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Structures of the dI₂dIII₁ Complex of Proton-Translocating Transhydrogenase with Bound, Inactive Analogues of NADH and NADPH Reveal Active Site Geometries^{†,‡}

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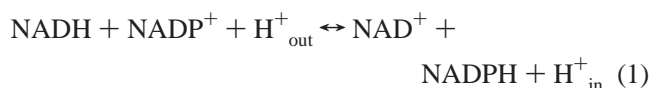
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ABSTRACT: Transhydrogenase couples the redox reaction between NADH and NADP⁺ to proton translocation across a membrane. The enzyme comprises three components; dI binds NAD(H), dIII binds NADP(H), and dII spans the membrane. The 1,4,5,6-tetrahydro analogue of NADH (designated H₂NADH) bound to isolated dI from *Rhodospirillum rubrum* transhydrogenase with similar affinity to the physiological nucleotide. Binding of either NADH or H₂NADH led to closure of the dI mobile loop. The 1,4,5,6-tetrahydro analogue of NADPH (H₂NADPH) bound very tightly to isolated *R. rubrum* dIII, but the rate constant for dissociation was greater than that for NADPH. The replacement of NADP⁺ on dIII either with H₂NADPH or with NADPH caused a similar set of chemical shift alterations, signifying an equivalent conformational change. Despite similar binding properties to the natural nucleotides, neither H₂NADH nor H₂NADPH could serve as a hydride donor in transhydrogenation reactions. Mixtures of dI and dIII form dI₂dIII₁ complexes. The nucleotide charge distribution of complexes loaded either with H₂NADH and NADP⁺ or with NAD⁺ and H₂NADPH should more closely mimic the ground states for forward and reverse hydride transfer, respectively, than previously studied dead-end species. Crystal structures of such complexes at 2.6 and 2.3 Å resolution are described. A transition state for hydride transfer between dihydronicotinamide and nicotinamide derivatives determined in ab initio quantum mechanical calculations resembles the organization of nucleotides in the transhydrogenase active site in the crystal structure. Molecular dynamics simulations of the enzyme indicate that the (dihydro)nicotinamide rings remain close to a ground state for hydride transfer throughout a 1.4 ns trajectory.

Transhydrogenase is found in the cytoplasmic membrane of bacteria and in the inner membrane of mitochondria from animal cells. It couples the transfer of hydride ion equivalents

between NAD(H)¹ and NADP(H) to the translocation of protons across the membrane.



Hydride transfer proceeds directly between bound nucleotides (1) and is stereospecific (2, 3): the *pro-R* hydrogen at C_{4N} of the dihydronicotinamide ring of NADH is transferred to C_{4N} on the *si* face of the nicotinamide ring of NADP⁺. In the “forward” direction, the enzyme is driven from left to right (eq 1) by the proton electrochemical gradient (Δp) generated by the respiratory (or sometimes photosynthetic) electron transport chain. One proton is translocated across the membrane through transhydrogenase per hydride ion transferred between nucleotides (4). There is broad agreement that conformational changes in the protein mediate coupling between the redox reaction and proton translocation (for reviews see refs 5–7). Kinetic (8) and mutagenesis (9–11) experiments indicate that binding changes at the NADP(H)-binding site are central in this process. Transhydrogenase produces NADPH for biosynthesis reactions and for the

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[‡] The two X-ray structures described in this paper are listed in the PDB as 2OOR (the dI₂dIII₁ complex with bound NAD⁺ and 1,4,5,6-tetrahydronicotinamide adenine dinucleotide phosphate) and 2O05 (the dI₂dIII₁ complex with bound 1,4,5,6-tetrahydronicotinamide adenine dinucleotide and NADP⁺).

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¹ Abbreviations: H₂NADH, 1,4,5,6-tetrahydronicotinamide adenine dinucleotide; H₂NADPH, 1,4,5,6-tetrahydronicotinamide adenine dinucleotide phosphate; AcPdAD⁺, acetylpyridine adenine dinucleotide (oxidized form); Nic⁺, nicotinamide; NicH, 1,4-dihydronicotinamide; H₂NicH, 1,4,5,6-tetrahydronicotinamide; NAD(H), NAD⁺ or NADH etc.; C_{4N}, the carbon-4 atom of the nicotinamide (dihydronicotinamide etc.) ring; dI, the NAD(H)-binding component of transhydrogenase; dIII, the NADP(H)-binding component; dII, the membrane-spanning component; dIII.E155W, the dIII component whose Glu155 is replaced by a Trp residue.

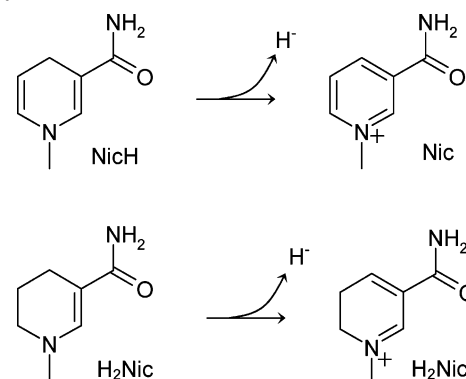
reduction of glutathione needed to minimize free radical damage (12, 13). Through a microcycle with the isocitrate dehydrogenases, it may participate in the control of the Krebs cycle in some cell types (14). Defects in the transhydrogenase gene lead to altered pathways of amino acid metabolism in *Escherichia coli* (15), to greater susceptibility to oxidative stress in *Rhodobacter sphaeroides* (16) and *Caenorhabditis elegans* (17), and to impaired control of glucose homeostasis in mice (18).

Two of the protein components of transhydrogenase, dI, which binds NAD(H), and dIII, which binds NADP(H), protrude from the membrane (on the cytoplasmic side in bacteria, the matrix side in mitochondria), and a third component, dII, spans the membrane [in 12–14 transmembrane helices, depending on species (19, 20)]. The enzyme is probably a “dimer” of two dI–dII–dIII “monomers”, though the polypeptide composition varies between species (see ref 21). The crystal structure of intact transhydrogenase is unavailable, but structures of isolated recombinant dI (22–24) and dIII (25–27) have been solved for several species.

Mixtures of isolated dI and dIII from *Rhodospirillum rubrum* form a heterotrimeric dI₂dIII₁ complex, which can catalyze a single turnover of rapid hydride transfer between bound NAD(H) and NADP(H) or their analogues, even in the absence of the membrane-spanning dII (1, 28–31). Crystal structures of the complex were obtained with bound NAD⁺ and NADP⁺ (32) and with bound NADH and NADPH (33). Having both nucleotides oxidized, or both reduced, prevents hydride transfer during crystallization. The structures of these “dead-end” complexes show that the hydride-transfer site is located at the dI/dIII interface. Of particular interest is the finding that the Nic(H) ring of the bound NAD(H) can occupy two positions, “distal” and “proximal” relative to the Nic(H) ring of the bound NADP(H). During the switch between the distal and proximal positions, the adenosine moiety of the NAD(H) remains fixed in its binding pocket, but rotations of bonds in the diphosphate and ribose phosphate region displace the Nic(H) ring. It was suggested that the positional switch of the Nic(H) ring of the NAD(H) may be important in gating the hydride-transfer reaction during proton translocation by the intact enzyme, and a plausible pathway linking the intermediate protein- and nucleotide-structural states involved in proton translocation was outlined (5). The asymmetries of the dI₂–dIII₁ complex indicate that the two dI–dII–dIII monomers of intact transhydrogenase proceed through alternating, coupled, out-of-phase conformational changes during turnover (21, 32).

In the ground state for hydride transfer, the active site will be occupied by NADH and NADP⁺ during forward transhydrogenation (eq 1) and by NAD⁺ and NADPH during reverse transhydrogenation. Because of their different charge balances, there is a possibility that the structures of NAD⁺/NADP⁺ and NADH/NADPH dead-end complexes might not accurately reflect the active site geometry in productive complexes. Recent attempts to crystallize dI₂dIII₁ with NADH (in excess) and NADP⁺ have led either to dead-end NADH/NADPH complexes or to complexes with nucleotides having an unknown redox state (34). In another attempt to determine the structure of the hydride-transfer site and the important conformational changes that occur during the redox reaction, we now describe the results of experiments

Scheme 1: Biological Oxidation of 1,4-Dihydronicotinamide (NicH) to Nicotinamide (Nic⁺) and of 1,4,5,6-Tetrahydronicotinamide (H₂NicH) to 5,6-Dihydronicotinamide (H₂Nic⁺)^a



^a NicH and Nic⁺ represent the reduced and oxidized nicotinamide moieties of the physiological nucleotides, NAD(P)(H). H₂NicH and H₂Nic⁺ represent the reduced and biologically oxidized (see text) moieties of the analogue nucleotides, H₂NAD(P)(H).

performed with 1,4,5,6-tetrahydro analogues of NADH and NADPH, in which the C5–C6 double bond of the dihydronicotinamide (NicH) ring of the physiological nucleotide has been reduced to a 1,4,5,6-tetrahydronicotinamide (H₂NicH) ring (see Scheme 1). These nucleotide analogues (H₂NADH and H₂NADPH, respectively) have been shown to bind like NAD(P)H to a number of soluble dehydrogenases but to be blocked in their on-enzyme redox chemistry (35–41). Crystal structures of liver alcohol dehydrogenase (40) and malate dehydrogenase (41) with H₂NADH show binding interactions and conformational changes similar to those seen with the physiological nucleotide, indicating that the analogue provides a good model for the active site. Loading the transhydrogenase dI₂dIII₁ complex with combinations of analogue and physiological nucleotides enables us to reconstruct the active site of this proton-translocating enzyme with a charge balance that more closely resembles the ground states for hydride transfer than is found in dead-end complexes.

EXPERIMENTAL PROCEDURES

Cultures of *R. rubrum* were grown under phototrophic conditions in RCV medium (42) supplemented with either biotin (in the case of the wild-type strain, S1) or biotin *plus* tetracycline [in the case of the transhydrogenase-overexpressing strain, RTB2 (4)]. Intracytoplasmic membrane vesicles (chromatophores) were prepared by French press treatment of harvested cells (43) and resuspended in 10% sucrose (w/v) and 100 mM Tris-HCl, pH 8.0. The bacteriochlorophyll concentration was measured using the *in vivo* extinction coefficient given (44).

Wild-type dI and the wild-type and E155W mutant of dIII from *R. rubrum* transhydrogenase were expressed from plasmids pCD1 (45), pNIC2 (46), and pJDV1 (47), respectively, in *E. coli* strains C600 (for dI) and BL21(DE3) (for dIII), and then purified by column chromatography and stored at –20 °C, as described in the earlier reports. The proteins were routinely >95% pure as judged by SDS–PAGE. Their concentrations were determined by the microtannin assay (48); values are expressed as monomers unless otherwise stated.

H₂NADH and H₂NADPH were prepared by hydrogenation of NADH and NADPH, respectively, in water at pH 8.5 (NaOH) using a 10% Pd/C catalyst (Fluka), essentially as described (35, 49). The reaction was terminated when 1.0 mol of hydrogen/mol of nucleotide was consumed. After removal of the catalyst by filtration through Celite, the H₂-NAD(P)H was purified by chromatography on Q-Sepharose Fast Flow (Amersham Biosciences) with a 4–400 mM gradient of NH₄HCO₃. The physiological nucleotides and the analogues were identified by their UV absorbance spectra (49). Purified fractions were lyophilized three times to remove NH₄HCO₃ and stored at –20 °C. The absorbance spectra of solutions used for experiments were checked before use (see ref 50). Unlike NAD(P)H, neither H₂NADH nor H₂NADPH was significantly fluorescent.

“Reverse” transhydrogenation was measured as the reduction of AcPdAD⁺, an analogue of NAD⁺, by NADPH from the absorbance change at 375 nm [$\epsilon = 6.1 \text{ mM}^{-1} \text{ cm}^{-1}$ (51)]. Measurements in the steady state were performed on a Shimadzu UV2401 and in the transient state after rapid mixing on a DX-17MV Applied Photophysics stopped-flow spectrophotometer. As normally prepared, isolated dIII is associated with tightly bound NADP⁺. For stopped-flow experiments, stored dIII (40 μM) was dialyzed overnight at 4 °C against 10 mM (NH₄)₂SO₄ and 20 mM Hepes, pH 8.0 (“SF buffer”). The protein was then concentrated to about 800 μM using a Vivaspin 20 kDa filter. A sample of this solution was mixed with H₂NADPH in SF buffer to give 100 μM dIII and 200 μM nucleotide, and this was incubated on ice for 2 h. An equal volume of a 200 μM solution of dI in SF buffer was added, and the mixture was loaded into one drive syringe of the stopped-flow machine; the other drive syringe contained 2.0 mM AcPdAD⁺ in SF buffer. Parallel experiments were performed as a control using NADPH instead of H₂NADPH. The stopped-flow machine was operated with 1:1 mixing at 20 °C; the dead time was 1.31 ms (*I*).

Fluorescence changes were recorded on a Spex Fluoromax. Isothermal titration calorimetry on isolated dI was initially carried out at 20 °C using the MicroCal MCS system, as described (52), but further experiments, including those with the dI₂dIII₁ complex, were performed (with guidance from Margaret Nutley and Alan Cooper at the University of Glasgow) on a MicroCal VP to improve resolution. One-dimensional ¹H NMR experiments on dI were performed using a Bruker AMX 500 MHz spectrometer equipped with an HCN 5 mm Z-PFG cryoprobe essentially as described (53). The protein for these experiments was washed in 10 mM [2H]Tris-HCl, pH 7.6, 1.0 mM dithiothreitol, 10 mM (NH₄)₂SO₄, and 0.025% NaN₃ in ²H₂O and concentrated to 405 μM in the same buffer by centrifugation through Vivaspin 10 kDa filters before being loaded into the NMR tube. Heteronuclear spin quantum coherence (HSQC) spectra of 200 μM ¹⁵N-labeled dIII protein in its NADP⁺ form, prepared as described (54), were recorded at 30 °C on a Varian Inova 800 MHz spectrometer equipped with a room temperature 5 mm ¹H/¹³C/¹⁵N Z-PFG probe. The bound NADP⁺ was then exchanged for either NADPH or H₂-NADPH, also as described (54), and HSQC spectra were recorded again. The data were processed using the AZARA 2.7 software suite (Boucher, <http://www.bio.cam.ac.uk/azara/>) and analyzed and visualized with the OpenGL version of

Table 1: X-ray Data Collection and Refinement Statistics

| | dI ₂ dIII ₁ | |
|--|---------------------------------------|--|
| | H ₂ NADH•NADP ⁺ | NAD ⁺ •H ₂ NADPH |
| data collection statistics ^a | | |
| unit cell parameters | | |
| <i>a</i> , <i>b</i> , <i>c</i> (Å) | 71.2, 101.0, 131.7 | 72.2, 74.4, 204.8 |
| α , β , γ (deg) | 90, 90, 90 | 90, 90, 90 |
| resolution (Å) | 62.6–2.6 (2.74–2.60) | 68.0–2.32 (2.45–2.32) |
| completeness (%) | 92.8 (94.4) | 91.4 (69) |
| multiplicity | 3.8 (3.1) | 4.2 (3.6) |
| <i>I</i> / σI | 15.1 (4.0) | 16.5 (3.4) |
| no. of observations | 103525 (12512) | 184225 (17038) |
| no. of unique observations | 27548 (4046) | 44362 (4726) |
| <i>R</i> _{sym} ^b | 7.3 (22.7) | 5.3 (31.7) |
| refinement statistics | | |
| Wilson <i>B</i> /average <i>B</i> ^c | 49.9/40.7 | 47.4/57.5 |
| no. of non-hydrogen atoms/waters | 7084/27 | 6984/101 |
| rmsd ^d bond (Å)/angle (deg) | 0.008/1.252 | 0.009/1.337 |
| Ramachandran ^e (%) | 91.5, 8.3, 0.1, 0.1 | 91.6, 7.9, 0.5, 0.0 |
| <i>R</i> / <i>R</i> _{free} | 21.9/27.5 | 24.7/27.5 |
| PDB code | 2OO5 | 2OOR |

^a Numbers in parentheses indicate data for the highest resolution shell. ^b $R_{\text{sym}} = \sum_j |I_j| / \sum_j I_j$, where I_j is the intensity of the *j*th reflection and $\langle I \rangle$ is the average intensity. ^c As calculated for all atoms. ^d rmsd is the root mean square deviation from ideal values. ^e The percentage of residues in the core, allowed, generously allowed, and disallowed regions, respectively, as measured by the program PROCHECK.

ANSIG (55). The combined chemical shift changes ($\Delta\delta$) in backbone ¹H and ¹⁵N atoms upon nucleotide replacement were calculated as described (56).

The dI and dIII proteins were prepared for crystallography by dialysis against 10 mM Hepes, pH 8.0, and then spin concentrated (see above). The dI₂dIII₁ complex in its H₂-NADH/NADP⁺ form was crystallized from equimolar solutions of dI and dIII containing 5 mM H₂NADH and 5 mM NADP⁺ in 100 mM Tris-HCl, pH 8.0, 75 mM Li₂SO₄, 16–20% 4K polyethylene glycol, and 10% glycerol. Before the dI₂dIII₁ complex was crystallized in its NAD⁺/H₂NADPH form, NADP⁺ on the dIII protein was exchanged by incubating the protein solution (300 μM) with 5 mM H₂-NADPH for 2 h. Crystals were then grown from equimolar solutions of dI and dIII containing 50 mM NAD⁺ and 5 mM H₂NADPH in 25–70 mM ammonium sulfate, 100 mM Mes, pH 6.5, 15–18% 4K polyethylene glycol, and 10% glycerol. Complete data sets of the H₂NADH/NADP⁺ crystals were collected to 2.6 Å on beamline ID14-4 at the European Synchrotron Radiation Facility at Grenoble, France, and of the NAD⁺/H₂NADPH crystals to 2.3 Å on beamline ID14-3. The data were integrated and scaled using the programs MOSFLM (57) and SCALA (58). The H₂NADH/NADP⁺ crystal had different cell dimensions (Table 1) to those previously reported for the NADH/NADPH crystal (PDB code 1U2D), and therefore, molecular replacement was used to determine the initial phases. The 1U2D structure was used as a search model against the data (up to 4 Å) for the H₂-NADH/NADP⁺ crystal in the program EPMR (59). The solution from molecular replacement was subjected to rigid body refinement in REFMAC5 (60). Simulated annealing protocols were also applied using the program CNS (61). The NAD⁺/H₂NADPH crystal was isomorphous with the

NAD⁺/NADP⁺ structure (PDB code 1HZZ), which was subsequently used to refine against the structure factor amplitudes of the NAD⁺/H₂NADPH crystal using REFMAC5. Simulated annealing protocols were applied using CNS before the model was adjusted to accommodate the new nucleotide combination. The two new structures were further subjected to constrained refinement with TLS in REFMAC5. The final models were assessed and adjusted according to the programs Whatcheck (62) and Procheck (63). Superposition of crystal structures was performed using LSQKAB from the CCP4 program suite, and domain rotations in dI were analyzed with the MORPH server (64) (Figure 8).

The molecular dynamics (MD) study was performed on the crystal structure of the dI₂dIII₁ complex (PDB code 2O05) using AMBER 8 (65). Atomic partial charges for the nicotinamide nucleotides were determined using the method (66). The atom-centered charges on the Nic⁺ and H₂NicH rings, and the diphosphate and 2'-phosphate groups, were derived in two stages coordinated by RED II (www.u-picardie.fr/labo/lbpd/RED). First, GAUSSIAN 98 (67), using the HF/6-31G* basis set, performed an ab initio structural optimization to calculate a three-dimensional molecular electrostatic potential. Then, in accordance with the RESP method (68), a two-stage fitting procedure was performed (RESP from AMBER 8) to calculate a model of atom-centered partial charges. The atom-centered charges of the adenine and ribose moieties were obtained from the RA3 R-adenosine unit in the AMBER ff94 force field (69). The structural parameters for the nicotinamide nucleotides were provided by the general AMBER force field (gaff) (70). The amino acids were described using the AMBER ff03 force field (71). The protein was stripped of crystallographic waters and then solvated with 32813 TIP3P waters (72). The solvent coordinates were then minimized, and the water positions were randomized by a short MD simulation of 20 ps at 300 K. Following this, the entire system (112709 atoms) was minimized and subjected to a 250 ps equilibration at 300 K. A time step of 2 fs was employed throughout, and atomic velocities were scaled using the Berendsen algorithm (73) to maintain the system temperature at 300 K. MD simulations were performed for 1 ns using the NVT ensemble, and coordinates were saved every 0.5 ps. Bonds between hydrogens and heavy atoms were constrained using the SHAKE algorithm (74), long-range electrostatics were calculated using the particle-mesh Ewald method (75), and nonbonding interactions were cut off at an 8 Å radius. All calculations were performed on the escience cluster at the University of Birmingham. Structures along the MD trajectory were manipulated, ring centers were defined, and ring tilts were determined, using GEOMALC from the CCP4 suite of programs.

Transition states for hydride transfer were determined using density function theory with GAUSSIAN98. A combination of Barone's 1-parameter modified Perdew-Wang 91 exchange functional and the Perdew-Wang 91 correlation functional (MPW1PW91) was used along with a 6-31G** basis set. Transition states were characterized by means of frequency calculations to confirm that they were genuine first-order transition states and that the transition vector corresponded to the hydride transfer reaction. *N*-Methylnicotinamide was used as a model for both NADP(H) and NAD(H). Starting structures were generated from initial

transition state calculations for hydride transfer between dihydropyridine and protonated pyridinium cation following the approach of Wu et al. (76). Amide groups were added at appropriate positions to a C₂ symmetrical structure (with dihydropyridine rings rotated by 120° around a C₄–C₄ axis, thus corresponding to the arrangement seen in the crystal structure) and a C_{2v} structure previously indicated to give the lowest energy transition state (76). The C_{2h} structure of Wu et al. (76) was not considered as this previous work had indicated that it led to transition states of considerably higher energy. Thus, starting points were generated for all possible combinations of amide position and orientation on a C₂ symmetrical core in addition to those corresponding to the “cis” and “trans” structures of Wu et al. (76). It should be noted, however, that the calculations described here are at a higher level of theory and involve a model modified from that of Wu et al. The results therefore are not strictly comparable (76). A further difference is that we have considered arrangements of amide bonds that generate asymmetric transition states.

The structure cartoons (Figures 3, 4, 7, and 8) were constructed using the program PYMOL (77) and the diagrams in Figure 5f,g using SWISS-PDV VIEWER. Amino acid residues are numbered according to the convention adopted (32).

RESULTS

Binding of H₂NADH to the dI Component of Transhydrogenase. To assess whether H₂NADH and H₂NADPH are good analogues of NADH and NADPH, respectively, their binding to the recombinant dI and dIII components of *R. rubrum* transhydrogenase was investigated. *R. rubrum* dI has a single Trp residue (at position 72) whose fluorescence emission is quenched upon addition of NADH but not NADPH (45). Addition of H₂NADH to a solution of isolated dI also led to the quenching of Trp72 fluorescence. *K_d* values calculated from titrations, after appropriate corrections for inner filtering effects, were similar: 18 μM for NADH (45) and 27 μM for H₂NADH (see Figure S1, Supporting Information). The binding of H₂NAD to isolated dI was also studied by isothermal titration calorimetry. The experiments revealed binding characteristics very similar to those of NADH (52). The *K_d* for H₂NADH was 24 μM [18 μM for NADH (52)], the enthalpy change was −45 kJ mol^{−1} dI monomer (−62 kJ mol^{−1} monomer for NADH), and for both the analogue and the natural nucleotide, the binding stoichiometry was approximately 1 per dI monomer (see Figure S2, Supporting Information).

In one-dimensional proton NMR experiments, the addition of NADH to isolated dI leads to line broadening and to changes in the chemical shift of relatively narrow resonances arising from a “mobile loop” (residues 220–240) in the protein (53, 78). The effects are attributable to a loss in mobility of residues in the loop as it closes down on the protein surface during nucleotide binding. The addition of H₂NADH to a solution of dI had very similar effects on the NMR spectrum: the same set of resonances was affected (see ref 78 for partial assignments) and in an equivalent way (Figure S3, Supporting Information). This indicates that the physiological and analogue nucleotides cause a similar conformational change in dI upon binding. The concentration

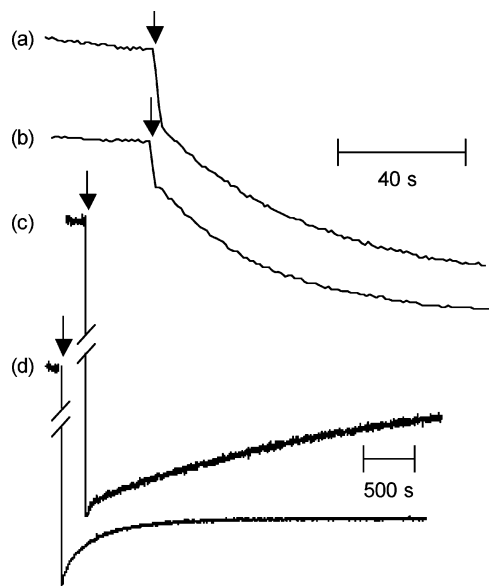


FIGURE 1: Changes in the fluorescence of Trp155 as dIII.E155W undergoes nucleotide exchange. (a) The fluorescence emission at 338 nm from excitation at 280 nm (4 nm band-pass) was recorded during addition of 10 μ M NADPH (arrow) to a solution of the NADP⁺ form of dIII.E155W (1.0 μ M) in 20 mM MOPS, pH 7.2 at 25 °C. (b) The fluorescence emission was recorded during addition of 10 μ M H₂NADPH to another sample of the same protein under similar conditions. (c) The NADP⁺ on dIII.E155W was first exchanged with NADPH as described in the text, and the 338–280 nm fluorescence change was then recorded during addition of 500 μ M NADP⁺ to a 1.0 μ M solution of this protein in the conditions stated above. The large, initial inner-filtering effect of the nucleotide has been partially removed from the trace where indicated. (d) As in (c) except that the NADP⁺ on dIII.E155W was first exchanged with H₂NADPH.

dependences were similar for NADH and for H₂NADH, but because the nucleotides are in intermediate exchange with the protein, the broadened resonances make the end points of the titrations difficult to judge accurately. Nevertheless, the data are consistent with K_d values in the order of 20 μ M for both NADH and H₂NADH.

It should be noted that NAD⁺ binds much more weakly to dI than either NADH or H₂NADH. The K_d for NAD⁺ is approximately 300 μ M (79, 80).

Binding of H₂NADPH to the dIII Component of Transhydrogenase. The isolated dIII component of *R. rubrum* transhydrogenase binds NADP(H) very tightly (28). The K_d values are too low to measure. To compare the binding of H₂NADPH with that of NADPH, we have performed experiments to measure rates of nucleotide exchange from dIII. In dIII.E155W, a unique Trp residue replaces the wild-type Glu155. The catalytic properties of the mutant protein are similar to those of wild-type dIII, but the fluorescence emission from the Trp usefully reflects nucleotide occupation (47) (and see ref 81). Thus, the Trp fluorescence is greater when NADP⁺ is bound to the protein than when NADPH is bound. In the experiments shown in panels a and b of Figure 1, NADPH and H₂NADPH, respectively, were added to solutions containing dIII in its NADP⁺ form. Following the rapid initial decrease due to the inner-filtering effect of the added nucleotide, the Trp155 fluorescence slowly declined over a period of > 100 s with approximately similar kinetics. According to earlier reasoning (47, 81), the results suggest that slow dissociation of bound NADP⁺ from dIII ($k_{off} \approx$

0.03 s⁻¹) is followed by the rebinding of either NADPH or H₂NADPH. Evidently, H₂NADPH can bind to dIII and quench Trp fluorescence in a similar way to the natural nucleotide.

In another set of experiments dIII.E155W was preincubated with either NADPH (Figure 1c) or H₂NADPH (Figure 1d) to load the protein with the respective nucleotide. A relatively high concentration of NADP⁺ was then added to each of the two protein samples. Following the prompt fluorescence decrease due to inner-filtering effects, there was a very slow increase in fluorescence in both experiments that is attributable to nucleotide exchange. The very slow rate of exchange is consistent with the tight binding of NADPH and H₂NADPH to dIII (i.e., slow rates of dissociation). When the concentration of added NADP⁺ is high enough, the exchange becomes limited by the dissociation of NADPH/H₂NADPH from the protein, and therefore, in the limit, the rate constant for the fluorescence increase corresponds to that for NADPH/H₂NADPH release. The dissociation rate constant for dIII.NADPH estimated from Figure 1c is similar to that described in an earlier report ($\approx 4 \times 10^{-4}$ s⁻¹), but that for dIII.H₂NADPH (though still small in absolute terms) is significantly greater than for the physiological nucleotide, about 3×10^{-3} s⁻¹ (Figure 1d).

In the HSQC experiment the spins of ¹H and ¹⁵N atoms in amide groups (both side chain and backbone) are correlated. The HSQC spectrum of isolated dIII from *R. rubrum* transhydrogenase in its NADP⁺-bound form is almost fully assigned (27). When the bound NADP⁺ is replaced by NADPH, there are chemical shift changes in distinctive subsets of the spectral peaks (54). Some of these chemical shift changes derive from amide groups of amino acid residues either close to or in contact with the NADP(H), but of particular interest are sensitive residues which extend some distance from the bound nucleotide into helix A, helix B, helix D/loop D, and loop E. The results indicate that there are changes in the magnetic environment within these regions, conformational changes in the protein, when the bound nucleotide becomes reduced. Similar data have been reported for isolated *E. coli* dIII (82). Figure 2 (top panel) shows the chemical shift changes in the HSQC spectrum of *R. rubrum* dIII when bound NADP⁺ was replaced by H₂NADPH. The changes are in the same regions of the polypeptide as those affected when bound NADP⁺ is replaced by NADPH, indicating that the two reduced nucleotides have analogous effects on the dIII structure. However, Figure 2 (lower panel) reveals a difference in the extent of the chemical shift changes caused by NADPH and H₂NADPH. Relative to the chemical shift changes induced by replacing NADP⁺ by NADPH, those induced by replacing NADP⁺ by H₂NADPH are less pronounced in helix D/loop D (residues 135–146) than in helices A and B and loop E. Thus, the character of the conformational change caused by the nucleotide substitution seems to be subtly different for H₂NADPH and for NADPH in helix D/loop D. Conformational changes are not observed in dIII crystal structures when NADP⁺ is replaced by NADPH (33, 83), and therefore, those detected by NMR may correspond only to small atomic displacements. However, in one set of crystallization conditions a conformational rearrangement is evident in the helix D/loop D region but is similar when either NADP⁺ or NADPH is bound (83). Changes in the redox state of bound

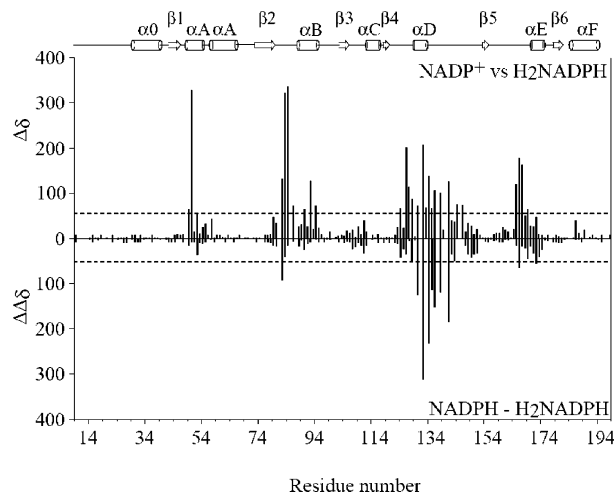


FIGURE 2: Chemical shift changes in dIII caused by substitution of bound NADP^+ with H_2NADPH and with NADPH . Upper panel: The HSQC spectrum of ^{15}N -labeled dIII in its NADP^+ form was recorded as described in the Experimental Procedures. Excess H_2NADPH was then added to the sample, and the spectrum was rerecorded (compare ref 54). The change in chemical shifts of backbone amides between the two spectra, $\Delta\delta$, is plotted as a function of residue number. Lower panel: An equivalent experiment was performed but using NADPH instead of H_2NADPH (as in ref 54). The $\Delta\Delta\delta$ values (i.e., the difference between $\Delta\delta$ from the NADP^+ minus H_2NADPH spectra and $\Delta\delta$ from the NADP^+ minus NADPH spectra) are plotted for each residue. The dashed lines correspond to twice the mean perturbation value (50 Hz) and indicate a significant change. The secondary structure of dIII is represented at the top of the figure.

NADP(H) are required to signal a reset in the proton translocation gate of dII (8), and the signal transmission pathway between dIII and dII might involve helix D/loop D (5). The small difference in structure between NADPH and H_2NADPH that is sensed by the chemical shift changes may be a reflection of this event.

H_2NADH and H_2NADPH Are Not Hydride Donors in the Transhydrogenase Reaction. Forward transhydrogenation (see eq 1) is conveniently measured by following the reduction of thio- NADP^+ (an analogue of NADP^+) with NADH . As in earlier experiments (84), the rate of this reaction was low in a sample suspension of darkened chromatophore membranes from *R. rubrum* strain RTB2 [$0.29 \mu\text{mol}$ of thio- NADP^+ reduced (μmol of bacteriochlorophyll) $^{-1} \text{ min}^{-1}$] but was accelerated about 6-fold upon illumination with near-infrared light. Stimulation results from the generation of Δp by the photosynthetic electron transport chain: the increased Δp drives H^+ back across the membrane through the transhydrogenase. In contrast to NADH , H_2NADH was unable to reduce thio- NADP^+ at a measurable rate in either darkened or illuminated chromatophore suspensions.

Reverse transhydrogenation can be monitored by following the reduction of AcPdAD^+ (an NAD^+ analogue) by NADPH (51). Wild-type chromatophore membranes catalyzed a rapid rate of this reaction [$2.7 \mu\text{mol}$ of AcPdAD^+ reduced (μmol of bacteriochlorophyll) $^{-1} \text{ min}^{-1}$] but failed to catalyze AcPdAD^+ reduction with H_2NADPH . In a separate series of experiments (Figure S4, Supporting Information), it was found that the reduction of AcPdAD^+ by NADPH was inhibited by H_2NADH and by H_2NADPH , consistent with a competitive interaction at the respective binding sites.

Mixtures of purified dI and dIII from *R. rubrum* transhydrogenase spontaneously form a dI_2dIII_1 complex which catalyzes a rapid, single turnover burst of hydride transfer (1, 28, 52). Further turnovers are heavily limited by the very slow rate of release of product NADP(H) from the dIII component of the complex. The kinetics of the hydride-transfer burst in dI_2dIII_1 complexes can be observed in stopped-flow experiments using NADPH and AcPdAD^+ as substrates; the apparent first-order rate constant (k_{app}) is $\approx 600 \text{ s}^{-1}$ (30). However, under conditions similar to those described earlier there was no detectable AcPdAD^+ reduction when NADPH was replaced by H_2NADPH . Within the experimental constraints k_{app} for this reaction is $< 3 \text{ s}^{-1}$ (data not shown).

The dI_2dIII_1 complex catalyzes a rapid, steady-state “cyclic transhydrogenation” reaction in which tightly bound NADP(H) is alternately reduced by NADH and oxidized by AcPdAD^+ (28). However, a combination of H_2NADH and AcPdAD^+ did not lead to any measurable cyclic transhydrogenation in dI_2dIII_1 complexes. This shows that H_2NADH cannot donate hydride equivalents to bound NADP^+ at a significant rate. In another set of experiments, isolated dIII ($3 \mu\text{M}$) in its NADP^+ -bound form was preincubated with either $100 \mu\text{M}$ NADPH or $100 \mu\text{M}$ H_2NADPH for 2 h. This leads to exchange of the bound NADP^+ with either NADPH or H_2NADPH , respectively (see above). After then adding dI to generate dI_2dIII_1 complexes, cyclic transhydrogenation was assayed with NADH and AcPdAD^+ . Where dIII had been preincubated with NADPH , the rate of the reaction was unabated, but where dIII had been preincubated with H_2NADPH , it was slowed to 7% of the control rate (measured in complexes with bound NADP^+). Consistent with the stopped-flow data, these results suggest that bound H_2NADPH cannot reduce AcPdAD^+ ; the low residual rate after preincubation with H_2NADPH can be attributed to NADP^+ that had not been exchanged from the complex.

There may be two reasons why H_2NADH and H_2NADPH are such poor hydride donors. First, their “enzyme-oxidized” forms (with H^- abstracted from C4_N) would have a 5,6-dihydronicotinamide ring (H_2Nic^+ ; see Scheme 1), which, unlike the nicotinamide ring of NAD(P)^+ , is not stabilized by a delocalized π -electron system. The redox potential of the analogue couples will, therefore, be higher than that of $\text{NAD(P)}^+/\text{NAD(P)H}$ ($E_{\text{m},7.0} \approx -0.32 \text{ V}$), and H_2NADH and H_2NADPH will be weaker hydride donors (from C4_N) than their physiological equivalents. Second, there may be steric factors that prevent proper juxtaposition of the H_2NicH ring and the Nic^+ ring of the partner nucleotide at the hydride-transfer site. Thus, the extra hydrogen atoms at positions 5 and 6, the altered bond lengths of H_2NicH , and the greater tendency of the ring to pucker might interfere with the reaction geometry. The unchanged binding constant, at least for H_2NADH relative to NADH , perhaps argues against this, and an increased redox potential may be the predominant reason for the failure of the analogues to participate in hydride transfer.

The Formation of Dead-End dI_2dIII_1 Complexes Is Disfavored. In previous work, microcalorimetry experiments revealed two different NADH -binding sites on the dI_2dIII_1 complex (52). The K_d of one site ($17 \mu\text{M}$) was similar in value to that of isolated dI and was, therefore, assigned to the dI polypeptide in the complex [designated dI(A)] which,

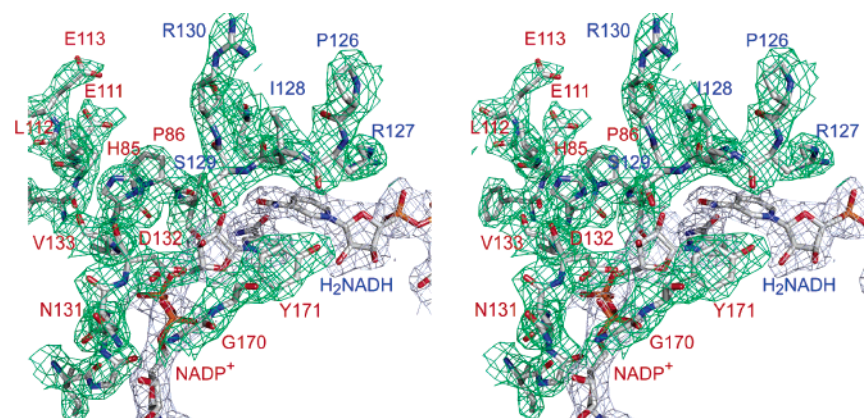


FIGURE 3: Electron density and atom coordinates of H₂NADH and NADP⁺ at the interface between dI(B) and dIII in the dI₂dIII₁ complex (PDB 20O5). The map (resolution 2.6 Å), shown in stereoview, is $2F_o - F_c$ contoured at 1σ . Electron densities of the nucleotides are shown in black, and those of amino acid residues are in green. Residues in dI are labeled in blue, and residues in dIII are in red.

in the crystal structure, makes few contacts with dIII (see Figure 4a). The K_d of the other site was elevated (to approximately 300 μ M) and was assigned to the dI polypeptide, dI(B), which interacts strongly with dIII to form the hydride-transfer pocket. To block hydride transfer, which would otherwise have led to complicated heat changes, and thus to distortion of the K_d , these titrations were performed on dI₂dIII₁ complexes preloaded with NADPH rather than NADP⁺ (apo-dIII from *R. rubrum* transhydrogenase is unstable). However, this left an ambiguity in the interpretation of the experimental result: it was not known whether the elevated K_d for NADH on dI(B) arose because dIII binding itself, or because the presence of the adjacent NADPH in the bound dIII, perturbed the site. Titrations with H₂NADH enable us to discriminate between the two possibilities. First, it was established that H₂NADH binds to NADPH-loaded dI₂dIII₁ complexes with parameters similar to those of NADH; there were indeed two binding sites, one with low (24 μ M) and one with high (176 μ M) K_d values (Figure S5a, Supporting Information). However, in subsequent H₂NADH titrations with NADP⁺-loaded complexes (an experiment which is possible because the H₂NADH is redox-inactive), the results were quite different: the data could be fitted by a single class of binding site, $K_d = 44 \mu$ M with a stoichiometry of 2.3 per dI₂dIII₁ complex (Figure S5b, Supporting Information). This suggests that perturbation by the nucleotide occupant of bound dIII, rather than dIII binding itself, has a more pronounced effect on the affinity of dI(B) for H₂NADH: bound NADPH lowers the binding affinity for H₂NADH (and presumably NADH) in dI(B). It seems that the dI₂dIII₁ complex can discriminate against the formation of dead-end species in which both bound nucleotides are reduced. A similar conclusion was drawn from experiments with the intact *E. coli* transhydrogenase when NADH binding to apoenzyme and to NADPH-loaded enzyme was compared (85).

Crystal Structures of dI₂dIII₁ Complexes. Samples of the dI₂dIII₁ complex loaded with NAD⁺ and H₂NADPH crystallized under conditions similar to those described for the complex loaded with NADH and NADPH. Somewhat different conditions were required for the crystallization of complexes loaded with H₂NADH and NADP⁺ (see Experimental Procedures). X-ray data sets for NAD⁺/H₂NADPH complexes were processed and refined to 2.3 Å (PDB accession number 20OR), and those for H₂NADH/NADP⁺

complexes, to 2.6 Å (PDB 20O5) (see Table 1). A $2F_o - F_c$ electron density map of the adjacent nicotinamide moieties of the bound H₂NADH and NADP⁺ and their surrounding amino acid residues in 20O5 is shown in Figure 3. An $F_o - F_c$ simulated annealing omit map of the region is presented in Figure S6 of the Supporting Information. Both 20OR and 20O5 have an essentially identical fold to that previously reported for the dead-end complexes (32, 33). In each case, the structure comprises two dI polypeptides (A and B) and one dIII. The dIII is more closely associated with dI(B) than with dI(A) (see Figure 4a). The A and the B polypeptides of dI have slightly different conformations to one another but are related by a pseudo-2-fold axis. Each dI has two domains (dI.1 and dI.2) which are separated by a deep cleft. Most of the contact surface between the dI monomers is provided by the two dI.2 domains, but a distinctive β -hairpin extends from the dI.2 of each monomer to make contact with dI.1 of the symmetry-related partner (Figure 4b). The dIII component comprises a single domain and is located at one end of the cleft of dI(B). It makes extensive contacts with dI.2(B), although there is also an important contact region between the RQD loop (residues 126–136) of dI.1(B) and helix D/loop D (residues 131–155) of dIII. The polypeptide and domain nomenclature that we adopt for the dI₂dIII₁ complex is illustrated in Figure 4a.

Intact transhydrogenase has two hydride-transfer sites per dimer. However, dI₂dIII₁ complexes, typified by 20OR and 20O5, have only one, located at the dI(B)–dIII interface (see Figure 4c). The second hydride transfer site of the dI₂-dIII₁ complex is incomplete because dI(A) is not associated with its own dIII (see below).

DISCUSSION

Detail at the Nucleotide-Binding Sites. To understand the mechanism of proton pumping by transhydrogenase, an accurate description of the hydride-transfer site is essential. In a first approach catalytically dead-end dI₂dIII₁ complexes were crystallized; the dI(B) nucleotide in structures with NAD⁺/NADP⁺ (1HZZ and 1U28) was disordered, but in a structure with NADH/NADPH (1U2D) the two NicH rings at the dI(B)–dIII interface were organized in a way that suggested we might have a good model for the hydride-transfer site (32, 33). In a second approach dI₂dIII₁ complexes were crystallized in the presence of NADP⁺ and excess

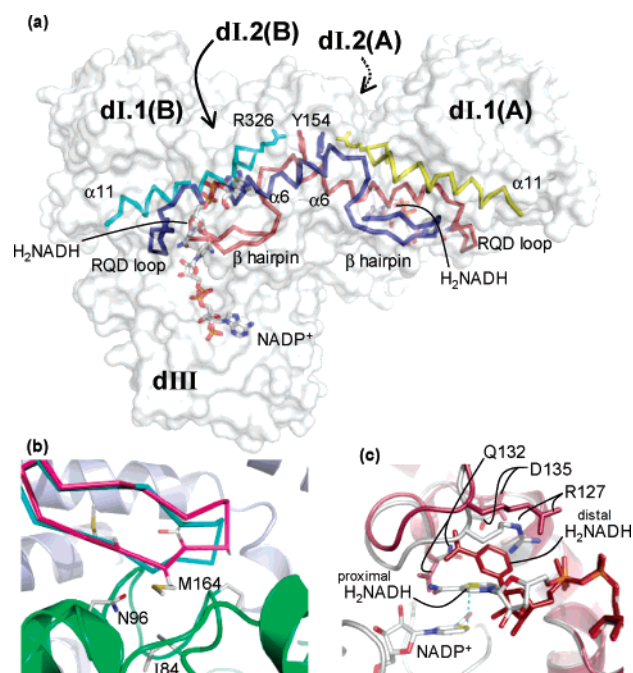


FIGURE 4: X-ray structure of the dI₂dIII₁ complex with bound H₂NADH and NADP⁺ (PDB 20O5). (a) Surface model of the dI₂-dIII₁ complex, also showing the bound nucleotides (in their conventional atom colors) and the C_α traces of the RQD loop, helix α6, the β-hairpin, and helix α11 of the A and B polypeptides of dI. The figure shows the domain and polypeptide nomenclature used in the text. (b) Interaction between dIII and the β-hairpin of dI. The pale blue ribbons represent dI(B), and the green ribbons represent dIII. Shown in red is the C_α trace of the β-hairpin of dI(A) with the side chain of its Met164 making van der Waals contact with those of Ile84 and Asn96 in dIII. The cyan C_α trace shows how the β-hairpin of dI(B) models into this site when dI.2 of the A polypeptide is superposed onto that of the B polypeptide (at residues 202, 308, and 312; see Experimental Procedures); this leads to clashes between the Met164(B) and Ile84 and Asn96 in dIII. (c) The white polypeptide chains, and the side chains and nucleotides in conventional atom colors, show the hydride-transfer site at the interface between dI.1(B) and dIII. Here, the H₂NicH ring of the H₂NADH is proximal to the Nic⁺ ring of the NADP⁺. The dashed cyan line shows the apposition of the two C₄N atoms (colored yellow). The amide group is shown with its O trans to C₂N as discussed in the text. The side chains of Arg127, Gln132, and Asp135 of dI(B) are shown; they participate in an H-bonded network similar but not quite identical to that seen in the equivalent site of 1U2D (33). The brown polypeptide, amino acid side chains, and nucleotide are those of dI(A) modeled by superposing dI.2 of the A and B polypeptides, as described in (b). Here, the H₂NicH ring is distal, and side chain movements have led to disruption of the H-bonded network between Arg127, Gln132, and Asp135.

NADH; hydride transfer during crystallization led to the formation of NADH/NADPH dead-end complexes and to complexes with undefined nucleotide composition (1XLT) (34). The hydride-transfer site at the interface of dI(H)-dIII(I) of 1XLT is somewhat similar to that of dI(B)-dIII in 1U2D. In a third approach we have now crystallized dI₂-dIII₁ complexes with unreactive nucleotide/nucleotide analogue combinations having a charge distribution similar to those in the productive reactions with natural substrates. Thus, the charge distribution on the nucleotides bound in the blocked hydride-transfer site of 20O5 (H₂NADH and NADP⁺) is similar to that in the intermediate which precedes the redox reaction during forward transhydrogenation with natural substrates (NADH and NADP⁺). The structure of the site in 20O5 closely resembles that in 1U2D and somewhat

resembles that in 1XLT. The convergence of the structures derived from the three different approaches strongly suggests that we have arrived at a reliable model for the hydride-transfer site of transhydrogenase. In 20O5, which is at an improved resolution, the H₂NicH ring of the H₂NADH and the Nic⁺ ring of the NADP⁺ are aligned such that the two C₄N atoms are apposed to one another (Figure 4c): the H₂NicH ring of the H₂NADH is held in a proximal position relative to the NADP⁺. This position is maintained by a network of H-bond interactions involving Gln132, Asp135, and Ser138 that hold Arg127 in a deep-lying location in the cleft between dI.1 and dI.2, preventing the H₂NicH from occupying a distal position (33, 86, 87). The site is sealed off from the solvent by the mobile loop (residues 220–240), whose Tyr235 also interacts with Arg127 and the nucleotide pyrophosphate. Note that the average *B*-factors for the bound H₂NADH (27.1 Å²) and NADP⁺ (36.1 Å²), and those for the 32 amino acid residues lying within 5 Å of the nicotinamide mononucleotide moieties of these nucleotides (30.8 Å²), are less than the average *B*-factor of the entire structure (40.7 Å²), thus supporting the view that this is a stable conformation of the hydride-transfer site. If NADH is modeled in place of the H₂NADH in 20O5, then the *pro-R* H on C₄N is positioned to transfer to the C₄N on the *si* face of NADP⁺, that is, in agreement with the “A–B” hydride transfer determined experimentally (2, 3).

The fact that both nucleotides are reduced in 1U2D and 1XLT (chains H and I) and that high concentrations of NADH are required to force occupation of the site in 1U2D (see above, Figure S5, and ref 52) evidently does not lead to large-scale distortion. There are, however, small differences between the hydride-transfer sites of 20O5 and those in the previously reported NADH/NADPH dead-end complexes. The two C₄N atoms are separated by only 3.4 Å in 20O5 compared with 3.6 Å in 1U2D and 4.6 Å in 1XLT (chains H and I), and the alignment of the (H₂)Nic(H) ring planes is slightly different in the three structures: in 20O5 and in 1U2D the two ring planes are almost parallel (especially in 1U2D), but in 1XLT they are somewhat tilted. Furthermore, the H-bond network involving Arg127, Gln132, Asp135, and Ser138 of dI and the (H₂)Nic(H) pyrophosphate group adopts a slightly different pattern in 20O5 and 1U2D and, particularly, in 1XLT. One striking difference is a flipped adenine position in 1XLT (chain H), not seen in any other dI polypeptide either in the isolated protein or in dI₂-dIII₁ complexes. It is not clear to what extent these differences are a consequence of the different resolutions of the structures (2.6 Å in 20O5, 3.0 Å in 1U2D, and 3.1 Å in 1XLT) and to what extent they reflect real structural differences arising from the different states of nucleotide occupation.

The charge distribution on the nucleotides bound in the blocked hydride-transfer site of 20OR (NAD⁺ and H₂-NADPH) is similar to that in the intermediate which precedes the redox reaction during reverse transhydrogenation with natural substrates (i.e., NAD⁺ and NADPH); see eq 1. However, the electron density for the NAD⁺ in dI(B) of 20OR is incomplete: that for the adenosine and pyrophosphate moieties of the nucleotide is clear but that for the Nic⁺ ring is weak. This suggests at least moderate occupancy of the site by NAD⁺, but disorder in the Nic⁺ of the nucleotide that does bind, and contrasts with structures having bound

NADH where the electron density of the dI(B) nucleotide is clear (see above). The structure comparisons show that disorder in the bound NAD⁺ of 1U28 is not simply a consequence of an unfavorable charge interaction between the Nic⁺ rings of NAD⁺ and NADP⁺, as was previously suggested (33), because it is also evident (in 2OOR) where H₂NADPH (with its uncharged H₂NicH ring) is bound to dIII. It must be concluded that the poor density for Nic⁺ in NAD⁺-loaded structures is an intrinsic characteristic of the way in which this oxidized nucleotide binds to the dI component of the complex. Movement of the Nic(H) ring in the hydride-transfer site is thought to be an important feature of the transhydrogenase mechanism (see below). The greater disorder of the Nic⁺ in bound NAD⁺ than that of the NicH in bound NADH may signify a smaller population of nucleotide in the proximal state for hydride transfer during reverse transhydrogenation and thus partly explain the much lower (>36-fold) hydride-transfer rate in this direction (88).

In all published X-ray structures of dI₂dIII₁ complexes including those presented here there is bound nucleotide (NADP⁺, NADPH, and H₂NADPH) in good electron density in the dIII polypeptide. Moreover, their conformations and those of neighboring amino acid residues are similar. The same is true of NADP(H) structures of isolated dIII, which are solved at higher resolution (33, 83, 89). Evidently, the changes in chemical shift that occur when NADP⁺ is exchanged with either NADPH or H₂NADPH (Figure 2 and refs 54, 82, and 90) are either predominantly electrostatic in origin or correspond to atomic displacements that are too small to be detected in X-ray structures at this resolution. The average coordinate error in 2O05 is 0.4 Å as calculated from the Luzzati plot.

Molecular Dynamics Simulation of the Hydride-Transfer Site in dI₂dIII₁ Complexes. In the intact transhydrogenase interactions between the peripheral dI/dIII components and the membrane-spanning dII gate the hydride-transfer reaction and regulate the binding and release of NADP(H) (8–11). Isolated dI₂dIII₁ complexes are locked in a conformation in which the redox reaction can take place at rapid rates (28, 31) and in which NADP(H) in dIII of dI₂dIII₁ complexes is occluded from the solvent by loop E (26). The X-ray structures of this complex are, therefore, a good model in which to study the protein dynamics of the intermediate state that is associated with hydride transfer in the intact enzyme. Figure 5 summarizes the results of an MD simulation of 2O05 along a 1.4 ns trajectory. The average root mean square deviation of the C_α atoms indicates that the protein continues to sample conformations close to the X-ray structure across this time scale (Figure 5a). The H₂NicH ring of the H₂NADH and the Nic⁺ ring of the NADP⁺ remain close to parallel during the trajectory: the tilt between planes through the C_{2N}, C_{4N}, and C_{6N} atoms of the two nucleotides in the time-averaged (0–1.4 ns) structure was 20.3° (compare 6.5° in 2O05).

The distance between the C_{4N} atoms of the H₂NicH and Nic⁺ rings fluctuates between only very narrow limits (Figure 5b). The reason for this is apparent from the data shown in Figure 5c–g. Thus, relative movements of the rings perpendicular to their planes (Figure 5e,g) are small in comparison with relative movements within their planes (Figure 5c,d,f): fluctuations tend to slide the rings past each other but not separate them. Presumably, separation motions

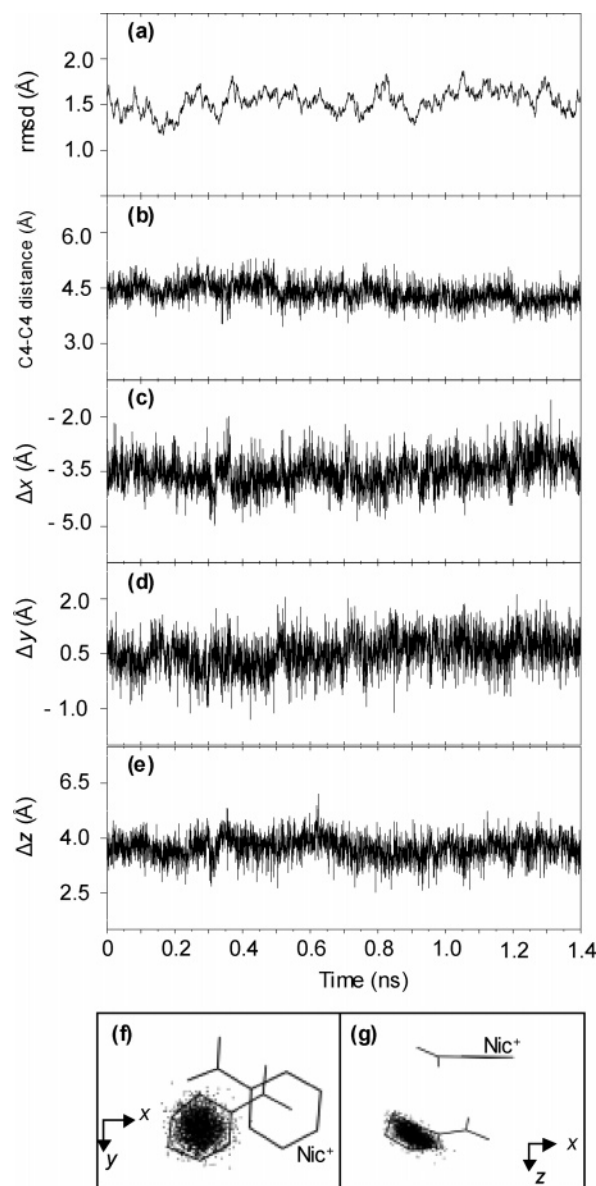


FIGURE 5: Molecular dynamics simulation of the dI₂dIII₁ complex, 2O05. (a) Time dependence of the root mean square deviation (rmsd) of the protein C_α atoms relative to those of the energy-minimized starting structure. (b) Time dependence of the distance between C_{4N} of the Nic⁺ ring of NADP⁺ and C_{4N} of the H₂NicH ring of H₂NADH. (c–e) The C_{2N}, C_{4N}, and C_{6N} atoms of the Nic⁺ ring of the NADP⁺ of each structure along the trajectory were superposed, and an x–y plane was defined by the superposed atoms with an origin at the ring centers. The time dependence of the distance between the centers of the Nic⁺ and H₂NicH rings along (c) the x coordinate, (d) the y coordinate, and (e) the z coordinate of the defined plane was determined from the superposed structures. (f, g) The lines show the projection of the Nic⁺ and H₂NicH rings in the time-averaged structure, and the small dots show the positions of the center of the H₂NicH ring relative to that of the superposed Nic⁺ rings of NADP⁺ during the trajectory. The arrows indicate the directions of the x, y, and z coordinates: z is normal to the page in (f) and y is normal to the page in (g). Note that a parallel trajectory with different starting atomic velocities gave equivalent results to those shown in panels a–e.

(in the z-direction) are restrained by overlap of the π -electron systems of the two rings and their amide groups. As will be seen in the following section, the continued exploration of these apposed configurations of NicH and Nic⁺ rings will readily lead to a reactive orientation for hydride transfer.

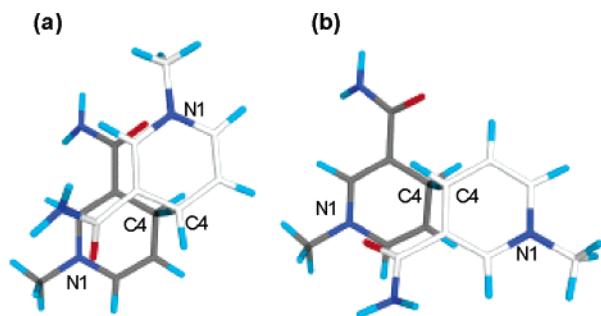


FIGURE 6: Transition-state structures of the hydride-transfer reaction between 1-methyl-1,4-dihydronicotinamide and the 1-methylnicotinamide cation. (a) Transition-state structure **1**. (b) Transition-state structure **7** (see Table S1). The C atoms of the upper ring are in white, and those of the lower ring are in gray. The transferred hydride can be just seen between the two C4 atoms in each structure.

The Transition State for Hydride Transfer between NAD(H) and NADP(H). Two transition-state structures for the C4–C4 hydride-transfer reaction between 1,4-dihydronicotinamide and protonated nicotinamide were identified by Wu et al. (76) using ab initio quantum mechanical methods (HF/6-31G*). Neither of these corresponds very well to the organization of the nicotinamide rings of nucleotides in the active site of transhydrogenase seen in the crystal structures. We have therefore extended the earlier calculations at a higher level of theory (dft/6-31G**), which includes a contribution for electron correlation. Furthermore, we have used 1-methyl analogues of dihydronicotinamide and protonated nicotinamide to provide a better model of the physiological reactants. Wu et al. (76) were working without the benefit of a crystal structure for this system and made the not unreasonable assumption that a symmetrical reaction ($A + B \rightarrow B + A$) would pass through a symmetrical transition state (at least C_2). As the crystal structure shows a C_1 arrangement of nicotinamide rings, we did not follow this constraint. A number of possible transition states have been identified, including structures equivalent to those described (76) and all combinations of amide position and orientation based on the arrangement of dihydropyridine cores observed in the crystal structure. Their geometries and their total and relative energies are summarized in Table S1, and the two lowest energy structures, which differ by only 5.7 kJ mol⁻¹, are illustrated in Figure 6. Of all the calculated transition states, that with the second lowest energy (structure **7**; see Figure 6b) most resembles the organization of the H₂-NicH and Nic⁺ rings of the H₂NADH and NADP⁺ in the X-ray structure. Thus, in **7**, and in 20O5, the planes of the two rings are close to parallel, the angle between the N1–C4 axes of the rings is approximately 120° (viewed from a normal to the ring planes as in Figure 6b), and while the C3–C7 bond of one amide group stacks on top of the C5–C6 bond of its partner ring, the other amide group points away from its respective partner. However, in the transition-state structure the two C4 atoms are in closer contact (2.67 Å), and both C4 atoms show partial sp³ hybridization consistent with hydride transfer. The O atoms of each of the two amide groups in **7** are *trans* to C2 (though rotated slightly out of the ring plane). The transition-state structure **7** is asymmetric (it has C_1 symmetry), and unlike the C_2 -symmetrical structure, **1**, it shows two different C4–H bond distances (1.29 and 1.38 Å compared with 1.34 Å, respectively). Asymmetric transition states for what is, in effect, a

symmetrical reaction are highly unusual and are normally associated with a two-step mechanism that passes through a symmetrical intermediate. That is clearly not the case here as both frequency and IRC calculations confirm that this structure is a first-order transition state for hydride transfer occurring as a single step. Transition-state structures **9** and **10** are similar to **7** but have the O atom of one amide *cis* to its C2; this raises the energy of the transition state by approximately 20 and 6 kJ mol⁻¹, respectively, possibly for the reasons put forward (76). Transition-state structure **8** has both amide O atoms *cis* to their C2s, and its energy is elevated by approximately 20 kJ mol⁻¹. The resolution of the X-ray structure is not enough directly to identify the rotation of the amide group about the C3–C7 bond, but the H-bonding pattern indicates that the O is *trans* to C2 in the amide of NADP⁺; equivalent clues are unfortunately lacking for the H₂NADH, and thus the crystal structure could correspond to either transition state **7** or **10** but not to either **8** or **9**. The possible significance of amide rotation in transhydrogenase turnover was discussed (89).

The similarity between the X-ray structure and a low-energy transition state may indicate that the transhydrogenase active site has evolved to optimize the relative positions of the Nic(H) rings and thus favor a rapid rate of hydride transfer from bound NADH to NADP⁺. For efficient coupling to proton translocation through the enzyme, there is also a need to switch on and switch off hydride transfer by moving the Nic(H) ring of the NAD(H) at appropriate times in the reaction cycle (see refs 33 and 86 and below). It would seem that this requirement has been managed during evolution without significantly compromising the need for an optimized position of the Nic(H) rings that will lead to a rapid redox reaction at the point of hydride transfer.

Coupled Asymmetric Motions in the dI Dimer: The Role of the β -Hairpin in the Asymmetric Binding of dIII. In the dI(A) polypeptides of all published wild-type dI₂dIII₁ structures the electron density of bound nucleotide is very clear, and the NAD(H) is seen to adopt a distinctively different conformation to that of NAD(H) in the dI(B)–dIII hydride-transfer site. The same is true for the NAD⁺ in dI(A) of 20OR and for the H₂NADH in dI(A) of 20O5, giving further support to the notion that the binding of the tetrahydro analogues leads to conformational changes similar to those resulting from the binding of physiological nucleotides. Thus, whatever the redox state, the nicotinamide moiety in all dI(A) polypeptides is pushed further back into the cleft between dI.1 and dI.2 than in dI(B). There are accompanying changes in the side chain conformations and H-bonding pattern of Arg127, Gln132, Asp135, and Ser138 in the RQD loop, and these share similar features in all structures and appear to be responsible for holding the nucleotide position (33, 86, 87). If the NAD(H) conformation seen in dI(A) were adopted in the dI(B)–dIII hydride-transfer site, the C4_N atom of the nucleotide would be too far from that of bound NADP(H) to allow the redox reaction (32, 33). This position of the Nic(H) was therefore described as distal, and a role for the distal–proximal switch in the gating of hydride transfer was suggested.

There are other asymmetric features in the A and B polypeptides of dI. Notably, these include a different degree of rotation of domains dI.1 and dI.2 and a different position of the β -hairpin. An analysis of the available structures as

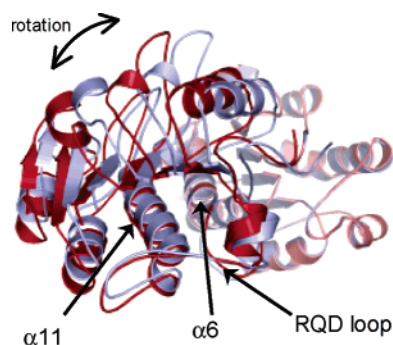


FIGURE 7: Difference in the relative rotation of the dI.1 and dI.2 domains between the A and B polypeptides of the dI₂dIII₁ complex, 2O05. The C α of residues 202, 308, and 312 of the dI.2 domain of the A polypeptide were superposed on the equivalent atoms of the B polypeptide (as described in Figure 4). The A polypeptide is in red, and the B polypeptide is in blue. The view is orthogonal to, and from the left of, that in Figure 4a. Thus, the dI.1 domains are at the front, and the dI.2 domains are superposed at the back.

outlined below and a consideration of recent work on a β -hairpin mutant (24) suggest that the asymmetries reflect structural changes that occur during operation of the enzyme. Specifically, we propose that these changes are involved in controlling the alternating interactions between dI and dIII at the two hydride-transfer sites.

There is a difference in the relative rotation of domains dI.1 and dI.2 in the two monomers of the dimer in crystals of isolated dI (22, 23). The axis of the rotation lies approximately along α helices 6 and 11 which link the two domains (see Figure 4a). There is an equivalent difference in the relative rotation of dI.1 and dI.2 in dI(A) and dI(B) in dI₂dIII₁ complexes (refs 32, 33, and 89 and this work). Importantly, this always has the same sense in dI(B), which binds dIII, relative to that in dI(A), which does not. The relative rotation between dI.1 and dI.2 in A and B of 2O05 is approximately 9° and is illustrated in Figure 7 and on the MORPH server (code 031760-24674) (64). In contrast, there are no detectable relative movements between the dI.2 domains of the A and B polypeptides; these appear to form a central core for the rotations of the peripheral dI.1 domains. It is concluded that the rotations of dI.1 and dI.2 are coupled and reciprocating and that dIII binds more tightly to the dI monomer that has the rotation found in the B polypeptide.

Figure 4b indicates that movements of the β -hairpin, residues 157–175 in dI, may be responsible for this binding selection. The β -hairpin of dI(A) extends around the contact site between dI(B) and dIII and allows a favorable interaction. However, the β -hairpin of dI(B) is in a different position, and this obstructs the docking of a dIII polypeptide onto dI(A). The positional change is small (1–2 Å at the β -turn of the hairpin) but can be seen in structures 1HZZ, 1U2D, 2OOR, and 2O05 of the dI₂dIII₁ complex and in 1F8G and 1L7E of the isolated dI dimer (not shown). It involves a hinged motion about residues Ala157 and Val175. The direction of the β -hairpin movement correlates with the direction of rotation of dI.1 and dI.2 indicating a linkage. Note that each β -hairpin makes extensive contact with the dI.1 of its symmetry-related partner polypeptide. It is suggested, therefore, that the domain rotations cause the change in β -hairpin position alternately to enable and to prevent the binding of dIII.

The role of the β -hairpin has also aroused the interest of other workers. Johansson et al. (24) recently constructed a mutant of *E. coli* dI carrying a 10-residue deletion in this element. The mutant protein folded as a dimer, had an affinity for *E. coli* dIII similar to wild-type dI, and the resulting complex had only a slightly decreased rate of hydride transfer. The similar affinity between dIII and mutant dI was thought to be “surprising” (24) because the β -hairpin contributes to the dI–dIII contact surface (32). In fact, these findings are entirely consistent with the suggestion advanced above: the β -hairpin is not required to promote dIII binding but rather to prevent it at the appropriate step in alternating-site catalysis. In the reported experiments (24) we infer that dIII will bind to either of the polypeptides (A or B) in the mutant dI to give a complex capable of good rates of hydride transfer; in contrast, dIII will bind only to dI(B) of wild-type dI. It is expected that a β -hairpin deletion in the intact enzyme (as distinct from dI–dIII complexes) would disrupt proton-translocating transhydrogenation.

How the reciprocating rotations of dI.1 and dI.2 in the A and B polypeptides are coupled is not yet understood. Nevertheless, the rotations seem to drive positional changes in the RQD loop, which shift the Nic(H) rings of bound NAD(H) between proximal and distal conformations to enable or block hydride transfer, as described (33, 86, 87) and, simultaneously, lead to changes in the β -hairpin position, which either enable or inhibit the binding of dIII. Because the rotations are reciprocating, the events in the A and B monomers are 180° out of phase in accordance with the “alternating-site” model of proton pumping by transhydrogenase (21, 32). It is widely accepted (following ref 8) that proton translocation through dII results in conformational changes, which manifest as NADP(H)-binding changes in dIII (6, 7, 9–11). The conformational changes in dI described above are subordinate to the conformational events in dII and dIII that are driven by proton translocation. The nature of these long-distance conformational changes is a matter of great interest.

The Structure of dI–dIII Complexes versus Crystal Packing. Recently, a previously unreported interaction between *R. rubrum* dI and dIII was described (34). In addition to the interactions discussed (32, 33) (and see above), it was observed that there is another contact surface in the crystal lattice of the dIdIII structure, 1XLT. This surface is between dI and the face of dIII opposite to that housing the NADP(H)-binding site (see Figure 8a). It was suggested that the interaction is not a crystallographic artifact but that it reflects a real contact surface in a hitherto unrecognized state of the intact enzyme; in this state, the NADP(H)-binding site of dIII would be brought into contact with the membrane-spanning dII. During catalysis, the entire dIII component would rotate by 180° between the originally described (32, 33) and the newly described (34) structural states. Were this to be true, it would represent an important new development in our understanding of transhydrogenase. However, in our view there is a distinct possibility that the “new” contact surface does only arise from crystal packing and that it does not reflect interactions either in the solution state of the dI₂–dIII₁ complex or in the intact enzyme.

The first reason to be skeptical about additional contacts between dI and dIII is that in stopped-flow, ultracentrifuge, microcalorimetry, and low-angle X-ray scattering experi-

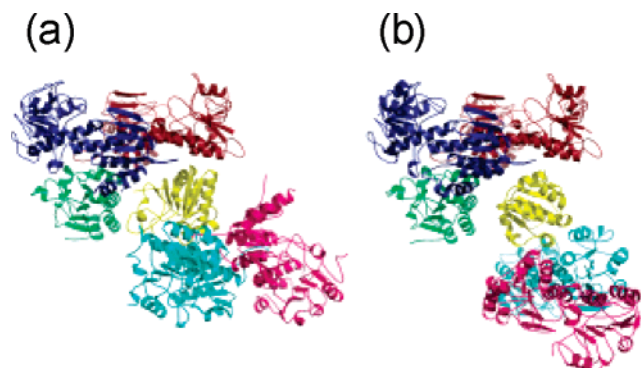


FIGURE 8: Crystal lattice contacts in the dI_2dIII_1 complex of 1XLT and 20O5. (a) The brown polypeptide is dI(G), the blue is dI(H), and the green is dIII(I) of 1XLT (34). This organization is similar to that of the dI_2dIII_1 complex derived from the earlier crystal forms, 1HZZ (32) and 1U2D (33), and is shown from the same perspective as in Figure 4a of this report. Also depicted are polypeptides dI(A) in pink, dI(B) in cyan, and dIII(C) in yellow, which form another dI_2dIII_1 complex in the unit cell (34). The yellow dIII(C) polypeptide is seen to make contacts with the brown dI(G) and the green dIII(I). (b) The brown polypeptide is dI(A), the blue is dI(B), and the green is dIII of 20O5, shown from the same perspective as G, H, and I of 1XLT in panel a. Another dI_2dIII_1 complex from the crystal lattice of 20O5 (related by $x + 1/2, -y - 1/2, -z$ crystallographic symmetry) is shown with dI polypeptides in pink and cyan and dIII in yellow. Again, there are contacts between the yellow dIII and the brown dI/green dIII, but note the different orientation of the yellow dIII polypeptides in (a) and (b).

ments it was established that the complex of dI and dIII, in solution, has a dI_2dIII_1 stoichiometry (30, 52). The K_d for the binding of the dI dimer to the dIII monomer is <60 nM. NMR experiments in the solution state show that a second dIII can bind to the dI_2dIII_1 complex but with a K_d that is >3 orders of magnitude higher than the first (54). Importantly, the distribution of chemical shift perturbations of amino acid side chains indicates that the second dIII binds to dI(A) in a manner that is roughly equivalent to the binding of the first dIII to dI(B); there were no chemical shift perturbations that can be attributed to an “inverted” binding of dIII.

A second reservation concerning the putative inverted contact between dI and dIII (34) is that there is an analogous but, in detail, distinctively quite different interaction between dI and dIII evident in the earlier structures of dI_2dIII_1 (PDB 1HZZ, 1PT9, 1U28, 1U2D) and in 2OOR and 2O05 reported here. In all of these structures there is, in the crystal lattice, a contact surface between dI and the face of dIII opposite to the nucleotide-binding site although, significantly, this is twisted approximately 90° relative to that described in 1XLT; the contact surface in 2O05 is shown in Figure 8b. The contact is very clear in the earlier published data, but because of the biophysical data (see above), we had privately attributed it to a crystal-packing artifact and rejected the idea that it is a significant structural feature. Following the logic of ref 34, it could alternatively be concluded that this contact is also real and that it too reflects an intermediate state during turnover of the intact enzyme. However, we should then arrive at the notion that, during catalysis, dIII undergoes two consecutive rotations on orthogonal axes. This is implausible not least because it is doubtful if the polypeptide linking dII and III is flexible enough to allow both sets of motions. It is also pertinent that transhydrogenases from protozoan parasites have a structure similar to bacterial and mammalian

enzymes, but they are joined, not only by a polypeptide linker between dII and dIII but also by a linker between dIII and dI (91–94). Even one set of rotational movements of the kind described (34) would present a formidable problem of polypeptide topology for these enzymes, which appear to operate by a mechanism similar to that of the bacterial and mammalian transhydrogenases (95, 96).

A final reason to be cautious about the inverted binding of dIII in the X-ray structures is that it is difficult to understand what feature of the enzyme mechanism would be favored by rotating the NADP(H) site, with its bound nucleotide, into the membrane domain. Whether or not there are large subunit rearrangements that take place during transhydrogenase turnover remains an important question, although good evidence for such changes is still lacking.

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SUPPORTING INFORMATION AVAILABLE

Table S1, energies and schematic structures of hydride-transfer transition states; Figure S1, quenching of Trp72 fluorescence of isolated dI by H_2NADH ; Figure S2, heat changes during titrations of isolated dI with H_2NADH ; Figure S3, chemical shift changes of Met239 in the mobile loop of isolated dI induced by H_2NADH ; Figure S4, effect of $H_2NAD(P)H$ on reverse transhydrogenation in chromatophore membranes; Figure S5, heat changes during titrations of dI_2dIII_1 complexes with H_2NADH and $NADH$; Figure S6, electron density omit map of the hydride-transfer site in 2O05. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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