

Cite this: *RSC Chem. Biol.*, 2022, 3, 227Received 6th August 2021,  
Accepted 25th October 2021

DOI: 10.1039/d1cb00160d

rsc.li/rsc-chembio

## Trapping and structural characterisation of a covalent intermediate in vitamin B<sub>6</sub> biosynthesis catalysed by the Pdx1 PLP synthase<sup>†</sup>

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**The Pdx1 enzyme catalyses condensation of two carbohydrates and ammonia to form pyridoxal 5-phosphate (PLP) via an imine relay mechanism of carbonyl intermediates. The I<sub>333</sub> intermediate characterised here using structural, UV-vis absorption spectroscopy and mass spectrometry analyses rationalises stereoselective deprotonation and subsequent substrate assisted phosphate elimination, central to PLP biosynthesis.**

Pyridoxal 5-phosphate (**10**, PLP, vitamin B<sub>6</sub>) is an essential cofactor in all living systems where it plays a central role in the stabilisation of carbanions adjacent to amines primarily in amino acid metabolism.<sup>1</sup> An enzyme complex consisting of Pdx1 and Pdx2 catalyses the conversion of glutamine, ribose 5-phosphate (R5P) and glyceraldehyde 3-phosphate (G3P) to PLP.<sup>2,3</sup> Protein structures show two conserved phosphate binding sites in Pdx1, spaced 22 Å apart.<sup>4–6</sup> During catalysis, the ribose phosphate group occupies the P1 phosphate binding site,<sup>5,7–9</sup> while the product PLP phosphate group binds in P2.<sup>9,10</sup> The Pdx1-associated glutaminase Pdx2 generates ammonia,<sup>2,3</sup> which diffuses through an internal channel towards the Pdx1 P1 site<sup>4</sup> where it then reacts with the covalent R5P adduct. This most intriguing catalytic step leads to formation of the double imine chromophoric intermediate I<sub>320</sub> (Fig. 1).

In this communication we study the Arabidopsis Pdx1 enzyme; the plant has three *pdx1* alleles,<sup>2,11</sup> of which we studied

Pdx1.3. Lys98 of Pdx1.3 was previously shown to form an imine with the R5P aldehyde,<sup>9,12</sup> enabling a series of sugar modifications in the ribose tailoring site that culminate in the addition of Lys166 to generate I<sub>320</sub><sup>9,12</sup> (**8**). The Lys98/Lys166 crosslinking intermediate forms after phosphate elimination;<sup>13</sup> however, the detailed mechanism is not understood. The vacated P1 site can then be occupied by the phosphate of the second substrate G3P, defining dual specificity for the Pdx1 active site.<sup>9</sup> Incorporation of G3P leads to formation of PLP<sup>13–16</sup> that is proposed to migrate to the PLP/P2 product site.<sup>9</sup> Our previous study trapped an intermediate prior to phosphate elimination in the Lys166Arg exchange variant of the Pdx1.1 paralog (5LNT, see ESI<sup>†</sup>), suggested to form from R5P modification with ammonia.<sup>9</sup> We now constructed the lysine/arginine exchange variant in the Pdx1.3 enzyme where structures of several other intermediates are available (reviewed in ESI<sup>†</sup>).<sup>9,12</sup> We characterised the intermediate that formed after addition of R5P and ammonia in this variant. For its characteristic absorbance we named the adduct I<sub>333</sub>, determined its mass and demonstrated ammonia incorporation. The crystallographic structure suggests key catalytic residues for I<sub>333</sub> formation, allowing us to propose a mechanism for deprotonation and phosphate elimination. The data rationalise how subsequent imine formation gives I<sub>320</sub>.

We used UV-vis absorption spectroscopy to characterise adducts formed by Pdx1.3 and its Pdx1.3<sup>K166R</sup> variant. R5P and (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> were added to Pdx1.3 (10 mM and 200 mM final concentrations respectively) leading to accumulation of the chromophoric I<sub>320</sub> intermediate over an hour at RT (**8**, λ<sub>max</sub> = 314 nm). Further addition of G3P (20 mM final concentration) leads to the formation of PLP in Pdx1.3 (λ<sub>max</sub> = 412 nm, Fig. 2). Pdx1.3<sup>K166R</sup> did not catalyse formation of the I<sub>320</sub> intermediate (compare<sup>14</sup>); instead yielding a product with λ<sub>max</sub> = 333 nm (Fig. 2). Addition of G3P to the I<sub>333</sub> complex did not lead to any significant changes in spectral signatures demonstrating that Pdx1.3<sup>K166R</sup>:I<sub>333</sub> does not convert to PLP. We therefore conclude that Lys166 is required for I<sub>320</sub> formation, consistent with recent structural proposals,<sup>9,12</sup> and that I<sub>320</sub> is a required intermediate for PLP biosynthesis.

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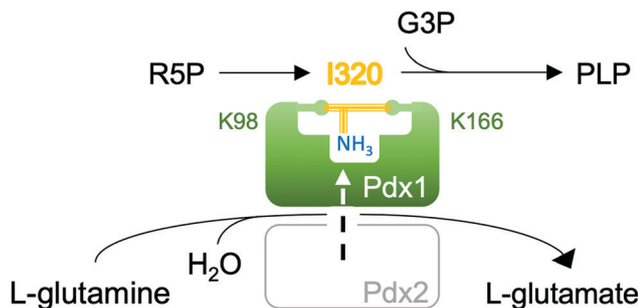
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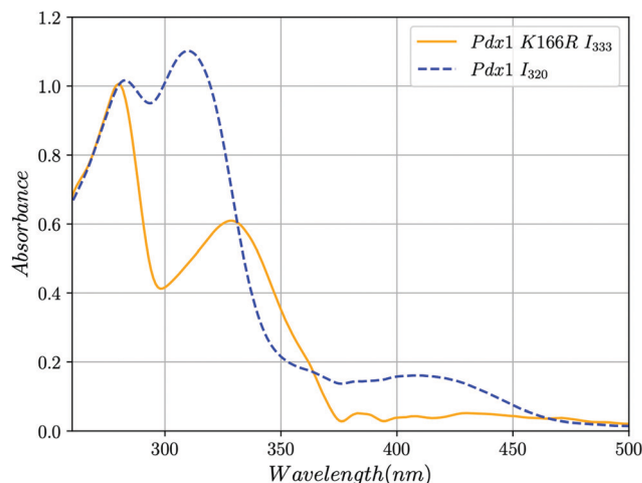
<sup>†</sup> Electronic supplementary information (ESI) available: Experimental procedures and MS analysis (PDF). See DOI: 10.1039/d1cb00160d





**Fig. 1** PLP synthase consists of the synthase Pdx1 and the glutaminase Pdx2 enzymes. Ammonia produced by glutamine hydrolysis is transferred to the Pdx1 active site where it is incorporated into the covalent ribose 5-phosphate (R5P) adduct. The I<sub>320</sub> intermediate formed bridges the catalytic Lys98 and Lys166 side chains. Addition of glyceraldehyde 5-phosphate (G3P) leads to conversion of I<sub>320</sub> to pyridoxal 5-phosphate (PLP).

To characterise intermediate formation by mass spectrometry, Pdx1.3 and Pdx1.3<sup>K166R</sup> were treated with substrates as indicated in Table 1. After tryptic digestion, the peptide mixtures were analysed by LC-ESI-TOF-MS. Comparison of the predicted and observed mass for the substrate free samples demonstrated that both Pdx1.3 and Pdx1.3<sup>K166R</sup> contained QAVTIPVMAK<sub>98</sub>, unmodified at lysine 98 ( $[M + H]^+ = 1057.6$  Da, Table 1 and Fig. S1, S2, ESI<sup>†</sup>). Addition of R5P to both samples leads to a mass difference of 212 Da, as seen in the samples of Pdx1.3 + R5P (Table 1 and Fig. S3, ESI<sup>†</sup>) and Pdx1.3<sup>K166R</sup> + R5P (Table 1 and Fig. S4, ESI<sup>†</sup>). For Pdx1.3<sup>K166R</sup>, the extra 16 Da on the Pdx1.3<sup>K166R</sup> peptide relative to Pdx1.3 is consistent with the oxidation of Met96 to its sulfoxide, which can occur during sample preparation.<sup>17</sup> The observation of the 212 Da adduct (**1**) is consistent with earlier observations.<sup>3,15</sup> From previous structural and MS analysis, it is known that I<sub>320</sub> is formed after the addition of R5P and ammonia<sup>14,16</sup> and crosslinks Lys98 and Lys166.<sup>9,12</sup> In the present study, instead of trapping I<sub>320</sub>, treatment of Pdx1.3 with R5P and ammonia followed by trypsin gave unmodified QAVTIPVMAK and TK<sub>166</sub>GEAGTGNIEAVR with a mass adduct of 96 Da (Table 1 and Fig. S5, ESI<sup>†</sup>), consistent with elimination of Lys98 from the I<sub>320</sub> adduct to give (**9**). In the key experiment, treatment of Pdx1.3<sup>K166R</sup> with R5P and ammonia followed by trypsin gave the I<sub>333</sub> modified QAVTIPVMAK<sub>98</sub> peptide with a mass adduct of 194 Da (Table 1 and Fig. S6, ESI<sup>†</sup>). The Pdx1.3<sup>K166R</sup> TR<sub>166</sub>GEAGTGNIEAVR peptide was not modified, demonstrating that it was not attached to I<sub>333</sub>. On the *Bacillus subtilis* Pdx1<sup>K149R</sup> variant (equivalent to Pdx1.3<sup>K166R</sup>) a 12-fold lower



**Fig. 2** UV-vis absorption spectra showing the formation of the I<sub>320</sub> and I<sub>333</sub> intermediates in solution at RT. Blue dashed line, addition of R5P, ammonia and G3P to Pdx1.3 results in the formation of the chromophoric intermediate I<sub>320</sub> ( $\lambda_{\max} = 315$  nm) and PLP ( $\lambda_{\max} = 412$  nm). Yellow line, addition of the same compounds to Pdx1.3<sup>K166R</sup> results in the formation of intermediate I<sub>333</sub> ( $\lambda_{\max} = 333$  nm) that does not convert to PLP after addition of G3P. The peak at  $\lambda_{\max} \sim 280$  nm represents protein. The discontinuity at 360 nm is caused by changeover of deuterium and halogen lamps. Spectra were smoothed and background correction for Rayleigh scattering was applied to account for protein aggregation as described in ESI.<sup>†</sup>

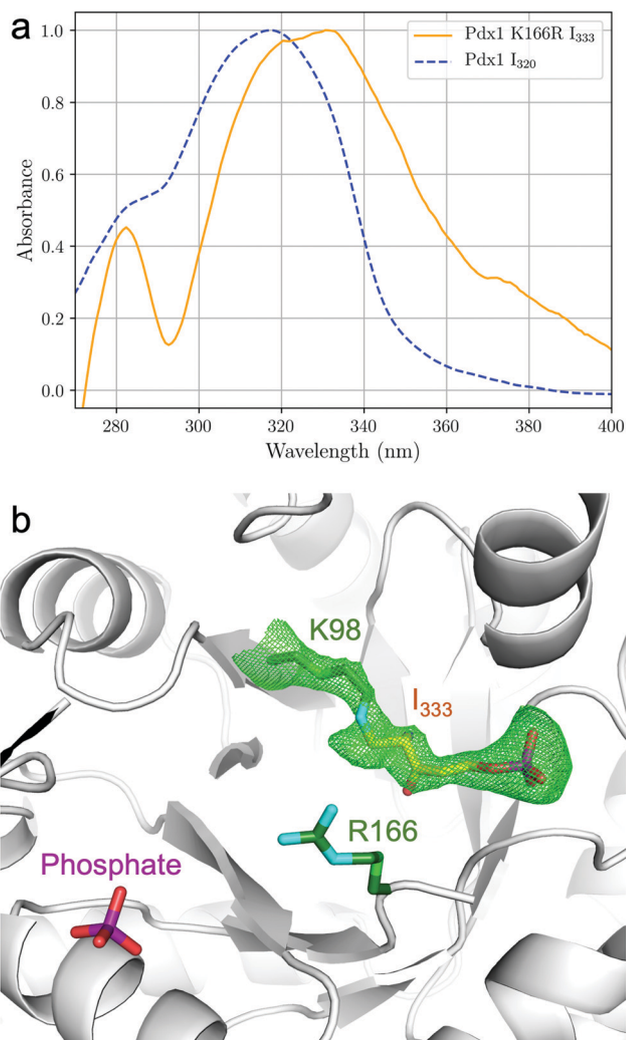
formation of I<sub>320</sub> was reported,<sup>14</sup> but the observation may indeed reflect formation of I<sub>333</sub> in the bacterial enzyme.

The I<sub>333</sub> adduct was further investigated by crystallographic analysis.<sup>‡</sup> Equivalent to our earlier study of the I<sub>320</sub> intermediate,<sup>9</sup> we soaked Pdx1.3<sup>K166R</sup> crystals with R5P and (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> where we used this protocol to generate the Pdx1.1<sup>K166R</sup>:preI<sub>320</sub> complex (5LNT, see ESI<sup>†</sup>). UV-vis spectra were taken *in crystallo*<sup>18</sup> to confirm formation of I<sub>333</sub> (Fig. 3a) and corrected for scattering at the crystal-solvent/air interface (see ESI,<sup>†</sup> 18,19). The observed absorbance at  $\lambda_{\max} = 333$  nm was consistent with the in-solution data (Fig. 2). The crystal structure shows I<sub>333</sub> as an adduct (**6**) between lysine 98 and C1 from R5P (Fig. 3b). Since the spectra indicated ammonia incorporation, the intermediate has been modelled with nitrogen in place of the oxygen at C2. This is consistent with previous NMR data showing that nitrogen incorporation precedes formation of the I<sub>320</sub> intermediate,<sup>15</sup> and is also consistent with the mass spectrometry analysis presented here (Table 1).

**Table 1** MS characterisation of covalent intermediates in Pdx1.3 and Pdx1.3<sup>K166R</sup> variant proteins after addition of R5P or R5P and ammonia substrates. Masses were determined for peptides bearing Lys98 or Lys166 that are covalently modified in intermediate formation. The molecular mass of the modification is determined as the difference between the observed mass ( $[M + H]^+_{\text{obs}}$ ) and the predicted mass ( $[M + H]^+_{\text{pred}}$ ) of the unmodified peptide

Subs	Pdx1.3	Pdx1.3 <sup>K166R</sup>
None	QAVTIPVMAK <sub>98</sub> $[M + H]^+_{\text{pred}} = 1057.6074$ $[M + H]^+_{\text{obs}} = 1057.6076$	QAVTIPVMAK <sub>98</sub> $[M + H]^+_{\text{pred}} = 1057.6074$ $[M + H]^+_{\text{obs}} = 1057.6085$
+R5P	QAVTIPVMAK <sub>98</sub> AR $[M + H]^+_{\text{pred}} = 1284.7457$ $[M + H]^+_{\text{obs}} = 1496.7546$ Modification ( <b>1</b> ) = 212.0089	QAVTIPVM(O)AK <sub>98</sub> AR $[M + H]^+_{\text{pred}} = 1300.7406$ $[M + H]^+_{\text{obs}} = 1512.7436$ Modification ( <b>1</b> ) = 212.003
+R5P +NH <sub>3</sub>	TK <sub>166</sub> GEAGTGNIEAVR $[M + H]^+_{\text{pred}} = 1515.8125$ $[M + H]^+_{\text{obs}} = 1611.8326$ Modification ( <b>9</b> ) = 96.0201	QAVTIPVMAK <sub>98</sub> AR $[M + H]^+_{\text{pred}} = 1284.7457$ $[M + H]^+_{\text{obs}} = 1478.7561$ Modification ( <b>6</b> ) = 194.0104

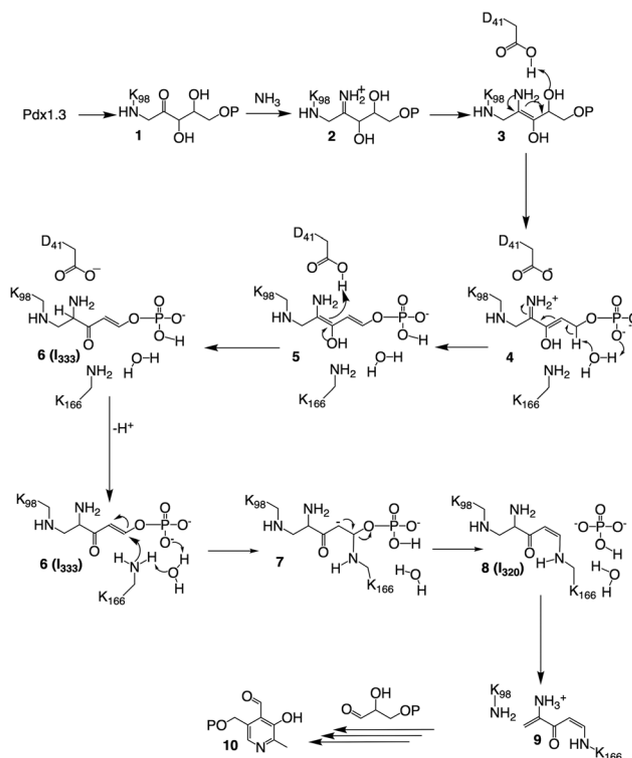




**Fig. 3** *In crystallo*<sup>18</sup> UV-vis absorption spectroscopy and structure of Pdx1.3<sup>K166R</sup>:I<sub>333</sub>. (a) UV-vis spectra normalised for the absorption maxima of the chromophores, collected from crystals of Pdx1.3:I<sub>320</sub> (blue dashed line) and Pdx1.3<sup>K166R</sup>:I<sub>333</sub> (orange line) after soaking with R5P and ammonia. Spectra were smoothed and background correction for light scattering at the crystal interface was applied as described in ESI†. (b) Structure of the ribose tailoring site of Pdx1.3<sup>K166R</sup>:I<sub>333</sub> showing bound I<sub>333</sub> (PDB:7NHE). The side chains of Lys98 and Arg166 are shown in stick format (carbon atoms green, nitrogen cyan). The I<sub>333</sub> adduct is distinguished by color (carbon atoms orange, oxygen red, nitrogen blue, phosphorous purple). The electron density is contoured at 2.5σ and was calculated using a simulated annealing protocol with Lys98–I<sub>333</sub> residues omitted from map calculation (see ESI†). A phosphate ion is seen in the PLP site.

The novel crystal structure and mass-spectrometry data lead to a revised mechanistic proposal for the formation of I<sub>320</sub> based on detection and characterisation of I<sub>333</sub>, as shown in Fig. 4. R5P forms an imine with Lys98, which then undergoes a tautomerisation to give **1**. Imine formation with ammonia gives **2**, which tautomerises to **3**. Elimination of water, assisted by acid catalysis by Asp41 gives **4**.

The crystal structure is particularly helpful for interpretation of catalytic steps required in I<sub>333</sub> formation (Fig. S8, ESI†). The base catalyst for the tautomerisation of **4** to **5** is a conserved



**Fig. 4** Mechanistic proposal for the formation of the I<sub>320</sub> and I<sub>333</sub> intermediates.

water molecule bound by hydrogen bonding to the amide NH of Gly230 and Val232 (β7α7 loop), the carbonyl oxygen of Phe250 (β8α8 loop) and the closest substrate phosphate oxygen, shown in Fig. 5. The active site bears a structure similar to a protease oxyanion hole.<sup>20</sup> Interactions with the amide NH of Gly252, Ser253 (β8α8 loop), Gly231 (β7α7 loop) and Gly170 (β6α6 loop) stabilise the negatively charged R5P phosphate, which itself polarises a water molecule activating it as a base in substrate-assisted catalysis; phosphate groups are frequently observed to perform such function.<sup>21</sup> The specific structure ensures stereospecificity of the C5 proS deprotonation reaction.<sup>13</sup> Tautomerisation of **5** to **6** is assisted by proton transfer from Asp41.

In the native enzyme, a conjugate addition of Lys166 is proposed to **6** to give **7**, where the substrate phosphate and the conserved water serve as catalytic base. Elimination of phosphate from carbanion **7** completes formation of I<sub>320</sub>, which is converted to **9** by elimination of Lys 98 in the presence of G3P to yield PLP, as previously described.<sup>9,15</sup>

## Conclusions

Our data are consistent with I<sub>333</sub> being on-pathway and serving a function in substrate-assisted catalysis during phosphate elimination. The active site structure allows rationalising stereoselective loss of the C5 hydrogen of R5P. The structure of I<sub>333</sub> elucidates the catalytic mechanism for the formation of the novel lysine crosslinking intermediate I<sub>320</sub> and is central to our understanding of the Pdx1-catalysed PLP biosynthesis.<sup>9,12</sup>



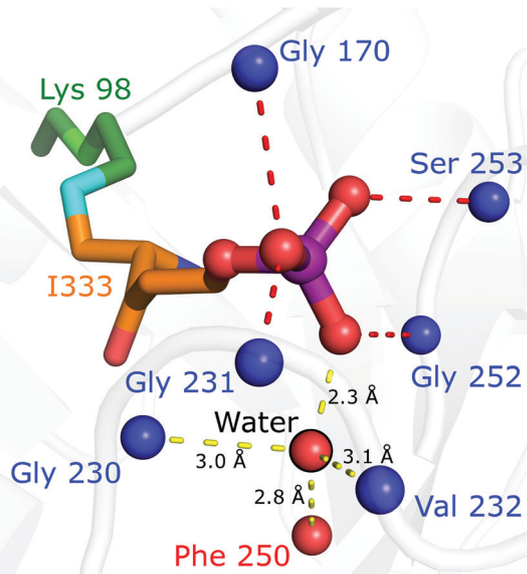


Fig. 5 The structure of the Pdx1.3<sup>K166R</sup>:I<sub>333</sub> complex (PDB:7NHE) argues for substrate assisted catalysis and provides an explanation for stereo-selective deprotonation. A water, conserved in position across different PLP synthase structures and proposed to act in catalysis, is shown as red sphere. Positions of intermediate coordinating main chain nitrogen/oxygen atoms are shown as blue/red spheres. The covalent I<sub>333</sub> adduct is shown in stick format (lysine carbon atoms green, I<sub>333</sub> carbon atoms orange, nitrogen blue, oxygen red, phosphorous purple).

## Author contributions

MJR, NG and IT conducted research, RAB, AR, YZ, TB and IT analysed data, YZ, TB, GE and SE were involved in data curation, IT and TB wrote the manuscript; all authors have reviewed and edited the manuscript.‡

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

We thank S. Findlow and C. Holes at the Macromolecular Crystallisation Facility, Biological Sciences, University of Southampton for support with protein crystallisation. We thank the staff at beamline ID23-1 and at the *icOS* Lab of the European Synchrotron Radiation Facility for their support with collection of X-ray diffraction and UV-vis absorption spectroscopy data. We thank Nicolas Caramello at the ESRF for help with processing of data obtained at the *icOS* Lab.

## Notes and references

‡ Protein structures were deposited with the PDB under accession codes 7NHE for the Pdx1.3<sup>K166R</sup>:I<sub>333</sub> complex and 7NHF for Pdx1.3<sup>K166R</sup>.

§ MJR and RAB were supported by joint studentships between Diamond Light Source and the University of Southampton. This work used the

*icOS* platform of the Grenoble Instruct-ERIC center (ISBG; UMS 3518 CNRS-CEA-UGA-EMBL) within the Grenoble Partnership for Structural Biology (PSB), supported by FRISBI (ANR-10-INBS-05-02) and GRAL, financed within the University Grenoble Alpes graduate school (Ecoles Universitaires de Recherche) CBH-EUR-GS (ANR-17-EURE-0003). NG and TPB were supported by the Robert A. Welch Foundation (A-0034). No competing financial interests have been declared.

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