

Biosynthesis of Thiamin Thiazole in Eukaryotes: Conversion of NAD to an Advanced Intermediate

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Abstract: Thiazole synthase catalyzes the formation of the thiazole moiety of thiamin pyrophosphate. The enzyme from *Saccharomyces cerevisiae* (THI4) copurifies with a set of strongly bound adenylated metabolites. One of them has been characterized as the ADP adduct of 5-(2-hydroxyethyl)-4-methylthiazole-2-carboxylic acid. Attempts toward yielding active wild-type THI4 by releasing protein-bound metabolites have failed so far. Here, we describe the identification and characterization of two partially active mutants (C204A and H200N) of THI4. Both mutants catalyzed the release of the nicotinamide moiety from NAD to produce ADP-ribose, which was further converted to ADP-ribulose. In the presence of glycine, both the mutants catalyzed the formation of an advanced intermediate. The intermediate was trapped with orthophenylenediamine, yielding a stable quinoxaline derivative, which was characterized by NMR spectroscopy and ESI-MS. These observations confirm NAD as the substrate for THI4 and elucidate the early steps of this unique biosynthesis of the thiazole moiety of thiamin in eukaryotes.

Introduction

Thiamin pyrophosphate is an essential cofactor in all living systems, where it plays an important role in amino acid and carbohydrate metabolism.^{1,2} Bacteria, fungi, and plants can biosynthesize this cofactor, whereas most other eukaryotes cannot. They need to obtain it as an essential vitamin from their diet (vitamin B₁). Thiamin was the first vitamin discovered and consists of a thiazole linked to a pyrimidine. The biosynthesis of thiamin involves the separate biosyntheses of the thiazole and pyrimidine moieties, which are then coupled.^{3–5} The biosynthesis of the thiazole moiety in prokaryotes is well-characterized and involves a complex oxidative condensation of 1-deoxy-D-xylulose-5-phosphate, glycine (or tyrosine), and cysteine, which is catalyzed by five different enzymes.³ In eukaryotes, the biosynthesis of the thiazole occurs by a different route, the details of which are only beginning to emerge.^{6–8}

Extensive mutagenesis in *Saccharomyces cerevisiae* has resulted in the identification of only one gene required for

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thiazole formation (THI4).⁹ We recently reported the characterization of an adenylated thiazole (ADT, **15**, Figure 1) tightly bound to the active site of THI4 (Figure 2, peak **D**).^{7,8} This identification suggested NAD **1** as the probable precursor to ADT **15** and provided key insights into the mechanism of thiazole biosynthesis in eukaryotes (Figure 1). Two other major adenylated metabolites were also found to be tightly bound to the active site (peaks **A** and **B**, Figure 2). These could not be characterized due to their instability. Peak **C** was always a minor component, and its abundance varied between different protein preparations. All attempts to reconstitute active THI4 by releasing the bound metabolites failed, making it impossible to experimentally confirm the novel proposal that NAD was the thiamin—thiazole precursor in *S. cerevisiae*.

In an attempt to generate a product-free enzyme able to catalyze the early steps of thiazole formation, several THI4 mutants were constructed based on the recently solved crystal structure.⁸ Here, we report the identification and characterization of two partially active mutants (C204A and H200N), which show informative activities. In the presence of glycine **5**, these mutants catalyzed the conversion of NAD **1** to one of the previously observed unstable THI4-bound metabolites (peak **A**, Figure 2) via the intermediacy of ADP-ribose **3** and ADP-

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Figure 1. Mechanistic proposal for the THI4 catalyzed formation of ADT, 15 (peak D, Figure 2).



Figure 2. Metabolites associated with THI4 that are released upon denaturing the protein. Peak **D** is ADT (15). Peaks **A** and **B** were unstable and degraded during the isolation process.

ribulose **4**, confirming NAD as the thiazole precursor. The unstable metabolite, corresponding to peak **A**, could be efficiently trapped with ortho-phenylenediamine (oPDA, **16**), suggesting that it contains a 1,2-diketone or equivalent functionality. A mechanistic proposal consistent with these observations is described (Figure 1).

Results

Mutagenesis Studies on THI4 to Identify Product-Free Partially Active Mutants. Extensive mutagenesis on wild-type THI4 was performed in an attempt to identify partially active mutants. Each mutant was analyzed for tightly bound metabolites by HPLC. The mutants were also incubated with NAD 1, and the reaction mixture was analyzed by HPLC. Mutants D238A, C205S, C204A, and H200N were shown to degrade NAD 1 (peak G) to produce ADPr 3 (peak E) and nicotinamide



Figure 3. Conversion of NAD (1, peak G) to ADPr (3, peak E), nicotinamide (2, peak H), and an unknown species (peak F) catalyzed by the partially active mutant C204A. Reaction 1 represents the HPLC analysis of the reaction mixture containing the mutant C204A and NAD. Control 1 represents an experiment where the mutant C204A, with no NAD, was treated identically to reaction 1. It represents the protein-bound small molecules (A, B, and D). Control 2 represents an enzyme-free reaction mixture, where NAD was incubated in the same reaction buffer and was treated identically to reaction 1. The condition (heat) applied to denature the protein caused minor hydrolysis of NAD to produce ADPr and nicotinamide. The ADPr and nicotinamide peaks were identified by comigration with ADPr and nicotinamide standards.

2 (peak **H**) as products (Figure 3 and Table 1). All the other mutants (H237N, H237A, R301Q, R301A, P304A, E97A, E97Q, and D207A) did not show any activity toward NAD and did not contain bound metabolites. These observations confirm that NAD **1** is one of the substrates for THI4 and demonstrate that the hydrolysis of NAD **1** to form ADPr **3** and nicotinamide **2** is the first step in the THI4 catalyzed biosynthesis of ADT **15**.

Table 1. Catalytic Properties of THI4 Mutants

mutant	bound metabolites	products (no. of turnovers after overnight incubation)
D238A	wild-type (A, B,	ADPr 3, ADPrl 4, and nicotinamide
	and D)	2 (<0.1)
C205S	none	ADPr 3, ADPrl 4, and nicotinamide
		2 (<0.1)
C204A	<wild-type (a,="" b,<="" td=""><td>ADPr 3, ADPrl 4, and nicotinamide</td></wild-type>	ADPr 3, ADPrl 4, and nicotinamide
	and D)	2 (~0.8)
H200N	only peak A	ADPr 3, ADPrl 4, and nicotinamide
		2 (~0.8)
other	none	none
mutants ^a		

^a H237N, H237A, R301Q, R301A, P304A, E97A, E97Q, and D207A.



Figure 4. Conversion of ADP-ribose (3, peak E) to ADP-ribulose (4, peak F) catalyzed by the THI4 C204A mutant and ribose-5-phosphate isomerase (RPi). Reactions 1 and 2 represent the reactions of ADP-ribose catalyzed by RPi and C204A, respectively. Control 1 represents a sample of the C204A THI4 protein that was denatured and analyzed for bound metabolites. It shows the enzyme-bound metabolites A-D. Control 2 shows the ADP-ribose control where ADP-ribose was incubated with the same reaction buffer and treated identically to reactions 1 and 2.

Enzyme Catalyzed Generation of ADP-Ribulose (ADPrl 4) from ADP-Ribose (ADPr 3). Incubation of the partially active mutants (D238A, C205S, C204A, and H200N) with NAD also resulted in the formation of a new adenylated species (Figure 3, peak F) along with ADPr 3. The same adenylated species could be generated directly from ADPr 3 (peak E), by incubating ADPr with these mutants (Figure 4). A standard ADP-ribulose sample (ADPrl, 4) was shown to comigrate with this new adenylated metabolite (Figure 4). Attempts at the direct isolation and characterization of the peak F compound failed, as the compound decomposed during the isolation process. However, it was possible to reduce the peak **F** compound with NaBD₄, and the resulting stable deuterated reduction product was characterized by NMR. These observations establish the isomerization of ADPr 3 to ADPrl 4 as the second step in the THI4 catalyzed biosynthesis of ADT 15 from NAD 1.

Glycine-Dependent Conversion of ADPr 3 and ADPrl 4 to a New Species. Prolonged incubation of NAD 1 or ADPr 3 with the C204A or H200N mutants resulted in a small increase in the intensity of one of the three Thi4-bound adenylated metabolites (peak A, Figure 2). This increase varied with different protein preparations. However, in the presence of glycine 5, incubation of these mutants with NAD 1 or ADPr 3 always resulted in a significant increase in the intensity of peak A and a concomitant decrease in the intensities of peak E (ADPr 3) and peak F (ADPrl 4) when compared to a similar reaction mixture without glycine (Figure 5). This observation indicates



Figure 5. C204A mediated conversion of NAD (1, peak G) to the proteinbound metabolite A, in the presence of glycine 5. HPLC chromatograms: reaction 1 represents a reaction mixture containing C204A and NAD (no glycine). Reaction 2 represents a reaction mixture containing C204A, NAD, and glycine. Control 1 represents the experiment where a sample of C204A was incubated in the absence of both NAD and glycine (to reveal proteinbound metabolites A, B, and D). Control 2 represents the experiment where NAD and glycine were incubated in the same reaction buffer, in the absence of protein, under the same conditions as for reactions 1 and 2. Peaks for ADPr (3, peak E), ADPrl (4, peak F), nicotinamide (2, peak H), NAD (1, peak G), and protein-bound molecules (peaks A, B, and D) are labeled. The glycine-dependent decrease in the intensity of E and F and a concomitant increase in the intensity of A are illustrated.



Figure 6. Trapping of the THI4-bound metabolite **A** with oPDA (16, peak **I**) as a new species **J**. The blue trace represents the reaction mixture where the metabolites released from THI4 (by heat denaturing) were incubated with an excess of oPDA at room temperature. The purple trace represents a control reaction where oPDA was not added (only THI4-bound metabolites).

that glycine is important for the further conversion of ADPrl 4 to the advanced intermediate represented by peak A in Figure 2.

Characterization of the Peak A Compound by Trapping with ortho-Phenylenediamine (oPDA). The attempts at direct isolation and characterization of the compound eluting as peak **A** (Figure 2) failed due to its instability. However, incubation with ortho-phenylenediamine (oPDA, **16**) resulted in its clean conversion to a stable new species, which was purified by HPLC (peak **J**, Figure 6). The other protein-bound metabolites were stable under the mild trapping conditions used. This new species had absorption maxima at 260 and 321 nm, suggesting the presence of an adenine moiety as well as a putative quinoxaline moiety (Figure 7A). Treatment of this derivative with nucleotide pyrophosphatase (pptase) followed by HPLC analysis revealed the formation of AMP and another species (peak **K**) that had an absorption maximum only at 321 nm (Figure 7B). This suggests a general structure where a



Figure 7. (A) UV-vis spectrum of the quinoxaline adduct, **J**, formed as a result of the trapping of the THI4-bound metabolite **A** by oPDA, **16** (peak **I**, Figure 6). (B) Purified peak **J** compound can be cleaved with nucleotide pyrophosphatase (pptase) to produce AMP and another species **K** (**19**) that has only one absorption maximum at 321 nm. The blue and purple traces represent the same reaction mixture containing the purified quinoxaline adduct **J** and pptase, observed at 261 and 321 nm, respectively. The green trace represents a control reaction, where pptase was not added, observed at 261 nm. (C) Proposed diketone (or equivalent) trapping with oPDA **16** and pyrophosphate hydrolysis of **18** with pptase.



Figure 8. Structure 20 represents the partial structure deduced from the NMR experiments and the observations suggesting the presence of a quinoxaline moiety. The five carbon atoms attached to ADP, derived from ADP-ribose, are labeled as u, v, w, y, and z. Structure 21 represents the actual structure deduced from additional negative mode ESI-MS experiments and other observations. Structure 21 must be generated from the 1,2-diketone containing species 22 (or equivalent) and oPDA (16).

quinoxaline moiety is attached to an ADP unit (structure 18, Figure 7C).

Analysis of ¹H. (¹H.¹H)-dqfCOSY, HMOC, and HMBC NMR spectra obtained for the peak J compound confirmed the presence of a 5'-phosphorylated adenosine (Figure 9, ¹H NMR of J). Additionally, a three proton spin system representing a -CHOH-CH₂-OP fragment (CH₂ at 4.24 and 4.35 ppm, m; CH at 5.40 ppm, dd), a methyl singlet (2.73 ppm), and a 1,2disubstituted benzene (7.77, m, 1H; 7.65, m, 1H; 7.62-7.58, m, 2H) were identified. In HMBC spectra, these four aromatic protons showed correlations to two quaternary carbons (139.4 and 139.8 ppm), one of which also showed a weak correlation to the singlet methyl protons. Furthermore, the HMBC spectra showed strong correlations of the methyl protons and the CHOH methine proton to two additional quaternary aromatic carbons (y, 153.06 ppm, and w, 152.91 ppm in structure **20**, Figure 8). As the cross-peaks representing the interaction of these two carbons with methyl protons in the HMBC spectrum overlapped considerably, the exact chemical shift values for these two carbons were determined from a ¹³C NMR spectrum. Spatial proximity of the methyl group to the CHOH-CH₂-OP spin system was confirmed by a ROESY spectrum that revealed a strong NOE between the methyl and the methine protons (on v) and a weaker NOE between the methyl and the methylene protons (on u). In summary, the analyses of the NMR spectroscopic data, coupled with the observations implying the presence of a quinoxaline moiety, suggested the partial structure 20 (Figure 8). The combined results of the mass spectrometric analysis (described next), the required presence of an ADP unit and partial structure 20, unambiguously identify 21 as the structure of the oPDA derivative (peak J) of the compound corresponding to peak A. In negative mode ESI-MS spectra (Figure 10), both the monoanionic (m/z = 612) and the dianionic (m/z = 305.6) species were visible, and extensive fragmentation analyses performed on both the monoanionic and the dianionic species further corroborated structure 21. Isolation of the nucleotide pyrophosphatase cleavage product of 21 (peak K) by HPLC, followed by ¹H NMR (Figure 11) analysis, revealed the expected quinoxaline 23 serving as a further confirmation for the structure 21.



Figure 9. ¹H NMR spectrum of the purified quinoxaline adduct (peak J). The peaks have been assigned as shown in the structure.



Figure 10. Negative mode ESI-MS analysis of the quinoxaline derivative 21. The m/z for the monoanionic species is 612 and for the dianionic species is 305.6. Peaks with m/z = 477, 426.1, 346.1, and 185 result from the fragmentation of 21 under spraying conditions.

Discussion

Recombinant wild-type THI4 copurifies with a set of tightly bound adenylated metabolites (peaks A-D, Figure 2). One of these was shown to be an adenylated thiazole derivative (ADT 15, peak D).⁷ This discovery provided the first clues to the enzymology of thiamin-thiazole formation in eukaryotes and suggested that the precursor to ADT 15 might be an adenylated pentose/pentulose such as ADP-ribose 3. It was reasonable to propose NAD 1 as the precursor to ADT 15 because NAD could be converted to ADPr 3 by a well-characterized reaction similar to that involved in protein ADP-ribosylation,¹⁰ and the conversion of ADPr 3 to ADT 15 could be accomplished using reasonable chemistry (Figure 1). In addition, sequence analysis of THI4 predicted a nucleotide binding motif. However, this proposal could not be tested using recombinant THI4 overexpressed in Escherichia coli or in S. cerevisiae because the tightly bound metabolites in the active site of this protein inhibited catalysis. All attempts to prepare active enzyme by releasing the bound metabolites, under a variety of denaturing conditions, failed.

We therefore carried out extensive mutagenesis on THI4 to identify a partially active mutant with empty active sites. This could either arise from a weaker affinity of a mutant toward its reaction products or from a catalytically impaired mutant that is unable to make the late intermediates or the product. The early steps of the complex reaction sequence leading to ADT **15** could be characterized if such a mutant retained some activity. The recently published crystal structure of THI4 enabled us to select a set of conserved active site residues for mutagenesis (Table 2 and Figure 12).

The D238A and C205S mutants showed very weak activities toward NAD 1 (peak G), the proposed substrate, to produce ADPr 3 (peak E) and nicotinamide 2 (peak H) as products (unpublished data). This same activity was much stronger for the C204A and H200N mutants (Figure 3). FAD and NADP were also tested as substrates for these two active mutants, but no reaction was detected. These observations confirm the hypothesis that NAD 1 is one of the substrates for THI4 and identify the hydrolysis of NAD 1 to form ADPr 3 as the first step of thiazole biosynthesis in eukaryotes.

In addition to the formation of ADPr **3** and nicotinamide **2** from NAD **1**, the partially active mutants also catalyzed the formation of a new adenylated species (peak **F**, Figure 3). The production of the same unknown species was observed when the C204A and H200N mutants were treated with ADPr (Figure 4). To determine if this intermediate was ADP-ribulose (ADPrl,

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Figure 11. ¹H NMR spectrum of the HPLC-purified quinoxaline phosphate unit (23) obtained from 21 by its hydrolysis catalyzed by nucleotide pyrophosphatase (pptase).

Table 2. Summary of All THI4 Mutants Constructed

amino acid	possible role(s)	mutants
E97	binding adenylated intermediates product via H-bonding	E97A, E97Q
	to the ribose part of the ADP moiety	
H200 ^a	catalysis of thiazole formation	H200N
$C204^{a,b}$	catalysis of thiazole formation	C204A
C205 ^a	Only conserved cysteine; catalysis of thiazole formation;	C205S
	possibly important in the sulfur transfer	
$D207^a$	catalysis of thiazole formation	D207A
H237	catalysis of thiazole formation	H237N, H237A
D238	catalysis of thiazole formation	D238A
R301	binding advanced intermediates via H-bonding to the carboxylate	R301Q, R301A
	unit coming from glycine; other catalytic role	
P304	providing structural rigidity to a loop spanning the active site	P304A

^{*a*} These conserved residues are part of a flexible loop that spans the active site of a different monomer in the octameric structure (see Figure 12). ^{*b*} This cysteine residue is replaced by a conserved serine in most THI4 orthologs.

4), the next intermediate on our proposed pathway (Figure 1), a reference sample of this compound was prepared by the ribose-5-phosphate isomerase (RPi) catalyzed isomerization of ADPr to ADPrl.¹¹ This reference compound comigrated with the compound corresponding to peak **F** generated by treating ADPr with either of the C204A or H200N mutants of THI4 (Figure 4). To further confirm the identity of the peak **F** compound as ADPrl **4**, it was reduced with NaBD₄ to produce a stable deuterated reduction product, which was isolated by HPLC and characterized by NMR. These observations establish the isomerization of ADPr to ADPrl as the second step in the THI4 catalyzed biosynthesis of ADT from NAD.

In the absence of glycine, the C204A and H200N mutants catalyzed low levels of conversion of ADPrl 4 to the compound corresponding to peak A after prolonged incubation. The amount of the peak A compound formed over the background was always low and varied between experiments using different protein preparations. In the presence of glycine, however, the same conversion occurred at a much higher level (Figure 5). This reaction, for the first time, led to the reconstitution of the

biosynthesis of one of the previously observed unstable THI4bound adenylated metabolites (peak A, Figure 2).

Direct characterization of the compound corresponding to peak A could not be achieved due to its instability. The mechanistic proposal outlined in Figure 1 suggested a possible presence of a 1,2-diketone or a 1,2-ketoimine functionality in this intermediate compound (e.g., structures 9 and 10). Facile decomposition of the isolated peak A compound to produce ADP as the major product is consistent with this prediction (unpublished results). Treatment of the reaction mixture with ortho-phenylenediamine (oPDA, 16), which traps reactive 1,2diketones as stable quinoxalines,^{12,13} resulted in the selective and efficient conversion of the peak A compound to a stable adduct, which was isolated and its structure determined as the quinoxaline 21 (discussed in the Results). While this assignment does not unambiguously determine the structure of the compound generating peak A, it is consistent with structure 9. The observation that the conversion of ADPrl 4 to the peak A

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Figure 12. Stereoview of the active site of THI4 with the ADT (15) carbon atoms labeled in cyan. Two separate monomers contribute to the ADT binding site. Carbon atoms from one monomer are labeled in green and those from the other are labeled in yellow. His200 is on a disordered region and is not shown.

compound is facilitated by glycine is consistent with the mechanism outlined in Figure 1, in which imine formation facilitates deprotonation reactions at the C3 carbon of the ribose. In this mechanism, imine formation between ADPrl 4 and glycine 5 gives 6, imine deprotonation at C3 gives 7, which eliminates water and tautomerizes to give 9, the proposed intermediate giving rise to peak A (Figure 1). Model studies have demonstrated that imine deprotonation is about 10^8 times faster than the corresponding ketone deprotonation.^{14,15} This pK_a enhancement strategy is used by many enzymes including the prokaryotic thiazole synthase.¹⁶

Much still remains to be understood about this protein and its function in the eukaryotic cell. None of the mutant THI4 enzymes are able to catalyze multiple turnovers presumably because the products in each case bind tightly to the enzyme. In addition, it was not possible to advance the thiazole biosynthesis beyond the compound corresponding to peak A (9). This could be due to the fact that the correct sulfur donor has not yet been identified. The following observations reported on this enzyme also require further investigation: THI4 appears to be involved in mitochondrial DNA damage prevention and other stress related pathways.¹⁷⁻²¹ THI4 in Neurospora crassa

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is highly expressed (1.5% of the entire proteome) in the exponential growth phase and seems to be post-translationally modified and forms a stable complex with a cyclophilin (cis/ trans-proline isomerase).22,23

The identification of reaction intermediates is a valuable tool in the investigation of an enzyme mechanism. In most cases, this involves extensive kinetic characterization and insightful design of substrate analogues to block the reaction at intermediate stages. THI4 allows a remarkable departure from this traditional strategy for investigating enzyme mechanisms. This protein copurifies with a set of tightly bound metabolites that identify its function as well as provide a series of molecular snapshots that reveal the mechanism of the complex reaction sequence that it catalyzes.

Experimental Procedures

All the chemicals and snake venom nucleotide pyrophosphatase were purchased from Sigma-Aldrich Corporation unless otherwise mentioned. For all the HPLC analysis, a Supelco LC-18-T (150 mm × 4.6 mm, 3 μ m i.d.) reverse phase column was used. The ribose-5-phosphate isomerase plasmid was a kind gift from Prof. Sherry L. Mowbray (Swedish University of Agricultural Sciences).

Construction of the THI4 Mutants. All the THI4 mutants were constructed by the Cornell Protein Purification Facility. Standard methods were used for DNA manipulations.24,25 Plasmid DNA was purified with the Qiagen Miniprep kit. E. coli strain MachI (Invitrogen) was used as a recipient for transformations during plasmid construction and for plasmid propagation and storage.

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Table 3. Primers Used in the Mutagenesis Process				
mutant	mutagenesis primer sequence, top strand	restriction site		
C205S	CCCAAGCTCACGGTACTCAATGTTCCATGGACCCTAACG	StyI		
R301Q	GGATGGATTAAACCAAATGGGACCAACTTTTGGAGCTATGGC	BsmFI		
R301A	CTGGATGGATTAAACGCCATGGGTCCAACTTTTGGAGCTATGGC	NcoI		
H237N	GGTGTCATTTTATCCACTACCGGCAATGATGGTCCATTTGGTGC	BsrDI		
H237A	GGTGTCATTTTATCCACTACCGGAGCGGATGGTCCATTTGGTGC	BsrBI		
D238A	GGTGTCATTTTATCCACTACCGGTCATGCGGGACCATTTGGTGC	<i>Bsm</i> FI		
C304A	GGATTAAACCGTATGGGCGCCACTTTTGGAGCTATGGC	KasI		
E97A	CTTGAAGGTTTGTATTATCGCGAGTTCAGTTGCACCAGGTGG	NruI		
E97Q	CTTGAAGGTTTGTATTATCCAGAGCTCAGTTGCACCAGGTGG	HgiAI		
C204A	CCAAGCTCACGGTACTCAAGCTTGCATGGACCCTAACG	HindIII		
H200N	GTTAGTTACCCAAGCTAACGGCACTCAATGTTGCATGG	N/A		
D207A	GGTACTCAATGTTGCATGGCCCCTAACGTAATTGAATTG	N/A		

 Table 4.
 Screening Primers Used Only for H200N and D207A

 Mutants
 Primers Used Only for H200N and D207A

mutant	screening primer sequence
H200N	TAGTTACCCAAGCTAACGGC
D207A	GGTACTCAATGTTGCATGGC

Site-directed mutagenesis was performed on pThi 4.28^7 by a standard PCR protocol using *Pfu*Turbo DNA polymerase per the manufacturer's instructions (Invitrogen) and *Dpn*I (New England Biolabs) to digest the methylated parental DNA prior to transformation.

For the C205S, R301Q, R301A, H237N, H237A, D238A, C304A, E97A, E97Q, and C204A mutants, primers were designed to introduce or remove a diagnostic restriction enzyme site to facilitate screening for a mutated clone. Only clones producing the anticipated restriction pattern were sequenced. For the H200N and D207A mutants, a third primer was designed to screen for the presence of the mutant by colony PCR with an appropriate vector specific primer. Only clones that produced a PCR product were sequenced. In every case, the mutagenesis primer pair consisted of the primer whose sequence and its reverse complement are in Tables 3 and 4.

Overexpression and Purification of THI4 and Its Mutants. E. coli BL21(DE3) containing the overexpression plasmid (containing a full length clone of THI4, or its mutant, in the pET28 vector) was grown in LB medium containing kanamycin (40 μ g/mL) with shaking at 37 °C until the OD₆₀₀ reached 0.6. Isopropyl- β -D-thiogalactopyranoside (IPTG) was then added (final concentration = 2 mM), and cell growth was continued at 15 °C for 16 h. The cells were then harvested by centrifugation, and the resulting cell pellets were stored at -20 °C until needed. To purify the protein, the cell pellets from 1 L of culture were resuspended in 20 mL of lysis buffer (10 mM imidazole, 300 mM NaCl, 50 mM NaH₂PO₄, pH 8) and lysed by sonication (Heat Systems Ultrasonics model W-385 sonicator, 2 s cycle, 50% duty). The resulting cell lysate was clarified by centrifugation, and the THI4 protein was purified on Ni-NTA resin following the manufacturer's (Qiagen) instructions. After elution, the protein was desalted using a PD-10 column (Amersham) pre-equilibrated with 50 mM potassium phosphate buffer, pH 8.0. All the THI4 mutant proteins expressed well and were well-behaved except for the E97A and E97Q mutants. These two mutants mostly precipitated immediately after purification.

HPLC Analysis of THI4 and Its Mutants for Bound Metabolites. The purified enzyme was denatured by heat (100 °C, 40 s) and then rapidly cooled on dry ice. The precipitated protein was removed by centrifugation. The supernatant was filtered through a 10 kDa MW cut-off filter (Microcon YM-10, Millipore), and 100 μ L of the filtrate was analyzed by reversed phase HPLC. The following linear gradient (method 1, in some experiments minor modifications to the method were made to obtain better resolution at certain parts of the chromato-gram) was used at 1 mL/min flow rate: solvent A is water, solvent B is 100 mM KP_i pH 6.6, and solvent C is methanol; 0 min 100% B, 5 min 10% A 90% B, 8 min 25% A 60% B 15% C, 14 min 25% A 60%

B 15% C, 19 min 30% A 40% B 30% C, 21 min 100% B, and 30 min 100% B. The absorbance at 220, 261, and 320 nm was monitored. The UV-vis spectra of the resolved components were analyzed by an inline diode-array UV detector.

Activity Assays with the Mutants. The purified mutants (100 μ M) were incubated with 300 μ M NAD or ADP-ribose (in the presence or absence of 300 μ M glycine) for 10 h at room temperature (25 °C). Control reactions were set up with the enzyme (without the substrate) or with the substrate(s) in the reaction buffer. Subsequently, the enzyme in the reaction mixture was heat denatured (100 °C, 40 s), and the precipitated enzyme was removed by centrifugation. The supernatant was filtered through a 10 kDa MWCO filter (Microcon YM-10, Millipore). The control reactions were treated identically. The filtrate was analyzed by HPLC as described previously.

Preparation of ADP-Ribulose. *E. coli* ribose-5-phosphate isomerase was overexpressed in *E. coli* strain BL21(DE3) containing the overexpression plasmid (containing a full length clone of the *E. coli* ribose-5-phosphate isomerase gene in the pET15b vector) in LB medium as described previously. The protein was purified on Ni-NTA resin following the manufacturer's (Qiagen) instructions and desalted into 50 mM Tris-HCl, pH 7.5 containing 100 mM NaCl. The 50 μ M protein was incubated with 250 μ M ADP-ribose for 5 h. The reaction mixture was heat denatured (100 °C, 40 s), and the precipitated protein was removed by centrifugation. The supernatant was filtered through a 10 kDa MWCO filter (Microcon YM-10, Millipore), and the filtrate was analyzed by HPLC as described previously.

Characterization of NaBD₄ Reduced Peak F Compound. The ADP-ribose (1 mM) was incubated with the C204A THI4 mutant (100 μ M) overnight at room temperature. The reaction mixture was heat denatured (100 °C, 40 s), and the precipitated protein was removed by centrifugation. The supernatant was filtered through a 10 kDa MWCO filter (Microcon YM-10, Millipore). From the filtrate thus obtained, the peak F compound (putative ADPrl) was purified by HPLC. The pooled fractions were combined (~20 mL), and directly reduced with NaBD₄ (2 mM, 20 min, room temperature) and neutralized with 1 M HCl, the entire sample was subjected to HPLC purification, and the collected fractions were lyophilized. The lyophilized product was dissolved in 0.5 mL of 100% D₂O (Sigma) and was relyophilized. The residue thus obtained was dissolved in about 0.25 mL of 100% D₂O, and the solution was placed in a Shigemi NMR tube (standardized for D₂O). NMR data were acquired on a Varian INOVA 600 MHz instrument equipped with a 5 mm triple-gradient inverse-detection HCN probe. The NaBD₄ reduction yielded two species observed by HPLC with an approximately 1:1 ratio, both of which were isolated. One of these was easily identified by NMR as the NaBD4 reduction product of ADPrl. The NMR analysis of the other product revealed very similar characteristics and most likely is the borate adduct of the reduction product.

Trapping of the Peak A Compound with ortho-Phenylenediamine (Analytical Sample). A total of 500 μ L of a concentrated wild-type THI4 sample was heat denatured to release all the bound metabolites,

and the precipitated protein was removed by centrifugation. The supernatant was filtered through a 10 kDa MWCO filter (Microcon YM-10, Millipore). To 150 μ L of the filtrate was added a freshly prepared aqueous solution of ortho-phenylenediamine to a final concentration of 4 mM. The mixture was incubated on ice for 15 min and analyzed by HPLC.

Purification of the Trapped Species. All the THI4 protein obtained from a 3 L culture (~150 mg) was concentrated to 6 mL (Amicon Utra-4, Millipore, 10 kDa cut-off) and divided into ten 600 μ L aliquots. Each aliquot was heat denatured (100 °C), and the precipitated protein was removed by centrifugation. The supernatants were combined and filtered through a 10 kDa MW cut-off filter (Amicon Ultra-4, Millipore). ortho-Phenylenediamine (oPDA) was added to the filtrate to a final concentration of 4 mM, and the mixture was incubated for 15 min on ice. A different HPLC method was developed for purification of the adduct because the buffer system had to be compatible with the ESI-MS analysis. The following linear gradient was used at a 1 mL/min flow rate (method 2): solvent A is ammonium acetate buffer (20 mM, pH 7) and 1% methanol and solvent B is 10% ammonium acetate buffer (20 mM, pH 7) and 90% methanol; 0 min 100% A, 3 min 100% A, 12 min 70% A 30% B, 14 min 70% A 30% B, 16 min 100% A, and 25 min 100% A. The peak corresponding to the adduct was collected, frozen with dry ice, and lyophilized. The product residue was redissolved in water and checked for purity by HPLC. It was found to be stable when kept cold (on ice) or frozen.

NMR Spectroscopy and Negative Mode ESI-MS Analysis of the Purified Adduct. The lyophilized product was dissolved in 0.5 mL of 100% D₂O (Sigma) and was relyophilized. The residue thus obtained was dissolved in about 0.25 mL of 100% D₂O, and the solution was placed in a Shigemi NMR tube (standardized for D₂O). The ¹³C NMR data was acquired on a Varian INOVA 500 MHz instrument. All other NMR data were acquired on a Varian INOVA 600 MHz instrument equipped with a 5 mm triple-gradient inverse-detection HCN probe.

For ESI-MS analysis, the lyophilized product was dissolved in 100 μ L of water. Right before spraying into the instrument (Esquire ion

trap instrument, Bruker), this aqueous solution was mixed with 100 μ L of methanol containing 0.1% triethyl amine. The sample was analyzed in the negative ion mode, and the peaks thus obtained were isolated and fragmented.

Cleavage of the Quinoxaline Containing Adduct 21 by Nucleotide Pyrophosphatase and the Isolation of the Quinoxaline Fragment. To a 100 μ L solution of the purified quinoxaline-ADP adduct (21) was added 200 μ g of lyophilized snake venom nucleotide pyrophosphatase (EC 3.6.1.9, Sigma) and MgCl₂ to a final concentration of 1 mM (20 mM Tris-HCl, pH 8.0). A control reaction was set up with only MgCl₂ and no enzyme. Both the samples were incubated at 37 °C for 2 h and filtered (10 kDa MW cut-off filter), and the filtrates were analyzed by HPLC (method 1 described previously). The same reaction was repeated on a large scale, and the quinoxaline fragment (23) peak was collected (method 2 described previously) and lyophilized. The product thus obtained was dissolved in 0.25 mL of 100% D₂O, and the solution was placed in a Shigemi NMR tube (standardized for D2O) and analyzed by ¹H NMR. ¹H NMR (600 MHz, D₂O): δ 8.12-8.06 (m, 1H, aromatic), δ 8.01–7.96 (m, 1H, aromatic), δ 7.87–7.79 (m, 2H, aromatic), δ 5.46 (dd, 1H, J = 3.66, 7.57 Hz), δ 4.16–4.03 (m, 2H), δ 2.83 (s, 3H).

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Supporting Information Available: NMR data for the characterization of ADPrl **4**, ESI-MS fragmentation analysis, 2-D NMR data, summary of chemical shifts, and NMR correlations for compound **21**. This material is available free of charge via the Internet at http://pubs.acs.org.

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