A Specific Two-pore Domain Potassium Channel Blocker Defines the Structure of the TASK-1 Open Pore*

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Two-pore domain potassium (K2P) channels play a key role in setting the membrane potential of excitable cells. Despite their role as putative targets for drugs and general anesthetics, little is known about the structure and the drug binding site of K2P channels. We describe A1899 as a potent and highly selective blocker of the K2P channel TASK-1. As A1899 acts as an open-channel blocker and binds to residues forming the wall of the channel pore, the drug was used to further our understanding of the channel pore. Using alanine mutagenesis screens, we have identified residues in both pore loops, the M2 and M4 segments, and the halothane response element to form the drug binding site of TASK-1. Our experimental data were used to validate a K2P open-pore homology model of TASK-1, providing structural insights for future rational design of drugs targeting K2P channels.

The acid-sensitive K2P4 channel TASK-1 (TWIK-related acid-sensitive K+) gives rise to instantaneous and non-inactivating potassium currents, which can be blocked by extracellular acidification (1). Five TASK channel family members have been described. The closest relatives of the TASK-1 channel are TASK-3 and TASK-5 (2, 3). The TASK-4 and TASK-2 channels show less homology and are functionally different as they are the acid-sensitive potassium channel.

K2P channels have a transmembrane topology that is unique among potassium channels. One subunit contains four transmembrane domains and two pore-forming domains. The functional channel is formed by a dimer. Although other potassium channels are tetramers with a pore formed by four identical transmembrane domains, the domains and symmetry of the K2P channel pore have long been a matter of speculation. The different topology of K2P channels does not allow a direct transfer of data from the crystal structures of other potassium channels without experimental proof of validity. The pore-forming domains and the rotation of these domains can be validated by an open-channel pore blocker as the amino acids of the binding site should face the water-filled pore cavity. We have identified the binding site of A1899 in the open pore of TASK-1 channels. In good agreement with recently published data (14), our results suggest that the M2 and M4 segments of two TASK subunits form the walls of the inner pore. Using A1899, we have confirmed the orientation of the M2 and M4 segments and provide a model of the mammalian K2P channel TASK-1 in the open state.

EXPERIMENTAL PROCEDURES

Site-directed Mutagenesis—Mutations in the human TASK-1 (KCNK3, NM_002246) cDNA were introduced with the QuikChange site-directed mutagenesis kit (Stratagene) according to the instructions of the manufacturer.

Oocyte Preparation, cRNA Synthesis, and cRNA Injection—Oocytes were obtained from anesthetized Xenopus laevis frogs and gland (8). In addition, they are also involved in peripheral chemosensation (9) and the regulation of the immune system (10). TASK channels are the molecular targets of volatile and local anesthetics (11, 12). Although studies of TASK channel knock-out mice have advanced our knowledge about the physiological function of these channels, a selective TASK channel blocker would be very valuable to perform future in vivo experiments. In the present study, we describe a highly potent blocker of the TASK-1 channel, A1899, active in the one-digit nanomolar range. As suggested by the high affinity, we found that A1899 is selective for TASK-1 channels. Although anandamide has been reported as a physiological blocker of TASK-1 channels (13), this lipid does not combine the high affinity and specificity as A1899.

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4 The abbreviations used are: K2P, two-pore domain potassium channel; TASK, TWIK-related acid-sensitive K+; MD, molecular dynamics; HRE, halothane response element; TWIK, tandem of P domains in a weak inwardly rectifying K+ channel; ERG, ether-a-go-go-related gene; dORK, Drosophila open rectifier K+ channel.

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and incubated in OR2 solution containing in mM: 82.5 NaCl, 2 KCl, 1 MgCl$_2$, 5 HEPES (pH 7.5) substituted with 2 mg/ml collagenase II (Sigma) to remove residual connective tissue. Subsequently, oocytes were stored at 18 °C in ND96 solution supplemented with 50 mg/liter gentamycin, 274 mg/liter sodium pyruvate, and 88 mg/liter theophylline. Human TASK-1 channels were subcloned into a pSGEM or pBF1 vector, and cDNA was linearized with NheI or MluI, respectively. cRNA was synthesized with mMESSAGE mMACHINE kit (Ambion). The quality of cRNA was tested using gel electrophoresis. Oocytes were each injected with 50 nl of cRNA.

Two-electrode Voltage Clamp Recordings—All two-electrode voltage clamp recordings were performed at room temperature (20–22 °C) with a TurboTEC 10CD (npi) amplifier and a Digidata 1200 Series (Axon Instruments) as analog/digital converter. Micropipettes were made from borosilicate glass capillaries GB 150TF-8P (Science Products) and pulled with a DMZ-Universal Puller (Zeitz). Recording pipettes had a resistance of 0.5–1.5 megaohms and were filled with 3M KCl solution.

Molecular Modeling—Based on the crystal structures of KcsA and KvAP (PDB IDs 1K4C and 1ORQ, respectively), homology models of TASK-1 were created using the Modeller 7 version 7 software. For ligand docking predictions, GOLD (version 4.0), AutoDock, and AutoDockTools were used to propose binding conformations for A1899 within the central cavity of TASK-1. Molecular dynamics (MD) simulations were performed with the Gromacs software. Detailed information on molecular modeling is provided in the supplemental Experimental Procedures, which includes details about the homology modeling, ligand docking experiments, and MD simulations.

Statistical Analyses—All values are expressed as means ± S.E. For all oocyte and CHO electrophysiological experiments, 3–12 cells were studied ($n = 3–12$). TRAAK, TWIK-related arachidonic-acid-stimulated K$^+$ channel.

### Statistical Analyses

- **A1899 Is a Selective Blocker of TASK-1 Channels**—We have tested the affinity of A1899 for TASK-1 channels expressed in *Xenopus* oocytes. Application of 40 nm A1899 led to a pronounced inhibition of TASK-1 currents (Fig. 1A). In oocytes, TASK-1 channels were blocked with an IC$_{50}$ of 35.1 ± 3.8 nm, analyzed at +40 mV (Fig. 1B).

### Results

**TABLE 1**

| Channel          | IC$_{50}$ (nM) |
|------------------|----------------|
| TASK-1           | 3.8            |
| TASK-3           | 120            |
| TASK-1 CHO       | 23.8           |
| TASK-3 CHO       | 8.4            |
| TASK-2           | 2.2            |
| TASK-4           | 2.3            |
| TREK-1           | 0.9            |
| TREK-2           | 2.7            |
| TRAAK            | 0.3            |
| THIK-1           | 2.0            |
| TRESK            | 0.0            |
| Kv1.1            | 0.0            |

Values were determined in *Xenopus* oocytes ($n = 3–12$) or when indicated in CHO cells ($n = 3–12$). TRAAK, TWIK-related arachidonic-acid-stimulated K$^+$ channel.

**Statistical Analyses**—All values are expressed as means ± S.E. For all oocyte and CHO electrophysiological experiments, 3–12 cells were studied ($n = 3–12$). Error bars in Figs. 1 and 3–5 represent S.E. values. Significance was assessed using two-tailed Student’s $t$ test. Asterisks indicate significance: *, $p < 0.05$; **, $p < 0.01$; ***, $p < 0.001$.
channels expressed in CHO cells was only 7 nM (Table 1), which is consistent with the notion that IC_{50} values determined in oocytes are mostly higher than those determined in mammalian cells. The IC_{50} for TASK-3 expressed in CHO cells was 70 nM (Table 1). Thus, TASK-3, the closest relative of TASK-1, binds A1899 with a 10-fold lower affinity. To further probe the selectivity of A1899, we expressed a set of different potassium channels in Xenopus oocytes and quantified the extent of current inhibition by 100 nM A1899 (Fig. 1C). Under these conditions, TASK-1 currents were blocked by 74.2 ± 3.4%. In contrast, the closest relative, TASK-3, was only blocked by 22.2 ± 1.6%. All the other channels tested showed only little (<11%) or no block (Fig. 1C). The IC_{50} values determined for several potassium channels (Table 1) indicate that A1899 is a specific TASK-1 blocker.

Subsequently, we recorded TASK-1 currents in the presence of an increased extracellular K⁺ concentration ([K⁺]_{out} = [K⁺]_{out}). Under these conditions, it was possible to test for a potential voltage dependence of block. We found that 400 nM A1899 blocked inward and outward currents to the same extent, indicating that the block of A1899 is not voltage-dependent (Fig. 1, D and E). However, the extent of block at +40 mV elicited by 400 nM A1899 in a bath solution with a high concentration of extracellular potassium was only 59.7% in comparison with about 90% in ND96 (Fig. 1, B and D), reflecting an almost 10-fold increase in IC_{50} of A1899. A reduced blocker affinity in bath solutions with elevated extracellular potassium concentrations is typical for an open-channel block mechanism (15, 16). Most open-channel blockers act from the intracellular side by binding in the water-filled central cavity (15, 17). To assess whether A1899 blocks TASK channels in the central cavity, we tested the time course of inhibition by A1899 using whole-cell and inside-out macropatch recordings from Xenopus oocytes. For these experiments, we tested inhibition of TASK-3 channels as these give rise to large macroscopic currents, necessary for inside-out macropatch recordings. In whole-cell recordings, the block of TASK-3 channels by A1899 developed very slowly. When the blocker was applied from the cytosolic phase during inside-out macropatch experiments, the block was faster by several orders of magnitude (Fig. 1F). These experiments suggest that A1899 acts as an open-channel blocker of TASK channels, presumably by binding to the inner cavity. If A1899 acts as an open-channel blocker, then the domains and the respective residues that interact with A1899 should form the walls of the central cavity. Hence, we continued to identify the drug binding site of A1899.

**A1899 Interacts with the M2 and M4 Transmembrane Segments of TASK-1 Channels**—As a first step toward identification of the drug binding site and the pore-forming transmembrane domains, we constructed chimeric channels between the drug-sensitive TASK-1 and the drug-insensitive TASK-4. We replaced one of the TASK-1 transmembrane segments at a time by a TASK-4 transmembrane domain (Fig. 2), expressed the chimeric channels in oocytes, and determined the IC_{50} values of A1899 (Fig. 2). Replacing the M1 or M3 segment of TASK-1 with TASK-4 did not cause a major reduction in the binding affinity of A1899. The IC_{50} values were 0.06 µM for the M1 exchange and 0.07 µM for the M3 exchange (Fig. 2), whereas the wild-type IC_{50} was 0.035 µM. In contrast, replacing the M2 or M4 transmembrane segments of TASK-1 increased the IC_{50} values of A1899 by several orders of magnitude (IC_{50} > 10 µM) (Fig. 2). We conclude from these findings that the M2 and M4 segments of TASK-1 are pore-forming domains that are crucial for the binding of A1899 and the subsequent block of the channel.

**The Halothane Response Element Is Part of the A1899 Binding Site**—Volatile anesthetics activate TASK-1 and TASK-3 channels via the halothane response element (HRE). Therefore, we tested A1899 on wild-type TASK-1 and a mutant TASK-1 channel with a deletion of the HRE sequence, [241VLRFMT] (indicated by a box). In both channels, in-frame deletion of the HRE (bottom) leads to a massive increase in IC_{50} values. rTASK, rat TASK-B, dose-response curves of A1899 on human TASK-1 WT (■) and an M247L (▲) mutant.

**FIGURE 2.** M2 and M4 segments of TASK-1 are part of the binding site for A1899. Four different chimeras, with TASK-4 transmembrane segments introduced into a TASK-1 background replacing M1, M2, M3, or M4, were analyzed for their IC_{50} of block by A1899. hTASK, human TASK.

**FIGURE 3.** The HRE influences A1899 binding. A, the sequence of the HREs of TASK-1 and TASK-3 differ in one amino acid (VLRF/M/LT) (indicated by a box). In both channels, in-frame deletion of the HRE (bottom) leads to a massive increase in IC_{50} values. rTASK, rat TASK-B, dose-response curves of A1899 on human TASK-1 WT (■) and an M247L (▲) mutant.
The A1899 binding site includes residues of the M2 and M4 segment and the selectivity filter. A–F, mutant TASK-1 channels were expressed in Xenopus oocytes, and block by 400 nM A1899 was analyzed. A and D, sample current traces for wild-type TASK-1 and mutated channels before (black) and after (gray) application of 400 nM A1899. B, C, E, and F, mutant TASK-1 channels with a reduced sensitivity to A1899 indicate residues as part of the binding site for A1899 (illustrated as striated bars). Threonine residues of the P1 and P2 pore loops are part of the A1899 binding site. In the M2 and M4 segment, residues of the binding site are located at a distance of 3–4 amino acids.

for wild-type TASK-1 to 117 nM (Fig. 3B), reflecting a 3.3-fold increase in IC₅₀ by converting the amino acid sequence to that of the TASK-3 HRE. The experiments indicate that the HRE is part of the A1899 drug binding site and that the sequence variations in the TASK-1 and TASK-3 halothane response elements might contribute to the different drug affinities of TASK-1 and TASK-3.

A1899 Is a Pore Blocker Binding to Threonine Residues of the Signature Sequence—Our data described above indicate that A1899 blocks TASK-1 channels from the intracellular side and that the M2 and M4 segments including the HRE are involved in A1899 binding. For different Na⁺, Ca²⁺, and K⁺ channels, it has been described that threonine residues of the signature sequence directly face into the central cavity and interact with open-channel blockers (18–21). If A1899 is in fact binding in the central cavity formed by M2 and M4 segments, the drug could interact with the corresponding threonine residues of the signature sequence to plug the channel pore. The positions of the analogous threonine residues (bold) are Val₂¹² and Ile₂³⁵ in the P2 loop and Thr⁹² and Thr⁹³ in the P1 region (Fig. 4B). Leu¹₂² and Thr¹₉⁹ of the M2 segment (Fig. 4C), Thr¹₉⁸ and Thr¹⁹⁹ of the P2 region (Fig. 4E), Ile²³⁵, Gly²³⁶, Leu²³⁹, and Asn²⁴⁰ of the M4 segment (Fig. 4F), and Val²⁴³ and Met²⁴⁷ of the HRE (Fig. 4F). The alanine scan revealed a helical pattern for the A1899 binding site in the M2 and M4 segment, meaning that substitutions by alanine that disturbed block were located in distances of about 3–4 amino acids. Consequently, the amino acids identified as parts of the binding site all point to the same direction of the respective α-helical transmembrane segment. As the identified residues should interact with the drug in the central cavity, our data can be used to validate the rotation of the M2 and M4 segment in a pore homology model.

To confirm the residues identified by our alanine scan, we introduced multiple mutations of the drug binding site simultaneously in the TASK-1 channels and tested these channels for their A1899 sensitivity (Fig. 5). Fig. 5A shows currents of the L122A/I235A and L122A/I239A double mutations and the L122A/I235A/M247A triple mutation, which all showed little or almost no block by 400 nM A1899. As expected, other TASK-1 constructs harboring multiple mutations at the A1899 binding site also showed a strong reduction in block (Fig. 5B). Summarizing, we have identified the residues of the A1899 binding site and conclude that the M2 and M4 segments build the pore of TASK-1 channels and that these amino acids forming the binding site of A1899 line the inner face of the pore.
Thr93 (cyan) and Thr199 (black) of the first and second pore loop, Ile118 (magenta) and Leu122 (yellow) of the M2 segment, Ile235/Gly236 (green/blue), Leu239/Asn240 (orange/white) of the M4 segment, and Val243 (sand) and Met247 (red) of the HRE element are facing into the central cavity (Fig. 6) or are, in the case of Met247, accessible on the way into the pore (Fig. 6). Thus, the A1899 binding site data validate our in silico TASK-1 pore homology model.

Confirmation of the A1899 Binding Site by Docking of the Drug to the Open Pore—A proportional ligand molecule was used to perform docking experiments. Dockings of the ligand molecule into the open state of TASK-1 yielded optimal binding scores. When bound, the drug clearly blocks the channel pore and impedes passage of K⁺ ions. In this conformation, the difluorobenzene end of the molecule lies closest to the selectivity filter, with the carbonyl oxygen of the amide linkage suggested to accept hydrogen bonds from residues of the signature sequence, Thr93 and Thr199 (Fig. 7, dotted lines). The majority of the other ligand-protein contacts are hydrophobic in nature, with Ile118, Leu122, Leu239 (the latter mutant did not express in oocytes), Ile235, and Leu239 all within 4 Å of the first three rings of the molecule. The fourth ring was found to be localized lower in the cavity toward the cytosolic entrance. At this position, the ring forms contacts with Leu122 and Val243, whereas the attached methoxy moiety could potentially form hydrogen bonds with Asn240. No interaction of the drug was predicted for Met247, which was located too low in the M4 segment. For a table listing the nature of the drug molecule interactions, see supplemental Fig. S2. It is noteworthy that the majority of the sites predicted by the docking experiments to the open state are in good agreement with our electrophysiological data as the model mostly highlights the same positions as the mutagenic screen (Figs. 4 and 6). Thus, the docking experiments confirmed the validity of the open-pore homology model and the data obtained by the alanine scanning approach.

A1899 Docking to a Closed-state Homology Model Supports a Binding to the Open State—Next, we compared the docking of A1899 to the open state (Fig. 7) with dockings to a closed-state model of TASK-1, which was generated on the basis of the KcsA structure (supplemental Fig. S3). Using the software GOLD, binding to the closed state was only possible if the M2 side chain and Phe125 were rotated so that this residue pointed up, toward the selectivity filter. This facilitates a strong binding mode that is in good agreement with the binding observed in the open-state model. In addition, the drug adopts the same orientation as in the open state. However, in this case, Phe125 would strongly interact with the drug, which is in disagreement with our experimental data (supplemental Fig. S3). Using the docking software AutoDock, the drug does not fit into the binding site, despite the displacement of Phe125 (as done for GOLD). In addition, using AutoDock, enabling a more flexible binding site
does not appear to improve the results. The experiments suggest that A1899 does not fit into a closed channel pore and that the channels have to open to allow an optimal binding of A1899.

**MD Simulations of A1899 in the Open Pore of TASK-1 Channels**—To test the binding of the drug molecule in more detail, MD simulations were performed to assess the dynamic interactions of the open-channel complex. Throughout all four of the 30-ns simulations, the complex remained intact, with the protein stabilizing to a typical Cα root mean square deviation of between 3 and 3.5 Å. The percentage of contacts, for each protein side chain within 4 Å of the drug molecule, was calculated at 10-ps intervals for all four simulations. Most contacts were observed for Thr92 and Thr93 of P1, Ile118 and Leu122 of M2, Thr198 and Thr199 of P2, and Ile235, Gly236, Leu239, and Asn240 of M4 (Fig. 8). Thus, overall, the majority of contacts remained the same as described for the original docking (Fig. 7 and supplemental Fig. S2). As a putative false positive, the MD simulations predicted an interaction with the residue Gln126, a residue that was not tested positive by our alanine scan. Similar to the initial docking experiments, the MD simulations did not predict an interaction with Phe125. The MD simulations (Fig. 8) and the original docking (supplemental Fig. S2) predict an interaction with Leu232. Unfortunately, introducing mutations at Leu232 led to non-functional channels. The MD simulations indicate a hydrogen bonding between drug and the side chains of Thr93 and Thr199 of the selectivity filter and Asn140 of the inner transmembrane helices. In addition to the H-bonding with Thr93 and Thr199, the uppermost carbonyl oxygen of the drug also appears to coordinate the K+ ion. Moreover, in two simulations tested, the ions starting at the S3 site move down to S4 to interact with the ligand (Fig. 8). It is noteworthy that open-channel blockers have been previously reported to interact with the K+ ion at site 4 of the selectivity filter (20–23). At depolarized potentials, when TASK-1 is open, the K+ ion seems to favor the occupancy at site 4 (22), which in turn appears to favor binding of the open-channel blocker A1899.

**DISCUSSION**

In recent years, K2p channels have been considered to represent promising drug targets. However, little is known about the structure of these channels, and in addition, there is a lack of specific blockers. We have identified A1899 as the most potent synthetic K2p channel blocker currently known. After showing that A1899 acts from the cytosolic phase as a pore blocker, we used a combined approach of chimeras and alanine scanning to identify the first drug binding site of a K2p channel. K2p channels have a unique transmembrane topology. Crystal structures of ion channels with different transmembrane topologies, including for example KcsA (24), rKv1.2 (25), and KirBac1.1 (26), suggest that the transmembrane segments following the selectivity filter are pore-forming inner helices. Our data support the idea that the M2 and M4 segments of K2p channels form the wall of the inner cavity. In addition, the interaction between the drug located in the central cavity and the amino acids of the M2 and M4 segments was assessed experimentally to provide data analyzing the rotation of the inner helices.

While we were in the process of finalizing our study, Kollewe et al. (14) proposed a structure for the Drosophila K2p channel dORK. The group utilized long range electrostatic compensation of positively charged amino acids in the selectivity region by negatively charged residues introduced into the M2 or M4 segment to identify pore-facing amino acids. This approach was used before in Kir2.1 channels by Chatelain et al. (27), who showed that charge and distance between the two mutations are decisive for a rescue effect and that electrostatic rescue does not work for residues that are located in close proximity or in great distance from each other. Because of this limitation, Kollewe et al. (14) could only test amino acids in a restricted region of the lower M2 and M4 segment. They probed residues corresponding to 119–127 of the TASK-1 M2 segment and 233–241 of the TASK-1 M4 segment (supplemental Fig. S4), whereas our more comprehensive approach scanned the whole length of those segments: 108–128 of the M2 and 227–248 of...
the M4 segment. Strikingly, five of the eight amino acids of dORK (Gly\textsuperscript{134}, Leu\textsuperscript{257}, Val\textsuperscript{258}, Met\textsuperscript{261}, and Thr\textsuperscript{262}), identified by Kollewe \textit{et al.} (14) as pore-facing, were also identified by our alanine scan. These residues correspond to the TASK-1 residues Leu\textsuperscript{122} in M2 and Ile\textsuperscript{235}, Gly\textsuperscript{236}, Leu\textsuperscript{239}, and Asn\textsuperscript{240} in M4 (supplemental Fig. S4). The data of dORK and TASK-1 can only be partially compared. In addition to the limited region of overlap, the mammalian TASK-1 channel shares only little sequence homology with dORK, and the structures of the two channels might differ as well. For example, we did not identify the M2 residues Thr\textsuperscript{121}, Phe\textsuperscript{125}, and Gln\textsuperscript{126} to be pore-facing as expected from the study with dORK (supplemental Fig. S4). In contrast, we identified Ile\textsuperscript{118}, a residue that was not in the scanning range of the electrostatic interaction approach, as pore-facing (supplemental Fig. S4). In addition, we identified two residues in the late M4 segment (HRE), namely Val\textsuperscript{243} and Met\textsuperscript{247}, which were also not within the spatial range for electrostatic interactions (supplemental Fig. S4).

The M2 and M4 segments of TASK-1 share low sequence homology (supplemental Fig. S4). Thus, not as expected from the primary sequence, the TASK-1 drug binding site has a remarkably high symmetry (Fig. 6). The primary binding site in the central cavity is formed by three rings of four identical residues. The first ring, located at the selectivity filter, is formed by four threonine residues, including two Thr\textsuperscript{93} residues of the P1 loop and two Thr\textsuperscript{199} residues of the P2 loop. The Ile\textsuperscript{118} of M2 together with Ile\textsuperscript{235} of M4 form a ring of four isoleucines, located underneath the selectivity filter in the central cavity. One helix turn below, there is a ring of four leucine residues formed by Leu\textsuperscript{122} of M2 and Leu\textsuperscript{239} of M4. In addition to these residues of a four-fold symmetrical binding site, we have identified Gly\textsuperscript{236} and Asn\textsuperscript{240} in the M4 segment and Val\textsuperscript{243} and Met\textsuperscript{247} in the HRE, which appear only twice in the binding site.

Interestingly, the HRE is a molecular prerequisite for the high-affinity block of A1899. This part of the channel is known to be the site of action for volatile anesthetics that activate TASK channels, and it has been described as a domain essential for the inhibition via G\textsubscript{q}-coupled receptors. However, it was not possible to identify single amino acids responsible for the action of anesthetics or the G-protein coupling within this region. In contrast, our alanine scan points out that two residues of the HRE, Val\textsuperscript{243} and Met\textsuperscript{247}, are responsible for the binding of A1899 to the HRE. Although the docking of A1899 to the pore homology model predicts an interaction of the HRE residue Val\textsuperscript{243} with the drug, Met\textsuperscript{247} seems to be too remote from the final binding site. Nevertheless, our mutagenesis data suggest that the difference in amino acid sequence at position 247 of the haloethane response element contributes to the different drug affinities of TASK-1 and TASK-3. An explanation is that Met\textsuperscript{247} is important for a trapping of the drug in the central cavity. Our 30-ns MD simulations, however, showed no evidence for a trapping of the drug through gate closure at Met\textsuperscript{247}, although this process likely requires MD simulations on a longer timescale. A more likely explanation lies within the fact that Met\textsuperscript{247} is located in the access pathway into the central cavity (Fig. 6). We suggest that position 247 is not part of the binding site but that it influences drug sensitivity because it regulates accessibility of the pore for A1899. The role of the HRE in high-affinity drug binding presented in this study suggests that the mechanism of synthetic modulators and endogenous mechanisms of TASK-1 regulation might be similar. Analogous actions of drug binding and endogenous G protein-coupled receptor modulation of TASK-1 might be the focus of future studies.

Our alanine scan does not suggest any impact of the F125A mutation on drug binding. The position Phe\textsuperscript{125} is analogous to Phe\textsuperscript{656} in human ERG channel, which is essential for the binding of most ERG blockers (19, 28). Former studies propose the corresponding amino acids in dORK (Phe\textsuperscript{137}) and TREK-1 (Leu\textsuperscript{174}) to line the inner pore (14, 29). However, in our computer models for the open state, Phe\textsuperscript{125} does not line the pore. An interaction of this residue with A1899 is only suggested in the closed-state model (supplemental Fig. S3). We therefore propose that the M2 segment is slightly rotated in the open state as compared with the closed-state, leading to a displacement of the aromatic residue of Phe\textsuperscript{125}. The importance of the exact orientation of the corresponding phenylalanine in human ERG has also been discussed by Chen \textit{et al.} (28).

Based on their TREK-1 model, Treptow and Klein (29) suggest that two leucines, one in the M2 and one in the M4 segment of TREK-1 (Leu\textsuperscript{174} and Leu\textsuperscript{289}), delimit the narrowest part of the inner pore and form a hydrophobic gate. These leucine residues align to Phe\textsuperscript{125} and Leu\textsuperscript{239} of TASK-1. Our data indicate that TASK-1 channels do not have the same or conserved hydrophobic gating ring. The hydrophobic gating ring might be the molecular correlate to the difference between the gating mechanisms of TREK-1 and TASK channels.

The selective TASK-1 blocker A1899 described here is a powerful tool for studying the physiological role of TASK-1 channels in native tissues. We took advantage of the blocker to probe the direct interaction with the amino acids of the inner pore and created the first model for a mammalian K\textsubscript{2p} channel with experimentally validated orientation of the inner helices. Thus, using A1899 as a chemical probe, we validated a model for the mammalian K\textsubscript{2p} channel TASK-1 in the open state and identified the first complete drug binding site in a K\textsubscript{2p} channel. The docking of A1899 into this open-pore homology model and the subsequent molecular dynamics simulations confirmed the amino acids of the drug binding site identified by our alanine scan. Our data provide an excellent starting point for future studies on the molecular pharmacology of other K\textsubscript{2p} channels. In addition, our data should influence the rational drug design of selective blockers of other K\textsubscript{2p} channels. As K\textsubscript{2p} channels have emerged as novel drug targets, the identification of a TASK-1-specific blocker together with the identification of a high-affinity drug binding site in a K\textsubscript{2p} channel is a promising finding for future drug design and for basic research in physiology.

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