Remodeling of the Binding Site of Nucleoside Diphosphate Kinase Revealed by X-ray Structure and H/D Exchange

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Supporting Information

ABSTRACT: To be fully active and participate in the metabolism of phosphorylated nucleotides, most nucleoside diphosphate kinases (NDPKs) have to assemble into stable hexamers. Here we studied the role played by six intersubunit salt bridges R80–D93 in the stability of NDPK from the pathogen Mycobacterium tuberculosis (Mt). Mutating R80 into Ala or Asn abolished the salt bridges. Unexpectedly, compensatory stabilizing mechanisms appeared for R80A and R80N mutants and we studied them by biochemical and structural methods. The R80A mutant crystallized into space group I222 that is unusual for NDPK, and its hexameric structure revealed the occurrence at the trimer interface of a stabilizing hydrophobic patch around the mutation. Functionally relevant, a trimer of the R80A hexamer showed a remodeling of the binding site. In this conformation, the cleft of the active site is more open, and then active His117 is more accessible to substrates. H/D exchange mass spectrometry analysis of the wild type and the R80A and R80N mutants showed that the remodeled region of the protein is highly solvent accessible, indicating that equilibrium between open and closed conformations is possible. We propose that such equilibrium occurs in vivo and explains how bulky substrates access the catalytic His117.

N nucleoside diphosphate kinases (NDPKs) are metabolic enzymes encoded by NME genes, also called NM23. They are present in all kingdoms of life and in certain viruses. NDPKs are responsible for the transfer of the γ-phosphate from nucleoside triphosphates (NTPs) to nucleoside diphosphates (NDPs), and in this way, they maintain the equilibrium between the pools of phosphorylated nucleosides (NTPs and NDPs).1 The γ-phosphate transfer reaction takes place in two steps according to a ping-pong mechanism, with transient phosphorylation of the enzyme on a 100% conserved histidine residue.2,7 NDPK recognizes with low specificity all tri- and diphosphate nucleosides.8 As expected from their intracellular concentrations, it has been shown that the major phosphate donor in the cell is the ATP while the major acceptor is the GDP.1 In mammals, by efficiently supplying with GTP the dynamin superfamily through direct interaction, researchers have recently shown that NDPK plays an important role in membrane remodeling events.5,6 Moreover, direct interaction between NDPK and different compounds or regulatory proteins from the cytoskeleton has been observed, indicating a clear role in nucleotide channeling and cell motility.5,6

Secondary enzymatic activities conserved from bacteria to humans have been established for NDPK. It can act as a protein histidine kinase,7 bind to single-stranded DNA, and exhibit 3′–5′ exonuclease activity, playing a potential role in DNA repair.8 Studies of Drosophila, zebrafish, and mice show a role of NDPK in early, larval, and embryo development.1 In mammals, 10 different NDPK isoforms exist. Isoforms 1–7 were characterized and present at least one of the enzymatic activities, while the function of isoforms 8–10 is less understood.1,8 Isoform 1, also known as NM23-A, is the most abundant one and was the first human metastasis suppressor to be reported.10–12 The molecular mechanism by which NM23-A inhibits in vivo the metastasis from spreading is largely unknown and has been linked to the role of NDPK in cell motility. However, the metastasis suppression function is dependent on the capacity of NDPK to perform its enzymatic activities:1,9 nucleoside diphosphate kinase,13 histidine kinase,7,14–16 and 3′–5′ exonuclease.8,17,18

To be functional, NDPK monomers have to autoassemble into hexameric or tetrameric complexes, sub-oligomeric species like monomers or dimers presenting at most 1–2% of the total activity.10 Hexamers are formed in eukaryotes, archaea, some viruses, and Gram-positive bacteria,19 while two different types

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of tetramers are present in Gram-negative bacteria. In terms of the subunit composition of complexes, NDPK hexamers as well as tetramers are homo-oligomeric or hetero-oligomeric.

Interestingly, hetero-oligomers are formed in vivo as well as in vitro, from different cellular isoforms, after infection from a mix of host and pathogen monomers, a potential way to have more interactions with other proteins and hence more cellular functions. All NDPK structures deposited at the Protein Data Bank (PDB) concern homo-oligomers. Figure 1A shows one monomer. The ferredoxin fold is conserved with two additional specific elements: the helix hairpin formed by helices $\alpha_2$ and $\alpha_3$ linking strands $\beta_1$ and $\beta_2$ and the Kpn loop (named from the Killer of prion lethal mutation in Drosophila) inserted between helices $\alpha_3$ and $\alpha_4$. All structures show that monomers are in the same conformation inside the hexamers or tetramers, in agreement with the idea that active sites are identical and independent.

Complete genome sequencing of Mycobacterium tuberculosis (Mt), the causative agent of tuberculosis, showed that it contains only one gene encoding NDPK (UniProtKB entry ASUSE1), and its crystal structure showed it forms hexamers [PDB entry 1k44 (Figure 1)]. In patients infected with tuberculosis, the bacterium is phagocytosed by macrophages but escapes degradation by preventing the fusion of the phagosome with the lysosome. The escape mechanism responsible for enhanced intracellular survival within macrophages is not fully understood but requires several thermolabile proteins, including NDPK. In this context, we previously studied the stabilizing mechanisms allowing Mt-NDPK to present a $15-20$ °C higher $T_m$ compared to those of NDPKs from other species. We showed that this unusually high stability is partially based on the presence of six intersubunit ionic bridges R80–D93 (Figure 1B). When the bridges were abolished by the D93N point mutation, the mutant was still hexameric and enzymatically active, but its $T_m$ dropped by 28 °C. Equilibrium analysis of the wild type (WT) and D93N mutant by hydrogen/deuterium exchange mass spectrometry (HDX-MS) allowed us to show that the R80–D93 bridge was not only important with respect to hexamer stability but also influenced its conformational dynamics.

We have continued our research to understand the role of ionic bonds in Mt-NDPK stability and conformational dynamics by mutating R80, the other amino acid involved in ionic bridge formation (Figure 1B). Mutants R80A and R80N (R80A/N) were produced, and their properties studied by enzymology, fluorescence, circular dichroism (CD), and size-exclusion chromatography (SEC). The structure of mutant R80A that crystallized into an unusual space group for NDPKs was determined. We noticed important conformational remodeling that has not been stated so far. It mainly concerned peripheral helices $\alpha_4$ and $\alpha_5$ that are involved in nucleotide binding and in it being locked inside the active site. In the current structure, the catalytic site is remodeled and becomes accessible for substrates larger than a nucleotide. To show that such a conformational change exists in solution, we performed at equilibrium HDX-MS measurements for WT, R80A, and R80N in the presence and absence of nucleotides. The HDX data that illuminated Mt-NDPK dynamics in solution were in perfect agreement with the structural remodeling observed in the crystal structure. We propose that the highlighted conformational change explains how bulky substrates access the catalytic His117.

**MATERIALS AND METHODS**

**Reagents.** Chemicals of the highest purity grade were bought from Sigma. Solutions of urea were freshly prepared for each experiment.

**Mutagenesis and Protein Purification.** The R80A gene mutation was introduced using a Transformer site-directed mutagenesis kit (Clontech), and the mutation was confirmed by nucleotide sequencing. The individual expression of WT, R80A, R80N, and D93N Mt-NDPK recombinant proteins was realized using a pET24 vector (Novagen) in the BL21-derived host strain BL21-CodonPlusH(DE3)-RIL (Stratagene). The 2YT culture medium contained 16 g/L bacto tryptone, 10 g/L bacto yeast extract, and 5 g/L sodium chloride, in the presence of 80 mg/mL kanamycin. The expression was induced with 1 mM isopropyl $\beta$-D-1-thiogalactopyranoside for 6 h at 37 °C, once the optical density reached 0.5–0.7 unit. As previously described, the purification steps were carried out at 4 °C. After being harvested, the Escherichia coli cells were sonicated and centrifuged to recover the soluble fraction containing Mt-NDPKs. The DNase-treated bacterial extract was loaded onto a Q-Sepharose column equilibrated in 100 mM Tris-HCl (pH 7.4). The enzyme was eluted at 0.5–0.6 M NaCl in a linear gradient from 0 to 0.8 M NaCl in the same buffer. Active fractions were precipitated with 80% saturated ammonium sulfate and further purified by salting-out chromatography on a Sepharose 6B column equilibrated with 80% ammonium sulfate and 100 mM Tris-HCl (pH 7.4). The protein was eluted by a linear gradient from 80 to 20% ammonium sulfate.

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Figure 1. View along the 3-fold axis of (A) a monomer and (B) a trimer of wild-type Mt-NDPK (PDB entry 1k44). The helix hairpin formed by helices $\alpha_2$ and $\alpha_3$ that link the second strand to the third strand of the central $\beta$-sheet is colored magenta. The intersubunit salt bridges R80–D93 contribute to the high stability of Mt-NDPK. The trimer interface is drawn as a dashed line. ADPs bound in the active sites are drawn as yellow surfaces by homology with NDPK-A. (C) Overall view of the hexamer. Two dimers are shown as pink and white surfaces, while the third is shown as a blue ribbon. The pink monomer shown in panels A and B is now at the bottom left of hexamer. This figure was drawn using PyMOL.
in the same buffer. The active fractions were pooled, dialyzed against 100 mM Tris-HCl (pH 7.4), and further purified on a Source 15Q column, under the conditions described for Q-Sepharose chromatography. The enzymes were precipitated by dialysis against a saturated solution of ammonium sulfate, recovered by centrifugation, and further purified by SEC on a Sephacryl S-200 column equilibrated with 0.2 M sodium phosphate buffer (pH 7.0). This step allowed the sample to be cleaned up, by eliminating aggregated and dissociated protein.

The enzymes were essentially pure as ascertained by polyacrylamide gel electrophoresis in the presence of sodium dodecyl sulfate. The concentrations of WT and mutant Mt-NDPKs were determined from the optical density at 280 nm using an extinction coefficient of 0.48 for 1 mg/mL, which was calculated from the amino acid composition. The molecular weight of proteins was checked by mass spectrometry.

Size-exclusion chromatograms were recorded on a Superose 12, 10/300 GL column equilibrated with 20 mM Tris-HCl (pH 7.0) and 100 mM sodium chloride to ensure that all proteins were homomeric. The following molecular markers were used: cytochrome c, 12.4 kDa; myoglobin, 17 kDa; carbonic anhydrase, 29 kDa; ovalbumin, 44 kDa; bovine serum albumin (BSA), 68 kDa; aldolase, 158 kDa; β-amylase, 200 kDa (GE Healthcare Life Sciences).

**Thermal Denaturation and Chemical Unfolding Experiments.** Thermal denaturation experiments were performed by incubating 10 μg/mL NDPK in a water bath and increasing the temperature at a rate of 1 °C/min. At different time points (corresponding to different temperatures), aliquots were taken and the enzyme activity was measured according to the NDPK enzymatic assay.

The chemical unfolding experiments were performed as previously described. A 10 μg/mL final concentration of native or unfolded Mt-NDPK was incubated for 16 h in 0–8 M urea or 0–5 M GuHCl and 20 mM phosphate buffer (pH 7.0) at 25 °C. Fluorescence intensities of the single tryptophan residue, Trp132, were measured at 335 nm with excitation at 295 nm.

**NDPK Enzymatic Assay.** The enzymatic activity of NDPK was measured using a coupled assay that involves the three independent reactions shown below. Enzymes performing the catalysis of each reaction are indicated in parentheses.

NDPK catalyzed the transfer of the γ-phosphate from ATP to 8-bromoinosine 5′-diphosphate (8-BrIDP). The reaction took place in 0.8 mL of an enzyme mix containing 1 mM ATP, 0.2 mM 8-BrIDP, 1 mM phosphoenolpyruvate (PEP), 0.1 mM NaH₂PO₄, 20 mM Tris-HCl (pH 7.5), 5 mM MgCl₂, 100 mM KCl, 1 mg/mL BSA, 2 units/mL pyruvate kinase, and lactate dehydrogenase. The disappearance of NADH was measured at 340 nm and 25 °C, using a PerkinElmer spectrophotometer.

\[
\text{ATP} + 8\text{-BrIDP} \rightarrow \text{ADP} + 8\text{BrITP (NDPK)}
\]

\[
\text{ADP} + \text{PEP} \rightarrow \text{ATP} + \text{pyruvate (pyruvate kinase)}
\]

\[
\text{pyruvate} + \text{NADH} \rightarrow \text{lactate} + \text{NAD}^+
\]

(lactate dehydrogenase)

**Crystallization and X-ray Diffraction Data Collection.** Recombinant Mt-NDPK-R80A was purified by Superdex 200 SEC in a buffer consisting of 100 mM NaCl and 20 mM Tris-HCl (pH 7.9). Collected peak fractions were pooled and concentrated to 12.1 mg/mL by ultrafiltration. Sparse-matrix screening of candidate crystallization conditions was set up on TTP Labtech IQ plates with mixes of 50 nL of protein and 50 nL of commercial precipitant solutions (Qiagen) and incubated at 20 °C. Monocrystalline 100 μm × 50 μm platelets of R80A Mt-NDPK grown for 5 weeks under condition JCSG IV 58 [0.8 M ammonium sulphate and 0.1 M HEPES-NaOH (pH 7.0)] were cryo-protected by addition of 25% glycerol and flash-frozen in liquid nitrogen. Two thousand images of a 0.1° rotation increment each were collected on the ID30A-3 MASSIF microfocus beamline at the European Synchrotron Radiation Facility (Grenoble, France) at 12.81 keV. Diffraction data were recorded on an Eiger X 4M detector. The data were indexed and integrated with IMOSFLM and SCALA in orthorhombic space group P2₁2₁2₁ with the following unit cell parameters: \(a = 113.1 \text{ Å}, b = 231.2 \text{ Å}, c = 150.71 \text{ Å}\). The data set was complete to >99% at a resolution of 2.2 Å. The structure was determined by a molecular replacement method with MOLREP using the highest-resolution available Mt-NDPK structure (PDB entry 4ane) as a search model. The unit cell contained six molecules in the asymmetric unit, resulting in a solvent content of 59% [Matthews coefficient (Vₘ) of 2.99 Å³ Da⁻¹].

**HDX-MS.** Stock solutions of 4.0 μM native Mt-NDPKs (WT and R80A/N) were prepared in 20 mM Tris-HCl (pH 4.0), 100 mM NaCl, and H₂O. Deuterium exchange was initiated by 20-fold dilution of the Mt-NDPK stock solution into 100 mM NaCl, 20 mM Tris-HCl (pH 7.0), and 99.9% D₂O at 10 °C. At different incubation times, a 100 μL aliquot was removed from the exchange reaction mixture (from 10 s to 245 min) and labeling was quenched at pH 2.5 by adding 10 μL of a solution of 1% formic acid and 5 M GuHCl at 0 °C. 100 μL of each acid-quenched sample was immediately injected into an HDX Waters nanoACQUITY ultra-performance liquid chromatography (UPLC) instrument. The sample passed through a Poroszyme-immobilized pepin 250 cartridge (Applied Biosystems) accommodated within the HDX manager at a flow rate of 100 μL min⁻¹ and 15 °C. Peptidic peptides eluting from the pepsin column were trapped and desalted for 3 min at a rate of 100 μL min⁻¹. Peptides were separated in 6 min with a 5 to 35% acetonitrile/water gradient in 0.1% formic acid, at a rate of 100 μL min⁻¹ with a 1.0 mm × 50.0 mm ACQUITY UPLC C18 HSS T3 column (Waters) containing 1.8 μm particles (back pressure of 13000 psi). All chromatographic elements were held at 0 °C for the entire time of the measurements. The calculated average amount of 261 back exchange was 31% on the basis of analysis of fully deuterated peptides of NDPK in GuDCl. Because all comparison experiments were performed under identical experimental conditions, there was no correction for back exchange. The error in the determination of the deuterium levels was ±0.20 Da in this experimental setup, consistent with previously obtained values. Mass spectra were recorded in HDMS mode using a Waters Synapt G2 Si instrument with a standard ESI source (Waters Corp., Milford, MA) over an m/z range of 50–2000. Mass accuracy was ensured by calibration with the Glu-fibrinogen peptide and was <10 ppm throughout all experiments. Identification of the peptic fragments was accomplished with at least four replicate HDMS analyses using PLGS 3.0 (Waters Corp.). Deuterium experiments were performed between two and four times. Mass spectra were processed using DynamX 3.0 (Waters). Automated selection of the isotope distribution was verified manually for all peptides and all charge states. The resulting relative deuterium levels...
were plotted versus the exchange-in time with respect to the described experimental conditions.

## RESULTS AND DISCUSSION

### Biochemical and Enzyme Properties of R80A/N Mt-NDPK

We previously studied the biochemical and enzyme properties of WT and D93N Mt-NDPK.\(^{20}\) Now, recombinant R80A and R80N Mt-NDPK were overexpressed in *E. coli* and purified without a tag. The mass of monomers was verified by mass spectrometry, and using SEC, we established that all proteins formed hexamers (Figure S1). Ultraviolet (UV), tryptophan fluorescence, and CD spectra were identical between WT and R80A/N. The enzyme specific activities of mutants were 350 and 320 units/mg, respectively (Table 1), close to that of the WT protein. Taken together, these results suggested that mutations did not significantly affect the overall three-dimensional protein structure and NDPK stability was studied next.

**Table 1**

<table>
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<th>PDB entry</th>
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<th>solvent content (%)</th>
<th>space group</th>
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<td>D93N</td>
<td>48.4</td>
<td>230</td>
<td>4anc</td>
<td>2.8</td>
<td>63</td>
<td>P4_320</td>
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<tr>
<td>R80N</td>
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<td>320</td>
<td>4ane</td>
<td>1.9</td>
<td>54</td>
<td>J2</td>
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<td>R80A</td>
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<td>350</td>
<td>6q22</td>
<td>2.2</td>
<td>59</td>
<td>J222</td>
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The enzymatic activity is strictly dependent on the presence of quaternary structure, meaning on correct assembled hexamers.\(^{19}\) Studying properties of natural mutants of NDPK from multiple species, different groups showed using SEC that such mutants can accumulate as stable monomers or dimers and present at most 1–2% of the hexamer activity.\(^{25,36–38,45}\) Figure 2B shows that the WT is inactivated at 5.3 M urea while R80A and R80N are inactivated at 4.2 M urea, indicating that hexamers of WT are more stable than those of the mutants.

Because the loss of activity and Trp fluorescence changes appear at similar urea concentrations for each NDPK, these results indicate that hexamers disassemble and unfold concomitantly, without accumulation of dissociated species. D93N hexamers exhibit a different behavior. As published, activity is lost at 0.3 M urea while a decrease in fluorescence appears at 2.5 M urea, indicating that hexamers disassemble before unfolding.\(^{20}\)

The stability of WT, R80A/N, and D93N hexamers was measured by the same procedure in the presence of GuHCl (Figure S2B). The WT was more stable than the three mutants. R80A and R80N exhibited similar stability but were more stable than D93N, a behavior similar to that determined in the presence of urea. Taken together, the results from both thermal stability and chemical stability show that mutating R80 or D93 to abrogate R80–D93 salt bridges has a different

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**Figure 2.** Unfolding pathway of WT, R80A, R80N, and D93N Mt-NDPK monitored by (A) intrinsic fluorescence and (B) residual enzymatic activity. Each sample was preincubated for 16 h in the presence of 0–7 M urea.
effect on NDPK stability. This suggested that compensatory effects exist for R80A/N but not for D93.

**Structural Remodeling of Three Subunits inside the ND PK Hexamer Is Revealed by the Structure of the R80A Mutant.** To understand the different effect of mutations R80A/N and D93N on hexamer stability, we next focused on the WT, R80N, and D93N (Table 1), and we determined here the structure of the R80A mutant at 2.2 Å resolution. Table 2 summarizes the X-ray data processing and refinement statistics. Structures were available for WT, R80N, and D93N (Table 1), and we determined here the 364 structure of the R80A mutant at 2.2 Å resolution. Table 2

<table>
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<tr>
<td>Rmerge on I (%)</td>
</tr>
<tr>
<td>Rmerge on I (%)</td>
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<tr>
<td>Rmerge on I (%)</td>
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<tr>
<td></td>
</tr>
<tr>
<td>Wilson B (Å²)</td>
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<tr>
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Model and Refinement

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<tr>
<td>all main/side chains</td>
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<tr>
<td>water/sulfate/Tris</td>
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aValues in parentheses are for the highest-resolution shell. Rmerge = \( \sum_{hkl} |I(hkl)| - \overline{|I(hkl)|} / \sum_{hkl} |I(hkl)| \), Rcryst = \( \sum_{hkl} |I(hkl)| - \overline{|I(hkl)|} / \sum_{hkl} |I(hkl)| \), Rfree = \( \sum_{hkl} |I(hkl)| - \overline{|I(hkl)|} / \sum_{hkl} |I(hkl)| \), and Rmerge (precision-indicating Rmerge) = \( \sum_{hkl} |I(hkl)| - \overline{|I(hkl)|} / \sum_{hkl} |I(hkl)| \), where N is the number of unique reflections. bThe average isotropic B includes TLS and residual B components. cRoot-mean-square deviation.

364 summarizing the X-ray data processing and refinement statistics. The crystals belong to the I222 space group with a single hexamer in the asymmetric unit. A one-dimensional infinite solvent channel with a cross section of 60 Å × 30 Å along the a 369 axis results in a high solvent content of 59% (Table 1 and Figure S3A). Only two other NDPK structures in the PDB (entries 4w98 and 4wbf) belong to this space group but in a different crystal packing. They both concern the NDPK from the bacterium *Acinetobacter baumannii*, a potential dimer from SEC experiments.36

On the 3-fold axis of the molecule, two Tris molecules are bound by Arg25, Arg28, and Asp106 side chains, and on the outer surface, sulfate ions are bound by each Arg86. A sulfate ion has also been modeled into each binding site as a ligand of His117, Arg104, and Arg86 (Figure S3B). Residual densities around this sulfate ion suggest the alternative presence of water, Tris, or HEPES molecules, but attempts to incorporate them into the model were not fully successful. Around mutation R80A, the trimer interface becomes highly hydrophobic, with a patch containing A80, A83, and A84 on one face and V95 and L109 on the other face (Figure S3C). R80A adopts a conserved ferredoxin fold of ~90 amino acids, formed by four antiparallel β-strands and two connecting α-helices (αt and αc) that cover one face of the β-sheet (Figure 3A). Residues 92–112 form the Kpn loop that contains a turn (amino acids 112–119) similar to the deep buried essential His117. In such an open conformation, moves to stack on the His117 imidazole ring 3.5 Å on the open one. NDPK structures containing a bound nucleotide show that it is sandwiched between the helix hairpin and the Kpn loop (Figure 1B,C). Amino acids involved in the nucleotide binding site are highly conserved, forming a well-defined cleft that is perfectly filled by one single phosphorylated nucleoside at a 4.2 Å resolution (Figure S3B). The active site is highly conserved, formed by crystal contacts, suggesting that in solution such conformations are possible. In trypanosomatid parasites, *Leishmania*, and *Giardia lamblia* NDPKs, these helices are not...
436 visible in the electron densities and were likely fully 437 disordered.38,39

HDX-MS Revealed Full Solvent Accessibility Only for 438 the Mt-NDPK Region That Can Be Remodeled.

Figure 3. (A) View of a monomer of chains A, C, and E and (C) view of a monomer of chains B, D, and F. The $\beta_2\alpha_2\alpha_2\beta_3$ region is remodeled in the latter monomers, and details are provided in panels B and D, respectively. Residues 40 and 41, 42−47, 48−57, 58−68, and 69 and 70 are colored blue, orange, cyan, green, and blue, respectively. The region covering residues 48−57 is not visible in the X-ray structure for monomers B, D, and F and is shown as a cyan dashed line in panels C and D. The $\alpha_2$ helix (B) and the successive $\beta$-turn structures (D) are shown as cartoons and ball-and-stick structures.

Figure 4. (A) Representative deuterium incorporation curves, including error bars, for peptides covering residues 39−45 and 46−62. The purple vertical line shows the maximal deuteration obtained experimentally for these peptides in the denatured state. (B) Top view showing the NDPK trimer. Residues covering amino acids 39−45 and 46−62 are colored magenta and brown, respectively. (C) Comparisons of relative deuterium exchange in WT vs R80A (left) and WT vs R80N (right). All relative differences are shown in daltons and color-coded according to the scale at the bottom. The left side shows the residue numbers of each peptide fragment, arranged in order from the N- to C-terminus (top to bottom, respectively). The deuterium incorporation graphs used to create this figure are shown in Figure S4B. Within both panels, the differences in uptake at various times, from 10 s to 120 min, are shown and indicated at the bottom of each panel.
question we answer is whether Mt-NDPK conformational changes observed by crystallography can be monitored in solution. HDX-MS is a method that has been applied to follow conformational changes and dynamics for many proteins, including serine, threonine, and tyrosine kinases. However, so far, only one study by our group used HDX-MS to study NDPK, which belongs to the histidine kinase family. We showed in this study that during folding dimers of D93N and WT-Mt-NDPK are stably formed. They are nativelike except for the minor subdomain (Kpn/α5), part of the active site, which is unfolded in the dimer and becomes structured only upon hexamerization. HDX-MS relies on the natural property that proteins rapidly exchange their backbone amide hydrogen atoms that are not involved in stable H-bonds and are accessible to a solvent water catalyst.

Exposing a protein of interest to a solvent containing D2O induces rapid replacement of the labile accessible backbone with deuterium (D). To spatially reveal where the exchange has happened, the incorporation of deuterium is measured by MS analysis after protein digestion and interpreted in light of the structure, if it is available.

The Mt-NDPK conformational changes revealed by crystallography concerned the region covering amino acids 40–70. We hypothesized that if in solution the NDPK shifts from one conformation to the other, maximal deuterium should be observed for this region. Here, we applied continuous labeling HDX-MS to native hexamers of WT, R80A, and R80N Mt-NDPK (see SEC in Figure S1). Samples were incubated at 10 °C from 10 s to 120 min in a buffer mimicking their physiological environment and containing a D2O final concentration of 95%. At different time points of deuteration, aliquots were taken, acid quenched, pepsin digested, separated by LC, and immediately introduced into a mass spectrometer for mass analysis.

There were 37 peptic peptides identified for the three proteins, and 15 of these peptides that were common to the three proteins were routinely analyzed by HDX (see the peptide map in Figure S4A). The region of amino acids 75–83 containing the site of mutation was covered for the WT and mutant proteins by three different peptides. The final sequence coverage was 91.2% for WT and R80N and 90.4% for R80A. The last six C-terminal residues covering amino acids 131–136 were not detected in these experiments with IMS. Previously, this region of the protein was maximally deuterated within 10 minutes (Mt-NDPK).

EX2 exchange was noticed with no exception for all peptides of WT and R80A/N (Figure S4B), in agreement with our previous study of WT and D93N. Deuterium incorporation plots were almost identical for the common peptides of the three proteins, and an increased deuteration level with time was observed for all analyzed peptides (Figure S4B for common peptides and peptides covering the mutation). When compared to values of maximal deuteration obtained from controls of fully denatured and deuterated NDPK in GuDCl, the region covering amino acids 39–62 was the only one for all three proteins to be flexible enough during the labeling experiment to reach maximal deuteration. Figure 4A illustrates the behavior of peptides 39–45 and 46–62 covering this region. Similar behavior was noticed for three redundant peptides covering this region (see peptides 37–45, 45–62, and 49–62 in Figure S4B). Because the maximal deuteration level appeared with a similar trend for these five peptides, this indicates that the region formed a subdomain showing high cooperativity. Importantly, this region is the one containing helices α5 and α6, which in the R80A crystal structure presented the conformational change (Figure 4B). Therefore, HDX-MS and crystallography give evidence of the high flexibility of this region of the protein that is essential for the full activity of NDPK. The other peptides of the three proteins showed for time points of <10 min modest deuteration, which was <50% of maximal deuteration (Figure S4B). The weakest exchange was noticed for peptides covering amino acids 2–36, which incorporated during 2 h <4 Da over 31 Da exchanged in the denatured peptides (<13% total deuteration). This indicates that a very stable core with modest dynamics exists in this area of the hexamer (amino acids 2–36). Interestingly, from an evolutionary point of view, this NDPK region is one of the most conserved.

We analyzed next the effect of the mutations R80A and R80N on the dynamics of the hexamer. The chicklet shown in Figure 4C indicates peptide by peptide the difference for all time points between WT and R80A and between WT and R80N. Interestingly, both mutants had dynamics very similar to those of the WT, except for the region covering amino acids 85–107 that was more dynamic in both mutants (Figure S4C). This region corresponds to the Kpn loop, a specific structure in NDPK known to be involved in hexamer stability and hexamer assembly. While a plateau of 7 Da incorporation was reached for this region after deuteration for 2 h for R80A/N, the same plateau was reached within 10 s for D93N. This indicates that mutation D93N that is thermodynamically more destabilizing than mutations R80A and R80N induced a higher flexibility not only around the mutation itself but also for the subdomain that contains it. Altogether, these results showed a strong correlation between protein stability and protein dynamics for the Mt-NDPK.

NDPK hexamers are fully phosphorylated by NTPs on catalytic histidine. We phosphorylated WT and R80A/N Mt-NDPK over 3 min by addition of ATP, followed by incubation in a D2O solution as previously described. Peptide deuterium plots for the phosphorylated versus nonphosphorylated proteins were identical, indicating that no conformational changes were induced by protein phosphorylation (Figure S4C). Similarly, no conformational changes were detected by continuous labeling HDX-MS after binding of AMP-PNP, a nonhydrolyzable analogue of ATP, to any of the three proteins (data not shown).

**CONCLUSIONS**

In most organisms, a majority of proteins are oligomers, their percentages increasing from bacteria to eukaryotes. Importantly, these assemblies have to be stable enough to perform their functions over different periods of time, which can stretch from minutes to years. Here we studied the role of intersubunit salt bridges R80–D93 in the stability of Mt-NDPK hexamers. Surprisingly, mutation of R80 or D93, each of which broke the salt bridge, did not have the same influence on hexamer stability, and compensatory mechanisms appeared. The D93N mutant was strongly destabilized, showing critical role of amino acids that belong to the Kpn loop (amino acids 92–112), while R80A and R80N were almost as stable as WT. The higher stability of the R80N mutant can be explained by the higher hydrogen bonds between the side chain of Asn80 and (i) the main chain carbonyl of Leu109 from the neighboring subunit and (ii) moreover the Asp93 carboxyl group via a water molecule. In the R80A mutant, these two bonds were absent.
and the hydrophobic patch around the mutation compensated for the loss of the salt bridge (Figure S3C).

During evolution, stabilization of NDPK complexes was performed by mechanisms that were different from organism to organism. In the NDPK from Dictyostelium discoideum, the quaternary structure counteracts the tendency of monomers to form molten globule structures. For the hyperthermophile bacteria Aquifex aeolicus, the stabilization of NDPK is realized through formation of intersubunit disulide bridges, while for the hyperthermophile archaea Pyrococcus aerophilum, it is performed with the help of two additional specific domains. Dimers of NDPK present a more modest stability compared to those of hexamers and tetramers. In this context, our study of Mt-NDPK showed a new stabilizing mechanism based on the presence of nonconserved intersubunit salt bridges R80–D93. Additionally, using HDX-MS methodology, we also showed a strong correlation for Mt-NDPK between protein stability and protein flexibility: the highest stability has the lowest flexibility.

According to its major enzymatic function in nucleotide homeostasis, the correct biochemical name of NDPK should be NTP/NDP transphosphorylase. Besides this function, NDPK has secondary functions as a protein histidine kinase, and a 3′–5′ exonuclease. For the histidine kinase activity, at least three defined substrates have so far been described: the β subunit of heterotrimeric G proteins, the intermediate conductance K⁺ channel K(Ca)3.1, and the Ca²⁺-dependent TRPV1 channel. With regard to the mechanism for the histidine kinase activity, a direct transfer of phosphate from the phosphorylated histidine (His117) in the active site of NDPK and the histidine in the substrate protein has been proposed. It is reasonable to assume that an opening of the active cleft must occur from the closed conformation, to allow the entry and the interaction of the target polypeptide with the NDPK catalytic His117. In all available structures of NDPK, the α₁−α₂ hairpin has atomic displacement parameters (B factors) higher than those of the rest of the protein. If the hairpin is involved in interhexamer contacts within the crystal, its electron density is strong. In some extreme cases, the density partially or totally disappears, the hairpin being fully unfolded. One NDPK presenting histidine kinase activity is the mammalian isoform Nm23-H2. Several structures of Nm23-H2 are available: one apo form, which is closed (PDB entry 1nue), one containing GDP, which is slightly more open (PDB entry 3bhf), and one with the most open cleft containing a dinucleotide (PDB entry 3bbb). The comparison of these structures of Nm23-H2 has highlighted the progressive opening of the α₁−α₂ hairpin without unfolding, and a model of interaction between Nm23-H2 and the c-myc gene has been proposed. For the R80A mutant of Mt-NDPK, the mutation induced an original crystal packing with a large solvent channel where the α₁−α₂ hairpin can unfold and refold in a new conformation with different secondary structures (Figure 3). The histidine kinase activity of Mt-NDPK has never been observed, but a secondary function of 3′–5′ exonuclease has been demonstrated. We overlapped the remodeled structure of the R80A mutant with three other NDPK structures: the apo WT from M. tuberculosis, the apo human WT isoform Nm23-H2, and the human WT isoform Nm23-H2 in complex with a dinucleotide (Figure S5A). Differences between these structures can be noticed, and they exemplify the dynamic behavior and the plasticity of the histidine kinase activity of NDPKs, we plan to perform H/D continuous labeling experiments with different human isoforms: H1 and H2, which present this activity, and isoforms H3 and H4, which do not. Applying HDX-MS technology to study how SK4 channels are activated by isoform H2 will be part of our future research program for better understanding the mechanism of the histidine kinase activity of NDPKs.

**ASSOCIATED CONTENT**

Supporting Information

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Figures S1–S5 (PDF)

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