The Transmission Interface of the *Saccharomyces cerevisiae* Multidrug Transporter Pdr5: Val-656 Located in Intracellular Loop 2 Plays a Major Role in Drug Resistance

Marianne T. Downes, Jitender Mehla, Neeti Ananthaswamy, Adina Wakschlag, Micheala Lamonde, Elliot Dine, Suresh V. Ambudkar, John Golin

Pdr5 is a major ATP-binding cassette (ABC) multidrug transporter regarded as the founding member of a fungal subfamily of clinically significant efflux pumps. When these proteins are overexpressed, they confer broad-spectrum ultraresistance. To better understand the evolution of these proteins under selective pressure, we exposed a *Saccharomyces cerevisiae* yeast strain already overexpressing Pdr5 to a lethal concentration of cycloheximide. This approach gave mutations that confer greater resistance to a subset of transport substrates. One of these mutations, V656L, is located in intracellular loop 2 (ICL2), a region predicted by structural studies with several other ABC transporters to play a critical role in the transmission interface between the ATP hydrolysis and drug transport domains. We show that this mutation increases drug resistance, possibly by altering the efficiency with which the energy from ATP hydrolysis is used for transport. Val-656 is a conserved residue, and an alanine substitution creates a nearly null phenotype for drug transport as well as reduced ATPase activity. We posit that despite its unusually small size, ICL2 is part of the transmission interface, and that alterations in this pathway can increase or decrease resistance to a broad spectrum of drugs.

The World Health Organization lists antimicrobial resistance as a major global concern that threatens advancements in medicine, national security, and economic development ([http://www.who.int/drugresistance/en/](http://www.who.int/drugresistance/en/)). Multidrug resistance remains a major clinical problem in the treatment of cancer and pathogenic infection (1). This phenomenon is attributable, in part, to the overexpression of ATP-binding cassette (ABC) efflux pumps that were initially identified and characterized in mammalian cells (2–5). Multidrug resistance was observed in *Saccharomyces cerevisiae* yeast in 1973, when it was attributed to a single-gene alteration (6), but the cloning and characterization of the efflux pumps mediating this phenomenon occurred much later (7–12).

Pdr5 is a highly promiscuous ABC transporter. Overexpression leads to hyperresistance for all known substrates (13–15). This efflux pump has a basic structure found in full-length members of this superfamily. It has two nucleotide-binding domains and two transmembrane domains (TMDs) made up of six alpha helices, each of which is connected by intra- and extracellular loops. In several significant ways, however, Pdr5 departs from the canonical architecture of ABC transporters. It has a deviant ATP-binding site that lacks an obvious catalytic residue in the Walker B motif; it also contains deviant sequences in the Walker A, signature region (C-loop) and Q-loop motifs. Furthermore, in contrast to most ABC transporters, the intracellular loops (ICLs) are very short. ICL2 and ICL4 are predicted to have only 10 and 7 residues, respectively (16). Important studies with Tap1/Tap2 antigen transporter (17) and P-gp (18) using cross-linking of cysteine-modified residues provide evidence that the X-loop, Q-loop, and intracellular loops are critical for establishing a transmission interface suggested by the atomic structure of the Sav1866 multidrug transporter (19). *In vivo* evidence for a similar interface was established in Pdr5 by using suppressor mutations (20, 21). In this transporter, a signal transmission-deficient mutation in transmembrane helix 2 (TMH2) (S558Y) was suppressed by second-site mutations in ICL1 (S597T, S597I) and the Q-loop region (N242K, E244G, D246A). These residues were implicated in intradomain signaling from the Sav1866 atomic structure.

Here we present new observations of the role of ICLs in the evolution of a highly asymmetric transporter; this investigation arose from a seemingly different issue concerning the evolution of ABC efflux pumps. Although overexpression of transporters creates hyperresistance to xenobiotic substrates, increasing the concentration beyond the 50% inhibitory concentration (IC$_{50}$) will kill even overexpressing cells. However, mutations that show even greater resistance than the original overexpressing strain can be selected. It is of great interest to understand how this occurs. We sought to determine whether such changes are limited to further increases in expression or to changes in the drug-binding pocket. We isolated mutants with increased resistance in an overexpressing strain on plates containing an otherwise lethal (5.4 μM) dose of cycloheximide (cyh). One of these mutations, V656L, is in ICL2 and confers increased resistance to Pdr5 substrates of diverse structure and function. We present evidence suggesting a novel mechanism of increased drug resistance in which a mutation in the transmission interface increases the efficiency by which the...
energy of ATP hydrolysis is used to power drug efflux. We also show that Val-656 is a conserved residue and that a V656A mutation has a phenotype characterized by markedly reduced transport and ATPase activity.

MATERIALS AND METHODS

**Chemicals.** All chemicals except 5-fluoroorotic acid (5-FOA) and G-418, which we obtained from Research Products International (Mt. Prospect, IL), aprotinin, which we purchased from Fisher Scientific (Waltham, MA), and clambazol, cyprodinil, cerulenin, cyproconazole, tebuconazole, and imazalil, which we obtained from LKT laboratories (St. Paul, MN), were purchased from Sigma-Aldrich (St. Louis, MO). All chemicals except cyh, which was dissolved in water, and 5-FOA and G-418, which were dissolved in sterilized medium, were dissolved in dimethyl sulfoxide (DMSO). [125I]iodoazidoarylprazosin (IAAP) was purchased from PerkinElmer Life Sciences (Waltham, MA).

**Yeast strains and plasmids.** All yeast strains used in this study are isogenic and are derived from R-1, a stock lacking all plasma membrane ABC transporters. It also contains an isogenic and are derived from R-1, a stock lacking all plasma membrane ABC transporters. It also contains a

**Preparation of purified plasma membrane vesicles.** Purified plasma membrane (PM) vesicles were prepared from 750 ml of exponentially growing yeast cells containing two copies of either WT or mutant PDR5. Following harvesting by centrifugation, the pellets were weighed and resuspended in homogenization buffer (50 mM Tris [pH 7.5], 2.5 mM EDTA, and a protease cocktail containing 1 mM phenylmethylsulfonylate, 10 μM aprotinin, 10 μg/ml tosyl phenylalanyl chloromethyl ketone [TPCK] and tosyllysine chloromethyl ketone [TLCK], and 1 μg/ml pepstatin and leupeptin) in a 1:2 ratio of buffer volume to pellet weight. The cells were lysed with glass beads in a bead beater (Biospec, Bartlesville, OK) using five 30-s intervals of lysing followed by 30 s of cooling as directed in the accompanying product manual. Whole cells and glass beads were pelleted at 1,000 × g for 7.5 min in an SS34 rotor at 4°C. The supernatant was spun at 100,000 × g in a Ti50 rotor for 1 h at 4°C to pellet the crude membranes. Following this, the crude membrane pellet was resuspended in 2 ml resuspension buffer (10 mM Tris, 0.5 mM EDTA, 10% glycerol with protease inhibitors) using a 7-ml Dounce homogenizer. The resuspension was applied to a discontinuous sucrose gradient made up of equal volumes of 53.5% (wt/vol) and 43.5% (wt/vol) sucrose. The supernatant was spun at 100,000 × g in an SW28 swinging bucket rotor, the purified PM vesicles were collected from the interface. The sucrose was diluted out by addition of resuspension buffer and centrifugation for 1 h at 100 × g in a Ti50 rotor. The pellets were resuspended in 2 to 3 ml of resuspension buffer, and 75-μl aliquots were frozen in 1.5-ml Eppendorf tubes at −80°C. The amount of protein in solubilized vesicles was determined with a bicinchoninic acid (BCA) protein determination kit (Thermo Scientific, Rockford, IL).

**Gel electrophoresis and Western blotting.** The presence and amount of Pdr5 in PM vesicles were confirmed by Western blotting. Proteins were separated by electrophoresis at 150 V on 7.5% Tris-acetate gels (Life Technologies, Grand Island, NY) before being transferred to a nitrocellulose membrane at constant current (400 mA, 1 h). The blots were blocked for 30 min in phosphate-buffered saline (PBS) buffer containing 1% Tween 20 and 5% nonfat milk. We obtained antibodies from Santa Cruz Biotechnology (Santa Cruz, CA). The Pdr5 antibody was diluted 1:500; the Pma1 antibody was diluted 1:2,000. Following overnight incubation at 4°C in primary antibodies, the blots were washed 3 times for 15 min in PBS plus 1% Tween 20. Following this, the blots were incubated in PBS with nonfat milk and 1% Tween containing a 1:1,000 dilution of donkey anti-goat IgG horseradish peroxidase antibody. The filters were washed 3 times for 5 min in PBS with 1% Tween 20. The chemiluminescent signal was developed by addition of 10 ml enhanced chemiluminescence (ECL) reagent (Life Technologies) for 1 min before exposure to X-ray film for 0.5 to 1 min. The ratio of Pdr5 to Pma1 signals was calculated with Image J software (rsweb.nih.gov/ij/) using scans with a resolution of 1,200 dpi.

**R6G transport assay.** Rhodamine 6G (R6G) transport assays were carried out with 5 μM R6G. Cells were grown overnight in SD medium supplemented with uracil and histidine to a concentration of ~10⁷ cells/ml. The cells were washed with 0.05 M HEPES buffer (pH 7.0) and concentrated to one-tenth of the starting volume in HEPES buffer. For each sample, 10⁷ cells were incubated in 0.05 M HEPES containing 5 μM R6G for 90 min at 30°C in a final volume of 100 μl. Following loading, the cells were spun down, the supernatant was removed, and the pellets were resuspended in 0.05 M HEPES and 1 mM glucose for 30 min. Following efflux, the cells were pelleted, the supernatant was removed, and the cells were resuspended in 400 μl HEPES for analysis by fluorescence-activated cell sorting. R6G fluorescence was measured with a Becton, Dickinson (Franklin Lakes, NJ) FACSort with an excitation wavelength of 488 nm.
and an emission wavelength of 620 nm. The data were analyzed with a CellQuest program. Retained fluorescence is expressed in arbitrary units.

**ATPase assays.** We assayed the Pdr5-specific ATPase activity with purified PM vesicles in an ATPase assay buffer containing 100 mM MOPS (morpholinepropanesulfonic acid) (pH 7.4), 50 mM KCl, 5 mM NaN₃, 2 mM EGTA (pH 7.0), 2 mM dithiothreitol (DTT), and 10 mM MgCl₂ at 35°C for 8 min as previously described (22). The V656L mutant increases resistance to cyh. We constructed V656L in pSS607 and placed it in R-1 (Δpdr5 strain) by yeast transformation. Following this, the original chromosomal mutation and the plasmid reconstruction were compared to WT and Δpdr5 isogenic controls for their relative resistances to cyh in liquid culture as described in Materials and Methods. All plots in this study were made using GraphPad Prism (San Diego, CA), n = 3. A, Δpdr5; ●, WT; ○, chromosomal V656L mutation; ◊, plasmid-reconstructed V656L mutation.

**RESULTS**

The V656L mutation appeared in two distinct genetic screens. We recovered the V656L mutation in two separate genetic screens, the first of which is illustrated in Fig. S1 in the supplemental material. We suppressed S558Y drug hypersensitivity by a series of second-site mutations, including several near or in the Q-loop, a region required for intradomain communication (20, 21). One of these mutants was N242K. When N242K is not coupled to S558Y, the former cannot be attributed to an increased level of the transmembrane (TM) residue to cyh, although it was not extensively characterized (21).

We carried out the second screen for our present study, exposing cells overexpressing WT Pdr5 (JG2015) to cyh at the MIC for the WT strain on solid medium (5.4 μM). The selected mutations were dominant. Therefore, standard complementation testing was not useful for determining whether these lie in PDR5. This was necessary because we expected to observe hyperresistant mutations both in and outside Pdr5. The genetic approach for distinguishing these possibilities in the JG2015 strain was recently described (20, 21). We identified five putative mutants in PDR5, and we estimate that they arose with a frequency of ~10⁻⁸.

We recovered the PDR5 gene from these mutants by PCR amplification and sequenced the DNA (Retrogen, San Diego, CA). Each mutant had a single-base-pair substitution resulting in a missense mutation. One of these was V656L. The other mutations were P596L in ICL1, L1367F in TMH12, A670S in TMH5, and I1063M in nucleotide-binding domain 2 (NBD2). In this study, we focus on Val-656 because of its location in a region strongly implicated in signal transmission by structural and cross-linking studies. We therefore recreated the V656L mutation in pSS607 and placed it into the Δpdr5 strain by yeast transformation. We compared these strains to the original chromosomal mutants. The data in Fig. 1 show that the cyh killing curves for the plasmid and chromosomal mutations are statistically indistinguishable. A two-way analysis of variance (ANOVA) comparing each of the mutant curves to the WT curve indicated that the likelihood that they are the same is <0.0001. We used the plasmid reconstruction in all of the subsequent experiments. Figure 2 shows a representative Western blot for the mutant strains used in this study. The V656L mutant protein is found at WT levels in purified plasma membrane (PM) vesicles. We determined the ratio of Pdr5 to Pma1 signals for all of the strains. This value allowed us to make a quantitative comparison between the WT and the mutants analyzed. The amount of Pdr5 protein in the V656L mutant was 0.89 ± 0.13 times that of the WT. Therefore, the increased resistance of the former cannot be attributed to an increased level of the transporter in the PM.

**Val-656 is one of four conserved residues in ICL2.** According to the bioinformatic analysis performed by Rutledge et al. (16) to build their atomic model of Pdr5, ICL2 is remarkable for its relatively short size. It is predicted to contain only 10 amino acids (654 to 663); in contrast, ICL1 is made up of 21 residues. They also observed two loops with analogous sizes in the second half of Pdr5.

We performed a sequence alignment using the Swiss Protein Database alignment tool (www.uniprot.org). We compared Pdr5 to 17 members of the Pdr5 subfamily, with amino acid identities ranging from 30.7% to 74.4%. These sequences were initially selected for use by Rutledge et al. (16) as part of an atomic modeling study. The alignment that we obtained is found in Fig. S2 in the supplemental material. The first and last amino acids in the loop are a highly conserved arginine and either a serine or threonine residue, respectively. The third position has a medium-to-large hydrophobic residue (V, I, L, M, F) followed by a small nonpolar
one (G or A) in the fourth. All of the residues located at the third position can be interchanged as a result of a single-nucleotide substitution. ICL4 is located in TMD2 and is predicted to comprise only seven amino acids (22), suggesting a correspondence to ICL2 because of its size similarity. In contrast to ICL2, however, none of the residues in ICL4 are conserved. This indicates that ICL2 and ICL4 may be functionally distinct. A recent report clearly demonstrates that ICL4 is necessary for drug resistance, although its role is unknown (23).

Val-656 is an essential residue for Pdr5-mediated drug transport. To determine whether Val-656 is essential for Pdr5 function, we used site-directed mutagenesis to create a V656A mutation and introduced it into the pdr5 strain. Ordinarily, a valine-to-alanine change is structurally neutral and a single-base-pair substitution can interchange these residues. However, no alanines are found in position 3 of ICL2 in the alignment sequences. This strongly suggested that such a substitution would be unfavorable, as indeed it is. The level of the V656A mutant protein in purified PM vesicles was, if anything, slightly higher (1.3 ± 0.1 times) than the level of the WT (Fig. 2). The relative cyh, clo, bifonazole, tamoxifen, and climbazole resistances of the V656A mutant were compared to those for the isogenic WT (JG2015) and pdr5 strains (Fig. 3A to E). The V656A and pdr5 mutations were equally hypersensitive to cyh, tamoxifen, bifonazole, and climbazole. Although the V656A mutant showed significantly more resistance to clo than did pdr5, it was considerably more sensitive than the WT (Fig. 3A). We also tested the ability of the V656A mutant to efflux R6G in a whole-cell assay. These data demonstrate that the V656A mutant and the pdr5 control were equally deficient in R6G efflux (Fig. 4).

ATPase activity is significantly reduced in the V656A mutant. A hallmark of loss-of-function mutations in the signal transmission interface of Pdr5 analyzed thus far is a reduction—but not an elimination—of ATPase activity. This feature does not fully reflect the severity of the mutation. For instance, although the E244G (deviant Q-loop) and S558Y mutants both showed

![FIG 3](image1)

![FIG 4](image2)
ICL2 of Pdr5 Modulates Multidrug Resistance

ATPase activity that is 3 to 4 times lower than that of the WT (mutants, ~50 nmol/min/mg; WT, ~200 nmol/min/mg), the E244G mutant retained significant transport capability and the S558Y mutant was phenotypically null (20, 21).

We performed similar experiments with the V656A mutant (Fig. 5A). Purified plasma membrane vesicles exhibited a reduced but significant level of ATPase activity. The $V_{\text{max}}$ obtained for three independent membrane preparations was 42 nmol/min/mg. This value is similar to those observed with the transmission interfaces mutants S558Y, E244G, and Q951G. The $K_{m}$ values were 1.6 mM for the WT and 0.8 mM for the V656A mutant, values within the range seen routinely in our experiments (for example, the WT data in Fig. 9C indicate a $K_{m}$ of 0.9 mM).

V656A mutant ATPase is significantly more resistant to clo than the WT enzyme. Although Pdr5 ATPase is not stimulated by drug substrates, some, such as R6G, clo, and ketoconazole, are inhibitors (22, 24). The residual ATPase activity found in the S558Y mutant is considerably more resistant to allosteric inhibition by clo than is the ATPase activity found in the WT (20). This strongly suggested that the inhibitory signal is not being properly transmitted in the mutant. To determine whether ICL2 is required for allosteric inhibition of ATPase, we tested the residual V656A mutant ATPase activity for its susceptibility to clo. The results show that the mutant ATPase exhibits significantly greater clo resistance than does the WT protein (Fig. 5B). The $I_{C_{50}}$ (clo) observed for the WT in this particular set of experiments was 2.2 \(\mu\)M, which is similar to the WT values obtained in the experiments described later in Fig. 9. In contrast, the mutant enzyme had a value of 9.9 \(\mu\)M. A two-way ANOVA indicated that the probability that the two curves are the same was less than 0.0001. These results strongly suggest that both transport and ATPase inhibition are highly dependent on Val-656.

V656A mutant drug binding is similar to that for WT Pdr5. The crystal structure of mouse P-gp showed that binding of its drug substrates takes place in the TMDs (25). It may be that the ICLs play an indirect role in drug binding and that the V656A mutant is deficient in this process. To find out, we compared the abilities of the WT and the V656A mutant to bind IAAP in the presence of the competing substrate clo (Fig. 6). Previous work in our laboratory demonstrated that both R6G and clo show a concentration-dependent inhibition of IAAP photoaffinity labeling (20, 26). This is significant because R6G is a competitive inhibitor of clo efflux (see Fig. S3 in the supplemental material). Substrates such as tripropyltin chloride or tritylimidazole that do not block R6G or clo efflux do not inhibit IAAP labeling (26). In addition, Pdr5 mediates IAAP efflux in a whole-cell transport assay (Z. Sauna and J. Golin, unpublished data). In the present experiments, we found that IAAP binds to the WT Pdr5 and the V656A mutant and yields a signal of similar intensity. Furthermore, the addition of clo yielded inhibition curves (percent IAAP incorpo-

FIG 5 ATPase activity of the V656A mutant. ATPase was measured as described in the experimental procedures using 16 \(\mu\)g of PM vesicle protein from double-copy strains (see reference 23). Reactions with a final volume of 100 \(\mu\)l were initiated by addition of ATP and incubated at 35°C for 8 min before being terminated with 2.5% (final volume) SDS. The nonspecific activity (<5%) present in the \(\Delta\)pdr5 strain is subtracted prior to calculating the activity. (A) Activity as a function of ATP concentration. Data represent the averages of at least two independent vesicle preparations per strain, and 3 independent experiments were performed. (B) Activity in the presence of clotrimazole (3.1 to 50 \(\mu\)M). Standard ATPase conditions were used. PM vesicles were incubated at room temperature for 5 min in the presence of clo before initiation of the reactions with the addition of ATP to 3 mM. ■, WT; ▲, V656A mutant. $n = 3$. 

FIG 6 The V656A mutant exhibits WT drug-binding behavior. We carried out photoaffinity labeling with IAAP in the presence and absence of clo as described in Materials and Methods. (A) Each lane contains 10.8 \(\mu\)g of solubilized PM vesicle protein. After electrophoresis, the gels were dried and exposed to an X-ray film. The radioactivity incorporated into the pdr5 band was quantified using a phosphorimagery and analyzed using GraphPad Prism. The autoradiogram described above shows IAAP binding to the WT pdr5 and the mutant pdr5 in the presence and absence of clo. The isogenic strain lacking Pdr5 (\(\Delta\)pdr5) was used as a control in the experiment. WT and V656A mutant strains were analyzed in the same experiments. (B) Plots of the experiment shown in panel A. ■, WT; ▲, V656A mutant. We performed this experiment an additional time with nearly identical results.
ration at a specific dose) that were indistinguishable from each other. In each case, the IC_{50} was about 15 μM. Thus, it is unlikely that V656A alters the transporter-drug substrate-binding interaction to a degree that would yield the observed null phenotype.

Response of the V656L mutant to Pdr5 substrates of varied strength. The Western blot in Fig. 2 demonstrates that the increased resistance of the V656L mutant to cyh is not attributable to a further increase in the expression of Pdr5. We considered several models to account for the observation that V656L increases resistance to cyh. The first, known as the kinetic selection model, was proposed by Ernst et al. (24) to explain the behavior of an H-loop mutation in Pdr5 that reduced R6G transport but exhibited WT resistance to other transport substrates. Pdr5 has a high basal ATPase activity that is not stimulated by drugs. Thus, hydrolysis may well be constitutive and independent of drug substrate stimulation. This model posits that mutations can alter the rate of switching from a drug-binding to a drug-effluxing conformation. While precise measurements of on- and off-rates of drugs would be required to rigorously establish or reject the model, this leads to a specific phenotypic prediction. If the Pdr5 pump is altered so that relative to the WT protein the mutant is more frequently in the drug-binding (according to the Sav1866 model, the inward-facing conformation) rather than the drug-effluxing state (the outward-facing model), modest or poor Pdr5 substrates that normally interact slowly will have a greater opportunity for binding.

The first model makes a specific prediction about substrate behavior: the V656L mutant exerts its strongest effect on substrates that interact relatively poorly with Pdr5. We therefore tested the response of the V656L mutant to a set of 10 transport substrates (Table 1). We determined the IC_{50}s for the V656L, WT, and pdr5 strains. The V656L mutant conferred greater resistance to eight compounds. These did not cluster according to the IC_{50} index but were evenly distributed. Furthermore, we observed a similar increase in resistance for each one. Thus, the V656L mutant is 1.4 to 2.1 times more resistant to the compounds than the WT. These results cast doubt on the kinetic selection model as an explanation for the observed ultraresistance of the V656L mutant.

### Table 1: Enhancement of drug resistance in the V656L mutant is independent of substrate strength

<table>
<thead>
<tr>
<th>Compound</th>
<th>IC_{50} (μM)</th>
<th>IC_{50} ratio</th>
<th>V656L mutant/ WT IC_{50} ratio</th>
<th>ATPase IC_{50} (μM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tamoxifen</td>
<td>1.7</td>
<td>13/18</td>
<td>1.4</td>
<td>Complete</td>
</tr>
<tr>
<td>Cyprodinil</td>
<td>2.0</td>
<td>140/140</td>
<td>1.0</td>
<td></td>
</tr>
<tr>
<td>Celmidin</td>
<td>1.0</td>
<td>14/28</td>
<td>2.0</td>
<td>No inhibition</td>
</tr>
<tr>
<td>Imazalil</td>
<td>10</td>
<td>4.3/7.7</td>
<td>1.8</td>
<td>Complete</td>
</tr>
<tr>
<td>Cycloheximide</td>
<td>17</td>
<td>7.5/16</td>
<td>2.1</td>
<td>No inhibition</td>
</tr>
<tr>
<td>Bifonazole</td>
<td>19</td>
<td>3.7/5.7</td>
<td>1.5</td>
<td>5.3</td>
</tr>
<tr>
<td>Clotrimazole</td>
<td>40</td>
<td>14/28</td>
<td>2.0</td>
<td>No inhibition</td>
</tr>
<tr>
<td>Tebuconazole</td>
<td>80</td>
<td>6.2/9.5</td>
<td>1.5</td>
<td>2.0</td>
</tr>
<tr>
<td>Cyproconazole</td>
<td>106</td>
<td>7.5/15</td>
<td>2.0</td>
<td>No inhibition</td>
</tr>
</tbody>
</table>

![Figure 7: Resistance of the V656L mutant to Pdr5 transport substrates.](http://aac.asm.org/downloaded.png)

**FIG 7** Resistance of the V656L mutant to Pdr5 transport substrates. We show plots of growth versus substrate concentration for six of the 10 compounds tested. Conditions for testing substrates in liquid culture are found in Materials and Methods. These vary in their substrate strength. (A to C) Data from substrates ranging from weak (tamoxifen) to strong (cyproconazole), to which the V656L mutant (purple plots) confers increased resistance. Strength is determined by comparing the IC_{50} of the WT (green) and Δpdr5 mutant (red) plots. (D to F) Data from substrates ranging from weak (cyprodinil) to strong (clo) that cause significant (>50%) inhibition of ATPase activity (see Fig. 9). All concentrations are in μM units. n = 3.
although it remains an important possibility for the H-loop mutant described by Ernst et al. (24).

We then considered a second and a third explanation for the behavior of the V656L mutant. The second asserts that Val-656 affects the level of Pdr5 ATPase. Increased resistance is due to an increased ATPase activity available for transport. A third model is that these mutants alter the transmission interface, causing a more efficient use of energy from rather than an increase in ATP hydrolysis.

Both of these models predict that if a transport substrate such as clo eliminates this activity before or at the WT IC50 for growth, the V656L mutant will not increase resistance. We therefore looked at the effect of substrates on ATPase activity.

**Effect of Pdr5 transport substrates on ATPase activity.** We tested the effect of the same set of compounds on the WT Pdr5 ATPase. Increased resistance is due to an increased ATPase activity available for transport. A third model is that these mutants alter the transmission interface, causing a more efficient use of energy from rather than an increase in ATP hydrolysis.

Both of these models predict that if a transport substrate such as clo eliminates this activity before or at the WT IC50 for growth, the V656L mutant will not increase resistance. We therefore looked at the effect of substrates on ATPase activity.

**Effect of Pdr5 transport substrates on ATPase activity.** We tested the effect of the same set of compounds on the WT Pdr5-mediated ATPase activity (Fig. 8A). Three of these, clo, bifonazole, and cyprodinil, caused strong (>50%) inhibition of WT ATPase activity. Although bifonazole and clo are similar in structure, cyprodinil is much smaller and it is not an imidazole derivative. The V656L mutant did not show an increase in resistance to cyprodinil or clotrimazole. The remaining transport substrates had a weak (≤30% inhibition) effect on ATPase activity or none at all. It would appear, then, that strong substrate inhibition of ATPase activity is seen with only a minority of the transport substrates. In some instances, an extremely weak stimulation of ATPase occurred at very high transport substrate concentrations. The V656L mutant showed increased resistance to all of the substrates that leave ATPase activity largely unaffected. However, it also exhibited increased resistance to bifonazole, which shows substantial inhibition of ATPase activity. Though this was initially unexpected, we observed that the IC50 for killing in WT whole cells is 3.7 μM while that of the V656L mutant (5.7 μM) is very close to the IC50 for ATPase inhibition by this drug (5.3 μM). This suggests that as long as the IC50 for allosteric inhibition is above the IC50 for killing of WT cells, the V656L mutation can promote increased resistance. An exact correlation between inhibition in vesicles and killing in whole cells cannot be made, however, without determining the effective intracellular concentration of the drug in question. Our observations are thus consistent with the second and third models, which invoke either an increase in or a more efficient utilization of ATPase activity. We next attempted to distinguish these possibilities.

**ATPase activity of the V656L gain-of-function mutation.** Suppressor mutations located very near (N242K) or in a Pdr5-specific residue defining the Q-loop that were isolated for their ability to restore resistance to the transmission interface-deficient mutant S558Y do not increase ATPase levels. Rather, they apparently improve the efficiency with which the signal from ATP hydrolysis is transmitted, and they restore noncompetitive inhibition of the Pdr5 ATPase by clo. Presumably, if the V656L mutant were to increase drug resistance without a concomitant increase in ATPase activity, we would have additional evidence that Val-656...
and, therefore, ICL2 play a role in transmitting the signal from the nucleotide-binding domains to the TMDs. We compared the ATPase activity of the V656L mutant to that of WT Pdr5 in purified plasma membranes (Fig. 8B). The $V_{\text{max}}$ of the V656L mutation (~200 nmol/min/mg) fell well within the range of WT values (216 nmol/min/mg in these experiments) but was certainly not greater. The $K_m$ values for the WT and mutant were 1.6 mM and 1.2 mM, respectively. The increased resistance that we observed is therefore not attributable to greater ATPase activity.

The ATPase activity of the V656L mutant is slightly more sensitive to clo than is that of the WT (Fig. 8C). According to a two-way ANOVA, the two curves are probably not equivalent ($P = 0.007$); however, the IC$_{50}$s are nearly the same. Any effect of V656L is therefore very small. The effects of climbazole, tamoxifen, and cyproconazole on V656L mutant ATPase activity were also tested. Climbazole and cyproconazole had identical effects on V656L mutant and WT ATPase activity (see Fig. S4 in the supplemental material). The V656L mutant ATPase activity, however, was significantly more sensitive to tamoxifen than was that of the WT (Fig. 8D). The IC$_{50}$ was about 32 $\mu$M, and the inhibition leveled out to about 70%. In V656L mutant cells, the IC$_{50}$ (tamoxifen) was about 18 $\mu$M (Table 1), a value not terribly different considering that the comparison is between measurements in PM vesicles and whole cells, which may not be equivalent. This observation may explain why the enhancement in resistance observed in the V656L mutant compared to the WT (~1.4×) was less than that obtained with some of the other compounds.

The V656L and V656A mutations therefore create opposite phenotypes that are consistent with altered signal transmission. The V656L mutation causes increased resistance to many drugs but increased sensitivity to allostery, at least in the case of tamoxifen. In contrast, the V656A mutant is markedly drug hypersensitive but more resistant to allosteric inhibition by clo.

The V656L mutant also suppresses E244G, a mutation in the residue defining the pdr-specific Q-loop. These data are consistent with the idea that Val-656 is part of the transmission interface. To test this idea further, we created an E244G, V656L double mutant to detect increased drug resistance and/or ATPase activity. E244G is the deviant Q-loop residue that is completely conserved in the Pdr5 subfamily. Recent work in our laboratory established its role in the transmission interface (20). When E244G is present in an otherwise WT strain, the result is reduced ATPase activity and increased drug sensitivity.

The Western blot in Fig. 9A shows that Pdr5 is present at similar levels in the PM vesicles of each strain. The data in Fig. 9B clearly demonstrate that the V656L mutant suppresses the significant cyh hypersensitivity of the E244G transmission interface mutant. In fact, the resistance observed in the suppressor strain is similar to that of the WT. The resistance of the double mutant is about four times that observed with the E244G mutant. At issue, however, is whether the suppression is attributable solely to better use of the residual ATPase activity in the E244G mutant or to an actual increase in the level of hydrolysis (Fig. 9C). Significantly, the ATPase activity of the double mutant was no different than the activity observed for the E244G mutant, which was in the range of 60 to 75 nmol/min/mg.

**DISCUSSION**

Multidrug resistance is often caused by overexpression of ABC efflux pumps, and it is a major problem in the clinical treatment of cancers and infection. Furthermore, Pdr5 defines an ABC subfamily found only in fungi; many of these are of agricultural significance. It is important to determine whether a further increase in resistance can be obtained under continued selective pressure in strains that are already overexpressing multidrug transporters. We found that mutations in the Pdr5 gene imparting resistance to cyh are recovered with a frequency of roughly $10^{-7}$. The V656L mutation leads to an increase in resistance to a broad spectrum of Pdr5 substrates that is 1.5 to 2 times that of the WT. The enhancement is not dependent on substrate strength. Although the increased resistance may seem modest, numerous chemotherapeutic and antifungal agents are toxic if the dose is doubled. Microevolution often works incrementally but effectively.

V656L is located in the very short ICL2 of TMD1. Thus, this residue is not in the drug-binding pocket, as might be expected, and this alteration does not result in an increased steady-state expression level of Pdr5 in purified PM vesicles. Rather, Val-656 is one of four conserved residues in ICL2. An alanine substitution has an effect that is entirely different from that of V656L. The V656A mutation is nearly phenotypically null to all Pdr5 substrates tested, although its drug-binding capability is intact. It is plausible that although valine and alanine are biochemically similar, the larger size of the former is required for interaction between ICL2 and the NBD(s). In fact, the behavior of the V656A mutant resembles that of the transmission-deficient mutant S558Y, which lies in TMH2. As was the case with the S558Y mutant, the noncompetitive inhibition of Pdr5-specific ATPase by clo is significantly diminished in the V656A mutant. These data show that the signals for drug transport and allosteric ATPase inhibition both use ICL2. V656L’s suppression of the E244G and
N242K signaling deficiencies in the Q-loop region also strongly implies a role for Val-656 and ICL2 in the transmission interface. This is further supported by the observation that although the V656L mutant increases resistance in either a WT or E244G mutant background, it does so without raising ATPase activity. These observations indicate that in the WT transporter, the energy from hydrolysis is not used to full capacity. From the standpoint of evolution, mutations that improve the efficiency of energy utilization should have a strong selective advantage in the presence of a xenobiotic agent. Thus, multidrug transporters can evolve to greater resistance by manipulating signal transmission. This is a novel mechanism of multidrug resistance not previously described. In the case of the V656L mutant, this results in additional resistance to a broad spectrum of Pdr5 substrates that appears limited only in cases in which ATPase activity is reduced by allosteric inhibition. The behavior of mutants in this study further emphasizes that it is impossible to link ATPase activity levels with the resistance phenotype in Pdr5.

The high-resolution structure of the Sav1866 homodimer shows physical contact between ICL2 and the X-loop and Q-loop residues of the same monomer. Cross-linking studies with other eukaryotic transporters also lend additional credence to a transmission interface arrangement in a trans orientation (17, 18). Sav1866 and P-gp were used heavily by Rutledge et al. (22) to construct a homology model of Pdr5. As a result, the Pdr5 interface is modeled in a trans conformation with ICL2 in close proximity to the Q-loop of NBD2. This model was extremely challenging to build, however, because there are big structural differences and relatively low amino acid identity between Pdr5 and these transporters. In Table S1 in the supplemental material, we list the residues of the Pdr5 transmission interface as defined by suppressor mutations. ICL1, ICL2, and the Q-loop region are all represented in the collection. All of these lie in the same half of the transporter. Our original collection of suppressors restored resistance to a transmission-deficient SS58Y mutant. Ser-558 is located at the extracellular end of TMH2, which is admittedly a good distance from the NBDs, so the interaction between these residues may be very indirect. The V656L and E244G residues, however, are presumably in much closer proximity to each other, and their observed interaction strengthens our contention that unlike other eukaryotic ABC transporters, ICL2 of Pdr5 (represented by Val-656) interacts with the cis Q-loop. We do not mean to imply that Val-656 and Glu-244 are necessarily a physically interacting pair in the WT, as their chemistry makes that implausible. It is possible that V656L and E244G create a new hydrophobic interaction that restores the signal to the interface. Alternatively, these mutant residues may never be in physical contact although they clearly are part of the same transmission pathway.

Although the E244G (Q-loop NBD1), Q951G (Q-loop NBD2), and V656A substitutions are deficient in intradomain communication, the Q-loop deficiencies show some drug specificity. For instance, the E244G mutant exhibits almost WT clo resistance (20). We also observed that the E244G mutant has R6G transport capability that is indistinguishable from that of the WT (J. Mehla, unpublished data). In contrast, an E244G, Q951G double mutant exhibits strong hypersensitivity to clo and cyh (20). Therefore, the Q-loop residues are overlapping in function. The broad, striking hypersensitivity of the V656A mutant and the absence of an obvious consensus sequence in ICL4 or ICL3 suggest the possibility that there is no equivalent residue in TMD2.

ACKNOWLEDGMENTS

This work was supported by PHS grant GM07721 and NSF grant 1048838 to J.G. and by the Intramural Research Program of the NIH, National Cancer Institute, Center for Cancer Research.

This article is dedicated to the memory of Seth Aryeh Golin Weisman (1981 to 2012), who isolated the first mutants with increased resistance in the summer of 2003 and continued to insist on the importance of their characterization.

REFERENCES


