Atrium-Specific Kir3.x Determines Inducibility, Dynamics and Termination of Fibrillation by Regulating Restitution-Driven Alternans

Running title: Bingen et al.; Kir3.x in atrial fibrillation

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Journal Subject Code: Basic science research:[132] Arrhythmias - basic studies
Abstract

Background—Atrial fibrillation (AF) is the most common cardiac arrhythmia. Ventricular proarrhythmia hinders pharmacological AF treatment. Modulation of atrium-specific Kir3.x channels, which generate a constitutively active current (I_{K,ACh-c}) after atrial remodeling, might circumvent this problem. However, it is unknown whether and how I_{K,ACh-c} contributes to AF induction, dynamics and termination. Therefore we investigated the effects of I_{K,ACh-c} blockade and Kir3.x downregulation on AF.

Methods and Results—Neonatal rat atrial cardiomyocyte cultures and intact atria were burst paced to induce reentry. To study the effects of Kir3.x on action potential characteristics and propagation patterns, cultures were treated with tertiapin or transduced with lentiviral vectors encoding Kcnj3- or Kcnj5-specific short hairpin RNAs. Kir3.1 and Kir3.4 were expressed in atrial but not in ventricular cardiomyocyte cultures. Tertiapin prolonged action potential duration (APD; 54.7±24.0 to 128.8±16.9ms, p<0.0001) in atrial cultures during reentry, indicating the presence of I_{K,ACh-c}. Furthermore, tertiapin decreased rotor frequency (14.4±7.4 to 6.6±2.0Hz, p<0.05) and complexity (6.6±7.7 to 0.6±0.8 phase singularities, p<0.0001). Knockdown of Kcnj3 or Kcnj5 gave similar results. Blockade of I_{K,ACh-c} prevented/terminated reentry by prolonging APD and changing APD and conduction velocity (CV) restitution slopes, thereby altering the probability of APD alternans and rotor destabilization. Whole heart mapping experiments confirmed key findings (e.g. >50% reduction in AF inducibility after I_{K,ACh-c} blockade).

Conclusions—Atrium-specific Kir3.x controls induction, dynamics and termination of fibrillation by modulating APD and APD/CV restitution slopes in atrial tissue with I_{K,ACh-c}. This study provides new molecular and mechanistic insights into atrial tachyarrhythmias and identifies Kir3.x as a promising atrium-specific target for antiarrhythmic strategies.

Key words: Kir3.x, I_{K,ACh-c}, atrial fibrillation, optical mapping, alternans
Introduction

Atrial fibrillation (AF) is the most common cardiac rhythm disorder in humans, and substantially contributes to morbidity, mortality and health care costs.1-3 Ablation techniques, breaking up AF circuits or triggers, have improved the outcome of AF in the past decades.4 While the success rate of AF ablation is relatively high in paroxysmal AF, in permanent AF ablation restores sinus rhythm only in a fraction of patients, due to altering underlying mechanisms and structural changes of the atrial myocardium. Importantly, there is a lack of consensus on the optimal ablation strategy in these patients, while this population is vastly growing.5,6 In addition, ablation procedures are associated with a risk of complications due to their invasive nature.7 Furthermore, long-term maintenance of sinus rhythm requires repeated procedures or continuation of anti-arrhythmic drugs in a significant proportion of chronic AF patients.8 Hence, first line treatment of AF is still performed pharmacologically.4 However, the use of anti-arrhythmic agents is hampered by potentially lethal ventricular pro-arrhythmia, lack of efficacy and serious side effects.9-13 Thus, research on AF treatment has focused on finding atrium-selective drugs, with higher efficacy in AF rhythm control but less side effects such as ventricular pro-arrhythmia.

In the heart of most mammals, including humans, the acetylcholine-activated potassium current (I_{K,ACH}) is found exclusively in the atrium.14 Hence, I_{K,ACH} is one of the novel candidates for atrium-specific drug treatment. Activating I_{K,ACH} by acetylcholine has been shown to shorten action potential (AP) duration (APD) in the atria.15 In patients suffering from persistent AF and atrial remodeling, I_{K,ACH} can become constitutively active.16,17 Whether this constitutively active acetylcholine-inducible current (I_{K,ACH-c}) affects AF induction, dynamics or termination, the mechanisms by which it could do so, as well as the channels involved in this process remain to be elucidated.
To test whether and how $I_{K,ACh-c}$ affects fibrillation, a 2D *in vitro* model of atrial tissue and a whole heart model of AF were developed using atrial neonatal rat cardiomyocytes (nrCMCs) and hearts with endogenous $I_{K,ACh-c}$. Using these models, fibrillation could be systematically and reproducibly studied. Inhibition of Kir3.1 or Kir3.4 activity by the highly specific drug tertiapin or lentiviral vector (LV)-mediated RNA interference (RNAi), was used to study the role of $I_{K,ACh-c}$ in AF initiation, dynamics and termination.

**Methods**

A detailed description of materials and methods can be found in the Supplementary Material online.

All animal experiments were approved by the Animal Experiments Committee of the Leiden University Medical Center and conformed to the Guide for the Care and Use of Laboratory Animals as stated by the US National Institutes of Health.

**Cell isolation and culture**

After careful separation of the atria and ventricles of neonatal Wistar rat hearts, cardiomyocytes were isolated by collagenase digestion and plated on fibronectin-coated, round glass coverslips (15-mm diameter) at a density of 2-8×10$^5$ cells/well in 24-well cell culture plates, as described previously.\(^{18}\) To restrict unwanted expansion of the remaining non-myocytes, cell proliferation was inhibited by incubation with Mitomycin-C (10 μg/ml; Sigma-Aldrich, St. Louis, MO) for 2 h at day 1 of culture.\(^{19}\)

**Western blotting**

Cardiomyocytes were lysed in 50 mM Tris–HCl (pH 8.0), 150 mM NaCl, 1% Triton X-100, 0.5% sodium deoxycholate, 0.1% sodium dodecyl sulfate. Three 15-mm wells, each seeded with
8×10^5 cells, were used for each sample, and each experiment consisted of at least 4 samples. The proteins in the lysate were size-fractionated in NuPage Novex 12% Bis-Tris gels (Life Technologies, Bleiswijk, the Netherlands) and transferred to Hybond polyvinylidene difluoride membranes (GE Healthcare, Diegem, Belgium). Membranes were blocked in Tris-based saline, 0.1% Tween-20, 5% bovine serum albumin (Sigma-Aldrich) for 1 h. Next, membranes were incubated with antibodies directed against Kir3.1 (Alomone Labs, Jerusalem, Israel), Kir3.4 (Santa Cruz Biotechnology, Dallas, TX) or glyceraldehyde 3-phosphate dehydrogenase (GAPDH; loading control; Merck Millipore, Billerica, MA) and corresponding horseradish peroxidase-conjugated secondary antibodies (Santa Cruz Biotechnology) for 1 h.

Chemiluminescence was detected using ECL Prime Western blot detection reagent (GE Healthcare).

**Optical mapping**

At day 9 of culture, investigation of AP propagation on a whole-culture scale by optical mapping (using di-4-ANEPPS [Life Technologies] as voltage-sensitive dye) and subsequent data analyses were performed as described previously.18 During optical mapping cells were stimulated electrically using a custom-made, epoxy-coated unipolar platinum electrode with square suprathreshold electrical stimuli at 1 and 2-20 Hz (2-Hz increments). Burst pacing with a cycle length of 20-100 ms was used to induce reentry. Complexity was defined as the number of phase singularities (PSs) per cm², determined by using the phase space method as described previously.18 The effect of several drugs (100 nmol/L tertiapin [Alomone Labs], 200 nmol/L atropine [Sigma-Aldrich] and 2 μmol/L carbachol [Sigma-Aldrich])16 was studied by pipetting them directly into the medium, dispersing them by gentle agitation, immediately followed by optical mapping.
Whole heart mapping was performed by incubating neonatal rat hearts with 2 μM di-4-ANEPPS in a tissue bath containing oxygenated Tyrode’s solution (comprising [in mM] NaCl 130, CaCl2 1.8, KCl 4.0, MgCl2 1.0, NaH2PO4 1.2, NaHCO3 24 and glucose 5.5 at pH 7.4). Prior to excision of the heart, the left ventricle was injected with Tyrode’s solution supplemented with 20 mM 2,3-butanedione monoxime (Sigma-Aldrich) to minimize motion artifacts with or without 200 nM tertiapin to block Kir3.x channels. During whole heart mapping, AF was induced by burst pacing at a cycle length of 20-100 ms using a custom-made bipolar platinum electrode.

RNAi

Kir3.1 and Kir3.4 expression in neonatal rat atrial cell cultures was selectively inhibited using self-inactivating LVs encoding short hairpin (sh) RNAs specific for rat Kcnj3 (LV-Kir3.1↓) and Kcnj5 (LV-Kir3.4↓), respectively. The shuttle constructs to generate these LVs are derivatives of plasmid SHC007 from the Mission shRNA library (Sigma-Adrich) in which the Aequorea victoria luciferase (PpLuc)-specific shRNA-coding sequence was replaced by a rat Kcnj3- or Kcnj5-specific shRNA-coding sequence and the marker gene cassette consisting of the human phosphoglycerate kinase 1 gene promoter and the puromycin-N-acetyltransferase-coding sequence was substituted by the human eukaryotic translation elongation factor 1 alpha 1 gene promoter and the Aequorea victoria enhanced green fluorescent protein-coding sequence. The negative control vector (LV-PpLuc↓) had the same genetic makeup, except that it contained the aforementioned PpLuc-specific shRNA-coding sequence.

Statistical analysis

Statistical analyses were performed using SPSS11.0 for Windows (SPSS, Chicago, IL). Comparison between two groups was performed using the Mann-Whitney U test or Wilcoxon
signed rank test or Fisher’s exact test where appropriate. Kruskall-Wallis testing with Bonferroni post-hoc correction was used for multiple groups and comparisons. Data were expressed as mean±standard deviation (SD) for a number (n) of observations. Differences were considered statistically significant if p<0.05. Non-linear regression curves were constructed by using a robust exponential one-phase decay curve fit. Accuracy of these curves was expressed as the coefficient of determination (R²). R² was calculated by the formula R²= 1-(SSreg/SStot) where SSreg is the regression sum of squares (the sum of the square vertical distances of individual points to the regression curve) and SStot is the total sum of squares (the sum of the square vertical distances to the mean of all Y values).

Results

Cell culture characteristics

A detailed characterization of atrial and ventricular nrCMC cultures by immunocytology and Western blotting can be found in the Supplementary Material online (Supplemental Figure 1).

Activation pattern characteristics in atrial nrCMC cultures

During optical mapping, atrial cultures showed uniform, convex and fast activation originating from the electrode upon 1-Hz stimulation (Figure 1A). After burst pacing, reentry was induced in the vast majority of atrial cultures. In 46 out of 49 of these arrhythmic cultures activation patterns remained stable during each cycle of reentry, while also the number and spatial dispersion of PSs did not change during 6 s of mapping (Figure 1B, Supplemental Movie 1).

Return mapping (plotting a peak-to-peak interval [PPI] against the subsequent PPI) and time series of the PPIs showed that through such a fixed activation pattern and PS position these arrhythmias had period-1 (P-1) oscillatory dynamics (i.e. all rotor periods had approximately the
same length) (Supplemental Figure 2A,C,E). Interestingly, the remaining 6% of cultures showed changes in activation pattern, PS number and PS position for each reentrant cycle (Figure 1C, Supplemental Movie 2). Return mapping and time series of the PPIs in these cultures showed period>1 or aperiodical oscillatory dynamics (i.e. the wavefront rotation alternated between >1 different periods or showed a different period during each cycle) (Supplemental Figure 2B,D,F).

During burst pacing-induced reentry (including both the stable and the unstable cases of reentry, n=49), rotor frequency showed a hyperbolic-like relationship with both APD₈₀ (R²=0.91) and wavelength (R²=0.63). No apparent relation was found between rotor frequency and conduction velocity (CV) (Figure 1D-F). Moreover, the number of rotors negatively correlated with wavelength (R²=0.73) and APD₈₀ (R²=0.70). Again CV, other than its effects on wavelength did not show a strong individual correlation with the number of PSs during reentry (Figure 1G-I). These results show that fibrillatory activation in this model can be maintained by P-1 or period>1/aperiodical reentry. Furthermore, the correlation analyses suggest that prolongation of atrial APD and wavelength during AF could decrease rotor frequency as well as complexity, possibly leading to prevention or termination of AF.

Blockade of atrium-specific Iₖ,ACH decreases rotor frequency, complexity and inducibility of reentry

Atrial nrCMCs abundantly expressed Kir3.1 and Kir3.4 in comparison to ventricular nrCMCs (100.0±6.3 vs 18.3±0.7 and 100±2.8 vs 7.8±2.0%, respectively, p<0.05, n=4 per group) as judged by Western blot analyses (Figure 2A). As such, Kir3.x could be a target for blockade in an attempt to selectively prolong atrial APD, thereby preventing or terminating AF. To test this hypothesis both atrial and ventricular nrCMC cultures were treated with 100 nM tertiapin, a
specific blocker of $I_{K,ACh}$.

Tertiapin significantly increased APD$_{80}$ in atrial nrCMCs during 1-Hz pacing (from 56.5±12.5 to 145.5±20.6 ms, $p<0.0001$, $n=33$) (Figure 2B), in the absence of exogenous acetylcholine, showing the presence of $I_{K,ACh-c}$ in these cells (see also Supplemental Results and Supplemental Figure 3). Tertiapin had no significant effect on APD in ventricular nrCMCs ($n=12$) (Figure 2C). Hence, the increase in APD$_{80}$ induced by tertiapin was significantly larger in atrial nrCMCs ($86.7±19.6$ vs $9.5±9.9$ ms in ventricular nrCMCs) (Figure 2D).

Blockade of $I_{K,ACh-c}$ by tertiapin increased APD$_{80}$ during reentry (from 54.7±24.0 to 128.8±16.9 ms, $n=42$) (Figure 2E-G), which significantly decreased rotor frequency (from 14.1±7.4 to 5.9±1.7 Hz) and complexity (from 6.7±7.7 to 0.58±0.83 PSs) (Figure 2H,I). Interestingly, the inducibility of reentry also decreased (from 89.2 to 27.2%) after tertiapin treatment (Figure 2J). Concomitantly, tertiapin led to complete termination of all rotors in 67.4% of cultures, while rotor termination occurred in only 7.0% of control cultures (Figure 2K).

**Kcnj3/5 knockdown in atrial nrCMC cultures**

To investigate the individual contribution of the molecular determinants of $I_{K,ACh-c}$ (Kir3.1 and Kir3.4) to the induction and dynamics of AF, expression of $Kcnj3$ and $Kcnj5$ was specifically downregulated in atrial nrCMC cultures by means of lentiviral, shRNA-mediated RNAi. Kir3.1 and Kir3.4 protein levels were significantly lowered in cultures transduced with LV-Kir3.1↓ and LV-Kir3.4↓, respectively, when compared to those in LV-PpLuc↓-treated control cultures (20.2±1.7 vs 100.0±2.4 and 34.6±3.5 vs 100.0±34.6%, respectively, $n=4$ per group) (Figure 3A-C). As expected, transduction with LV-Kir3.1↓ ($n=11$) and LV-Kir3.4↓ ($n=20$) resulted in significant APD prolongation when compared to PpLuc↓-treated control cultures ($n=11$) (Figure
3D). APD$_{80}$ was significantly increased throughout all pacing cycle lengths (PCLs) (Figure 3E). Also, CV was lowered significantly by both Kcnj3 and Kcnj5 knockdown (Figure 3F), possibly caused by a depolarizing effect of long-term I$_{K,ACh}$ downregulation on the resting membrane potential.

**Effect of Kcnj3/5 knockdown on reentry in atrial nrCMC cultures**

Next, reentry was induced in cultures transduced with LV-Kir3.1↓(n=14), LV-Kir3.4↓ (n=12) or the control LV (n=19) to investigate the effect of Kir3.x-dependent APD prolongation on spiral waves. As expected, reentrant cycle length was significantly increased after knockdown of either Kcnj3 or Kcnj5 (Figure 4A-D,F) consistent with prolongation of APD$_{80}$ during reentry (Figure 4E). Similarly, activation frequency and complexity were significantly decreased (Figure 4G,I, Supplemental Movie 3). In addition, inducibility of reentry was clearly reduced by Kcnj3 or Kcnj5 knockdown (Figure 4H). Together, these results show that the effect of tertiapin on reentry induction and dynamics can be reproduced by RNAi-mediated reduction of Kir3.1 or Kir3.4 protein levels.

**I$_{K,ACh-c}$ blockade prevents APD alternans by decreasing APD and CV restitution slope**

To investigate how I$_{K,ACh-c}$ blockade or downregulation prevents reentry induction we studied the effect of tertiapin on APD and CV restitution. During pacing at 1-20 Hz Kir3.x blockade by tertiapin increased the APD$_{80}$ throughout all PCLs when compared to control cultures (n=8 per group) (Figure 5A). CV was unaltered following 1-2 Hz pacing after tertiapin treatment.

However, at higher pacing frequencies, CV was significantly lower in cultures treated with tertiapin (Figure 5B). This was possibly attributable to a change in the maximal diastolic potential, which gets more depolarized at higher pacing frequencies if the APD is prolonged by I$_{K,ACh-c}$ blockade. Furthermore, wavelength was significantly increased by tertiapin at all
activation frequencies (Figure 5C). Interestingly, the slope of the APD and CV restitution curves were strongly flattened by tertiapin treatment as a consequence of significantly increased minimal APD_{80} and decreased maximal activation frequency (Figure 5D,E). As the restitution curves naturally become steeper at shorter diastolic intervals (Figure 5A-C), prolonging minimal APD (and thereby decreasing maximal activation frequency) prevents steepening in APD and also CV restitution. Tertiapin treatment decreased the maximal APD restitution slope from 1.0±0.4 to 0.3±0.2 (Supplemental Figure 4A,B), while the percentage of cultures with a maximal APD restitution slope above the critical value of 1.21 decreased from 71.4% to 0% (Supplemental Figure 4C).

As a consequence of restitution moderation, spatial (21.7±8.0 vs 56.5±4.4% in control) and temporal (7.8±6.5 vs 38.1±8.0% in control) dispersion in APD were significantly decreased by Kir3.x blockade (Figure 5F-I). Since steepening in APD and CV restitution causes small PCL changes to result in large APD and CV changes, APD alternans occurred in control but not in tertiapin-treated cultures. During APD alternans at stable PCL in a control culture, as a consequence of the relationship between APD and diastolic interval (see also Figure 5A) a long APD is repeatedly followed by a short APD, because the diastolic interval is shortened after the long APD (Figure 5G). Because the short APD follows after incomplete repolarization during the long AP, amplitude alternans occurs (likely due to inactivation of Na^{+} channels as a result of incomplete repolarization). Therefore blockade of I_{K,ACH-c} leads to a significant decrease in temporal amplitude dispersion (Figure 5G,J). Altogether, these findings suggest that I_{K,ACH-c}-dependent alternans is linked to restitution kinetics (see also Supplemental Results, Supplemental Figure 5).

Role of APD alternans in reentry
In control atrial cultures, reentry initiation after burst pacing was found to be a consequence of highly incident APD alternans, during which a long AP was repeatedly followed by a short AP (Figure 6, left panel). Typically, the long AP was uniformly propagated throughout the culture (Figure 6, right panel, long AP). However, because of the spatial heterogeneity in APD, the short AP frequently underwent conduction block when propagated from an area with short-long APD to an area with long-long APD. This caused PSs to arise adjacent to the area of conduction block (Figure 6, left and right panel, short AP). The AP was subsequently propagated around the PS. This could lead to reentry if the area in which the conduction block occurred repolarized before return of the wavefront, which then circled the PS in a retrograde fashion (Figure 6, right panel, short AP). These results show that APD alternans, which can be prevented by blockade of $I_{K_ACh-c}$, is a major factor in reentry initiation.

**Mechanism of rotor termination after $I_{K_ACh}$ blockade**

In addition to preventing reentry induction, blockade of $I_{K_ACh}$ terminated a large portion of rotors initiated by burst pacing. Atrial cultures were mapped during addition of tertiapin, to study how $I_{K_ACh}$ blockade led to rotor termination. Interestingly, during tertiapin incubation rotors with P-1 oscillatory dynamics destabilized into period-2 (P-2) and aperiodical reentry before termination (Figure 7 and Supplemental Figure 6). As shown in Figure 5A, incubation of atrial cultures with tertiapin led to an increase in minimal APD and a decrease in the maximal slope of the APD restitution curve. The higher steepness of the restitution slope at-long PCLs in the tertiapin-treated cultures suggests that this slope becomes critically steep before tertiapin has completely increased minimal APD. This caused APD alternans in vulnerable spots during reentry, leading to P-2 oscillatory dynamics (Figure 7, middle panel and Supplemental Figure 6B,D).

Subsequently, APD prolonged further and refractory periods around the PSs increased.
Consequently, if the refractory period got critically long the reentrant wave had to alter its path from the previous cycle in order to sustain. Therefore, PSs tended to shift position after incubation with tertiapin, leading to aperiodical reentry dynamics (Figure 7, lower panel, Supplemental Figure 6C,D), which simultaneously increased the chance of these PSs to meet a boundary, followed by rotor termination (Figure 7, lower panel, Supplemental Figure 6C,D). As tertiapin increased wavelength and decreased the incidence of APD alternans after full incubation, new rotors are not formed, resulting in a net decrease in complexity of conduction patterns after tertiapin treatment, ultimately leading to termination of AF (Supplemental Movie 4). These results show that apart from affecting reentry initiation and global reentry characteristics, $I_{\text{KACh-c}}$ also determines the period dynamics and propensity towards termination by controlling the onset of APD alternans and PS drift during reentry.

**Kir3.x blockade in whole heart mapping**

Our novel 2D model of atrial tissue appeared to be crucial for a mechanistic understanding of the role of $I_{\text{KACh-c}}$ in rotor formation, dynamics and termination, as these events are likely to occur subepicardially in the intact atria, which precludes their direct read-out and interpretation. However, whole heart data are needed to show the effects of $I_{\text{KACh-c}}$ on actual AF in a more complex and relevant setting. We thus studied the effects of tertiapin in a whole heart model of AF using neonatal rat hearts. In these hearts, immunocytological analyses confirmed the expression of myosin light chain 2a (MLC2a) in the atria only (Supplemental Figure 7A). The atria consisted of 37.9±12.1% non-myocytes, which were predominantly fibroblasts as judged by $\alpha$-actinin/collagen-I double staining (Supplemental Figure 7B,C). Consistent with the in vitro results, Western blot analyses showed a significantly higher expression of Kir3.1 and Kir3.4 in the atrium when compared to the ventricle (100.0±21.3 vs 13.7±6.0 % p<0.05 and 100.0±24.2 vs...
22.4±5.2 % p<0.05 respectively) (Supplemental Figure 7D).

During optical mapping of whole neonatal rat hearts blockade of Kir3.x channels by tertiapin significantly increased atrial APD (62.8±20.9 vs 111.0±32.9 ms, p<0.05) in sinus rhythm while the APD in the ventricles was not significantly altered (219.7±57.6 vs 211.8±42.7 ms, p<0.05) (n=10 per group) (Figure 8A-C). Atropine treatment had no significant effect on atrial APD (54.7±13.9 vs 62.8±20.9 ms in control hearts) (Supplemental Figure 8 A-C), confirming the M2-receptor independent, constitutive activation of I_{KAC} in neonatal rat atria. After burst pacing, AF was maintained by period-1 oscillatory dynamics in both control and tertiapin-treated hearts. In hearts treated with tertiapin, APD during AF was significantly longer when compared to control hearts (54.8±14.2 ms vs 38.8±7.9, p<0.05) (Figure 8D,E). As a consequence, AF cycle length was significantly increased (106.5±10.3 vs 81.3±11.3 ms, p<0.05) (Figure 8F), while the inducibility of AF showed a significant decrease after tertiapin incubation (90% vs 40%, p<0.05) (Figure 8G). Together, these results support the notion that Kir3.x determines the initiation and maintenance of AF in the whole heart.

**Discussion**

The key findings of this study are, (1) the acetylcholine-inducible potassium current (I_{KAC}), mediated by Kir3.x, is highly atrium-specific and constitutively active in neonatal rat atrial cell monolayers and intact atria. (2) In the presence of this current, sustained reentry can be easily induced electrically (~90% incidence), while the incidence decreases strongly after I_{KAC} blockade. (3) After induction, tachyarrhythmias in atrial cultures are maintained by stable period-1, shifting period>1 or aperiodical rotors, and result from restitution-driven alternans, whereas they are terminated by alternans-mediated PS drift. (4) Mechanistically it is shown by pharmacological and genetic interventions that Kir3.x is a key regulator of these processes of
rotor induction, dynamics and termination by controlling APD and APD alternans through APD restitution steepening. (5) Kir3.x represents a promising atrium-specific target for anti-arrhythmic strategies.

Kir3.x in models of AF

Previous studies showed that I_{K,ACh} may play an important role in the onset of AF, as acetylcholine activates I_{K,ACh} during parasympathetic stimulation and thereby shortens APD.\textsuperscript{15} However, after atrial remodeling, which occurs for instance during permanent AF, APD is shortened independently of parasympathetic activation, partly because I_{K,ACh} has become constitutively active.\textsuperscript{16} While alteration of atrial electrophysiology after atrial remodeling has been established, most of the insights into the electrophysiological mechanisms and complex dynamics of wave propagation in AF have come from detailed investigations in computer and animal models of AF, which did not take into account the molecular consequences of atrial remodeling.\textsuperscript{22} Hence, these results may be applicable to paroxysmal AF, but translation to persistent AF and AF after atrial remodeling remains difficult. In the current study, models have been used that include one of these molecular consequences, being constitutive activation of I_{K,ACh}. As such, the present models might provide a novel means to link the molecular biology of (persistent) AF to its basic electrophysiological mechanisms.\textsuperscript{22}

We found that the detrimental effects of I_{K,ACh,c} are strongly dependent on Kir3.x expression, as is the case when I_{K,ACh} is activated by increased parasympathetic tone.\textsuperscript{14,15} As Kir3.1 homomers do not to form a functional channel,\textsuperscript{21,24} the preventive effect of Kcnj3 knockdown is likely attributable to diminished density of Kir3.1/Kir3.4 heteromers at the sarcolemma, while Kir3.4 downregulation affects both Kir3.4 homomers and Kir3.1/Kir3.4 heteromers. This might explain why a 3-fold reduction in Kir3.4 level produced similar results as
a 5-fold decrease in Kir3.1 abundance. While earlier studies show that there is very little contribution of Kir3.4 homomers to IK,ACh,25 our results suggest that after Kcnj3 knockdown there is still residual IK,ACh-c, most likely provided by Kir3.4 homomers, the formation of which is inhibited after Kcnj5 knockdown. This could imply that the contribution of Kir3.x subunits to IK,ACh and to its constitutively active counterpart differs.

Nevertheless, the effects of tertiapin on APD, rotor frequency, arrhythmia complexity and inducibility were found to be larger than those of Kir3.1 and Kir3.4 downregulation. As the atrial nrCMCs have a higher tolerability for short-term IK,ACh-c blockade by tertiapin as opposed to long-term IK,ACh-c blockade by LV-mediated RNAi, tertiapin treatment could provide a somewhat more efficient blockade of IK,ACh-c at the moment of electrophysiological analysis. Furthermore, long-term downregulation of Kir3.1 or Kir3.4 seemed to induce a depolarization of the cardiomyocyte membrane potential, decreasing CV even at long PCLs. The conduction-slowing effect of Kcnj3/5 knockdown by shRNAs strongly diminishes its wavelength-prolonging effect. Therefore, we used tertiapin as an alternative means to study the role of IK,ACh-c in AF in whole hearts. The fact that Kir3.1 and Kir3.4 downregulation still strongly decreased inducibility of reentry in atrial cell cultures shows that the effects of IK,ACh-c inhibition on the slopes in the restitution curves prevails over its effects on wavelength.

**Mechanisms of AF maintenance**

For years there has been an ongoing discussion on whether AF is the consequence of single or multiple ectopic focal discharges, the result of reentrant waves or of randomly appearing and disappearing wavelets.22 Isolating the pulmonary veins (PV) can be successful in preventing paroxysmal AF, which could be interpreted as evidence for the focal discharge theory.26 Nevertheless, reentrant sources generating fibrillatory conduction as a consequence of a
dominant frequency gradient have been demonstrated in chronic AF.\textsuperscript{27} Importantly, in patients with a long history of AF, PV isolation has a low success rate.\textsuperscript{5,6} Therefore, at least in permanent AF, reentry seems to play an important role. The presence of a dominant frequency gradient also makes the multiple wavelet hypothesis less plausible, as a hierarchy in frequencies defies randomness as postulated in this theory.\textsuperscript{28}

As demonstrated in our models, constitutive activation of I\textsubscript{K,ACH} may be one of the determinants of the seemingly increasing role of reentry in the maintenance of AF over time. We show that the constitutively active I\textsubscript{K,ACH} causes APD alternans, making the atrial tissue prone to wavebreak and reentry initiation. In most cases, after reentry is initiated, multiple stable rotors maintain fibrillation. Without ongoing electrical remodeling, emergence of such rotors in the left atrium would lead to fibrillatory conduction to the right atrium. However, in several cultures we observed AF maintained by shifting aperiodical rotors. As the APD and CV restitution can become critically steep if the I\textsubscript{K,ACH} is constitutively active, causing both spatial and temporal heterogeneity in repolarization, rotors can meander and break up. Therefore, in the remodeled atrium (c.q. with constitutively active I\textsubscript{K,ACH}) random wavelets (i.e. random rotors of which the propagated wavefront appear as wavelets at the atrial surface) could maintain AF. This could also explain the possibility of AF originating from a rotor in the right atrium with fibrillatory conduction to the left atrium (for instance after left atrium ablation and permanent AF)\textsuperscript{28}, even though the refractory period is shorter in the left atrium. The fibrillatory aspect of conduction could here be provided by the seemingly random breakup, appearance and disappearance of rotors in the right atrium as opposed to an APD gradient.

**APD alternans and AF**

Disturbed repolarization is thought to play an important role in producing spatial heterogeneity
causing fibrillation in the ventricles. In theory, if fast ectopic firing occurs, for instance from the PVs, spatial heterogeneity in repolarization would also favor reentry initiation in the atria. Hence, it has recently been proposed that repolarization alternans could also play an important role in AF. Indeed, it has been shown that atrial repolarization alternans occurs frequently before initiation of AF.\textsuperscript{29-31} In the diseased atrium changes in calcium handling have been shown to cause the APD to alternate as a consequence of calcium instabilities.\textsuperscript{32} During calcium-dependent APD alternans, APD restitution is not necessarily altered. We show here for the first time that APD alternans underlying reentry can be caused by constitutive activation of $I_{K,ACH}$ which steepens APD restitution. Hence, we confirm that disturbed atrial repolarization is associated with the onset of AF. In addition to this association we show experimentally that wavebreak and resultant reentry are the direct consequence of APD alternans. Earlier \textit{in silico} work predicted that discordant APD alternans leads to wavebreak if the short AP in a long-short sequence reaches the refractory tail of a long AP in a short-long sequence. In accordance with these predictions, we found that propagation of a short AP of an alternating long-short sequence halts when it meets the refractory tail of a long AP. However, in our model the long AP was usually not the consequence of discordant APD alternans, \textit{e.g.} a short-long sequence. Instead, it was the result of a large APD dispersion causing areas with APD alternans to border areas with solely long APDs. Thus, in our model, we provide an extension to the aforementioned theory by showing that spatial APD dispersion in combination with APD alternans (while not being classically discordant) can cause AF in cells with constitutively active $I_{K,ACH}$.

\textbf{Atrial Fibrosis in AF}

Atrial fibrosis is thought to be an important component of AF substrates.\textsuperscript{33} In this study, atrial cell cultures consisted of approximately 17\% cardiac fibroblasts as deduced by collagen type I
immunostaining. Previous studies showed that atrial conduction abnormalities as a consequence of fibrosis strongly depend on the pattern of fibrosis.\textsuperscript{34} It was found that long fibrotic strands of tissue could cause significant conduction abnormalities, and thereby contribute to AF, while diffuse fibrosis only marginally affected conduction. In our in vitro model, fibroblasts are diffusely spread. Also, the percentage of fibroblasts in this model is lower than in intact neonatal rat atria (see also Supplemental figures 1F and 7B,C). In spite of this difference in fibroblast content the atrial cell culture and whole heart models yielded very similar results in terms of the inducibility of AF and the ability to suppress AF by I\textsubscript{K,ACh-c} blockade. Furthermore, in both our in vitro and whole heart model, pathological conditions that promote fibrosis are absent. It thus appears that in our models a possible contribution of fibroblast to AF induction is overshadowed by the strong pro-arrhythmic effects of I\textsubscript{K,ACh-c}.

\textbf{Study limitations}

The present study made use of nrCMCs and neonatal rat hearts, which differ electrophysiologically from the more clinically relevant adult human cardiomyocytes and hearts. Use of cardiomyocytes from human adults is hampered by the difficulties to (i) obtain human cardiac tissue of sufficient quality for the isolation of cardiomyocytes and (ii) maintain human cardiomyocytes in a differentiated state during culture. In addition, healthy atrial nrCMCs were used to facilitate the study of Kir3.x and I\textsubscript{K,ACh-c}. It should be noted, however, that constitutive activation of I\textsubscript{K,ACh} in human atrial myocytes is usually preceded by significant atrial remodeling and hence may have a different origin than in our model. Hence, our study focused on a proof-of principle, investigating the role of I\textsubscript{K,ACh} on APD alternans, AF prevention and termination irrespective of its onset. As such, the results may not be directly extrapolatable to the clinical setting.
Conclusions

In neonatal rat atrial cell monolayers and intact atria, the acetylcholine-inducible potassium current is constitutively active and plays a crucial role in the initiation of sustained tachyarrhythmias. Mechanistically it is shown in atrial cell monolayers that I_{KACH-c} is mediated by Kir3.x and not only regulates the initiation but also the maintenance and termination of these arrhythmias by controlling APD and APD alternans through APD restitution steepening. Accordingly, this study provides insights into the molecular basis of atrium-specific I_{KACH-c} and revealed the crucial role it could play in pro- and anti-arrhythmic mechanisms in atrial tissue. These novel insights could contribute to the development of mechanistically driven and atrium-specific, anti-arrhythmic strategies.

Acknowledgments: We wish to thank Huybert J.F. van der Stadt for excellent technical support.

Funding Sources: This work was supported by the Dutch Heart Foundation (E. Dekker grant [2012/T023 to B.O.B.]) and the Netherlands Organisation for Scientific Research (NWO; VENI grant [91611070 to D.A.P].

Conflict of Interest Disclosures: None.

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**Figure Legends:**

**Figure 1.** Activation maps (top, 6-ms isochrone spacing) and corresponding phase maps (bottom) of atrial nrCMC cultures during (A) 1-Hz activation showing uniform propagation, (B) three subsequent reentrant cycles after burst pacing showing a stable activation pattern and PS density and localization during each cycle and (C) five subsequent reentrant cycles after burst pacing showing changes in activation pattern and PS density and localization during each cycle. The direction of activation is indicated by the white arrows, PSs are depicted as white circles. Relationship between rotor frequency and (D) APD\(_{80}\), (E) CV and (F) wavelength and between complexity of reentry and (G) APD\(_{80}\), (H) CV and (I) wavelength.

**Figure 2** (A) Western blot analysis of Kir3.1 and Kir3.4 levels in atrial and ventricular nrCMC cultures using GAPDH as loading control. (B,C) Typical optical AP records of untreated (red) and tertiapin-treated (blue) atrial and ventricular nrCMC cultures, respectively. (D) Quantification of ΔAPD\(_{80}\) after tertiapin treatment in atrial vs ventricular nrCMC cultures. (E) Typical optical signal records of untreated (red) and tertiapin-treated (blue) atrial nrCMC cultures after induction of AF by burst pacing. (F) Typical phase maps of untreated (left) and tertiapin-treated (right) cultures after burst pacing. White circles indicate PSs. Quantification of (G) APD\(_{80}\) during reentry, (H) rotor frequency, (I) complexity of reentry, (J) inducibility of reentry and (K) complete rotor termination in control and tertiapin-treated atrial nrCMC cultures. #: p<0.05 vs atrial cultures, *: p<0.05 vs control, a.u.: arbitrary units.

**Figure 3.** Western blot of (A) Kir3.1 levels in LV-PpLuc\_ (i.e. control vector)- and LV-Kir3.1\_
transduced atrial nrCMC cultures and (B) Kir3.4 levels in LV-PpLuc↓- and LV-Kir3.4↓-transduced atrial nrCMC cultures using GAPDH as loading control and (C) their quantification. (D) Typical optical signal in atrial nrCMC cultures transduced with the control vector LV-PpLuc↓ or with LV-Kir3.1↓ or LV-Kir3.4↓. Quantification of (E) APD and (F) CV restitution in atrial nrCMC cultures transduced with the control vector LV-PpLuc↓ or with LV-Kir3.1↓ or LV-Kir3.4↓. *: p<0.05 vs LV-PpLuc↓, a.u.: arbitrary units.

**Figure 4.** (A) Activation maps and corresponding phase maps during reentry in atrial nrCMC cultures transduced with the control vector LV-PpLuc↓ or with LV-Kir3.1↓ or LV-Kir3.4↓. White arrows indicate the direction of AP propagation, white circles depict PS position. Optical signal traces during reentry in atrial nrCMC cultures transduced with (B) LV-PpLuc↓, (C) LV-Kir3.1↓ or (D) LV-Kir3.4↓. Quantification of (E) APD, (F) cycle length, (G) activation frequency, (H) inducibility of reentry and (I) complexity of reentry after burst pacing in control (red), LV-Kir3.1↓-transduced (blue) and LV-Kir3.4↓-treated (black) atrial nrCMC cultures. *: p<0.05 vs LV-PpLuc↓, a.u.: arbitrary units.

**Figure 5.** Restitution of (A) APD$_{80}$, (B) CV and (C) wavelength during 1-20 Hz pacing in control (red) and tertiapin-treated (blue) atrial nrCMC cultures. Red and blue dotted lines indicate the maximal slopes in the restitution curves, the solid black line indicates slope=1. Quantification of (D) minimal APD and (E) maximal activation frequency in control and tertiapin-treated atrial nrCMC cultures. (F) APD$_{80}$ maps of untreated (left) and tertiapin-treated (right) atrial nrCMC cultures. (G) Typical optical signal traces during pacing at maximal activation frequency in control cultures (left) showing alternating APDs (red double arrows) on
equal PCLs (green double arrows) and tertiapin-treated cultures showing stable APDs (red double arrows) on equal PCLs (green double arrows). Quantification of (H) temporal APD_{80} dispersion, (I) spatial APD_{80} dispersion and (J) temporal amplitude dispersion. *: p<0.05 vs control, a.u.: arbitrary units.

Figure 6. Typical optical signal traces (graph at left side) in a 9-mm^2 square in a control atrial nrCMC culture during APD alternans leading to reentry. Green and red dotted lines indicate PCL and alternating APD, respectively. Corresponding phase maps (panels at right side) during uniform propagation of the long AP (top panel) and conduction block followed by circular propagation of the short AP (lower panel). Arrows, double lines (blue in the optical signal traces and black in the phase maps) and white circles indicate the direction of AP propagation, conduction block and the PS position, respectively. The positions of points I and II in the culture are indicated in the upper left phase map.

Figure 7. Phase map sequence of an atrial nrCMC culture during tertiapin incubation. The upper row shows the initial P-1 reentrant arrhythmia rotating around a single stable PS and its corresponding optical signals. The second row shows the change into a P-2 reentry (approximately 4 s after tertiapin addition), with meandering, disappearing and reappearing PSs and its corresponding optical signals showing APD alternans in critical areas. The lower 2 rows show the shift in aperiodical reentry (approximately 9 s after tertiapin addition) followed by the termination of rotors as a consequence of PS drift towards the edge of the culture as the wavefront increasingly meets refractory tissue. The white circles indicate PSs, the translucent white circles indicate the PS position in the previous frame. The lower right graph shows the
corresponding AP between point I and II in the culture, where the blue arrow indicates the propagation of the final AP leading to conduction block. a.u.: arbitrary units.

**Figure 8.** (A) Typical examples of a control neonatal rat heart during optical mapping, showing the real image (left panel; left atrium, right atrium ventricles are indicated by the letters LA, RA and V, respectively), a map of atrial activation sequence during sinus rhythm (middle panel, 6-ms isochrone spacing) and a map of the atrial activation sequence during AF induced by burst pacing (right panel, 6-ms isochrone spacing) showing circular activation in the atrial epicardium (white arrow). The white squares indicate the areas from which typical optical signals were derived. (B) Examples of the optical signal traces in the atrium (red) and ventricle (black) in control (left) and tertiapin-treated hearts (right). Quantification of (C) the APD$_{80}$ in atria and ventricles of control and tertiapin-treated hearts during sinus rhythm and (D) of the atrial APD$_{80}$ during AF. (E) Examples of the optical signal traces during AF after burst pacing in the atrium (red) and ventricle (black) in control and tertiapin-treated hearts. Irregularly appearing ventricular traces are indicative of AF. Quantification of (F) AF cycle length and (G) inducibility of AF by burst pacing in control and tertiapin-treated neonatal rat hearts. *: p<0.05 vs control NS: non significant.
Figure 2
Figure 4
Figure 5
Figure 7
Atrium-Specific Kir3.x Determines Inducibility, Dynamics and Termination of Fibrillation by Regulating Restitution-Driven Alternans

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Circulation, published online September 24, 2013;

Circulation is published by the American Heart Association, 7272 Greenville Avenue, Dallas, TX 75231
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Print ISSN: 0009-7322. Online ISSN: 1524-4539

The online version of this article, along with updated information and services, is located on the World Wide Web at:
http://circ.ahajournals.org/content/early/2013/09/24/CIRCULATIONAHA.113.005019

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Methods

All animal experiments were approved by the Animal Experiments Committee of the Leiden University Medical Center and conformed to the Guide for the Care and Use of Laboratory Animals as stated by the US National Institutes of Health.

Cell isolation and culture

Ventricular and atrial cardiomyocytes (CMCs) were isolated from 2-3 old neonatal Wistar rats by collagenase digestion as described previously. Isoflurane inhalation (4-5%) was used to anaesthetize animals. After adequate anesthesia had been confirmed by the absence of pain reflexes, hearts were excised. Subsequently, large vessels were removed and atria were separated from ventricles. Atrial and ventricular tissue was gently minced and digested using collagenase type 1 (450 U/ml; Worthington, Lakewood, NJ) and DNase I (18,75 Kunitz/ml; Sigma-Aldrich, St. Louis, MO) during 2 subsequent 30-min digestion steps with agitation in a water bath at 37°C. Cell suspensions were pre-plated on Primaria-coated culture dishes (Becton Dickinson, Breda, the Netherlands) for 120 min to allow for preferential attachment of non-myocytes. Next, the unattached cells (mainly CMCs) were passed through a nylon cell strainer with a mesh pore size of 70 µm (Becton Dickinson) to remove cell aggregates and, after counting, the cells were plated isotropically on fibronectin (Sigma-Aldrich)-coated, round glass coverslips (15-mm diameter) in 24-well plates (Corning Life Sciences, Amsterdam, the Netherlands). Cell densities of 2-8×10^5 cells/well were used depending on
To restrict unwanted expansion of the remaining non-myocytes, cell proliferation was inhibited by incubation with Mitomycin-C (10 µg/ml; Sigma-Aldrich) for 2 h at day 1 of culture as described previously. All cultures were refreshed daily with Dulbecco’s modified Eagle’s medium (DMEM)/HAM’s F10 (1:1, v/v; both from Life Technologies, Bleiswijk, the Netherlands) supplemented with 5% horse serum (Life Technologies) and cultured in a humidified incubator at 37°C and 5% CO2.

Immunocytology

Cells were stained for the markers of interest after several rinses with ice-cold Error! Not a valid link. to wash out the culture medium, fixation with 1% formaldehyde in PBS and permeabilization with 0.1% Triton X-100 in PBS. Primary antibodies (1:200 dilution in PBS+5% fetal bovine serum [FBS; Life Technologies]) and corresponding Alexa Fluor 488/568-conjugated secondary antibodies (1:400 dilution in PBS+5% FBS; Life Technologies) were left on the cells for 16 and 2 h, respectively, at 4°C. Cultures were stained using antibodies directed against α-actinin (Sigma-Aldrich) as CMC marker, myosin light chain 2a (MLC2a; a gift from Dr. S.W. Kubalak, Charleston, SC) and natriuretic peptide precursor A (NPPA, Merck Millipore, Billerica, MA) as markers for atrial CMCs and myosin light chain 2v (MLC2v; Synaptic Systems, Goettingen, Germany) to identify ventricular CMCs. Non-myocytes were characterized immunostaining using antibodies specific for collagen type I (fibroblasts; Abcam, Cambridge, MA), platelet endothelial cell adhesion molecule-1 (PECAM-1; endothelial cells; Abcam) and smooth muscle myosin heavy chain (smMHC; smooth muscle cells; Abcam). Primary antibodies specific for connexin40 (Cx40; Santa Cruz Biotechnology, Dallas, TX) and connexin43 (Cx43; Sigma-Aldrich) were used to determine expression of gap junction proteins. Counterstaining of the nuclei was performed.
by a 5-min incubation at room temperature with 10 µg/ml Hoechst 33342 (Life Technologies) in PBS+5% FBS. Cells were rinsed twice with PBS+5% FBS after incubation with primary antibodies, secondary antibodies and Hoechst 33342. Coverslips were mounted in Vectashield mounting medium (Vector Laboratories, Burlingame, CA) to minimize photobleaching. Images were taken at equal exposure times between compared groups using a fluorescence microscope equipped with a digital color camera (Nikon Eclipse 80i; Nikon Instruments Europe, Amstelveen, the Netherlands). Dedicated software (NIS Elements [Nikon Instruments Europe] and ImageJ [version 1.43; National Institutes of Health, Bethesda, MD]) were used to store and quantify immunofluorescence signals, respectively. All proteins of interest were studied in at least 3 different cultures per experimental group, from which at least 15 representative images were taken.

**Immunohistology**

Neonatal rat hearts were rinsed with PBS, fixed overnight using 4% formaldehyde in PBS and dehydrated by immersion in 70% ethanol (3 h), 96% ethanol (3 h), 100% ethanol (3 h) and 1-butanol (overnight), respectively. Hearts were embedded in paraffin, cut in 5-µm-thick sections and mounted on StarFrost adhesive microscope slides (Knittel Glass, Braunschweig, Germany). Next, sections were deparaffinized in xylene and rehydrated by the subsequent immersion in 100% ethanol, 96% ethanol, 70% ethanol and PBS for 5 min each. Antigen retrieval was performed by incubating the slides with 0.05% trypsin and 0.1% CaCl$_2$ in dimineralized water at pH 7.8 for 15 min at 37°C and 15 min at 20°C, respectively. Sections were immunostained overnight with the aforementioned primary antibodies directed against MLC2a, α-actinin and collagen type I diluted 1:100 in PBS with 1% bovine serum albumin (BSA; Sigma-Aldrich) and 1% Tween-20 (PBSBT). Corresponding secondary antibodies
Alexa Fluor 488/568-conjugated antibodies, Life Technologies) diluted in PBSBT were incubated for 2 hours, after which the nuclei were counterstained with 10 µg/ml Hoechst 33342 in PBSBT. Image acquisition, processing and analysis were done using the fluorescence microscope and software described above.

**Western blotting**

CMC cultures were rinsed twice with ice-cold PBS to wash out the culture medium. Next, the cells were lysed in RIPA buffer (50 mM Tris–HCl [pH 8.0], 150 mM NaCl, 1% Triton X-100, 0.5% sodium deoxycholate, 0.1% sodium dodecyl sulfate supplemented with protease inhibitors [cOmplete, Mini Protease Inhibitor Cocktail Tablet; Roche Applied Science, Penzberg, Germany]). Lysates were subsequently flash frozen in liquid nitrogen, thawed and centrifuged at 4°C and 21,130×g for 15 min to get rid of undissolved material. The protein concentration in the supernatant was determined using BCA Protein Assay Reagent (Thermo Fisher Scientific, Etten-Leur, the Netherlands). Proteins (10 µg per sample; ≥3 samples per group) were then size-fractionated in NuPage Novex 12% Bis-Tris gels (Life Technologies) and transferred to Hybond polyvinylidene difluoride membranes (GE Healthcare, Diegem, Belgium) by wet electroblotting. Membranes were blocked in Tris-based saline, 0.1% Tween-20 (TBST) supplemented with 5% BSA for 1 h at room temperature. Next, the blots were incubated with primary antibodies directed against Cx40 (1:1,000), Cx43 (1:100,000), Kir3.1 (1:1,000; Alomone Labs, Jerusalem, Israel), Kir3.4 (1; 1,000; Santa Cruz Biotechnology), MLC2a (1:200,000), MLC2v (1:5,000), NPPA (1:5,000) and glyceraldehyde 3-phosphate dehydrogenase (1:120,000; loading control; Merck Millipore) for 1 h at room temperature in TBST+5% BSA. Following 3 rinses with TBST, the blots were incubated with appropriate horseradish peroxidase-conjugated secondary antibodies (Santa Cruz Biotechnology) diluted
1:1,000 in TBST+5% BSA for 1 h at room temperature. After another 3 rinses with TBST, membranes were immersed in ECL Prime Western blot detection reagent (GE Healthcare) and chemiluminescence was captured using the ChemiDoc XRS imaging system (Bio-Rad Laboratories, Veenendaal, the Netherlands).

For Western blotting of whole hearts, hearts were excised, atria and ventricles were carefully separated and rinsed in ice-cold PBS before lysis in RIPA buffer using a TissueLyser LT (QIAGEN, Benelux, Venlo, the Netherlands). The whole heart lysates were subsequently processed in the same manner as the lysates of cultured cells.

**Construction of self-inactivating lentivirus (SIN-LV) shuttle plasmids**

To repress rat *Kcnj3* expression, a SIN-LV shuttle construct encoding a short hairpin (sh) RNA targeting mouse *Kcnj3*, and matching perfectly with the coding sequence of the rat *Kcnj3* gene, was obtained from the MISSION shRNA library (Sigma-Aldrich; clone TRCN0000069736). To knock down rat *Kcnj5* expression, the hybridization product of oligodeoxyribonucleotides

5’ CCGGGACCACAAGAAGATCCCAAAACTCGAGTTTGGGGATCTTTGTTGCTTT
TTG 3’ and

5’ AATTCAAAAAAGACCACAAGAAGATCCCAAAACTCGAGTTTGGGGATCTTTGTTGCTTT
GGTC 3’ was inserted in between the unique SgrAI and EcoRI recognition sites of SHC007 (MISSION shRNA library; Sigma-Aldrich) to replace its Photinus pyralis luciferase (*PpLuc*)-specific shRNA-coding sequence. Next, the marker gene cassette in these constructs and in SHC007, which consisted of the human phosphoglycerate kinase 1 gene promoter and
the puromycin-N-acetyltransferase-coding sequence was replaced by the human *eukaryotic translation elongation factor 1 alpha 1* gene promoter and the *Aequorea victoria* enhanced green fluorescent protein (eGFP)-coding sequence. This yielded the SIN-LV shuttle plasmids pLKO.1-mKcnj3-shRNA.hEEF1a1.eGFP, pLKO.1-rKcnj5-shRNA.hEEF1a1.eGFP and pLKO.1-PpLuc-shRNA.hEEF1a1.eGFP, which were used to generate LV-Kir3.1↓, LV-Kir3.4↓ and LV-PpLuc↓ particles, respectively.

**SIN-LV production**

Vesicular stomatitis virus G protein-pseudotyped SIN-LV particles were generated by transfecting subconfluent monolayers of 293T cells with the packaging plasmids psPAX2 (Addgene, Cambridge, MA; plasmid number: 12260) and pLP/VSVG (Life Technologies) and one of the 3 aforementioned SIN-LV shuttle constructs at a molar ratio of 2:1:1. The 293T cells were cultured in high-glucose DMEM containing 10% FBS. The transfection mixture, which consisted of 40 µg of plasmid DNA and 120 µg of polyethyleneimine (Polysciences Europe, Eppelheim, Germany) in 2 ml of 150 mM NaCl per 175-cm² cell culture flask (Greiner Bio-One, Alphen aan den Rijn, the Netherlands) was directly added to the culture medium. Sixteen hours later, the transfection medium in each flask was replaced by 15 ml of DMEM supplemented with 5% FBS and 25 mM HEPES-NaOH (pH 7.4). At 40-48 h after the start of the transfection procedure, the culture medium was harvested and cleared from cellular debris by centrifugation for 10 min at 3,000×g and filtration through a 33-mm diameter, 0.45-µm pore size polyethersulfone syringe filter (Millex Express; Merck Millipore). To concentrate and purify the SIN-LV particles, 30 ml of vector suspension were layered onto a 5-ml cushion of 20% (wt/vol) sucrose in PBS and centrifuged at 15,000 rotations per min for 2 h at 4°C in an SW32 rotor (Beckman Coulter, Fullerton, CA). Next,
the supernatant was discarded and the pellet containing the SIN-LV particles was suspended in 500 μl of PBS-1% BSA by gentle rocking overnight at 4°C. The concentrated vector suspension was divided in 50-100 μl aliquots and stored at -80°C until use. The 3 SIN-LVs were applied to the atrial CMC cultures at doses that resulted in transduction of essentially all cells in the cultures. The transduction level was assessed using a Zeiss Axiovert 200M inverse fluorescence microscopy to visualize eGFP fluorescence.

**In vitro optical mapping**

At day 9 of culture, action potential (AP) propagation was investigated on a whole-culture scale by optical mapping using the voltage-sensitive dye di-4-ANEPPS (Life Technologies) as described previously.\(^1\)\(^2\) During optical mapping cells were stimulated electrically using a custom-made, epoxy-coated unipolar platinum electrode with square suprathreshold electrical stimuli at 1 and 2-20 Hz (2-Hz increments). Fibrillation was induced by burst pacing with a cycle length of 20-100 ms. A specialized electrical stimulus module with corresponding software (Multichannel Systems, Reutlingen, Germany) was used to perform electrical stimulation. Data analysis and construction of activation maps were performed with specialized software (BrainVision Analyzer 1101; Brainvision, Tokyo, Japan) after pixel signals were averaged with 8 of its nearest neighbors, to minimize noise artifacts. Conduction velocity (CV) in cultures with uniform or reentrant activation patterns was calculated perpendicular to the activation wavefront, between two 3 by 3 pixel grids typically spaced 2-8 mm apart. CV, activation frequency, APD during maximal paced activation frequency (*i.e.* minimal APD) and APD during 1 Hz pacing were determined at 6 different locations equally distributed throughout the culture and averaged before further analysis. APD was determined at 80% of repolarization (*APD\(_{80}\)*) because of the rat AP shape. Wavelength was calculated by
multiplying average CV with APD_{80} (for uniform propagation) or reentrant cycle length.\(^1\)

Complexity was defined as the number of phase singularities (PSs) per culture, determined by using the phase space method as described previously.\(^1\)

The effect of several drugs (100 nmol/L tertiapin [Alomone Labs], 200 nmol/L atropine (Sigma-Aldrich) and 2 µmol/L carbachol (Sigma-Aldrich)\(^6\) was studied by pipetting them directly into the medium, dispersing them by gentle agitation, immediately followed by optical mapping.

**Whole heart mapping**

Neonatal (2-3 days old) Wistar rats were anesthetized by isoflurane inhalation (4-5%) and adequate anesthesia was confirmed by the absence of pain reflexes. Subsequently, the thoracic wall was cut and lifted to expose the heart. Oxygenated Tyrode’s solution (comprising [in mM] NaCl 130, CaCl\(_2\) 1.8, KCl 4.0, MgCl\(_2\) 1.0, NaH\(_2\)PO\(_4\) 1.2, NaHCO\(_3\) 24 and glucose 5.5 at pH 7.4) supplemented with 20 mM 2,3-butanedione monoxime (Sigma-Aldrich, BDM) with or without 200 nM tertiapin to minimize motion artifacts and to block Kir3.x channels, respectively, was carefully injected in a 200 µL bolus into the left ventricle using a 30-G needle. The heart was excised just prior to absence of visible contractions, and submersed in Tyrode’s solution with BDM and tertiapin to remove remaining blood. Next, hearts were incubated in Tyrode’s solution with BDM and tertiapin containing 2 µM di-4-ANEPPS for 2 min at 37°C, after which the heart was rinsed and submersed in Tyrode’s solution with BDM and tertiapin and placed on top of a 37°C heating plate under the optical mapping camera. AF was induced by burst pacing at a cycle length of 20-100 ms using a custom-made bipolar platinum electrode. Control hearts were treated in an identical manner.
except that tertiapin was left out during the entire procedure. Typical optical mapping experiments were performed within 6 min after excision of the heart.
Results

Cell culture characterization

Immunocytological analysis at day 9 of culture showed that 100% of the CMCs (i.e. α-actinin-positive cells) in the atrial cell cultures were MLC2a-positive (i.e. of atrial origin) while in the ventricular cell cultures no MLC2a-positive CMCs were detected (Supplemental Figure 1A,C). In addition, NPPA levels were much higher in the atrial than in the ventricular CMC cultures (Supplemental Figure 1B). Western blot analyses confirmed these results and showed that MLC2v was exclusively present in the ventricular cell cultures. (Supplemental Figure 1D). Atrial CMC cultures contained more Cx40 than the ventricular CMC cultures, while Cx43 levels were higher in ventricular CMCs as judged by both Western blotting (Supplemental Figure 1D) and immunocytology (Supplemental Figure 1E). Atrial cell cultures were also analyzed by collagen type I, PECAM-1 and smMHC staining. While none of the cells contained detectable amounts of PECAM-1 or smMHC, 17.0±2.5% of the cells stained positive for collagen-I (Supplemental Figure 1F) suggesting that the non-myocytes in atrial cell cultures consisted mainly, if not exclusively, of fibroblasts.

Constitutive activation of $I_{K\text{ACH}}$ in neonatal rat atrial CMC cultures

During constitutive activation $I_{K\text{ACH}}$ becomes activated independently of acetylcholine or the acetylcholine receptor. Therefore, to confirm constitutive $I_{K\text{ACH}}$ ($I_{K\text{ACH-c}}$) activity in our neonatal rat atrial CMC cultures, atropine (a non-selective muscarinergic receptor antagonist) was used to block the acetylcholine receptor during optical mapping to study the dependence of APD on acetylcholine receptor activation. Atropine had no significant effect on the APD$_{80}$ (38.5±3.9 vs 38.6±3.9 ms in control cultures, p=ns), whereas subsequent treatment with
tertiapin significantly increased APD$_{80}$ (to 110.9±13.1 ms, p<0.0001) (Supplemental Figure 3A,C). This illustrates that after blockade of the acetylcholine receptor there is still current flowing through Kir3.x channels shortening APD. Thus, our cultures of neonatal rat atrial CMCs indeed possess I$_{KACH}$ activity. Nonetheless, treatment of neonatal rat atrial CMCs with carbachol (a non-selective muscarinergic receptor agonist) induced a significant shortening of APD$_{80}$ (from 39.6±4.2 to 29.3±3.9 ms, p<0.0001). After subsequent atropine treatment APD$_{80}$ rose to 45.6±6.9 ms, (p<0.0001 vs carbachol-treated cells) abolishing the carbachol-induced APD shortening, while addition of tertiapin to the carbachol- and atropine-treated CMCs again greatly increased APD$_{80}$ (to 119.8±18.5, p<0.0001 vs [carbachol- and] atropine-treated cells) (Supplemental Figure 3B,D). Together, these results show that the short AP in neonatal rat atrial CMC cultures is caused by a tertiapin-sensitive current which is independent of ligand-induced muscarinergic receptor activation and therefore constitutively active. The cells, however, also still possess muscarinergic receptor stimulation-dependent Kir3.x activity given the APD shortening-effect of carbachol treatment and its inhibition by atropine.

**APD alternans in neonatal rat atrial CMCs is independent of intracellular [Ca$^{2+}$]**

To confirm the restitution-based nature of APD alternans, neonatal rat atrial CMC cultures were treated with the cell-permeable Ca$^{2+}$ chelator BAPTA-AM (10-100 μmol/L; Life Technologies) to rule out the contribution of intracellular Ca$^{2+}$ to APD alternans and reentry induction. Successful buffering of intracellular Ca$^{2+}$ was confirmed by phase contrast microscopy, showing absence of visible contractions after incubation with 10 μmol/L BAPTA-AM during 1-Hz pacing (data not shown). Treatment with BAPTA-AM did not
prevent APD alternans (Supplemental Figure 5A), while the frequency of reentry induction after burst pacing remained equal for all tested concentrations of BAPTA-AM (84.6% in controls vs 91.7%, 83.3% and 83.3% after treatment of the cells with 10, 50 or 100 µmol/L BAPTA-AM, respectively) (Supplemental Figure 5B). This suggests that APD alternans and consequential reentry induction in our model are driven by the $I_{K_{ACh-c}}$-induced steepness in the APD/CV restitution curve.
Figures and Legends

A

DNA

α-actinin

MLC2a

Merge

atrial culture

DNA

α-actinin

MLC2a

Merge

ventricular culture

B

DNA

α-actinin

NPPA

Merge

atrial culture

DNA

α-actinin

NPPA

Merge

ventricular culture

C

% MLC2a of CMCs

0 20 40 60 80 100

Atrial culture Ventricular culture

ND

D

NPPA

MLC2v

MLC2a

Cx43

Cx40

GAPDH

Protein Level (fold change)

0 30 60 90 120 150

NPPA MLC2v MLC2a Cx43 Cx40

#

#

#

#

#

E

Cx40 α-actinin

atrial culture

ventricular culture

10 μm

F

Collagen I α-actinin

PECAM-1 α-actinin

MLC2a α-MHIC

% cells in atrial culture

0 20 40 60 80 100

FB EC SMC
Supplemental Figure 1. Typical examples of immunocytological double staining of (A) MLC2a (green) and α-actinin (red) and (B) NPPA (green) and α-actinin (red) in atrial (upper panels) and ventricular (lower panels) neonatal rat CMC cultures. (C) Quantification of MLC2a and α-actinin double-positive cells in atrial and ventricular neonatal rat CMC cultures. (D) Western blot analysis (left) and quantification (right) of NPPA, MLC2v, MLC2a, Cx43 and Cx40 levels in atrial and ventricular neonatal rat CMC cultures using GAPDH as loading control. (E) Immunocytological double staining of Cx40 (green) and α-actinin (red) and of Cx43 (green) and α-actinin (red) in atrial (1st and 3rd panel from the left, respectively) and ventricular (2nd and 4th panel from the left, respectively) neonatal rat CMC cultures. (F) Immunocytological double staining of collagen type I (green, left) and PECAM-1 (green, middle) with α-actinin (red) and of MLC2a (green, right) with smMHC (red, right). The corresponding quantification of fibroblasts (FB), endothelial cells (EC) and smooth muscle cells (SMC) is depicted on the far right. #: p<0.05 vs ventricular cultures, ND: not detected.
Supplemental Figure 2. Typical examples of the optical signal in neonatal rat atrial CMC cultures with (A) period-1 and (B) aperiodical reentry after burst pacing. Return maps of the peak-to-peak interval (PPI) sequences in an atrial CMC culture after burst pacing with (C) period-1 reentry, characterized by monofocal clustering in the return map and (D) aperiodical reentry, characterized by the absence of clustering in the return map. Time series of the PPIs in an atrial CMC culture with (E) period-1 (corresponding with Supplemental Figure 2A,C and Figure 1B in the main manuscript) and (F) aperiodical reentry (corresponding with Supplemental Figure 2B,D and Figure 1C in the main manuscript). The red dotted lines indicate clustering of PPI sequences and PPIs in the return maps and time series, respectively.
Supplemental Figure 3. (A) Typical examples of optical signal traces in a neonatal rat atrial CMC culture before (control) and after the cumulative treatment with 200 nmol/L atropine and 100 nmol/L tertiapin during 1-Hz pacing. (B) Typical examples of optical signal traces in a neonatal rat atrial CMC culture before (control) and after the cumulative treatment with 2 µmol/L carbachol, 200 nmol/L atropine and 100 nmol/L tertiapin during 1-Hz pacing. (C) Quantification of APD$_{80}$ at a 1-Hz pacing frequency in untreated atrial CMC cultures and in atrial CMC cultures treated with atropine and tertiapin, respectively. (D) Quantification of APD$_{80}$ at a 1-Hz pacing frequency in untreated atrial CMC cultures and in atrial CMC
cultures treated with carbachol, atropine and tertiapin, respectively. *: p<0.05 vs control, #: p<0.05 vs atropine, ‡: p<0.05 vs carbachol.
Supplemental Figure 4. (A) Typical examples of APD restitution curves of a single control (left) and a single tertiapin-treated (right) culture of neonatal rat atrial CMCs, showing that the maximal slope of the curve in the control culture exceeds 1, while the slope in the tertiapin-treated culture remains well below 1. Black dotted lines indicate slope=1. Colored dotted lines indicate the maximal slope in control (red) and tertiapin-treated (blue) cultures. (B) Quantification of the maximal slope in the APD restitution curve of control and tertiapin-treated cultures. (C) Quantification of the percentage of cultures with a maximal slope in the APD restitution curve ≥1.
Supplemental Figure 5. (A) Typical examples of optical signal traces of control (left) and BAPTA-AM-treated (right) neonatal rat atrial CMC cultures at maximal activation frequency in areas showing APD alternans (B) Quantification of reentry inducibility in control cultures and in cultures treated with 10 µmol/L, 50 µmol/L or 100 µmol/L BAPTA-AM showing no difference in the induction of reentry by burst pacing during buffering of intracellular Ca²⁺.
Supplemental Figure 6. Return maps of the PPI sequences in an atrial CMC culture during (A) typical period-1 reentry showing a single cluster in the return map, (B) the early phase of tertiapin incubation leading to a change into period-2 reentry showing two clusters (red) as a result of APD alternans (See Figure 7 in the main manuscript, second row) and a third cluster (blue) corresponding to the subsequent prolonged reentrant cycle length (temporary 2:1 conduction) starting at the arrow in both subfigures B and D, (C) the late phase of tertiapin incubation leading to aperiodical reentry lacking any clustering in the return map. Dotted red circles indicate clustering of PPI sequences. (D) Time series of the PPIs during the incubation with tertiapin, showing the progression of period-1, to period>1, to aperiodical reentry and
eventually termination. The return maps in A-C are each based on a part of the time series separated by the gray rectangles.
Supplemental Figure 7. Immunohistological double staining of whole neonatal rat hearts for (A) MLC2a (green) and α-actinin (red) and (B) collagen type I (green) and α-actinin (red). (C) Quantification of the number of α-actinin-positive (i.e. CMCs) and -negative cells (i.e. non-myocytes) as judged by immunohistology. (D) Western blot analysis and quantification of Kir3.1 and Kir3.4 levels in the neonatal rat atrium and ventricle using GAPDH as loading control. LA: left atrium, RA: right atrium, LV: left ventricle, RV: right ventricle, *: p<0.05 vs control.
Supplemental Figure 8. Typical optical signal traces in the atria (red trace) and ventricles (black trace) of (A) control and (B) atropine-treated neonatal rat hearts. (C) Quantification of APD$_{80}$ in the atria of control and atropine-treated neonatal rat hearts. NS: non-significant vs control. a.u.: arbitrary units.
**Movie 1:**

Typical example from an optical mapping experiment in a neonatal rat atrial CMC culture after burst pacing showing period-1 reentry dynamics. The first part shows the high-pass-filtered optical signal exemplifying the repeating activation pattern during period-1 reentry. The second part displays the phase map progression of the same optical mapping experiment, showing the wave propagation around multiple PSs with fixed positioning throughout the experiment.

**Movie 2:**

Typical example from an optical mapping experiment in a neonatal rat atrial CMC culture after burst pacing showing aperiodical reentry dynamics. The first part shows the high-pass-filtered optical signal exemplifying the transient activation pattern during aperiodical reentry. The second part displays the phase map progression of the same optical mapping experiment, showing multiple instances of PS formation and disappearance leading to aperiodical dynamics.

**Movie 3:**

Typical example from an optical mapping experiment in neonatal rat atrial cultures investigating the effect of treatment with LV-Kir3.1↓, LV-Kir3.4↓ or the control vector LV-PpLuc↓ after reentry induction by burst pacing. The high-pass-filtered optical signal shows less complex conduction patterns and a lower activation frequency in cell cultures that had
been transduced with the Kcnj3- or Kcnj5-specific shRNA-coding SIN-LV than in those exposed to the control vector.

**Movie 4:**

Typical movie from optical mapping experiment investigating the effect of tertiapin on reentry dynamics and termination. Part 1 shows the high-pass-filtered optical signal in an atrial culture after burst pacing during the first 500 ms after tertiapin incubation, which is characterized by persisting rotors and period-1 dynamics. The second part shows the high-pass-filtered optical signal in an atrial culture after a few seconds of tertiapin incubation, characterized by drifting rotors and aperiodical dynamics, which is followed by termination (part 3) when the last rotor collides with the culture boundary.
Reference List

