The role of action potential alternans in the initiation of atrial fibrillation in humans: a review and future directions

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This review highlights the role of atrial monophasic action potential duration (APD) in understanding atrial electrical properties in paroxysmal, persistent, and permanent atrial fibrillation (AF) states. Alternans of APD and rate maladaptation in a spatially divergent way appear mechanistically involved in AF initiation, development, and persistence. The underlying pathophysiology warrants further investigation.

Keywords: Atrial fibrillation • Electrical remodelling • Monophasic action potential (MAP) • Action potential duration (APD) • Electrical restitution curve (ERC) • Rate maladaptation • Action potential duration (APD) alternans • Alternans • Electrical heterogeneity

Introduction

Atrial fibrillation (AF) currently affects 1–2% of the population with an expectation to increase over the next several decades.1 Atrial fibrillation serves as an adverse prognostic marker and is associated with, among other things, increased rates of death, stroke, and heart failure events.1 Both the prevalence of AF and the number of strokes attributable to AF increases in each decade of life.2,3 It is predicted that the prevalence of AF-related hospitalizations in the USA will exceed 3.8 and 5.6 million by 2025 and 2050, respectively.4

Several mechanisms of AF have been described. The prevailing concept has focused on interplay between triggers that induce and substrate that maintain arrhythmia.5 Pulmonary vein triggers are currently thought to be integral in the development of AF. Studies suggest that AF can be initiated by pulmonary vein focal impulses that propagate to atrial tissue, and that radiofrequency ablation of these foci can terminate AF.6 This review focuses on the role of action potential duration (APD) alternans as a ubiquitous preamble to AF initiation.

The role of monophasic action potential recording in studying the substrate of atrial fibrillation

Electrical remodelling of the atria and other structural changes are recognized as substrates essential for persistence of AF. Recording of monophasic action potentials (MAPs) by the ‘Franz’ contact electrode catheter have facilitated the investigation of both normal properties and pathological states of the atria, including AF. Recently, investigation of atrial MAP recordings in human AF resulted in a theory positing APD alternans as a dynamic substrate for AF.7 Moreover, APD alternans was noted to precede all AF episodes and was absent when AF did not initiate. To appreciate the role of the human atrial action potential in the initiation of AF, it is appropriate to review the MAP and how its relationship with AF continues to evolve.

Monophasic action potential recording

Localized myocardial depolarization and repolarization in the human heart in situ can only be investigated by recording MAPs.8,9 Under optimal conditions, MAPs record the repolarization time course of transmembrane action potentials (TAPs) with high fidelity.10 Monophasic action potentials can be recorded from contact with cardiac surface in contrast to TAPs, which require tedious in vitro preparations. As the MAP electrode diameter is prohibitively large from entering a single cardiac cell, the probable origination of the MAP is from the electrical gradient between the myocardium subjacent and adjacent to the catheter tip.10,11

Action potential duration

Precise measurement of total MAP duration is difficult because of the asymptotic end of repolarization. As a result, the MAP duration is usually determined at a repolarization level of 90% (or another fraction) with respect to the MAP amplitude. The MAP amplitude
is defined as the distance from the baseline to the crest of the MAP plateau voltage, not its upstroke peak.\textsuperscript{10,12} The intersection between the diastolic baseline and a tangent placed on phase 3 of the action potential may also determine MAP duration, but typically yields more arbitrary results.

In normal states, APD is dependently related to changes in heart rate, for which physiological advantages exist. A longer cycle length (CL) causes a requisite APD lengthening, providing a longer period of excitation–contraction coupling. Conversely, a shorter CL causes APD shortening and maintenance of a sufficient diastolic interval (DI), preserving coronary flow and ventricular filling.\textsuperscript{13} This APD–CL relationship can be expressed by the electrical restitution curve (ERC), which describes APD recovery time course as a function of the DI or CL from the most premature beat to long intervals following a steady-state APD.\textsuperscript{14–17} An ERC is obtained by pacing at a constant S1–S1 interval, followed by an S2, a single premature stimulus. S1–S2 coupling interval is progressively shortened until block of the premature pulse, or the effective refractory period. Figure 1 exhibits an example of an ERC, a graphical relationship between APD 90\% (or another fraction) to its preceding DI or CL.\textsuperscript{17–19} A modified technique for describing the APD–CL relationship, referred to as the ‘dynamic’ restitution curve, exhibits the relationship between APD and different steady CLs determined during the early phases of rate adaptation.\textsuperscript{15,17}

### Substrate for arrhythmia

It is possible that many mechanisms combine to initiate arrhythmia.\textsuperscript{20} Cycle length-dependent changes in APD are suspected to participate in arrhythmogenesis and, in particular, fibrillation.\textsuperscript{15–17} As ERCs describe, progressive shortening of CL and DI accentuate changes in APD. Increasing APD ‘alternans’ leads to larger wavelength oscillations and points of wavebreak resulting in temporal heterogeneity and reentry. The mechanisms by which APD alternans may contribute to electrical instability have been described utilizing computer and numeric simulations.\textsuperscript{20–22} The demonstration of APD alternans at high stimulus rates provides mechanistic support for this concept.\textsuperscript{23–26} Action potential duration alternans has also been linked to ventricular fibrillation (VF) vulnerability.\textsuperscript{27–29} Notably, there is now compelling evidence that APD alternans may lead directly to AF and its transition from atrial flutter (AFl) in animal models,\textsuperscript{30,31} in numerical simulations\textsuperscript{32} and directly in humans.\textsuperscript{7,33–35}

Pictorial assessment of APD alternans is typically performed spectrally. Utilizing validated software, APs of a contiguous selection are each denoted by beat number and time sample. A fast-Fourier transform is used to compute power spectra across beats for each time sample, and then summation of spectra across the AP is obtained.\textsuperscript{26} As shown in Figure 2, the magnitude of APD alternans is represented by the dimensionless k score utilizing the following equation: $k = \frac{\sum T - \mu_{\text{noise}}}{\sigma_{\text{noise}}}$, where
\[ \Sigma T \] is the spectral magnitude at 0.5 cycles/beat, and \( \mu_{\text{noise}} \) and \( \sigma_{\text{noise}} \) are the mean and standard deviation of noise. A \( k > 0 \) indicates that alternans exceeds noise.

It is postulated that the degree of APD alternans, which is related to the slope of the ERC, estimates wavebreak and the substrate for reentry. A steeper ERC slope represents a relatively small change in DI or CL yet renders large changes in APD and refactoriness. The ‘restitution hypothesis’ suggests that when the ERC slope > 1, augmented APD ‘alternans’ results in larger wavelength oscillations and points of wavebreak. Alternatively, when ERC slope < 1, oscillations progressively decrease and ultimately achieve a new steady state.16,27,35,37 In addition, studies have shown that drugs that flatten the restitution curve decrease the chance of VF induction and stabilize VF to ventricular tachycardia.38–40 These findings make this hypothesis quite an attractive explanation for the correlation between APD alternans and arrhythmogenesis, and suggest that a ‘flatter than normal’ ERC may be antiarrhythmic.

This raises the question, however, as to whether the steep early phase of the ERC is indeed hazardous and a predictor of VF (or AF). In normal myocardium, APD alternans in this early ERC phase dams up over a few beats as a rapidly adapting change in APD follows a sudden change in rate.14 This physiological APD rate adaptation leads to shortening of the APD as it adapts to the new steady-state CL, which allows the DI to get longer and the APD to move quickly onto the flat portion of the ERC and away from the vulnerable window and threshold for greater alternans.15 The steep initial slope of the ERC may, in fact, be protective as it facilitates APD ‘movement’ to a flatter, safer portion of the ERC. Flattening of the ERC by medications will indeed flatten the slope and essentially extinguish APD alternans. This is, however, not physiological and may have other adverse effects on cardiac performance. Of note, myocardial ischaemia also flattens the ERC slope, which obviously is not an antiarrhythmic scenario.41

**Atrial electrophysiological properties and atrial fibrillation**

The normal atrial AP is triangular with a relatively short APD compared with the ventricular myocardial tissue.42,43 Depolarizing currents, both inactivating Na\(^+\) current (\( I_{\text{Na}} \)) and the L-type Ca\(^{2+}\) current (\( I_{\text{CaL}} \)), are the same as in ventricular myocytes. Atrial myocytes, however, express certain ion channels and channel subunits nearly absent on ventricular myocytes, including ultra-rapid delayed rectifier current (\( I_{\text{Kur}} \)), which initiates atrial repolarization faster than the ventricular myocyte predominant rapid delayed rectifier current (\( I_{\text{Kr}} \)).44 The transient outward K\(^+\) current (\( I_{\text{TO}} \)), which greatly contributes to early rapid phase 1 repolarization, is more heavily distributed on normal atrial than ventricular myocytes and likely facilitates a shorter atrial APD.45 In addition, Ca\(^{2+}\)-activated potassium channel (\( I_{\text{BKCa}} \)) was discovered to have greater effect and density in the atria compared with the ventricle.46 These and other atrial electrophysiological features afford complex conduction patterns at rapid atrial rates.45 Further, the atrial \( I_{\text{KATP}} \) channel is of great importance, but is beyond the scope of this review.

Atrial electrophysiological remodelling as a consequence of AF was significantly advanced by work conducted in animal models.47,48 Further studies also suggested that AF induced atrial APD shortening and action potential plateau phase depression in animals.49,50 as well as humans.50–53 Alterations in ion channels have been proposed to contribute to these electrophysiological observations.4 AF induces abnormal cellular calcium handling that may promote compensatory downregulation of inward L-type Ca\(^{2+}\) current (\( I_{\text{CaL}} \)).53–56 It is theorized that this reduction in \( I_{\text{CaL}} \) activity mitigates the plateau phase and largely explains the accentuated shortening of atrial APD.45,57 Transient outward K\(^+\) current (\( I_{\text{TO}} \)) is also reduced in chronic AF,43,45,58,59 which seems puzzling but may help explain additional effects of AF on APD. Chronic AF also exhibits an increase in inward rectifying K\(^+\) current \( k_{1} \)59,60 which primarily controls the resting potential of the atrial myocyte.61 In sum, a complex interplay of currents in atrial AF myocytes renders a shorter APD and mitigation of the plateau phase.

While the time course of the changes in atrial refractoriness may well be explained, at least in large part, by ionic changes, it typically occurs before the development of persistent AF.57,49 This suggests that factors other than atrial electrophysiological changes contribute to the development of AF.56,62 This ‘second factor’ hypothesis includes several potential contributors, including atrial dilation, stretch, and connexin and gap junction remodelling.56,63–65

An interesting phenomenon, first noted in goats by Wijffels et al.,47 was an alteration to physiological rate adaptation. As expected, normal goats in sinus rhythm exhibited atrial APD shortening at shorter pacing intervals. Goats artificially maintained in AF over 24–48 h, however, lost physiological adaptation and exhibited either constant or shorter APD at slower heart rates. Further, normal heart rate adaptation was restored within a few days after cardioversion to sinus rhythm. A later study by VanDerVelden et al.66 added that increasing pacing rates rendered longer APD in a post-cardioversion chronic AF goat model. This constitutes an AF-induced, paradoxical rate adaptation of the atrial APD to changes in heart rate. This maladaptation of the atrial APD had also been observed in prior human investigation57,68 as well as animal.49,69–71 models. More, recently, the studies by Narayan et al.74 also found this to be present in human atria.

**What is the mechanism?**

Ionic mechanisms for this rate-maladaptive response have been proposed. As noted earlier, \( I_{\text{TO}} \) is heavily distributed in normal atrial myocytes but is significantly reduced in atrial myocytes affected by longer periods of AF. This decrease in \( I_{\text{TO}} \) lengthens early rapid repolarization and may influence the rate maladaptive response.54 Downregulation of the \( \alpha_{1C} \) subunit of the L-type Ca\(^{2+}\) channel66 and general cytosolic Ca\(^{2+}\) overload50 may contribute while ryanodine may be protective,59 suggesting that abnormal intracellular Ca\(^{2+}\) handling and Ca\(^{2+}\) overload may be complicit in the maladaptive rate response.

Narayan et al.34 documented atrial APD alternans in humans during progressive disorganized alteration of AF to AF, and noted that the presence of rate maladaptation leads to APD alternans preceding all such transitions. Conversely, much less APD alternans was seen in patients in whom AF did not develop. A critical observation, suggested in other investigations, is of spatial heterogeneity within the atria.33,74 and electrical heterogeneity of rate maladaptation in AF and AF contributing to AF development and
Compared with other atrial sites, the isthmus represented an area in which faster atrial rates resulted in less robust APD shortening, a hallmark of maladaptation. In these maladaptive areas, pacing stimuli approached a partially or fully refractory myocardium more often than in ‘normal’ myocardium. The immediately following APD and DI then underwent compulsory lengthening, facilitating APD alternans in the isthmus. Similar observations subsequently were made in the left human atrium, near the pulmonary veins.55

It was also shown that continued incremental pacing enhanced APD alternans until beat-to-beat conduction was unable to be maintained physiologically, leading to 2 : 1 conduction block, which was also noted in prior investigation.19 Narayan et al. posited that, as a result of a maladaptive APD rate response, spatially heterogeneous APD alternans at increasing heart rates leads to wavefront fractionation and wavebreak, conduction block, and the eventual transition to AF.34

Narayan et al.7 further evaluated the concept of heterogeneity of APD maladaptation and alternans during initiation of AF in the electrophysiology laboratory in patients with and without clinical AF. As noted above, APD alternans is largely reported at fast heart rates and may conceptually be explained by restitution. His group hypothesized that AF-induced atrial remodelling may cause APD alternans at slower heart rates. Using an incremental pacing technique, the group studied left and right atrial APD90, APD restitution curve, APD alternans, and non-alternating complex APD oscillations on transitions to AF in 12 patients with persistent AF, 13 patients with paroxysmal AF, and 8 control patients without clinical AF.

The development of APD alternans was elicited at near-resting rates in patients with persistent AF, at intermediate rates in those with paroxysmal AF, and only at very rapid rates just prior to AF in controls. Moreover, the amplitude of the APD alternans also varied by group, and was greatest in those with permanent AF, intermediate in those with paroxysmal AF, and smallest just before AF induction in controls. A representative patient with persistent AF is shown in Figure 3, in whom marked left atrial APD alternans was seen at CL 600 ms (100 b.p.m.) and CL 500 ms (120 b.p.m., illustrated). At faster rates (shorter CLs), APD alternans disorganized to complex APD oscillation immediately preceding AF transition. In stark contrast, Figure 4 illustrates that APD alternans in control subjects did not arise until very fast rates (CL 200–250 ms), just prior to AF onset. No control subject had APD alternans at CL ≥250 ms and, when exhibited, alternans had very small magnitude. Moreover, the APD rate–response relationship and APD alternans directly indicated AF ‘substrates’, presumably caused by electrical remodelling. The CL onset of APD alternans and CL range varied among the three groups. As delineated in Figure 5, persistent and paroxysmal AF subjects exhibited APD alternans at all rates while controls only exhibited APD alternans around CL 250 ms.

The suggestive role of APD alternans and heart rate was also evaluated. The ‘restitution hypothesis’, described earlier, is the concept that an ERC slope > 1 indicates greater APD alternans at a faster rate and will facilitate wavebreak and arrhythmia development. This theory was supported by the initiation of APD alternans in some control subjects as well as a few paroxysmal AF
subjects, in whom spontaneous premature atrial complexes triggered AF by increasing the ERC slope.

Groups with paroxysmal and permanent AF did not exhibit similar restitution kinetics. The ERC slope in all paroxysmal and persistent AF subjects was \( \leq 1 \) at the onset of APD alternans, even if maximum slope was \( > 1 \) at faster rates. The observation that persistent AF leads to APD alternans at slower heart rates with ERC slopes \( \leq 1 \) prompts inquiry into the mechanistic explanation of this phenomenon. As noted earlier, AF is thought to affect several aspects of atrial myocyte character and function, including abnormal intracellular Ca\(^{2+}\) handling, membrane ion current remodelling changes, and atrial conduction slowing. These and other mechanisms could potentially contribute to the observations of this study and require additional investigation.

Corroborating prior investigation, atrial APD alternans was noted to precede every AF transition, suggesting that APD alternans may be mechanistically coupled to transitions to AF. Accordingly, in addition to spontaneous premature atrial complexes, APD alternans amplification or complex oscillation preceding AF transition in fast rates may represent multiplying periods in non-linear systems, another suspected trigger for AF. Proposed mechanisms to elucidate this phenomenon are somewhat nebulous at this time and require further investigation.

**Conclusion**

Atrial fibrillation is common in our ageing population and is recognized as an adverse prognostic marker. The prevailing concept of AF mechanism focuses on interdependence of triggers and substrate, and MAP recordings illustrate differences between normal myocardial properties and AF substrate. While prior investigation...
suggests an arrhythmogenic nature of APD alternans in the ventricle, work highlighted in this review supports this concept in the atria and reports the role of APD alternans as a vital precursor to AF development. This idea, along with support from corroborating efforts, will likely alter our current notion of AF development, uncover new areas for exploration, and hopefully advance the formulation of therapeutic options.

Conflict of interest: S.M.N. reports being co-inventor on intellectual property owned by the University of California and licensed to Topera Medical, Inc. S.M.N. holds equity in Topera. Topera has not sponsored any research, including that presented here. S.M.N. also reports having received honoraria from Medtronic, St. Jude Medical and Biotronik Corporations and grant support from Biosense-Webster. All other authors have declared no conflict of interest.

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