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Comparative chemical genomics reveal that the spiroindolone antimalarial KAE609 (Cipargamin) is a P-type ATPase inhibitor

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The spiroindolones, a new class of antimalarial medicines discovered in a cellular screen, are rendered less active by mutations in a parasite P-type ATPase, *PfATP4*. We show here that *S. cerevisiae* also acquires mutations in a gene encoding a P-type ATPase (*ScPMA1*) after exposure to spiroindolones and that these mutations are sufficient for resistance. KAE609 resistance mutations in *ScPMA1* do not confer resistance to unrelated antimicrobials, but do confer cross sensitivity to the alkyl-lysophospholipid edelfosine, which is known to displace ScPma1p from the plasma membrane. Using an *in vitro* cell-free assay, we demonstrate that KAE609 directly inhibits ScPma1p ATPase activity. KAE609 also increases cytoplasmic hydrogen ion concentrations in yeast cells. Computer docking into a ScPma1p homology model identifies a binding mode that supports genetic resistance determinants and *in vitro* experimental structure-activity relationships in both *P. falciparum* and *S. cerevisiae*. This model also suggests a shared binding site with the dihydroisoquinolones antimalarials. Our data support a model in which KAE609 exerts its antimalarial activity by directly interfering with P-type ATPase activity.

The spiroindolones, a novel class of orally bioavailable antimalarials discovered in a phenotypic whole-cell screen, have been shown in Phase II clinical trials to rapidly clear parasites from adults with uncomplicated *P. vivax* and *P. falciparum* malaria¹. KAE609 (cipargamin or NITD609), a representative compound, works twice as fast in patients as the current gold standard treatment, artemisinin. KAE609 possesses both the potency (average IC₅₀ of 550 pM against asexual blood-stage *P. falciparum*)² and favorable pharmacokinetics (elimination half-life of ~24 hours in humans)³ needed for a single-dose cure, a feature that could help slow the onset of parasite resistance and that is not shared by existing, approved antimalarial drugs. KAE609 is also unique in its ability to block transmission to mosquitoes⁴.

Despite promising activity in both cellular and organismal assays, the spiroindolones act by a relatively uncharacterized mechanism. Directed-evolution experiments in parasites have shown that resistance is conferred by mutations in the gene encoding the parasite plasma membrane P-type ATPase, *PfAtp4p*. Biophysical studies have shown that parasites treated with KAE609 are not only unable to extrude intracellular sodium, but also

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exhibit changes in intracellular pH⁵. On the other hand, *PfATP4* mutations also confer resistance to a variety of unrelated chemical scaffolds with antimalarial activity^{6–9}, suggesting that *PfATP4* may be a multidrug resistance gene instead of the true spiroindolone target.

Although direct work on *PfAtp4p* would be desirable, our attempts to study recombinant *PfAtp4p* have not been successful (our unpublished data) nor is a structure available. To better understand the mechanism of action of spiroindolones, we used directed evolution experiments in baker's yeast (*Saccharomyces cerevisiae*), a more genetically tractable model system.

Results

KAE609 inhibits *S. cerevisiae* growth. To determine whether yeast could be used to study the function of KAE609, we tested the compound in a cellular, phenotypic assay. Using yeast proliferation as a readout (OD600), we found the half maximal inhibitory concentration (IC₅₀) of KAE609 against a wild-type strain (SY025) to be prohibitively high for drug-selection studies (IC₅₀ = 89.4 ± 18.1 μM, 9 observations). Reasoning that the yeast cells might be expelling KAE609 via drug efflux pumps, we next tested a strain that lacks 16 genes encoding ATP-binding cassette (ABC) transporters, termed “ABC₁₆-Monster”¹⁰. As predicted, KAE609 was more potent against ABC₁₆-Monster (IC₅₀ = 6.09 ± 0.74 μM), suggesting that this yeast strain could be a useful surrogate for malaria parasites.

KAE609 resistance is conferred by mutations in *ScPMA1*, an ortholog of *PfATP4*. We next sought the KAE609 target in *S. cerevisiae* using the same *in vitro* evolution and whole-genome scanning method that previously identified *PfATP4* as a KAE609 resistance gene². ABC₁₆-Monster cells were exposed to increasing KAE609 concentrations in three clonal cultures. In all three cultures, compound resistance emerged after two rounds of selection, with new IC₅₀ values of 20.4 ± 2.2, 29.1 ± 2.6, and 26.4 ± 4.6 μM, respectively. After an additional three rounds of selection, two of the cultures developed additional resistance (40.5 ± 4.7 and 61.5 ± 7.1 μM) (Fig. 1a). To determine the genetic basis of this *in vitro* resistance, we prepared genomic DNA from clonal strains of the terminal selection. Samples were fragmented, labeled, and sequenced with >40-fold coverage (Supplemental Table 1). The sequences were then compared to the sequence of the parental clone.

Sequencing revealed 5–8 single nucleotide variants (SNVs) in each line and no additional copy number variants (CNVs) beyond the 16 ABC₁₆-transporter deletions and selection-marker insertions characteristic of the strain. Among the SNVs, there were 2–3 missense mutations in protein-coding genes per clone (Table 1). The transcription factor *ScYRR1* was mutated in two lineages. *ScPMA1* was the only gene mutated in all three clones. *ScPMA1*, a gene that encodes a P-type ATPase responsible for maintaining hydrogen-ion homeostasis across the plasma membrane in yeast¹¹, was the only essential gene among those identified. It is also a homolog of a KAE609 resistance-conferring gene in *P. falciparum*, *PfATP4* (Fig. 1b). The identified *ScPMA1* mutations (Pro339Thr, Leu290Ser, and Gly294Ser) are clustered in the E1-E2 ATPase domain, in a region that is homologous to *PfATP4*. These mutated amino acids are positioned near or at the same homologous residues that confer parasite resistance to both the spiroindolones and the dihydroisoquinolones, another compound class predicted to inhibit *PfAtp4p*⁷. Sanger sequencing of *ScPMA1* and *ScYRR1* at each round of selection was used to determine when each mutation arose in its respective lineage. This same sequencing also identified an additional clone in Lineage 2 with its own distinct *ScPMA1* mutation (Asn291Lys). Mutations in *ScPMA1* and *ScYRR1* each correlate with increased KAE609 resistance (Fig. 1a).

***ScPMA1* alleles are sufficient to confer resistance to KAE609.** To further investigate the contribution of different alleles to the resistance phenotypes, we determined whether the mutations we found were specific to the spiroindolones. We performed 103 additional directed-evolution experiments in ABC₁₆-Monster against 26 different compounds with blood-stage *P. falciparum* activity. None of the 103 genomes sequenced had *ScPMA1* mutations. However, 22 clones resistant to six unrelated compounds also had *ScYRR1* mutations (Supplemental Table 2). These findings suggest that *ScPMA1* is the spiroindolone target, and *ScYRR1* is a more general resistance gene.

To separate out the individual alleles' contribution to resistance, genetic validation using the CRISPR/Cas system was performed. These experiments confirmed that mutations in *ScPMA1* and *ScYRR1* both cause a 2.5 fold increase in KAE609 resistance and that they have a multiplicative effect, as observed in the directed-evolution experiments. However, *ScYrr1p* does not appear to be the primary KAE609 target. When we deleted *ScYRR1*, the resulting strain was viable, demonstrating that *ScYRR1* is not essential. Furthermore, KAE609 potency increased in the deletion mutant, further suggesting that *ScYrr1p* confers resistance through an indirect mechanism (e.g., by activating the transcription of proteins that can detoxify KAE609, Fig. 2a).

***ScPMA1* mutation confers sensitivity to edelfosine, but not other antimicrobials.** Having shown that *ScPMA1* mutations are sufficient to generate KAE609 resistance, we next sought to determine the specificity of the resistance. We tested the L290S CRISPR-engineered *ScPMA1* mutant against a set of known antimicrobials with mechanisms unrelated to *ScPma1p*. We also included the alkyl-lysophospholipid edelfosine as a positive control because it reduces *ScPma1p* plasma-membrane concentrations by selectively displacing *ScPma1p* for endosomal degradation^{12–15}. None of the unrelated antimicrobials demonstrated cross-resistance or sensitivity in the *ScPMA1* mutant. However, the mutant did demonstrate 7.5-fold increase in sensitivity to edelfosine, suggesting that the mutation confers a fitness cost to the functioning of *ScPma1p* (Fig. 2b).

KAE609 exposure increases intracellular acidity in yeast. Although homology modeling and functional evidence indicates that *PfAtp4p* maintains sodium homeostasis⁵, *ScPma1p* functions as the major proton pump in *Saccharomyces* and is essential for cell viability¹⁶. It is responsible for maintaining pH homeostasis by pumping protons out of the cell; disruption of *ScPma1p* function produces a drop in intracellular pH as hydrogen

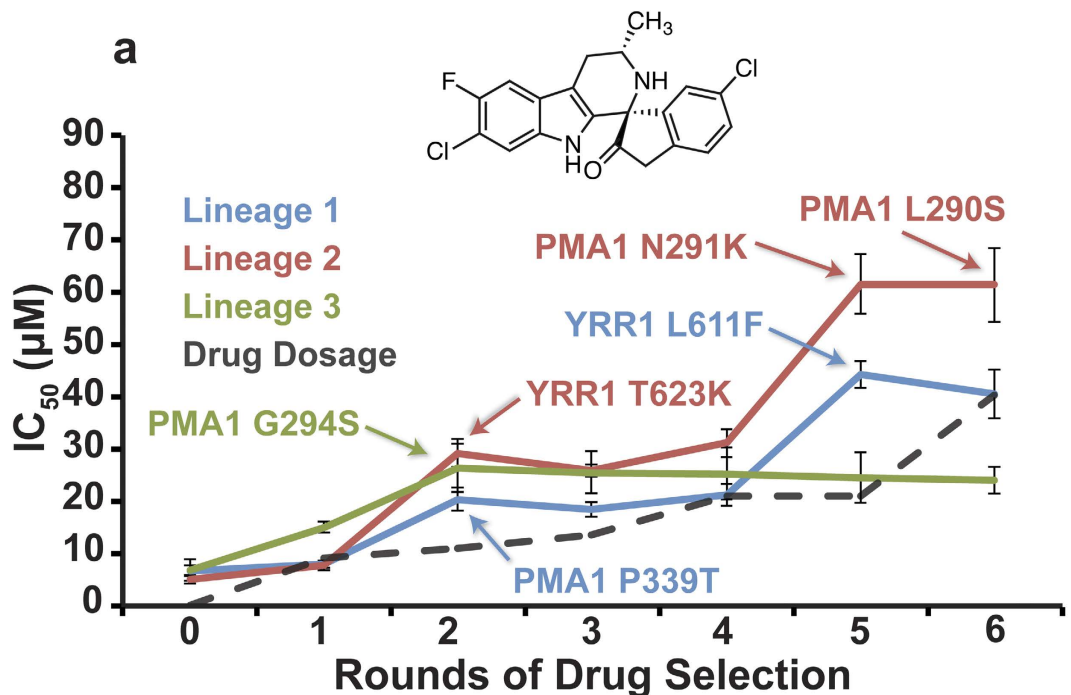


Figure 1. KAE609 directed evolution produces mutations in *ScPMA1*. (a) An IC_{50} analysis of ABC₁₆-Monster resistant lines showed that resistance developed over multiple rounds of drug selection. Sanger sequencing of *ScPMA1* and *ScYRR1* at each round was used to determine when each mutation (highlighted) arose in its respective lineage. (b) Alignment of *PfAtp4p* to *ScPma1p* (S288c) using PSI-BLAST (Position-Specific Iterated BLAST)⁶⁸. The figure shows conserved domains (*PfAtp4p* amino acid 139–462;*ScPMA1* 87–389; *PfATP4* 744–1032;*ScPMA1* 527–784). *ScPma1p* and *PfAtp4p* are homologous only in the E1-E2 ATPase domain. The alignments of *PfAtp4p* amino acids 314–429 and *ScPma1p* 252–356 are shown below. Residues that confer resistance in *Plasmodium* when mutated are colored based on the compound class used: red for dihydroisoquinolones and blue for spiroindolones (see³⁸ for a review).

ions accumulate in the cytosol^{17,18}. If KAE609 inhibits *ScPma1p*, we would expect a drop in cytosolic pH after drug exposure. Therefore, intracellular pH was measured using a strain of *S. cerevisiae* expressing the pH-sensitive green fluorescent protein (pHluorin)¹⁹. Since these cells are not ABC₁₆-transporter multi-knockouts, higher KAE609 dosages were required. After 3 hours of treatment with 200 μ M KAE609, the cytoplasmic pH dropped from 7.14 ± 0.01 to 6.88 ± 0.04 . This equates to an 80.6% increase in the cytoplasmic hydrogen ion concentration ($p = 0.0024$) (Fig. 2c), consistent with the hypothesis that KAE609 prevents *ScPma1p* from pumping hydrogen ions from the cytoplasm to the extracellular space.

Selection	Gene Name	Nucl. change	AA Change	Description
Lineage 1	<i>PMA1</i>	Cca/Aca	P339T	Plasma membrane P2-type H ⁺ -ATPase
Lineage 1	<i>YOS9</i>	aaC/aaA	N490K	ER quality-control lectin
Lineage 1	<i>YRR1</i>	ttG/ttC	L611F	Zn2-Cys6 zinc-finger transcription factor
Lineage 2	<i>PMA1</i>	aaC/aaG	N290S	Plasma membrane P2-type H ⁺ -ATPase
Lineage 2	<i>SLA1</i>	Aat/Gat	N788D	Cytoskeletal protein binding protein
Lineage 2	<i>YRR1</i>	aCg/aAg	T623K	Zn2-Cys6 zinc-finger transcription factor
Lineage 3	<i>GCD1</i>	Gtt/Ctt	V450L	Gamma subunit of the translation init. factor eIF2B;
Lineage 3	<i>PMA1</i>	Ggt/Agt	G294S	Plasma membrane P2-type H ⁺ -ATPase

Table 1. Nonsynonymous changes identified by whole-genome sequencing. Clones were sequenced to 40–90X coverage using paired-end reads and aligned to the S288c reference genome. *PMA1* mutations are shown in bold. No intergenic mutations near *PMA1* were identified. In addition, PCR analysis of nonclonal cultures identified an additional L291K *PMA1* substitution in Lineage 2, Round 5, derived from a parent containing the *YRR1* L611F mutation. This genotype was confirmed by whole-genome sequencing. Nonsynonymous coding changes in retrotransposons and flocculation genes (*FLO1*, *FLO4*, *FLO2*, *FLO9*) were also observed but were considered nonspecific.

KAE609 homology model explains mutations. To investigate KAE609 binding in more detail, we created a homology model of wild-type ScPma1p (UniProt ID: P05030) in the E2 (cation-free) state and mapped the four identified mutations (Leu290Ser, Pro339Thr, Gly294Ser, and Asn291Lys) onto the protein. The altered amino acids line a well-defined, cytoplasm-accessible pocket within the membrane-spanning domain that is large enough to accommodate a small molecule (Fig. 3). The computer-docking program Glide XP^{20–22} was next used to position KAE609 within the predicted wild-type pocket, with minimal manual adjustments. The docked pose placed the tricyclic tetrahydropyridoindole (THPI) moiety of the ligand between two residues that were mutated during directed evolution (Leu290 and Pro339, Fig. 3a,b), providing a plausible explanation for why these residues are so critical for KAE609 binding.

The docked pose suggests that receptor–ligand interactions are governed predominantly by high shape complementarity and hydrophobic contacts, provided by residues such as Leu102 (Fig. 3b,c). That both the Leu290Ser and Pro339Thr mutations substituted nonpolar with polar amino acids further supports the hypothesis that hydrophobicity plays a critical role. We note that Pro339Thr in ScPMA1p corresponds to Pro415Thr in *PfAtp4p*. In malaria parasites, the latter amino-acid modification is known to confer resistance to the dihydroisoquinolones, another class of antimalarials believed to target *PfAtp4p* by binding into the same pocket that we here identify as the KAE609 binding site⁷. Interestingly, our data showed that Leu290Ser ABC₁₆-Monster yeast were sensitized to the action of (+)-SJ000571311 (also known as SJ311), a representative dihydroisoquinolone⁷ resulting in a small shift from 133.4 ± 23.4 to 83.65 ± 7.98 μ M (Supplemental Table 3).

KAE609 may also form multiple hydrogen bonds with Ser364 and Leu363. The backbone amino groups of both these residues are predicted to form hydrogen bonds with the Ser364 ketone oxygen atom at the 2 position. Depending on the configuration of the Ser364 side chain, the Ser364 hydroxyl group may function as a hydrogen-bond acceptor bonded to the Ser364 nitrogen at the 1 position (as shown in Fig. 3b), or as another hydrogen-bond donor to the ketone oxygen atom at the 2 position. A hydrogen bond may also form between Thr343 and the KAE609 fluorine atom at the 6' position (Fig. 3b). H-F hydrogen bonds have been known to contribute to small-molecule binding in some contexts²³, though some argue that they are rare and typically weak²⁴.

KAE609 potently blocks ScPma1p ATPase activity in a cell-free assay. The homology model suggested that KAE609 is a direct ScPma1p inhibitor. To test for direct inhibition, ScPma1p-coated vesicles were harvested from *S. cerevisiae* cells with a temperature-dependent defect in secretory-vesicle/plasma-membrane fusion and bearing a *ScPMA1* overexpression plasmid. This combination results in high levels of ScPma1p accumulation in vesicles with little detectable ATPase activity in empty vector controls or from mitochondrial or vacuolar ATPases. This ATPase assay measures the hydrolysis of ATP to Pi²⁵. In these experiments, KAE609 had an IC₅₀ of 81.1 nM (SE = 1.2 nM), ten times more potent than inorganic orthovanadate, the non-specific ATPase inhibitor used as a positive control (IC₅₀ = 842 nM, SE = 1.20 nM, Fig. 3d). Artemisinin, an unrelated antimalarial that served as a negative control, showed no activity. KAE609 inhibition of ScPma1p in a cell-free context is consistent with the hypothesis that ScPma1p is the direct KAE609 target and inconsistent with an indirect mechanism by which *ScPMA1* mutations confer KAE609 resistance or an indirect mechanism by which KAE609 affects ScPma1p functioning.

To verify the hypothesis that a hydrophobic interaction at the 7' position is key to potency, we tested the activity of KAE609 derivatives²⁶ with substitutions at both the 6' and 7' positions (Fig. 4a,b). In the cell-free ScPma1p assay (Fig. 4c), substituting fluorine for the chlorine at position 7' results in a 10-fold decrease in potency. The

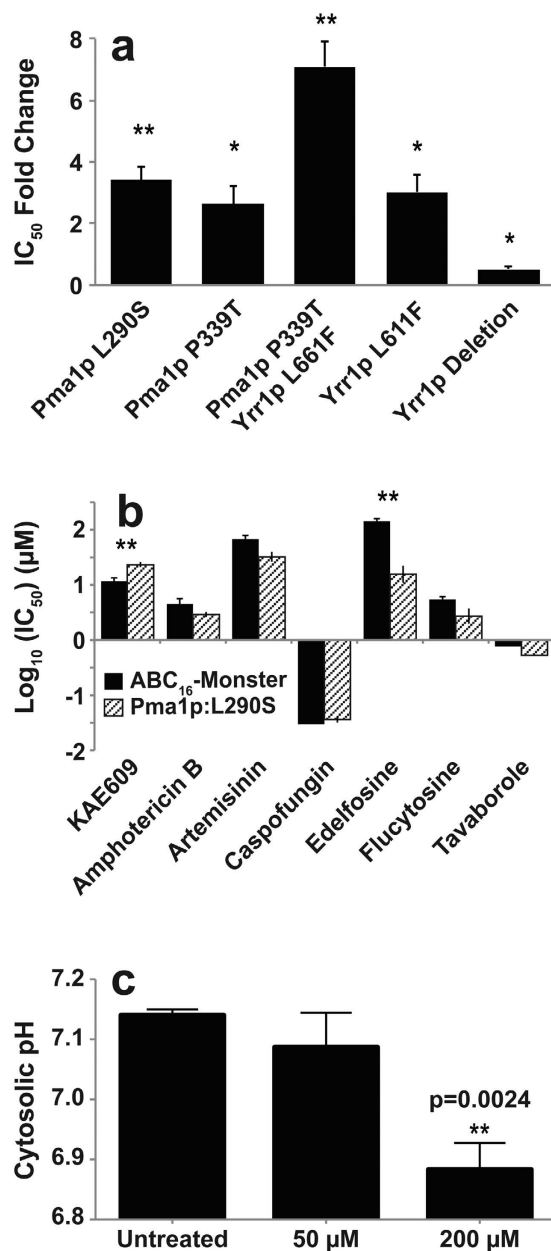


Figure 2. Mutations in *ScPMA1* confer resistance to KAE609, sensitize yeast to edelfosine, and treatment with KAE609 leads to a change in cytosolic pH. (a) Changes in ABC₁₆-Monster KAE609 resistance associated with mutations introduced using the CRISPR/Cas9 system, relative to the unmodified parent line. (b) The CRISPR mutant containing the *ScPma1p*:L290S amino acid change was tested for cross resistance or sensitivity to a range of antimicrobials. It shows 7.5-fold cross sensitivity to edelfosine, a chemical known to indirectly inhibit *ScPma1p* function. (c) The effect of KAE609 treatment on cytosolic pH was measured using cytosolically expressed ratiometric pHluorin, a pH-sensitive GFP¹⁹. Cytosolic pH was measured for three biological replicates. pH values are represented as mean \pm standard deviations. P values for IC₅₀ fold-changes were determined using a one-tailed ratio paired t-test comparing the ratio of the mutant-strain IC₅₀ value to that of the parental strain. For all bar graphs, (*) indicates $p < 0.05$ and (**) indicates $p < 0.01$.

removal of both the 6' and 7' halides causes a 100-fold decrease in potency. As the presence of fluorine at the 6' position has little effect on potency, the removal of the 7' halide appears to be exclusively responsible for the 100-fold decrease. Similar structure-activity relationships (SAR) are also observed in the whole-cell assay (Fig. 4d). The high degree of correlation between these two assays ($r = 0.93$) further supports our hypothesis that the cytotoxic effect of KAE609 is primarily due to direct *ScPma1p* inhibition.

Discussion

Phenotypic screening is a powerful tool for discovering anti-infective and antiproliferative compounds²⁷. Phenotypic cellular screens are efficient because they effectively multiplex many critical targets, identifying

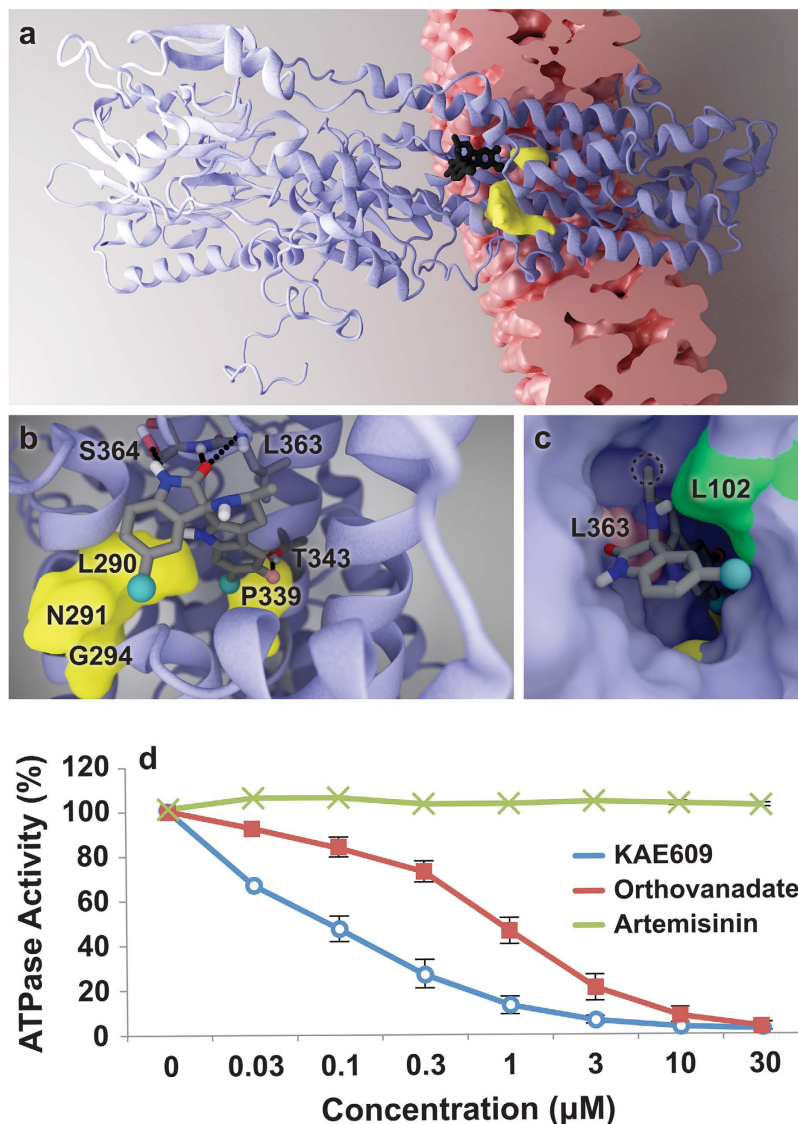


Figure 3. Functional and docking analyses support the hypothesis that KAE609 is a direct *PfAtp4p* inhibitor. (a–c) Illustrations of the ScPma1p homology model and KAE609 docked pose. (a) The docked ligand is shown in black. Amino acids associated with evolved resistance are shown in yellow. The lipid-bilayer location was predicted using PDBID 1MHS⁶⁹, CHARMM-GUI⁷⁰, and the OPM database⁷¹. (b) The ligand with predicted hydrogen-bond partners (Ser364, Thr343, and Leu363). (c) The receptor in surface representation. Leu102 (green) is predicted to form hydrophobic contacts with KAE609. The position of Leu363 (pink) explains why the chirality at the 3' carbon atom is critical to potency; if inverted, the attached methyl group (circled) would clash sterically with Leu363. (d) KAE609 inhibition of ScPMA1p in the vesicle-based assay²⁵. Orthovanadate, a non-specific ATPase inhibitor, and artemisinin, an unrelated antimalarial, are included as controls. Two independent experiments were performed.

cytotoxic inhibitors of DNA replication, protein synthesis, secretion, transcription, or other cellular processes via a single, miniaturized assay. Although not necessarily critical for compound development, discovering a specific protein target can inform subsequent drug development, enabling both structure-guided drug design and a better understanding of off-target activity.

Existing techniques for target deconvolution are limited. For example, “omic methods” such as haploinsufficiency screens may produce an overwhelming amount of difficult-to-interpret data (e.g. at least 2,452 gene deletions confer resistance or sensitivity to cycloheximide, according to the *Saccharomyces* Genome Database)²⁸. Proteomics approaches, which detect compound-binding proteins from a cell lysate, also produce high false-positive rates²⁹, as do computational approaches such as proteome-wide virtual screening³⁰. Additionally, transcriptional profiling is only rarely used for target identification^{31,32}.

Directed evolution, on the other hand, has been used for many years to discover targets in bacteria. For example, the target of nalidixic acid was discovered to be DNA gyrase by mapping mutations in resistant lines³³. More recently, directed evolution has also been used to find the targets of a novel antifungal³⁴. However, its use

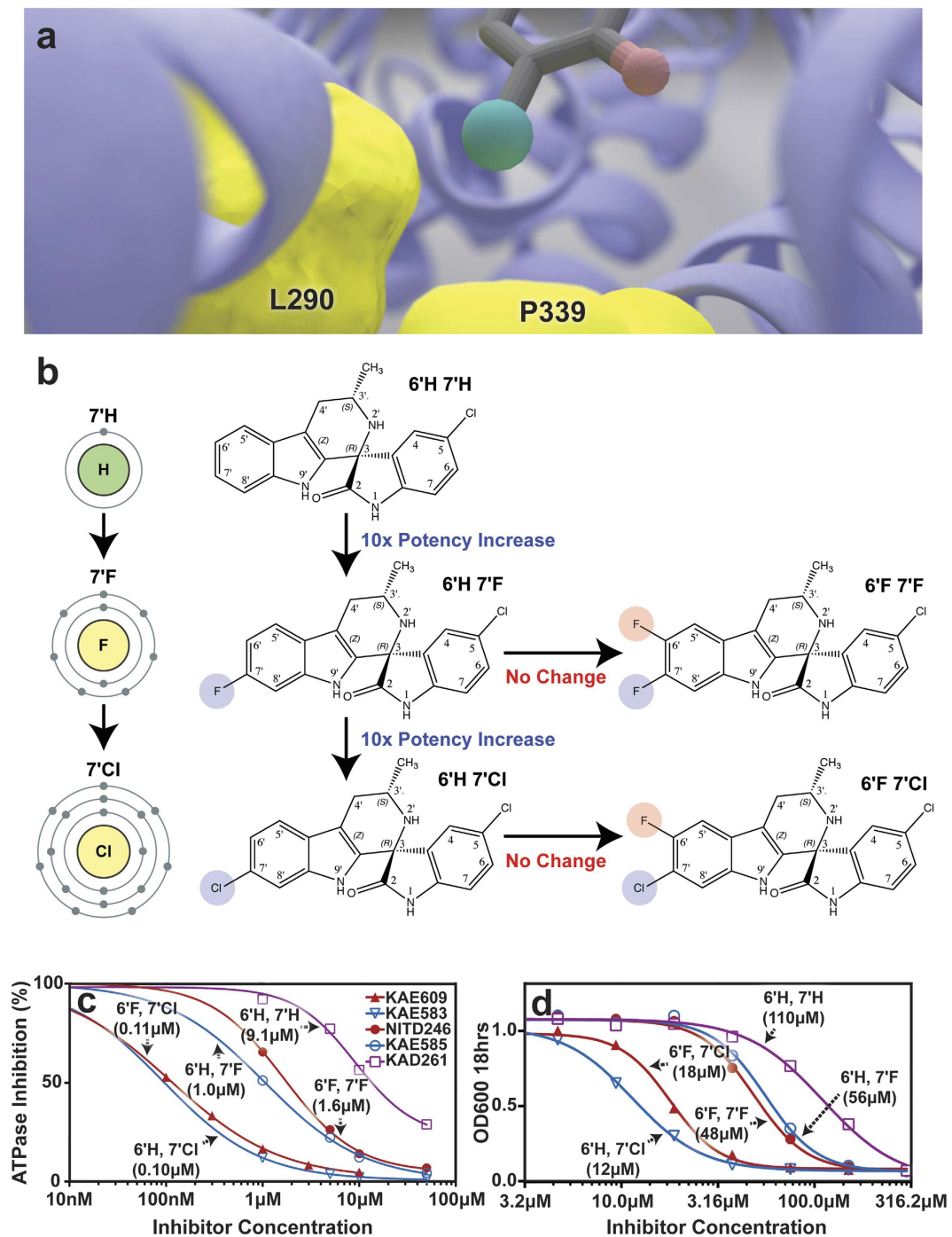


Figure 4. KAE609 halogenation. (a) An illustration of the predicted binding pose, which positions the 7' chlorine atom (green) between two residues that were altered during directed evolution (L290S and P339T). Unlike the polar mutant residues, the nonpolar wild-type residues may stabilize the interaction with the 7' chlorine atom. (b) In yeast, halogenation at the 7' position (blue) has a substantial impact on potency, but halogenation at the 6' position (red) has little impact. Note that KAE609 is the molecule designated 6'F 7'Cl. (c) The blue, red, and purple curves indicate 7', 6'/7', and no halogenation. In the cell-free assay, 7' chlorination yields ~10- and ~100-fold greater potency than fluorination and no halogenation, respectively. (d) Structure-activity relationships showing correlation between direct inhibition of ScPMA1p in the vesicle-based assay²⁵ and activity in a whole-cell yeast replication assay (see methods). The *in vitro* IC₅₀ values against blood-stage *P. falciparum* NF54 (in nM) were 13 ± 2.2 (6'H 7'H), 4.3 ± 0.4 (6'H 7'F), 0.33 ± 0.06 (6'F 7'F), 5.6 ± 0.7 (6'H 7'Cl), and 1.2 ± 0.2 (6'Cl 7'F, KAE609)⁷². The following naming conventions were used by Yeung *et al.* in their paper concerning spiroindolone SAR: 6'H 7'H = (+)-1, 6'H 7'F = (+)-5, 6'F 7'F = (+)-6, 6'H 7'Cl = (+)-3, and 6'Cl 7'F (KAE609) = (+)-7²⁶.

has mostly relied on the availability of rapid methods for mapping the resistance locus to a given chromosomal position. The advent of whole-genome tiling arrays and whole-genome sequencing has made the method more practical in other single-celled organisms such as *Mycobacterium tuberculosis* bacilli³⁵, malaria parasites³⁶, trypanosomes³⁷, and yeast^{38–40}. Unlike other essential gene products that are often described as possible drug targets, drug targets discovered through directed evolution are known from the outset to be druggable and are considered “chemically validated.” The method has been successfully used to identify the targets of several antimalarial compounds initially found through phenotypic screens, as well as to identify genes contributing to the resistome^{2,41,42}.

On the other hand, directed evolution followed by genome sequencing is still not practical in organisms that are multicellular, sexual, or difficult to culture. Many organisms also have lengthy cell cycles, large genomes that are prohibitively expensive to sequence, and/or multidrug efflux pumps that render them insensitive to many compounds of interest. Furthermore, resistance in multiploid organisms often requires simultaneous mutations in all alleles. Additionally, directed evolution is not possible for compounds that target non/low-replicative stages of pathogen life-cycles, such as next generation antimalarials currently under investigation that specifically target the liver or sexual stages. These compounds may, however, be cytotoxic to drug-sensitive yeast.

Here we demonstrate that directed evolution in surrogate species can identify the target class of a given drug, if not the actual target, opening up the method for use in other species. *S. cerevisiae* has advantages over many other organisms, including a well-characterized genome, a rich scientific literature, and a fast growth rate that produces resistant, engineered, or recombinant yeast in 24 hours versus 24 weeks (i.e., a nonconservative estimate of the time needed to perform selections and confirmation in pathogens such as *P. falciparum*). However, it should be acknowledged that yeast, even with a number of drug pumps removed, is still better at detoxifying xenobiotics than many microbes and that higher concentrations of compound will likely be needed to achieve inhibition.

Our data support the hypothesis that *PfAtp4p* is the direct target of the spiroindolones and by extension the dihydroisoquinolones, which share the same binding site. We show direct inhibition of the *ScPma1p* enzyme in a cell-free system and study the SAR of the spiroindolones against their specific target, rather than the whole cell. We show that *ScPma1p* inhibition leads to a cytosolic decrease in pH, consistent with the hypothesis that *ScPma1p* is the KAE609 target. Our results are in agreement with studies showing that spiroindolones significantly reduce Na⁺-dependent ATPase activity in isolated membranes derived from parasitized erythrocytes⁵.

Our findings may also help explain the recent controversial observation that *PfATP4* mutants have either increased resistance or sensitivity to a broad range of unrelated antimalarials (reviewed in ref. 38), raising the possibility that *PfATP4* is a general drug resistance gene. The finding that a single amino acid change in *ScPma1p* sensitizes the cell to edelfosine by 750% suggests that these mutations confer a fitness cost to the protein, reducing either the ability to pump hydrogen ions or to stay associated with plasma-membrane lipid rafts. It stands to reason that amino-acid changes in *PfAtp4p* may similarly affect function, leading to changes in cytosolic pH and membrane potential, with differing effects on xenobiotic uptake, deactivation, modification, degradation, or export.

An open question is whether or not *PfAtp4p* and *ScPma1p* have the same function. The function of *PfAtp4p* in malaria parasites remains uncertain, but experiments have shown that it may affect both sodium and proton homeostasis⁵. In contrast, *ScPma1p* is only involved in maintaining proton homeostasis. Although highly homologous ($P(n) = 4.60E-38$ per BLAST-P), *PfAtp4p* and *ScPma1p* share only the E1-E2 ATPase domains, not the cation ATPase C terminal region (pfam00689) or the hydrolase domain (Fig. 1b). *PfAtp4p* shows higher homology to *ScPmr1p*, *ScEna1p*, *ScEna2p*, and *ScEna5p* than to *ScPma1p*. However, with the exception of *ScEna5p*, these proteins are nonessential in *Saccharomyces* and thus are unlikely KAE609 targets. In addition, we show that KAE609 treatment results in acidification of the yeast cytosol, but alkalization in *P. falciparum* was reported previously⁵. An explanation may be that both proteins pump hydrogen ions, but are in opposite orientation in the plasma membrane. This seems unlikely: although the orientation of *PfAtp4p* in the plasma membrane is not known, it is likely that the ATP-binding domain, as with *ScPma1p*, is located in the cytosol⁴³. Given that parasites lose the ability to extrude sodium after KAE609 treatment⁵, it seems likely that the two proteins do not have the same cellular function.

On the other hand, the *PfAtp4p* binding pose is likely similar to the *ScPma1p* pose presented here, especially given that the evolved yeast mutations are in a highly conserved region of the protein that is shared with *P. falciparum*. Both the 6' and 7' halides were originally added during the drug optimization process to reduce CYP2C9-mediated hepatic clearance, and the discovery that these modifications also increase potency up to 45-fold in whole-cell malarial assays was fortuitous²⁶. The current work suggests a structure-based explanation for the enhanced antimalarial activity of the halogenated compounds. However, further study of the malarial protein is warranted. There are some differences between the two proteins, as reflected in subtle differences in the ABC₁₆-Monster and malarial whole-cell SAR. For example, 6' halogenation has little effect on ABC₁₆-Monster inhibition. In contrast, it increases potency against whole-cell *P. falciparum*²⁶. Additionally, 7' chlorination reduces malarial potency versus 7' fluorination, but the opposite is true in yeast²⁶.

Although our data strongly suggest that *PfAtp4p* is the direct target of both the spiroindolones and, by analogy, the dihydroisoquinolones, it is not yet clear that *PfAtp4p* is the direct target of the aminopyrazoles⁶ or pyrazole amides⁹. An aminopyrazole (GNF-Pf4492) was tested and had little measurable effect on ABC₁₆-Monster growth (see Supplemental Table 3). Aminopyrazole resistance-conferring and -sensitizing mutations in *PfATP4* are distant from the sites identified in yeast. It is possible that these compounds bind to another pocket, or that these *PfATP4* mutations confer resistance through another mechanism.

Despite these questions, this work supports the hypothesis that *PfAtp4p* is the direct KAE609 target in malaria and provides an additional method for characterizing P-type ATPase inhibitors. The data also support previous work suggesting that *ScPma1p* may be an attractive target for novel antifungals^{44–46}. Recent successes in targeting P-type ATPases in *Plasmodium* may be translated into new strategies against other pathogens for which novel drug classes are badly needed.

Methods

Whole-Cell *S. cerevisiae* IC₅₀ Assay. To measure compound activity against whole-cell yeast, each yeast generation was established using cells taken from single colonies on agar plates and inoculated into 2 mL of media in 5 mL snap-cap culture tubes. The tubes were cultured overnight at 250 RPM in a shaking incubator at 30 °C (Controlled Environment Incubator Shaker, Model G-25, New Brunswick Scientific Co., Inc.). Cultures were extracted when they were in mid-log, as judged by an OD600 (600 nm) reading between 0.1 and 0.5.

After being diluted to OD600 0.01, 30 μ L of the cells were added to the wells of a 384 well plate. Fifteen 1:2 serial dilutions were subsequently performed, in duplicate for each clone, with starting IC₅₀ values of 150 μ M. Following an initial reading of OD600 (time 0 hours), the plate was placed in an incubator at 30 °C for 18 hours. After incubation, the plates were placed in a Synergy HT spectrophotometer, shaken for one minute on the “high” setting, and immediately read at OD600.

IC₅₀ values were determined first by subtracting OD600 values at time 0 hours from time 18 hours. Nonlinear regression on log(inhibitor) vs. response with variable slope (four parameters) was then performed using Graphpad Prism. Minimum values were constrained to 0.0. For resistance assays, three biological replicates of technical duplicates were used to calculate each final IC₅₀ value. P-values for IC₅₀ fold-changes were determined using a one-tailed ratio paired t-test, comparing the ratio of the mutant and parental IC₅₀ values. For the cross-resistance/cross-sensitivity assay, at least three biological replicates of technical triplicates were used to calculate each final IC₅₀ value. P-values for IC₅₀ fold-changes were determined using a two-tailed ratio paired t-test, since we did not have a prior hypothesis as to whether the IC₅₀ should increase or decrease.

Directed-Evolution Selection Protocol. ABC₁₆-Monster cells (10 μ L) were taken from a saturated culture and inoculated into 30 mL of YPD media in a 50 mL conical tube, with the desired KAE609 concentration. The tubes were then cultured at 250 RPM in a shaking incubator at 30 °C. Cultures that achieved OD600 values greater than 0.7 were advanced to the next generation of directed evolution.

DNA Extraction, PCR, and Sanger Sequencing. DNA extractions were performed using the YeaStar Genomic DNA kit. To confirm all non-synonymous SNVs identified in open-reading frames by whole-genome sequencing, genomic DNA was PCR amplified in a 20 μ L reaction volume with PrimeSTAR Max polymerase (Takara Bio Inc, Otsu, Shiga, Japan) using the forward primer (5'-CTT CCA CTG TTA AGA GAG GTG AAG G-3') and the reverse primer (5'-CTG GAA GCA GCC AAA CAA GCA GTC-3') (Allele Biotechnology & Pharmaceuticals, Inc). PCR products were sequenced directly (Eton Bioscience Inc, San Diego, CA).

Whole-Genome Sequencing and Analysis. Genomic DNA libraries were normalized to 0.2 ng/ μ L and prepared for sequencing using the Illumina Nextera XT kit whole-genome resequencing library, according to the manufacturer's instructions (see the Illumina protocol of tagmentation followed by ligation, v. 2013, Illumina, Inc., San Diego). DNA libraries were clustered and run as 2 \times 100 paired end reads on an Illumina HiSeq, according to the manufacturer's instructions. Base calls were made using CASAVA v1.8.2. After sequencing, initial alignment was done using the Platypus software⁴⁷. Briefly, reads were aligned to the reference *S. cerevisiae* genome using BWA, and unmapped reads were filtered using SAMTools. SNPs were then initially called using GATK and filtered using Platypus. Copy number variants were analyzed using Control-FreeC⁴⁸. Sequencing statistics can be found in Supplementary Table 1.

Site-Directed Mutagenesis. Point mutations were introduced in the ABC₁₆-Monster strain using the Clustered Regularly Interspaced Short Palindromic Repeats (CRISPR) and CRISPR-associated (Cas) system from *Streptococcus pyogenes*⁴⁹. Plasmids p414-TEF1p-Cas9-CYC1t (p414) and p426-SNR52p-gRNA.CAN1.Y-SUP4t (p426) were obtained from Addgene (Cambridge, MA). Plasmid transformation markers were replaced with markers that are compatible with the yeast strains used in the current study. The *TRP1* marker of the p414 plasmid was replaced with the *MET15* marker. As the centromere-autonomous replication sequence (CEN-ARS) region of this plasmid was difficult to amplify using Q5 DNA polymerase (New England Biolabs, Ipswich, MA), the CEN-ARS region of the original plasmid was also replaced with a CEN-ARS sequence from *Mycoplasma mycoides* JCVI-syn1.0⁵⁰. To create a p414 fragment containing the mycoplasma CEN-ARS sequence, three PCR products were generated: 1) an upstream 289-base-pair fragment amplified from p414 using the primers 414_VecF and 414_GB_3R, 2) a 525-base-pair CEN-ARS fragment amplified from genomic DNA of the mycoplasma strain using the primers 414_GB_2F and 414_GB_2R, and 3) a downstream 644-base-pair fragment amplified from p414 using the primers 414_GB_1F and Pre-p426_Halfway_R (see Supplementary Table 4). These three fragments were combined using crossover PCR to generate a single, contiguous fragment (fragment 1). To prepare the remaining parts, a 2.1-kb fragment containing the *MET15* gene was amplified from a *MET15*-containing plasmid⁵¹ using the primers 414_MET15F and New_414_MET15_R (fragment 2). The two additional fragments generated from the p414 plasmid were a 1.7-kb fragment amplified using the primers p426_Halfway_F and New_414_Vec_Only_HalfR (fragment 3), and a 5.6-kb fragment amplified using the primers 414_Vec_Half_F and 414_VecR (fragment 4). Fragments 1–4 were combined using homologous recombination in yeast to make the plasmid p414-TEF1p-Cas9-CYC1t-MET15.

The *URA3* marker of the p426 guide RNA plasmid was replaced with the *LEU2* marker. The p426 backbone was amplified as two fragments, a 2.2-kb fragment amplified with the primers Attempt2_426_Vec_F and p426_Halfway_R (fragment 5), and a 3.1-kb fragment amplified with the primers p426_Halfway_F and New_426_Vec_R (fragment 6). A 2.1-kb *LEU2* fragment was amplified from a plasmid derived from pRS315 (ref. 52; AM & YS, unpublished result) using the primers New_426_LEU2_F and Attempt2_426_LEU2_F (fragment 7). Fragments 5–7 were combined using Gibson assembly⁵³ to make the plasmid p426-SNR52p-gRNA.CAN.Y-SUP4t-LEU2.

For the introduction of each mutation, a unique 20-base-pair guide RNA (gRNA) sequence was required. To generate the required variants of the gRNA plasmid, p426-SNR52p-gRNA.CAN.Y-SUP4t-LEU2 was PCR-amplified using a high-fidelity DNA polymerase PrimeSTAR Max (2 × Master Mix, Takara Bio, Mountain View, CA) and primers containing the appropriate 20-base-pair gRNA sequence at the 5' end (see Supplementary Table 5). The two ends of the PCR product shared 25 base pairs of homology, replacing the original gRNA sequence.

After purification using NucleoSpin Gel and the PCR Clean-up kit (Macherey-Nagel, Bethlehem, PA), the linearized plasmid was introduced into High Efficiency NEB 5-alpha chemically competent cells (New England Biolabs, Ipswich, MA), per the manufacturer's protocol^{54,55}. Transformed colonies were selected on LB-ampicillin agar plates. Colonies were then cultured in liquid LB-ampicillin medium, and plasmids were isolated using a miniprep kit (Qiagen, Valencia, CA). The correct gRNA sequence was verified using Sanger sequencing. Repair fragments were prepared from complementary 90-base-pair oligonucleotides (IDT, Coralville, IA), which included the desired mutation (see Supplementary Table 6). To prepare double-stranded repair fragments, complementary oligonucleotides were mixed in equimolar amounts, denatured at 100 °C for five minutes, and allowed to cool to 25 °C with a ramp rate of 0.1 °C per second.

The ABC₁₆-Monster strain was initially transformed with ~200 ng of the p414-TEF1p-Cas9-CYC1t-MET15 plasmid using a standard lithium acetate transformation method⁵⁶ and selected on synthetic complete (SC) minus Met agar plates. To introduce a desired mutation, Cas9-expressing cells were cultured overnight in SC-Met. Using lithium acetate transformation⁵⁶, the cells were co-transformed with ~500 ng of the appropriate gRNA plasmid and 2.5 nmol of the corresponding double-stranded repair fragment. All of the transformed cells were selected on SC-Met-Leu agar plates.

To verify the correct introduction of the mutations, the genomic region around the mutation was PCR-amplified using a high-fidelity DNA polymerase Q5 (2 × Master Mix, New England Biolabs) and sequenced using Sanger sequencing (see Supplementary Table 7). Before phenotype testing, strains were cultured overnight in a YPDA liquid medium, the resulting cells were plated on a YPDA agar medium to form isolated colonies, and the loss of both plasmids was confirmed by transferring the colonies to both rich and selective media.

ScPma1p ATPase Assay. ATP hydrolysis was assayed at 30 °C in 0.5 mL of 50 mM MES/Tris, pH 6.25, 5 mM Na₃, 5 mM Na₂, ATP (Roche), 10 mM MgCl₂, and an ATP regenerating system (5 mM phosphoenolpyruvate and 50 µg/mL pyruvate kinase). The reaction was terminated after 20 minutes by the addition of Fiske and Subbarow reagent, and the release of inorganic phosphate was measured at 660 nm after 45 minutes of color development²⁵.

Cytosolic pH Measurement. To measure cytosolic pH-changes the yeast strains containing a cytoplasmically expressed ratiometric pHluorin plasmid (UCC9633) was used (see Supplemental Table 8). The plasmid was created by amplifying a fragment of plasmid pADH1pr-RMP⁵⁷ using primers URA3-tTA-intChr1 F and R to integrate ratiometric pHluorin¹⁹ into an empty region of chromosome 1 (17068–17161) under the control of the ADH1 promoter. Cells were cultured in YEPD (1% yeast extract, 2% peptone, 2% glucose) exponentially for 4 hours prior to 3 hours of treatment with 200 µM KAE609. For pH measurement, cells were transferred to low fluorescence medium⁵⁸ at a density of 3 × 10⁷ cells/mL after rinsing them in an equal volume of low fluorescence medium. Calibration curves were calculated as described in ref. 57.

To quantify cytosolic pH and generate calibration curves, pHluorin fluorescence emission was measured at 512 nm using a SpectraMax M5 microtitre plate spectrofluorometer (Molecular Devices, Sunnyvale, CA), providing excitation bands of 9 nm centered around 390 and 470 nm. Background fluorescence for wild-type cells not expressing pHluorin (UCC4925) was subtracted from the measurements. The ratio of emission intensity resulting from excitation at 390 and 470 nm was calculated. Ratios were fitted to a calibration curve to derive cytosolic pH from three biological replicates for each pH measurement. pH values are represented as mean ± sd.

ScPma1p Homology Modeling. To better understand the KAE609 binding pose, a homology model of ScPma1p was built with Schrödinger's Prime software⁵⁹ using the UniProt⁶⁰ sequence P05030 and a structure of the highly homologous *Sus scrofa* sodium-potassium pump (PDBID: 3N2F, chain C)⁶¹. The 3N2F and P05030 amino-acid sequences were first aligned using ClustalW⁶². The model was then constructed using Schrödinger's knowledge-based method. The structure was further processed with Schrödinger's Protein Preparation Wizard⁶³. Hydrogen atoms were added at pH 7.0 using PROPKA⁶⁴, water molecules were removed, and disulfide bonds were appropriately modeled. The system was then subjected to a restrained minimization using the OPLS_2005 forcefield^{65,66}, converging the heavy atoms to an RMSD of 0.30 Å.

KAE609 Docking. A three-dimensional KAE609 model was prepared using Schrödinger's LigPrep module. The protonation state was determined for pH values in the 5.0–9.0 range using Epik⁶⁷. The most likely tautomeric state and low-energy ring conformations were similarly determined. The molecular geometry was optimized using the OPLS_2005 forcefield^{65,66}.

Glide XP^{20,21} was used to dock KAE609 into the modeled E2-ScPma1p pocket near the amino acids associated with evolved resistance. The ligand diameter midpoint box was 14 Å cubed. Glide was instructed to use flexible ligand sampling, to sample nitrogen inversions and ring conformations, to bias torsion sampling for amides only, and to add Epik state penalties to the docking scores. The van der Waals radii of atoms with assigned partial charges less than or equal to 0.15 e were scaled to 80%.

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Author Contributions

G.M.G., S.O., J.D.D. and E.A.W. designed the experiments and wrote the manuscript. Preliminary feasibility studies were performed by Y.S. and C.W.M. Directed-evolution experiments were performed by G.M.G. IC₅₀ experiments were performed by G.M.G., S.O., E.V., J.Y., J.S., R.S. and M.P. Whole-genome sequencing was performed by the UCSD Institute for Genomic Medicine Core Facility. Sequence analysis was performed by G.M.G., F.G. and M.J.M. PCR studies were designed and performed by G.M.L., E.V., J.S. and G.M.G. CRISPR mutant strains were designed and engineered by M.K., Y.S., A.M., M.N. and G.L. Intracellular pH studies were designed and performed by K.H. Cell-free ATPase assays were designed and performed by K.E.A. and C.W.S. Computational docking studies were designed and performed by J.D.D. and R.E.A.

Additional Information

Accession codes: Sequences have been placed in the short read sequence archive (<http://www.ncbi.nlm.nih.gov/sra>) under Accession code SRP074482.

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