The association of low voltage atrial electrograms and the myocardial substrate for atrial fibrillation

Abstract

Abnormal human atrial muscle has elevated resting membrane potential, depressed maximal amplitude of the action potential, and decreased upstroke velocity. These electrical abnormalities are histologically correlated to degenerative changes of the atrial muscle structure. Investigations based on the recording of abnormally prolonged and fractionated atrial endocardial electrograms during sinus rhythm have provided important knowledge about the electrophysiological properties of the pathological atrium. Human atrial tissues, where fractionated electrograms are recorded, consist of muscle fibers that are widely separated and their orientation distorted by connective tissue. These histological changes result in decreased intercellular connections, which might increase resistance to current flow, and thereby slow conduction. The potential clinical significance is that detection of abnormal atrial endocardial electrograms during sinus rhythm in patients with atrial vulnerability may help to identify those with greater atrial conduction defects and higher incidence of induced and spontaneous episodes of AF. 3D electroanatomic mapping (EAM) has been utilized in clinical electrophysiological studies for substrate description in atrial arrhythmias. Using this 3D EAM system low bipolar endocardial voltage can be identified and localized for guiding catheter ablation. Voltage mapping with 3D electroanatomic mapping is based on the correlation of the registration of low-voltage areas (<0.5 mV) in the left atrium with endocardial scar and/or structural defects as an atrial substrate. These low-voltage zones are the target for individualized substrate ablation or modification. Patients with low-voltage substrate have lower success rates after pulmonary vein ablation. However, this low success rate can be significantly improved by targeting atrial substrate zones with low voltage areas. Although more clinical evidence should be further generated, patient specific substrate ablation targeting atrial fibrosis estimated by bipolar voltage mapping is a promising alternative strategy for additional substrate modification beyond PVI ablation for AF.

Keywords: electroanatomic mapping, PVI ablation, atrial muscle, atrial fibrillation, 3D, abnormal atrial electrograms

Abbreviations: EAM, electroanatomic mapping; AF, atrial fibrillation; AAE, abnormal atrial electrograms; LVZ, low voltage zone

Introduction

Atrial fibrillation (AF) is a common arrhythmia associated to lethal complications such as clinical outcomes. Pathological changes in the atrial myocardium may lead to electrophysiological produce decreasing local voltage, conduction block and low electrogram amplitude. Indeed, atrial degenerative changes of the atrial muscle structure. Investigations based on the recording of abnormally prolonged and fractionated atrial local electrograms during sinus rhythm have provided important knowledge about the electrophysiological properties of the pathological atrium. Human atrial myocardium where abnormal atrial electrograms are recorded, consist of muscle fibers that are widely separated and their orientation distorted by collagen. These histological changes result in decreased intercellular connections, which might increase resistance to current flow, and thereby slow conduction. Therefore, abnormally prolonged and fractionated atrial electrograms result in an irregular atrial conduction. This irregularity is characterized by a non-homogeneous local electrical activity, related to an anisotropic, non-uniform and delayed conduction through a pathological atrial myocardium. In such altered atrial substrate arrhythmias may develop due to reentry. The 3D electroanatomic mapping (EAM) revealed anatomic and structural atrial abnormalities that were associated with a change in the nature of sinus pacemaker activity. In addition, there were loss of the normal multi-centric pattern of onset of sinus impulse, shift of the pacemaker complex to low crista terminalis sites, and the presence of abnormal and circuitous conduction around lines of conduction block. 3D electroanatomic bipolar voltage mapping is a diagnostic auxiliary method that has been utilized in clinical electrophysiological studies for substrate description in atrial arrhythmias. Recent advances in 3D EAM systems have facilitated catheter manipulation within the atrium at reduced fluoroscopy exposure time. Using this 3D EAM system low bipolar endocardial voltage can be identified and localized for guiding catheter ablation. The electrical information obtained can be used in different ways to record different information. For example, certain map points recorded can be utilized for the color-coded display of the electrical activation sequence known as “activation mapping”. In addition, the display of post-pacing intervals known as “entrainment mapping” can be recorded, as well as, the
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display of unipolar/bipolar electrograms as part of “fractionation” or “voltage mapping”. Therefore, we aim to analyze the association of low voltage atrial electrograms and the myocardial substrate for the development of atrial fibrillation.

Abnormal atrial electrograms and the myocardial substrate

During an atrial depolarization the axial current flows from one myocardial cell to the adjacent one through the intercalated discs, which normally have a relatively low resistance. Under normal tissue conditions some mild disturbances in conduction of cardiac impulses may occur. The conduction perpendicular to the longitudinal axis of the atrial muscle fibers can be delayed because in that direction the effective axial resistance is greater. This greater axial strength results in part from the scarce amount and short length of the intercalated discs in the lateral-lateral direction of the muscle fibers, compared to the terminal-terminal direction. There is an added alteration to the electrical conduction when the atrial muscle fibers are separated by connective tissue. In this context, the conduction properties can be further altered due to the effects on the axial resistance creating a discontinuous and anisotropic propagation. In this regard, Spach et al., have demonstrated in a nicely done work that the slowing of the conduction velocity causes a decrease in the amplitude and, an increase in the duration of the extracellular electrogram of the canine Purkinje system. In a computerized model of electrogram generation study, Lesch et al., has shown that the decreased conduction velocity was responsible for the increase in electrogram duration, while increased intracellular resistance was responsible for the fractionation of the electrogram.

Figure 1 Sites of endocardial catheter mapping of the right atrium are depicted.

The atrial electrograms were recorded from 12 endocardial sites in each patient. The sites listed from A to D correspond to the upper part of the right atrium while the sites from E to H are from the middle part, and those from I to L are from the lower part of the right atrium. The sites marked A, E and I correspond to the anterior region, sites B, F and J to the lateral region, sites C, G and K to the posterior region, and sites D, H and L correspond to the medial region of the right atrium. SVC: superior vena cava. IVC: inferior vena cava. Ao: Aorta. PA: pulmonary artery. LA: left atrium. RV: right ventricle. LV: left ventricle.

In the lower part (A) there is an abnormal endocardial atrial electrogram with a duration of 130 ms and 10 fragmented deflections. (B) shows a normal endocardial atrial electrogram that lasts 80 ms in duration and has two fragmented deflections. Reprinted with permission from Centurion OA et al. Influence of advancing age on fractionated right atrial endocardic electrograms. Am J Cardiol 2005; 96: 239-242.

More than a quarter of century ago, Centurión et al., designed a study to assess the importance and significance of recording of abnormal atrial electrograms (AAE) during sinus rhythm in AF patients (Figure 1). We showed that patients who had AAE had a significantly increased atrial vulnerability compared to those with normal electrograms. Abnormal atrial electrograms showed a very good specificity and positive predictive value when evaluating the induction of sustained AF. The specificity demonstrated was 94% with a positive predictive value of 93%. We have also demonstrated a significantly longer conduction time of sinus impulses in AF patients who possess AAE within the right atrium. In our study the mean intra-atrial (54 ms) and the mean interatrial conduction time (100 ms) of sinus impulses were significantly longer in the AAE (+) Group.

In addition, AAE also proved to be a useful marker to differentiate patients with more extensive atrial electrical abnormalities who are more likely to have inducible AF. Prolonged and fractionated AAE may reflect inhomogeneous local electrical activity related to a delayed and non-uniform anisotropic conduction through a diseased atrial muscle (Figure 2). The potential clinical significance is that detection of AAE during sinus rhythm in patients with atrial vulnerability may help to identify those with greater atrial conduction defects and higher incidence of induced and spontaneous episodes of AF.

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3D electroanatomic mapping (EAM) has been utilized in clinical electrophysiological studies for substrate description in atrial arrhythmias. Using this 3D EAM system low bipolar endocardial voltage can be identified and localized for guiding catheter ablation. Voltage mapping with 3D electroanatomic mapping is based on the correlation of the registration of low-voltage areas (<0.5 mV) in the left atrium with endocardial scar and/or structural defects as an atrial substrate (Figure 3). These low-voltage zones are the target for individualized substrate ablation or modification. Patients with low-voltage substrate have lower success rates after pulmonary vein ablation. However, this low success rate can be significantly improved by targeting atrial substrate zones with low voltage areas. In a study with 3D electroanatomic mapping, it was found an age-related evidence of generalized conduction slowing in the atria. They also demonstrated increasing numbers of fractionated electrograms and double potentials in the atrium but along the crista terminalis with aging in patients prone to develop AF. This fact may be important because functional conduction delay at this structure has been implicated in the development of a range of atrial arrhythmias. In this regard, Sanders et al. designed a study with 3D electroanatomic mapping in order to search for evidence of diffuse atrial abnormalities in patients with a propensity to develop AF. They performed 3D EA mapping to assess regional conduction velocity, double potentials, fractionated electrograms, regional voltage, and areas of scar tissue. They demonstrated significant increase in atrial ERP at all right atrial sites, increased atrial conduction time along the lateral RA and CS, prolongation of the P-wave duration, and greater number and duration of double potentials along the crista terminalis. The authors described a significant regional conduction slowing with double potentials and fractionation associated with areas of low voltage and scar. These regions of low voltage and fibrous tissue were particularly marked along the crista terminalis. These abnormalities resulted in propagation detours and colliding activation wave fronts of atrial activation.

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Indeed, their patients with prolonged sinus pauses had a higher percentage of complex electrograms, higher right atrium volumes, and lower voltage in the high lateral right atrium than control patients regardless of age. In previous studies it was found remodeling changes of both atria in lone-AF patients, including structural abnormalities, lower conduction velocities, increased effective refractory periods, abnormal atrial histologic results, and inflammation in patients with sinus pauses greater than 3 seconds or unexplained sinus bradycardia (<40 beats/min).36–38

It is well known that AF is associated with structural, electrical, and contractile remodeling of the atria. Atrial fibrosis certifies structural remodeling of the atrium and is considered to be the substrate for AF maintenance.39–43 Atrial fibrosis yields the pathological substrate to promote AF by interfering depolarization wave continuity through the atrial myocardium, developing conduction alterations, and causing anisotropic tissue conduction.44 Human histopathological results demonstrated that large extent of atrial fibrosis is found in the atria of AF patients and, the extension of atrial fibrosis was significantly associated with AF and duration of the arrhythmia.44–46 Atrial septal biopsies obtained from patients with lone paroxysmal AF have revealed inflammatory infiltrates consistent with myocarditis and fibrosis.45

Low-voltage electrograms are produced by diverse mechanisms and, the primary mechanism has been suggested to be atrial fibrosis.45–52 These atrial changes may reflect poor fiber coupling, non-homogeneous conduction, and anisotropic tissue depolarization caused by atrial myocardial fibrosis. The definition of low voltage zone (LVZ) still remains controversial, mainly due to lack of histopathological studies. Sanders et al.32,33 tried to define areas of low voltage. They performed electroanatomical maps of the right atrium by pacing the distal coronary sinus or mapping during sinus rhythm in order to study the electrophysiological atrial substrate for AF patients. They identified electrically silent areas, namely, scar defined by the absence of recordable activity or a bipolar voltage amplitude ≤0.05 mV; and low-voltage areas defined by atrial zones with bipolar voltage ≤0.5 mV. Later on, Verma et al.,29 utilized the same definition criteria for the left atrium. They demonstrated that pre-existent left atrial fibrosis is a significant independent predictor of recurrence after PVI. Yamaguchi et al.,29 contrasted the long-term clinical outcomes after PVI alone in patients with and without LVZs defined as <0.5 mV. The incidence of AF free- survival after PVI alone was significantly lower in patients with LVZs, and the extent of the LVZ was identified to be an independent risk factor. Masuda et al.,53 also demonstrated similar results showing that the presence of LVZ predicts AF recurrence. The DECAAF multicenter and prospective clinical trial showed that atrial fibrosis estimated by LGE-MRI was independently associated with the likelihood of AF recurrence in patients undergoing AF ablation.54

It is undeniable the useful help of delayed-enhancement MRI in detecting atrial fibrosis. The group of Marrouche et al.,14 reported on the efficacy and utility of LGE-MRI in identifying areas of fibrosis in the left atrium, and observed that an enlarged size of left atrial wall enhancement was significantly related to recurrence after PVI ablation. They also found a significant correlation between enhancement on LGE-MRI and low-voltage areas on voltage-guided mapping. It is very interesting that left atrial wall biopsies showing fibrosis were associated with zones of left atrial wall enhancement on LGE-MRI. On the contrary, areas with normal biopsy tissue were correlated with non-enhanced regions, demonstrating how accurate LGE-MRI is for atrial fibrosis detection.

There are several studies showing that AF ablation targeting atrial fibrosis by voltage mapping may improve clinical results. Indeed, several investigations demonstrated the effectiveness of voltage mapping of the atrium in persistent AF patients.55–59 However, the limitation is that most of these investigations were small and observational studies in nature. A couple of years ago, Yang et al.60 demonstrated similar success results in the one year and a half follow-up, with lower procedure time and fluoroscopy exposition and durations, as well as, shorter energy delivery time. It is very interesting the fact that over half of non-paroxysmal AF patients did not need further ablation beyond pulmonary vein isolation.55–59 Last year, Kircher et al.,61 also observed the effectiveness of AF ablation guided by voltage mapping in a randomized single-center research. The START Trial is an ongoing large, multicenter, randomized study that was designed to analyze the safety and efficacy of left atrial voltage-based ablation comparing two AF ablation strategies, namely, PVI plus LVZ homogenization Vs. PVI alone in patients with LVZ (https://www.umin.ac.jp/ctr/index.htm, UMIN000022119). This ongoing trial will provide further knowledge into this AF ablation strategy.62

Conclusion

In conclusion, investigations based on conventional atrial catheter mapping with the recording of abnormally prolonged and fractionated atrial local electrograms during sinus rhythm, as well as, their characteristic distribution within the right atrium have provided important knowledge about the electrophysiological properties of the pathological atrial myocardium in AF patients. The detection of abnormal atrial electrograms during sinus rhythm identifies a group of patients with increased atrial vulnerability and a significantly higher incidence of spontaneous or induced episodes of atrial fibrillation. The 3D EAM revealed anatomic and structural atrial abnormalities and the presence of abnormal and circuitous conduction around lines of conduction block. Although more clinical evidence should be further generated, patient specific substrate ablation targeting atrial fibrosis with low voltage zones estimated by bipolar voltage mapping is a promising alternative strategy for additional substrate modification beyond PVI ablation for AF.

Acknowledgements

None.

Conflicts of interests

Author declares that there is no conflict of interest.

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