Structural bases for the different anti-fibrillatory effects of chloroquine and quinidine

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Aims
Chloroquine, an anti-malarial quinoline, is structurally similar to quinidine. Both drugs have been shown to block ion channels. We tested the hypothesis that chloroquine's mode of interaction with the vestibule of the cytoplasmic domain of the inward rectifier potassium channel Kir2.1 makes it a more effective Ik1 blocker and anti-fibrillatory agent than quinidine.

Methods and results
We used comparative molecular modelling and ligand docking of the three-dimensional structures of quinidine and chloroquine in the intracellular domain of Kir2.1. Simulations predicted that chloroquine effectively blocks potassium flow by binding at the centre of the ion permeation vestibule of Kir2.1. In contrast, quinidine binds the vestibular side, only partially blocking ion movement. We tested the modelling predictions in Kir2.1-expressing human embryonic kidney (HEK)-293 cells. The half-maximal inhibitory concentration for chloroquine block of Ik1 was 1.2 μM, while that of quinidine was 57 μM. Finally, we used optical mapping of Langendorff-perfused mouse hearts with cardiac-specific Kir2.1 up-regulation to compare the anti-fibrillatory effects of the drugs. In five of six hearts, 10 μM quinidine slowed the frequency but did not terminate the tachyarrhythmia. In five of five hearts, 10 μM chloroquine terminated the arrhythmia, restoring sinus rhythm.

Conclusion
Quinidine only partially blocks Ik1. Chloroquine binds at the centre of the ion permeation vestibule of Kir2.1, which makes it a more effective Ik1 blocker and anti-fibrillatory agent than quinidine. Integrating the structural biology of drug-ion channel interactions with cellular electrophysiology and optical mapping is an excellent approach to understand the molecular mechanisms of anti-arrhythmic drug action and for drug discovery.

Keywords
Inward rectifier potassium current • Chloroquine • Quinidine • Fibrillation

1. Introduction
The pharmacologic treatment of cardiac fibrillation is inadequate. Arguably, this is in part because of our relatively incomplete understanding of the mechanisms of arrhythmias, and our poor knowledge of the molecular bases of drug effects. Recent advancements in the structural biology of membrane ion channel proteins provide new opportunities to better understand drug-ion channel interactions and hopefully improve anti-arrhythmic therapy.

Quinidine is a class-1 anti-arrhythmic quinoline-type drug that is only partially effective as an anti-fibrillatory agent, although recently it had a resurgence as being potentially useful in the prevention of arrhythmias associated with the short QT syndrome (SQTS).

Here, we aimed at shedding light on the molecular bases on quinidine’s anti-arrhythmic action by comparing its effects with those of the anti-malarial chloroquine, a related quinoline, used briefly in the past to treat arrhythmias in patients. More recently, chloroquine was shown to be a potent blocker of the strong inward rectifier potassium channel Kir2.1, which is the major carrier of Ik1 in mammalian ventricles. It was also demonstrated to effectively terminate fibrillatory activity in three different experimental models. On the other hand, the gain of function mutations in Kir2.1 has been shown to result in both SQTS type 3 and familial atrial fibrillation (AF) in humans.

The three-dimensional crystal structure of the intracellular domain of Kir2.1 has been solved. It has also been shown that amino acids
E224, D259, E299, F254, and D255 of the Kir2.1 intracellular domain, which form a highly electronegative ring, are critical for chloroquine’s blocking action. This new information allowed us to directly compare the structural and electrophysiological properties of chloroquine-mediated Kir2.1 block to those of quinidine. Chloroquine and quinidine are closely related in their chemical structure; both have a quinoline ring base but different functional groups. We tested the hypothesis that chloroquine is a more effective blocker of IC\textsubscript{50} than quinidine because the geometry of the intracellular ion-permeation pathway in the Kir2.1-chloroquine complex is markedly different from that of Kir2.1:quinidine due to the dissimilarities in the functional groups of the drugs.

We employed a numerical and experimental approach integrating protein structure, single cell, and whole organ electrophysiology. We compared chloroquine’s specific mode of interaction with the vestibule of the Kir2.1 cytoplasmic domain and the resulting geometry of the vestibule—chloroquine complex with those of quinidine.

We substantiated the molecular modelling predictions using patch clamping to compare the effects of chloroquine with those of quinidine on BA\textsuperscript{2+} sensitive currents in human embryonic kidney (HEK)-293 cells transiently expressing Kir2.1. Finally, we used isolated Langendorff-perfused hearts from transgenic mice with cardiac-specific over-expression of Kir2.1 to compare the anti-arrhythmic actions of the two drugs.

2. Methods

2.1 Molecular modelling

Numerical studies were based on the co-ordinates for chloroquine and quinidine developed in the small molecule topology generator PRODRG and the cytoplasmic domain of Kir2.1 (PDB ID: 1U4F). Docking simulations were performed in Autodock 4.0.1 employing the genetic algorithm for 10 runs with a population of 150. Prior to these calculations, the ionization of the ligands was adjusted for physiological pH. The same protonated protein model, grid size, and grid placement were used for all docking studies, and the ligands were initially positioned adjacent to the protein model outside of the channel centred at (0,0,0). During calculations, the protein was held rigid and the ligands were allowed to flex. The resulting complexes were ranked according to the binding energy. The complex with the lowest binding energy was further refined by slow-cooled simulated annealing followed by minimization in Crystallography and the nuclear magnetic resonance (NMR) system program. During the simulated annealing runs, both protein and ligand were flexible. The complex was heated to 1000 K and then cooled to 300 K in 25 K steps with a surrounding dielectric constant of 80. Solvent accessible surface areas for the complexes were calculated using Access from CCP4 (The CCP4 Suite: Programs for Protein Crystallography) and Solvent accessible surface maps of the complexes were calculated using a 1.4 Å probe and displayed in Pymol with the APBS plug-in (PyMOL Molecular Graphics System 2008; DeLano Scientific, Palo Alto, CA, USA).

2.2 Experimental

2.2.1 Molecular biology and patch clamping

Kir2.1 cDNA was subcloned into pcDNA3.1(+) plasmid (Invitrogen) and expressed in HEK-293 cells (ATCC, Virginia). Macroscopic currents were recorded as detailed earlier and Ba\textsuperscript{2+}-sensitive (2 mM) current voltage (IV) relationships were constructed. For the determination of the half-maximal inhibitory concentration (IC\textsubscript{50}) for the IC\textsubscript{50} block by chloroquine (0.3–10 μM) or quinidine (3–1000 μM), the fractional block of current was plotted as a function of drug concentration ([D]) and the data fit with the Hill equation: 1/[1 – (IC\textsubscript{50}/[D])\textsuperscript{m}]. Analysis of membrane currents through the action potential (AP) voltage clamp protocol was performed in HEK-293 cells by using the AP recorded from isolated mouse ventricular myocytes as the command signal. Currents elicited in response to the AP voltage clamp were recorded under control conditions, and in the presence of 10 μM chloroquine or quinidine. Kir2.1 currents are represented as Ba\textsuperscript{2+}-sensitive currents elicited by the AP voltage clamp. Alanine mutagenesis of Kir2.1 residues was conducted as detailed earlier and the current at −50 mV was recorded in transfected HEK cells in response to voltage steps from −80 mV to −50 mV, and then to −100 mV, in the absence or presence of 100 μM quinidine.

2.2.2 Optical mapping

All animal studies conformed to the Guide for the Care and Use of Laboratory Animals published by the US National Institutes of Health (NIH Publication No. 85-23, revised 1996) and to the University of Michigan Committee on the Use and Care of Animals (protocol 09985). Adult, Kir2.1-overexpressing transgenic mice were heparinized and anaesthetized. The excised heart was Langendorff-perfused with warm (36 ± 1°C), and oxygenated Tyrode’s solution, and optical mapping was carried out as described earlier. Recording rate: 600–1000 frames per second, and resolution: 109 μm.

2.2.3 Arrhythmia induction

To induce ventricular tachycardia/fibrillation (VT/VF), we used a bipolar, silver tip stimulation electrode placed at the apex of the heart to deliver short bursts of 3 ms square pulses at a cycle length of 10–20 ms.

2.2.4 Dominant frequency maps

We used fast Fourier transform analysis on each pixel of the epicardial optical signals to generate dominant frequency (DF) maps and constructed maps of spatial DF distribution as previously described.

2.2.5 Statistics

Results are mean ± standard deviation. ANOVA and Fisher’s exact and t-tests were used as appropriate. Significance was at P < 0.05.

3. Results

3.1 Chloroquine and quinidine interact differently with the tetrameric Kir2.1 intracellular domain

Pegan et al. elucidated the three-dimensional crystal structure of Kir2.1. Since there is no crystal or NMR structure of Kir2.1 or any family member complexed with a drug-like molecule, Rodriguez-Menchaca et al. and Noujaim et al. used modelling to predict the interaction of chloroquine and Kir2.1. Here, we compared the docking of chloroquine and quinidine in the ion permeation of the tetrameric Kir2.1 intracellular domain. Chloroquine and quinidine are structurally related, in that they both have a quinoline ring base. However, their functional groups are different. Consequently, as shown in Figure 1, modelling studies revealed that chloroquine and quinidine have different modes of binding to the Kir2.1 intracellular domain. In Figure 1A, chloroquine represented in green sticks, acts as a plug, fitting in the centre of the tetrameric channel. The drug is suspended in a band of negative charges formed by the acidic amino acids E224, D259, and E299. As shown in Figure 1C, E224 forms charge–charge interactions with chloroquine. D259 is within 4 Å of the drug, and is involved in van der Waals interactions. H226 and R260 form hydrogen bonds with the protonated nitrogens of chloroquine, and F254 from three subunits provide a hydrophobic...
base for the molecule. E299 is located >5 Å from the chloroquine molecule. Quinidine, yellow sticks in Figure 1B and D, binds in the same zone of negative charge. However, it forms hydrogen bonds and van der Waals interactions with specific residues from two subunits of Kir2.1 (Figure 1D), causing it to be off-centred (in contrast to the centred position of chloroquine in Figure 1A and C) and drawn to one side of the channel (Figure 1B and D). The carboxylate of D259 from one subunit forms a hydrogen bond with the protonated quinoline ring of quinidine, while the carboxylate from E224 of an adjacent subunit forms hydrogen bonds with the caged nitrogen. In addition to the hydrophilic interactions, F254 from two subunits (the unlabelled pink and blue residues underneath quinidine in Figure 1D) form hydrophobic interactions with the quinoline ring and the carbon cage along with the acyl chain of R260 from two subunits. Due to the four-fold symmetry of Kir2.1, the top four docking results were equivalent, simply separated by a 90° rotation about the channel.

3.2 Chloroquine and quinidine have different effects on the geometry of the aqueous vestibule of Kir2.1

To unveil the consequences of the different interactions of chloroquine and quinidine with the channel, solvent accessible areas and maps were calculated using CCP4 and Pymol, respectively (Figure 2). Maps were constructed by probing the channels with a probe of 1.4 Å in radius, corresponding to the radius of a water molecule. The probe traces the water accessible regions of the channel. The resulting shell structure is three-dimensional, but for visualization, a slab was shown to expose the shells’ insides. The yellow colours denote the exterior of the ‘solvent accessible shell’ and black is the inside of the shell’s cavity or the ion-permeation pathway. The figure shows differences in the cavity’s geometries depending on whether chloroquine or quinidine was bound to the ion-permeation pathway. The total surface area decreased by 162 and 217.5 Å² upon binding of quinidine and chloroquine, respectively. The surface maps defining the channel show that Kir2.1 has a cavity that runs through the centre of the tetramer, but the channel is disrupted upon compound binding. Chloroquine causes a break in the ion pathway (Figure 2 white broken lines), while quinidine partially blocks the channel, but allows room for ions to still pass through albeit at a slower pace (Figure 2 white arrow). It is possible for more than one molecule of quinidine to bind to the electronegative band in the cytoplasmic tail of Kir2.1. However, the restricted space imposes an extremely tight fit, rendering this scenario unlikely.

3.3 Chloroquine is a more potent blocker of IK₁ compared with quinidine

Molecular modelling predicted that chloroquine was a better blocker of IK₁ than quinidine. We tested this prediction using patch clamping. Figure 3 shows whole-cell current traces in Kir2.1-expressing HEK-293 cells before and after 10 μM chloroquine (Figure 3A) or 10 μM quinidine (Figure 3C). The IV curves illustrate the ability of chloroquine to block most of the outward component of IK₁ (n = 5) (Figure 3B), whereas the effect of 10 μM quinidine was much less pronounced (n = 5) (Figure 3D). Figure 3E shows that the fraction of blocked outward current at −60 mV in Kir2.1-expressing HEK-293 cells by
chloroquine was $0.90 \pm 0.03$ vs. $0.09 \pm 0.2$ in response to quinidine ($n = 5$, *P < 0.01). In Figure 3F, we examined the whole-cell $I_{K1}$ current in Kir2.1-expressing HEK-293 cells in response to a mouse AP clamp protocol. The top traces are voltage-clamp profiles and the bottom traces are the control and chloroquine- (left) and quinine- (right)-sensitive currents. The arrows indicate the maximum outward $I_{K1}$, which was markedly reduced by $10 \mu M$ chloroquine, compared with the more subtle effect of $10 \mu M$ quinidine. Figure 3G shows the dose–response curves of the fractional blocked peak outward current in response to the AP voltage clamp with chloroquine and quinidine. Solid lines are Hill equation best fits. IC$_{50}$: chloroquine = $1.2 \pm 0.2 \mu M$, quinidine = $57 \pm 3.8 \mu M$. Chloroquine is 48 times more potent as an $I_{K1}$ blocker than quinidine.

### 3.4 Chloroquine but not quinidine terminates $I_{K1}$-mediated tachyarrhythmias in the mouse heart

Up-regulation of $I_{K1}$ in the mouse heart via cardiac-specific Kir2.1 over-expression leads to stable and fast re-entrant VT/VF. Here, we compared the effects of chloroquine with those of quinidine on VT/VF in this mouse model. On the basis of the molecular simulations, and the patch clamp experiments, we postulated that at the same dose, intracoronary perfusion of $10 \mu M$ chloroquine, but not quinidine, should terminate VT/VFs and restore sinus rhythm, even though other currents, such as $I_{Na}$, are affected by both chloroquine and quinidine.

We then proceeded to evaluate the anti-fibrillatory properties of quinidine compared with chloroquine in six mouse hearts. VT/VF was induced by burst pacing the apex of the heart. Figure 4A, top row, is $DF_{\text{max}}$ maps showing the frequency of the tachyarrhythmia during the course of 30 min perfusion of $10 \mu M$ quinidine. The tracings below each map are 500 ms single pixel recordings obtained from the same location. At the onset, the frequency was 48 Hz. It decelerated progressively to 28 Hz after 30 min of quinidine. However, sinus rhythm was not restored. In the bottom row of Figure 4A, after a washout of 20 min, the arrhythmia accelerated to 46 Hz. Thereafter, chloroquine perfusion started, and after 7 min, the tachyarrhythmia slowed to 23 Hz, and sinus rhythm resumed. Figure 4B shows a composite data from six experiments. The time

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**Figure 2** Stereoviews of solvent accessible surface maps of the intracellular pore of Kir2.1 in complex with chloroquine and quinidine. Views oriented with respect to a $90^\circ$ rotation of Figure 1A and B. Kir2.1 cytosolic side is on the left in each figure. Outer surfaces: yellow; cavities: black. Slabbed surfaces reveal the cavity of the inner vestibule of the intracellular domain of Kir2.1. (A) Surface map for Kir2.1. Surface map of Kir2.1 in complex with chloroquine (B) and quinine (C). Orange arrow: ion-permeation pathway in Kir2.1 alone. Broken white lines: break in the ion-permeation pathway. White arrow: narrowing of the ion-permeation pathway.
elapsed between individual data points after the application of 10 μM quinidine or chloroquine is 1 min. With quinidine, tachyarrhythmias terminated in one heart (filled symbol) at 8 min of quinidine perfusion but not in five other hearts even after 30 min of continuous quinidine perfusion (open symbols). In these hearts, the DF max decreased by a factor of 0.6 ± 0.1 without stopping, and after a washout period of 20 min, the DF max recovered back to 0.94 ± 0.05 of control. We then perfused five of these hearts with 10 μM chloroquine. The VT/VF frequency reduced by a factor of 0.55 ± 0.035, and sinus rhythm was restored after an average time of 8 min of chloroquine perfusion. Chloroquine restored sinus rhythm in five of five hearts, while quinidine did so in one of six hearts (χ² = 0.015, Fisher’s test).

Since 10 μM is below the IC₅₀ for quinidine block of IK₁ (Figure 3G), we tested the effects of 60 μM quinidine on induced tachyarrhythmias in three isolated hearts. This high dose of quinidine terminated the sustained tachyarrhythmia; however, in all hearts, the restored sinus beats were repeatedly interrupted by spontaneous, non-sustained bursts of VT/VF. On the other hand, hearts perfused with chloroquine were in normal sinus rhythm after chemical defibrillation. Figure 3A and B shows examples of 10 s electrocardiogram (ECG) runs illustrating that while 10 μM chloroquine restored sinus rhythm, 60 μM quinidine was pro-arrhythmic.

### 3.5 Mutation of specific residues suggested by modelling to be involved in quinidine’s interaction with Kir2.1 makes the channel less susceptible to block

Figure 5C and D shows that amino acids E224, F254, and D259 proposed by the model to interact with chloroquine are experimentally involved in quinidine’s block of the channel. These residues have also been shown earlier to be important for chloroquine’s block of Kir2.1. In Figure 5C, we show representative currents recorded in HEK cells transfected with WT Kir2.1 or Kir2.1 in the presence or absence of 100 μM quinidine. Quinidine blocks the WT channel; however, in the presence of the F254A mutation, the ability of quinidine to block the outward current was reduced. Figure 5D is the quantification of the fraction of blocked...
chlordiazepoxide terminated VF/VT in a total of five of five mouse hearts. Quinidine terminated arrhythmias in one of six hearts (χ² = 0.015).

Integration of information, from the molecule to the organ, derived from the study of the effects of currently available drugs on cardiac electrical properties may help set the stage for the development of new, more effective and safer therapeutic approaches, with the ultimate goal of preventing sudden cardiac death. We have therefore conducted studies at three complementary levels of integration to investigate the anti-fibrillatory effects of chloroquine, an anti-malarial quinoline, which has been suggested to have anti-arrhythmic properties, in comparison with quinidine, another quinoline, widely used in the past.

Recently, quinidine was suggested to be beneficial in certain cases of short QT syndrome. In addition, quinidine was shown to have superior anti-arrhythmic properties in the treatment of VF in animal models of pharmacologically induced short QT compared with sotalol and flecainide because it prolongs refractoriness and reduces dispersion of repolarization. Similar to quinidine, chloroquine also prolongs the AP duration and refractoriness in the atria and ventricles by effectively blocking inward rectifier potassium currents, including IK1, the acetylcholine-activated potassium current (IK,ACH), and the ATP-dependent potassium current (IK,ATP). Our results using a transgenic mouse model of Kir2.1 overexpression demonstrate that the anti-fibrillatory effects of chloroquine are substantially greater than those of quinidine in IK1-mediated tachyarrhythmias. Those effects correlate strongly with the preferential ability of chloroquine to block IK1 when compared with quinidine. Similarly, by blocking other inward rectifier potassium currents, chloroquine was shown to terminate both VT/VF and AF in three different species, including mouse, rabbit and sheep.

It seems clear that the anti-fibrillatory effects of chloroquine in the Kir2.1-overexpressing mouse model are due in large measure to preferential IK1 blockade. IK1 is of interest because it plays a significant role in the stability and frequency of re-entrant VT/VF and in AF. From the clinical standpoint, IK1 is important because mutations in the Kir2.1 protein have been shown to be associated with at least two different inherited arrhythmogenic diseases in humans. However, one should not underestimate the potential contribution of chloroquine’s inhibition of other channels to the mechanism of VF termination. Compared with other inward and outward currents, chloroquine preferentially blocks IK1 (IK1 > IKr > IKCaL > IKs > IKur), but while similar comparisons on the relative effectiveness of inhibition on different ionic channels have not been made for quinidine, there is evidence that it also blocks IKCaL, IKCaL, IKr, IKr, IKs, and IK1.

IK1 is important in the control of the rate and frequency of functional re-entry. Experiments and numerical simulations demonstrated that IK1 mediates electrotonic interactions between the rotor core and its immediate surroundings. When IK1 is increased, the rate of the terminal phase of repolarization of the AP is accelerated, and the resting membrane potential is slightly hyperpolarized. The space constant is also decreased resulting in a smaller core size. The small core, combined with the short AP duration, reduces the likelihood of wave front-wave tail interactions, allowing a fast and relatively stable rotor. The opposite occurs when IK1 is reduced, such as by chloroquine or BaCl2, both of which lead to slowing and eventual termination of rotor activity.

Figure 4 Effects of chloroquine and quinidine on ventricular arrhythmia. (A) Top row: DF maps at the onset of pacing-induced VT (DFmax = 48 Hz), 4 min (35 Hz), 7 min (31 Hz), and 30 min (28 Hz) after starting 10 μM quinidine. VT slowed down but did not terminate. Bottom row: DF maps of VT after 20 min quinidine washout (DFmax = 46 Hz), then at 4 min (33 Hz) and 7 min (23 Hz), followed by sinus rhythm restoration (4 Hz), in the presence of 10 μM chloroquine. Single pixel recordings (500 ms) are shown below each map. (B) Plot of normalized DFmax in six hearts during VT. In five of six hearts (open symbols) 30 min of 10 μM quinidine slowed down but did not terminate VT/VF. In one heart tachyarrhythmia terminated (filled symbol). After 20 min of washout, 10 μM chloroquine applied to five tachyarrhythmic hearts terminated the arrhythmias and sinus rhythm resumed.

4. Discussion

We have taken advantage of knowledge derived from the solved crystal structure of the Kir2.1 intracellular domain to provide a structural basis for the electrophysiological effects of two different quinolines (chloroquine and quinidine) on IK1. Molecular modelling predicts that chloroquine should be a more effective IK1 blocker than quinidine. Although both compounds are envisaged to bind to the negatively charged ring within the vestibule, they have different binding modes, which alter the size of the ion pathway. While chloroquine introduces a break in the ion-permeation pathway, quinidine’s interactions with the channel results in a partial block. This was further verified experimentally by patch clamping and optical mapping. The IC50 of chloroquine’s block of IK1 was 1.2 μM while that of quinidine was 57 μM. Consequently, in a mouse model of IK1 up-regulation, current at −50 mV in the presence of E224A, F254A, or D259A mutations, which made the channel resistant to the quinidine block.
critical for chloroquine blockade. Noujaim et al.\textsuperscript{9} also showed that chloroquine interacts with electronegative residues in the intracellular domains of Kir2.1, Kir3.1, and Kir6.2, leading to $I_{K1}$, $I_{KAC}$, and $I_{KATP}$ blockade, and termination of arrhythmias mediated by these three inward rectifier currents in the mouse, rabbit, and sheep hearts. This study elucidates the structural basis for the different $I_{K1}$ blocking efficacies of chloroquine and quinidine. Similarly to the case of chloroquine, E224, D259, F254, H266, R260, and D255 are also critical for a partial block of Kir2.1 by quinidine. The atoms of E299 are not solvent accessible, therefore the role of E299 is most likely to maintain the structural integrity of the protein and to offset the positive charges of R228 and H226. Such residues help to form an electronegative band that interacts with the protonated quinoline ring and caged nitrogen of quinidine. Hence, both quinidine and chloroquine block Kir2.1 by interacting with the band of negative charge located in the inner vestibule of the intracellular pore.\textsuperscript{12} However, chloroquine is a more potent $I_{K1}$ blocker, and is thus more effective than quinidine in terminating $I_{K1}$-mediated re-entrant VT/VF.

Drug-ion channel interaction studies that span multiple levels of integration that include structure/function relationships, up to the whole heart may be a helpful strategy in furthering our understanding of current anti-arrhythmics, and the development of novel, potentially useful anti-fibrillatory pharmacophores.

4.1 Limitations

Molecular simulations cannot account for all the drug-channel interactions that can take place or the possible effects of modulatory agents, such as PIP2, on Kir channels.\textsuperscript{25} Crystallization and determination of the 3D structure of the drugs in complex with the Kir channels would be essential to understand all aspects of drug-channel interactions.

The mouse is a useful animal model. However, several differences exist between the assemblage of ion channels underlying the AP of the mouse compared with other species.\textsuperscript{26,27} Care should be taken in translating to the human, basic electrophysiological findings in the mouse. Chloroquine and quinidine are not specific $I_{K1}$ blockers. Their effects on other currents can contribute to the results we observe in the heart.

While reducing $I_{K1}$ may be antifibrillatory, there is also evidence that $I_{K1}$ loss-of-function, such as in Anderson–Tawil syndrome (ATS), may be proarrhythmic.\textsuperscript{28} However, whilst ATS patients may develop ventricular tachyarrhythmias, including torsades de pointes, sudden cardiac death is a very rare event.\textsuperscript{28–30} Does the above suggest that the ventricles of patients with ATS are incapable of undergoing stable, long-standing, functional re-entry? And does our study suggest that compared with chloroquine, quinidine’s relatively lower efficacy as an anti-fibrillatory agent is due to its relatively modest effect on $I_{K1}$? Further studies in animal models of ATS, together with pharmacologic analyses of the type presented here should help answer this question. Such knowledge, together with the new information derived from our study, may lead to novel integrative and multidisciplinary approaches in search of new anti-fibrillatory pharmacophores that increase the inward going rectification of $I_{K1}$, preferably in a cardiac-specific manner.

Conflict of interest: none declared.
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