

Predicting the Three-Dimensional Structure of Human P-Glycoprotein in Absence of ATP by Computational Techniques Embodying Crosslinking Data: Insight into the Mechanism of Ligand Migration and Binding Sites

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ABSTRACT P-glycoprotein is a membrane protein involved in the phenomenon of multidrug resistance. Its activity and transport function have been largely characterized by various biochemical studies and a low-resolution image has been obtained by electron microscopy. Obtaining a high-resolution structure is, however, still remote due to the inherent difficulties in the experimental determination of membrane protein structures. We present here a three-dimensional (3D) atomic model of P-glycoprotein in absence of ATP. This model was obtained using a combination of computational techniques including comparative modeling and rigid body dynamics simulations that embody all available cysteine disulfide crosslinking data characterizing the whole protein in absence of ATP. The model features rather well most of the experimental interresidue distances derived both in the transmembrane domains and in the nucleotide binding domains. The model is also in good agreement with electron microscopy data, particularly in terms of size and topology. It features a large cavity detected in the protein core into which seven ligands were successfully docked. Their predicted affinity correlates well with experimental values. Locations of docked ligands compare favorably with those suggested by cysteine-scanning data. The finding of different positions both for a single ligand and for different ligands corroborates the experimental evidence indicating the existence of multiple drug binding sites. The interactions identified between P-glycoprotein and the docked ligands reveal that different types of interactions such as H-bonds, π - π and cation- π interactions occur in agreement with a recently proposed pharmacophore model of P-glycoprotein ligands. Furthermore, the model also displays a lateral opening located in the transmembrane domains connecting the lipid bilayer to the central cavity. This feature supports rather well the commonly admitted mechanism of substrate uptake from the lipid bilayer. We propose that this 3D model may be an important tool to understand the structure-function relationship of P-glycoprotein. *Proteins* 2006;63:466–478. © 2006 Wiley-Liss, Inc.

Key words: ABC transporters; computer-assisted molecular modeling; drug transport; membrane proteins; multidrug resistance; P-gp

INTRODUCTION

The P-glycoprotein (P-gp), product of the *mdr1* gene in humans, is an ATP-dependent transporter that extrudes a large range of structurally diverse compounds out of eucaryotic cells. P-gp is found at the level of elimination organs, like intestines, kidney, and liver, where it is involved in the secretion of drugs and metabolites, as well as at the blood-brain barrier, where its efflux activity prevents the accumulation of cytotoxic agents in the central nervous system.^{1,2} The overexpression of P-gp is also associated with a multidrug resistance phenotype in various forms of cancer,³ causing suboptimal outcomes in chemotherapy.

P-gp is a member of the ATP-binding cassette (ABC) family of transporters.^{4,5} It is a single polypeptide of 1280 residues, organized as two homologous halves (which are 43% identical in human P-gp) of ~ 610 residues joined by a ~ 60 residues linker. Each half consists of six transmembrane (TM) segments followed by a cytoplasmic nucleotide-binding domain (NBD). The two halves of P-gp are essential for the activity of the transporter. They cannot act autonomously and instead appear to function coordinately, suggesting that they interact in some fashion. Several helices of the TM domains have been proposed to

Abbreviations: ABC, ATP-binding cassette; EM, electron microscopy; MTS, methanethiosulfonate; NBD, nucleotide-binding domain; P-gp, P-glycoprotein; TM, transmembrane; 3D, three-dimensional.

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contain the binding site(s) and consequently are believed to form the pathway through which the substrates cross the membrane. On the other hand, the NBDs couple the energy associated with ATP binding and hydrolysis to drug transport.⁶ The linker of ~ 60 residues does not appear to be essential to the protein function though when present its flexibility appears to be a prerequisite.⁷

The P-gp mediated-transport mechanism has been extensively studied but remains controversial. Briefly, it is believed^{8,9} that the transport cycle is initiated by substrate binding in the TM domains of P-gp, which increases the ATP affinity for the protein. After binding and/or hydrolysis of a first ATP molecule in one of the NBDs several restructurings occur in the TM domains of the protein. This conformational change allows the release of the drug to the extracellular medium. After hydrolysis of a second ATP molecule P-gp returns to its original configuration. The protein is then reset for another cycle.

A first low-resolution structure obtained by electron microscopy (EM) on detergent solubilized P-gp suggests that the TM domains form a cone-shaped chamber in the membrane, open towards the extracellular face.¹⁰ A second structure was obtained by EM for the protein reconstituted in a lipid bilayer.¹¹ This structure, which is the only one representative of the protein in its native environment, presents P-gp in a conformation in which the two NBDs are in close interaction. A third structure, derived at a resolution of 8 Å for P-gp bound to a nonhydrolysable analog of ATP,¹² provides valuable data on the spatial arrangement of the TM helices and the NBDs. Mutagenesis,¹³ photoaffinity labeling,¹⁴ and cysteine scanning and crosslinking studies^{15–25} have identified several residues involved in drug binding in the TM domains and have permitted to obtain structural information on the interactions between the TM domains and between the NBDs. These studies, however, do not supply with a detailed atomic structure of P-gp, which would be of considerable help in elucidating the molecular mechanism for substrate binding and release.

Crystallization of membrane proteins to obtain direct information on three-dimensional (3D) structure is still a difficult task. At this time three different 3D models of P-gp at an atomic level have been elaborated,^{26–28} but none of them fully takes into account the available experimental structural data. We therefore present here a new 3D model that was elaborated by the combination of different computational methods such as comparative modeling and rigid body dynamics. We used the structure of *E. coli* MsbA²⁹ as a template to generate an initial model. However, we did not exploit the structure of the dimer as its structural features do not match those expected for P-gp structure. Instead we resorted to the structure of each monomer independently to model the two halves of P-gp. To feature the close association of the two NBDs in P-gp as witnessed by several experimental studies, the 3D structure of BtuCD³⁰ was also used. Rigid body dynamics involving distance-restrained potentials was then applied to the initial model. This step was performed to include in our model cysteine-scanning mutagenesis

data describing helices orientation and interresidue distances. The validity of the model is then discussed in regard to the available experimental data.

The interaction of compounds with P-gp is clearly a complex process. A general pharmacophore model of P-gp drugs has been recently proposed,³¹ and involves two hydrophobic points, three H-bond acceptor points and one H-bond donor point organized in a precise geometry. Our 3D model was therefore used to dock several ligands in the central cavity harbored by the TM domains. The geometry of interaction of the ligands and their predicted affinity are examined and compared with the experimental data.

MATERIALS AND METHODS

Sequences Alignment

All protein sequences were obtained from the Swiss-Prot data base. The sequences of the N- and C-terminal halves of P-gp (residues 1–630 and 690–1280, respectively) were aligned pairwise with the sequence of *E. coli* MsbA monomer using the program ClustalW.³²

The sequences of the NBD of each P-gp half (residues 386–630 and 1029–1280) were aligned with NBDs of closely related ABC transporters (*E. coli* BtuCD, *E. coli* MsbA, *P. fur.* Rad50, and MJ0796) using ClustalW.

The alignment of the sequences used in the comparative modeling process is shown in Figure 1.

Comparative Modeling

E. coli MsbA is the protein with the highest percentage of sequence identity to P-gp whose 3D structure is known. However, for reasons explained in the Results and Discussion section, we did not use the dimeric structure because of its complete discrepancy with the experimental data on P-gp. Nevertheless we resorted to each MsbA monomer as a template to generate an initial model of each P-gp half using the sequence alignment shown in Figure 1.

The X-ray structure of MsbA²⁹ (PDB code: 1JSQ) contains the C α atom positions only. The structure of *E. coli* MsbA was then completed by using MaxSprout,³³ a database algorithm to generate the missing protein backbone and sidechains. Additional templates for P-gp modeling were searched with the Blast program³⁴ for the tiny portions for which there is no corresponding 3D structure in MsbA (see Fig. 1). Three structures were found (PDB codes: 1F3M, 1JJ7, 1FB3).

Modeling of the P-gp NBD dimer was performed using as templates the NBD dimer of BtuCD and each NBD monomer of MsbA following the alignment given in Figure 1. The BtuCD structure³⁰ (PDB code: 1L7V) is a good template to model the close association of the dimeric structure of the NBDs and the relative orientation of each P-gp monomer, as observed experimentally.

Comparative modeling was performed with the Modeller 5 software,³⁵ using the model routine with the alignments described on Figure 1. Ten models were generated, and the structure with the most favorable intramolecular energy, as described by the objective function of the program, was selected to represent an initial model.

P-gp	112	TRYAYYSYGIGAGVLVAAYIQVSWFCWLAAGRQIHKIRKQFFHAIMRQEIWFVDFVHVGELNRLTDDVSKINEGIGDKIGMFFQSMATFFTFIVGFTRG	211
MsbA	65	VWMLPVVIGLMILRGITSYVSSYICISWVSGKVVMTMRRRLFGHMMGMPVSVFFDKQSTGTLRSRITYDSEQVASSSSGALITVVREGASIIGLFIMMFYYS	164
BtuD		-----	
1F3M		-----	
1JJ7		-----	
P-gp	212	WKLTLVILAIISPVGLSAAVWAKILSSFTDKELLAYAKAGAVAEVLAARTVIAFGGQKKELERYNKNLEEAKRIGIKKAITANISIGAAFLLIYASYA	311
MsbA	165	WQLSIIILIVLAPIVSIARVSVSKRFRNISKNMQNTMGQVTTSA-----RLQGMKMSASSISDPIIQLIASLALA	264
BtuD		-----	
1F3M	C1	-----SDEEILEKLSRISVSGDPKKKYTRFEKIQGASG-----	C34
1JJ7		-----	
P-gp	312	LAFWYGTTLVLSGEYSIGQVLTVFFSVLIGAFVSVGQASPSIEAFANARGAAEYIFKIDNKPSIDSYSKSGHKPDNIKGNLEFRNVHFSYPSRKEVKILK	411
MsbA	265	FVLYAASFPSVMSLDTAGTITVVFSSMIALMRPLKSLTNVNAQFQRGMAACQTLFTILDSEQEKD--EGKRVIERATGDVE-----	343
BtuD	2	-----SIVMQLQDVA-----ESTRLG	17
1F3M		-----	
1JJ7	A14	-----VQFQDVSFAYPNRPDVLVLQ	A34
P-gp	412	GLNLKVSQGTVALVGNSSGCGKSTTVQLMQRLYDPTGEMVSDGGDIRTINVRFLREIIGVVSQEPVL-FATTIAENIRYGR-ENVMTDEIEKAVKEANA	509
MsbA	419	-----VALVSQNVHL-FNDTVANNIAYARTEQYSREQIEEAARMAYA	459
BtuD	18	PLSGEVVRAGEILHLVGPNGAGKSTLLARMAGMTSGK-GSIQFAGQPLEAWSATKLALHRAVLSQQQTPPFATPVWHYLTLHQ-HDKRTELLNDVAGALA	115
1F3M		-----	
1JJ7	A35	GLTFTLRPGVETALVGPNGSGKSTVAALLQNLQYQPTGGQLLDDGKPLPQYEHRYLHRQVA-----	A94
P-gp	510	YDFIMKLPKFDTLVGERGALSGGQKQRIAIARALVR----NP--KILLDEATSALDTESEAVVQVALDKA--RGRRTIVIAHRLS-TVRNADVIA	599
MsbA	460	MDFINKMDNGLDVTIGENGVLLSGGQQRQRIAIARALLR----DS--PILILDEATSALDTESERAIQALDEL--QKNRTSLVIAHRLS-TIEKADEIV	549
BtuD	116	-----LDDKLGRSTNQLSGGEWQRVRLAAVVLIQITPQANPAGQLLLEDPMNSLDVAQQSALDKILSALCQ-QGLAIVMSSHDNLNHLRHAHRA	204
1F3M		-----	
1JJ7		-----	
P-gp	510	GFDDGVIVEKGNHDE	614
MsbA	550	VVEDGVIVERGTHND	564
BtuD	205	LLKGGKMLASGRREE	219
1F3M		-----	
1JJ7		-----	
<hr/>			
P-gp	753	NLFSLLFLALGIIISFITFFLQGFVFGKAGEILTCLRMYMFRSMLRQDVSWFDDPKNTTGALTRRLANDAAQVKAIGASRLAVITONIANLGTGIIISFI	852
MsbA	65	VWMLPVVIGLMILRGITSYVSSYICISWVSGKVVMTMRRRLFGHMMGMPVSVFFD--KQSTGTLRSRITYDSEQVASSSSGALITVVREGASIIGLFIMMFY	162
BtuD		-----	
1FB3		-----	
1JJ7		-----	
P-gp	853	YGWQLTLLLLAIVPIIAIAGVEMKMLSGQALKDKKELEGAGKIATEAIENFRTVVSLTQEQ--KFEHMYAQSLOVQPYRNSLRKAHIFGITFSFTQAMM	949
MsbA	163	YSWQLSIIILIVLAPIVSIARVSVSKRFRNISKNMQNTMGQVTTSA-----RLQGMKMSASSISDPIIQLIA	259
BtuD		-----	
1FB3	A201	-----MKEKAPENFRLDFAVSREQTNEKGEKMYIQTRMAQYA-----	A237
1JJ7		-----	
P-gp	950	YFSYAGCFRFGAYLVAHKLMSFEDVLLVFSVAVFGAMAVGVSSFAPDYAKAKISAAHIIMIIEKTPILDSYSTEGLMPTLEGVNTFGEVVFNYPTRPD	1049
MsbA	260	SLALAFVLYAASFPSVMSLDTAGTITVVFSSMIALMRPLKSLTNVNAQFQRGMAACQTLFTILDSEQEKD--EGKRVIERATGDVE-----	343
BtuD	2	-----SIVMQLQDVA-----E	12
1FB3		-----	
1JJ7	A15	-----VQFQDVSFAYPNRPD	A29
P-gp	1050	IPVLQGLSLEVKKQTLALVGSNGCGKSTVVQLLERFYDPLAGKVLVDGKEIKRLNVQWLRALHGVSVQEPIL-FDCSIAENIAYGDNRSVVSQEEIVRA	1148
MsbA	419	-----VALVSQNVHL-FNDTVANNIAYARTEQ-YSREQIEEA	453
BtuD	13	STRLGPLSGEVVRAGEILHLVGPNGAGKSTLLARMAGMTSGK-GSIQFAGQPLEAWSATKLALHRAVLSQQQTPPFATPVWHYLTLHQHD--KTRTELLND	109
1FB3		-----	
1JJ7	A30	VLVLQGLTFTLRPGVETALVGPNGSGKSTVAALLQNLQYQPTGGQLLDDGKPLPQYEHRYLHRQVA-----	A94
P-gp	1149	AKEANIHFIESLPNKYSTKVGDKGTQLSGGQKQRIAIARALVR-----QPHILLDEATSALDTESEKVVQEAALDKA--REGRTCIVIAHRLS-TIQ	1238
MsbA	454	ARMAYAMDFINKMDNGLDVTIGENGVLLSGGQQRQRIAIARALLR----DSPILILDEATSALDTESERAIQALDEL--QKNRTSLVIAHRLS-TIE	543
BtuD	110	V--AGAL-----ALDDKLGRSTNQLSGGEWQRVRLAAVVLIQITPQANPAGQLLLEDPMNSLDVAQQSALDKILSALCQ-QGLAIVMSSHDNLNHLR	198
1FB3		-----	
1JJ7		-----	
P-gp	1239	NADLIVVFQNGRVKEHGTHQQ	1259
MsbA	544	KADEIVVVEDGVIVERGTHND	564
BtuD	199	HAHRAWLLKGGKMLASGRREE	219
1FB3		-----	
1JJ7		-----	

Fig. 1. Alignment of the sequences used in the comparative modeling procedure. (Top panel) Alignment to the N-terminal half of P-gp. (Bottom panel) Alignment to the C-terminal half of P-gp. The first line is the P-gp sequence to be modeled; second line is the sequence of MsbA; third line is the sequence of BtuCD (used for the modeling of the NBDs); and the last two lines are sequences of protein structures used to model tiny portions of P-gp for which structure is missing in MsbA.

The initial model of each P-gp half was then fitted on the NBD dimer model. The fitting was performed using the positions of the C $_{\alpha}$ atoms.

Rigid Body Molecular Dynamics

Starting from the model issued from the comparative modeling and fitting procedure, a P-gp 3D model was

generated by resorting to multibody molecular dynamics simulations. This approach is a multigranular modeling whereby different parts of the molecule are modeled at different levels of detail.³⁶ The molecular system is substructured in rigid bodies and atomistic-described regions. The molecular dynamics simulation was performed with the CHARMM c27b3 software.³⁷ The transmembrane helices and the NBDs were substructured as separate rigid bodies. These bodies are allowed to undergo large motions relative to each other but no motion occurs within a body. The remaining loops connecting helices and NBDs were treated as atomic particles or single atom bodies.

For the modeling of the partial model (i.e., helices 4, 5, 6, 10, 11, 12, NBD1, and NBD2), three successive molecular dynamics simulations of 30,000 steps were performed. For the modeling of the final model (adding helices 2, 3, 8, and 9 to the previous model), a single molecular dynamics simulation of 30,000 steps was performed. All the simulations were performed using the Lobatto integrator with a time step of 1 fs. The temperature of the bath was set to 50 K in order to avoid excessive heating of the system.

Force-field interactions were obtained by a conventional all-atom CHARMM force field (CHARMM22).³⁸ Distance-based restraint potentials were added to the classical potentials so as to account for structural experimental evidence. These distances were derived from cysteine crosslinking experiments with two types of reactants.^{15,16,18–21,27} When crosslinking between two residues was observed with $\text{Cu}^{2+}(\text{phen})_3$ as oxidant at a temperature of 4°C, a distance of 8 to 10 Å between the two C α atoms involved was used. For the experiments performed at 21°C or 37°C, the maximum distance value was increased to 15 Å in order to take into account the increased mobility of the protein. When crosslinking was observed with methanethiosulfonate (MTS) compounds as substrates, a 3D model of their structure was generated using the Corina program.³⁹ The distances were then derived by measuring the end-to-end distance of the 3D structure of these molecules. The distances measured this way may somewhat differ from those given in the papers of Loo and Clark.^{15,16,18–21} Table I lists the distance values assigned in the rigid body dynamics procedure.

Each rigid body molecular dynamics was followed by an energy minimization of 500 steps with the steepest descent algorithm.

In order to proceed with the docking experiments positioning of the sidechains was performed again with the specialized program SCWRL.⁴⁰

Docking

In order to assign the correct protonation state of each molecule, pKa values were obtained either from experimental data or predicted using the ChemSilico program.⁴¹ According to these values all the ligands bear a positive charge at physiological pH. Verapamil was built as R-verapamil (also known as dexverapamil). It is indeed known that racemic verapamil and R-verapamil have identical inhibition constant values.⁴² The 3D structures of the ligands were generated with the Corina program.

Their atomic partial charges were computed with MOPAC using the semiempirical MNDO Hamiltonian.⁴³ The atomic partial charges for the protein were taken from the CHARMM19 force field.³⁷

Docking was performed with the Autodock program (v. 3.0),⁴⁴ using the implemented genetic algorithm and the default parameters. Twenty-five independent docking runs were performed for each ligand.

RESULTS AND DISCUSSION

Modeling of P-gp Structure

Because the structure of P-gp undergoes several conformational changes along its catalytic cycle, it is important to decide which conformation to model, and carefully sort the experimental data describing it. We chose to produce a model reflecting the structure of P-gp in absence of ATP. This structure has been extensively studied by cysteine scanning mutagenesis and cysteine disulfide crosslinking experiments,^{15–25} which permitted to obtain important structural information about the relative positions of different parts of the TM domains, NBDs, and on the substrate binding site(s).

The search for proteins of known structure which share at least 20% of global sequence identity with P-gp, produced only two possibilities: the structures of MsbA of *E. coli*²⁹ (PDB code: 1JSQ) and of *V. cholera*⁴⁵ (PDB code: 1PF4). Both structures were recently determined by X-ray spectroscopy to a 4.5 and 3.8 Å resolution, respectively. The two proteins are homodimers of 1164 residues that exhibit 34% and 27%, respectively, of sequence identity with human P-gp, which makes them potential candidates to elaborate a 3D model for P-gp using comparative modeling. However, these two structures could not be directly used as a template at least in their dimeric form. There is indeed considerable support from cysteine disulfide crosslinking studies,²¹ EM,¹¹ and fluorescence energy transfer studies⁴⁶ for the two NBDs of P-gp to be in close association. This feature is also well supported by recent X-ray structural characterization of the NBDs of other ABC transporters such as Rad50,⁴⁷ MJ0796,⁴⁸ and BtuCD.³⁰ In contrast the two NBDs in *E. coli* MsbA structure are as far as 50 Å. The MsbA dimer structure of *V. cholera* shows an arrangement in which the two NBDs are in close association, seemingly in accordance with experimental data on P-gp. However, a closer look at the structure reveals that the two ATP binding sites are located on the faces opposite to the dimer interface, which appears to us as an unnatural configuration as it is in disagreement with several experimental studies.^{11,21,30,46–48}

Though MsbA seemed, in terms of sequence identity, a good template candidate to model the P-gp atomic structure, neither *E. coli* nor *V. cholera* could be used in their dimeric form. In order to circumvent this problem and to generate a model for P-gp in agreement with the largest number of experimental data, we used a combination of computational techniques. The different steps of the modeling procedure are explained below. In a first step, each P-gp half was modeled independently using the structure of MsbA monomers from *E. coli*. P-gp is indeed composed of two homologous halves having 36% and 32 % of sequence

TABLE I. Interresidues C_α-C_α Distances Used in Rigid Body Dynamics

		Initial	Min. Restr.	Max Restr.	Final	Delta			Initial	Min. Restr.	Max Restr.	Final	Delta
TM 4-10	S222 1868	81.4	13	23.5	23.6	0.1		L531 S1071	13.6	8	15	12	0
	S222 G872	77.7	13	23.5	25.8	0.3		L531 S1072	11.9	8	15	10.8	0
4-12	L227 S993	71.2	8	15	21.4	6.4		L531 G1073	10.7	8	10	8.5	0
	V231 S993	66.5	8	15	18.4	3.4		L531 C1074	13.7	8	10	11	1
	W232 S993	62.7	8	15	15.9	0.9		L531 G1075	15.9	8	15	13.7	0
	A233 S993	62.6	8	10	16.5	6.5		L531 K1076	18.6	8	15	16.8	0.8
5-10	1306 1868	77	15	23.5	18.1	0		S532 S1072	10.7	8	15	10.7	0
	1306 G872	72.6	15	23.5	12.9	2.1		S532 G1073	9.8	8	10	9.2	0
								S532 C1074	12.1	8	15	10.9	0
5-11	1306 T945	81.5	15	23.5	24.1	0.6		S532 G1075	15	8	15	14.2	0
5-12	A295 S993	66.4	8	15	18.4	3.4	NBD- NBD	G533 S1072	10.1	8	15	9.7	0
	1299 S993	69.1	8	10	15.2	5.2		G533 G1073	10.6	8	15	9.6	0
	1306 V982	76.6	15	23.5	17.5	0		G533 C1074	12.8	8	15	11.7	0
	1306 G984	77.1	15	23.5	21.9	0		G534 S1071	13.9	8	15	13.3	0
6-10	P350 V874	68.1	8	15	20.4	5.4		G534 S1072	13.6	8	15	12.9	0
	P350 E875	64.4	8	15	17.5	2.5		G534 S1072	13.6	8	15	12.9	0
	P350 M876	64.1	8	15	19.3	4.3		Q535 S1071	16.8	8	15	16.1	1.1
	L339 1868	72.6	15	23.5	15.4	0		S429 L1176	13.8	8	15	13	0
	L339 G872	68.2	15	23.5	15.2	0		G430 L1176	11.8	8	15	10.3	0
	L332 Q856	92.5	13.5	19	24.8	5.8		C431 L1176	14.6	8	10	12.5	2.5
6-11	P350 G939	64.5	8	15	18.1	3.1		G432 L1176	16.4	8	15	14.8	0
	L339 F942	71.4	19.5	23.5	25.6	2.1		S429 S1177	12.2	8	15	12.2	0
	L339 T945	74.4	14.5	23.5	23.6	0.1		G430 S1177	10.7	8	15	10.1	0
6-12	F343 M986	70.7	8	15	15.9	0.9		C431 S1177	13	8	15	12.2	0
	G346 G989	63.1	8	15	16.6	1.6		G432 S1177	15.5	8	15	15	0
	P350 S993	65.2	8	15	18.1	3.1		S429 G1179	13.8	8	15	13.8	0
	L339 V982	70.6	8	15	11.1	0							
	L339 V982	69.4	13.5	23.5	11.5	2		L531 C1074	13.7	8	10	11	1
	L339 A985	67.8	14.5	23.5	14.6	0		C431 L1176	14.6	8	10	12.5	2.5
TM 2-11	Y117 G955		8	17.5	16.3	0	TM 5-8	G317 N753		8	17.5	20.8	3.3
	Y117 C956		8	17.5	16.7	0		G317 L754		8	17.5	20.6	3.1
	Y117 F957		8	17.5	15.5	0		G317 F755		8	17.5	19	1.5
	Y118 G955		8	17.5	13.9	0		T318 N753		8	17.5	21.7	4.2
	Y118 C956		8	17.5	15.1	0		T318 L754		8	17.5	20.7	3.2
	Y118 F957		8	17.5	14.3	0		T318 F755		8	17.5	17.2	0
	S119 G955		8	17.5	15.4	0		T319 N753		8	17.5	18.3	0.8
	S119 C956		8	17.5	16.9	0		T319 L754		8	17.5	17.6	0.1
	S119 F957		8	17.5	15.4	0		T319 F755		8	17.5	14	0

In each panel, the first column indicates the pair of TM helices (or NBDs) involved; the second and third columns give the residue number in each pair; the third column denotes the initial distance value measured in the initial model (before the rigid body dynamics); the fourth and fifth columns indicate the minimal and maximal values respectively of the restraints applied in the dynamics (see the Materials and Methods section); the sixth column shows the distance value in the final model and the seventh column gives the deviation either to the minimum or the maximum distance value in the restraint potentials. All distances are in Angströms (Å).

Top left panel: residues in helices 4, 5, 6, 10, 11, and 12.

Top right panel: residues in the two NBDs.

Bottom panels: residues involved in helices 2, 5, 8, and 11.

identity, respectively, with each MsbA monomer from *E. coli*. Moreover, each MsbA monomer contains six helices as in each half of the P-gp TM domains. The choice of *E. coli* MsbA, despite a better resolution of the *V. cholera* struc-

ture, was mainly guided by the fact that the sequence identity to P-gp is higher for the *E. coli* protein. Notwithstanding the influence of this template on the final model of P-gp will be strongly altered in the following steps which

will transform the initial model by accounting for structural data from P-gp.

In a second step we built an initial model of the P-gp NBD dimer. For this purpose, we resorted to comparative modeling, using two templates: the NBD dimer of *E. coli* BtuCD, the vitamin B₁₂ importer, and each NBD monomer of *E. coli* MsbA. The working hypothesis behind this step is that there is considerable support that the NBDs in P-gp form an interface in its resting state, though possibly not a tight one, as shown by cysteine disulfide crosslinking experiments realized in absence of ATP.^{21,49} In that respect, BtuCD provides a good template for the relative position and orientation of the NBDs in P-gp. Each MsbA NBD was used as a template to model the internal conformation of each NBD monomer of P-gp.

Each P-gp half, modeled in the first step, was thereafter fitted on the modeled P-gp NBD dimer. The resulting model describes a structure in which the NBDs are in close contact as expected but in which the helices of the two TM domains are as far as 50 Å at their largest separation. This feature, however, is in complete disagreement with the experimental data obtained from the cysteine disulfide cross-linking studies that demonstrate that distance values between the two TM domains should not exceed 25 Å.¹⁹ Moreover in this model some of the residues proposed to be involved in the drug binding were not oriented towards the protein interior.

To solve these discrepancies between the model and the experimental data, molecular dynamics simulations were performed on the model reduced to portions involved in the drug and nucleotide binding, namely the TM helices 4, 5, 6, 10, 11, and 12 and the two NBDs. Each of these portions was defined as a rigid body to keep their internal structure as derived from the comparative modeling stage. Those rigid bodies were allowed to move and reorient subjected to a classical force field and to additional distance-based restraint potentials. These distances were derived from cysteine crosslinking studies in which a single cysteine was introduced in each P-gp half and either an oxidant (Ref. 16, and references therein) or MTS-specific crosslinkers containing 2 to 17 atoms spacer arms^{18,19} were used. In the TM domains, 28 double mutants were found to be crosslinked, revealing distances from 8 to 23.5 Å. For the NBDs, 27 crosslinked proteins were observed from which restraints were derived, with distances from 8 to 15 Å (see Methods section for a detailed description of the derivation of the restrained distances and Table I).

This partial model was then completed by adding missing portions using experimental data whenever available. More recent crosslinking data have shown that TM helix 2 is close to TM helix 11 at both intracellular and extracellular ends, with distances value ranging from 8 to 17.5 Å.²⁷ This observation also holds true for TM helices 5 and 8. To position helix 3, we took into consideration that the extracellular loop between helices 3 and 4 is very short and thus that helix 3 on that side of the cell cannot, therefore, be far from helix 4. The same reasoning was made for helix 9 and 10. These structural data, summarized in Table I, were introduced into a second run of rigid body molecular

dynamics simulation to produce a new model. The model obtained in the previous step was defined as a rigid body, and TM helices 2, 5, 8, and 11 formed four additional rigid bodies. Finally the loops connecting the TM helices to the NBDs were added by comparative modeling using as additional templates segments from the initial model of each P-gp half.

The TM1 and TM 7 helices that are not involved in the substrate binding were not modeled because no experimental data were available. Likewise the linker region between the two P-gp halves, which consists of a segment of ~ 60 residues, was not modeled. It has been experimentally shown that its absence does not affect the protein function.⁷

The final model is shown in Figure 2. Its stereochemistry was assessed with Procheck NMR⁵⁰ and is consistent with X-ray structures determined at a resolution of 2.0 Å. All the interresidue distances involved in restraints were checked in regard to the experimentally derived data (see Table I for final distance values). The agreement is quite good as the average deviation on all distances is 1.2 Å and the maximum deviation observed is 6.5 Å. The interresidue distances values that deviate most from the restraint values belong to the TM helices. These discrepancies could in part arise from the fact that the restraints were not derived from a single experiment, but from different disulfide crosslinking involving each a particular double mutant. As a result it is possible that all experimental restraints could not be fully satisfied at the same time in one structure. An example of this involves the subsequent residues V231, W232, and A233 (on helix 4) which are experimentally observed to be at a similar distance from residue S993 (on helix 12). All these three residues belong to a helix and consequently can not simultaneously face the P-gp central cavity and S993. This suggests that possible conformational changes involving rotation of the TM helices along their axis may occur as this was shown in P-gp upon ATP hydrolysis.¹⁸

Visual inspection shows that the model presents a bundle of parallel helices, with a pseudo twofold symmetry axis in the middle of the bundle, parallel to the helices (Fig. 2). The TM domains forms a volume of roughly 65 × 56 × 40 Å, which is in very good agreement with the recent EM structure of P-gp whose resolution is 8 Å,¹² and whose TM domain dimensions are 70 × 60 × 40 Å. Some openings are observed between the helices, notably between TM helices 5 and 8 on one hand and TM helices 2 and 11 on the other hand. They both lead to a large cavity which occupies the center of the helix bundle and is open towards the extracellular side. It is important to note that the openings are not due to the absence of TM helices 1 and 7 in the model. Based on mutagenesis experiences on P-gp⁵¹ and on the crystal structure of MsbA,²⁹ those two helices are believed to be in the vicinity of TM helices 6 and 12, respectively [see Fig. 2(B)]. The existence of such lateral openings has been observed in EM studies of P-gp.⁹ It has been proposed that their role is to allow the P-gp substrates to move from the bilayer inner leaflet into the protein central cavity.⁵² The NBDs are in close interaction, and extend on a surface

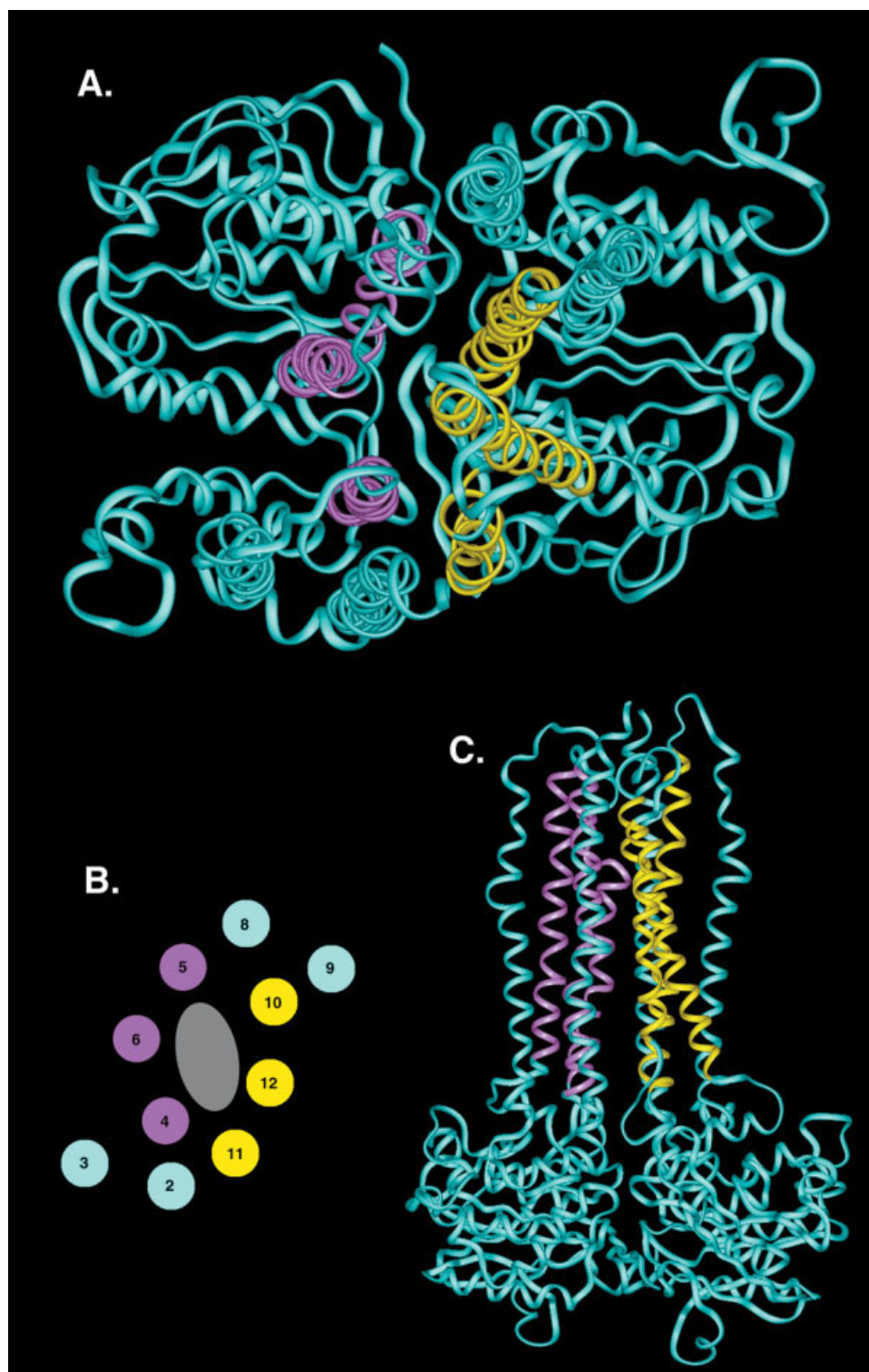


Fig. 2. Ribbon representation of the 3D model of P-gp. Helices 4, 5, and 6 are colored in pink while helices 10, 11, and 12 are colored in yellow. Those six helices contribute to the cavity wall and are involved in drug binding. (A) Top view from the extracellular side. Note the pseudo twofold symmetry axis, with the axis perpendicular to the figure. (B) Diagram showing the relative positions of the TM helices in the model. The grey oval in the middle of the helices suggests the position of the drug binding site. (C) Longitudinal view across the membrane of the bundle of helices.

TABLE II. List of the Interactions between Atoms of the Docked Ligands (partner 1) and Residues in the P-gp Model (partner 2)

Partner 1	Partner 2	Type of Interaction	Notes	
Rhodamine 1	>N ⁺ = —COO ⁻	Tyr316 aromatic moiety Ser979 —OH	Cation- π H-bond	In the vicinity of Leu975 and Val982
Rhodamine 2	>N ⁺ = Ar2 O in the cycle	Tyr953 aromatic moiety Trp232 aromatic moiety Trp232 NH	Cation- π π - π stacking H-bond	In the vicinity of Ile340, Phe343, Val981 and Val982
Rhodamine 3	>N ⁺ = Ar2 O in the cycle	Tyr953 aromatic moiety Trp232 aromatic moiety Trp232 NH sidechain	Cation- π π - π stacking H-bond	In the vicinity of Ile340, Phe343, Val981 and Val982
Verapamil 1	Ar2 —O—CH ₃ on Ar1 —O—CH ₃ on Ar2	Phe343 aromatic moiety Tyr310 —OH Ser228 —OH	π - π stacking H-bond H-bond	In the vicinity of Leu339 and Ala342

See Figure 3 for a representation of the chemical structure of the docked ligands with the numbering of their cycles. The last column specifies the residues experimentally known to be involved in the ligand binding and which are found to interact with the docked structures. For rhodamine numbers 1, 2, and 3 refer to three different positions for the docked molecule.

of roughly $74 \times 47 \text{ \AA}$, which is in excellent agreement with the $68 \times 45 \text{ \AA}$ measured on the P-gp structure obtained by EM in a lipid environment.¹¹ The interface between the TM domains and the NBDs in ABC transporters is likely to play a role in the transmission of the conformational change produced by ATP binding and (or) hydrolysis. In our model, this interface is predominantly formed by the Q-loop (residues 475–483 in NBD1 and 1018–1026 in NBD2), a protein region that has been proposed to act as a γ -phosphate sensor prone to change its conformation upon ATP binding and (or) hydrolysis.^{53,54} It is interesting to note that the Q-loop in our model is indeed located in the vicinity of the Walker A motif, where ATP binding occurs. This packing is similar to that observed in the BtuCD X-ray structure.³⁰

Other P-gp Models

An atomic level structure of P-gp has been modeled by Seigneuret and Garnier-Suillerot.²⁶ It was determined using comparative modeling using the dimer structure of *E. coli* MsbA as a sole template. As a consequence, the resulting model resembles the structure of the MsbA dimer with a large inner chamber open to the cytoplasmic side. The presence of this chamber, accessible from the lipid bilayer, is supported by an EM study.¹⁰ This is, however, the only point of concordance because the EM structure features a V form, inverted relative to this P-gp model, with a large opening to the extracellular side that is closed at the cytoplasmic face of the membrane. The authors of the model suggested that the model could be a representation of an open structure of P-gp opposed to a closed conformation. However, their modeled structure, with a distance as large as 50 \AA between the two NBDs and the cytoplasmic ends of the TM domains, is in total disagreement with the cysteine disulfide crosslinking experiments. It is only upheld by the shape of the dimeric structure of *E. coli* MsbA. One can however not completely exclude the possibility that during the catalytic cycle conformational changes lead to an open conformation as in

the MsbA structure though no structural data on P-gp has backed up this type of structure yet. While this model comprises the full P-gp sequence, it was derived based solely on MsbA structure and therefore includes no structural information from P-gp.

Another 3D atomic model was developed by Stenham and colleagues.²⁷ In their work, an initial structure of P-gp was generated using the *E. coli* MsbA dimer structure. These authors then applied global translations and rotations on each modeled P-gp half to close the structure. It has indeed been shown by Lee and colleagues¹¹ that it is only when the two MsbA monomers are rotated to bring their NBDs together that the X-ray structure of MsbA is consistent with the EM projection structure of P-gp. Energy minimization was then performed to optimize the packing of the two TM domains. The resulting model is in relatively good agreement with a number of crosslinking data. However, a few inter-TM domains residue distances were not correctly reproduced by the model and other crosslinking data involving TM residues were not checked against the model. Also no inter-NBD experimental residue distances were considered to validate their structure. Furthermore, each P-gp half structure was modeled from each MsbA monomer without any alteration. We showed that in such a model some residues involved in substrate binding are located on helix faces opposite to the drug binding cavity suggesting that MsbA monomer cannot be used as a rigid structure to model each P-gp half.

Pajeva and colleagues²⁸ presented in a recent work a partial model of the TM domain of P-gp, describing the protein in the bound-nucleotide state. Each helix was first independently built by modeling. All helices were then subjected to an energy minimization including restraints between some of the TM residues involved in crosslinking. Though no real docking was performed, this model was used to identify potential binding sites for two different ligands including rhodamine. This model was elaborated based only on structural information derived from P-gp. Furthermore, it includes only a restricted number of TM

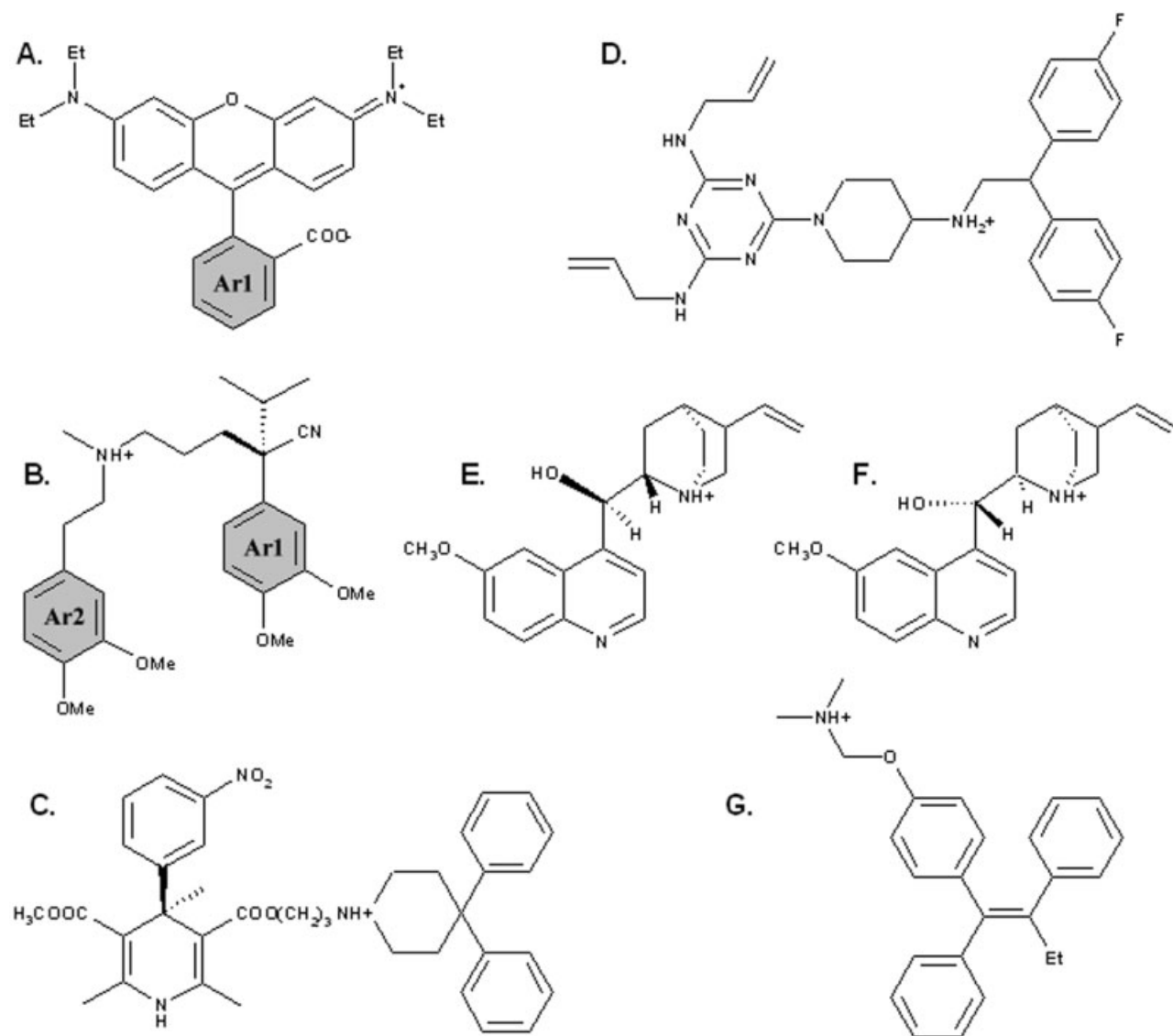


Fig. 3. Chemical structures of the different ligands. (A) Rhodamine; (B) verapamil; (C) dexniguldipine; (D) S9788; (E) quinine; (F) quinidine; and (G) tamoxifen. The Ar1 and Ar2 in rhodamine and verapamil describe specific aromatic moieties involved in interactions with P-gp model upon docking (see Table II).

helices, those directly involved in drug binding, and lacks the NBDs.

Ligand Docking into P-gp Model

In order to further assess the quality of the model, we performed the docking of a number of P-gp ligands. The scope is double. First, to verify whether not only the cavity, but also the lateral holes, are large enough to accommodate P-gp ligands, we performed the docking of two known P-gp ligands: verapamil and rhodamine (Fig. 3). Protein residues in contact with these two compounds have been identified.^{17,22} Second, we selected six inhibitors for which K_i values have been reported in a comparative study.⁴² Docking was performed for dexniguldipine, S9788, quinidine, verapamil, tamoxifen, and quinine (Fig. 3). Only first and second generation inhibitors of P-gp have been se-

lected, as third generation inhibitors are not competitive inhibitors, that is, their binding doesn't occur in the cavities of the TM domains.

For verapamil and rhodamine, 25 independent docking runs were performed in a zone surrounding the lateral opening between helices 5 and 8 on one hand, and in the central cavity on the other hand.

The position of the mass center of the molecules docked in the central cavity reveals that each ligand can occupy different positions. For verapamil, the positions of the mass centers are spread throughout the whole cavity while for rhodamine three different clusters are found, which contain 18 of the 25 mass centers. The geometry of interaction of the docked ligands with P-gp was examined in minute detail for five structures of each ligand selected on the basis of their best affinity.

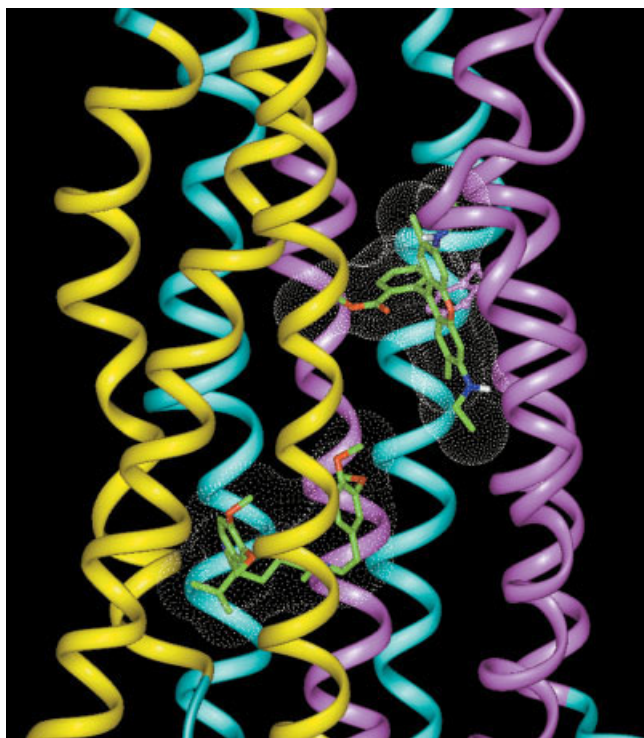


Fig. 4. Ribbon representation of the P-gp model: longitudinal view across the membrane. Two different ligands could be accommodated in the central cavity (rhodamine in the upper right; verapamil in bottom left). Helices 4, 5, and 6 are colored in pink; helices 10, 11, and 12 in yellow, and the other helices in cyan. The docked ligands are depicted as sticks and colored following their atom type (green for carbon; red for oxygen; blue for nitrogen; and white for hydrogen). Their molecular surface is also shown.

Certain residues of P-gp involved in rhodamine binding have been identified by cysteine scanning mutagenesis.²² They are located on TM helices 6, 9, 11, and 12. One of the five docked structures is located close to Leu975 and Val982 in TM12. Two other rhodamine molecules are close to residues Ile340, Phe343 (TM6), Val981, and Val982 (TM12). All these residues have been experimentally found to be involved in the binding of the drug. Table II describes the interactions between these three structures and P-gp. Each forms one H-bond and at least one π - π or one cation- π interaction. The residues involved in the verapamil binding have also been identified by cysteine scanning mutagenesis.¹⁷ They pertain to helices 4, 6, 10, 11, and 12. The inspection of the positions of the five structures docked into P-gp reveals that only one of them is close to binding site residues Leu339 and Ala342. Interactions with the P-gp model are reported in Table II.

The drug binding site cavity in the model shows that one rhodamine molecule cannot bind at the same time to all residues experimentally proposed to be involved in binding: the distances between some of them are just too large. This also holds for verapamil. Several studies have presented evidence for the existence of multiple drug interaction sites. Dey and colleagues,⁵⁵ for example, reported that iodoarylazidoprazosin has two nonidentical drug interaction sites in P-gp. In another P-gp model, Pajeva and colleagues²⁸ discussed the existence of two distinct sites of

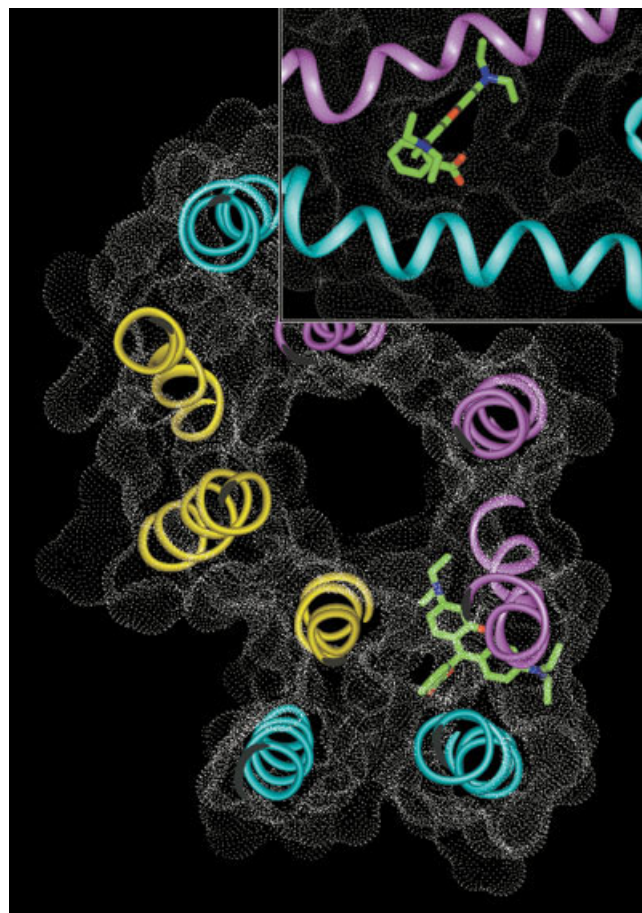


Fig. 5. Top view from the extracellular side of a cross-section of the P-gp model depicted as a ribbon and by its molecular surface. The central cavity prone to harbor ligands is located in the middle of the protein. A molecule of rhodamine has been docked between helices 5 and 8 in a lateral opening leading to the central cavity. Helices 4, 5, and 6 are colored in pink; helices 10, 11, and 12 in yellow; and the other helices in cyan. The docked rhodamine molecule is colored following its atom type (green for carbon; red for oxygen; and blue for nitrogen). Inset: Lateral view, illustrating the lateral opening between helices 5 and 8 and a docked rhodamine molecule. Note that the aromatic cycles and one of the amines are in the opening, while the other amine group lies on external surface of the protein.

binding for rhodamine, one in the vicinity of TM6 and the other in the vicinity of TM12. Furthermore, it has been shown that the binding cavity is large enough to accommodate two molecules at the same time. By resorting on drug binding measurements, Pascaud and colleagues⁵⁶ showed that while verapamil inhibits vinblastine binding, it has no effect on nifedipine binding, meaning that those two ligands have distinct specific sites of binding. By cysteine scanning experiments, Loo and colleagues²³ revealed that upon binding of a derivated rhodamine binding of verapamil is still possible. The P-gp model we present here does allow positioning these two ligands in the central cavity of our model (Fig. 4). Among the five docked verapamil structures, three are compatible with this double-ligand binding and all of them form at least two H-bonds with P-gp.

Rhodamine (see Fig. 5) and verapamil were successfully docked into the lateral opening between helices 5 and 8. More

TABLE III. Inhibition Constants Measured Experimentally⁴² and Calculated by Docking (Average over the 25 Runs)

K _i /M	Dexniguldipine	S9788	Quinidine	Verapamil	Tamoxifen	Quinine
Experiment	3.7 10 ⁻⁸	2.5 10 ⁻⁷	2.7 10 ⁻⁷	6.0 10 ⁻⁷	8.0 10 ⁻⁷	1.9 10 ⁻⁶
Docking	1.5 10 ⁻⁶	8.4 10 ⁻⁶	1.6 10 ⁻⁵	3.6 10 ⁻⁵	4.1 10 ⁻⁵	4.0 10 ⁻⁵

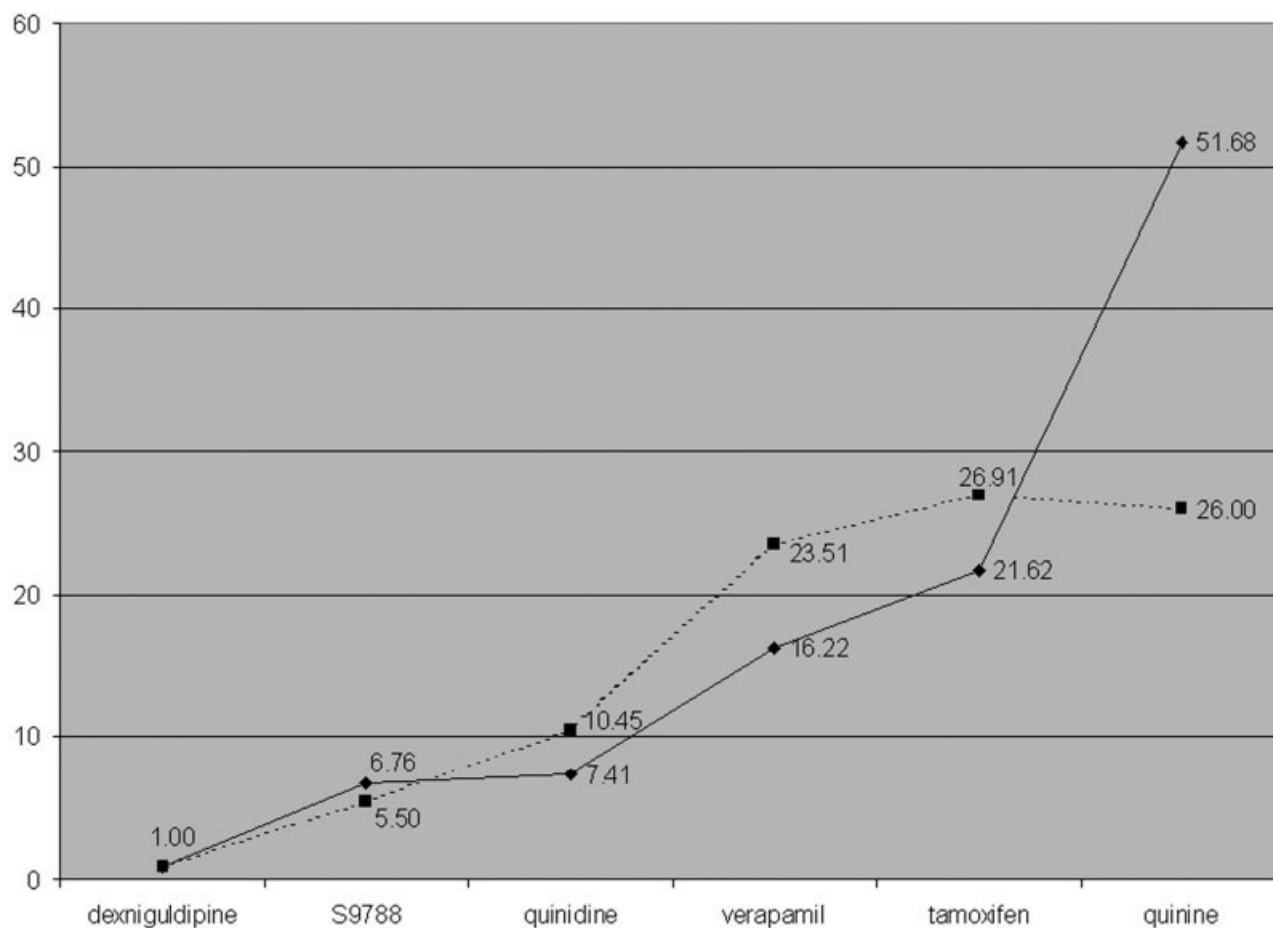


Fig. 6. Comparison of experimental and calculated affinities for known P-gp inhibitors, expressed as relative K_i. Experimental data values are shown as a solid line and diamonds; calculated docking values are shown as a dashed line and squares.

precisely for rhodamine, the two aromatic cycles (Fig. 5) are engaged in the hole, while one of its amines lies on the external surface of the protein. The capacity of the 3D model to feature a possible passage of the ligand from the bilayer into the central cavity is another important feature, because it has been proposed that, after the transfer from the cytoplasm to the inner bilayer, the drug diffuses through a lateral opening from the bilayer to the binding site.⁵²

For dexniguldipine, S9788, quinidine, verapamil, tamoxifen, and quinine, 25 independent docking runs were performed in the central cavity for each inhibitor. The average affinity for P-gp of each docked compound, expressed as a K_i, is compared to the experimental value in Table III. All calculated K_i values underestimate the inhibition. This is not surprising, as such a systematic error on the evaluation of affinity often occurs in docking experiments. A calibration of

the evaluation function on a set of known ligands can circumvent this. To do so we expressed the calculated K_i values relative to dexniguldipine, the strongest inhibitor. The results are shown in Figure 6. A clear correlation between experimental and calculated values is obtained. For example, tamoxifen is predicted to be 26.91 less active than dexniguldipine, to be compared with an experimental value of 21.62. The only exception is quinine whose predicted relative affinity is underestimated. This result is somewhat surprising as the affinity of its diastereoisomer, quinidine, is better predicted.

One can argue that performing docking on this particular 3D model is a bit perilous. However, our concern was first to verify whether the model is prone to accommodate known P-gp ligands in the lateral openings. Second, the docking of different ligands into the central cavity of the

protein was found to agree pretty well with experimental data. As such these docking results add credit to the validity of the model.

CONCLUSION

Unlike other models, the model described here was generated using a combination of various *in silico* methods including all possible structural information on both the TM domains and the NBDs derived from crosslinking studies on P-gp. This atomic level model of the protein in absence of ATP is quite complete (with 10 helices and both NBDs) and coherent because experimental data for P-gp related to this structure were carefully selected. The modeled structure is in good agreement with the structure of P-gp obtained by EM in a lipid environment. Like the EM-derived structure, our model features a closed conformation with the two NBDs in interaction. Its size and shape are also in good accordance with these EM data. In addition to these characteristics, the 3D model presented here shows the existence of an internal cavity prone to accommodate different P-gp ligands. The mode of association observed for the docked ligands favors the existence of multiple binding sites in the protein, a feature supported by several experimental studies. Predicted and experimental affinities correlate well. For each of the ligands, one or more positions were found to involve interactions with residues identified by cysteine-scanning experiments as binding these drugs. In particular, several types of interactions with these residues, such as H-bonds, π - π or cation- π interactions, were recognized and are consistent with a pharmacophore model elaborated from P-gp ligands. Furthermore, the central cavity is connected to the outside by a lateral opening which is large enough to allow the transit of the three ligands studied here. This finding corroborates the commonly admitted mechanism of substrate uptake from the lipid bilayer.

This model may, therefore, constitute a useful starting point for the understanding of the complete structural picture of P-gp along its catalytic mechanism.

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