

Noncanonical role for the binding protein in substrate uptake by the MetNI methionine ATP Binding Cassette (ABC) transporter

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Contributed by Douglas C. Rees, August 26, 2018 (sent for review June 26, 2018; reviewed by Bert Poolman and Nieng Yan)

The Escherichia coli methionine ABC transporter MetNI exhibits both high-affinity transport toward L-methionine and broad specificity toward methionine derivatives, including p-methionine. In this work, we characterize the transport of D-methionine derivatives by the MetNI transporter. Unexpectedly, the N229A substrate-binding deficient variant of the cognate binding protein MetQ was found to support high MetNI transport activity toward p-selenomethionine. We determined the crystal structure at 2.95 Å resolution of the ATP_γS-bound MetNIQ complex in the outwardfacing conformation with the N229A apo MetQ variant. This structure revealed conformational changes in MetQ providing substrate access through the binding protein to the transmembrane translocation pathway. MetQ likely mediates uptake of methionine derivatives through two mechanisms: in the methionine-bound form delivering substrate from the periplasm to the transporter (the canonical mechanism) and in the apo form by facilitating ligand binding when complexed to the transporter (the noncanonical mechanism). This dual role for substrate-binding proteins is proposed to provide a kinetic strategy for ABC transporters to transport both high- and low-affinity substrates present in a physiological concentration range.

ATP Binding Cassette transporter | alternating access transport mechanism | methionine transporter | transinhibition

he substrate specificity of a transporter represents one of the most basic functional attributes and often serves as the defining characteristic. Transporters can exhibit both high affinity for a particular substrate and broad specificity in translocating related compounds. An example is the Escherichia coli MetNI methionine importer established by Kadner et al. to mediate the high-affinity uptake of L-methionine, while transporting D-methionine with lower affinity; other methionine derivatives including selenomethionine are also transported (1-4). MetNI is a member of the ATP Binding Cassette (ABC) family of transporters (5–9), consisting of two transmembrane subunits (MetI) that form a translocation pathway and two nucleotide-binding subunits (MetN) that couple ATP binding and hydrolysis to transport (10, 11). As for other ABC importers (5, 12-15), MetNI requires an additional component, MetQ (8, 10, 11, 16-18), which is a substrate-binding protein (SBP), generally viewed to deliver substrates to their cognate transporters.

The role of SBPs has been intriguing since their original identification as osmotic sensitive components that were required for the transport of certain nutrients (19). Investigations of a number of SBPs support a model where binding of substrate by free apo-SBP is accompanied by a "Venus flytrap" conformational change (20, 21), followed by delivery of ligand to the ABC transporter (22–25); in this canonical model, the SBP delivers the substrate to the transporter and is predominantly responsible for the substrate specificity of transport. However, these sequential events have been challenged by biochemical and single-molecule studies (26–30), including an early demonstration by Shuman that binding protein-independent variants of the

MalFGK₂ maltose transporter could be isolated (31, 32). Different roles of the binding protein were also inferred by Merino and Shuman (33), who reported that the unliganded maltosebinding protein, despite its inability to bind lactose, is required for a MalFGK₂ variant to transport lactose efficiently. The properties of these variants suggested substrate specificity was determined in the transmembrane regions of the transporter itself (31, 34). Participation of both the SBP and the transporter in determining substrate specificity was reported in a structural analysis of the maltose transporter with various bound substrates (35). More recently, studies have suggested a role for substrates binding directly to a preformed complex between the SBP and transporter, rather than delivered to the transporter by free SBP (28, 30).

Although these studies have provided evidence for a noncanonical role of SBPs in facilitating substrate uptake by ABC importers, no supporting structural evidence has been reported.

Significance

The high-affinity methionine importer MetNI belongs to the ATP Binding Cassette (ABC) family of transporters that carry out the ATP-dependent uptake of substrates into cells. As with other ABC importers, MetNI requires a soluble binding protein (MetQ) that in the canonical mechanistic model delivers substrates to the transporter. We made the unexpected observation that a MetQ variant with significantly impaired ligandbinding properties supports D-selenomethionine uptake at a higher rate than wild-type MetQ. A crystal structure of MetNIQ in the outward-facing conformation reveals access channels through the binding protein to the transmembrane translocation pathway. These studies support a noncanonical role for the binding protein in facilitating the uptake of certain substrates directly through the transporter–binding protein complex.

The authors declare no conflict of interest.

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Data deposition: The atomic coordinates and structure factors have been deposited in the Protein Data Bank, www.wwpdb.org (PDB ID code 6CVL). The MetNI and MetQ constructs generated in this work have been deposited in Addgene [ID codes 118268 (N229A MetQ) and 118269 (N229A E166Q MetNI) for the structural work, and ID codes 118253, 118254, 118256–118261, and 118581 for the transport assay].

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This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10. 1073/pnas.1811003115/-/DCSupplemental.

Published online October 23, 2018.

Author contributions: P.T.N. and D.C.R. designed research; P.T.N., J.Y.L., A.T.L., and J.T.K. performed research; P.T.N., J.Y.L., and A.T.L. contributed new reagents/analytic tools; P.T.N., J.Y.L., A.T.L., J.T.K., and D.C.R. analyzed data; and P.T.N. and D.C.R. wrote the paper.

Reviewers: B.P., University of Groningen; and N.Y., Princeton University.

In the course of characterizing the transport properties of MetNI, we made an unexpected observation that the N229A MetQ variant with greatly reduced ligand binding affinity supported D-selenomethionine uptake at a higher rate than wild-type MetQ. We solved a crystal structure of the ATP γ S-bound MetNI in complex with the ligand free N229A variant of MetQ. This structure reveals unprecedented features of the outward-facing conformation that support a noncanonical transport mode of the MetNI transporter in which MetQ facilitates access of substrate to the translocation pathway through the MetNIQ complex. The ability of intracellular methionine to inhibit the uptake of extracellular methionine (transinhibition), a key component of MetNI regulation described by Kadner (3), is also structurally detailed.

Unexpected High Transport Rates for a Substrate-Binding Deficient MetQ

To address the transport function of MetNI, we carried out in vivo transport assays using a $\Delta metNIQ$ knockout E. coli strain. There are two transport systems in E. coli that are responsible for methionine uptake (4): the higher-affinity MetD transport system (MetNIO operon) and the lower-affinity, and as yet uncharacterized, MetP system (36). The uptake of L-methionine is mediated by both systems, whereas D-methionine is specifically transported by MetD (2, 4, 36). To selectively study MetNI transport, we took advantage of the exclusive role of MetNI in Dmethionine transport and used a D-methionine derivative, Dselenomethionine (D-semet), as the transport substrate for quantification by inductively coupled plasma-mass spectrometry (ICP-MS) (Fig. 1A). Kinetic constants for the transport activity of D-semet derived from Michaelis-Menten analysis (Fig. 1B) are shown in Table 1. The maximal uptake rate of the wild-type MetNIQ is 6.3 ± 0.4 nmol·min⁻¹·mg⁻¹ of transporter, with K_m = 1.8 μ M (n = 3) (Fig. 1B and Table 1). Because L-methionine is present in the cytoplasm at 0.1-0.3 mM (1, 37), to bypass the potential transinhibition effect, an N295A mutation at the



Fig. 1. In vivo uptake assay of MetNIQ and its variants. (A) Schematic of whole-cell uptake assay for MetNIQ variants. (B) Michaelis–Menten plot of initial velocities for D-semet uptake versus substrate concentrations for MetNIQ and its variants. The corresponding V_{max} and K_m values are listed in Table 1.

Table 1. Summary of kinetic constants for transport activity of MetNIQ variants

MetNIQ variants	Uptake rate V _{max} *	K _m , μΜ
MetNIQ	6.3 ± 0.4	1.8 ± 0.4
MetNI	0.2 ± 0.1	N/A
N295A (EcN) MetNIQ	10 ± 0.5	1.7 ± 0.3
N229A (EcQ) MetNIQ	10.5 ± 0.9	7.4 ± 0.5
N295A, N229A MetNIQ	15.4 ± 1.3	4.8 ± 0.7
N295A, F103A (Ecl) MetNIQ	—	—
N295A, M107A (Ecl) MetNIQ	—	—
N295A, Y160A (Ecl) MetNIQ	14.0 ± 1.3	4.1 ± 0.2
N295A, M163A (Ecl) MetNIQ	4.7 ± 0.5	1.4 ± 0.2

The maximum uptake rate (V_{max}) and Michaelis–Menten constant (K_m) were determined from the plot shown in Fig. 1B. N/A, not applicable; dash, not detectable. The molar ratio of the MetQ:MetNI variants in these studies was measured to be ~4.

*nmol·min⁻¹·mg⁻¹ of transporter.

C2 domain of the MetNI was introduced that reduced the binding of L-methionine to undetectable levels as assessed by isothermal titration calorimetry (ITC). This variant results in a higher maximal uptake rate of 10 ± 0.5 nmol·min⁻¹·mg⁻¹ of transporter (n = 9) with no change in K_m (Fig. 1B and Table 1). We infer this represents the uptake rate of D-methionine by MetNI in the absence of transinhibition regulation by the internal pool of L-methionine.

Deletion of MetO significantly reduced the uptake rate of Dsemet to $0.2 \pm 0.1 \text{ nmol·min}^{-1} \cdot \text{mg}^{-1}$ of transporter (n = 3), which indicates that MetQ is required for the transport; however, an unanswered question is whether MetQ must bind substrate and deliver it to the transporter. To address this question, we generated a substrate-binding deficient variant of MetQ, N229A (NA MetQ), that reduces the affinity of MetQ for D-methionine by ~20-fold and for D-semet by approximately eightfold compared with the wild-type MetQ (SI Appendix, Fig. S1). Unexpectedly, the N229A MetNIQ mutation stimulates the maximal transport rate for D-semet ($V_{max} = 10.5 \pm 0.9 \text{ nmol·min}^{-1} \cdot \text{mg}^{-1}$ of transporter) (n = 6) by ~1.5-fold compared with the wild-type MetNIQ ($V_{max} = 6.3 \pm$ 0.4 nmol·min⁻¹·mg⁻¹ of transporter), with an approximately four-fold increase in K_m to 7.4 μ M (Fig. 1*B* and Table 1). A similar stimulation is observed for the MetQ N229A variant in combination with the MetN N295A MetNIQ ($V_{max} = 15.4 \pm 1.3 \text{ nmol·min}^{-1} \cdot \text{mg}^{-1}$ of transporter) (n = 6) compared with the MetN N295A MetNIQ ($V_{max} = 10 \pm 0.5 \text{ nmol·min}^{-1} \cdot \text{mg}^{-1}$ of transporter) (Fig. 1B and Table 1). These results indicate that the binding protein can facilitate substrate uptake through mechanisms in addition to the canonical role where the free SBP delivers the substrate to the transporter.

Conformational Changes in the Outward-Facing State

To complement our functional data, we determined the 2.95-Å resolution crystal structure of the ATP γ S-bound E166Q N295A MetNI in complex with N229A MetQ (Fig. 2A). As anticipated, MetNI adopts an outward-facing (OWF) conformation exhibiting significant conformational changes relative to the uncomplexed MetNI in the inward-facing (IWF) state (Fig. 2 B-D). A convenient reference frame is the molecular twofold axis that relates the pairs of MetNI subunits in each conformation. Relative to the IWF conformation, the MetI subunits in the MetNIQ complex have rotated $\sim 31^{\circ}$ toward the molecular twofold axis (SI Appendix, Fig. S2) opening the translocation pathway toward the periplasm and closing it at the cytoplasm. Relative to the transmembrane domains, the nucleotide binding domains (NBDs) are rotated $\sim 18^{\circ}$ toward the molecular twofold axis in going from the IWF to OWF conformation, facilitating dimerization of two NBDs in the presence of ATP_γS.



Fig. 2. Crystal structure of the MetNIQ complex. (*A*) Side-view representation of the MetNIQ. MetQ subunit is colored slate, MetI subunits are forest and pale yellow, and MetN subunits are cyan and firebrick. C-regulatory (C2) domains are at the C termini of MetN subunits. Conformational changes of (*B*) the transmembrane MetI subunits, (C) the nucleotide-binding MetN subunits, and (*D*) the C2 domains between their OWF (colored as described in *A*) and IWF conformations (PDB ID code 3TUJ) (colored gray). One subunit of MetI (colored forest) or MetN (colored cyan) of the OWF conformation was overlaid to that of the IWF conformation to highlight the differences in the relative placement of the opposite subunit (*B* and C). Superposition of MetNIQ and MetNI reveals the rotation of the C2 domains around the molecular twofold axis between the two structures (*D*).

MetNI-catalyzed transport is regulated by the intracellular concentration of methionine, a phenomenon known as transinhibition (1-3), which is mediated by the C2 domains located at the C-terminal end of the MetN subunits (8, 38, 39). Between the IWF [Protein Data Bank (PDB) ID codes 3DHW, 3TUI, and 3TUZ] and OWF conformations, the C2 domains rotate ~32° around the molecular twofold axis (SI Appendix, Fig. S2); this rearrangement is associated with the binding of L-methionine across the interface between C2 domains in the IWF conformation (SI Appendix, Fig. S34). The transition to the OWF conformation is accompanied by a repositioning of the C2 domains arising from a register shift (38) in the hydrogen-bonding network across the antiparallel β-sheet stabilizing the C2 dimer (SI Appendix, Fig. S3B). The toggling of the C2 dimer between these two states in the absence and presence of methionine shifts the separation between the two H-motif catalytic residues H199 from 13 to 24 Å (C α to C α), respectively. Although H199 and the adjacent 200-helix (residues 200-205) are part of the NBDs, each 200-helix maintains a fixed position with respect to a single C2 subunit (Fig. 3). Consequently, the shift in β -sheet register leads to a significantly increased separation between the H199 residues in the presence of methionine.

A Substrate Access Channel in the Outward-Facing State

Significant rearrangements are also evident in the conformation of MetQ in the MetNIQ complex relative to the methioninebound form. In contrast to the Venus flytrap-type of hinge bending motion that separates the two lobes of the MalE and BtuF SBPs in complexes with MalFGK₂ and BtuCD, respectively, MetQ in the MetNIQ complex shows a 24° twist around an axis perpendicular to the interface between the two lobes (Fig. 44). Significantly, this twisting motion creates an opening for substrate access through MetQ that is not present in the MalFGK₂E and BtuCDF complexes. The rmsd between the methionine-bound form of MetQ and the conformation of MetQ in complex with MetNI is 4.4 Å, which, despite the smaller ligand size, is larger than observed between the corresponding liganded and transporter-bound forms for MalE (40) and BtuF (41, 42) with rmsds of 3.2 and 1.0 Å, respectively. The outward hinge-type movements of the MalE and BtuF SBP lobes, together with insertions of periplasmic or scoop loops from the transmembrane subunits (Fig. 5A), have been proposed to facilitate substrate delivery from the SBP to the maltose and vitamin B12 transporters (22, 41). In contrast, scoop loops are not present in the MetNIQ complex (Fig. 5A). The combination of the large distortion of MetQ upon complex formation with MetNI and the relatively small size of the methionine substrate may allow exogenous methionine to access the MetNIQ complex through MetQ. To explore this possibility, the CAVER 3.0.1 program (43) was used to identify several possible substrate entry pathways in MetNIQ, using a shell radius of 3 Å. Identification of a continuous channel starting at the cleft between the two lobes of the complexed MetQ suggests that methionine cannot only access the MetQ substrate binding site but can also reach the translocation pathway through MetI in the MetNIQ complex (Fig. 4B). This structure therefore provides evidence for a substrate entry pathway in an ABC importer-SBP complex through the SBP.

Gating Mechanisms

Interconversion of the inward- and outward-facing conformations of MetNI is associated with rearrangements of the cytoplasmic and periplasmic gates (Fig. 6). At the periplasmic gate, residues Y177 and M163 move away from the center of the translocation pathway (Fig. 6*A*), compared with those in the IWF conformation of MetNI (Fig. 6*B*). On the other side, gating residue M107 (TM3) from both MetI subunits form a cytoplasmic gate, not seen in the IWF conformation of MetNI (Fig. 6*D*), restricting the translocation pathway (Fig. 6*C*). Although substitution of the periplasmic gate M163 to A reduces maximal transport activity (V_m= 4.7 ± 0.5 nmol·min⁻¹·mg⁻¹ of transporter)



Fig. 3. Rotation of the C2 domains facilitates ATP-induced dimerization of MetN subunits. Comparison of the C2 domains following superposition of one of the two C2 domain subunits. C2 domains are presented in the locked transinhibited IWF state (colored wheat; PDB ID code 3TUZ), the unlocked IWF state (colored light blue; PDB ID code 3TUJ), and MetNIQ in the OWF state [colored firebrick, structure 3; PDB ID code 6CVL (this study)]. The superimposed subunit is to the right in each panel. This comparison highlights the significant differences in the two subunits between these states. Of note, the separation between the 200-helices (cyan ellipses, *Left*) in the unlocked OWF MetNIQ (NIQ) and IWF DM states is significantly shorter than in the IWF locked state (CY5) in the presence of methionine. As a consequence, the catalytic H-motif H199 residues from both NBDs are too far apart in the locked state to form the ATPase active form of the NBDs. L-methionine bound to the regulatory sites on the C2 domains are depicted as yellow spheres.



Fig. 4. Conformational changes in MetQ creates substrate entry pathways in MetNIQ. (A) Superposition of the substrate-bound (colored cyan) and MetNI complexed MetQ (colored slate) shows a 24° twist between the two lobes that potentially provides access to potential substrate entry pathways. (*B*) Possible substrate entry pathways (shown in gray spheres) to the MetNIQ assembly were calculated by CAVER 3.0.2 pymol plugin with shell radii of 3 Å (43).

(n = 3), the cytoplasmic gate mutation M107A severely disrupts transport activity (Table 1).

A cavity is present between the MetI subunits in the OWF conformation (Fig. 5B), whose boundary is defined by residues Y160 and F103 at the top and bottom of the cavity, respectively (Fig. 7A), along the translocation pathway. This cavity is smaller than those observed in the maltose and vitamin B12 transporters (40, 42) (Fig. 5B) and could accommodate a methionine. No binding of methionine was observed, however, in the translocation pathway of MetNIQ either in the original structure determination or after soaking crystals in selenomethionine, or as inferred biophysically by ITC measurements using the preformed MetNIQ complex with N229A MetQ to bypass binding of D-methionine to SBP MetQ. This is in contrast to the maltose MalFGK₂ and arginine ArtQN transporters where substrates are found in the translocation pathway (40, 44, 45). Together with the gating residues M163 and M107, Y160 and F103 residues from both MetI subunits line the translocation pathway as if to form a selectivity filter, M- Ω - Ω -M (Ω , aromatic ring residue), for methionine selection (Fig. 7B). Aromatic rings from these residues may favor a selection toward the sulfur atom of the methionine residue, as seen in the binding pocket of holo-MetQ (17).

A Noncanonical Transport Mechanism for D-Methionine

Detailed structural mechanisms have been developed for the transport cycles of MalFGK₂E by Chen and coworkers (22) and BtuCDF by Locher and coworkers (23), reflecting a canonical role for the SBP in delivering substrate to the transporter. Our observations on the uptake of D-(seleno)methionine by MetNI support the presence of a distinct (noncanonical) mechanism where the initial substrate binding event is proposed to occur directly to the MetNIQ complex. The noncanonical and canonical models are compared in Fig. 8; a critical parameter distinguishing these pathways will be the affinity of the SBP for substrate, with the noncanonical pathway preferentially utilized by lower-affinity substrates such as D-(seleno)methionine and the

canonical pathway favored by higher-affinity substrates such as Lmethionine. Two intermediates are common to both pathways in this scheme, the IWF MetNI state 1 and the OWF MetNIQ state 3 containing bound substrate. The transition between state 3 and state 1 is accompanied by the translocation of substrate across the membrane. The distinction between the two pathways is provided by the precise sequence of events for the association of MetQ and substrate with the transporter. In the noncanonical pathway, unliganded MetQ associates with the ATP-bound form of MetNI to form state 2, corresponding to the MetNIQ structure determined in this work. D-(seleno)methionine from outside the cell then binds to generate the transport-competent state 3. In contrast, the canonical pathway proceeds through the association of liganded MetQ to MetNI to generate state 2', which consequently contains both SBP and substrate. State 2' has been modeled after the pretranslocation (pre-T) state of the maltose and molybdate transporters where binding of the liganded SBP precedes binding of ATP (22, 25); state 3 is then formed upon binding of ATP.

The regulation of the MetNI transport cycle through transinhibition arises from the binding of intracellular L-methionine to the C2 domains, resulting in transport termination. As established previously (38), the presence or absence of methionine toggles between two conformational states related by a register shift in the β -sheet hydrogen-bonding pattern at the dimer interface between C2 domains. In the transinhibited state 4, L-methionine bound to one C2 domain hydrogen bonds with N295 of the other (*SI Appendix*, Fig. S34), helping stabilize a hydrogen-bonding network including between the peptide NH and CO groups of the two A299 residues on adjacent β strands.



Fig. 5. Translocation pathways of different ABC importers. (A) Surface slab views of the outward-facing conformations of three ABC importers, including MetNIQ (this work), MaIFGK2 (PDB ID code 2R6G), and BtuCDF (PDB ID code 4FI3), reveal potential substrate-binding pockets (red box in *B*) with different sizes and shapes in the translocation pathways. Although MetI subunits form a small cavity, larger cavities are observed in the case of maltose and vitamin B12 transporters. Only maltose (shown in ball-and-stick) was structurally observed in the cavity. (*B*) Although a scoop loop was proposed to facilitate substrate handoff between the maltose- and vitamin B12-binding proteins with their transporters, no scoop loop is present in the methionine transporter.



Fig. 6. Conformational changes of the translocation pathway gates of the MetNIQ complex. Structure comparison of the MetI subunits in their outward-facing conformation (*A* and *C*) (colored forest and pale yellow) versus inward-facing conformation (*B* and *D*) (colored gray) reveals major rearrangements of the periplasmic gate residues, including Y177 and M163 (cyan; shown in ball-and-stick representation) (*A* and *B*), and of the cytoplasmic gate residues, including F103, M107, and Y160 (black; shown in ball-and-stick representation).

In the absence of bound L-methionine (states 1–3), the β -sheet hydrogen-bonding network between C2 domains shifts two residues, so that A299 is now paired with M301 (*SI Appendix*, Fig. S3B). As discussed above, the two conformations of the C2 domains are coupled to significant changes in the separation between the catalytic H-motif residues H199 and ultimately between the NBDs (Fig. 3). Transinhibition consequently reflects the increased distance between H199 residues in the presence of bound methionine, thereby precluding formation of the catalytically competent NBD dimer for ATP hydrolysis.

A Kinetic Transport Model for both High-Affinity and Low-Affinity Substrates

The ability of SBPs to mediate substrate uptake through both free (in the canonical model) and transporter-bound (in the noncanonical model) forms may represent a strategy to transport both high- and low-affinity substrates in physiologically relevant concentration ranges. A basic kinetic model (Fig. 9A) capturing these properties can be devised where the SBP (Q) can bind to the transporter (E_1) in either substrate-bound (Q_m) or substratefree (Q) to form $E_{2'}Q_m$ and $E_{2}Q$, respectively. These complexes can then convert into a transport-competent form E_3Q -m by isomerization (rate constant k_1) and binding of substrate [rate constant $k_2(m)$, which after transport returns to the initial E_1 state. An informative comparison can be made between the two cases where the substrate can bind to EQ and when it cannot (the noncanonical and canonical pathways, defined by $k_2 > 0$ and $k_2 = 0$, respectively), using a typical substrate concentration of 10^{-5} M. For high-affinity substrates (K_d << 10^{-5} M), there is little difference between these models, but for weakly bound substrates, the ability of E2Q to bind substrates allows transport to occur under condition of high Kd where Qm and E2'Qm would only be present at low concentrations (Fig. 9*B*). Although the details will depend on the specific parameter values used, evaluation of K_m as a function of K_d reveals that K_m plateaus above a threshold K_d in this model (Fig. 9*C*), so that transport can occur at physiological concentrations even for substrates that are weakly bound by the SBP. Together with reports from the maltose and vitamin B₁₂ uptake systems supporting the binding of substrate by the SBP–transporter complex (28, 30), these observations highlight an active role for both the SBP and transporter in the establishing the specificity and affinity of substrates that are imported by ABC transporters (33).

Materials and Methods

Cloning, Expression, and Purification. The *metN* and *metI* genes were amplified from *E. coli* K12 genomic DNA and cloned into a pET-modified dual vector, under two T7 promoters (pTN2) with the N-terminal hexahistidine tag sequence at the 5' end of *metN* as described previously (8, 46). The *metQ* gene encoding for a mature MetQ with no signal sequence was cloned into a separate pET19b (+) plasmid with N-terminal decahistidine tag followed by an enterokinase-cleavage site. All of the mutations, including E166Q, N295A in MetN, and N229A in MetQ, were done by site-directed mutagenesis (Stratagene). The cloned plasmids were expressed separately in *E. coli* BL21-gold (DE3) cells (EMD) at room temperature in ZY media (47).

Purification of MetNI mutants was carried out by homogenizing 10 g cell paste in 100 mL of lysis buffer containing 20 mM Tris-HCl, pH 8, 100 mM NaCl, 10% glycerol, 5 mM β-mercaptoethanol (βMe), 20 µg/mL DNase I, 200 µg/mL lysozyme. The cells were broken by three passes through a cell disruptor, M110L pneumatic microfluidizer (Microfluidics). C12E7 was added at a final concentration of 0.5% (vol/vol) to solubilize membranes at 4 °C for 2 h. Unlysed cells and cell debris were removed by centrifugation at $100,000 \times g$, 4 °C, and 30 min. The supernatant was collected. 70 mM imidazole, pH 8.0. was added, and the sample was then loaded onto a 5-mL Ni-Sepharose HP column (GE Healthcare) equilibrated with the MetNI purification buffer containing 20 mM Tris HCl, pH 7.5, 100 mM NaCl, 0.1% C12E7, 70 mM imidazole. After sample loading, the column was washed with 12 column volumes of the purification buffer. Two column volumes of elution buffer containing 20 mM Tris·HCl, pH 7.5, 100 mM NaCl, 0.1% C12E7, 300 mM imidazole were used to elute protein off the Ni column. The eluate was then subjected to size-exclusion chromatography. The eluted peaks were collected, concentrated, ultracentrifuged (267,000 \times g, 20 min, 4 °C) to remove aggregation, and adjusted to 12 mg/mL. Three hundred-µL-per-tube aliquots of MetNI were flash frozen in liquid nitrogen and stored at -80 °C.

Purification of signal sequence cleaved MetQ was done by resuspending 10 g cell paste in 100 mL of lysis buffer as mentioned above. Cell lysis was done by freezing and thawing for three cycles in liquid nitrogen and in a 42 °C water bath, respectively. Clearing of cell lysates was done by centrifugation at 37,500 \times g, 30 min, and 4 °C. After the supernatant was collected, 70 mM



Fig. 7. Residue rearrangement and potential substrate-binding cavity along the translocation pathway. (A) Movement of TM3 and TM4 of the Metl subunits form a small cavity (shown in Fig. SA), whose boundary is defined by residues Y160 and F103 along the translocation pathway. (B) Superposition of Metl subunits in the OWF and IWF conformation reveals a rearrangement of a potential selectivity filter, M-Ω-Ω-M (Ω, aromatic ring residues), along the translocation pathway of the OWF MetNIQ. Side chains in the OWF and IWF conformations are colored cyan and black, respectively.



Fig. 8. Mechanistic model for the MetNI-catalyzed transport of methionine derivatives. The relationship between the canonical and noncanonical pathways in the MetNI transport cycle is illustrated utilizing available structures of MetNI, MetQ, and MetNIQ represented as cylinders. Methionine, ADP, and ATP are shown as ball-and-sticks. Conformational states of the two C2 domains are shown in cyan and firebrick colored surface representations. State 1 represents a resting state of MetNI (PDB ID code 3TUJ). In the noncanonical pathway, state 2 represents an ATP-bound MetNIQ in the absence of methionine (this study; PDB ID code 6CVL). D-methionine can subsequently access the translocation pathway of the MetNIQ assembly through MetQ to form state 3 (also modeled as PDB ID code 6CVL). The canonical pathway proceeds through state 2' depicted as a hypothetical pre-T conformation of L-methionine–bound IWF MetNIQ, based on structural studies of the maltose and molybdate transporters (22, 25). The transition from state 3 to state 1 is common to both pathways and associated with methionine transport into the cell. During the transport cycle, the C2 domains are in unlocked conformations that can accommodate the changes between IWF and OWF states. State 4 represents the transinhibited state of MetNI transporter (PDB ID code 3TUZ) with the C2 domains in a locked conformation stabilized by methionine binding and rearrangement of the β -sheet hydrogen bonding pattern at the interface between the C2 domains.

imidazole, pH 8.0, was added before loading onto a 5-mL Ni-Sepharose HP column (GE Healthcare) equilibrated with purification buffer containing 20 mM Tris-HCl, pH 7.5, 100 mM NaCl, 70 mM midazole. After sample loading, the column was washed with 12 column volumes of the same buffer. Two column volumes of elution buffer containing 20 mM Tris-HCl, pH 7.5, 100 mM NaCl, 300 mM imidazole were used to elute protein off the Ni column. The eluate was then subjected to size-exclusion chromatography. The eluted peaks were pooled, incubated with enterokinase (NEB) to cleave the His tag, and passed through a handpacked Ni column again to remove uncut MetQ. The His-tag-cleaved MetQ was then concentrated to 20 mg/mL and flash-frozen in aliquots of 300 μ L per tube.

Selenomethionine-Substituted Proteins. To prepare selenomethionine-substituted MetNI and MetQ proteins, the cloned plasmids were transformed in *E. coli* auxotroph B834 (DE3) cells (EMD), which were then grown in phosphate-, amino acids-, and selenomethionine (PASM)-containing autoinduction media (47), containing 125 μ g/mL selenomethionine for 3–5 d at room temperature.

Crystallization of the *E. coli* MetNIQ Protein Complex. The native MetNIQ and selenomethionine-substituted MetNIQ complexes were prepared as de-

scribed previously (17). Crystallization screens for MetNIQ were conducted at different concentrations, 5–20 mg/mL, by vapor diffusion in hanging drops at 20 °C using three different commercial kits, including Mem-Gold2, MORPHEUS (Molecular Dimension), and PEGRx (Hampton Research). MetNIQ was crystallized in a reservoir containing 0.1 M Mes, pH 6, and 22% PEG 400, using 5 mg/mL protein at a ratio of protein to reservoir of 2:1. Crystals appeared after 2–3 d, fully grew after 5–7 d, and shattered after 10–14 d. Crystals were harvested at day 7 and cryoprotected by increasing PEG 400 concentration to 25, 30, and then 35%, followed by flash-freezing for data collection. Selenomethionine-substituted MetNIQ crystals often diffracted better than the native ones.

Heavy-Metal Derivatives. MetNIQ crystals were soaked in crystallization buffer (0.1 M Mes, pH 6, and 22% PEG 400) containing 1 mM K₂HgI₄ for an hour before back-soaking, cryoprotecting, and flash-freezing.

Data Collection and Structure Determination. All X-ray diffraction datasets were collected at the Stanford Synchrotron Radiation Laboratory (SSRL) beamline 12-2 equipped with a PILATUS 6M pixel array detector. Diffraction



Fig. 9. Kinetic analysis of canonical and noncanonical substrate uptake by ABC transporters. (A) Kinetic scheme modeling canonical and noncanonical pathways in substrate import by the transporter, E, where E1, E2, E2, and E3 represent different states of the transporter as shown in Fig. 8. The methionine ligand (m) binds to Q to form the liganded species Q_m with a dissociation constant K_d. Q_m and Q can bind to E₁ to form E₂·Q_m and E₂Q, with effective dissociation constants K1 and K2, respectively. Transport occurs from the state E3Q m that is generated either by isomerization of E2Qm with the first-order rate constant k1 or by binding of m to E2Q with a second-order rate constant k2. The steady-state solution to this scheme for the case where E, E2Qm, E2Q, Q, and Q_m are at equilibrium is described in Materials and Methods. (B) Dependence of the transport rate for a substrate at 10 µM concentration on various parameters of the kinetic scheme in A, as a function of the K_d for binding of substrate to the SBP. Curves *i*, *ii*, and *iii* correspond to k₂ = 5,000, 500, and $0 \text{ M}^{-1} \text{ s}^{-1}$, with $Q_{tot} = 10^{-4} \text{ M}$, $K_1 = K_2 = 10^{-4} \text{ M}$, and $k = k_1 = 0.01 \text{ s}^{-1}$. For substrates with high affinity to the SBP ($K_d < \sim 10^{-6} \text{ M}$) the models are equivalent, whereas for more weakly bound substrates ($K_d > \sim 10^{-4}$ M) the transport rate depends critically on the value of k_2 . Depending on the value of k_2 , the rate of transport of a weakly bound substrate can exceed that of a substrate that binds tightly to the SBP (compare curves i and ii). When k₂ = 0 (curve iii), transport does not occur under these conditions, which corresponds to the case where substrate is only delivered to E₁ by binding to the SBP. (C) Dependence of the K_m for transport on the dissociation constant K_d for binding of substrate to the SBP. The red and black circles represent experimental data points for the methionine (this study) and maltose transporter (30), respectively. The curves are generated with the following parameters: i, $k = 0.01 \text{ s}^{-1}$, $k_1 = 0.1 \text{ s}^{-1}$, and $k_2 = 0.01 \text{ s}^{-1}$, $k_2 = 0.01 \text{ s}^{-1}$, $k_1 = 0.1 \text{ s}^{-1}$, $k_2 = 0.01 \text{ s}^{-1}$, $k_2 = 0.01 \text{ s}^{-1}$, $k_3 = 0.01 \text{ s}^{-1}$, $k_4 = 0.01 \text{ s}^{-1}$, $k_5 = 0.01 \text{ s}^{-1}$, $k_5 = 0.01 \text{ s}^{-1}$, $k_5 = 0.01 \text{ s}^{-1}$, $k_6 = 0.01 \text{ s}^{-1}$, $k_8 = 0.01 \text{ s}^$ 1,000 M⁻¹·s⁻¹; *ii*, k = 0.1 s⁻¹, k₁ = 0.1 s⁻¹, and k₂ = 100 M⁻¹·s⁻¹; and *iii*, k = 0.1 s⁻¹, k₁ = 0.1 s⁻¹, and k₂ = 0 M⁻¹·s⁻¹, with Q_{tot} = 10⁻⁴ M and K₁ = K₂ = 10⁻⁴ M for all K₁ = 0.1 s⁻¹, k₁ = 0.1 s⁻¹, k₁ = 0.1 s⁻¹, k_2 = 0 M⁻¹·s⁻¹, k_1 = 0.1 s⁻¹, k_2 = 0 M⁻¹·s⁻¹ curves. Curves were not explicitly fit to the experimental data but reflect parameter values that approximate the data. For this kinetic scheme, K_m reaches a plateau as K_d increases, and depending on the parameter values, this plateau can be tuned to the physiological concentration range. For curve iii, where substrate does not bind to E_2Q , a plateau region is not reached, so substrates with poor affinities for Q (high K_d) would have correspondingly high K_m values.

images were processed and scaled with X-ray Detector Software (XDS) (SI Appendix, Table S1).

Several datasets diffracting anisotropically to 4-3 Å resolution were collected at wavelengths of 1.0000 Å (native crystals), 0.9794 Å (Semetsubstituted crystals), and 1.00582 Å (Hg-soaked crystals). Ellipsoidal truncation and anisotropic scaling were applied to the MetNIQ diffraction datasets using the University of California, Los Angeles, Molecular Biology Institute Diffraction Anisotropy server (services.mbi.ucla.edu/anisoscale) (48) (SI Appendix, Fig. S4). All crystals belonged to space group P3₂21 and showed partial twinning with twinning fractions ranging from 0.10 to 0.28 determined by Xtriage (Phenix) (49); therefore, a twin law with operator -h, -k, I was applied. Initial phases were obtained by molecular replacement-single anomalous dispersion (SAD) (AutoSol, Phenix) using SAD data from a 3.8-Å selenomethionine derivative crystal and a MalFGK₂-E (PDB ID code 2R6H)derived model of MetNI in an OWF conformation as a partial model. SAD experimental phasing (Phaser-EP; Phenix) using heavy-atom (selenium) sites was carried out to reduce model bias (SI Appendix, Fig. S5). These phases were extended to 3.5 Å resolution and then 2.95 Å resolution using two other higher-resolution datasets, the latter collected from a K₂HgI₄-soaked crystal. Coupled with density modifications (AutoSol: Phenix), the latter showed clear electron densities for MetQ, which allowed us to manually dock the two individual lobes of MetQ. Validation of the sequence register was obtained from the selenium positions and the location of mercury sites near cysteines in MetN observed from a K2Hgl4-soaked MetNIQ crystal. The MetNIQ model was refined using a combination of Phenix.refine (49), phenix.rosetta_refine (50), and Refmac5 (Collaborative Computation Project Number 4) (51). Although the densities of MetN and MetI subunits are well defined, that of MetQ is weaker (SI Appendix, Fig. S6). The absence of crystal contact in the MetQ of the MetNIQ crystal may result in some flexibility or even some orientational disorder. In general, the final model of MetNIQ fit the density calculated from the 2.95-Å resolution dataset well, and the final R_{work}/R_{free} = 0.21/0.23 (refinement statistics are in *SI Appendix*, Table S1).

Software used in the project was installed and configured by SBGrid (52).

Structure Comparisons. The OWF conformation of MetNI determined in this work (PDB ID code 6CVL) was compared with IWF conformations determined previously (8, 38): unlocked C2 domains in the presence of the detergent n-decyl-β-D-maltoside (DM; PDB ID code 3TUJ) and locked C2 domains in the presence of the detergent cyclohexyl-pentyl-β-D-maltoside (CY5; PDB ID code 3TUI) and with bound selenomethionine (PDB ID code 3TUZ). The

structure of methionine-bound MetQ (PDB ID code 4YAH) is described in Nguyen et al. (17).

In Vivo Transport Assays. Whole-cell uptake assays were conducted following an established protocol by the Poolman laboratory (24). In brief, different arabinose-inducible (pBAD) plasmids that carry variants of metN, metI, and metQ genes were transformed into *AmetNIQ E. coli* strain BW2115. Cells were grown in LB media to an OD_{600} of 1.5, induced for 2 h with 0.2% arabinose, and harvested by centrifuging for 10 min at 5,000 imes g, 4 °C. The cell pellets were then washed twice in 50 mM sodium phosphate buffer, pH 6.5, and resuspended to $\mathsf{OD}_{600}\sim10$ in the same buffer. Before transport assays, cells were energized with 0.5% glucose and 5 mM MgCl₂ for 10 min at 37 °C. The transport assays were started by adding D-selenomethionine (Dsemet) at different concentrations at 37 °C, shaking at 300 rpm. The reactions were then stopped by centrifugation at 18,000 \times g, 4 °C, for 1 min, followed by discarding the supernatant. The pellet was washed three more times in the same buffer before homogenizing in 20% HNO₃. The samples were incubated at 85 °C overnight and diluted 10-fold for quantifying selenium content by ICP-MS (8800 QqQ; Agilent) [Environmental Analysis Center, California Institute of Technology (Caltech)]. The amount of MetNI and MetQ in each reaction was quantified by Western blot using an anti-His antibody against His-tagged MetN and MetQ, respectively. A correlation between different substrate concentrations and uptake rate was shown in a final Michaelis-Menten plot (Fig. 1B) and a summary table (Table 1). Data were processed using SigmaPlot 10.0 (SigmaPlot).

Isothermal Titration Calorimetry. MetQ variants were dialyzed overnight in their purification buffers using a Slide-a-lyser mini dialysis device (Thermo Scientific). The samples were then ultracentrifuged at 267,000 × g, 4 °C, for 20 min to remove aggregates and the protein concentration adjusted to 100 μ M. Substrate D-methionine was prepared in the last dialysis buffer of each sample. Titrations of D-methionine to MetQ variants were done in triplicate on a MicroCal iTC-200 calorimeter at 25 °C. Titration curves were shown in *Sl Appendix*, Fig. S1. Data were processed using Origin v7.0 (OriginLab).

Steady-State Rate Solution to Kinetic Scheme in Fig. 9. E_1 , E_2Q , E_2Q_m , Q, and Q_m are assumed to be in equilibrium with dissociation constants defined in Fig. 9.A. The total SBP concentration, $Q_{tot} = (Q) + (Q_m)$, is assumed to be

much larger than the total transporter concentration, $E_{tot} = (E_1) + (E_2Q) + (E_2Q_m) + (E_3Q \cdot m)$, and so the contributions of (E_2Q) , $(E_2 \cdot Q_m)$, and $(E_3Q \cdot m)$ to Q_{tot} are neglected.

The concentrations of $(E_3Q \cdot m)$ and $(X) = (E_1) + (E_2Q) + (E_2 \cdot Q_m)$ may be determined using the steady-state treatment detailed in Segel (53) and Cha (54), yielding

$$\begin{split} E_{tot} &= \quad (E_1) \quad + (E_2 Q) \quad + (E_2 \cdot Q_m) \quad + (E_3 Q \bullet m) \\ & \propto \{ k(K_1 K_2 (K_d + (m))) \} + \{ k K_1 K_d Q_{tot} \} + \{ k K_2 Q_{tot} (m) \} \\ & \quad + \{ (k_1 K_2 + k_2 K_1 K_d) Q_{tot} (m) \}. \end{split}$$

The rate of transport $[v=k(E_3Q\text{-}m)]$ relative to the maximum possible rate $(V^\circ=kE_{\rm tot})$ is

$$\begin{split} & \frac{v}{V^{o}} = \frac{\kappa(t_{3}Q \bullet m)}{kE_{tot}} \\ & = \frac{(k_{1}K_{2} + k_{2}K_{1}K_{d})Q_{tot}(m)}{\{k(K_{1}K_{2}(K_{d} + (m)))\} + \{kK_{1}K_{d}Q_{tot}\} + \{kK_{2}Q_{tot}(m)\} + \{(k_{1}K_{2} + k_{2}K_{1}K_{d})Q_{tot}(m)\}} \end{split}$$

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Putting this into standard form allows the evaluation of V_{max} and K_m (V_{max} in this context represents the fraction of the maximum possible rate V°):

$$\begin{split} & \frac{v}{V^{o}} \equiv \frac{V_{max}(m)}{K_{m} + (m)} \\ \Rightarrow & V_{max} = \frac{(k_{1}K_{2} + k_{2}K_{1}K_{d})Q_{tot}}{(k_{1}K_{2} + k_{2}K_{1}K_{d})Q_{tot} + kK_{2}(K_{1} + Q_{tot})} \\ \Rightarrow & K_{m} = \frac{kK_{1}K_{d}(K_{2} + Q_{tot})}{kK_{1}K_{2} + kK_{2}Q_{tot} + k_{1}K_{2}Q_{tot} + k_{2}K_{1}K_{d}Q_{tot}} \end{split}$$

In the limits as $K_d \rightarrow 0$ and $K_d \rightarrow \infty,$ the values of K_m approach

$$\begin{split} \mathsf{K}_{d} &\rightarrow 0, \ \mathsf{K}_{m} \rightarrow \frac{k \ \mathsf{K}_{1}(\mathsf{K}_{2} + \mathsf{Q}_{tot})}{k \ \mathsf{K}_{1}\mathsf{K}_{2} + k \ \mathsf{K}_{2}\mathsf{Q}_{tot} + \mathsf{k}_{1}\mathsf{K}_{2}\mathsf{Q}_{tot}}\mathsf{K}_{d} \\ \mathsf{K}_{d} &\rightarrow \infty, \ \mathsf{K}_{m} \rightarrow \frac{k \ (\mathsf{K}_{2} + \mathsf{Q}_{tot})}{k_{2}\mathsf{Q}_{tot}}, \end{split}$$

so that for high-affinity substrates (K_d \to 0), K_m \propto K_d, and for low-affinity substrates, K_m plateaus to a constant value.

This scheme focuses on the interactions of the transporter, SBP, and ligand m and omits many features of the transport cycle, including the binding of ATP, ATP hydrolysis, and dissociation of the products ADP and Pi, and the interconversion of inward- and outward-facing conformations. Because there

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is a strong nucleotide state dependence to SBP binding, the dissociation constants in scheme Fig. 9A have been approximated as effective dissociation constants described by the ratio of the dissociation rate constants for E₂Q or E₂·Q_m and the association rate constant between E₁ and Q or between E₁ and Q_m. For MetNIQ, the association rate constant is ~10³ M⁻¹·s⁻¹, independent of whether MetQ has bound methionine or not and independent of whether MetNI is in the ATP or nucleotide free states (29, 55). Under turnover conditions, dissociation of the complex will be set by the rate of ATP hydrolysis, which has been measured to be ~25 min⁻¹ or ~0.3 s⁻¹ (39) so that as an order of magnitude estimate, K₁ ~ K₂ ~ 10⁻⁴ M⁻¹. For comparison, the dissociation constants for the binding of unliganded and liganded MetQ to the ATP-bound form of MetNI are 27 nM and 1 µM (17), respectively, when ATP hydrolysis is prevented through either mutation or absence of Mg⁺². The calculations for Fig. 9 were performed with Mathematica version 9.0.0.

The transport rate determined for the N295A MetNI transporter with pselenomethionine [~10 nmol·min⁻¹·mg⁻¹ (Table 1)] is equivalent to a turnover time of ~0.02 s⁻¹, which provides an estimate for the rate constant k; this rate is consistent with measurements made on other ABC transporters, with turnover times of ~0.002–0.03 s⁻¹ (24, 26, 56–58).

Data and Materials Availability. Coordinates and structure factors have been deposited in the Protein Data Bank of the Research Collaboratory for Structural Bioinformatics, with PDB ID code 6CVL. The MetNI and MetQ constructs have been deposited with Addgene, with IDs 118269 (N295A E166Q MetNI) and 118268 (N229A MetQ) for the structural work, and IDs 118253, 118254, 118256–118261 and 118581 for the transport assay (Table 1).

ACKNOWLEDGMENTS. We thank Dr. Raymond Liu for critical reading of the manuscript and Dr. Nathan Dalleska and Stefan Petrovic for assistance with ICP-MS and ITC, respectively. A Vietnam International Education Development scholarship from the Vietnam Ministry of Education and Training (to P.T.N.) is gratefully acknowledged. We gratefully acknowledge the Gordon and Betty Moore Foundation and the Beckman Institute at Caltech for their generous support of the Molecular Observatory at Caltech and the staff at Beamline 12-2, SSRL, for their assistance with data collection. SSRL is operated for the US Department of Energy and supported by its Office of Biological and Environmental Research and by the NIH, National Institute of General Medical Sciences (P41GM103393), and the National Center for Research Resources (P41RR001209). We thank the Center for Invironmental Microbial Interactions for their support of microbiology research at Caltech. D.C.R. is an investigator at Howard Hughes Medical Institute.

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