Testing novel pharmacological strategies for the management of atrial fibrillation in a large animal experimental model

Summary of PhD Thesis

Viktor Juhász, MD

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# ABBREVIATIONS

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>APD&lt;sub&gt;25&lt;/sub&gt;, APD&lt;sub&gt;50&lt;/sub&gt;, APD&lt;sub&gt;90&lt;/sub&gt;</td>
<td>Action potential duration at 25, 50 and 90% of repolarization</td>
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<tr>
<td>AERP</td>
<td>Atrial effective refractory period</td>
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<td>AF</td>
<td>Atrial fibrillation</td>
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<tr>
<td>BCL</td>
<td>Basic cycle length</td>
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<td>ERP</td>
<td>Effective refractory period</td>
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<td>I&lt;sub&gt;K, ACh&lt;/sub&gt;</td>
<td>Acetylcholine activated potassium current</td>
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<tr>
<td>I&lt;sub&gt;Kr&lt;/sub&gt;</td>
<td>Rapid component of the delayed rectifier potassium current</td>
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<tr>
<td>I&lt;sub&gt;Kur&lt;/sub&gt;</td>
<td>Ultrarapid outwardly rectifying potassium current</td>
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<tr>
<td>I&lt;sub&gt;Na&lt;/sub&gt;</td>
<td>Voltage dependent sodium current</td>
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<tr>
<td>NFAT</td>
<td>Nuclear factor of activated T-cells</td>
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<td>TdP</td>
<td>Torsades de Pointes chaotic ventricular tachycardia</td>
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INTRODUCTION

Atrial fibrillation (AF) is the most frequently encountered chronic arrhythmia, associated with increased morbidity and mortality due to thromboembolic complications and concomitant heart failure. Its incidence and prevalence is rapidly increasing with the aging of the population and the number of cases will probably double by 2060 in the EU. There is a great unmet need for safer and more effective pharmacological AF therapy, since drugs currently used for rhythm control may significantly increase the risk for Torsades de Pointes arrhythmias due to their ventricular electrophysiological effects; can promote adverse vascular events; can exhibit reduced efficacy in persistent AF; and the most effective antiarrhythmic drug, amiodarone, has serious extracardiac side effects following chronic administration. In addition, as part of the pathological electrical remodeling accompanying AF, the expression of numerous ion channels and exchangers is altered that can modify the arrhythmia substrate and increase triggered activity resulting in AF to become self-sustaining, and also significantly altering potential drug targets.

One of the possible approaches to improve pharmacotherapy of AF is the identification of drug targets ideally expressed only in atrial tissue, since atrial selective ion channel modulation would be devoid of ventricular proarrhythmic adverse effects. The $K_{v}1.5$ channel that conducts the ultrarapidly activating outwardly rectifying $I_{Kur}$ represents such a target, since its block prolongs the atrial action potential duration (APD) and effective refractory period (ERP). However, data regarding changes in $I_{Kur}$ expression are inconsistent in animal experimental models of AF and human, with some clinical studies showing downregulation of the channel in patients with chronic AF. Another target with relative atrial selectivity is the acetylcholine-receptor activated inwardly rectifying $I_{K,ACh}$ ($K_{ir3.1}/K_{ir3.4}$), that is found in larger amounts in the atria than in the ventricles. Importantly, in contrast to the fact that $I_{K,ACh}$ downregulation was also observed in chronic AF, a constitutively active component of $I_{K,ACh}$ was detected in patients with persistent AF. Therefore, $I_{K,ACh}$ block represents a promising target for the treatment of AF. The selective inhibition of $I_{K,ACh}$ was shown to exhibit beneficial effects against AF in different animal models anesthetized with a combination of isoflurane and thiopental.
However, thiopental and isoflurane have well known ion channel modulating properties. Thiopental blocks $I_{Ca,L}$, $I_{K1}$, $I_{Ks}$ and isoflurane blocks $I_{Ca,L}$ and $I_{Ks}$, modulates mitochondrial calcium-activated $K^+$ channels. These different effects on cardiac ion channels can profoundly influence the outcome of atrial and ventricular arrhythmia studies depending on the anesthetic used, including thiopental and isoflurane.

Given the complex aetiology of AF, it has been suggested that drugs targeting multiple pathways involved in AF development may be more effective. Over the past fifteen years, studies on the cellular pathways in AF have revealed potential therapeutic targets to develop new antiarrhythmic drugs for AF management. In addition to the ion channels mentioned above, voltage-gated sodium channels, two-pore potassium channels, oxidative stress and activation of the transcription factor NFAT have all been implicated in AF development. To date, AF drugs have been exclusively targeted towards ion channels. However, non-ionic remodeling events are also likely to contribute to the initiation and maintenance of AF. It has been further suggested that targeting maladaptive remodeling events („upstream therapy”) in AF may also be required for effective control of AF.

**AIMS OF THE STUDIES**

1. To establish a large animal *in vivo* atrial fibrillation model in the Department of Pharmacology and Pharmacotherapy for the testing of potential novel drug candidates for AF management, and to provide remodeled atrial tissue for AF mechanism studies for *in vitro* investigations.

2. To test the effects of an atrial selective ion channel blocker in the established model on inducibility of AF, on the duration of AF episodes, on the right atrial effective refractory period (AERP) in conscious dogs with AF and investigation of the mechanisms of action on atrial trabeculae isolated form these animals.

3. To investigate experimental compounds with multiple mechanisms of action with parallel targeting of ion channels and cellular pathways implicated in AF in the chronic right atrial tachypacing induced conscious dog atrial fibrillation model.
METHODS

Ethical issues, experimental animals

The studies were conducted in compliance with the Guide for the Care and Use of Laboratory Animals (NIH Publication No. 85-23, Revised 1996), and the protocol was approved by the Ethical Committee for the Protection of Animals in Research of the University of Szeged, Hungary (I-74-5-2012) and by the Department of Animal Health and Food Control of the Ministry of Agriculture (XIII/1211/2012).

Establishment of the large animal model of chronic atrial tachypacing induced AF in conscious dogs in the laboratory

Male Beagle dogs weighing 12-16 kg were used for the experiments. The dogs were accommodated to experimental personnel and equipment, every day for a week before the start of the studies. The pacemaker and pacemaker electrode implantation procedures were performed under ketamine (induction: 10 mg/kg i.v., maintenance: 2 mg/kg, every 20 min) + xylazine (induction: 1 mg/kg, maintenance: 0.2 mg/kg, every 20 min) anesthesia. The implantations were carried out under proper antibiotic coverage as follows: amoxicillin/clavulanic acid (1000 mg/200 mg i.v.) and gentamicin (40 mg i.v.) given before the operation, amoxicillin/clavulanic acid (500 mg/125 mg orally, twice a day for 5 days; Augmentin 500mg/125mg®) given following the operation. For peri-operative analgesia metamizole sodium (1000 mg i.v., 1g/2ml) and tramadol (50 mg i.v.) were administered. Two bipolar pacemaker electrodes (Synox SX 53-JBP and Synox SX 60/15-BP) were positioned into the right atrial appendage and apex of the right ventricle, respectively, and the electrodes were connected to pacemakers (Logos DS and Philos S) in subcutaneous pockets in the neck area, followed by radiofrequency catheter ablation of the AV node. The pacemakers were programmed by the ICS 3000 Programmer (Biotronik Hungary Ltd.). Following recovery from surgery (3-5 days), right atrial tachypacing was started at 400 beats/min, maintained for 6 to 7 weeks before the experiments to allow electrical remodeling of the atria (monitored by the measurement of the right atrial effective refractory period (AERP) every second day). The AERPs were measured at basic cycle lengths (BCL) of 150 and 300 ms with a train of 10 stimuli (S1)
followed by an extrastimulus (S2), with the AERP defined as the longest S1-S2 interval that did not produce a response.

On the day of the experiment atrial pacing was stopped, continuous recording of the electrocardiogram started using precordial leads and the AERP was measured. A control set (25 times) of 10-second-long rapid atrial bursts (800 beats/min, at twice threshold) were performed in order to induce atrial fibrillation in conscious dogs preceded by a bolus infusion of vehicle in 15 min. Following the measurement of AERP, additional sets of atrial bursts (25-25 times) were applied subsequent to either tertiapin-Q (Tocris Bioscience, 18 µg/kg then 56 µg/kg), or dofetilide (Sigma-Aldrich, 25 µg/kg), or propafenone (RYTMONORM, 0.3 mg/kg then 1 mg/kg) i.v. administration. At least 4 days were allowed for washout between in vivo experiments with different compounds.

In another set of experiments (on other PM-instrumented animal group), similarly performed tests were done in order to measure the in vivo effects of Compound 1. In those tests the bolus infusion of vehicle was 20 mL of a mixture of DMSO + β-hydroxypropyl-cyclodextrin + saline (all from Sigma-Aldrich), and during the control 25 bursts and subsequent AF episodes, a continuous infusion of vehicle was maintained (in a volume of 1.7 mL/kg/min). Following the measurement of AERP, Compound 1 (C1) was infused in a dose of 0.3 mg/kg (in 15 min bolus + maintenance) and AF was again induced 25 times. An identical procedure was repeated in every dog with 1 mg/kg dose of C1. All intravenous infusions were performed using a programmable infusion pump (Terufusion TE-3, Terumo Europe, Leuven, Belgium). The ECG was recorded using precordial leads, was digitized and stored on a computer for off-line analysis using SPEL Advanced Haemosys software (version 3.2, MDE Heidelberg GmbH). The incidence of AF, the total duration of AF, the average duration of AF episodes were measured and calculated along with changes in AERP and QT interval. QT intervals were measured on dogs with pacemaker implantation before the 12th burst and were not corrected for heart rate since QT measurements were made at the heart rate set to 80 beats/min by the ventricular pacemaker. Experiments were performed in freely moving conscious dogs so that any effects of anesthetics on AERP and AF could be ruled out.
Action potential recordings from canine right atrial trabeculae with the conventional microelectrode technique

The dogs from the in vivo AF studies were used. Following sedation (xylazine, 1 mg/kg, i.v. and ketamine, 10 mg/kg, i.v.) and anesthesia (pentobarbital, 30 mg/kg i.v.), the heart was rapidly removed through right lateral thoracotomy. The hearts were immediately rinsed in oxygenated modified Locke’s solution containing (in mM): NaCl 128.3, KCl 4, CaCl$_2$ 1.8, MgCl$_2$ 0.42, NaHCO$_3$ 21.4, and glucose 10. The pH of this solution was set between 7.35 and 7.4 when saturated with the mixture of 95% O$_2$ and 5% CO$_2$ at 37 ºC. Isolated right atrial trabeculae were obtained and individually mounted in a tissue chamber with a volume of 50 ml. The preparations were stimulated through a pair of platinum electrodes in contact with the preparation using rectangular current pulses of 2 ms duration. The stimuli were delivered at a constant BCL of 500 ms for at least 60 min allowing the preparation to equilibrate before the measurements were initiated. Transmembrane potentials were recorded using conventional glass microelectrodes, filled with 3M KCl and having tip resistances of 5-20 MΩ, connected to the input of a high impedance electrometer (type 309, MDE Heidelberg GmbH, Heidelberg, Germany) which was coupled to a dual beam oscilloscope. The conduction time, maximum diastolic potential, action potential amplitude, and action potential duration measured at 25%, 50% and 90% of repolarization (APD$_{25}$, APD$_{50}$ and APD$_{90}$, respectively) were evaluated off-line using a custom made software running on an IBM compatible computer equipped with an ADA 3300 analogue-to-digital data acquisition board (Real Time Devices Inc., State Collage, PA, USA) having a maximum sampling frequency of 40 kHz. Stimulation with a constant BCL of 500 ms was applied during the course of the experiments. We aimed at maintaining the same impalement throughout each experiment, however, in case the impalement became dislodged, adjustment was performed and the experiment continued if AP characteristics of the re-established impalement deviated less than 5% from the previous measurement.

Measurement of different ionic currents implicated in atrial fibrillation in the studies on the resveratrol derivative C1
The detailed characterization of the effects of resveratrol and its derivatives, C1-C4 and in more detail, of C1, on \( I_{Kur} \), \( I_{K,ACh} \), \( I_{Na,peak} \) and \( I_{Na,late} \), as well as \( I_Kr \), were performed by our colleagues in Canada. In brief, \( K_v,1.5 \), hERG and \( Na_v,1.5 \) currents were measured from tsA201 cells expressing the human heart genes encoding the \( K_v,1.5 \), \( K_v,11.2 \) and \( Na_v,1.5 \) channels using the patch clamp technique in the whole cell configuration. Carbachol-induced \( I_{K,ACh} \) currents were measured in rat atrial cardiomyocytes.

**Cell shortening and calcium transient recordings**

Ventricular myocytes were prepared from adult Sprague-Dawley rats killed by an overdose of pentobarbital (150 mg/kg, i.p.). The hearts were removed and right ventricular myocytes were then obtained by enzymatic dissociation using standard protocols, which have been described previously by our group. Cell shortening by field stimulation and edge detection and calcium transients using the calcium-sensitive fluorescent probe Calcium Green-1AM were measured during standard procedures as published previously by our group.

**NFAT reporter assay**

Neonatal rat ventricular myocytes were isolated from the hearts of 1-3-day-old neonatal rat pups and cultured for 40 hours. The cells were infected with Ad.GFP or Ad.NFAT-Luc-Promoter adenovirus. Cells were treated 24 h postinfection with vehicle or varying concentrations of C1 or resveratrol and/or angiotensin II (1 \( \mu \)mol/L for 24 h). Cells were harvested with the reporter lysis buffer supplied in the luciferase assay system kit (Promega, Madison, WI, USA) and luminescence according to the manufacturer’s instructions.

**RESULTS**

**1st Aim**

In cooperation with and with the critical help of our colleagues (Dr. László Sághy, Dr. Róbert Pap) at the 2nd Department of Internal Medicine and Cardiology Centre, University of Szeged, the large animal model of chronic atrial tachypacing induced atrial fibrillation was established in the
In Vivo Electrophysiology Laboratory at the Department of Pharmacology and Pharmacotherapy, University of Szeged.

The author of this thesis played an essential role in the manual establishment of the large animal model of chronic atrial tachypacing induced AF in conscious dogs in the laboratory of his supervisor, participating in setting up the model as well as taking part in almost every pacemaker and pacemaker electrode implantation surgery for the studies performed in the laboratory of his supervisor.

**2nd Aim**

*Effects of the \(I_{K, ACh}\) blocker tertiapin-Q, the \(I_{Kr}\) blocker dofetilide and the \(I_{Na}\) blocker propafenone on right atrial effective refractory period (AERP) in conscious dogs*

Before the commencement of right atrial tachypacing, right AERP was 117 ± 5.8 and 127 ± 6.4 ms in conscious dogs (n = 6; at basic cycle lengths of 150 and 300 ms, respectively). Chronic rapid right atrial pacing markedly shortened right AERP. AERP was significantly and dose dependently prolonged by tertiapin-Q at both cycle lengths, yielding the following values at 150 ms BCL: 82.3 ± 1.48 ms in control vs. 93.3 ± 3.33 ms (n=6, p<0.05) following 18 µg/kg and 106.7 ± 2.11 ms (n=6, p<0.05) following 56 µg/kg, respectively. The AERP was also significantly prolonged by dofetilide. Only the larger dose of propafenone increased AERP at the BCL of 150 ms, while the AERP was significantly increased by both propafenone doses at the cycle length of 300 ms.

*Effects of tertiapin-Q, dofetilide and propafenone on burst-induced AF in conscious dogs*

Rapid right atrial bursts at 800/min did not induce any AF in any of the animals before the commencement of atrial tachypacing. Infusion of tertiapin-Q dose dependently and robustly reduced the incidence of right atrial burst induced AF (90.2 ± 1.89% in control vs. 12.4 ± 8.69% following 18 µg/kg and 1.0 ± 0.99% following 56 µg/kg, respectively [n=6, all p<0.05]), the total duration of AF (log10 total duration was 3.9 ± 0.19 in control vs. 1.9 ± 8.69 following 18 µg/kg and 0.3 ± 0.23 following 56 µg/kg, respectively [n=6, all p<0.05]) and the average duration of AF episodes (log10 average duration was 2.3 ± 0.25 in control vs. 0.7 ± 0.32 in
18 µg/kg and 0.2 ± 0.18 in 56 µg/kg, respectively [n=6, all p<0.05]) in conscious dogs. The antiarrhythmic effect of tertiapin-Q was then compared to dofetilide and propafenone, both used in clinical settings for rhythm control in AF management. Both dofetilide and propafenone reduced AF incidence, the total duration of AF and the mean duration of AF episodes. These results clearly show that tertiapin-Q exhibits marked antiarrhythmic effect against AF in conscious dogs, and this effect seemed to be stronger than those of dofetilide and propafenone.

Effect of tertiapin-Q, dofetilide and propafenone on the QT interval in conscious dogs

Importantly, none of the investigated doses of tertiapin-Q prolonged the QT interval in conscious dogs, yielding 283.0 ± 10.36 ms in control vs. 281.8 ± 13.29 ms (n=6, p>0.05) following 18 µg/kg and 268.2 ± 17.75 ms (n=6, p>0.05) following 56 µg/kg, respectively. Dofetilide (25 µg/kg) significantly prolonged the QT interval in conscious dogs, from 265.8 ± 8.68 ms in control to 302.8 ± 10.53 ms (n=6, p<0.05). Propafenone did not influence the duration of the QT interval.

Effects of tertiapin-Q, dofetilide and propafenone on action potentials in atrial trabeculae isolated from dogs with AF

Right atrial trabeculae were isolated from the dogs used for the in vivo AF studies, allowing wash-out of the last compound tested. All measurements were performed at the cycle length of 500 ms. Tertiapin-Q significantly prolonged the action potential at all percentage of repolarization (APD$_{25}$, APD$_{50}$ and APD$_{90}$) in right atrial trabeculae from dogs with AF. Tertiapin-Q (30 nM) did not influence conduction, action potential amplitude, diastolic potential. Dofetilide (100 nM) significantly prolonged the action potential duration only at 90% of repolarization. Dofetilide did not alter conduction time, action potential amplitude, diastolic potential. Propafenone (1 µM) did not prolong the atrial action potential, did not influence the action potential amplitude or diastolic potential but significantly increased conduction time.

The results of these studies are detailed in publications No. I and III. (Juhász et al., 2017; Jost et al. 2011).

3rd Aim
Modulation of multiple ion channels and cellular pathways implicated in AF: effects of Compound 1 (C1), a novel experimental compound

Drugs developed for AF management have been mostly targeted towards ion channels previously. However, structural and other, non-electric remodeling processes also contribute to the initiation and maintenance of AF. It has been suggested that targeting pathological remodeling in AF may also be useful for more effective management of AF. Based on the available data in the literature, an ideal anti-AF compound should exhibit the following effects: (i) $K_v1.5$ ($I_{Kur}$) inhibition in a frequency dependent manner; (ii) $I_{K,ACh}$ block; (iii) $I_{Na,late}$ inhibition; (iv) lack of $I_{Kr}$ inhibition; (v) display atrial specificity: no effect on ventricular repolarization and on excitation–contraction (EC) coupling in ventricular tissue; (vi) antioxidant properties; (vii) NFAT inhibition. In this regard, a novel multifunctional small molecule, related to the structure of resveratrol, was developed at the University of Alberta and University of Manitoba, and was characterized and tested in cooperation with our group.

Effects of C1 and the parent molecule, resveratrol on different ionic currents implicated in atrial fibrillation

The inhibitory effects on $K_v1.5$ currents of four novel resveratrol derivatives were compared to resveratrol. Resveratrol proved to be a weak inhibitor of $K_v1.5$ currents ($IC_{50}$=66 μmol/L). Four resveratrol derivatives were synthesized, and C1 was the most potent blocker of $K_v1.5$ ($IC_{50}$s = 0.36 μmol/L and 0.11 μmol/L for peak and late current inhibition, respectively). The other related compounds (C 2–4) displayed intermediate $K_v1.5$ peak current inhibition (8.3, 10.9 and 11.2 μmol/L, respectively). Therefore, only C1 was selected for further studies. In single atrial cells isolated from rat hearts, $I_{K,ACh}$ currents were induced by carbachol. C1 significantly inhibited carbachol-induced rat atrial $I_{K,ACh}$ currents with an $IC_{50}$ of 1.9 μmol/L. To test the effects of C1 on the peak and late recombinant sodium current, $Na_v1.5$ whole-cell currents were measured. 3 μmol/L C1 resulted in a 50% inhibition of peak current. To obtain concentration–response curves for peak and late $Na_v1.5$ currents, cells were treated with the sea anemone toxin (ATX-II; 3 nmol/L) to induce the late-current component and to test the inhibitory effects of C1 at different concentrations. C1 preferentially inhibited the late current when compared
with peak current \((\text{IC}_{50}s = 1.1 \, \mu\text{mol/L vs.} \, 3.2 \, \mu\text{mol/L, respectively})\).

Inhibition of the hERG \((I_Kr, K_v11.1)\) may result in arrhythmogenic QT prolongation, therefore, the effects of C1 on recombinant whole-cell hERG channel currents were tested. Construction of concentration–inhibition curves revealed that C1 was a weak inhibitor of peak and tail hERG currents with IC\(_{50}\)s of 30 and 25\(\mu\text{mol/L, respectively, and these values were 100-fold higher than those inhibiting peak and late K_v1.5 currents.}\)

The antioxidant and NFAT inhibitory effects of C1

Antioxidant therapy has been shown to reduce the incidence of post-operative AF in patients, and reactive oxygen species (ROS) also directly activate K_v1.5 current. As resveratrol is a known antioxidant, we compared the antioxidant properties of C1 with resveratrol. At 10 \(\mu\text{mol/L, resveratrol and C1 displayed significant antioxidant effects (0.59 \pm 0.04 vs. 0.77 \pm 0.02 of maximal DPPH 517 nm absorbance signal). At 100 \(\mu\text{mol/L, these values were 0.09 \pm 0.02 and 0.19 \pm 0.02 for resveratrol and C1, respectively. It has been shown that resveratrol inhibits NFAT activation induced by phenylephrine contributing to a reduction in maladaptive hypertrophy in neonatal rat ventricular myocytes. Accordingly, we tested the effects of C1 on NFAT activation in these cells. Treatment with 0.01–25 \(\mu\text{mol/L of C1 resulted in a significant reduction in NFAT activity when compared to no treatment. These results revealed that C1 significantly reduced NFAT activity at concentrations of 0.1 \(\mu\text{mol/L and higher, with 1 \(\mu\text{mol/L of C1 exhibiting half-maximal inhibition. C1 also significantly reduced Angiotensin-II-induced activation of NFAT.}\)

Effects of C1 on right atrial effective refractory period (AERP) and atrial fibrillation in conscious dogs

Right atrial effective refractory period (AERP) measurements before the start of rapid atrial pacing at 400 beats/min yielded values of 118 \pm 3.7 and 130 \pm 3.2 ms in conscious dogs \((n = 5; \text{ at basic cycle lengths of 150 and 300 ms, respectively}).\) Rapid right atrial pacing for 6–7 weeks resulted in a significant decrease of right AERP, as AERP decreased below 80 ms in all five animals, and were defined as 79 ms for the following reason: the lower adjustable limit for S1–S2 intervals in the pacemakers available for this study was 80 ms, therefore the exact AERP after 7 weeks of rapid atrial pacing and immediately before C1 administration could not be determined
(measured AERP values were less than 80 ms in all dogs at both basic cycle lengths (BCLs) as an S1–S2 interval of 80 ms still evoked a P wave), although statistical comparison of AERP values was not possible. AERP measurements following C1 administration yielded 87.5 ± 2.50 ms at 150 ms BCL in four animals (in one animal AERP was less than 80 ms) and 90 ± 3.16 ms at 300 ms BCL, respectively, following 0.3 mg/kg C1; and 90.0 ± 3.16 ms at 150 ms BCL, 100 ± 3.16 ms at 300 ms BCL, respectively, following 1 mg/kg C1 (n = 5; with the exception of AERP measurement at 150 ms BCL following 0.3 mg/kg C1). The incidence of AF was not influenced by administration of C1 (86.4 ± 6.4% in control vs. 71.2 ± 15.7% and 66.4 ± 15.7% following 0.3 and 1 mg/kg C1, respectively, both p>0.05). However, the total duration of AF was significantly reduced by 1 mg/kg C1 administration and the average duration of AF episodes was also significantly decreased by 1 mg/kg C1. These results clearly demonstrate in vivo efficacy of C1 against AF in conscious dogs. HPLC analysis of blood plasma collected within 5 min of i.v. bolus injection showed that we obtained concentration ranges of 0.32–0.79 and 0.7–3.0 μmol/L for the 0.3 and 1 mg/kg doses, respectively. None of the investigated doses exhibited any QT interval prolonging effect in five conscious dogs, yielding 261.7 ± 9.37 ms in control vs. 260.9 ± 8.77 ms (p = 0.15) following 0.3 mg/kg C1 and 257.3 ± 9.05 ms (p = 0.55) following 1 mg/kg C1.

The results of these studies are detailed in publication No. II. (Baczkó et al., 2014).

**DISCUSSION**

There is an unmet need for the safer and more efficacious pharmacological management of AF with compounds that lack ventricular cardiac electrophysiological proarrhythmic adverse effects. In the majority of in vivo studies characterizing drug candidates against AF, animals anesthetized with volatile and/or intravenous anesthetics were used. These anesthetic agents have their own relatively well identified effects on cardiac ion channels that can significantly influence the results of these antiarrhythmic studies. In the work that serves as the basis for the thesis, the effects of the atrial selective I_{K,ACH} inhibitor tertiapin-Q, and the multichannel blocker C1 on AF were investigated in freely moving,
conscious dogs, and the results were compared to those with dofetilide and propafenone, drugs used in the clinical setting for rhythm control in patients with AF. Also, for the first time, the effects of some of these compounds on atrial action potential configuration and parameters were compared in right atrial trabeculae isolated from dogs with chronic right atrial tachypacing induced AF. Rapid atrial pacing in dogs is an established large animal AF model where tachypacing leads to electrical and structural remodeling in the atria. In our studies, the electrical remodeling was monitored as the gradual decrease in AERP over the course of chronic tachypacing in our animals.

*Atrial selective ion channel modulation and AF: effects of the $I_{K,ACH}$ blocker tertiapin-Q on AF and action potential configuration in remodeled atrial trabeculae*

In our conscious *in vivo* canine AF model, tertiapin-Q markedly and dose dependently reduced the incidence of AF, the total and average duration of AF episodes, and this effect was paralleled by a significantly increased right AERP following acute intravenous tertiapin-Q administration. The significant prolongation by tertiapin-Q of the action potential duration at all percentages of repolarization was most likely responsible for the increased AERP in right atrial trabeculae isolated from these animals. Tertiapin-Q is a honey bee venom toxin peptide derivative that is a highly selective inhibitor of GIRK ($K_{ir}$3) channels carrying the acetylcholine-activated potassium current, $I_{K,ACH}$. This channel is activated via muscarinic receptors following vagal stimulation leading to atrial action potential shortening and increased atrial dispersion of repolarization, suggesting an important role for this channel in creating an arrhythmia substrate for AF. Although $I_{K,ACH}$ downregulation was found in AF patients, a constitutively active component independent of muscarinic receptor activation was later identified in patients with chronic AF. In a dog model of atrial tachypacing induced AF, constitutive $I_{K,ACH}$ was also observed. Inhibition of $I_{K,ACH}$ by tertiapin-Q increased atrial action potential duration in atrial tachycardia-remodeled canine coronary-perfused left atrial preparations and decreased atrial tachycardia inducibility, similarly to the APD prolongation observed in right atrial trabeculae and the *in vivo* antiarrhythmic activity following tertiapin-Q application in our study. $I_{K,ACH}$ inhibition proved to be beneficial in previous, other canine models of AF – like aconitine and vagal nerve
stimulation induced AF, however, in these studies the effects of $I_{K, ACh}$ inhibition were tested during isoflurane and/or combined isoflurane+thiopental anesthesia. Thiopental significantly prolonged AERP in a concentration dependent manner and caused an increase in atrial wavelength in guinea pig hearts, and isoflurane was found to have antifibrillatory effects in canine atria. Although $I_{K, ACh}$ is also present in the ventricles, it is important to note that in conscious dogs tertiapin-Q did not prolong the QT interval in this study, suggesting that selective $I_{K, ACh}$ block is unlikely to provoke ventricular arrhythmias based on repolarization prolongation.

In summary, we found that the selective $I_{K, ACh}$ inhibitor tertiapin-Q significantly decreased the incidence of AF, reduced the duration of AF episodes and prolonged atrial effective refractory period in conscious dogs with chronic right atrial tachypacing induced atrial remodeling. In this model, similar effects on AF and AERP were observed following the administration of the class IC antiarrhythmic drug propafenone, and the class III compound dofetilide, both used in the clinical management of AF. In right atrial trabeculae isolated from these dogs with AF, atrial action potential durations were prolonged by tertiapin-Q and dofetilide, but not by propafenone, which increased atrial conduction time. Importantly, tertiapin-Q did not affect the QT interval, suggesting that the beneficial effects against AF are not accompanied by adverse effects on ventricular repolarization, therefore, selective $I_{K, ACh}$ inhibitors may be promising atrial selective compounds in the future management of AF.

The other possible path for more efficacious AF management: the parallel modulation of multiple ion channels and cellular pathways implicated in AF - effects of C1, a novel compound related to resveratrol

Inhibition of the $I_{Kur}/K_v1.5$ repolarizing potassium channel that is predominantly expressed in atria rather than the ventricles has been identified as an attractive therapeutic target in recent years. It was found that C1 was an effective inhibitor of this ion channel. The calculated IC$_{50}$ values for $K_v1.5$ channels were in the 0.11–0.36 μmol/L range for late and peak $K_v1.5$ current inhibition, 180- to 600-fold lower than observed for the parent molecule, resveratrol (66 μmol/L). Inhibition of peak sodium current in a frequency-dependent manner may also be a useful strategy to manage
AF by reducing the occurrence of premature action potentials and reduce the incidence of AF. Induction of the Na\textsubscript{v}1.5 late current may not only increase the risk of early after depolarization (EAD)-induced arrhythmias but also might lead to excessive sodium loading within cells that can promote chronic calcium loading, a primary trigger for EADs and calcineurin-mediated activation of the NFAT gene transcriptional pathway leading to pathological hypertrophic remodeling observed in chronic AF. Our results indicate that C1 inhibited peak and late components of recombinant human heart Na\textsubscript{v}1.5 currents more potently than resveratrol. I\textsubscript{Kr} (hERG) inhibition can lead to excessive action potential prolongation and serious TdP arrhythmias. Importantly, we observed that C1 was a weak inhibitor of hERG and it did not prolong the QT interval in conscious dogs. Investigations on cellular pathways implicated in AF generation and maintenance revealed that C1 maintained antioxidant and NFAT-inhibitory properties comparable to its parent molecule, resveratrol. In our \textit{in vivo} conscious dog model with chronic atrial tachypacing induced atrial remodeling and atrial fibrillation, C1 increased AERP and significantly reduced the duration of AF episodes. These results provide \textit{in vivo} evidence that C1 may be beneficial in the treatment of certain forms of AF.

**CONCLUSIONS, NEW RESULTS AND THEIR POTENTIAL SIGNIFICANCE**

1. In cooperation with our clinician colleagues, the author of this thesis has played a key role in establishing a large animal model of chronic right atrial tachypacing induced atrial fibrillation. This model is suitable for the \textit{in vivo} testing of novel drug candidates for the management of atrial fibrillation and also provides essential cardiac tissue for \textit{in vitro} measurements aiming at the investigation of mechanisms responsible for AF initiation and maintenance.

2. We found that the selective I\textsubscript{K,ACh} inhibitor tertiapin-Q significantly decreased the incidence of AF, reduced the duration of AF episodes and prolonged atrial effective refractory period in conscious dogs with chronic right atrial tachypacing induced atrial remodeling. In this model, similar but somewhat less pronounced effects on AF and AERP were observed following the administration of the class IC antiarrhythmic
drug propafenone, and the class III compound dofetilide, both used in the clinical management of AF. Importantly, tertiapin-Q did not affect the QT interval, suggesting that the beneficial effects against AF are not accompanied by adverse effects on ventricular repolarization, therefore, selective \( I_{\text{K,ACH}} \) inhibitors may be promising atrial selective compounds in the future management of AF.

3. In right atrial trabeculae isolated from dogs with AF, atrial action potential durations were prolonged by tertiapin-Q and dofetilide, but not by propafenone, which increased atrial conduction time. The prolongation of repolarization and slowing of conduction would prevent or decrease atrial reentry following the administration of the investigated compounds.

4. A novel resveratrol derivative small molecule developed by our colleagues at the University of Alberta, Edmonton, Canada, exhibited \textit{in vivo} efficacy in our conscious dog AF model and its effects were based on an advantageous combination of effects on multiple ion channels and cellular pathways implicated in AF generation and maintenance. Therefore, C1 can be a starting point for further development of compounds modulating multiple targets for improved pharmacological treatment of AF.
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