enzymes is not known, and our data on insertion mutants do not support the explanation that this is due to the shorter active-site motif (HECCH).

The biochemical characterization of DPP III activity revealed significant differences between the two plant enzymes: PpND showed lower pH optimum than AtND (at pH 7.4 vs. pH 7.9). In addition, they differed in their substrate specificity. Among the substrates of X-Arg-2NA series, the *Physcomitrella* enzyme moderately preferred Arg<sub>2</sub>-2NA. In contrast, the

*Arabidopsis* enzyme showed low activity for Arg<sub>2</sub>-2NA, and pronounced preference for Ala-Arg-2NA (Tables 1 and 2).

Screening of PpND and AtND against a library of potential Nudix hydrolase substrates by phosphate sensor assay (Figure 3, Supplementary Table 2) revealed that these plant proteins possess Nudix hydrolase activity as well. Thus, experimental finding of dual activity of PpND and AtND confirmed our starting hypothesis, based on their primary structures. These two plant enzymes show both Nudix hydrolase and dipeptidyl peptidase III activity, catalysing the hydrolysis of phosphate bonds, and of peptide bonds. To our knowledge, this is the first report on Nudix and DPP III activity contained in a monomeric enzyme. Nevertheless, we were unable to express the individual domains of PpND separately, leading us to conclude that both domains are needed for proper folding and activity of this protein.

The existence of two distinct active sites (Nudix and DPP III) is supported by site-directed mutagenesis. Glu592 from the active-site motif H<sup>591</sup>ECCH<sup>595</sup> (Figure 1) is essential for DPP III activity of PpND. The importance of the glutamic acid residue from the characteristic zinc-binding motif HELLGH has been already established for the catalytic activity of mammalian and yeast DPP III (Fukasawa *et al.*, 1999; Baral *et al.*, 2008; Bezerra *et al.*, 2012). Mutational analysis further revealed that Glu92 from the Nudix box motif is crucial for the Nudix hydrolase (IPP hydrolase) activity of PpND. This residue aligns with Glu53 in *E. coli* MutT (multiple sequence alignment, MSA, not shown). The major role of Glu53 as a general base in MutT catalysis has been confirmed (Mildvan *et al.*, 2005).

Results of site-directed mutagenesis on essential glutamic acid residues (E92 and E592) in PpND indicate two distinct active sites (Nudix and DPP III).

Our results provide clear evidence that both Nudix-DPP III (PpND and AtND) have phosphohydrolase activity against IPP though its physiological relevance is unproven. Since the addition of inorganic pyrophosphatase in the assay did not have an effect on this hydrolytic activity, we inferred that both plant enzymes catalyze the hydrolysis of the terminal phosphate bond. This was confirmed by LC-MS/MS analysis of the reaction mixture in which the formation of isopentenyl monophosphate was identified. Isopentenyl diphosphate (IPP) and its isomer dimethylallyl diphosphate (DMAPP) are elemental (C<sub>5</sub> building blocks) for the biosynthesis of isoprenoids, naturally occurring compounds, produced by all living organisms. They are considered to be the largest and most diverse group of natural products

(George *et al.*, 2015), composed of over 50,000 compounds which include primary metabolites (sterols, carotenoids) and secondary metabolites. Isoprenoid compounds *in vivo* perform a large variety of functions (Berthelot *et al.*, 2012). In plants they are broadly represented and play essential roles acting as pigments, hormones, pollinator attractors, insecticides, pesticides. IPP and DMAPP are synthesized by two independent pathways, the mevalonic acid (MVA) and methylerythritol phosphate (MEP) pathways. Most recently, it was reported that, in a modified MVA pathway in the cytosol of plant cells, isopentenyl phosphate kinase (IPK) catalyses the formation of IPP from IP (Henry *et al.*, 2015). An IPP phosphatase was postulated. The authors suggest that IP/DMAP formation is likely an adaptive trait, and not just the consequence of general phosphatase activity. Both AtND and PpND are cytosolic proteins. Their ability to hydrolyze IPP distinguishes them from the majority of known Nudix hydrolases, though few others have been tested for such activity. Moreover, no IPP hydrolase has been characterized which has a level of activity that could be physiologically relevant. Our results support that PpND and AtND could participate in the plant isoprenoid metabolic network. These enzymes may have uses in biotechnology where IPP phosphohydrolase activity is useful.

Most recently, Magnard *et al.*, (2015) have shown that the cytosolic Nudix hydrolase RhNUDX1 from rose possesses geranyl diphosphate diphosphohydrolase activity *in vitro* and contributes to the biosynthesis of monoterpene alcohol geraniol in plants. Our findings on PpND and AtND hydrolytic activity towards IPP thus corroborates the idea that isoprenoid diphosphates may be natural substrates of Nudix hydrolases.

In summary, for the first time, two atypical members of the DPP III (M49) family, containing a Nudix box motif, were identified, heterologously expressed and biochemically characterized: *Physcomitrella patens* protein (PpND) and *Arabidopsis thaliana* protein (AtNUDT3, AtND). Our results revealed that both plant proteins possess dual enzymatic activity, acting as a dipeptidyl peptidase III and Nudix hydrolase, in agreement with our starting hypothesis based on the bioinformatic analysis. Two glutamic acid residues, E592 and E92, catalytically important for DPP III and Nudix hydrolase activity, respectively, were confirmed by site-directed mutagenesis. AtND and PpND cleaved DPP III substrates with different specificity. Both enzymes hydrolyzed isopentenyl diphosphate (IPP), a universal precursor for the biosynthesis of isoprenoid compounds, with similar catalytic efficiency. The activity was significant but insufficient to be certain of its physiological role. Future studies are needed to prove whether PpND and AtND participate in plant isoprenoid metabolism, and

whether PpND is involved in protection against oxidative stress in the moss *Physcomitrella patens*, both suggested by the biochemical rationale of our present results.

## Materials and methods

### Cloning and mutagenesis

The full-length cDNA clone for the *P. patens* A9TLP4 (PpND) was generated by GeneArt AG (Regensburg, Germany). The codon usage was adapted to the codon bias of *Escherichia coli* genes. The cDNA was cloned into the pET21a\_A028 bacterial expression vector between *NdeI* and *XhoI* restriction sites adding His-tag codons at the 3' end. The full-length cDNA clone for the *Arabidopsis thaliana* protein Q8L831, which we designated as «AtND», was prepared from *A. thaliana* cDNA clone U16680 obtained from the Arabidopsis Biological Resource Center (Yamada *et al.*, 2003) in the pLATE31 plasmid (aLICator kit, Thermo Fisher Scientific, Waltham, MA, USA).

Mutants of PpND were constructed using the QuikChange® XL Site-Directed Mutagenesis Kit (Stratagene, La Jolla, CA, USA). The template used for mutagenesis was the pET21a-A9TLP4-His<sub>6</sub> construct (GeneArt AG). Domain fragments of PpND were cloned into pLATE31 with a C-terminal His-tag. The Nudix domain fragment contained residues 1 to 206, and the DPP III domain fragment contained residues 207-770. The primers, listed in Supplementary Table 3 were custom synthesized by Sigma-Aldrich (St. Louis, MO, USA) and Macrogen (Seoul, South Korea).

### Production, purification and characterization of recombinant PpND and AtND proteins

Cells of *Escherichia coli* BL21- Codon Plus (DE3) - RIL were transformed with the appropriate plasmid (constructs containing PpND wild-type, mutant or domain fragment of PpND, or AtND), grown in Luria Bertani broth with ampicillin to OD<sub>600</sub> 0.6, and were then cooled to 18°C. The culture was supplemented with magnesium and zinc sulfate to final concentrations of 1mM MgSO<sub>4</sub> and 10µM ZnSO<sub>4</sub>. Expression was induced with 0.25 mM IPTG (isopropyl β-D-1-thiogalactopyranoside), and cells were harvested after 20 h at 18°C and 130 rpm. Cell harvesting and protein purification from bacterial pellet was performed according to Jajčanin-Jozić *et al.*, (2010). In brief, cell pellets were resuspended in lysis buffer

(50 mM Tris-HCl pH 8.0, 300 mM NaCl, 10 mM imidazole) and lysed enzymatically by lysozyme as well as mechanically with sonication. Proteins were purified using Ni-NTA affinity chromatography and gel filtration on the column of HiLoad 16/60 Superdex 200 prep grade. Protein purity was confirmed by SDS-PAGE according to Laemmli (1970) on a BioRad Mini-Protean Tetra Cell, and by SDS-PAGE with PhastGel Homogeneous 12.5 plates on PhastSystem (Pharmacia, Uppsala, Sweden). Molecular mass of purified proteins was determined by SDS-PAGE using PhastGel 8-25% gradient plates and low molecular weight standard mixture from Pharmacia for reference proteins. Isoelectric focusing for pI determination, and native-PAGE with PhastGel Gradient plates were performed on PhastSystem following manufacturer's procedures. Protein bands were stained with Coomassie Brilliant Blue R-250.

In order to check the secondary structure similarity between the PpND and AtND proteins, circular dichroism (CD) spectra were recorded on a Jasco J-815 spectropolarimeter (JASCO, Easton, MD, USA) with automatic temperature control using a cylindrical quartz cuvette of 0.1 mm or 1 mm path length. Enzyme samples prepared for CD measurements were exchanged into 10 mM Tris-HCl buffer, pH 7.4. The protein concentrations used were from 0.25 to 0.40 mg ml<sup>-1</sup>.

## **Enzyme activity determination and kinetic analysis**

### **Dipeptidyl peptidase activity**

DPP III activity was determined by a standard assay with Arg-Arg-2-naphthylamide (Arg<sub>2</sub>-2NA) as substrate, at 37 °C and at pH 7.4 (for PpND), or at pH 7.9 (for AtND) in Tris-HCl buffer (I=0.01) containing 75 μM CoCl<sub>2</sub>, using the colorimetric method (Špoljarić *et al.*, 2009).

The influence of metal cations on PpND hydrolytic activity towards Arg<sub>2</sub>-2NA was examined by adding metal salts (CoCl<sub>2</sub>, MnCl<sub>2</sub>, Mg-acetate, or Zn-acetate) at concentrations between 2 μM to 100 μM in the preincubation mixture with the enzyme, and after 3 min at 37°C the reaction was started with substrate. The effect of peptidase inhibitors was investigated by preincubation of the enzyme in 50 mM buffer with selected effectors for 30 min at 25°C, followed by the determination of the activity with the standard assay (15 min at 37°C, in the presence of CoCl<sub>2</sub>).



Kinetic parameters ( $K_m$  and  $k_{cat}$ ) for hydrolysis of Arg<sub>2</sub>-2NA and of other dipeptidyl-2-naphthylamides were determined with initial rate measurements (Jajčanin-Jozić *et al.*, 2010) at 25°C and at pH 7.4 (PpND), or at pH 7,9 (AtND) in Tris-HCl buffer (ionic strength 0.01) in the presence of the activator, 75  $\mu$ M CoCl<sub>2</sub>.

Protein concentrations were determined by the Bradford method (1976).

### **Nudix hydrolase activity**

Nudix hydrolase activity was determined by a continuous fluorescence assay (Xu *et al.*, 2013), using fluorescently tagged *E. coli* phosphate binding protein (PBP) as a phosphate sensor. The phosphate sensor was produced and purified using the methods described by Xu *et al.* (2013). The assay contained 10 mM Tris HCl pH 7.6, 1 mM MgCl<sub>2</sub>, and 5-10  $\mu$ M phosphate sensor. In the case where pyrophosphate was a possible product of the reaction, 0.05 U/ml inorganic pyrophosphatase (PPase) was added and where a nucleoside monophosphate was a possible product, 1 U/ml of alkaline phosphatase (APase) was added.

The enzymes were screened using a GENios microplate reader ( $\lambda_{ex}$  425 nm,  $\lambda_{em}$  465 nm, gain 50, 100 cycles). The reactions were recorded over 25 minutes at 37°C to determine substrates with significant kinetic activity. The substrate concentrations were kept at 5  $\mu$ M and screened against enzyme concentration of 50 nM. Kinetic parameters were found via FluoroMax-4 spectrofluorometer ( $\lambda_{ex}$  430 nm, slit width 1-2.5 nm,  $\lambda_{em}$  465 nm, slit width 1-5 nm). The concentration of Nudix enzyme ranged from 20–344 nM. These reactions were measured over 5 minutes at 37°C over varying concentrations of substrates. The slope of the standard curve of the sensor was measured to be  $1.8 \times 10^6$  RFU/ $\mu$ M phosphate by Fluoromax and  $2.5 \times 10^4$  RFU/ $\mu$ M phosphate by GENios.

The enzyme was initially screened against a library of 73 possible substrates. See Supplementary Table 1 for the full list of substrates and their groupings. Initial screening used 11 individual substrates and 11 substrates in pooled groups according to the structural similarity. Further screening of individual substrates within groups was conducted as needed. Specifically, PpND was additionally screened against ADP, ATP, dADP, dATP, 8-oxo-dATP, 2'-O-Me-ATP, N1-Me-ATP, N6-Me-ATP, dGDP, dGTP, GDP, GTP, p4G, 3'-dGDP, 8-oxo-dGTP, 2'-O-Me-GTP, 8-oxo-GTP, and N1-Me-GTP. The substrates with notable activity were then measured by careful kinetic analysis. The plots of  $v_i/[E_0]$  over different substrate concentrations were measured up to 20  $\mu$ M, in which saturation of the enzyme was

not achieved for any of the substrates for either PpND or AtND. Therefore, this substrate concentration range was deemed low enough ( $s < K_m$ ), to justify simplifying the Michaelis-Menten equation to the linear form at a fixed intercept at 0:

$$\frac{v_i}{[E_0]} = \frac{k_{cat}}{K_m} [S]$$

The ratio  $k_{cat}/K_m$  was obtained via linear regression for all reactions.

In order to identify isopentenyl monophosphate (IP) as a product of phosphatase activity of PpND towards IPP, liquid chromatography coupled to tandem mass spectrometry (LC-MS/MS-MRM) was performed. Standard chemicals IPP and IP were purchased from Sigma Aldrich. IPP (50  $\mu$ M) was incubated at 37°C for 10 min with PpND, 60 nM or 600 nM, in reaction mixture (total volume 100  $\mu$ l) containing 10 mM Tris-HCl buffer pH 7.6 and 1 mM MgCl<sub>2</sub>. Enzyme reaction was stopped by addition of 5 mM EDTA. As a control, IPP was incubated under the identical conditions, without the enzyme. All reaction mixtures were subjected to custom service analysis at BIOCenter, Zagreb, Croatia. In brief, reaction products were analyzed by liquid chromatography coupled to tandem MS/MS (Agilent Technologies 1290 Infinity LC System and Agilent Technologies 6460 Triple Quadrupole mass spectrometer, Agilent, San Jose, CA, USA) on an Eclipse Plus C18 RRHD column (50 mm length, 2.1 mm internal diameter, and 1.8  $\mu$ m particle size). Products of enzyme reactions were eluted isocratically with a mix of phase A (20 mM NH<sub>4</sub>HCO<sub>3</sub> containing 0.1% triethylamine) and phase B (80% acetonitrile, v/v, containing 0.1% triethylamine) in ratio 70:30, with a flow-rate of 0.2 ml min<sup>-1</sup>. The effluent (1  $\mu$ l) was introduced into the ion source of a 6460 Triple Quadrupole mass spectrometer. The capillary voltage was set to - 3500 V, desolvation gas flow was 5 l min<sup>-1</sup>, and desolvation temperature was 300°C. Negative ESI tandem mass detection was run in the multiple reaction monitoring (MRM) mode.

The Malachite Green Phosphate Detection Kit (R&D Systems, Minneapolis, MN, USA) was used to determine the liberated inorganic phosphate when Nudix hydrolase (phosphatase) activity of the PpND mutants was investigated towards IPP (Figure 4). Enzyme (60-70 nM) was incubated with IPP (50  $\mu$ M) for 10 min at 37°C in 500- $\mu$ l reaction mixture containing 10 mM TrisHCl (pH 7.6) and 1 mM MgCl<sub>2</sub>, and 125  $\mu$ l of 5 mM EDTA was added to stop the reaction.

### Isopentenyl diphosphate isomerase (IDI) assay

In order to test for potential IDI activity, a color complementation assay (Wang *et al.*, 2009) was performed. In this assay, *E. coli* XL1Blue cells are transformed with pAC-BETA plasmid carrying four genes necessary for production of  $\beta$ -carotene in *E. coli*. With addition of a plasmid carrying a gene for an IDI, the production of  $\beta$ -carotene in *E. coli* is significantly increased, resulting in visible change of colony color from yellow (without IDI) to orange. We obtained pAC-BETA and pTrcAtipi plasmids (containing the *IDI* gene from *A. thaliana*) as a generous gift from Dr. F. X. Cunningham. The pTrc plasmid was produced from pTrcAtipi by cutting out the AtIDI gene with *Pst*I enzyme and recircularization of the empty plasmid. Empty pTrc plasmid represented a negative control. The AtND and PpND sequences were cloned into the pTrc plasmid using specific primers (Supplementary Table 3), and restriction enzymes *Bam*HI and *Xba*I, so that the ORF began with MSRSAAGGS and continued with full length PpND and AtND. The variant PpND\_A76C was prepared by site-directed mutagenesis using pTrcPpND as the template. The double mutant PpND\_A76C\_E92A was generated using plasmid pTrcPpND\_E92A as template for Ala76Cys mutagenesis. To test *Physcomitrella* and *Arabidopsis* NDs for IDI activity, *E. coli* cells carrying pAC-BETA plasmid (Cm<sup>R</sup>) were transformed with one of the prepared pTrc plasmids (empty pTrc as a negative control, pTrcAtND, pTrcPpND, pTrcPpND\_A76C and pTrcPpND\_A76C\_E92A to be tested, and pTrcAtIDI as a positive control, all Amp<sup>R</sup>). Cells were grown on LB plates supplemented with chloramphenicol and ampicillin overnight at 37 °C and 5 days at room temperature for the development of color.

### Bioinformatics methods

Sequences of PpND homologues from other species were obtained from UniProt KB (July 2015, [www.uniprot.org](http://www.uniprot.org)) (The UniProt Consortium 2011) following BLASTP search (Altschul *et al.*, 1997). Multiple sequence alignment was done using Clustal O 1.2.1 (Sievers *et al.* 2011) at <http://www.ebi.ac.uk/Tools/msa/clustalo/>. Online databases of plant genomes were searched for homologues of the *Physcomitrella* A9TLP4 protein (PpND) using Conserved Domain Architecture Retrieval Tool (July 2015, Conserved Domain Database v. 3.14) (Geer *et al.*, 2002).

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## Tables and figures

**Table 1** Hydrolytic activity<sup>a</sup> of *P. patens* A9TLP4 (PpND) and *A. thaliana* NUDT3 (AtND) towards peptidase substrates.

Substrate	Relative hydrolysis rate (%)	
	PpND	AtND
Arg-Arg-2NA	100	100
Ala-Arg-2NA	62	684
Ala-Ala-2NA	40	366
Gly-Arg-2NA	16	197
Pro-Arg-2NA	14	622
Phe-Arg-2NA	12	411
Asp-Arg-2NA	0	0
Gly-Pro-2NA	0	0
Gly-Phe-2NA	0	0
Arg-2NA	0	0
Leu-2NA	0	0
Ala-2NA	0	0
BANA	0	0

<sup>a</sup>The enzymes were incubated at 37°C in Tris-HCl buffer (ionic strength = 0.01) pH 7.5 (PpND), respectively 7.9 (AtND) containing 75 µM CoCl<sub>2</sub>, with the substrates at a final concentration of 0.04 mM (100 % = 131.4 nmol, or 17.2 nmol of Arg<sub>2</sub>-2NA hydrolyzed per min and milligram of PpND or AtND, respectively). 2NA: 2-naphthylamide; BANA: N $\alpha$ -benzoyl-L-Arg-2-naphthylamide

**Table 2** Kinetic parameters<sup>a</sup> for the hydrolysis of dipeptidyl-2-naphthylamide substrates catalyzed by the *Arabidopsis thaliana* Nudix-DPP III (AtND).

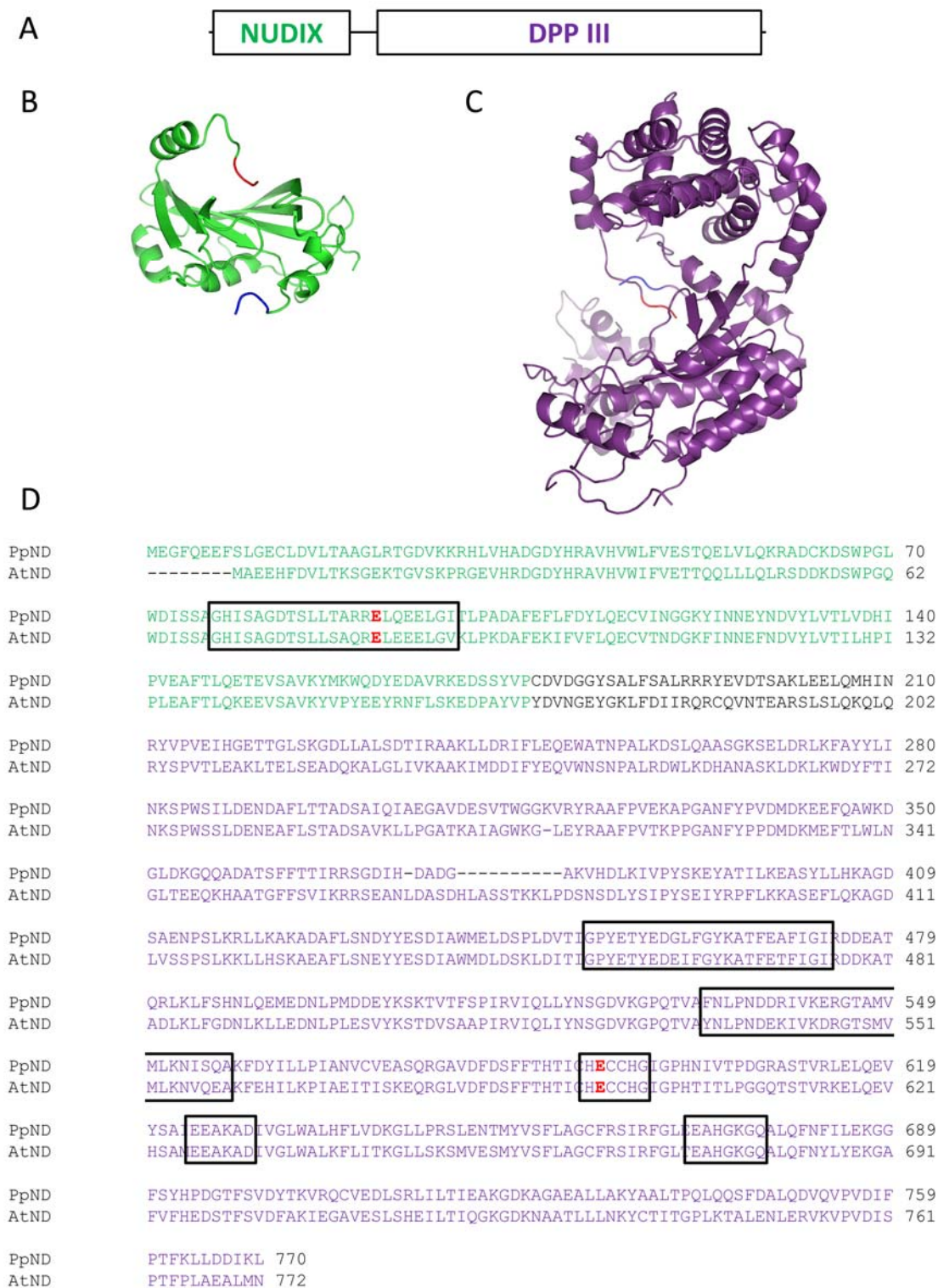
Substrate	$K_m$ (µM)	$k_{cat}$ (min <sup>-1</sup> )	$k_{cat}/K_m$ (M <sup>-1</sup> s <sup>-1</sup> )
Ala-Arg-2NA	5.0 ± 0.3	2.49 ± 0.21	8300.0
Pro-Arg-2NA	40.85 ± 12.09	2.23 ± 0.3	908.2
Arg-Arg-2NA	3.25 ± 1.12	0.120 ± 0.001	615.4
Ala-Ala-2NA	240.2 ± 69.4	2.28 ± 0.41	158.2

<sup>a</sup>The kinetic parameters were determined from the initial reaction rates at 25°C, as described in materials and methods.  $K_m$  and  $k_{cat}$  values are presented as mean ± SD.

**Table 3** Kinetic parameters for Nudix hydrolase catalyzed reactions.

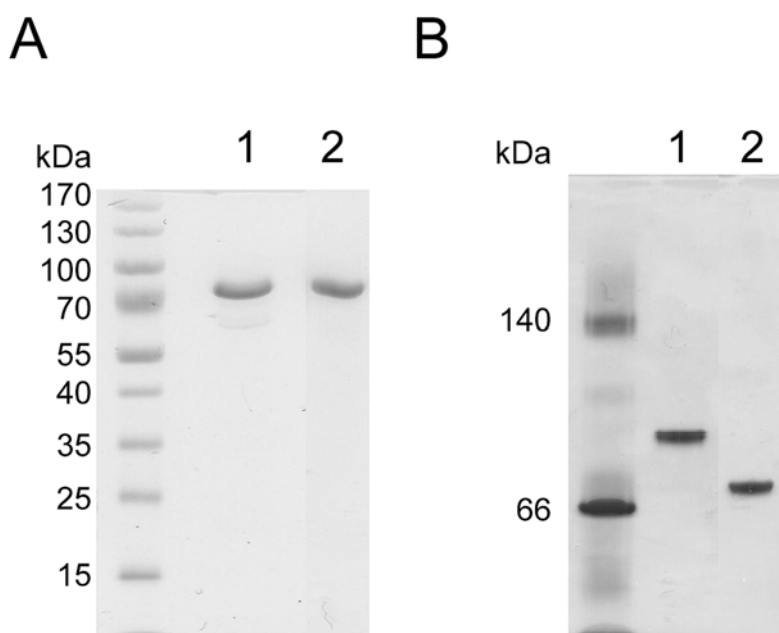
Enzyme	Substrate	$k_{cat}/K_m$ (M <sup>-1</sup> s <sup>-1</sup> )
PpND	IPP	$(1.9 \pm 0.16) \times 10^4$
	8-oxo-dATP	$(1.1 \pm 0.64) \times 10^4$
	ADP	280 ± 26
	dGDP	170 ± 18
	dGTP	185 ± 15
AtND	IPP	$(1.8 \pm 0.13) \times 10^4$

## New plant hydrolase with IPP dephosphorylation ability



**Figure 1** (A) a schematic representation of plant Nudix-DPP III proteins, where the first 200 amino acids form a Nudix fold domain, and the last 550 amino acids form a DPP III fold; (B) PpND model (residues 5-177) based on human IDI1 structure (PDB: 2i6k; Zhang *et al.*, 2007); (C) PpND model (residues 210-760) based on human DPP III structure (PDB: 3fvy; Bezerra *et al.*, 2012). First three amino acids in the models are colored blue to indicate N-

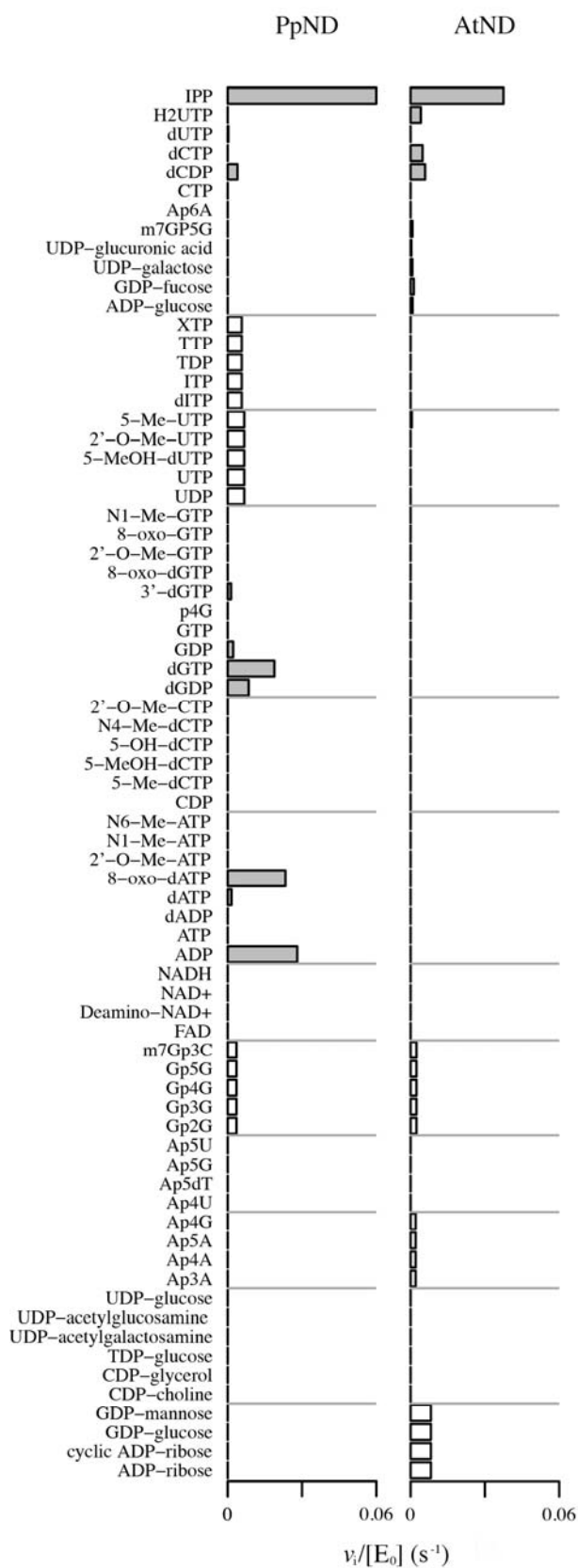
terminus, last three amino acids are colored red to indicate C-terminus. Models were made using Phyre2 server ([www.sbg.bio.ic.ac.uk/phyre2](http://www.sbg.bio.ic.ac.uk/phyre2)) and visualized using PyMOL ([www.pymol.org](http://www.pymol.org)); (D) alignment of protein sequences of *Physcomitrella patens* and *Arabidopsis thaliana* Nudix-DPP III, with framed variant Nudix box and five conserved regions of the DPP III family. Catalytic glutamates – E92 for Nudix hydrolase and E592 for peptidase activity – are given in red. In all panels, Nudix domain is shown in green and DPP III domain in purple. Putative linker sequence is shown in black.



**Figure 2** Polyacrylamide gel electrophoresis (PAGE) of affinity and size-exclusion purified AtND (lane 1) and PpND (lane 2).

(A) SDS-PAGE performed under reducing conditions on PhastGel Homogeneous (12.5 %) plate; (B) native-PAGE performed on PhastGel Gradient (8-25%) plate. Proteins were visualized by Coomassie Blue staining.

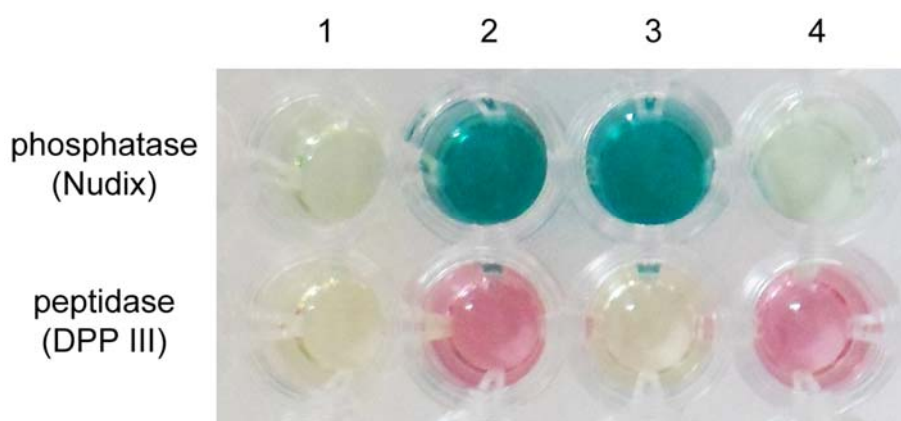
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**Figure 3** Substrate specificity screening of PpND and AtND against a 73-compound library by phosphate sensor assay.

## New plant hydrolase with IPP dephosphorylation ability

The  $v_i/[E_0]$  values ( $s^{-1}$ ) are graphed for every substrate in the library that was found to have detectable activity against the enzymes. These reactions were measured at pH 7.6 and 37°C, with 5  $\mu$ M substrate and 50 nM enzyme. Estimated activities for substrates that were screened individually are recorded in gray, while the activities of substrates within a group are recorded in white.



**Figure 4** Activity of the wild-type and PpND mutants towards IPP (phosphatase) and Arg<sub>2</sub>-2NA (peptidase) determined using colorimetric assays described under materials and methods.

1, negative control (reaction mixture without the enzyme); 2, wild-type PpND; 3, PpND E592A, disrupting DPP III motif; 4, PpND E92A, disrupting variant Nudix box.

## Supplementary table and figures

**Supplementary Table 1** Substrates used for Nudix hydrolase activity screening.

Group	Abbreviation	Name	2nd Enzyme
1	ADP-ribose	Adenosine 5'-diphosphoribose	APase
1	cyclic ADP-ribose	Cyclic adenosine diphosphate-ribose	APase
1	GDP-glucose	Guanosine 5'-diphosphoglucose	APase
1	GDP-mannose	Guanosine 5'-diphospho-D-mannose	APase
2	CDP-choline	Cytidine 5'-diphosphocholine	APase
2	CDP-glycerol	Cytidine 5'-diphosphoglycerol	APase
2	TDP-glucose	Thymidine-5'-diphospho- $\alpha$ -D-glucose	APase
2	UDP-acetylgalactosamine	Uridine 5'-diphospho-N-acetylgalactosamine	APase
2	UDP-acetylglucosamine	Uridine 5'-diphospho-N-acetylglucosamine	APase
2	UDP-glucose	Uridine 5'-diphosphoglucose	APase
3	Ap3A	P1,P3-Di(adenosine-5') triphosphate	APase
3	Ap4A	P1,P4-Di(adenosine-5') tetraphosphate	APase
3	Ap5A	P1,P5-Di(adenosine-5') pentaphosphate	APase
3	Ap4G	P1(-5'-Adenosyl)-P4-(5'-guanosyl)-tetraphosphate	APase
4	Ap4U	P1(-5'-Adenosyl)-P4-(5'-uridyl)-tetraphosphate	APase
4	Ap5dT	P1(-5'-Adenosyl)-P5-[5'-(2'-deoxy-thymidyl)]-pentaphosphate	APase
4	Ap5G	P1(-5'-Adenosyl)-P5-(5'-guanosyl)-pentaphosphate	APase
4	Ap5U	P1(-5'-Adenosyl)-P5-(5'-uridyl)-pentaphosphate	APase
5	Gp2G	P1(-5'-guanosyl)-P2-(5'-guanosyl)-diphosphate	APase
5	Gp3G	P1(-5'-guanosyl)-P3-(5'-guanosyl)-triphosphate	APase
5	Gp4G	P1(-5'-guanosyl)-P4-(5'-guanosyl)-tetraphosphate	APase
5	Gp5G	P1(-5'-guanosyl)-P5-(5'-guanosyl)-tetraphosphate	APase
5	m7Gp3C	P1(-5'-7-methyl-guanosyl)-P3-(5'-cytidyl)-triphosphate	APase
6	FAD	Flavin Adenine Dinucleotide	APase
6	Deamino-NAD <sup>+</sup>	Nicotinamide hypoxanthine dinucleotide	APase
6	NAD <sup>+</sup>	Nicotinamide Adenine Dinucleotide	APase
6	NADH	Nicotinamide Adenine Dinucleotide-Reduced	APase
7	ADP	Adenosine 5'-diphosphate	PPase
7	ATP	Adenosine-5'-Triphosphate	PPase

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7	dADP	2'-Deoxyadenosine-5'-diphosphate	PPase
7	dATP	2'-Deoxyadenosine 5'-triphosphate	PPase
7	8-oxo-dATP	8-Oxo-2'-deoxyadenosine-5'-Triphosphate	PPase
7	2'-O-Me-ATP	2'-O-Methyladenosine-5'-Triphosphate	PPase
7	N1-Me-ATP	N1-Methyladenosine-5'-Triphosphate	PPase
7	N6-Me-ATP	N6-Methyladenosine-5'-Triphosphate	PPase
8	CDP	Cytidine-5'-diphosphate	PPase
8	5-Me-dCTP	5-Methyl-2'-deoxycytidine-5'-Triphosphate	PPase
8	5-MeOH-dCTP	5-Hydroxymethyl-2'-deoxycytidine-5'-Triphosphate	PPase
8	5-OH-dCTP	5-Hydroxy-2'-deoxycytidine-5'-Triphosphate	PPase
8	N4-Me-dCTP	N4-Methyl-2'-deoxycytidine-5'-Triphosphate	PPase
8	2'-O-Me-CTP	2'-O-Methylcytidine-5'-Triphosphate	PPase
9	dGDP	2'-Deoxyguanosine-5'-diphosphate	PPase
9	dGTP	2'-Deoxyguanosine 5'-triphosphate	PPase
9	GDP	Guanosine-5'-diphosphate	PPase
9	GTP	Guanosine 5'-triphosphate	PPase
9	p4G	Guanosine 5'-tetraphosphate	PPase
9	3'-dGTP	3'-Deoxyguanosine-5'-Triphosphate	PPase
9	8-oxo-dGTP	8-Oxo-2'-deoxyguanosine-5'-Triphosphate	PPase
9	2'-O-Me-GTP	2'-O-Methylguanosine-5'-Triphosphate	PPase
9	8-oxo-GTP	8-Oxoguanosine-5'-Triphosphate	PPase
9	N1-Me-GTP	N1-Methylguanosine-5'-Triphosphate	PPase
10	UDP	Uridine 5'-diphosphate	PPase
10	UTP	Uridine-5'-Triphosphate, Trisodium Salt	PPase
10	5-MeOH-dUTP	5-Hydroxymethyl-2'-deoxyuridine-5'-Triphosphate	PPase
10	2'-O-Me-UTP	2'-O-Methyluridine-5'-Triphosphate	PPase
10	5-Me-UTP	5-Methyluridine-5'-Triphosphate	PPase
11	dITP	2'-Deoxyinosine-5'-Triphosphate	PPase
11	ITP	Inosine-5'-Triphosphate	PPase
11	TDP	Thymidine-5'-diphosphate	PPase
11	TTP	Thymidine-5'-triphosphate	PPase
11	XTP	Xanthosine-5'-Triphosphate	PPase
Individual	ADP-glucose	Adenosine-5'-diphosphoglucose	APase
Individual	GDP-fucose	Guanosine 5'-diphospho-β-L-fucose	APase
Individual	UDP-galactose	Uridine 5'-diphosphogalactose	APase
Individual	UDP-glucuronic acid	Uridine 5'-diphosphoglucuronic acid	APase
Individual	m7GP5G	P1-(5'-7-methyl-guanosyl)-P5-(5'-guanosyl)-pentaphosphate	APase

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Individual Ap6A	P1(-5'-Adenosyl)-P6-(5'-adenosyl)-hexaphosphate	APase
Individual CTP	Cytidine 5'-triphosphate	PPase
Individual dCDP	2'-Deoxycytidine-5'-diphosphate	PPase
Individual dCTP	2'-Deoxycytidine 5'-triphosphate disodium salt	PPase
Individual dUTP	2'-Deoxyuridine-5'-Triphosphate	PPase
Individual H2UTP	5,6-Dihydrouridine-5'-Triphosphate	PPase
Individual IPP	Isopentenyl Pyrophosphate	NA

APase: alkaline phosphatase; PPase: inorganic pyrophosphatase.

When the substrates are initially screened for Nudix hydrolase activity, most are tested in mixtures. All substrates that have the same group number are first screened in the same batch, and only if there is activity detected are the substrates within that group then tested individually.

**Supplementary Table 2** (to complement Figure 3 depicting Nudix library screening).

Substrate	PpND activity $v_i/[E_0]$ ( $s^{-1}$ )	AtND activity $v_i/[E_0]$ ( $s^{-1}$ )
ADP-ribose	0	$8 \times 10^{-3}$
cyclic ADP-ribose	0	$8 \times 10^{-3}$
GDP-glucose	0	$8 \times 10^{-3}$
GDP-mannose	0	$8 \times 10^{-3}$
CDP-choline	0	0
CDP-glycerol	0	0
TDP-glucose	0	0
UDP-acetylgalactosamine	0	0
UDP-acetylglucosamine	0	0
UDP-glucose	0	0
Ap3A	0	$2 \times 10^{-3}$
Ap4A	0	$2 \times 10^{-3}$
Ap5A	0	$2 \times 10^{-3}$
Ap4G	0	$2 \times 10^{-3}$
Ap4U	0	0
Ap5dT	0	0
Ap5G	0	0
Ap5U	0	0
Gp2G	$4 \times 10^{-3}$	$2 \times 10^{-3}$
Gp3G	$4 \times 10^{-3}$	$2 \times 10^{-3}$
Gp4G	$4 \times 10^{-3}$	$2 \times 10^{-3}$
Gp5G	$4 \times 10^{-3}$	$2 \times 10^{-3}$
m7Gp3C	$4 \times 10^{-3}$	$2 \times 10^{-3}$
FAD	0	0
Deamino-NAD <sup>+</sup>	0	0



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NAD+	0	0
NADH	0	0
ADP	$3 \times 10^{-2}$	0
ATP	0	0
dADP	0	0
dATP	$2 \times 10^{-3}$	0
8-oxo-dATP	$2 \times 10^{-2}$	0
2'-O-Me-ATP	0	0
N1-Me-ATP	0	0
N6-Me-ATP	0	0
CDP	0	0
5-Me-dCTP	0	0
5-MeOH-dCTP	0	0
5-OH-dCTP	0	0
N4-Me-dCTP	0	0
2'-O-Me-CTP	0	0
dGDP	$8 \times 10^{-3}$	0
dGTP	$2 \times 10^{-2}$	0
GDP	$2 \times 10^{-3}$	0
GTP	0	0
p4G	0	0
3'-dGTP	$1 \times 10^{-3}$	0
8-oxo-dGTP	0	0
2'-O-Me-GTP	0	0
8-oxo-GTP	0	0
N1-Me-GTP	0	0
UDP	$7 \times 10^{-3}$	0
UTP	$7 \times 10^{-3}$	0
5-MeOH-dUTP	$7 \times 10^{-3}$	0
2'-O-Me-UTP	$7 \times 10^{-3}$	0
5-Me-UTP	$7 \times 10^{-3}$	$5 \times 10^{-4}$
dITP	$6 \times 10^{-3}$	0
ITP	$6 \times 10^{-3}$	0
TDP	$6 \times 10^{-3}$	0
TTP	$6 \times 10^{-3}$	0
XTP	$6 \times 10^{-3}$	0
ADP-glucose	0	$8 \times 10^{-4}$
GDP-fucose	0	$1 \times 10^{-3}$
UDP-galactose	0	$7 \times 10^{-4}$
UDP-glucuronic acid	0	$3 \times 10^{-4}$
m7GP5G	0	$8 \times 10^{-4}$
Ap6A	0	0

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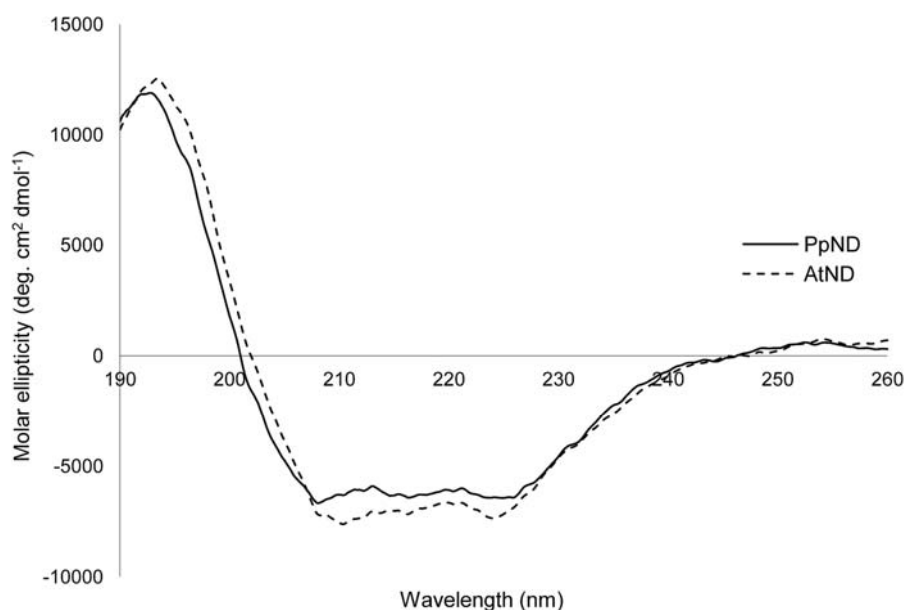
CTP	0	0
dCDP	$4 \times 10^{-3}$	$6 \times 10^{-3}$
dCTP	0	$5 \times 10^{-3}$
dUTP	$2 \times 10^{-4}$	0
H2UTP	0	$4 \times 10^{-3}$
IPP	$6 \times 10^{-2}$	$4 \times 10^{-2}$

Highlighted numbers indicate the screened activity for a substrate that was assayed individually against the enzyme. Otherwise, the activity against a substrate was measured as part of group mixture. Screening values are very approximate.

**Supplementary Table 3** Primers (synthesized by Sigma-Aldrich and Macrogen) used for cloning and/or mutagenesis.

<b>AtND cloning</b>	
AtND_F	AGAAGGAGATATAACTATGGCGGAGGAGCACTTCGACG
AtND_R	GTGGTGGTGTGGTGTGGCCGTTTCATCAATGCTTCTGCTAATGGAAATGTCGG
<b>PpND active site mutants</b>	
PpND_595G_F	CATCTGCCATGAATGCTGTGGCCATGGTATTGGTCCGC
PpND_595G_R	GCGGACCAATACCATGGCCACAGCATTTCATGGCAGATG
PpND_C594L_F	CCATGAATGCCTGGGCCATGGTATTGG
PpND_C594L_R	CCAATACCATGGCCCAGGCATTTCATGG
PpNd_E92A_F	GACCGCACGTCGTGCGCTGCAAGAAGAAGT
PpND_E92A_R	CAGTTCTTCTGTCAGCGCACGACGTGCGGTC
PpND_E592A_F	CCATACCATCTGCCATGCGTGCTGTCATGGTATTGG
PpND_E592A_R	CCAATACCATGACAGCACGCATGGCAGATGGTATGG
PpND_A76C_F	GGGATATTAGCAGCTGCGGTCATATTAGTGCC
PpND_A76C_R	GGCACTAATATGACCCGAGCTGCTAATATCCC
<b>M49 fragment cloning</b>	
PpND_DPP3frag_F	AGAAGGAGATATAACTATGCACATTAATCGTTATGTTCCGG
PpND_DPP3frag_R	GTGGTGGTGTGGTGTGGCCAGTTTGATGTCTGCCAGC
<b>Cloning into pTrc for IDI assay</b>	
PpNDTrc_F	GCCGGATCCATGGAAGGCTTTC AAGAGG
PpNDTrc_R	GCCTCTAGATTACAGTTTGATGTCTGCCAGCAG
AtNDTrc_F	GCCGGATCCATGGCGGAGGAGCACTTC
AtNDTrc_R	GCCTCTAGATTAGTTCATCAATGCTTCTGCTAATGG

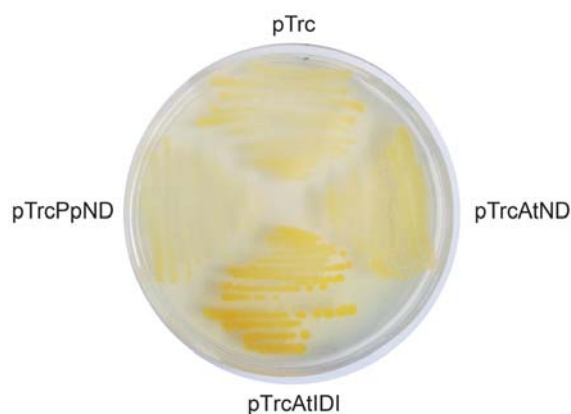
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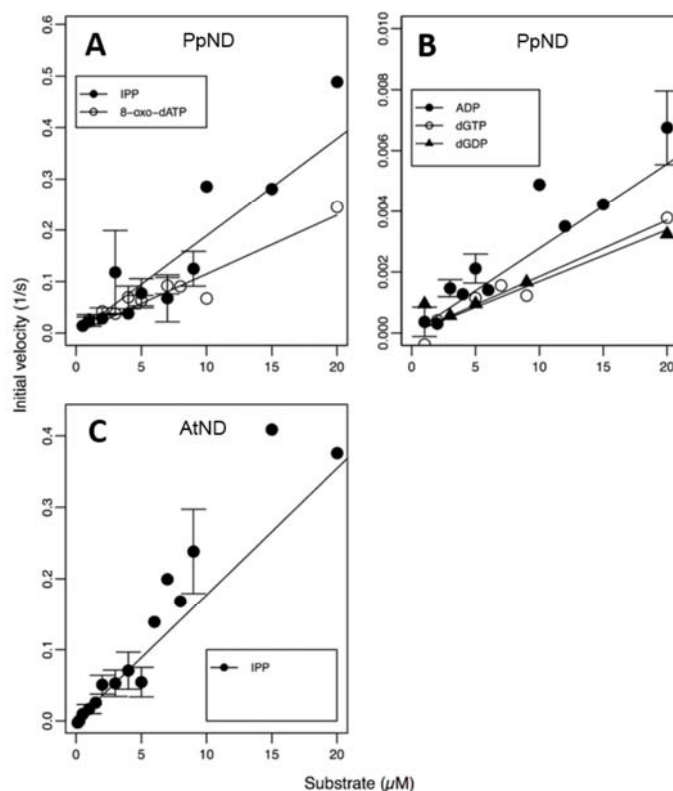
**Supplementary Figure 1** Circular dichroism spectra of wild-type PpND and of AtND proteins.

Q13907 IDI1_HUMAN	-----MPEINTNHLDKQQVQLLAEMCILIDENDNKIGAETKKNCHLNENIEKGL	49
Q46822 IDI_ECOLI	-----MQTEHVILLNAQGVPPTGTL-----KYAAHTADTR	30
Q9HP40 IDI_HALSA	MRDSMSEADRSSPGSGKTDREDETAENATQDVIIVTPDDERTGLAN-----RLDAHTGDGV	56
A9TLP4 A9TLP4_PHYPA	-----MEGFQEEFSLGECLDVLTAAAGLRTGDVK-----KRHLVHADGD	38
Q8L831 NUDT3_ARATH	-----MAEEHFVLTKSGEKTGVSK-----PRGEVHRDGD	30
Q13907 IDI1_HUMAN	LHRAF <sup>●</sup> SVFLFNTENK-LLLQQRSDAKITFPGCFTNTCCSHPLSNPAELEESDALGVRRAA	108
Q46822 IDI_ECOLI	LHLAFSSWLFNAKGQ-LLVTRRALS <sup>●</sup> KAWPGVWTNSVCGHPQLGESNE-----DAV	80
Q9HP40 IDI_HALSA	RHRAFTCLLFDEEDGR-VLLAQRADKRLWDTHWDGTVASHPIEQSQV-----DAT	106
A9TLP4 A9TLP4_PHYPA	YHRAVHVWLFVESTQELVLRADCKDSWPGLWDISSAGHISAGDTSL-----LTA	89
Q8L831 NUDT3_ARATH	YHRAVHVWIFVETTQQLLQLRSDDKDSWPQWDISSAGHISAGDTSL-----LSA	81
Q13907 IDI1_HUMAN	QRRLKAE <sup>●</sup> LGIPLEEVPEEINYLTRIHYKAQSDGIWGEHEIDYILLVRK-----NVTLNP	163
Q46822 IDI_ECOLI	IRRCRYELGVEITPPES----IYPDFRYRATDPSGIVENEVCPVFAART----TS-ALQI	131
Q9HP40 IDI_HALSA	RQLAEELGIEPHQYDKLE--ITDRFEYKRRYLDEGLEWEVCAVLQATL----HDTSFDR	160
A9TLP4 A9TLP4_PHYPA	RRELQEEELGITLPADAFEF--LFDYLQECVINGGKYINNEYNDVYLVTLVDHIPVEAFTL	147
Q8L831 NUDT3_ARATH	QRELEEE <sup>●</sup> LGVKLPKDAFEK--IFVFLQECVTNDGKFINNEFNDVYLVTLHPIPLEAFTL	139
Q13907 IDI1_HUMAN	DPNEIKSYCVSKEELKELKKAASG--EIK----ITPWFKIIAATFL-FKWARD----NL	212
Q46822 IDI_ECOLI	NDDEVMDYQ---WCDLADVLHGIDA--TPWAF----SPWMVMQATNREARK-----	173
Q9HP40 IDI_HALSA	DPEEVGGAM---WVDYEDLYENPRY-YRQLRL----CPWFEIAMRRDFEGDAD-----	205
A9TLP4 A9TLP4_PHYPA	QETEVSAVKYMKWQDYEDAVRKEDSSYVPCDVGGSALFSAL-RRRYEVDTSAKLEELQ	206
Q8L831 NUDT3_ARATH	QKEEVSAVKYVPYEEYRNFLSKEDPAYVPYDVNGEYKGLFDII-RQRQVNTTEARSLSLQ	198

**Supplementary Figure 2** Multiple sequence alignment of human, *Escherichia coli* and *Halobacterium salinarium* IDIs with PpND (A9TLP4) and AtND (Q8L831) proteins (made using Clustal O 1.2.1.). Residues important for IDI activity (known from human and *E. coli* IDI structures) are highlighted yellow, the catalytic cysteine is marked with a black circle, and the IDI active-site motif (NTCCSH) with a line above the sequence.



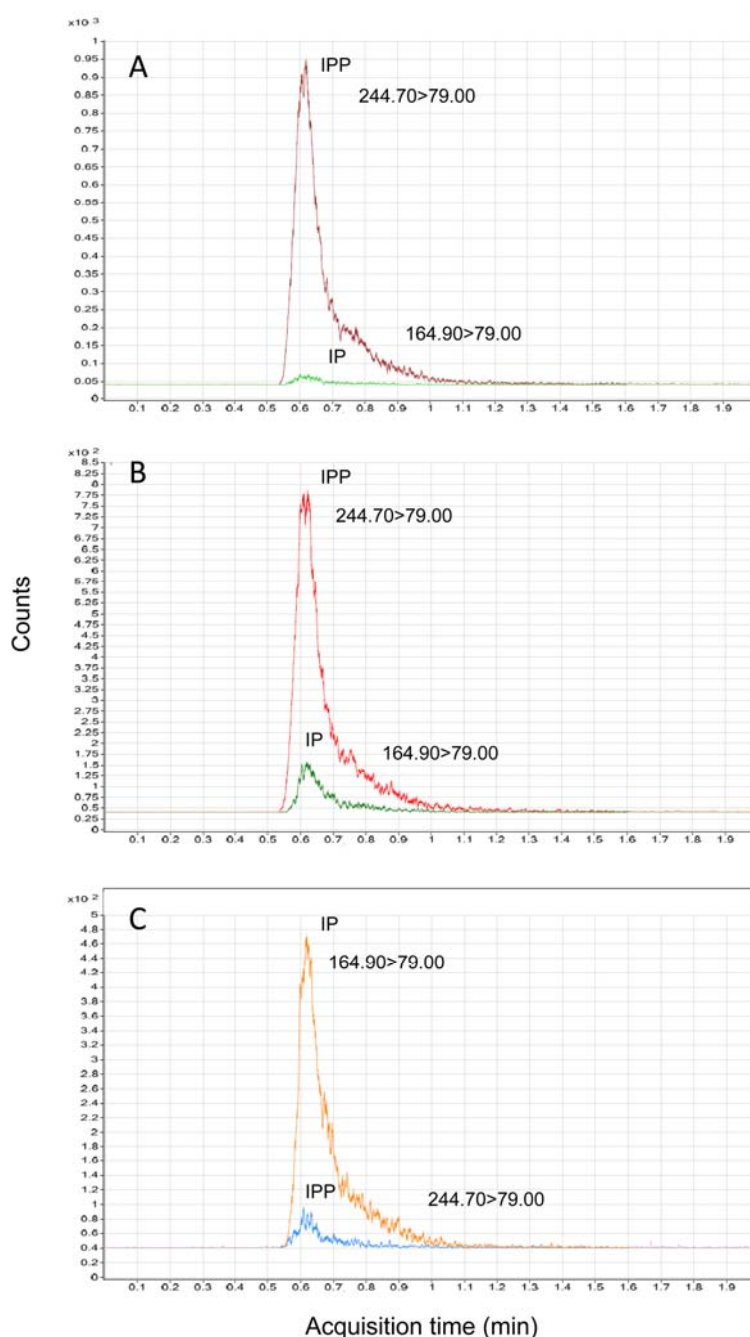
**Supplementary Figure 3** IDI color complementation assay. pTrc is a negative control, pTrcAtIDI is a positive control containing IDI gene from *Arabidopsis thaliana*, pTrcPpND contains Nudix-DPP III (PpND) from *Physcomitrella patens*, and pTrcAtND contains Nudix-DPP III (AtND) from *A. thaliana*.



**Supplementary Figure 4** Kinetic characterization of PpND and AtND against their most reactive substrates. The reactions against the substrates with the highest recorded activity identified in the screening were measured using kinetic analysis. These reactions were measured at pH 7.6 and 37°C, with enzyme concentrations from 20 to 344 nM. Error bars are

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shown where justified. In all cases, linear regression was used to determine the ratio of  $k_{\text{cat}}/K_m$ . The reported values of the kinetic parameters for enzyme and substrate are recorded in Table 3.



**Supplementary Figure 5** Extracted LC-MS chromatograms of ion product  $m/z=79.00$  of IPP (244.70) and IP (164.90): (A) control, IPP incubated at 37°C for 10 min, no enzyme added; (B) IPP incubated at 37°C for 10 min with PpND (60 nM); (C) IPP incubated at 37°C for 10 min with PpND (600 nM).