An Asymmetric Post-Hydrolysis State of the ABC Transporter ATPase Dimer

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Abstract

ABC transporters are a superfamily of enzyme pumps that hydrolyse ATP in exchange for translocation of substrates across cellular membranes. Architecturally, ABC transporters are a dimer of transmembrane domains coupled to a dimer of nucleotide binding domains (NBDs): the NBD dimer contains two ATP-binding sites at the intersubunit interface. A current controversy is whether the protomers of the NBD dimer separate during ATP hydrolysis cycling, or remain in constant contact. In order to investigate the ABC ATPase catalytic mechanism, MD simulations using the recent structure of the ADP-Pi-bound MJ0796 isolated NBD dimer were performed. In three independent simulations of the ADP+Pi/apo state, the NBD dimer remained close to the crystal structure, and no opening of either active site occurred. The results thus showed allosteric coupling between the active sites, mediated by intrasubunit conformational changes. Opening of the apo site is exquisitely tuned to the nature of the ligand, and thus to the stage of the reaction cycle, in the opposite site. In addition to this, in also showing how one active site can open, sufficient to bind nucleotide, while the opposite site remains occluded and bound to the hydrolysis products ADP+Pi, the results are consistent with a Constant Contact Model. Conversely, they show how there may be no requirement for the NBD protomers to separate to complete the catalytic cycle.

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Introduction

ABC transporters comprise the major superfamily of primary active membrane translocases. They are found in all kingdoms of life, and are engaged in multifarious cellular processes [1–3]. A subgroup of ABC exporters mediate resistance to structurally diverse chemotherapeutic drugs; a phenomenon known as multidrug resistance (MDR) [4–6]. Other ABC transporters are involved in serious and prevalent genetic diseases in humans, such as cystic fibrosis [7,8]. In bacteria, ABC importers are essential for cell survival through their role in nutrient uptake and osmoregulation, while in pathogenic microbes and parasites, ABC exporters are involved in virulence [9]. An understanding of the molecular mechanism of ABC transporters is critical to the development of treatments for a wide range of pathological conditions affecting not only human health, but also productivity in agriculture.

A conserved core architecture of ABC transporters consists of two transmembrane domains (TMDs) and two nucleotide binding domains (NBDs). The two transmembrane domains together form a translocation channel and substrate binding sites, while the NBDs, which are located in the cytosol, bind and hydrolyse ATP [2,10,11]. In contrast to the TMDs, the NBDs are highly conserved, and contain a number of characteristic sequence motifs, including the Walker A and B consensus motifs for ATP binding, the LSGGQ signature sequence or C-motif, which is diagnostic of the ABC superfamily, as well as the D-, Q-, and H-loops, which are named according to conserved residues [12–14].

The NBD has a bilobal structure, consisting of a nucleotide binding core subdomain, and a helical subdomain, which includes the ABC signature sequence C-motif and is unique to the ABC ATPase fold. A feature of emerging importance to the functioning of the NBD is the flexible nature of the connection between the core and helical subdomains. This was inferred from comparative crystallographic analysis [15] and also observed in MD simulations [12,16–18]. Elastic network analysis of the ABC NBD monomer and dimer also revealed that relative motions between the core and helical subdomains comprise the principal dynamic modes of the subunit. This indicates such motions are intrinsic and fundamental to the contact topology of the subunit, consistent a functional role [17,18]. More recently, experiments using the maltose permease provided biophysical evidence that significant conformational changes between the helical and core subdomains occur during the catalytic cycle [19].

The NBDs interact to form two composite active sites, in which the C-motif of one monomer engages nucleotide bound to the Walker A and B sequences of the opposite monomer [20–22]. Symmetrical sandwich dimers, in which two nucleotides are bound
at the NBD intermonomer interface, have been observed in numerous crystal structures, both of isolated NBDs and whole transporters [Figure 1A; 23–28]. The prevalent view is that the NBD sandwich dimer represents a physiological state in the ABC catalytic cycle, in which the NBDs are proposed to undergo cycles of ATP binding and dimer formation, followed by sequential hydrolysis of both ATPs, and then dimer dissociation, a scheme known as the Processive Clamp or Switch Model [Figure 1B; 13,27,29]. Nevertheless, alternative schemes have been proposed in which the NBDs remain in contact in an asymmetric state, with only one nucleotide tightly bound at any time, referred to as Constant Contact Models [Figure 1C; 2,12, 30–36].

A current question thus involves the nature of the catalytic cycle and the role of the relative movements between the core and helical subdomains. A number of studies of ABC exporters have used agents such as vanadate, beryllium fluoride and aluminium fluoride, which combine with ADP in the active site to form a static analogue of the transition state of the hydrolysis reaction. This arrests the transporter in a state in which one active site is occluded while the other is accessible [37]. MD simulations of the ATP/apo state using the structure of the bacterial ABC exporter Sav1866, which is a homologue of well characterised mammalian MDR ABCs, revealed how an asymmetric closed/open state occurred by way of intrasubunit rotations between the helical and core subdomains [18,38]. In respect of this, we have proposed a Constant Contact Model in which a closed/open asymmetry between the active sites is mediated by relative rotations between the core and helical subdomains [30,36].

Recently, a crystal structure of the MJ0796 (E171Q) ABC isolated NBD dimer was published [39]. This dimer was very close in structure to that of the previously reported MJ0796 ATP sandwich dimer [22], but contained the hydrolysis products ADP and phosphate (Pi) in each active site; and is the only published structure of this state for an ABC transporter NBD dimer (Figure 1A). In our Constant Contact Model, the conformation of the ADP+Pi/apo state of the NBD dimer is predicted to be asymmetric, with one site occluded and containing ADP+Pi, and the other open by way of rotation of the empty (apo) core subdomain, primed to bind ATP. In order to investigate the ADP+Pi/apo state, here we performed MD simulations of the isolated NBD dimer using the recent ADP+Pi-bound MJ0796 dimer structure.

Figure 1. The ABC NBD Dimer. (A) NBD dimer structure. Structural figure prepared from PDB entry 3TIF, used as the starting structure in the simulations. Cartoon representation with core subdomains coloured “wheat” and helical subdomains “slate”; separate monomers are situated top and bottom. ADP+Pi molecules are bound at the intermonomer interface, and shown in stick representation, with carbon yellow, oxygen red, nitrogen blue, and phosphorous orange. Catalytic metal is shown as a purple sphere. (B) ABC Switch Model. Schematic representation of the Switch Model for the catalytic cycle of ABC transporters. Separate NBD monomers are situated top and bottom in each frame, with the larger core subdomains coloured blue and the smaller helical subdomains pink. (1) Resting state. NBDs are separated with helical domains outwardly rotated. (2) ATP binds to each NBD. (3) ATP binding induces inward rotation of helical subdomains. (4) NBDs move together forming a sandwich dimer with two occluded ATPs. (5–6) ATPs are hydrolysed processively in the sandwich dimer, resulting in movement apart of NBDs, product release and return to resting state (1). (1) Constant Contact Model. Schematic representation of the Constant Contact Model for the catalytic cycle of ABC transporters. Separate NBD monomers are situated top and bottom in the frame, with the larger core subdomains coloured green and the smaller helical subdomains yellow. (1) ATP is occluded in left site. Opposite site is empty and open by way of outward rotation of its core subdomain, and has low affinity for nucleotide. (2) ATP hydrolysis induces switch to high affinity for ATP in opposite open core subdomain. (3) ATP binds to empty high affinity site. (4) ATP binding to the empty open core subdomain induces its inward rotation and occlusion of ATP. Occlusion of ATP induces opening of opposite site and release of hydrolysis products. (5–6) are repeated steps in the opposite phase with respect to the two active sites to complete the cycle.

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Methods

System Setup

The starting coordinates for MD simulations were taken from the X-ray structure of MJ0796 (E171Q; PDB 3TIF; [39]). The E171Q mutation was reversed. Unresolved C-terminal residues 233–235 were modelled using Modeller 8v1 [40]. NBD protomers are referred to as A and B, according to the chain identifiers in the Protein Data Bank entry 3TIF. Two systems were prepared: ADP+Pi/apo and ATP/apo, with the nucleotide bound to the Walker A motif in protomer A. The active site sodium ion was replaced by a magnesium ion. Rotamer A for active site residue Q90 was used. In order to generate starting coordinates for the ATP γ-phosphate moiety, as well as for a canonical water molecule which mediates the interaction of the Walker aspartate and the catalytic magnesium ion, the Cx coordinates of the P-loop of ATP-bound MJ0796 [PDB 1L2T; residues 32–49; [22] were root mean square (RMS) fitted to the equivalent atoms in monomer A in the simulation starting structure. All crystallographic water molecules located within 10 Å of the protein, nucleotide, or Pi group were retained and used in the starting structure.

Each complex was solvated in explicit water using TIP3 water molecules. A truncated octahedral periodic cell was used with a minimum of 20 Å between periodic images of the protein, and the system neutralised with 0.2 M NaCl counterions. Histidine 141 was neutral and protonated at the ε nitrogen and active site histidine 204 was ionized. All other ionisable residues were in the default ionization state, with the exception of E171 in protomer A in the ADP+Pi/apo system, which was neutral.

In order to simulate the ADP+Pi-bound active site, the protonation state of the phosphate ion must be set. In solution at pH 7, the ratio of single to double protonated phosphate is about 2:3; however the pKa of the ion is likely to be altered by the electrostatic environment in the active site. Whilst a doubly protonated Pi would account for the stoichiometry of the hydrolysis reaction, the exact mechanism by which stoichiometry is maintained with respect to the protons derived from the nucleophilic water is unknown. Default parameters for ATP in MD forcefields, and MD studies of ABCs generally, assume that the γ-phosphate of ATP is unprotonated. In this case, upon nucleophilic attack by an activated water molecule, initially a singly protonated phosphate ion would be formed. Moreover, it is generally expected that in ABC ATPases, the conserved glutamic acid residue, immediately C-terminal to the Walker B aspartate, acts as a general base, releasing a proton from the attacking nucleophile [38].

In view of the above, in the ADP+Pi/apo system, the phosphate was modeled as singly protonated, and the proximal conserved glutamic acid residue as neutral, being protonated at the sidechain carboxyl oxygen proximal to the Pi hydroxyl oxygen. The hydroxyl oxygen of Pi was chosen to be that which, in the crystal structure, is within hydrogen bonding distance of the backbone carboxyl oxygen of D-loop residue A175 from the opposite monomer (Figure S1), thus being consistent with their proximity (2.7 Å). This geometry is also consistent with a canonical hydrolysis scenario [22,38].

Simulation Parameters

MD simulations were performed using NAMD version 2.9 [41] with the CHARMM27 force field [42], including φ/ψ cross-term map corrections, and the TIP3P model for water [43]. Partial charges for hydrogen phosphate were from [44]. Unprotonated phosphate oxygens were assigned the CHARMM27 ON3 type, and the hydroxyl oxygen ON4; parameters for the ON3-P-ON3 angle were k = 104.0, θ = 108.23. The SHAKE and SETTLE algorithms were used to constrain bonds containing hydrogens to equilibrium length [45]. A cutoff of 11 Å (switching function starting at 9.5 Å) for van der Waals and real space electrostatic interactions was used. The particle-mesh Ewald method was used to compute long-range electrostatic forces with a grid density of approximately 1/A˚3 [46]. An integration time step of 1.5 fs was used with a multiple time stepping algorithm; interactions involving covalent bonds and short-range nonbonded interactions were computed every time step, while long-range electrostatic forces were computed every two time steps. A Langevin piston was used to control pressure with a target of 1 atm, a decay period of 100 fs and a damping timescale of 50 fs. Langevin dynamics was utilized to maintain a constant temperature of 343 K with a friction coefficient of 5 ps⁻¹ on all non-hydrogen atoms. The natural habitat of M. januscheii varies between 48–94 °C with an optimal growth temperature of 85 °C In a previous MD simulation analysis of MJ0796 [17], we used a temperature of 85°C. Here, a lower temperature of 70°C was used, which was considered to strike a better balance between ensuring conditions that would enable significant conformational changes to occur on the simulated timescale, and the optimal temperature of the MD forcefield.

Equilibration

The solvated starting structure was minimized using conjugate gradient minimization to a 0.5 kcal/mol Å RMS gradient during which all non hydrogen atoms of the protein and nucleotide were fixed. Water molecules, NaCl ions, and hydrogens were then further minimized during a 50 ps MD run at 343 K, in which all protein and Mg²⁺ATP/ADP+Pi heavy atoms were again fixed. This starting model was then minimized with harmonic positional constraints on the NCSCO backbone. A 100 kcal/mol Å² force constant was used to minimise the system to a 0.5 kcal/mol Å RMS gradient. The constraints were gradually removed by subsequent minimizations to a 0.1 kcal/mol Å RMS gradient, scaling the initial force constants by factors of 0.5, 0.15, 0.05, and 0. The minimized structure was then heated from 43 K to 343 K in steps of 25 K using velocity reassignment during a 30 ps MD run. Three separate minimizations were performed for each system, using different random seeds for the initial velocity assignments.

Production Runs

The six equilibrated systems described above were used for the production runs without constraints. For each of the two states, 3 simulations were performed, one for 150 M integration steps (225 ns) and two for 100 M steps (150 ns). All simulations remained stable to completion. For analysis, the coordinates were saved every 5 K steps (7.5 ps).

Analysis

Principal component analysis of the Cx atom trajectories was performed using the GROMACS package, with frames taken at 0.075 ns intervals over the entire trajectory, and aligned to the starting coordinates [47]. The program Hingefinder [48] was used to analyse domain rotations: this analysis used Cx coordinates, the slow partitioning algorithm and a tolerance of 40%. VMD [49], Xplor-NIH [50] and Simulaid [51] were used to prepare the system and analyse MD trajectories. Calculation of the RMSD of individual core subdomains used Cx atom coordinates of residues 1–90 and 164–228; this excludes a highly mobile unstructured C-terminal tail of 7 residues (residues 229–235). Calculation of the RMSD of individual helical subdomains used Cx atom coordi-
nates of residues 94–111 and 117–161; this excludes the α3–4 loop (residues 112–116) joining the first and second α-helices in the helical subdomain (see Results). These calculations also exclude the Q-loop (91–93) and Pro-loop (161–163), which are not assigned to either subdomain. All structural figures were prepared using PyMol (http://www.pymol.org/pymol/), except for Figure S2, which was prepared using VMD.

Results

To investigate the catalytic cycle of the ABC transporter NBDs, MD simulations of the ADP+Pi/apo state of the MJ0796 dimer were performed, using the recent ADP+Pi-bound MJ0796 dimer structure. Simulations of the ATP/apo state were also performed, under the same conditions. Three unrestrained simulations of each state were performed, one for 150 M integration steps (225 ns) and two for 100 M steps (150 ns), totaling six independent simulations, each originating from independently equilibrated starting structures. The simulations of the ADP+Pi/apo state are referred to as ADPPi-1, ADPPi-2 and ADPPi-3 for the 225 ns and two 150 ns runs respectively, and the simulations of the ATP/apo state are referred to as ATP-1, ATP-2 and ATP-3 for the 225 ns and two 150 ns runs, respectively.

To assess the stability of the protein, Figure 2 shows the time series of the RMS deviation (RMSD) of Cα atom coordinates from the initial structure, for all simulations. In Figure 2A the RMSD for run ADPPi-1 is plotted; this indicates that the dimer undergoes a relatively large change with respect to the crystal structure. Also plotted in Figure 2A is the RMSD for each NBD monomer separately; this shows that the monomers remain relatively close to the starting structure. Thus, together the data in Figure 2A indicate that the large RMSD observed in the dimer is due to relative movements between the monomers. Figure 2B shows that the large conformational change that occurred in run ADPPi-1, did not occur in any of the simulations of the ATP/apo state, which remained stable and relatively close to the starting structure. Figure 2C shows, in contrast, that in the two separate 150 ns runs of the ADP+Pi/apo state, relatively large changes in RMSD occurred, as in run ADPPi-1.

In order to investigate the stability of the protein more closely, Figure 2D shows the time series of the RMSD for the individual core and helical subdomains in each NBD for run ADPPi-1. The calculation for the helical subdomains excludes the five residue loop joining the first and second α-helices in the helical subdomain (α3–4 loop, residues 112–116), which varies in structure between the monomers in the crystal structure, and was observed to be highly mobile in MD simulations [17,52]. This analysis shows that these subdomains remain very close to the starting conformation; these data, and in particular the very low RMSD for the monomer A helical subdomain, indicate clearly that the high temperature of the simulation (70°C) did not induce significant instability in the protein.

Visual examination of the simulation trajectories revealed that in the three simulations of the ADP+Pi/apo state, the empty active site opened, while the ADP+Pi-bound site remained closed. Opening of either active site did not occur in any of the three ATP/apo runs. This is illustrated in Figure 3, where the time series of the distances between opposing helical and core subdomains are plotted for each simulation.

To further define the conformational changes that occurred in the simulations of the ADP+Pi/apo state, Principal Component Analysis (PCA) of the Cα atom coordinate trajectories was performed. PCA distinguishes concerted global motions from random thermal fluctuations, through analysis of pairwise correlations between atoms, over a simulation trajectory. The

![Figure 2. Stability and conformational changes in the simulations.](http://www.plosone.org/doi/fig/10.1371/journal.pone.0059854.g002)
Figure 3. Opening of the empty active site in the ADP+Pi/apo dimer. Changes in the active sites monitored by the distance between the geometric centres of the Cα atoms of Walker A residues 43–46 (GKST) and C-motif residues 147–150 (SGGQ) of opposing monomers. (A-C) show ADP Pi runs 1–3 respectively, (D-F) show ATP runs 1–3 respectively. Nucleotide-bound active site is red and empty active site black. Sampled at 75 ps intervals. doi:10.1371/journal.pone.0059854.g003

Figure 4. Global change in the ADP+Pi/apo simulation. Projection of the Cα atom trajectory for run ADPPi-1 onto PCA mode 1. Sampled at 75 ps intervals. doi:10.1371/journal.pone.0059854.g004
derived Principal Components (PCs) are ranked according to the amplitude of the motions they describe.

For run ADPPi-1, the first PCA mode (PC1) represented 60% of all Cα atom fluctuations, while PC2 and PC3 represented 11.4% and 5.5% respectively; PC1 thus describes a clearly dominant motion. Figure 4 shows the time series of the projection of the Cα atom trajectory on to PC1 derived from run ADPPi-1. The projection illustrates the extent to which the protein has moved along the transition described by the mode; Figure 4 thus depicts a transition of the protein from one conformation toward another.

For run ADPPi-1, the time series of the projection of PC1 correlated with the time series of the distance between the opposing helical and core subdomains comprising the empty active site (Figure 3), with a correlation coefficient of 0.991; this indicates that PC1 embodies the global motion by which the empty active site opens.

To characterise PC1 from run ADPPi-1 in terms of domain motions, the program Hingefinder was used to identify rotating domains between the maximum and minimum projection structures of PC1; these structures represent the two extremes between which the transition described by the mode occurs. Figure 5 shows the trajectory frame corresponding to the maximum projection structure for PC1, which occurs at t = 167 ns. This illustrates the open apo site, as well the rotating

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**Figure 5. Opening of the empty active site via relative movements of core and helical subdomains.** Stereo view of the trajectory frame corresponding to the maximum projection of PC1 in run ADPPi-1, showing rotating domains and their hinge axes as defined by the Hingefinder analysis of the maximum and minimum projection structures of PC1. Axes are longer red, blue and green lines, with the two shorter same-coloured lines connecting the centre of mass of the rotating domain to the pivot point on the axis. Cartoon image of the protein with monomer A bottom and monomer B top. Rotating core subdomain of monomer B (residues 1–88, 165–174, 180–234) is coloured cyan and its hinge axis is in red. Rotating helical subdomain of monomer B (residues 90–164) is coloured magenta and its hinge axis is in green. Rotating helical subdomain of monomer A (residues 87–163) is coloured orange and its hinge axis is in blue. Rotating domain residues are as determined by Hingefinder.

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**Figure 6. Extent to which the predominant global changes in run ADP+Pi-1 occurs in the other simulations.** Projection of PCA mode 1 derived from run ADP+Pi-1 onto the Cα atom trajectory for each run. ADPPi-1 green, ADPPi-2 yellow, ADPPi-3 blue, ATP-1 red, ATP-2 mauve, ATP-3 grey. The sign for the values for run ADPPi-1 are reversed with respect to Figure 4, to facilitate the comparison. Sampled at 75 ps intervals.

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domains in PC1, as detected and defined by the analysis, and their rotation axes and pivot points, as well as the distance moved by the geometric centre of the respective rotating domain.

This analysis reveals that the largest change is undergone by the core subdomain of monomer B, which rotates outward and away from the opposite helical subdomain, thereby opening the unliganded active site (Figure 5; Figure S2). In addition, a rotation of the monomer A helical subdomain contributes to opening of the empty active site. The helical subdomain that engages the nucleotide also pivots, changing its orientation with respect to the opposite P-loop. Nevertheless, interactions of the nucleotide with the conserved serine and glutamine residues of the monomer B LSGGQ C-motif, present in the crystal structure, are essentially maintained in the simulation (Figure S3).

To examine the other simulations with respect to the dominant motion observed in run ADPPi-1, the Cz atom coordinate trajectory from each simulation was projected onto PC1 from run ADPPi-1 (Figure 6). This shows that, while the two 150 ns runs of the ADP+Pi/apo dimer sampled the configurational space defined by this mode to a significant extent, comparatively, the ATP/apo runs clearly did not. The degree to which the other simulations sampled the configurational space defined by PC1 from run ADPPi-1 was also examined by calculating the subspace overlap coefficients between this mode and PCs 1–2 calculated from the Cz atom coordinate trajectory of each simulation: these were ADPPi-2, 0.819; ADPPi-3, 0.804; ATP-1, 0.296; ATP-2, 0.011; ATP-3, 0.381. This further indicates that the configurational space sampled in runs ADPPi-2 and ADPPi-3 clearly overlapped with that of run ADPPi-1, while simulations of the ATP-bound dimer did so to a significantly lesser degree.

Notably, for run ADPPi-3, PCs 1 and 2 corresponded approximately to PCs 2 and 1, respectively, from run ADPPi-1. This indicates that for run ADPPi-3, the mode described by ADPPi-1 PC1 was not the largest amplitude motion that occurred. This, however, is consistent with the observation that opening of the active site in ADPPi-3 did not occur to as great an extent as in the other two ADPPi runs. Finally, the correlation coefficients between the distance the active site opened, and the projection onto ADPPi-1 PC1 were 0.995 for run ADPPi-2 and 0.818 for run ADPPi-3, further illustrating that motions described by PC1 from run ADPPi-1 were well correlated with active site opening in runs ADPPi-2 and ADPPi-3.

**Discussion**

In this study, to further understand and characterise the functioning of the ABC ATPase, unrestrained MD simulations of the ADP+Pi/apo and ATP/apo states of the isolated NBD dimer of the bacterial ABC transporter MJ0796 were performed. In three independent simulations of the ADP+Pi/apo state, comprising a total of >0.5 µs of simulated time, the empty active site opened, clearly sufficient to allow binding of nucleotide to the unliganded Walker A motif. This opening was achieved substantially by way of rotation of the empty core subdomain away from the opposite monomer. Rotation of the helical subdomain engaging the empty active site also contributed toward its opening. ADP+Pi remained bound in the active site, and the LSGGQ motif maintained contact with the nucleotide, throughout all simulations of the ADP+Pi/apo state. In contrast, in three independent simulations of the ATP/apo state, also comprising a total of >0.5 µs of simulated time, the core and helical subdomains forming the empty active site did not alter in conformation from the sandwich dimer starting structure, and no opening of either active site was observed.

The fact that the empty active site did not open in the simulations of the ATP/apo state suggests that active site opening in the ADP+Pi/apo state was not a random fluctuation. This is also supported by the reproducibility of the opening as shown by the PCA data (Figure 6). Crystallographic and MD simulation analyses of the ABC NBD monomer suggested that conformational changes involving the helical and core subdomains, between the ATP-bound, and ADP-bound or empty NBDs, are determined simply by the nature of the nucleotide, or its absence, in the Walker A and B motifs [15,17,53]. However, here we find that removal of nucleotide from one monomer did not result in conformational changes in the ATP/apo state. This indicates that the outward rotation of the core subdomain observed in the ADP+Pi/apo state was contingent upon hydrolysis products being in the active site in which the unliganded monomer C-motif engages. Thus the apo NBD monomer also utilized the C-motif to sense and respond dynamically to events in the opposite core subdomain.

The opening of the empty active site in the ADP+Pi/apo state is consistent with our Constant Contact Model, with the conformation observed here corresponding well to stages 2 and 5 in Figure 1C. It is also consistent with experimental data showing that in the vanadate trapped state, which is expected to be a transition state analogue close to the ADP+Pi state, one site is occluded while the other is accessible [37]. However, the fact that the empty active site did not open in the ATP/apo state does not appear consistent with our model, which predicts the empty site to be open (stages 1 and 4 in Figure 1C). In regard to the ATP/apo result, it should be noted that the Constant Contact Model is based simply on two premises [30]: an alternating function between the active sites, and a sequence for each active site as follows: empty (open, high affinity), ATP-bound (open), ATP-bound (closed), ADP+Pi (closed), ADP+Pi (open), empty (open, low affinity). The important feature of the ATP/empty state, predicted on this basis, is that the empty active site has low affinity. Thus, it may be that the active site opens to release ADP+Pi, and then closes to produce a low affinity site. However, in our MD simulations of the ATP/apo state of the Sav1066 cytosolic domains, the empty active site opened, notably, however, to a lesser extent than observed here for the ADP+Pi simulations [16,38].

How might the data from our MD studies be reconciled with the Constant Contact Model? Studies of ABCB1, nucleotide-trapped using different transition state analogues, suggested that the degree of opening of the opposite, non-occluded active site is a function of the point along the reaction coordinate that is mimicked by the particular transition state analogue [37]. Moreover, it can be inferred that the ABC transporter undergoes a significant global transition that pivots about the hydrolysis reaction transition state [2]. In our simulations of the Sav1066 ATP/apo state, a progression toward a putative hydrolysic-competent state was observed, whereas in the present simulations, this did not occur. Thus, our MD data are consistent with the notion that in the ATP/apo state, the opening of the empty active site is dependent on the position of the ATP-bound site along the hydrolysis reaction continuum. Furthermore, since the Sav1066 system included the TMD intracytosolic domains, together our MD data are consistent with the idea that the TMDs propagate signals which stimulate hydrolysis, and that in their absence, the empty active site remains closed, corresponding to the ground or reactant state of the opposite ATP-bound site. This is supported by the generally low ATP hydrolysis rates exhibited by isolated ABC NBDs compared to full transporters.
On the basis of the present and previous data, we thus propose a modification to our model such that, following ADP+Pi release, the empty active site is closed in the ATP/apo state, producing a low affinity apo state. From this ATP/apo state, the empty site opens along a continuum that is a function of the point the ATP-bound state is along the hydrolysis reaction coordinate, becoming fully open in the ADP+Pi/apo state.

It might also be argued that the present results support the Switch Model. Thus, the stability of the ATP/apo simulation opens along a continuum that is a function of the point the ATP-bound active site while the post-hydrolytic state destabilizes it. It is important to note, however, that the ATP/apo and ADP+Pi/apo states per se are not part of the Switch Model, so the interpretation of the results in terms of the Switch Model is not clear. Nevertheless, some relevant questions do arise. Thus, how might the NBD dimer’s apparent ability to exchange nucleotide at one active site, before the NBD protomers dissociate, be reconciled with the Switch Model?

The salient feature of the results of this study is the ability of the NBD dimer to respond in a dramatically different way to the presence of either ATP, or the hydrolysis products ADP+Pi, within the active site. This shows that the protein is able to detect and respond to relatively small changes in the nucleotide occupancy of the active site. This is in accord with the notion that ATP hydrolysis induces conformational changes that, in turn, are transmitted to the TMDs, resulting in substrate translocation.

The present findings also suggest that if the conformational differences between the two states observed here indeed embody functional changes, then the free energy released upon ATP hydrolysis may not be transmitted directly to the protein to produce a high energy state, that then relaxes via a functional transition to perform the mechanical work of substrate translocation. Rather, it suggests that the equilibrated, or lowest energy conformation of the protein is different for any particular state of the transport cycle, with conformational changes between states corresponding to those that implement the translocation process. In this view, conformational changes between states occur through thermal fluctuations and are reversible; and the vectorial nature of the transport cycle is determined simply by the energy gradient for ATP hydrolysis.

Supporting Information

Figure S1 Structure of the ADP+Pi-bound active site in the MJ0796 NBD Dimer. Structural figure prepared from PDB entry 3TIF, used as the starting structure in the simulations. Orientation of the monomers is approximately as in Figure 1A; as are the stick forms and colours of the nucleotide and sidechains. Backbone atoms of the C-motif (left) and D-loop (right) of the trans monomer are shown in stick form, with carbon green, oxygen red, and nitrogen blue. A hydrogen bond between the backbone carbonyl oxygen of D-loop residue A175 and the Pi molecule is shown as a red dashed line. Weaker interactions between the conserved glutamine (left) and the E171Q glutamine (centre) are shown as green dashed lines. (PDF)

Figure S2 PGA mode 1 from simulation ADPPi-1. (A) to illustrate the mode, the maximum and minimum projections of PC1 along the trajectory, together with 8 interpolated structures, are superimposed by rms fitting using all Cα atom coordinates. Cα atom coordinates only are shown, with core subdomains blue and helical subdomains green. Red arrows indicate P-loops and black arrows the LSGQQ motif. (B) and (C) stereo views derived from (A) using the trajectory smoothing function in VMD to reduce range of motion and illustrate relative degrees of motion. Arrows as in (A). Colours indicate progression in time with blue near t = 0 and red near t = 150 ns. (TIF)

Figure S3 Interaction of the C-motif with the nucleotide in the ADP+Pi/apo simulations. (A–C) Time series of the distance between the C-motif serine hydroxyl oxygen and the proximal nucleotide β-phosphate oxygen for ADPPi runs 1–3 respectively. (D–F) Time series of the minimum distance between either the oxygen or nitrogen of the C-motif glutamine (LSGQQ) sidechain amino, and the proximal nucleotide ribose hydroxyl oxygen, for ADPPi runs 1–3 respectively. (PDF)

Author Contributions

Conceived and designed the experiments: PMJ. Performed the experiments: PMJ. Analyzed the data: PMJ. Contributed reagents/materials/analysis tools: PMJ. AMG. Wrote the paper: PMJ. AMG.

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