Inhibition of Small-Conductance Calcium-Activated Potassium Current ($I_{K,Ca}$) Leads to Differential Atrial Electrophysiological Effects in a Horse Model of Persistent Atrial Fibrillation

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Background: Small-conductance Ca$^{2+}$-activated K$^+$ ($K_{Ca}$2) channels have been proposed as a possible atrial-selective target to pharmacologically terminate atrial fibrillation (AF) and to maintain sinus rhythm. However, it has been hypothesized that the importance of the $K_{Ca}$2 current—and thereby the efficacy of small-conductance Ca$^{2+}$-activated K$^+$ current ($I_{K,Ca}$) inhibition—might be negatively related to AF duration and the extent of AF-induced remodeling.

Experimental Approach and Methods: To address the hypothesis of the efficacy of $I_{K,Ca}$ inhibition being dependent on AF duration, the anti-arrhythmic properties of the $I_{K,Ca}$ inhibitor NS8593 (5 mg/kg) and its influence on atrial conduction were studied using epicardial high-density contact mapping in horses with persistent AF. Eleven Standardbred mares with tachypacing-induced persistent AF (42 ± 5 days of AF) were studied in an open-chest experiment. Unipolar AF electrograms were recorded and isochronal high-density maps analyzed to allow for the reconstruction of wave patterns and changes in electrophysiological parameters, such as atrial conduction velocity and AF cycle length. Atrial anti-arrhythmic properties and adverse effects of NS8593 on ventricular electrophysiology were evaluated by continuous surface ECG monitoring.

Results: $I_{K,Ca}$ inhibition by NS8593 administered intravenously had divergent effects on right and left AF complexity and propagation properties in this equine model of persistent AF. Despite global prolongation of AF cycle length, a slowing of conduction in the right atrium led to increased anisotropy and electrical dissociation, thus increasing AF complexity. In contrast, there was no significant change in AF complexity in the LA, and cardioversion of AF was not achieved.
**INTRODUCTION**

Atrial fibrillation (AF) is the most common sustained cardiac arrhythmia, which poses a serious public health issue in Western societies as its prevalence increases drastically with age (Heeringa et al., 2006). It is estimated that one in four adults over the age of 40 in Europe and the United States will develop AF during the remainder of their lifetime, which is associated with a twofold increase in all-cause mortality and a markedly impaired quality of life (Kirchhof et al., 2016, 2020). Hence, the current unmet need for effective and safe pharmacological treatment options (Waks and Zimetbaum, 2017) vindicates further research efforts in the development of novel pharmacological treatment strategies (Milhes et al., 2012; El-Hau et al., 2015; Ravens, 2017).

Small-conductance Ca$^{2+}$-activated K$^+$ (K$_{Ca}$2) channels have recently been proposed as a possible atrial-selective target to pharmacologically terminate AF (Diness et al., 2010). K$_{Ca}$2 channel inhibition has indeed been shown to affect atrial repolarization in healthy human atrial myocytes and increase the atrial effective refractory period (aERP) in dissected atrial tissue strips (Skibsbye et al., 2014). However, the role of K$_{Ca}$2 channels in persistent and permanent AF pathophysiology is still unclear.

A genome-wide association study revealed a possible relationship between gene variants encoding the K$_{Ca}$2.3 channel and AF without detectable cause (Ellinor et al., 2010, 2012). Moreover, the atrial $I_{K_{Ca}}$ current was shown to be enhanced after short-term atrial tachypacing in dogs (Qi et al., 2014). However, Skibsbye et al. (2014) reported a down-regulation of K$_{Ca}$2.2 and K$_{Ca}$2.3 mRNA in atrial cardiomyocytes from chronic AF patients, resulting in the absence of action potential duration (APD) and the aERP prolonging effect of K$_{Ca}$2 inhibition. This was further supported in a canine model of heart failure and AF without detectable cause (Ellinor et al., 2010, 2012).

The same horses were also included in a previous study, as this study is part of a series on the mechanisms underlying persistent AF conduction, inter-atrial heterogeneity, equine/horse model.

**Conclusions:** Intra-atrial heterogeneity in response to $I_{K_{Ca}}$ inhibition by NS8593 was observed. The investigated dose of NS8593 increased the AF cycle length but was not sufficient to induce cardioversion. In terms of propagation properties during AF, $I_{K_{Ca}}$ inhibition by NS8593 led to divergent effects in the right and left atrium. This divergent behavior may have impeded the cardioversion success.

**Keywords:** persistent atrial fibrillation, atrial selectivity, SK/K$_{Ca}$2 channels, NS8593, epicardial contact mapping, AF conduction, inter-atrial heterogeneity, equine/horse model

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**MATERIALS AND METHODS**

**Animals**

A total of 11 Standardbred mares (461 ± 56 kg, height 158 ± 4 cm) with a mean age of 7 ± 3 years (range 3–12) were included in the study. The horses had no history of cardiovascular disease prior to the study, as confirmed by clinical examination, cardiac auscultation, 24-h Holter ECG monitoring, and echocardiographic examination. The specific inclusion protocol has been described previously (Carstensen et al., 2017).

The same horses were also included in a previous study, as this study is part of a series on the mechanisms underlying persistent AF remodel.
AF and novel atrial-selective, target-based treatment strategies (Fenner et al., 2020). When enrolled in the present study, all horses had been under the influence of tachypacing-induced persistent AF for 42 ± 5 days (Figure 1).

While 11 horses entered the course of this study, five horses had to be excluded from mapping data analysis due to deficiencies in experimental conduct, which potentially influenced cardiovascular stability and thereby cardiac electrophysiology.

All experiments were performed at the Large Animal Teaching Hospital, Department of Veterinary Clinical Sciences, University of Copenhagen, Taastrup, Denmark.

The study was approved by the local ethical committee at the Department of Veterinary Clinical Sciences, University of Copenhagen and by the Danish Animal Experiments Inspectorate (license number 2016-15-0201-01128), and was performed in accordance with the European Commission Directive 86/609/ECC.

AF Induction
A subcutaneous implantable cardioverter defibrillator in pacing mode (ICD; Maximo® II, Concerto®; Medtronic Inc., Minneapolis, MN, United States; equipped with a specialized high rate pacing algorithm) and two right atrial bipolar pacing leads (Tendril™ STS Pacing Leads, 100 cm, St. Jude Medical Inc., St. Paul, MN, United States) were implanted in all horses. Following implantation, all horses were treated with antibiotics (Benzylpenicillin natr., Panpharma, Luitré, France), and non-steroidal anti-inflammatory drugs (Finadyne Vet. 50 mg/ml, Flunixin, IV, MSD, Intervet International B.V. AN Boxmeer, Netherlands) for a minimum of 3 days and were allowed a recovery period of 14 days. Subsequently, AF was induced by high-rate pacing (10 Hz) for ≥48 h or until AF was self-sustained. Longitudinal assessment of AF stability and progression was achieved by manual AF cycle length (AFCL) measurements from intra-atrial electrograms (Supplementary Figure 1) on days 3, 5, 11, 17, 29 (Fenner et al., 2020), and ~40 after AF induction. The AF induction and maintenance protocol was part of a preceding study investigating the efficacy of pharmacological I\(K_{\text{Ca}}\) inhibition in cardioverting AF of varying duration by serial cardioversion over a period of 1 month (Fenner et al., 2020).

Open-Chest Mapping Procedure
The electrophysiological effects of K\(_{\text{Ca}}\) inhibition by NS8593 were evaluated in an open-chest experiment. The horses were intravenously premedicated with flunixin-meglumine (Finadyne®vet., MSD, Segré, France, 1.1 mg/kg), detomidine (Domosedan®vet., Orion Pharma Animal Health, Copenhagen, Denmark, 0.01 mg/kg), acetylpromazine (Plegicil®vet., Dechra Veterinary Products A/S, Uldum, Denmark, 0.03 mg/kg), morphine (Morfín DAK 20 mg/ml, Takeda Pharma A/S, Taastrup, Denmark, 0.06 mg/kg), and butorphanol (Torbugesic®, Orion Pharma Animal Health, Copenhagen, Denmark, 0.01 mg/kg). General anesthesia was induced by zoletapam combined with tiletamine (Zoletil®, Virbac Denmark A/S, Kolding, Denmark, 1.5 mg/kg i.v.) and maintained by isoflurane (IsoFlo Vet., Orion Pharma Animal Health, Copenhagen, Denmark, 1.4%). A constant-rate infusion of rocuronium (Rocuronium, Hameln Pharmaceuticals, Hameln, Germany, 0.3 mg/kg/h) was used for muscle relaxation. ECG and blood pressure were continuously monitored. Aortic pressure (P\(A_o\)) was measured using a pressure catheter (Pressure sensor, Sentron Europe BV, Leek, Netherlands) positioned in the ascending aorta via carotid artery access. A left-sided thoracotomy enabled simultaneous recording of atrial electrical activity by high-density contact mapping (249 electrodes, 2.5 mm inter-electrode distance; Supplementary Figure 2) on the epicardial surface of both atria (Adler et al., 2020). Unipolar AF electrograms were recorded at a sampling rate of 1,039 Hz. The signals were hardware filtered (16th order high pass filter at 0.56 Hz, followed by a 16th order low pass filter at 408 Hz) before digitization using
a 16-bit analog-to-digital converter. To avoid interference of ventricular far-field, doubtful waveforms were detected (using a ventricular electrogram) and removed using averaged beat QRST-template cancellation.

**Drug Administration and Measurement of the Electrophysiological Parameters**

The negative $K_{Ca}$ channel modulator NS8593 ($5 \text{ mg/kg i.v., Acesion Pharma ApS}$) was administered over a 10-min period (Haugard et al., 2015). Plasma levels in venous blood samples were measured at the time points $T = -5$, 0, 5, 10, and 30 min (where $T = 0$ was the start of drug administration) and subsequently analyzed by Syngene International Ltd., Bangalore, India. Plasma protein binding of NS8593 was determined using a standard plasma protein binding assay with tolbutamide as internal standard and warfarin as a reference compound. The assay is based on rapid equilibrium dialysis and subsequent compound detection by quantitative mass spectrometry (additional information in the Supplementary Material).

AF high density maps and ECG parameters were analyzed at the time points $T = -5$, 10, and 20 min. Mapping files of 10–60 s (dependent on availability) were analyzed to quantify the effects of NS8593 on conduction properties during AF. Local activation times (AT) were identified in all electrogams using a probabilistic algorithm (Zeemering et al., 2012). Based on the work of Kléber and Rudy (2004), we considered conduction block to occur if a putative conduction velocity (CV) of $<20 \text{ cm/s}$ was measured. Given the interelectrode distance of 2.5 mm, a maximal AT difference of 12 ms in the horizontal/vertical direction and a difference of 17 ms in the oblique direction was deducible. Conduction velocity was determined by plane-fitting through all activation time-points of the direct neighboring electrodes given the limits of 12 or 17 ms. Moreover, ATs were used to reconstruct fibrillatory waves. Wave boundaries were defined as the edge of the electrode or areas of conduction block. Fibrillation waves were classified as “peripheral” if the earliest activation site was located at the periphery of the array, and as a breakthrough if it was within the mapped area and could not be explained by epicardial conduction. Waves and breakthroughs were normalized to the cycle length. Re-entrant activity was identified based on conduction paths. Conduction paths were determined as the shortest continuous trajectory between a starting and end point of a wave, presuming a $CV \geq 20 \text{ cm/s}$. If the trajectory had $\geq1$ self-intersection after $\geq75\%$ of the mean AFCL it was defined as a local re-entry. Isolated activations of $<3$ adjacent electrodes were considered to be prone to noise or occasional mis-assignment of an AT. These waves were therefore not included in the analysis.

The anisotropy of conduction direction was determined at each electrode and was based on the circular variance of conduction vectors (Maesen et al., 2013). ECG analysis to determine the heart rate (HR), QRS, and rate-corrected QT interval (QTc) was conducted for 10 subsequent RR intervals. The QTc was corrected for HR using a piecewise linear regression model tailored to horses (Pedersen et al., 2016).

**Statistical Analysis**

All data are displayed as median with IQR. Analyses were performed using GraphPad Prism 8 software (GraphPad Prism, RRID:SCR_002798) with $P \leq 0.05$ considered significant. Parameters were tested for normality using a Shapiro–Wilk test. Conditions with $<4$ observations were not included in the analysis. The statistical method used was 2-way repeated measures ANOVA with time point as fixed and horse ID as a random variable to account for correlations over time. In this repeated-measures design, multiple comparisons were performed using Sidák's test for multiple comparisons. The post hoc test was conducted only if the measure of matching effectivity achieved the necessary level of statistical significance ($P \leq 0.05$), and significant variance inhomogeneity was not evident. A piecewise linear regression model has been used to correct QT intervals for HR on the data presented in Supplementary Figure 3.

**RESULTS**

**Atrial Fibrillation Maintenance and Atrial Remodeling**

On average, AF became sustained after 6 ± 4 days of pacing at 10 Hz (Figure 2A). AF stabilization was accompanied by a progressive reduction in AFCL, from 209 (55) ms on day 3 to 169 (17) ms on day 29 ($p < 0.05$; Figure 2B), which indicated the progression of atrial electrical remodeling (Fenner et al., 2020).

**NS8593 Pharmacology and Pharmacokinetics**

A maximal total plasma concentration ($C_{\text{max}}$) of $\sim 7,200 \text{ ng/ml (27.4 } \mu \text{M})$ was reached at the end of infusion (Figure 3A). NS8593 had a plasma protein binding rate of 90.5% in equine plasma, corresponding to a free unbound plasma concentration of approximately 2.6 $\mu$M (9.5% of total plasma concentration) at $C_{\text{max}}$. A rapid decrease in free plasma concentration to 0.33 $\mu$M (free unbound concentration) was observed over a period of 20 min after the end of infusion (Figure 3A). During those 20 min, the atrial fibrillatory process was monitored continuously using body surface ECG and atrial contact ECG recording to assess whether cardioversion to sinus rhythm occurred.

**Effect of NS8593 on Global and Local Conduction Properties During AF**

NS8593 significantly prolonged AFCL by $\sim 50$ ms [Figure 3B: RA: 185 (44) ms to 228 (81) ms; $p < 0.05$]. Interestingly, AFCL reached values similar to those measured 48 h after initial AF induction, when most animals had not yet developed sustained AF [See Figure 2B – day 3: 209 (55) ms and Figure 3B – T30, 228 (81) ms]. Despite this prolongation of AFCL, cardioversion of AF was not achieved.

Wave patterns were determined to elucidate changes in conduction during AF. The recordings displayed various patterns, i.e., wavefront collision and fusion, breakthrough, and re-entrant circuits. These patterns were characterized
by spatiotemporal instability, leading to a variety of patterns observed in horses and atria, respectively. The combined number of waves/AF cycle, breakthrough, and re-entries were quantified to capture this variety of patterns. We found no significant change in the total number of waves/AF cycle (LA + RA; Figure 4A) or in the relative number of breakthrough waves (Figure 4B). During a 60-s recording (Figure 4C), between 0 and 2 re-entries were observed, limited to 1.3 (4.7)% of the time (Figure 6C). The stability of observed re-entrant circuits was not affected by NS8593 (Figure 4D).

Representative activation maps from the right and left atrium are shown in Figure 5 before and after the administration of NS8593. Activation videos of 10 s of AF before and after drug administration are presented in the Supplementary Figure 4.

When stratifying the changes in AF properties, we observed differences in behavior between the LA and RA. In the LA, the number of waves decreased after NS8593 infusion, while right atrial activation maps displayed narrower waves compared to the baseline. Prior to NS8593 infusion, the number of waves/cycle and breakthroughs displayed higher numbers in the LA [waves/cycle: 3.8 (7); breakthroughs: 1.6 (3.6)] compared to the RA [waves/cycle: 2.7 (4); breakthroughs: 0.7 (1.7); Figures 6A,B], however not statistically significant. There was no significant change in the number of waves and breakthroughs in the LA in response to NS8593 administration [waves/cycle: 3.6 (5); breakthroughs: 1 (1.6)]. In the RA, however, the number of waves and breakthroughs increased significantly [waves/cycle: 2.7 (4) to 6.6 (6), p < 0.05; breakthroughs: 0.7 (1.7) to 2.8 (3), p < 0.05], illustrating

**FIGURE 2** | AF stabilization over 1 month. (A) Percentage (%) of horses in stable, self-sustained AF (≥ 24 h) after induction by high-rate atrial pacing (10 Hz). Induction of stable AF required 6 ± 4 days of pacing. (B) Longitudinal assessment of AF stability and progression by AF cycle length (AFCL) from right atrial intra-atrial electrogram recordings 3, 5, 11, 17, 29, and ∼40 days after AF induction. The red dotted line indicates the time point of the terminal high-density atrial mapping experiments. Statistical significance is defined as p < 0.05 and is marked with an asterisk (*).

**FIGURE 3** | NS8593 pharmacokinetics and its effect on the atrial fibrillation cycle length (AFCL). (A) Plasma concentration of NS8593 measured at time points T = –5, 0, 5, 10, and 30 min relative to the start of drug infusion. (B) Influence of NS8593 on right atrial tissue refractoriness in persistent AF (blue) and acutely induced AF (gray). The depicted data on acute AF (gray) are derived from unpublished data from Haugaard et al. The shown data points represent the AF cycle length (AFCL) measured at the start of drug infusion (T0) and at the end of the 20-min observational period following the end of drug injection (T30, persistent AF study), as well as immediately before cardioversion (acute AF study), respectively. Inf., infusion. Statistical significance is defined as p < 0.05 and is marked with an asterisk (*).
FIGURE 4 | Global influence of NS8593 on conduction patterns contributing to AF maintenance. (A) No significant change in the total number of waves/cycle. (B) No significant change in the number of breakthrough waves relative to the number of waves/cycle. (C) No significant change in the total number of re-entrant circuits/second. (D) No significant change in the re-entrant circuit stability (average number of re-entrant circuit revolutions).

Influence of NS8593 on Ventricular Electrophysiology and Hemodynamics

The RR interval, QRS duration, QTc interval, and mean PAo were assessed in 5-min intervals. A significant shortening of QTc (Supplementary Figure 3B, p < 0.05) was observed, associated with a trend toward shorter RR intervals (Supplementary Figure 3A). There was no significant change in the duration of the QRS complexes (Supplementary Figure 3C). In addition, no significant effects on systemic blood pressure or cardiovascular stability were noted (Supplementary Figure 3D).

DISCUSSION

To our knowledge, this is the first study to investigate the effect of antiarrhythmic drugs on conduction patterns in a horse model of sustained AF. Based on our previous work,
we anticipated a strong possibility that $I_{K_{Ca}}$ inhibition by NS8593 would terminate AF (Haugaard et al., 2015). However, restoration of sinus rhythm was not achieved in any of the investigated horses following NS8593 administration. The strong AFCL prolonging effect, beyond the point where AF previously had been unstable, suggests that a sufficient dose of NS8593 was administered to affect the electrical substrate in the horses, yet this effect was not accompanied by a reduction in AF complexity (number of waves). Interestingly, we were able to demonstrate differing effects of NS8593 on right- and left-atrial conduction.
properties. The effect on LA conduction was limited, while in the RA, conduction slowed down, anisotropy increased, and AF became more complex. It is conceivable that these effects on RA conduction impeded successful cardioversion.

Influence of Pharmacological \(I_{K, Ca}\) Inhibition by NS8593 on Atrial Tissue Refractoriness

NS8593 has been shown to exert an \(I_{K, Ca}\) inhibitory effect by negative allosteric modulation of two specific amino acid residues located in the inner pore interacting with channel-specific gating structures (Jenkins et al., 2011). The resulting shift in \(Ca^{2+}\) sensitivity leads to a decrease in potassium outward current and thereby to APD and aERP prolongation (Simó-Vicens et al., 2017). Furthermore, \(K_{Ca2}\) channel inhibition has been suggested to reduce the fast sodium current: a mechanism that may explain the observed decrease in conduction velocity in the RA. Earlier it had been proposed that Na\(^+\) channel availability is indirectly influenced via the slight positive shift in the resting membrane potential; however, very recently, a direct inhibition of the sodium current by NS8593 in canine atria has been observed (Skibsbye et al., 2015; Burashnikov et al., 2020). This combination of these class I and III effects in NS8593 has previously been reported effective in horse with acute AF (Haugaard et al., 2015). However, in the present study, none of the horses cardioverted after 40 days of AF, even though a comparable unbound free fraction of NS8593 [2.6 \(\mu M\) (9.5%) at \(C_{max}\)] was attained. The unbound free fraction of NS8593 was about \(\sim 3\) times higher than the \(IC_{50}\) for \(K_{Ca2.2}\) and \(K_{Ca2.3}\) in human atrial cells (Skibsbye et al., 2014), which may suggest that targeting \(K_{Ca2}\) channels alone is not sufficient in horses with persistent AF.

Despite the inability to terminate AF in these horses, NS8593 increased the AFCL substantially by \(\sim 50\) ms, corresponding to an AFCL prolongation, which effectively shortened and terminated AF paroxysms in acutely induced AF (Figure 3B). It can therefore be hypothesized that AF stability was further facilitated by additional remodeling processes. It is well known

![Graph A](image1.png)
![Graph B](image2.png)
![Graph C](image3.png)
![Graph D](image4.png)
that the efficacy of currently available anti-arrhythmic drugs (AADs) to convert AF reduces with the progression of the atrial substrate (Eijsbouts et al., 2006; Kirchhof et al., 2016; Carstensen et al., 2018), which also seems to be the case for \( K_{\text{Ca}} \) channel inhibition.

The apparent lack of cardioversion success despite the global AFCL prolongation of \(~50\) ms to values similar to early AF progression agrees with previous observations in goats that the critical AFCL required for pharmacological cardioversion might increase substantially in longer-lasting AF (Eijsbouts et al., 2006). It could be hypothesized that further increasing the NS8593 concentration might have led to a sufficient increase in AFCL. However, this may result in a higher probability of non-specific ion channel block and subsequent loss of atrial specificity (Skibsbye et al., 2014).

When considering the pharmacological selectivity profile of NS8593 on relevant cardiac ion currents (Skibsbye et al., 2014), it is likely that some of the reported effects on atrial and ventricular electrophysiology can be attributed to not only the indirect but possibly also the direct class I drug effect (Burashnikov et al., 2020), as the free unbound \( C_{\text{max}} \) of 2.6 \( \mu \text{M} \) is comparable to the compound’s IC\(_{50}\) on Na\(_v\)1.5 of 5 \( \mu \text{M} \).

From the present study, we cannot conclude whether the lack of cardioversion was due to insufficient \( K_{\text{Ca}} \) channel block, down-regulation of \( K_{\text{Ca}} \) channels, or further substrate remodeling (including structural changes) to maintain persistent AF.

### AF Complexity and Inter-Atrial Heterogeneity

Similar to the apparent lack of any anti-arrhythmic effect of \( I_{\text{K,\text{Ca}}} \) inhibition in the present study, contrasting efficacy in pre-clinical drug testing has previously been reported for the class III AAD dofetilide. Dofetilide was highly effective in terminating “coarse” atrial fibrillatory patterns, whereas AF of higher complexity could not be terminated, even though AFCL was equally increased. This led to the assumption that class III...
drug efficacy might be significantly influenced by the underlying mechanism perpetuating AF (Li et al., 2000).

When investigating the coherence of increasing AF stability and the declining efficacy of currently available AADs, it has been shown that AF conduction patterns in the atrial free walls dissociate widely and thereby stabilize over the course of AF (Verheule et al., 2010). The presented differences in anti-arrhythmic efficacy of NS8593 in acute and persistent AF in horses therefore seem to be in agreement with the mechanistic findings of Verheule et al. as we also reported increased dissociated conduction in the right atrial free wall. The inter-atrial heterogeneity in AF complexity seen in response to NS8593 treatment in the persistent AF model is further supported by the class III AAD dofetilide influencing atrial electrophysiology toward stable and persisting AF patterns, particularly maintained by right atrial activity (Li et al., 2000). Similarly, NS8593 increased AFCL equally in both atria but did not affect complexity in the LA, while the RA complexity increased.

In accordance with several studies elaborating inter-atrial differences in AF propensity (Li et al., 2001; Verheule et al., 2010; Embi et al., 2014), LA activation maps reflected distinctively higher electrical complexity prior to drug infusion, suggesting that AF perpetuation was initially driven by left atrial electrical activity. In response to drug administration, however, CV decreased in the RA and remained unaffected in the LA, possibly preventing a global organization of the AF pattern. It, therefore, seems that the class III drug effect exerted by SK channel inhibition in the setting of persistent AF influences right atrial conduction in a way that supports AF perpetuation.

In a recent study investigating clinical AF cases characterized by left-to-right frequency gradients, it has been proven that elimination of the inter-atrial heterogeneity and AF complexity by ablation results in long-term SR maintenance (Atienza et al., 2009). This highlights the clinical importance of atrial specific investigation of pharmacological effects on cardiac electrophysiology in terms of conduction velocity and AF complexity in pre-clinical drug development as well as prospective clinical studies.

**Atrial Size and AF Perpetuation**

Using the horse model of persistent AF raises the question of whether atrial size, and thereby substrate dimension, constitutes an important factor in AF stabilization and complexity (Kaese and Verheule, 2012). Comparisons made between mapping data in the goat and horse model of AF show a similar degree of AF complexity within the mapped area. Nevertheless, normalization of wave and path length to absolute atrial circumference suggests a relatively higher number of waves/cycle in the horse atria (Gatta et al., 2018). However, it seems unlikely that the evident lack of cardioversion by NS8593 treatment in this study is based on an initially higher relative AF complexity. On the contrary, the observed global increase in AFCL would give us reason to expect an increase in wave length (WL), effectively abrogating re-entrant circuits initially responsible for AF maintenance (Wang et al., 1993). However, as WL is the product of ERP and CV (WL = ERP × CV), the increase in global refractoriness seems to be equated by distinct uniatrial (RA) conduction slowing, ultimately preventing the anticipated increase in WL. Given the atrial size in horses, the lack of WL increase seems to allow for continuous activity, with exceptionally stable and more complex wavefronts observed in the RA contributing to AF perpetuation.

We further hypothesize that the observed local right atrial slowing of conduction, putatively due to indirect and possibly also direct sodium channel and/or gap junction blockade, facilitates the persistence of atrial fibrillatory activity. Right atrial activation maps display complex conduction patterns, further stabilized by increased anisotropy and lateral conduction failure in the mapped epicardial plane, known as “longitudinal dissociation” (Myerburg et al., 1973). Analogous conduction patterns of longitudinal dissociation, enhancing the AF complexity and stability, were likewise observed in a mapping study in long-term AF patients (Allessie et al., 2010). In these patients, lines of conduction block ran parallel to the right atrial pectinate muscles, offering a potential explanation for the RA-LA gradient in direction dependence, as left atrial trabeculae are oriented more randomly. It is conceivable that the same principle applies to the present study, as the anatomical structure of equine and human pectinate muscles has been reported to be comparable (Bright and Marr, 2010).

**LIMITATIONS**

The present study contributes knowledge to experimental electrophysiology and provides novel insights into the in vivo pharmacology and electrophysiological properties of NS8593 inhibiting atrial $I_{K,\text{Ca}}$ current. Using the horse as a new large animal model in cardiac electrophysiological research has both advantages and disadvantages. The size of the animal, and, as a consequence, the size of the atria, as well as the fact that horses are one of the few mammalian species besides humans that spontaneously develop AF mean that this species offers good translational value (Schüttler et al., 2020). However, due to the size of the species, studies are often limited to a relatively small number of animals. In the present study, a higher number of animals would have allowed us to set up a sham-operated control group to assess the effect of anesthesia and the open-chest setting on cardiovascular stability in order to differentiate between drug effect and unwanted interference with cardiac electrophysiology in the specific experimental setting. However, Haugaard et al. (2015) did not report any changes in aERP due to anesthesia.

Furthermore, a larger sample size might have allowed for species-specific dose-response experiments. However, given the well-known pharmacokinetic properties and the reported in vitro pharmacology, we would expect a significant portion of the $I_{K,\text{Ca}}$ current to be inhibited.

Additionally, one technical limitation associated with atrial size must be considered – although both atria were mapped simultaneously, the mapped area was limited to the size of the electrode grid. As a result, it is possible that events might have taken place outside the field of view.
As mentioned in the result section, activation time videos illustrated that conduction patterns in AF are characterized by a high instability of spatiotemporal behavior (Supplementary Figure 4). This might affect the perception of the arrhythmia in short time samples, pilot analyses, however, revealed that the investigated parameters were not impacted by the reported minimal recording length of 10 s.

Lastly, future investigations to fully elucidate the effect of NS8593 and its derivatives in the presence of re-entrant circuits would be beneficial to further support our conclusions.

CONCLUSION

A new open-chest in vivo model, including high-density contact mapping on the equine heart, has been developed. It allows detailed electrophysiological measurements and comparison between the LA and RA and was used to study the effect of \( K_{Ca2} \) channel inhibition by NS8593, exhibiting both class I and III anti-arrhythmic effects during AF.

In conclusion, our results have shown that selective inhibition of \( K_{Ca2} \) channels in horses with persistent AF leads to a global slowing of fibrillation frequency. However, the administered dose of NS8593, which successfully terminated acute AF, was not sufficient to lead to cardioversion in any of the included animals with persistent AF.

The observed differential effect on CV and AF complexity between the atria indicates inter-atrial differences in susceptibility for the indirect and direct class I drug effect of NS8593. In combination with the apparent coherence between an RA-LA frequency gradient in clinical AF and the capability of maintaining SR following cardioversion attempts (Atienza et al., 2009), this study’s findings highlight the importance of atrial specific investigation of pharmacological effects on cardiac electrophysiology in terms of basic conduction properties, such as conduction velocity, complexity and AF cycle length in pre-clinical drug development as well as prospective clinical studies.

Finally, as it seems like targeting \( K_{Ca2} \) channels alone is not sufficient to achieve a relevant prolongation of atrial tissue refractoriness in horses with persistent AF, experimental investigation of combinations of atrial-selective AADs may be considered in the future.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

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ETHICS STATEMENT

The animal study was reviewed and approved by the Danish Animal Experiments Inspectorate (license number 2016-15-0201-01128).

AUTHOR CONTRIBUTIONS

MF, GG, AV, and RB contributed to the conception of the study. MF, GG, SS, TJ, SV, AV, and RB contributed to the design of the study. MF, GG, SS, MK, EH, DA, MS, SV, and AV contributed to the acquisition of data. MF, GG, SS, MK, EH, DA, MS, USc, USø, JD, TJ, SV, AV, and RB contributed to the intellectual content of the work by revising the draft until all authors gave their final approval for this version to be published. All authors agreed to be accountable for all aspects of the work and for ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fphys.2021.614483/full#supplementary-material
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**Conflict of Interest:** USø and JD are co-founders of Acesion Pharma ApS and USø was one of the inventors of NS8593.

The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest, as the anti-arrhythmic compound NS8593 was provided free of charge solely for academic purposes.

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