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Introduction

Currently, more than 170 RNA modifications are known.¹ Among the least explored RNA modifications are 5' RNA caps. A typical example of a noncanonical RNA cap is nicotinamide adenine dinucleotide (NAD), which has been detected in all domains of life.^{2–5} Other caps, such as NADH (a reduced form of NAD),⁶ coenzyme A (CoA),⁷ and flavin adenine dinucleotide (FAD),⁸ have been detected only in some organisms. Recently, we have discovered an entirely new class of RNA caps in bacteria with the structure of dinucleoside polyphosphates (Np_nNs).⁹ The presence of free Np_nNs molecules in cells of various organisms has been known more than 50 years.^{10–12} Their intracellular concentration increases under stress conditions,¹³ thus they are often called alarmones. For instance, Ap₃A and Ap₄A are known to act as alarmones in *Arabidopsis*

Arabidopsis thaliana NudiXes have RNA-decapping activity[†]

Maria-Bianca Mititelu, ^b^{ab} Oldřich Hudeček,^a Agnieszka Gozdek,^c Roberto Benoni,^a Ondřej Nešuta, ^b^a Szymon Krasnodębski,^c Joanna Kufel^c and Hana Cahová ^{*}

Recent discoveries of various noncanonical RNA caps, such as dinucleoside polyphosphates (Np $_{\sigma}$ N), coenzyme A (CoA), and nicotinamide adenine dinucleotide (NAD) in all domains of life have led to a revision of views on RNA cap function and metabolism. Enzymes from the NudiX family capable of hydrolyzing a polyphosphate backbone attached to a nucleoside are the strongest candidates for degradation of noncanonically capped RNA. The model plant organism Arabidopsis thaliana encodes as many as 28 NudiX enzymes. For most of them, only in vitro substrates in the form of small molecules are known. In our study, we focused on four A. thaliana NudiX enzymes (AtNUDT6, AtNUDT7, AtNUDT19 and AtNUDT27), and we studied whether these enzymes can cleave RNA capped with NprNs (Ap2-5A, Gp3-4G, Ap3-5G, m⁷Gp3G, m⁷Gp3A), CoA, ADP-ribose, or NAD(H). While AtNUDT19 preferred NADH-RNA over other types of capped RNA, AtNUDT6 and AtNUDT7 preferentially cleaved Ap₄A-RNA. The most powerful decapping enzyme was AtNUDT27, which cleaved almost all types of capped RNA at a tenfold lower concentration than the other enzymes. We also compared cleavage efficiency of each enzyme on free small molecules with RNA capped with corresponding molecules. We found that AtNUDT6 prefers free Ap₄A, while AtNUDT7 preferentially cleaved Ap₄A-RNA. These findings show that NudiX enzymes may act as RNA-decapping enzymes in A. thaliana and that other noncanonical RNA caps such as Ap₄A and NADH should be searched for in plant RNA.

> *thaliana* (*A. thaliana*),¹³ although their mechanism of action is unknown. The discovery of these molecules as RNA caps opened new perspectives on their function and metabolism. Free molecules such as Np_nNs, NAD, and CoA are usually cleaved by NudiX enzymes. Plants especially have a high number of NudiX-encoding genes, with a total of 28 in *A. thaliana*;¹⁴ by comparison, there are 24 in human cells and only 13 in *Escherichia coli*.¹⁵ Nevertheless, it is still unclear why so many NudiX enzymes are necessary for cellular metabolism and why many are redundant, *i.e.* two or more enzymes cleave the same substrates *in vitro*. Importantly, some NudiX enzymes have also been reported to have RNA-decapping activity.^{16–20} Identification of the new decapping role of some NudiX enzymes will help us to reveal new potential RNA caps and explore their biological role.

> In our search for new noncanonical RNA caps, we explored *A. thaliana* NudiX proteins and their substrate specificity. In particular, we examined four enzymes, cytosolic AtNUDT6, AtNUDT7, and chloroplastic AtNUDT19 and AtNUDT27, since small molecules such as NAD(H), CoA, ADP-ribose, and various Np_nNs are known to be their substrates *in vitro*.¹⁴ In addition, these proteins are the closest plant homologs of mammalian Nudt2 and Nudt12, which have been demonstrated to have RNA non-canonical decapping activity *in vitro*, namely deFADding and deCoAding, respectively.^{20,21} We therefore tested whether

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^a Institute of Organic Chemistry and Biochemistry of the CAS, Flemingovo náměstí 2, Prague 6, Czechia. E-mail: cahova@uochb.cas.cz

^b Charles University, Faculty of Science, Department of Cell Biology, Viničná 7, Prague 2, Czechia

^c Institute of Genetics and Biotechnology, Faculty of Biology, University of Warsaw, Pawinskiego 5a, Warsaw 02-106, Poland

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Results and discussion

First, we screened the *in vitro* activity of AtNUDT6, AtNUDT7, AtNUDT19, and AtNUDT27 on RNA capped with NAD(H), CoA, ADP-ribose, canonical $m^7Gp_3A(G)$, or various dinucleoside polyphosphates (Ap₂₋₅A, Gp₃₋₄G, Ap₃₋₅G). We prepared ³²P labelled RNA by *in vitro* transcription with T7 RNA polymerase in the presence of cap precursors using two DNA templates (Table S1, ESI†). Template 35A has at +1 position A and was used in experiments with Ap₂₋₅A, Gp₃₋₄A, NAD(H), CoA, ADP-ribose and m^7Gp_3A . Template 35G has at position +1 G and was used in experiments with Gp₃₋₄G, Ap₃₋₅G, and m^7Gp_3G . Asymmetrical Np_nNs such as Ap₃₋₅G can be used with both templates in order to obtain RNA with flanking A or G, Ap₃₋₅G-RNA, or Gp₃₋₅A-RNA, respectively.

Following *in vitro* transcription, we purified the capped RNA from uncapped by treatment with 5' polyphosphatase and TerminatorTM exonuclease. This treatment guaranteed that the amount of capped RNA in each reaction was in average 96 \pm 4% (only in case of m⁷Gp₃A(G)-RNA the amount of capped RNA was 58 \pm 3%). Afterwards, we treated capped RNA with NudiX enzymes (Fig. 1A) and followed the cleavage with PAGE analysis (Fig. 1B and Fig. S1 and S2, ESI†). We observed the highest enzymatic activity on the capped RNA at 500 nM concentration for AtNUDT27

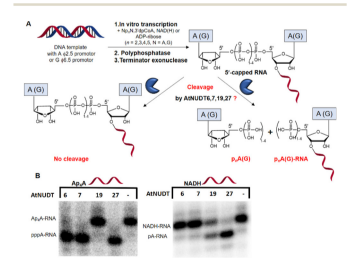


Fig. 1 Screening of the RNA-decapping activity of AtNUDT6, 7, 19, and 27. (A) Scheme of the experimental setup. ³²P labelled RNA was prepared by *in vitro* transcription with T7 RNA polymerase in the presence of the small molecules Np_nNs, NAD(H), CoA, or ADP-ribose. The side product, uncapped triphosphate RNA, was degraded by treatment with 5' polyphosphatase and TerminatorTM exonuclease. (B) A representative example of PAGE analysis of the RNA-decapping activity of AtNUDT6, 7, 19, and 27 on 5'-capped Ap₄A-RNA and NADH-RNA.

(Fig. S1 and S2, ESI^{\dagger}). AtNUDT6 and AtNUDT7 had a similar substrate specificity, with the Ap₄A-RNA being the best substrate for both of them (Fig. 1B).

In addition, AtNUDT6 also partially cleaved (less than 50%) Ap₃A-, Ap₅A- and Ap₄G-RNA. AtNUDT7 cleaved Ap_{2,3,5}A-RNA, Ap_{4,5}G-RNA, and (to some extent) Gp₄A-RNA (Table 1 and Fig. S3, ESI†). The main product of the Ap₄A-RNA-decapping reaction by AtNUDT6 and AtNUDT7 is a triphosphate (ppp)-RNA. AtNUDT27 cleaved Ap₄A-RNA into a mixture of ppp-RNA and monophosphate (p-)RNA (Fig. S4, ESI†). While AtNUDT6 and AtNUDT7 had quite wide substrate specificities, AtNUDT19 efficiently cleaved only the NADH-RNA, with the NAD-RNA being partially cleaved (Fig. 1B and Table 1 and Fig. S3, ESI†). Surprisingly, at a tenfold lower concentration (50 nM), AtNUDT27 still partially hydrolyzed the majority of capped RNA with a relatively high cleavage efficiency (60–90% after 1 h), except for Ap₃A-RNA (less than 30%) (Table 1 and Fig. S3, ESI†).

While some of the small molecules are known to be *in vitro* substrates of these NudiX enzymes, the entire set of recently discovered noncanonical caps has not yet been tested as free compounds. Therefore, we screened all the noncanonical RNA caps in their free form as substrates for the four NudiX proteins (Table 2 and Fig. S5–S8, ESI†). AtNUDT6 and AtNUDT7 had a similar substrate specificity on both the small molecules and

Table 1Cleavage of RNA capped with different small molecules in vitroby AtNUDT6 (500 nM), AtNUDT7 (500 nM), AtNUDT19 (500 nM), andAtNUDT27 (50 nM). Values are in % of cleaved capped RNA and represent amean of three independent replicates \pm standard deviations

AtNUDT	6	7	19	27		
	(500 nM)	(500 nM)	(500 nM)	(50 nM)		
5'-cap-RNA	(%)	(%)	(%)	(%)		
Ap ₂ A-	x	53 ± 10	х	73 ± 9		
Ap ₃ A-	48 ± 7	55 ± 7	х	26 ± 8		
Ap ₄ A-	80 ± 11	73 ± 5	х	87 ± 4		
Ap ₅ A-	15 ± 8	65 ± 17	х	90 ± 5		
Ap ₃ G-	x	х	х	74 ± 11		
Ap ₄ G-	35 ± 10	70 ± 20	х	85 ± 5		
Ap ₅ G-	x	61 ± 12	х	85 ± 5		
Gp ₃ G-	x	х	х	83 ± 5		
Gp ₄ G-	x	х	х	83 ± 1		
NAD-	x	x	20 ± 4	60 ± 7		
NADH-	x	x	57 ± 5	65 ± 10		
CoA-	x	х	х	82 ± 5		
Gp ₃ A-	x	х	х	80 ± 5		
Gp ₄ A-	x	30 ± 6	х	84 ± 5		
Gp ₅ A-	x	х	х	90 ± 3		
m ⁷ Gp ₃ A-	x	x	x	80 ± 7		
m ⁷ Gp ₃ G-	x	x	x	70 ± 6		
ADP-ribose	x	х	х	70 ± 2		
				1		
0–25%	26–50%	51-75%	76–100%			
	capped RNA cleaved					

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Table 2 Cleavage of different small molecules (400 μ M) *in vitro* by AtNUDT6 (500 nM), AtNUDT7 (500 nM), AtNUDT19 (500 nM), and AtNUDT27 (500 nM). Values are in % of cleaved substrate and represent a mean of three independent replicates \pm standard deviations

AtNUDT	6	7	19	27		
small	(500 nM)	(500 nM)	(500 nM)	(500 nM)		
molecule	(%)	(%)	(%)	(%)		
Ap ₂ A	80 ± 9	64 ± 7	16 ± 3	х		
Ap ₃ A	18 ± 5	18 ± 1	x	x		
Ap ₄ A	31 ± 1	40 ± 3	x	99 ± 0.5		
Ap ₅ A	х	31 ± 1	x	99 ± 0.5		
Ap ₃ G	х	x	x	х		
Ap ₄ G	20 ± 2	x	x	96 ± 5		
Ap ₅ G	х	x	x	89 ± 1		
Gp ₃ G	х	x	x	x		
Gp ₄ G	х	x	x	98 ± 1.5		
NAD	x	52 ± 1	x	х		
NADH	99 ± 1	83 ± 4	56 ± 7	х		
3'-dpCoA	53 ± 4	18 ± 6	x	х		
ADP-ribose	55 ± 10	45 ± 3	x	х		
				1		
0–25%	26–50%	51–75%	76–100%			
		small molecule cleaved				

the capped RNA. Surprisingly, their best substrate was not free Ap₄A, as expected based on the results from assays on capped RNA (Table 2 and Fig. S5–S8, ESI[†]) but rather Ap₂A and NADH, both of which have a diphosphate bridge. As in the case of capped RNA, AtNUDT19 hydrolyzed free NADH with the highest efficiency (Table 2 and Fig. S5–S8, ESI[†]). AtNUDT27 had a very high activity at 500 nM concentration and quantitatively cleaved all the free tetraphosphates (Ap₄A, Ap₄G, and Gp₄G) and Ap₅A (Table 2 and Fig. S5–S8, ESI[†]). To better observe the selectivity of the enzyme towards small free molecules, we decreased its concentration to 50 nM (Fig. S9, ESI[†]). Under these reaction conditions, AtNUDT27 preferred dinucleoside tetraphosphates (*i.e.* Ap₄A, Ap₄G, and Gp₄G) over other non-canonical RNA caps.

In addition, we compared Michaelis–Menten kinetic parameters for AtNUDT6, 7, and 27 and Ap₄A as a substrate (Fig. 2A). These parameters were already known for AtNUDT19 and NADH as a substrate.²² Although $K_{\rm M}$ values were comparable for all three enzymes, $V_{\rm max}$ and $K_{\rm cat}$ of AtNUDT27 were both significantly higher (P < 0,05) compared to AtNUDT6 and 7 (Fig. S11a and b, ESI†). This finding may explain observed behaviour of AtNUDT27 to be more efficient in cleaving free Ap₄A than AtNUDT6 and 7 (Table 2).

Next, we wanted to determine whether NudiX enzymes prefer free small molecules or capped RNA as substrates especially since this could reflect the situation *in vivo*. Due to the need to use a very large amount of RNA substrate for the entire series of tests, we compared only one set of substrates, free Ap₄A and Ap₄A-RNA. We observed the cleavage of both substrates at a concentration of 2.5 μ M and 15- and 60 minute time points.

Α	Protein	Substrate	к _м (µМ)	V _{max} (µM min ⁻¹)	K _{cat} (min ⁻¹)	K _{cat} /K _M (min ⁻¹ μM ⁻¹)
	AtNUDT6	Ap₄A	6.8 ± 1.5	0.5 ± 0.0	9.3 ± 0.2	1.4 ± 0.3
	AtNUDT7	Ap ₄ A	11.3 ± 4.9	0.4 ± 0.2	8.8 ± 3.5	0.8 ± 0.2
	AtNUDT27	Ap ₄ A	22.8 ± 10.4	3.6 ± 1.3	71.7 ± 25.6	3.2 ± 0.6

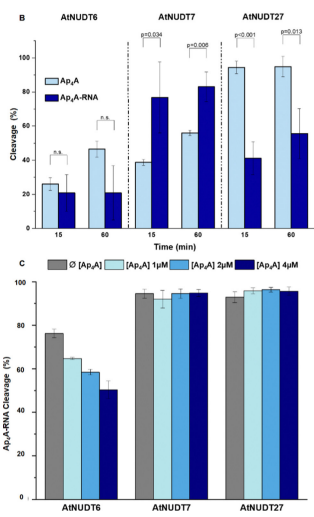


Fig. 2 (A) Kinetic parameters of AtNUDT6, AtNUDT7 and AtNUDT27 cleaving Ap₄A expressed as mean \pm standard deviation. (B) Cleavage efficiency of free Ap₄A (2.5 μ M) and Ap₄A-RNA (2.5 μ M) by AtNUDT6, AtNUDT7, and AtNUDT27 (50 nM) at 15- and 60 min time points. (C) Inhibition study of Ap₄A-RNA (1 μ M) cleavage by AtNUDT6 (500 nM), AtNUDT7 (500 nM) and AtNUDT27 (50 nM) with addition of 1, 2 or 4 μ M Ap₄A.

The cleavage of Ap_4A was analysed by HPLC, while decapping of Ap_4A -RNA was measured by quantification of corresponding bands of capped RNAs resolved on PAGE (Fig. 2B and Fig. S10, ESI†). Comparison of the substrates showed that AtNUDT27 preferred the small molecule (Ap_4A) and cleaved it more efficiently than Ap_4A -RNA under the chosen conditions. AtNUDT6 cleaved comparably well both substrates – Ap_4A and Ap_4A -RNA. AtNUDT7 had higher activity on Ap_4A -RNA than on free Ap_4A . This finding may explain the redundancy of NudiX enzymes observed *in vivo*, *i.e.* some enzymes are used by the cell to hydrolyze free small molecules, while others cleave non-canonically capped RNA.

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In order to understand whether free dinucleoside polyphosphates can inhibit the cleavage of capped RNA, we measured cleavage of Ap₄A-RNA (1 μ M) by AtNUDT6, 7 and 27 in presence of free Ap₄A (1,2 and 4 μ M). We observed that only AtNUDT6 activity was affected in dose-dependent manner. The activity of other two enzymes AtNUDT7 or AtNUDT27 was not affected by addition of free Ap₄A (Fig. 2C and Fig. S12, ESI†). The difference between AtNUDT6 and AtNUDT7, which have comparable kinetic parameters, could be explained by AtNUDT7 preference for Ap₄A-RNA instead of free Ap₄A as we demonstrated in Fig. 2B. On the other hand, AtNUDT27 showed almost 8 times higher K_{cat} for Ap₄A than AtNUDT6 and 7 suggesting that added free Ap₄A would be metabolised at high rate leaving enough enzymatic capacity to hydrolyse Ap₄A-RNA as well and thus the inhibition effect was not observed.

Conclusions

In light of recent discoveries of noncanonical RNA caps in various organisms, we focused on potential RNA-decapping NudiX enzymes from A. thaliana. We explored the cleavage properties of four enzymes, cytosolic AtNUDT6, AtNUDT7,²³ and chloroplastic AtNUDT19 and AtNUDT27.14,22 First, we tested whether they could cleave canonical and noncanonical caps from RNA using in vitro transcribed RNA capped with Np_nNs, CoA, NAD(H), m⁷Gp₃A(G), and ADP-ribose. We found that AtNUDT19 was very specific and cleaved NADH-RNA almost exclusively. Also, AtNUDT6 and AtNUDT7 were relatively specific and cleaved mainly Ap₄A-RNA. In contrast, AtNUDT27 effectively hydrolyzed almost all types of capped RNA at a tenfold lower concentration than the other tested enzymes, making it potentially a general RNA-decapping enzyme. As the in vitro substrates were not known for all of these enzymes, we also tested their substrate specificity on free small molecules that can form RNA caps (Np_nNs, 3'-dpCoA, NAD(H), $m^{7}Gp_{3}A(G)$, and ADP-ribose). Again, AtNUDT19 was very specific towards NADH, which is in accordance with role of AtNUDT19 in regulating NADH and NADPH levels in chloroplasts, as previously reported.²² Under our experimental conditions AtNUDT19 cleaved only free NADH but not NAD. Interestingly, when capped RNA was used as substrate, AtNUDT19 cleaved NADH-RNA and also partially NAD-RNA. This may indicate its role as an NAD/NADH-RNA-decapping enzyme. Contrary to its promiscuity in the case of capped RNA, AtNUDT27 hydrolyzed dinucleoside tetraphosphates and pentaphosphates almost exclusively. Our observed activity of AtNUDT27 against dinucleoside tetraphosphates is in contrast with previously reported results.²² There, authors observed cleavage of only Ap₅A but not of Ap₄A. Nevertheless, the presence of reducing agents may positively influence the activity of these enzymes.²⁴ Thus, this discrepancy may be explained by the presence of a higher concentration of reducing agent DTT (2 mM) in our reaction in comparison with their 1 mM DTT concentration. AtNUDT6 and AtNUDT7 mainly cleaved free molecules with the diphosphate bridge. In fact, both AtNUDT6 and AtNUDT7 were demonstrated to be cytosolic NADH pyrophosphohydrolases

participating in stress responses through modulation of intracellular NADH levels in *A. thaliana*.²⁵ Ap₄A is also known to be involved in stress signalling in many organisms, and because AtNUDT6, AtNUDT7, and AtNUDT27 efficiently cleaved both the free Ap₄A and the Ap₄A-RNA, we compared their preferences on cleavage of these substrates. We observed that AtNUDT27 and AtNUDT6 cleaved the free Ap₄A more efficiently, while AtNUDT7 favoured the Ap₄A-RNA. These differences in substrate specificities suggest that *in vivo* some NudiX proteins may function in degradation of free small molecules, while others serve as specific decapping enzymes, probably during regulated RNA decay.

It has been shown that diadenosine polyphosphates such as Ap₃A and Ap₄A behave like alarmones in *A. thaliana* and trigger a cascade of reactions to yield various protective compounds.¹³ Our experiments show that plant NudiX enzymes have RNA-decapping activity and that they can cleave various noncanonical RNA caps. This finding supports the notion that plant RNA also contains other unknown RNA caps in addition to the canonical m⁷G structure and NAD cap.^{3,26} Altogether, the existence of specific enzymes that cleave NADH- and Ap₄A-RNA suggest that these RNA caps may play an important role in plant RNA metabolism, signalling, or stress reaction.

Moreover, the specific cleavage properties of the individual NudiX enzymes characterized in this work can be used to develop capturing strategies for noncanonically capped RNA and thus help us to identify the sequence and features of these RNA species.

Materials and methods

Preparation of RNA

In vitro transcription with T7 RNAP. All chemicals were either purchased from Merck or Jena Biosciences and used without further purification. Oligonucleotides were purchased from Generi Biotech. In vitro transcription was performed using a modification of a previously published method,⁹ in a 25 µL mixture containing: 80 ng µL⁻¹ of template DNA (35A for Ap₂₋₅A, Gp₃₋₄A, NAD(H), CoA, ADP-ribose and m⁷Gp₃A, or 35G for Gp₃₋₄G, Ap₃₋₅G, and m⁷Gp₃G), 1 mM UTP, 1 mM CTP, 1 mM ATP, 0.8 mM GTP and 0.5 μ L α ³²P GTP (activity: 9.25 MBg in 25 μ L), 1.6 mM of Np_nNs (Ap₂₋₅A, Gp₃₋₄G, Ap₃₋₅G, m⁷Gp₃G, m⁷Gp₃A, ADP-ribose), or 8 mM of cofactors (3'-dpCoA, NAD and NADH), 5% DMSO, 0.12% triton X-100, 12 mM DTT, 4.8 mM magnesium chloride and $10 \times$ reaction buffer for T7 RNAP (40 mM Tris-HCl, 6 mM MgCl₂, 1 mM DTT, 2 mM spermidine, pH 7.9 at 25 °C) and 62.5 units of T7 RNAP. The mixture was incubated for 2 h at 37 °C.

DNAse I treatment. The DNA template was digested by DNAse I to obtain pure RNA. 25 μ L of the transcription mixture, 3 μ L of 10× reaction buffer for DNAse I (100 mM Tris-HCl, 25 mM MgCl₂, 5 mM CaCl₂, pH 7.6, supplied with the enzyme), and 4 units of DNAse I (NEB) were incubated at 37 °C for 60 min. The enzyme was heat deactivated at 75 °C for 10 min followed by immediate cooling on ice. All samples were purified on RNA mini Quick Spin Columns (Merck) for further use.

RNA 5'-**polyphosphatase treatment.** The products from the previous reaction were treated with 20 units of 5'-polyphosphatase (Epicentre) in the reaction buffer (50 mM HEPES-KOH, pH 7.5, 100 mM NaCl, 1 mM EDTA, 0.1% β -mercaptoethanol, and 0.01% Triton X-100) for 1 h at 37 °C.

TerminatorTM 5'-**phosphate-dependent exonuclease treatment.** Because of the incompatibility of the buffers, all samples were purified on RNA mini Quick Spin Columns before the reaction. The RNA was treated with 1 unit of TerminatorTM 5'-phosphate-dependent exonuclease (Epicentre) in reaction buffer A (supplied with the enzyme) at 30 °C for 1 h.

Purification of recombinant NudiX enzymes

A. thaliana NUDT6, 7, 19, and 27 genes were cloned into the pET26b vector with the C-terminal His tag (Novagen). Due to the problem with expressing the full-length AtNUDT27, it was cloned without the first 44 aa containing the chloroplast signal sequence. Proteins were overexpressed in Escherichia coli Rosetta DE3 cells grown for 16 hours in auto-inducing medium.²⁷ Cells were lysed by sonication in buffer A (20 mM Tris-HCl, pH 8.0, 300 mM NaCl, 10% glycerol and 5 mM β-mercaptoethanol) and recombinant proteins were purified on TALON Metal Affinity Resin (Clontech), eluted in buffer A supplemented with 250 mM of imidazol and fractionated by gel filtration chromatography on an ENrichTM SEC70 column (BioRad). The purified proteins were concentrated to 50 µM in a buffer containing 20 mM Tris-HCl, pH 8.0, 200 mM NaCl, 2 mM DTT, and 10% glycerol, flash frozen in liquid nitrogen, and stored at -80 °C. Protein concentration was determined at 280 nm using the molar extinction coefficient calculated on the basis of protein composition with the Prot-Param tool (https://web.expasy.org/protparam/) (Fig. S13, ESI⁺).

Cleavage of capped RNA by A. thaliana NudiX enzymes

The decapping assay was performed at 37 °C in 10 μ L reaction. To test the cleavage of the 5'-caps, the RNA samples were divided into two parts. The positive control contained 100 ng of the pure RNA (*in vitro* transcription, DNAse I treatment, purified on RNA mini Quick Spin Columns), 1 μ L of 10× buffer (100 mM Tris-HCl, pH 7.5, 1 M KCl, 20 mM MgCl₂, 20 mM MnCl₂, 20 mM DTT), and 500 nM of the recombinant AtNUDT6, 7, 19, and 50 nM AtNUDT27. Water was used as the negative control. The mixtures were incubated at 37 °C for 1 h and purified on RNA mini Quick Spin Columns. Then 3 μ L of the purified samples were mixed with 2× RNA loading dye (NEB) for analysis by 12% PAGE (600 V, 3.5 h). Typhoon FLA 9500 was used as a visualization imaging system.

Calculations of RNA decapping

The experiments were performed in triplicates, and the amount of capped RNA was calculated from PAGE analysis using the software ImageJ.²⁸ Areas under the peaks (AUC) corresponding to the 5'-capped RNA (Ar_p) in the control without enzyme and 5'-capped RNA (Ar_{cap}) in samples with enzyme were calculated. The percentage of 5'-capped RNA species was calculated according to: (Ar_{cap}/Ar_p) × 100.

Enzymatic assay and kinetics assay on small free molecules

The hydrolytic activities of AtNUDT proteins towards different Np_nNs (Ap₂₋₅A, Gp₃₋₄G, Ap₃₋₅G, m⁷Gp₃G, m⁷Gp₃A), 3'-dpCoA, ADP-ribose, and NAD(H) were performed using 500 nM of the AtNUDT6, 7, 19, and 27 and 400 µM of substrate in 10 mM Tris-HCl (pH 7.5), 100 mM KCl, 2 mM magnesium acetate, 2 mM DTT, at 37 °C. The enzyme was heat deactivated at 75 °C for 10 min followed by cooling on ice. The mixture was then analysed by monitoring absorbance at 259 nm and retention time compared to standards on high-performance liquid chromatography using a Kinetex C18 column (4.6 \times 150 mm, 5 μ m, Phenomenex) at a flow rate of 1 ml min⁻¹. The mobile phase consisted of 100 mM triethylammonium acetate pH 7 (buffer A) and Acetonitrile (buffer B). Separation was achieved using a 25 minute gradient: 100% A for 2 min; linear decrease to 95% A over 4 min; linear decrease to 70% A over 9 min; maintaining 70% A for 3 min; returning linearly to 100% A over 1 min. Chromatograms were integrated, and the percentage of substrate cleavage was calculated from corresponding AUC compared to the control without enzyme.

Kinetic parameter calculation

Kinetic parameters of AtNUDT enzymes were calculated from experiments using increasing concentrations of Ap₄A (0, 5, 10, 20, 30, 50, 70, and 100 μ M) and 50 nM of AtNUDT6, 7, and 27. The 500 μ L mixture was incubated at 37 °C, and aliquots of 100 μ L were collected at 0, 2.5, 5, 10, and 15 min and immediately heated to 75 °C for 10 min to stop the reaction. All samples were analysed using the HPLC method described above. Calculated initial reaction velocities were plotted against Ap₄A concentration and fitted to the Michaelis–Menten model to obtain $K_{\rm m}$ and $V_{\rm max}$ values using Origin software (Northampton, MA). Student *t*-test and analysis of variance (ANOVA) with *post hoc* Bonferroni analysis were performed in Microsoft Excel to assess significant differences between measured parameters where needed.

Inhibition test with Ap₄A for AtNUDT enzymes

To test the inhibition of AtNUDT6, AtNUD7 and AtNUDT27 enzymes in the presence of Ap₄A we measured the cleavage of Ap₄A-RNA at 37 °C in 10 μ L reaction using 1 μ M of purified Ap₄A-RNA and increased concentrations of Ap₄A (1, 2 and 4 μ M). The concentration used for AtNUDT6 and AtNUD7 was 500 nM while for AtNUDT27 was 50 nM. The reaction was stopped using 2× RNA Loading Dye (NEB).

Author contributions

M.-B.M., O.H., A.G., S.K. performed experiments. M.-B.M. and O.H. studied RNA and small molecule behaviour and designed experiments, A.G. and S.K. purified plant proteins. R.B. and O.N. designed kinetic assays. J.K. and H.C. designed experiments and coordinated study. M.-B.M., O.N., and H.C. wrote the manuscript.

Conflicts of interest

Authors declare no conflicts of interests.

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