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Structural basis for the magnesium-dependent activation of transketolase from *Chlamydomonas reinhardtii*

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Abstract

Background: In photosynthetic organisms, transketolase (TK) is involved in the Calvin-Benson cycle and participates to the regeneration of ribulose-5-phosphate. Previous studies demonstrated that TK catalysis is strictly dependent on thiamine pyrophosphate (TPP) and divalent ions such as Mg^{2+} .

Methods: TK from the unicellular green alga *Chlamydomonas reinhardtii* (CrTK) was recombinantly produced and purified to homogeneity. Biochemical properties of the CrTK enzyme were delineated by activity assays and its structural features determined by CD analysis and X-ray crystallography.

Results: CrTK is homodimeric and its catalysis depends on the reconstitution of the holo-enzyme in the presence of both TPP and Mg^{2+} . Activity measurements and CD analysis revealed that the formation of fully active holo-CrTK is Mg^{2+} -dependent and proceeds with a slow kinetics. The 3D-structure of CrTK without cofactors (CrTK_{apo}) shows that two portions of the active site are flexible and disordered while they adopt an ordered conformation in the holo-form. Oxidative treatments revealed that Mg^{2+} participates in the redox control of CrTK by changing its propensity to be inactivated by oxidation. Indeed, the activity of holo-form is unaffected by oxidation whereas CrTK in the apo-form or reconstituted with the sole TPP show a strong sensitivity to oxidative inactivation.

Conclusion: These evidences indicate that Mg^{2+} is fundamental to allow gradual conformational arrangements suited for optimal catalysis. Moreover, Mg^{2+} is involved in the control of redox sensitivity of CrTK.

General significance: The importance of Mg^{2+} in the functionality and redox sensitivity of CrTK is correlated to light-dependent fluctuations of Mg^{2+} in chloroplasts.

Keywords:

Transketolase; *Chlamydomonas reinhardtii*; magnesium; thiamine pyrophosphate; crystal structure; CD spectroscopy

1. Introduction

Transketolase (TK) is a ubiquitous enzyme operating in the non-oxidative branch of the pentose phosphate pathway (PPP) and in the Calvin-Benson cycle (CBC) in photosynthetic organisms [1-4]. It catalyzes the transfer of a glycolaldehyde fragment from a ketose donor onto an aldose acceptor. D-xylulose-5-phosphate (X5P), fructose-6-phosphate (F6P) and sedoheptulose-7-phosphate (S7P) are typical donor substrates for TK, while glyceraldehyde-3-phosphate (G3P), D-ribose-5-phosphate (R5P) and D-erythrose-4-phosphate (E4P) are typical acceptor substrates [1,2]. Hence, within the CBC, TK catalyzes the conversion of fructose-6-phosphate/sedoheptulose-7-phosphate and glyceraldehyde-3-phosphate into erythrose-4-phosphate/ribose-5-phosphate and xylulose-5-phosphate, contributing to the regeneration of ribulose-5-phosphate [5]. All possible reactions are reversible and require the presence of both thiamine pyrophosphate (TPP) and divalent metal ions as TK cofactors. In the three-dimensional structure of TK from *Saccharomyces cerevisiae*, a Ca^{2+} ion is stabilized by TPP, the conserved residues Asp157, Asn187, and Ile189 and a water molecule [6,7]. Earlier studies on the same enzyme indicated that Ca^{2+} , Mg^{2+} , Mn^{2+} , Co^{2+} and Ni^{2+} can replace each other without affecting the maximal catalytic activity [8]. Although several three-dimensional structures from different sources have been solved, the chloroplastic TK from *Zea mays*, containing Mg^{2+} and TPP in the active site, is the only one belonging to a photosynthetic organism reported to date [9].

Several CBC enzymes (sedoheptulose-1,7-bisphosphatase, SBPase; phosphoribulokinase, PRK; glyceraldehyde-3-phosphate dehydrogenase, GAPDH; fructose-1,6-bisphosphatase, FBPase; and phosphoglycerate kinase, PGK) have a low activity in the dark and are activated in the light through reduction of regulatory disulfide bonds by the ferredoxin-thioredoxin system [10,11]. This system, composed of ferredoxin, ferredoxin-thioredoxin reductase (FTR) and thioredoxin (TRX), converts a photosynthetic signal received from photosystem I-reduced ferredoxin into a thiol signal that is transmitted to TRX through the action of FTR [12,13]. Under illumination, TRX becomes reduced and catalyzes the reduction of regulatory disulfide bonds controlling the activation state of CBC target enzymes [5,14].

TK is a potential candidate for redox regulation, being involved in multiple metabolic pathways as the connection between PPP and glycolysis, carbon fixation, and the biosynthesis of coenzymes, fatty acids, amino acids, glutathione and vitamins. However, there is no direct evidence that TK activity might be modulated by redox regulatory mechanisms [15], although proteomic surveys suggested that this enzyme could be a potential target of TRX [16-19].

In this work, we focused on the structural and biochemical characterization of chloroplastic TK from the green microalga *Chlamydomonas reinhardtii* (CrTK). In order to study the properties of this key enzyme, CrTK was heterologously expressed in *E. coli* and purified by metal affinity chromatography. Biochemical features of the enzyme were defined and its structure was determined by X-ray crystallography. Regulation by oxidative modifications and Mg^{2+} concentration were analyzed by activity assays and circular dichroism (CD). As observed for other TKs, we found a strong influence of Mg^{2+} on the reconstitution of the active holo enzyme. Our results suggest that fluctuations of the Mg^{2+} concentration have a strong impact on the CrTK activity and participate in the redox control of the enzyme by changing its propensity to be inactivated by oxidation.

2. Materials and Methods

2.1 Materials

All chemicals were obtained from Sigma-Aldrich unless otherwise specified. Desalting columns (NAP5 and PD-10) were purchased from GE Healthcare. Crystallization plates and silanized cover slips were purchased from Molecular Dimensions.

2.2 Plasmid construction for expression of CrTK and EcE4PDH in Escherichia coli

The cDNA-encoding chloroplastic transketolase from *C. reinhardtii* (CrTK) was amplified by PCR using a forward primer introducing an *NdeI* restriction site at the start codon (5'-AACGTGCATATGGCTCAGGCTGCCCCGCT-3'); and a reverse primer introducing a *BamHI* restriction site downstream of the stop codon (5'-ACCATGGGATCCTTAGTGCTGCAGGGTGGC-3'). CrTK is encoded by the gene at locus Cre02.g080200 and appears to be the only TK in *C. reinhardtii*. CrTK was cloned in a modified pET-3c vector containing additional codons upstream of the *NdeI* site to express a His-tagged protein with seven N-terminal histidines. This modified vector (pET-3c-His) was generated by restriction with *XbaI* and *NdeI* of pET-3c to insert the following sequence: 5'-TCTAGAAATAATTTTGTTTAACTTTAAGAAGGAGATATACAATGCACCACCACCA**CCACCACCATATG**-3' (*NdeI* italicized, start codon underlined and seven His codons in bold). The recombinant CrTK contains 692 residues with an estimated molecular weight of 75,169.6 Da; it starts at the N-terminus with the introduced MHHHHHHHM peptide followed by the mature protein

sequence (*i.e.* upon removal of the chloroplast targeting sequence), beginning with Ala36 (Fig. S1). The plasmid encoding erythrose-4-phosphate dehydrogenase from *Escherichia coli* (Ece4PDH) was used as in [20].

2.3 Protein purification

Recombinant CrTK was produced in *E. coli* with the pET-3c-His/BL21 expression system. Bacteria were grown at 37 °C in LB medium supplemented with 50 µg mL⁻¹ ampicillin; the expression of the protein was induced with 0.2 mM isopropyl-β-D-thiogalactopyranoside (IPTG) at 25 °C for 16-18 h when the culture reached an Abs₆₀₀ of 0.5. Cells were then harvested by centrifugation, re-suspended in 20 mM Tris-HCl (pH 7.9), 0.5 M NaCl, 5 mM imidazole (binding buffer), and disrupted by a double passage in a French Press Cell set at 1000 PSI. RNase and DNase (10 µg mL⁻¹) were then added to the suspension and let to digest nucleic acids for 20 min at room temperature. Residual cell debris were removed by centrifugation at 30,000 × *g* for 45 minutes. The supernatant was then applied onto a Ni²⁺ NTA column (Chelating Sepharose Fast Flow resin, GE Healthcare) pre-equilibrated with binding buffer. The column was washed with the same buffer containing increasing concentrations of imidazole (25, 60 and 100 mM) to remove contaminants. Finally, CrTK was eluted with binding buffer supplemented with 250 mM imidazole. Eluted fractions were desalted using PD10 columns (GE Healthcare) pre-equilibrated with 50 mM Tris-HCl (pH 7.9), 1 mM EDTA. Fractions were analyzed by SDS-PAGE and those containing pure CrTK concentrated by ultrafiltration in Centricon cartridge (Amicon, 10 kDa cut-off) and stored at -80 °C. The protein concentration was determined experimentally based on the UV absorption at 280 nm (*A*₂₈₀); the molar extinction coefficient of CrTK ($\epsilon_{280} = 89,270 \text{ M}^{-1} \text{ cm}^{-1}$) was calculated assuming that cysteine residues are not involved in disulfide bonds [21]. Ece4PDH was purified following the same protocol described above, except that its expression was induced by addition of 0.2 mM IPTG at 30 °C for 16-18 h when the culture reached an Abs₆₀₀ of 0.5. The concentration of Ece4PDH was determined by UV spectroscopy ($\epsilon_{280} = 32,430 \text{ M}^{-1} \text{ cm}^{-1}$).

2.4 Liquid chromatography-mass spectrometry analysis

The molecular weight of CrTK was evaluated by LC-ESI-MS analysis. The chromatographic analysis was performed on an Agilent 1200 HPLC system (Walbronn, Germany) using a monolithic CIMac C4 Analytical column (5 mm × 5.3 mm I.D.; BIA Separations, Ljubljana, Slovenia). Mobile phases A [water–acetonitrile-FA (99:1:0.1, v/v/v)] and B [water–acetonitrile-FA (2:98:0.1, v/v/v)] were used

to develop a gradient. The optimized mobile phase gradient was set as follows: A–B from (80:20, v/v) to (50:50, v/v) in 6 min; from (50:50, v/v) to (20:80, v/v) in 1 min; (20:80, v/v) for 1 min. The column was equilibrated with the mobile phase composition of the starting conditions for 3 min before the next injection. The flow rate was set at 0.5 mL/min. The injection volume was 2 μ L corresponding to 0.7 μ g of protein solubilized in 5 mM ammonium bicarbonate buffer at pH 7.9. MS analysis was carried out on a Q-ToF Micro quadrupole time-of-flight (Q-TOF) hybrid analyzer (Micromass, Manchester, UK) with a Z-spray electrospray ion source (ESI). The ESI-Q-TOF source temperature was set at 100 °C, the capillary voltage at 3.3 kV and the cone voltage at 35 V. The scan time was set at 2.0 s and the inter scan time at 0.1 s. The cone gas flow was set at 120 L h⁻¹ and the desolvation gas at 500 L h⁻¹. Chromatograms were recorded in total ion current (TIC), within 800 m/z and 2000 m/z . The CrTK baseline-subtracted spectrum (m/z 1,300–2,000) was deconvoluted onto a true mass scale using the maximum entropy (MaxEnt1)-based software supplied with MassLynx 4.1 software. Output parameters were set as follows: mass range 20,000–100,000 Da; resolution 2 Da/channel. The uniform Gaussian model was used, with 0.5 Da width at half height.

2.5 Gel filtration

Gel filtration analysis was performed on a Superdex 200 HR10/300 GL column (GE Healthcare) connected to an ÅKTA Purifier system (GE Healthcare), previously calibrated with standard proteins, as described in [22]. The column was equilibrated with 50 mM Tris-HCl (pH 7.5), 150 mM KCl and 1 mM EDTA. The volume of loaded CrTK samples was 0.25 mL at a concentration of 1 mg mL⁻¹ (12 μ M). The protein was eluted at a flow rate of 0.5 mL min⁻¹. A control run was conducted using the tetrameric cytoplasmic glyceraldehyde-3-phosphate dehydrogenase from *Arabidopsis thaliana* (isoform 1, AtGAPC1) [23].

2.6 Reconstitution of the holoenzyme and activity assay

Before activity assays, CrTK samples in the apo-form (hereafter CrTK_{apo}) were desalted using NAP5 columns (GE Healthcare) pre-equilibrated with 50 mM Tris-HCl (pH 7.9). CrTK_{apo} (2.9 mg mL⁻¹) was then incubated at 25 °C for variable times (5–300 min) in the presence of 0.7 mM TPP supplemented or not with different concentrations of MgCl₂ ranging from 0.05 to 8 mM (hereafter CrTK_{TPP/Mg} and CrTK_{TPP}, respectively). The catalytic activity of CrTK forms was detected spectrophotometrically by a coupled enzymatic assay with EcE4PDH [24], in a mixture containing 50 mM Tris-HCl (pH 7.9), 0.1 mM TPP, 2.5 mM β -NAD, 15 mM MgCl₂, 5 mM CaCl₂, 2 mM DL-G3P,

12 mM F6P, 2.7 μM EcE4PDH. The reaction was started by the addition of pre-incubated CrTK samples (10-50 nM). Activity was monitored for 2 min and calculated from the increase in absorbance at 340 nm (NADH production) with a Jasco V-550 UV/Vis spectrophotometer. For each protein sample, we assessed the linear dependency of CrTK activity upon CrTK concentration and the rate of NADH production was only measured within linearity. The catalytic activity, expressed as $\text{mol}_{\text{NADH}}/\text{mol}_{\text{TK}} \text{ s}^{-1}$, was calculated from the $\Delta\text{Abs}_{340}/\text{min}$ considering a molar extinction coefficient for NADH of $6.22 \text{ M}^{-1} \text{ cm}^{-1}$.

2.7 Oxidant treatments

Before oxidative treatments, CrTK_{TPP} and CrTK_{TPP/Mg} were prepared as described above and diluted seven-times in 50 mM Tris-HCl (pH 7.9). After dilution, CrTK samples were incubated at 25 °C in the presence of 20 or 50 mM trans-4,5-dihydroxy-1,2-dithiane (DTTox). At the indicated times, an aliquot was withdrawn to assay the activity as described above. Control CrTK samples were directly diluted in buffer without the addition of DTTox.

2.8 Dynamic light scattering

Dynamic Light Scattering (DLS) measurements were performed employing a Malvern Nano ZS instrument equipped with a 633 nm laser diode. Samples consisted of CrTK_{apo}, CrTK_{TPP} (0.7 mM TPP) and CrTK_{TPP/Mg} (0.7 mM TPP plus 8 mM MgCl₂) and CrTK_{apo} pre-incubated in the presence of 50 mM DTTox. All samples were at 12 μM CrTK in 50 mM Tris-HCl (pH 7.9). Protein samples were used in disposable polystyrene cuvettes (100 μL) of 1 cm optical path length. The reported hydrodynamic radii (R_h) have been averaged from the values obtained from five measurements, each composed of 10 runs of 10 seconds.

2.9 Circular dichroism spectroscopy

The CD analysis on CrTK samples was performed on a Jasco J-810 spectropolarimeter equipped with a PTC-423S Peltier-type temperature control system; measurements were carried out in a Hellma 115B-QS micro cell (1 cm pathlength, 400 μL volume) using a 2 sec data integration time, a 2 nm spectral bandwidth and an accumulation cycle of 3 spectra per measurement. CrTK_{apo} sample (6.3 μM) was prepared in 5 mM Tris-HCl (pH 7.9) supplemented with 0.1 mM Na₂EDTA and 2 mM MgCl₂. CrTK_{TPP/Mg} was prepared by direct addition of TPP to the CrTK_{apo} sample inside the micro cell (final concentrations: [CrTK] = 5.6 μM , [TPP] = 55.6 μM). Samples were then analyzed at 25 °C

in the 400–250 nm spectral range using a 20 nm min⁻¹ scanning speed.

The kinetics of TPP binding to CrTK was evaluated on a CrTK_{TPP/Mg} sample (17.8 μM) prepared in 5 mM Tris-HCl (pH 7.9) supplemented with 220 μM TPP and 2.5 mM MgCl₂. The sample was analyzed at 25 °C in the 400–300 nm spectral range using a 50 nm min⁻¹ scanning speed; measurements were performed at different times after TPP addition over a 20 h period. The effect of Mg²⁺ depletion on TPP binding to CrTK_{TPP/Mg} was then evaluated by adding an excess of Na₂EDTA (8 mM); the measurement was carried out after 2 h incubation at 25 °C.

CD spectra were converted to molar units ($\Delta\epsilon$) based on the total concentration of CrTK; the insurgence of induced CD (ICD) signals [25,26] upon TPP binding was evaluated by subtraction of the CD spectrum of CrTK_{apo} from the CD spectra of CrTK_{TPP/Mg} samples.

2.10 Crystallization and data collection

CrTK_{apo} (without cofactors) and CrTK_{TPP/Mg} (protein plus 0.25 mM TPP and 2 mM MgCl₂) were concentrated to 4.5 mg mL⁻¹ in 50 mM Tris-HCl (pH 7.9), 1 mM EDTA and crystallized by the hanging drop vapor diffusion method at 293 K. The Structure screen 1 from Molecular Dimensions and the Crystallization Extension kit for Proteins from Sigma Aldrich, both based on the sparse matrix screen elaborated in [27] were used as starting screening. A protein solution aliquot of 2 μL was mixed to an equal volume of reservoir, and the prepared drop was equilibrated against 800 μL of reservoir. Small crystals and crystalline aggregates grew from various reservoirs of the Extension kit (*i.e.* 22: 12 % w/v PEG 20K, 0.1 M MES, pH 6.5; 30: 10 % w/v PEG 6K, 5 % v/v MPD, 0.1 HEPES, pH 7.5; 37: 10 % w/v PEG 8K, 0.1 M HEPES, pH 7.5; and 38: 20 % w/v PEG 10K, 0.1 M HEPES, pH 7.5). The crystallization conditions were optimized and the best crystals grew from a reservoir containing 10 % w/v PEG 6K, 5 % v/v MPD and 0.1 M MES pH 6.5-7.0 or 0.1 M HEPES pH 7.0-8.0. Crystals were mounted from the crystallization drop into cryo-loops, briefly soaked in a cryo-protectant solution containing 12 % w/v PEG 6K and 15 % v/v MPD, and then frozen in liquid nitrogen. Diffraction images were recorded at 100 K using a synchrotron radiation at Elettra (Trieste, beam line XRD1) for CrTK_{TPP/Mg} and at the European Synchrotron Radiation Facility (Grenoble, beam line ID30B) for CrTK_{apo}. The data at a resolution of 1.74 Å for CrTK_{TPP/Mg} and 1.58 Å for CrTK_{apo} were processed using XDS [28] and scaled with SCALA [29]. The correct space group was determined with POINTLESS [29] and confirmed in the structure solution stage. The data collection statistics are reported in Table 1.

2.11 Structure solution and refinement

The CrTK_{TPP/Mg} structure was solved by molecular replacement using the program MOLREP from CCP4 package [30]. The coordinates of TK from *Zea mays* (PDB code 1ITZ) [9], deprived of TPP, Mg²⁺ and water molecules, were used as search probe. Initial stages of the refinements were performed with CNS1.3 [31] selecting 5 % of reflections for R_{free}. The residual electron density map clearly showed the position of TPP and Mg²⁺, which were added to the model. Water molecules were automatically added and, after a visual inspection, they were conserved in the model only if contoured at 1.0 σ on the (2F_o - F_c) map and if they fell into an appropriate hydrogen bonding environment. In the final stages of the refinement performed with REFMAC 5.5.0109 [32], alternate conformations, if visible, were inserted. The structure of CrTK_{TPP/Mg} without TPP, Mg²⁺ and waters, was used as probe to solve the CrTK_{apo} structure by molecular replacement using MOLREP [30]. The refinement was performed as described above for CrTK_{TPP/Mg}. The refinement statistics are reported in Table 1. The manual rebuilding was performed with Coot [33]; protein superimpositions were performed by LSQKAB from CCP4 package [30].

2.12 Accession numbers

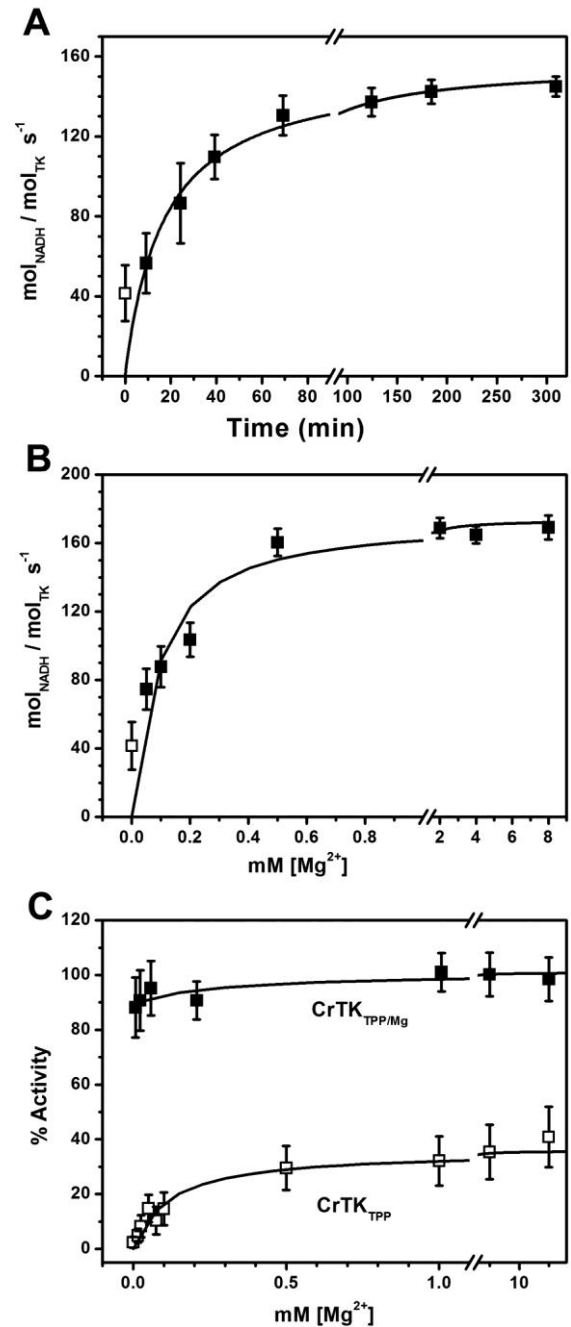
The atomic coordinates and structure factors of CrTK_{TPP/Mg} and CrTK_{apo} structures have been deposited in the Protein Data Bank with the accession numbers 5ND5 and 5ND6.

3. Results

3.1 Magnesium and TPP are both fundamental to reconstitute fully active CrTK

The reconstitution of the fully active TKs isolated from different organisms shows a strong dependence on divalent cations and TPP [8,34,35]. Since the cofactor TPP and Mg²⁺ were lost during the purification procedure (see Material and Methods and below), we first analyzed the time-dependence of CrTK catalysis on CrTK reconstitution in the presence of over-saturating concentrations of TPP and Mg²⁺. Samples of CrTK without cofactors (thereafter indicated CrTK_{apo}) were incubated at 25 °C in reconstitution buffer containing 50 mM Tris-HCl (pH 7.9), 0.7 mM TPP and 8 mM MgCl₂. Activity was evaluated over-time (5-300 min) with a coupled enzymatic assay using fructose-6-phosphate (F6P) and glyceraldehyde-3-phosphate (G3P) as substrates. As shown in Fig. 1A, the enzyme activity increased over time, reaching its maximal value after ~3 h (~150-160

Fig. 1. Reconstitution of the CrTK_{TPP/Mg} active enzyme. (A) Activities of CrTK samples reconstituted with 0.7 mM TPP and 8 mM MgCl₂. At the indicated times, an aliquot was withdrawn to measure CrTK activity following the procedure described in Materials and Methods. White square: activity of CrTK incubated in the absence of MgCl₂. The experimental data have been fitted with an arbitrary function and represent the mean \pm SD of three replicates (n = 3). (B) Activities of CrTK samples reconstituted with 0.7 mM TPP and increasing Mg²⁺ concentrations. After 3 h incubation, the activity was measured following the procedure described in Materials and Methods. The experimental data have been fitted with an arbitrary function and represent the mean \pm SD of three replicates (n = 3). The white square symbol refers to CrTK_{TPP} reconstituted in the absence of Mg²⁺ (40 mol_{NADH}/mol_{TK} s⁻¹). (C) Activities of CrTK reconstituted with 0.7 mM TPP (CrTK_{TPP}, white squares) or with 0.7 mM TPP supplemented with 8 mM MgCl₂ (CrTK_{TPP/Mg}, black squares). Activities of CrTK samples have been plotted as a function of Mg²⁺ concentrations in the assay buffer and normalized to the maximal activity (130 mol_{NADH}/mol_{TK} s⁻¹) measured for the fully reconstituted enzyme (CrTK_{TPP/Mg}). The experimental data have been fitted with an arbitrary function and represent the mean \pm SD of three replicates (n = 3).



mol_{NADH}/mol_{TK} s⁻¹). We then evaluated the dependence of the CrTK catalytic activity on CrTK reconstitution with over-saturating TPP and variable amount of Mg²⁺. Samples of CrTK_{apo} were incubated for 3 h at 25 °C in reconstitution buffer containing 50 mM Tris-HCl (pH 7.9), 0.7 mM TPP and different MgCl₂ concentrations ranging from 0 to 8 mM. After incubation, the activity was

measured as described above. As shown in Fig. 1B, incubation in the presence of Mg^{2+} significantly stimulated the activity of CrTK, and the enzyme attained its full activation ($\sim 150\text{-}160 \text{ mol}_{\text{NADH}}/\text{mol}_{\text{TK}} \text{ s}^{-1}$) at Mg^{2+} concentration $\geq 1 \text{ mM}$. This value exceeds almost 20 times the stoichiometric amount required on the basis of monomer concentration. By contrast, the activity of CrTK_{apo} incubated in the presence of TPP without Mg^{2+} (thereafter indicated CrTK_{TPP}) was $40 \text{ mol}_{\text{NADH}}/\text{mol}_{\text{TK}} \text{ s}^{-1}$ (Fig. 1B, white square), a value four times lower compared to fully reconstituted CrTK (thereafter indicated CrTK_{TPP/Mg}). In agreement with what reported about the activation of yeast TK [36], when CrTK_{TPP} was not pre-incubated in the presence of Mg^{2+} the enzyme exhibited a lower activity, even if exposed to saturating concentrations of divalent cations (*i.e.* Mg^{2+} and Ca^{2+}) during the activity assays.

As the coupled assay used to measure CrTK activity was conducted at high concentrations of Mg^{2+} and Ca^{2+} , we have also determined the dependence of CrTK_{TPP} and CrTK_{TPP/Mg} activity as a function of Mg^{2+} concentration in the assay buffer deprived of calcium ions. As shown in Fig. 1C (black squares), CrTK_{TPP/Mg} was not responsive to the presence of Mg^{2+} , showing 90% of maximal activity even in the absence of Mg^{2+} in the assay buffer. A different scenario is presented by CrTK_{TPP} that displayed a strong sensitivity to Mg^{2+} in the assay buffer (Fig. 1C, white squares). Moreover, the activity of CrTK_{TPP} measured under assay conditions deprived of Mg^{2+} was $\sim 4 \text{ mol}_{\text{NADH}}/\text{mol}_{\text{TK}} \text{ s}^{-1}$ (Fig. 1C, first white square), a value representing 10% of the maximal activity reached at saturating Mg^{2+} concentration (Fig. 1C, white squares). As reported by Heinrich *et al* [8], we cannot exclude the presence of trace contaminant cations in the assay buffer due to the addition of substrates, or that the enzyme retains some activity even in the absence of Mg^{2+} . However, when we compared the activities of CrTK_{TPP} and CrTK_{TPP/Mg} in assay conditions deprived of Mg^{2+} , the former activity is extremely low, being only 3% of the activity measured for CrTK_{TPP/Mg} (Fig. 1C, first white and black squares, respectively).

Altogether, these results indicate that maximal catalytic activity requires long-time incubation (at least 2 hours) of CrTK with over-saturating TPP and Mg^{2+} concentration. Indeed, CrTK reconstituted in the presence of TPP without Mg^{2+} (*i.e.* CrTK_{TPP}) does not reach the optimal catalysis activity even in the presence of over-saturating Mg^{2+} in the assay buffer.

3.2 Induced CD spectra of CrTK

Upon binding of TPP to TK from *S. cerevisiae*, an increase in absorption can be observed in the near UV range (285–360 nm) [1,37-39]. A peculiar bisignate induced CD (ICD) signal, closely related to the increase in UV absorption, is also observed across a variety of TPP-dependent enzymes [1,39,40].

CrTK_{TPP/Mg} displays a bisignate ICD signal in the 380–260 nm range (Fig. 2A), with a negative band centered at 325 nm and a positive band centered at 275 nm; the ICD profile is very similar to those reported for other variants of TK. The binding of TPP to CrTK displayed a relatively slow kinetics: the intensity of the ICD signal increased with time, reaching saturation 3 h after addition of the cofactor (Fig. 2B) consistent with activity assays (Fig. 1A). Treatment of CrTK_{TPP/Mg} with excess of the chelating agent Na₂EDTA caused the complete disappearance of the ICD signal of TPP (Fig. 2B): this behavior clearly demonstrated that Mg²⁺ is necessary for the effective formation of a CrTK/TPP complex in the most catalytically active configuration.

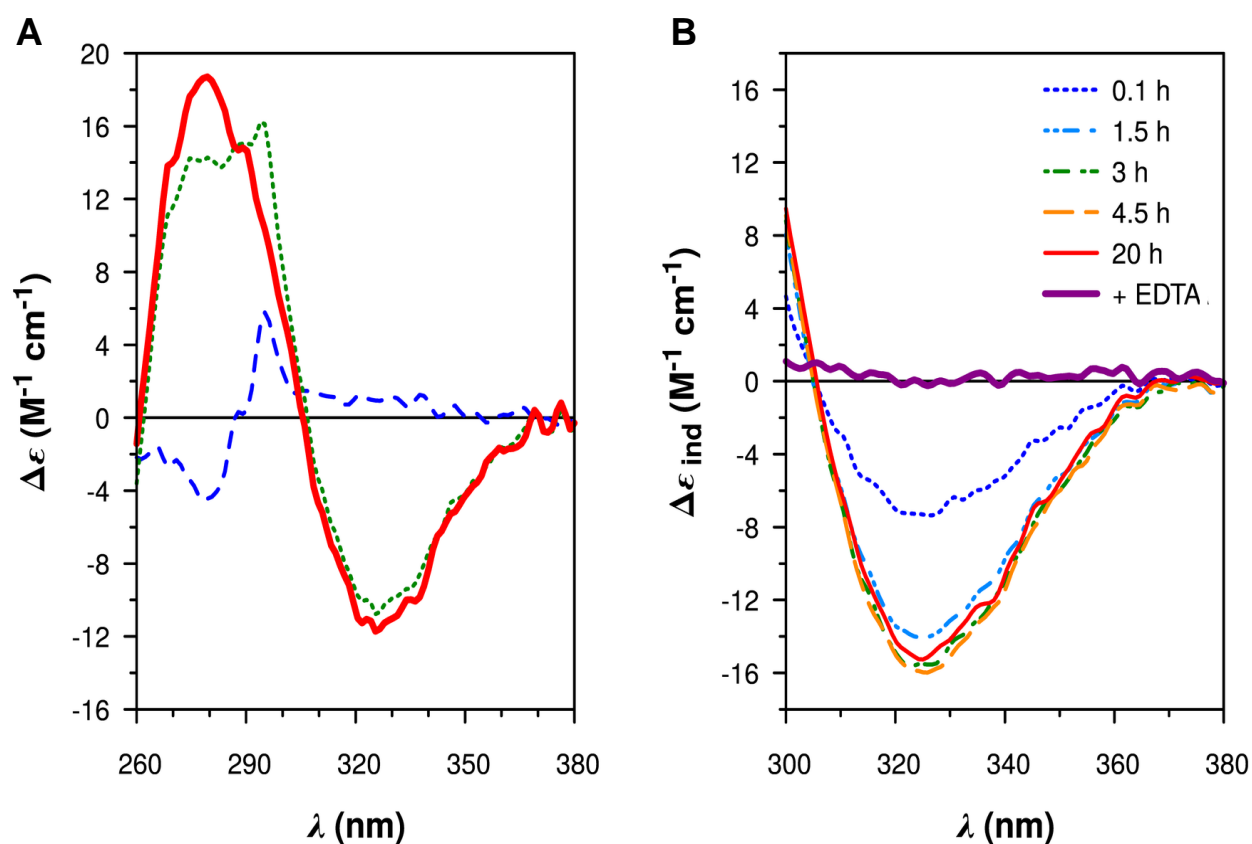


Fig. 2. ICD spectra for TPP binding to CrTK

(A) CD spectra for CrTK_{apo} (6.3 μM; blue thin line) and CrTK_{TPP/Mg} (5.6 μM, [TPP] = 55.6 μM; green dotted line); the resulting ICD signal for the binding of TPP to CrTK is shown (bold red line). (B) ICD spectra for the binding of TPP (220 μM) to a sample of 17.8 μM CrTK, 2.5 mM MgCl₂ at different times after TPP addition (thin lines) and after addition of 8 mM Na₂EDTA (bold purple line).

3.3 Quaternary structure of CrTK_{apo} and CrTK_{TPP/Mg}

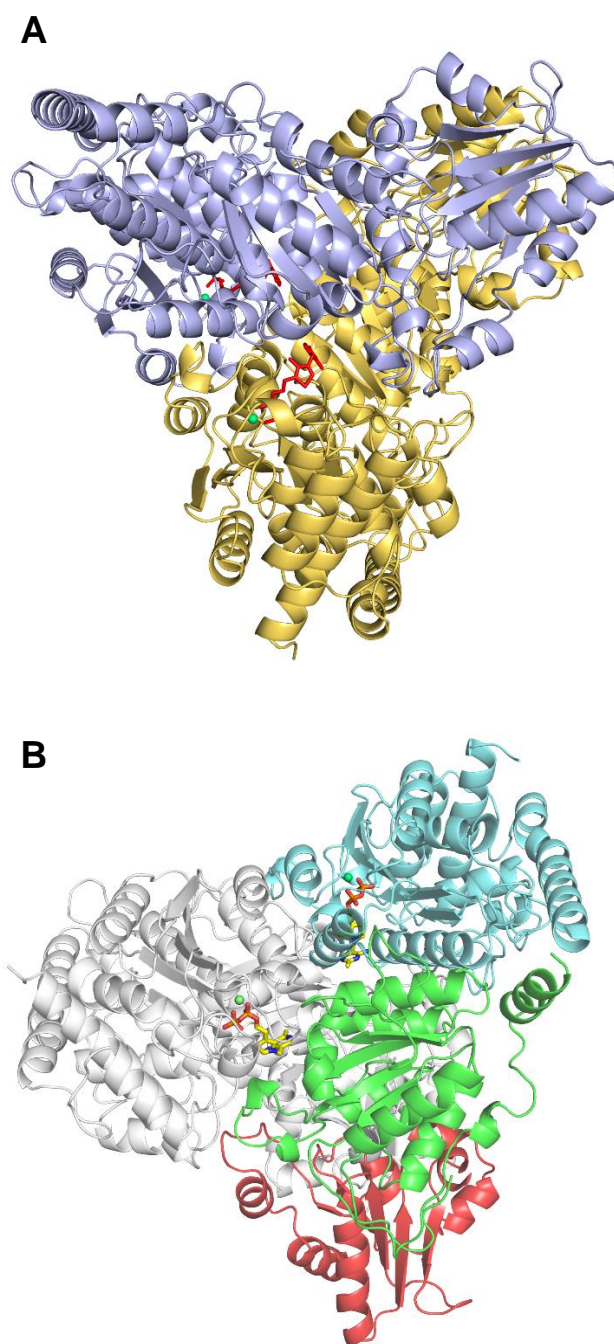
Based on the available structures of TKs, the enzyme is a homodimer composed by two identical subunits [6,9]. In order to investigate the oligomerization state of CrTK, we performed gel filtration experiments and DLS analysis. Even though that the purified CrTK_{apo} is deprived of the cofactors Mg²⁺ and TPP (*i.e.* CrTK_{apo}), it eluted as a single symmetric peak with an apparent molecular mass of 156 kDa when analyzed by size-exclusion chromatography (Fig. S2). The elution behavior is similar to the well-known tetrameric enzyme AtGAPC1 (160 kDa) [23]. Since the molecular weight of the CrTK monomer determined by mass spectrometry is 75170 Da (Fig. S3), the CrTK appears as a homodimeric protein. The dimeric state of CrTK_{apo} in solution was further confirmed by DLS measurements that reported a hydrodynamic radius of 5.1 ± 0.1 nm, corresponding to an apparent molecular mass of 151.6 ± 7.4 kDa (Table S1). DLS analysis was also employed to evaluate the effect of TPP and Mg²⁺ on the overall structure of CrTK. Under these conditions, CrTK remains homodimeric (Table S1).

3.4 Crystal structure of CrTK_{apo} and CrTK_{TPP/Mg}

CrTK_{apo} and CrTK_{TPP/Mg} were crystallized as described in the Materials and Methods section. Both forms produced different polymorphs, belonging to C2 or P1 space groups (not shown). One crystal for each form was selected for the final structure determination on the basis of resolution and quality of the data (Table 1). Despite differing for unit cell dimensions, crystals of CrTK_{apo} and CrTK_{TPP/Mg} both belong to C2 space group and show a similar Matthews coefficient (2.32 and 2.42, respectively) and solvent content (47.1 % and 49.2 %, respectively). In both crystals, the asymmetric unit contains two protein chains related by a non-crystallographic two-fold axis and forming the V-shaped homodimer (Fig. 3A). In both forms, each monomer is formed by 670 amino acids, starting from Ser49 (the first residue visible in the electron density map), to the last C-terminal residue (His718) that could be modeled only in one subunit of CrTK_{apo}. To avoid misinterpretation with previous studies [5,41], amino acids of CrTK were numbered following the full-length enzyme (mature enzyme plus transit peptide; Fig. S1). As previously reported for other TK structures [6,9], each CrTK monomer is composed of three consecutive α/β domains all containing a central five stranded β -sheet (Fig. 3B). The N-terminal domain (residues 49-372) folds in a parallel β -sheet composed of strands β 1, β 2, β 3, β 6 and β 7 surrounded by several α -helices (α 1- α 12; Fig. S4). It binds the Mg²⁺ ion and the pyrophosphate moiety of TPP, therefore it is called PP domain (Fig. 3B). The middle domain

(residues 373-582) is also composed of a parallel β -sheet ($\beta 9$ - $\beta 13$) embedded in several helices from $\alpha 13$ to $\alpha 18$ (Fig. S4). This domain binds the cofactor pyrimidine ring of the adjacent monomer and is called Pyr domain (Fig. 3B). The C-terminal domain (residues 583-718) is the smallest domain and it folds in a mixed β -sheet with four parallel strands ($\beta 16$ - $\beta 19$), one antiparallel strand ($\beta 15$), and several α -helices ($\alpha 19$ to $\alpha 25$). This domain is not involved either in dimer interface or in the cofactor binding (Fig. 3B). The CrTK structure is characterized by long α -helix regions but it contains also several short 3_{10} and π helices connecting different secondary structure elements.

Fig. 3. Ribbon representation of CrTK_{TPP/Mg} structure (A) View of the V-shaped dimer including the two cofactors TPP and Mg²⁺ present in each monomer. The two monomers are differently colored. The TPP (red) is shown as sticks and the Mg²⁺ (green) as sphere. (B) View of the dimer including TPP shown as sticks and the Mg²⁺ as green sphere. The three domains of one monomer are differently colored: N-terminal or PP domain in cyan, the middle or Pyr domain in green and the C-terminal domain in red. One TPP interacts with the PP domain (cyan) by the pyrophosphate groups; the second TPP interacts with the Pyr domain (green) by the pyrimidine ring. Atom colors: yellow (C), blue (N), red (O), purple (P) and green (Mg). The figure was prepared with Pymol (The PyMOL Molecular Graphics System, Version 1.5.0.5 Schrödinger, LLC).



3.5 Comparison of CrTK with other TK structures

The three-dimensional structure of TK from *Zea mays* (PDB code 1ITZ) [9] is the unique TK structure from a photosynthetic organism present in PDB. CrTK shares with this protein a sequence identity of about 65%. Superimposition of C α atoms based on secondary structure matching, results in a root mean square deviations (rmsd) of 0.9 Å on 1320 aligned residues (whole dimer) and 0.8 Å on 663 aligned residues (single monomer) indicating that the TK structure is highly conserved between the two organisms. While the internal β -sheets of N-terminal and middle domains are well superimposed, some deviations can be observed in correspondence to the solvent exposed helices α 12 and α 13 and the C-terminal domain (Fig. S5A). Moreover, the N-terminal and the middle domains of the two structures superimpose with rms deviations of 0.6 Å (on 322 aligned residues) and 0.7 Å (on 209 aligned residues), respectively, while the C-terminal domain does with a slightly higher rmsd (0.9 Å on 132 aligned residues).

In the case of the yeast enzyme (PDB code 1TRK) [6], the sequence identity with CrTK is about 53%. The rmsd from the superimposition of C α atoms based on secondary structure matching increases to 1.2 Å on 1295 aligned residues (whole dimer) and 1.1 Å on 651 aligned residues (single monomer) with respect to the previous comparison. Differences are mainly observed in the correspondence to some exposed regions of the N-terminal domain, in particular in the region comprising helices α 10, α 11, α 12 and α 13 (Fig. S5B). In this case, the superimposition of N- and C-terminal domains gives higher deviations (1.1 Å on 313 aligned residues and 1.2 Å on 127 aligned residues, respectively) with respect to that one of middle domains (0.9 Å on 210 aligned residues).

3.6 CrTK_{TPP/Mg} structure reveals multiple interactions between TPP, Mg²⁺ and active site residues

In the CrTK_{TPP/Mg}, each subunit binds one TPP molecule and one Mg²⁺ ion in the active site cleft at the dimer interface. The TPP and the Mg²⁺ are set in place by hydrogen bonds involving residues of both subunits and water molecules (Fig. 4 and Table S2).

the same subunit and Leu433 of the adjacent subunit, while the pyrimidine ring is stacked between Leu168 of the same subunit and Phe490 of the adjacent subunit (Fig. 5A). Moreover, the hydrophobic cavity is also completed by Pro167, Val461, Leu433, Phe487 and Tyr493 from the two subunits (Fig. 5A). The N1 nitrogen atom of the pyrimidine ring is hydrogen bonded to the carboxyl oxygen OE2 of the strictly conserved residue Glu463 (Fig. 4 and Table S2) [42,43]. In addition, the N4 atom of the pyrimidine ring forms an intramolecular interaction with the C2 carbon of the thiazolium ring (Fig. 5A and Table S2) which favors the V-like conformation of TPP after binding and the C2 deprotonation required for catalysis [9,44].

Mg²⁺ shows an octahedral coordination with six oxygen atoms as electron donor, belonging to the pyrophosphate groups of TPP (O1B and O2A), the carboxylic group of Asp208 (OD2), the amide group of Asn238 (OD1), the main chain carbonyl group of Ile240 and a water molecule (Fig. 5B).

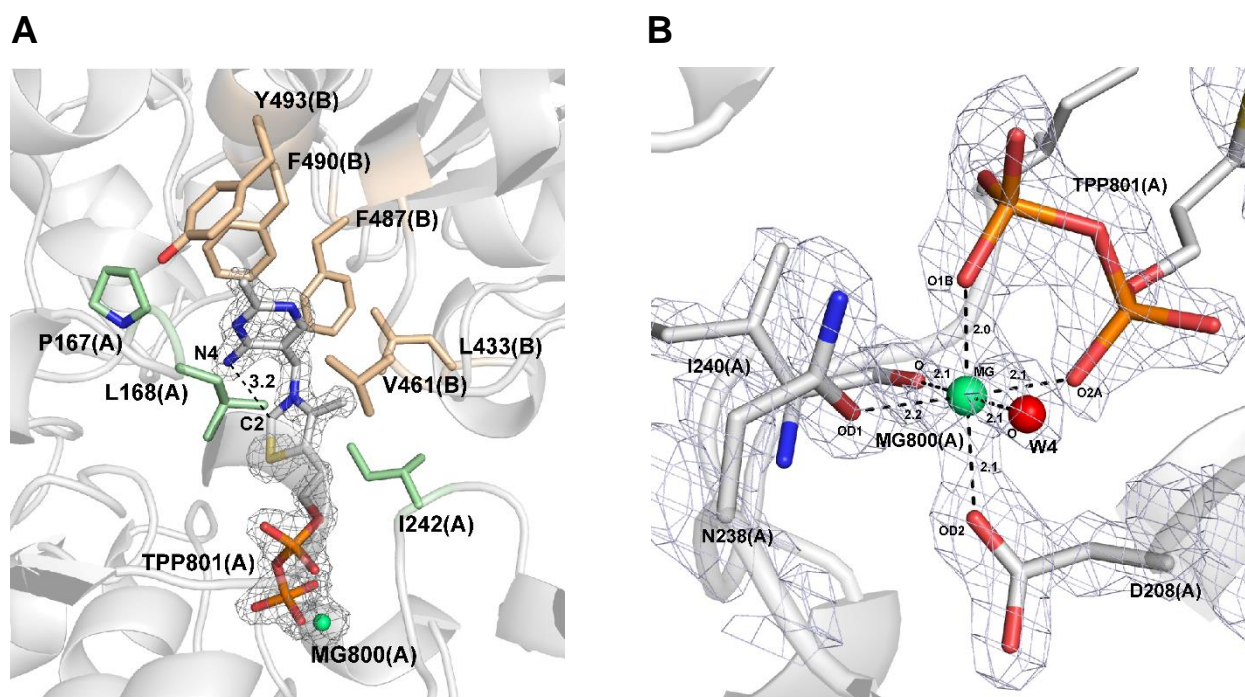


Fig. 5. Hydrophobic interactions of TPP and coordination site of Mg ion. (A) $2F_o - F_c$ electron density map contoured at 2σ around TPP. The pyrimidine and the thiazolium rings are stabilized by several protein residues of both subunits (A in light green, B in pale brown) forming a hydrophobic cavity. The side chain of Phe490 is in stacking interaction with the TPP pyrimidine ring. (B) $2F_o - F_c$ electron density map contoured at 1.5σ around Mg²⁺ and its ligands. The Mg²⁺ shows an octahedral coordination with six oxygen atoms as electron donors. Mg²⁺ is represented as a green sphere, the ligands as ball-and-stick. Atom colors: grey (C), blue (N), red (O) and orange (P). The figure was prepared with Pymol (The PyMOL Molecular Graphics System, Version 1.5.0.5 Schrödinger, LLC).

3.7 Comparison between CrTK_{apo} and CrTK_{TPP/Mg}

In the CrTK_{apo}, the cleft displays no electron density attributable to TPP and Mg²⁺ supporting the fact that the purified protein before reconstitution is cofactor(s) free. By superimposing C_α atoms of CrTK_{apo} and CrTK_{TPP/Mg} monomers (chains A and B) and dimers, rmsd of 0.3 Å and 0.4 Å (on 667 and 668 superimposed atoms) and 0.4 Å (on 1335 superimposed atoms) are observed, respectively. Despite the high similarity between the two forms, in the CrTK_{apo} two regions ranging between residues 239-247 and 433-437 (Fig. S4) are completely deprived of electron density, indicating that these regions are likely disordered and flexible (Fig. 6A). In CrTK_{TPP/Mg} structure both regions are involved in TPP and Mg²⁺ binding (Fig. 6B) as for example the main chain carbonyl group of Ile240 which is hydrogen bonded to both TPP pyrophosphate O1B atom and Mg²⁺ (Fig. 4, Fig. 5B and Table S2) or the side chains of Ile242 and Leu433 stabilizing the TPP aromatic rings by hydrophobic interactions. CrTK_{apo} structure shows that TPP is replaced by several water molecules located in the region of phosphate groups (W214, 450, 526, 557 and 751 in chain A and W366, 493 and 643 in chain B) and pyrimidine ring (W284, 496 and 559 in chain A and W172, 300 and 530 in chain B), whereas the Mg²⁺ site is empty (Fig. 6C). These molecules contribute mainly by hydrogen bonds to the stabilization of active site residues lying almost in the same position in the CrTK_{TPP/Mg} structure, except for the two protein portions mentioned above. Both protein regions, disordered in CrTK_{apo} structure, are also involved in dimer interface in CrTK_{TPP/Mg} structure. Indeed, Ser241, Ile242 and Asp243 of one subunit interact with the carboxylate group of Asp432, the main chain nitrogen of Ala434 and the amino group of Arg458 of the other subunit (Fig. 6D). The absence of TPP probably induces a disorder in the region from residues 239 to 247 causing the impairment or the loss of the hydrogen bonds with residues 433-437 of the other subunit. In consequence, this second region more exposed to the solvent becomes flexible and disordered. To further investigate whether the presence/absence of TPP/Mg²⁺ (*i.e.* absence/presence of disordered regions) influences the protein stability, we evaluated the thermal-induced denaturation of CrTK_{apo} and CrTK_{TPP/Mg}. Protein activities have been measured after 30 min exposure to increasing temperature ranging from 20 °C to 60 °C. CrTK_{apo} has a T₅₀ (temperature causing 50% inactivation) of around 37 °C while this value is shifted to ~45 °C for CrTK_{TPP/Mg} (Fig. S6). Moreover, at 40 °C the activity of the CrTK_{TPP/Mg} sample was still maximal while the CrTK_{apo} was almost completely inactivated. This indicates that the presence of TPP/Mg²⁺ in the active site stabilizes the enzyme when exposed to denaturing temperatures, suggesting that a significant decrease of the overall disorder/flexibility is due to TPP and Mg²⁺ binding.

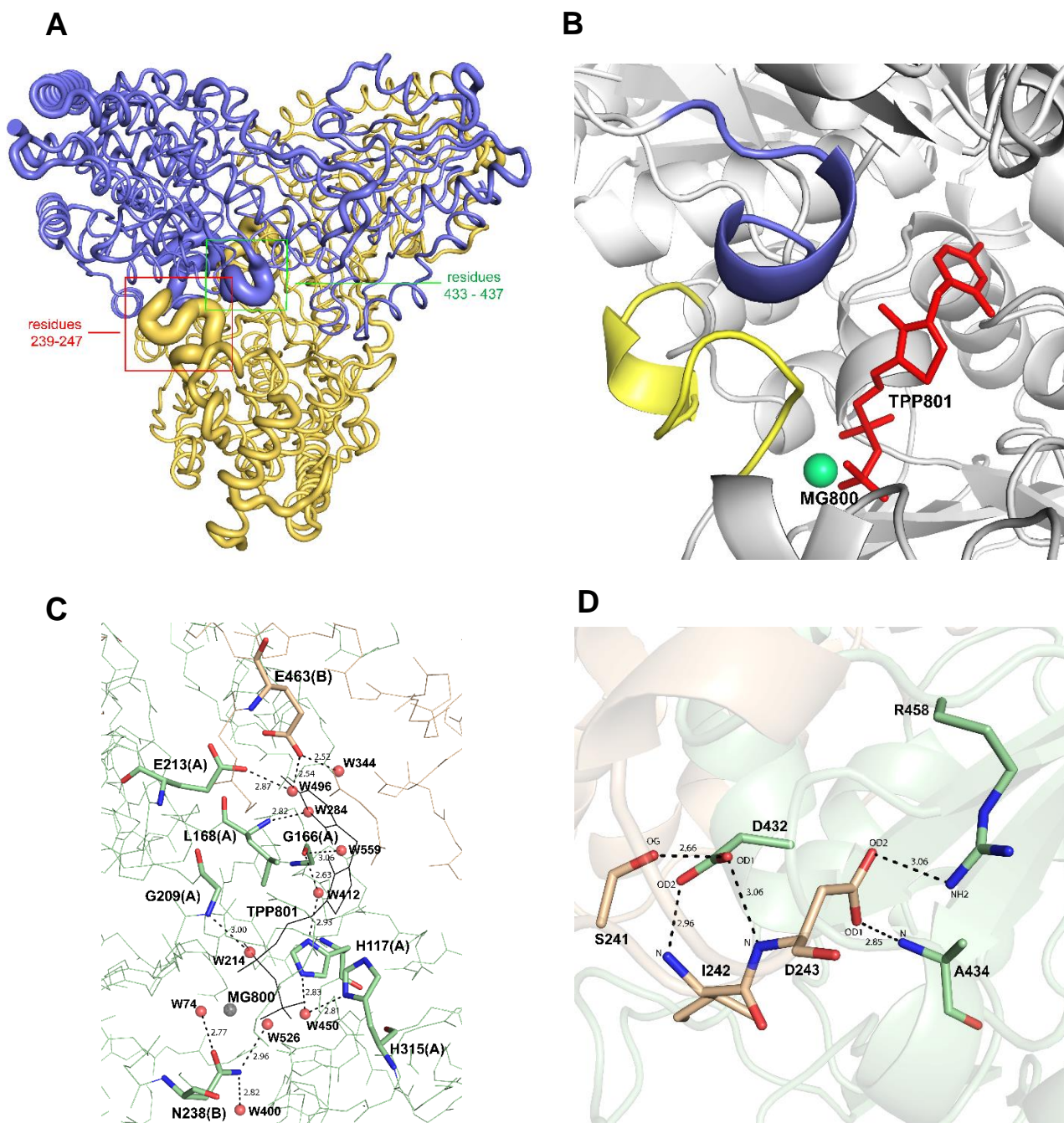


Fig. 6. Comparison between CrTK_{apo} and CrTK_{TPP/Mg} structures. (A) Representation of the CrTK_{apo} dimer structure. The thickness is correlated to the atom thermal factor (B-factor). The highlighted protein regions (239-247 and 433-437) are disordered and flexible. (B) CrTK_{TPP/Mg} structure reveals that the same protein regions have an ordered conformation and interact with TPP and Mg²⁺. (C) CrTK_{apo} active site structure shows that water molecules occupy the position of TPP in CrTK_{TPP/Mg} structure (represented by grey thin lines for TPP and sphere for Mg²⁺) and stabilize catalytic residues by hydrogen bonds. (D) Dimer interface in CrTK_{TPP/Mg} structure showing the interactions between the residues belonging to regions 239-247 and 433-437, disordered in CrTK_{apo} structure. The figure was prepared with Pymol (The PyMOL Molecular Graphics System, Version 1.5.0.5 Schrödinger, LLC).

3.8 Effect of DTT oxidation on CrTK activity

Plant TK was previously identified as putative target of TRX-dependent regulation [5] but no biochemical studies confirmed the sensitivity of this enzyme to dithiol/disulfide exchange reactions. Mature CrTK (CrTK sequence deprived of chloroplast transit peptide) enumerates twelve cysteines (Fig. 7A) mainly distributed in the PP and Pyr domains (Cys58, 84, 173, 204, 210, 220 and 366 in the PP domain; Cys386, 470, 484 and 582 in the Pyr domain) with only Cys638 located in the C-terminal domain (amino acids numbering refers to the full-length CrTK, Fig. S1). Among them, only cysteine residues at position 84, 210, 220, 470 and 484 are conserved in photosynthetic organisms (Fig. S4). Crystal structures of both CrTK_{apo} and CrTK_{TPP/Mg} show that all cysteines are in a reduced state making CrTK a putative target of oxidative regulation.

The analysis of CrTK_{TPP/Mg} structure shows that the thiol sulfur atoms of Cys470 and Cys484 are located only 4.0 Å apart, while the distances Cys173 - Cys220 and Cys204 - Cys210 are 6.6 Å and 7.0 Å, respectively (all distances refer to the sulfur atoms; Fig. 7B). The thiol sulfur atoms of all other cysteine pairs lie at distances higher than 10 Å. These distance values do not significantly vary in CrTK_{apo} structure (Cys470 SG - Cys484 SG = 4.1 Å, Cys173 SG - Cys220 SG = 6.7 Å and Cys204 SG - Cys210 SG = 7.0 Å). Thus, these residues may participate to the formation of regulatory disulfide bonds.

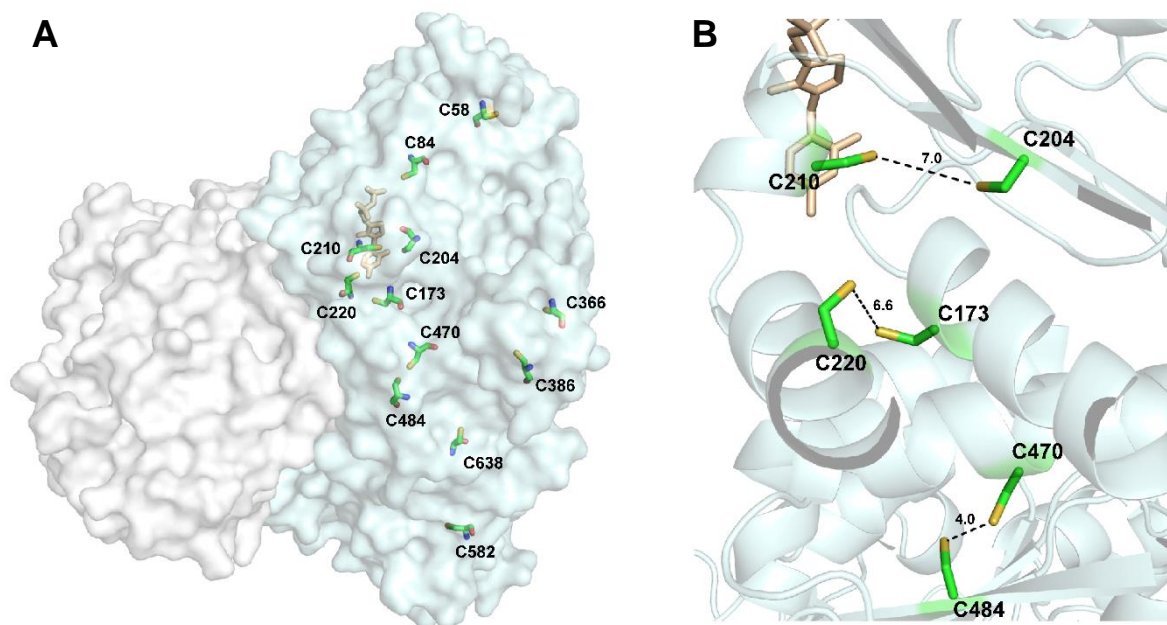


Fig. 7. Cysteine residues in CrTK_{TPP/Mg} structure

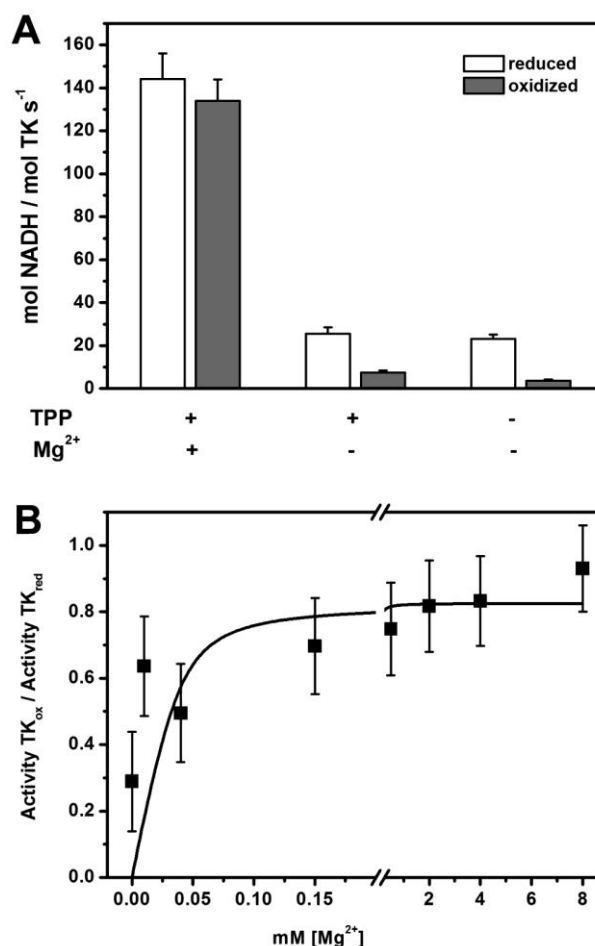
(A) Position of cysteine residues in CrTK structure. Cys58, 84, 173, 204, 210, 220 and 366 are located in PP domain;

Cys386, 470, 484 and 582 in Pyr domain and Cys638 in C-terminal domain. Cysteines and TPP are represented in ball-and stick, the two protein subunits as surface (chain A in light-cyan and chain B in gray). (B) Three pairs of cysteines belonging to chain A, show the thiol sulfur atoms groups at a distance $< 10 \text{ \AA}$. Atom colors: green (C), blue (N), red (O) and yellow (S). The figure was prepared with Pymol (The PyMOL Molecular Graphics System, Version 1.5.0.5 Schrödinger, LLC).

To test the sensitivity of CrTK activity to oxidative conditions, we exposed CrTK_{apo}, CrTK_{TPP} and CrTK_{TPP/Mg} to oxidized DTT (DTTox), an oxidant molecule known to induce disulfide formation. After 3 h incubation, the activity of CrTK_{TPP/Mg} was almost unaffected, retaining more than 90% of the control (reduced) enzyme activity (Fig. 8A, left bars).

Fig. 8. Oxidation of CrTK samples.

(A) CrTK_{apo}, CrTK_{TPP} and CrTK_{TPP/Mg} samples were prepared following the procedure described in Material and Methods. After incubation, each sample was divided in two aliquots and diluted seven-times in 50 mM Tris-HCl (pH 7.9), in the presence (gray bars) or absence (white bars) of 50 mM DTTox. After 3 h incubation, the activity was measured following the procedure described in Materials and Methods. Data represent the mean percentage \pm SD ($n = 3$). (B) CrTK_{apo} was reconstituted in 50 mM Tris-HCl (pH 7.9) in the presence of 0.7 mM TPP, and variable Mg²⁺ concentrations. After 3 h incubation, each sample was treated as described in (A) and after additional incubation (3 h), the activity was assayed following the procedure described in Materials and Methods. The ratios between the activity of the oxidized and reduced (control) sample are plotted as a function of the Mg²⁺ concentration. The experimental data have been fitted with an arbitrary function and represent the mean \pm SD of three replicates ($n = 3$).



Thus, the oxidation induced by DTTox does not substantially influence the catalytic activity of the

fully reconstituted enzyme. Although its activity is 4-fold lower compared to the fully reconstituted enzyme (CrTK_{TPP/Mg}, Fig. 1B), CrTK_{TPP} is strongly inhibited by oxidation, retaining only 20-30 % of its control activity when exposed to 50 mM DTTox (Fig. 8A, central bars and Fig. S7). A similar result was observed for the CrTK_{apo}, being 80% inactivated following incubation in the presence of identical amount of DTTox (Fig. 8A, right bars). Incubation of oxidized CrTK_{TPP} with the strong reducing agent tri-carboxy-ethyl-phosphine (TCEP) [45] completely restored the activity, indicating that enzyme inhibition is due to reversible modifications such as disulfide bond(s) (Fig. S7). To better investigate the influence of the Mg²⁺ on the DTTox-induced inactivation of the enzyme, CrTK_{apo} was reconstituted in the presence of TPP and variable concentrations of Mg²⁺. Protein samples were then incubated in the absence or presence of DTTox. By plotting the ratios between oxidized and control activities against Mg²⁺ concentration, we noticed that the decrease of the Mg²⁺ amount in the reconstitution buffer is at the same time followed by an increased sensitivity of the enzyme to the oxidizing conditions (Fig. 8B). Taken together, these results indicate that only the CrTK_{apo} and CrTK_{TPP} forms are prone to oxidation-induced inhibition, and suggest that fluctuations of the Mg²⁺ concentration can modulate the extent of the enzyme sensitivity to oxidative regulation by influencing the relative amount of CrTK_{TPP/Mg}.

4. Discussion

The activity of TK isolated from diverse organisms is strictly dependent upon the presence of divalent cations and TPP [8,34,35,46]. The aim of the present study was to investigate the role of Mg²⁺ in the catalysis and regulatory mechanisms of CrTK and to get insight into the structural features of the enzyme. We focused our attention on the importance of Mg²⁺ ions because, in photosynthetic organisms, one of the documented effects induced by the transition from dark to light conditions is an increase of the magnesium concentration in the chloroplast stroma [47,48].

The reconstitution of fully active CrTK in the presence of an excess of TPP and Mg²⁺ requires long-time incubations (Fig. 1A) and is dependent on Mg²⁺ concentration in the pre-incubation media (Fig. 1B). Moreover, the activity of CrTK_{TPP/Mg} is not affected by the absence of Mg²⁺ in the reaction assay (Fig. 1C). By contrast, the activity of CrTK_{TPP} is sensitive to increasing concentrations of Mg²⁺ in the assay buffer but its maximal activity was 4-fold lower compared to the activity of CrTK_{TPP/Mg} (Fig. 1B). This suggests that the transition from the partially active CrTK_{TPP} to the fully active CrTK_{TPP/Mg} is a slow process that does occur during long incubation of the enzyme with

over-saturating concentration of both TPP and Mg^{2+} . Indeed, short exposure of the CrTK_{TPP} to Mg^{2+} ions during activity assay (2 min) is not sufficient to allow the enzyme to adopt suited conformation for optimal catalysis. This agrees with previous studies that analyzed the rate of transition from the apo-form of yeast TK to the fully active holo-form, demonstrating that the long time needed to reach the maximal activity is strongly decreased upon pre-incubation of the protein with both TPP and Mg^{2+} [8,36]. The formation of fully reconstituted TPP-dependent enzymes can be monitored by the appearance of UV and ICD signals upon TPP binding [39]. ICD spectra stem from the perturbations to the symmetry and electronic structure of a non-chiral ligand (*i.e.* TPP) interacting with a chiral guest [25,26]. In the case of TPP, these perturbations are due to the stabilization of the 4'-imino-1',4'-dihydropyrimidine tautomer of TPP, which is considerably less stable in aqueous solution than the 4'-aminopyrimidine tautomer, but is stabilized by the hydrophobic environment inside the catalytic site [38,49] and by the interaction with the highly conserved Glu463 (Fig. 4 and Fig. S4; Glu418 in yeast TK, [40]). Upon binding, TPP assumes a V-like conformation dictated by the optimal mutual orientation of the thiazole and pyrimidine rings for the effective deprotonation at C2 on the thiazole ring [50-52]. Consequently, these spectroscopic features arise from the catalytically active configuration of TPP and their intensities are linearly correlated to the catalytic activity of the enzyme [53,54]. A bisignate ICD signal in the 380–260 nm range was observed upon TPP addition to CrTK incubated in the presence of over-saturating amounts of $MgCl_2$ (Fig. 2A). In agreement with activity data (Fig. 1A), this ICD signal evolves with time, reaching its maximum in about 3 hours (Fig. 2B), showing that the transition from CrTK_{apo} to CrTK_{TPP/Mg} is a slow process. The chelating agent EDTA fully abolished the ICD signal (Fig. 2B), indicating that Mg^{2+} is fundamental for the correct binding of TPP inside the active site. Indeed, the CrTK_{TPP/Mg} crystal structure shows that the metal ion is coordinated by 2 oxygen atoms of the TPP pyrophosphate moiety (Fig. 4 and Fig. 5A) as already reported for yeast and maize TKs [9,55]. Taken together, the biochemical and structural results indicate that Mg^{2+} is strictly required for optimal enzyme functioning and stabilizes the binding interaction between TPP and the enzyme, allowing the active site to assume the optimal configuration for the catalytic activity. By contrast, TPP alone is not sufficient to allow CrTK to reach its fully active state (Fig. 1B).

The comparison between the two limit structures in terms of activity (*i.e.* CrTK_{apo} and the fully active CrTK_{TPP/Mg}) showed that the general fold is highly conserved except for two different portions (residues 239-247 and 433-437) which are completely disordered and flexible in CrTK_{apo} (Fig. 6A) as previously reported for yeast TK [56]. The CrTK_{TPP/Mg} structure highlighted that these protein

regions are involved in cofactors binding and they fold in an ordered conformation composed by short helices and random coil segments only in the presence of TPP and Mg^{2+} (Fig. 6B). Moreover, these two regions are located at the dimer interface and they form inter-chain interactions in $CrTK_{TPP/Mg}$ (Fig. 6D) that are absent in $CrTK_{apo}$. The results of the thermal stability comparison between $CrTK_{apo}$ and $CrTK_{TPP/Mg}$ agree with the structural observations. Notably, $CrTK_{TPP/Mg}$ showed a T_{50} significantly higher than $CrTK_{apo}$, indicating that the presence of TPP and Mg^{2+} in the active site decreases CrTK flexibility and stabilizes the protein towards thermal denaturation (Fig. S6). Therefore, we can propose that Mg^{2+} -dependent reconstitution/activation of CrTK is a slow process accompanied by protein conformational changes that allow a correct orientation of the cofactor TPP inside the catalytic center of the enzyme. Consistently, the formation of $CrTK_{TPP/Mg}$ can support maximal catalysis (Fig. 1). It has been already reported for yeast TK that the transition from the apo-protein to the fully active enzyme is a multi-step process, having a rate determined by slow rearrangements of the protein due to cofactor binding [1].

The dithiol/disulfide interchanges represent the main mechanism involved in the control of the activity of Calvin-Benson cycle (CBC) enzymes during light-dark transitions [5]. The formation of regulatory disulfides is accompanied by structural changes that modulate the activity of CBC enzymes allowing partial or complete inactivation. The regulation of CBC enzymes also involves other factors including light-dependent fluctuations of Mg^{2+} concentration [47,48,57,58]. In the last decades, several proteomic studies identified plant TK as a putative target of TRX-mediated regulation [16,17,19,59], suggesting the possible existence of regulatory cysteine residue(s). However, no biochemical study confirmed the TK sensitivity to redox transitions. The structures of CrTK here presented, reveal three putative cysteine couples (Cys173-Cys220, Cys204-Cys210 and Cys470-484) whose thiol sulfur atoms lie at distance lower than 10 Å, potentially involved in the formation of regulatory disulfide bonds (Fig. 7B). Moreover, we demonstrated that CrTK is sensitive to oxidative deactivation *in vitro* but this regulatory process is dependent on Mg^{2+} . Indeed, this metal preserves the fully active $CrTK_{TPP/Mg}$ from the inactivation induced by DTTox treatment (Fig. 8), likely precluding the formation of one or more regulatory disulfide bond(s). By contrast, both $CrTK_{apo}$ and $CrTK_{TPP}$, which lack Mg^{2+} in their active sites, are strongly sensitive to reversible inhibition by oxidation (Fig. 8 and Fig. S7). The role of Mg^{2+} in modulating such oxidative regulation is also suited to the physiological context. In fact, it has been previously reported that stromal Mg^{2+} concentration is in the low millimolar range [60], and varies according to several factors including light conditions [61,62]. It is well known the stromal pH increases upon illumination, and the transfer

of protons across the thylakoid membrane is believed to be electrically compensated by Mg^{2+} whose concentration increases in the stroma since the chloroplast envelope is relatively impermeable to this metal ion [63-65]. The stromal Mg^{2+} concentration is considered to increase 3- to 4-fold upon illumination [48,66]. This light-dependent fluctuation perfectly correlates with the strong dependency of CrTK activation upon increasing Mg^{2+} and oxidative inactivation observed under decreasing Mg^{2+} concentrations. The Mg^{2+} -dependent activation of CrTK and the role of Mg^{2+} in the modulation of oxidative sensitivity are reminiscent of the regulation of two other CBC enzymes, namely fructose-1,6-bisphosphatase (FBPase) and sedoheptulose-1,7-bisphosphatase (SBPase). Indeed, both enzymes are well known to be activated by Mg^{2+} , and this feature participates in the light-dependent activation of the enzymes [57,72]. This regulation was also found to act in concert with TRX-dependent modulation of FBPase redox state, indicating that both factors are undoubtedly important to control the activity of this enzyme. In addition, it has been reported that Mg^{2+} protects FBPase from oxidative deactivation, due to the stabilizing role of the complex between the substrate FBP and Mg^{2+} [73].

In conclusion, we can assume that fluctuations in Mg^{2+} content occurring in the stroma might affect at multiple extents the activity and regulation of CrTK. Indeed, this metal ion is implicated (i) in the reconstitution of the fully active form, (ii) in the formation and stabilization of the holo-form, and finally (iii) in the modulation of the redox sensitivity of the enzyme. Further studies will be required on CrTK and other CBC enzymes to investigate the molecular mechanisms underlying the crosstalk between magnesium-induced and redox-induced light-dependent activation of photosynthetic carbon fixation.

Table 1. Data collection and refinement statistics for CrTK_{apo} and CrTK_{TPP/Mg} structure

	CrTK _{TPP/Mg}	CrTK _{apo}
<i>Data collection</i>		
Unit cell (Å)	201.49, 75.93, 103.76, 90.00, 109.99, 90.00	165.21, 74.31, 133.72, 90.00, 119.30, 90.00
Space group	C2	C2
Resolution range* (Å)	97.52 - 1.74 (1.82-1.74)	116.60 – 1.58 (1.64 – 1.58)
Unique reflections	114961	193102
Completeness* (%)	76.2 (64.6)	98.1 (97.9)
R _{merge} *	0.053 (0.240)	0.049 (0.567)
I/σ(I)	11.9 (1.7)	12.7 (1.9)
Multiplicity	2.1	3.3
<i>Refinement</i>		
Resolution range* (Å)	97.52- 1.74 (1.78-1.74)	116.61 – 1.58 (1.62 – 1.58)
Reflection used	109213 (6421)	180084 (13980)
R/R _{free} *	15.9/19.9 (21.3/25.4)	15.9/19.2 (30.4/31.8)
rmsd from ideality (Å, °)	0.02, 2.04	0.021, 1.97
<i>N° atoms</i>		
Non-hydrogen atoms	10815	11118
Protein atoms	10223	10219
Solvent molecules	538	841
Hetero atoms	54	58
<i>B value (Å²)</i>		
Mean	17.7	26.2
Wilson plot	25.7	28.8
Protein atoms	17.5	26.1
Hetero atoms [§]	16.1	47.4
Solvent molecules	22.27	30.7
<i>Ramachandran plot (%)</i>		
Most favoured	97.7	97.1
Allowed	2.3	2.2
Disallowed	0.0	0.7

*The values in parenthesis refers to the last resolution shell.

§ The hetero atoms are 2 TPP molecules and 2 Mg²⁺ in CrTK_{TPP/Mg} and 1EDO (ethylene glycol) molecule in CrTK_{apo}.

Acknowledgements

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