Predicting arrhythmias in primary prevention heart failure patients: picking up the fragments

Nathan Engstrom,1 Hayley Louise Letson †,1 Kevin Ng,2 Geoffrey Phillip Dobson1

ABSTRACT
Identifying patients with high-risk heart failure (HF) who would benefit from an implantable cardioverter-defibrillator (ICD) remains controversial. A potential marker for arrhythmic sudden death is fragmented QRS (fQRS). fQRS is the notching and slurring of the QRS complex in a 12-lead ECG and it indicates abnormal ventricular depolarisation and myocardial scarring and fibrosis. However, before fQRS complex can be included into selection criteria for ICD therapy, more complete reporting is required on their association with malignant arrhythmias, left ventricular remodelling and myocardial scarring/fibrosis in patients with HF. The molecular basis of the fQRS-arrhythmia-fibrosis connection in HF also needs to be explored. It is not widely appreciated that changes in the QRS complex and phases 0 and 1 of the ventricular action potential occur before contraction and predetermine Ca2+ release during contraction and later Ca2+ sparks. It is currently not known whether different zig-zag patterns of the QRS are associated with aberrant Ca2+ cycling and arrhythmogenic sparks in patients with HF.

INTRODUCTION
Despite major advances in treating patients with severe heart failure (HF), deciding who should receive an implantable cardioverter-defibrillator (ICD) remains challenging. Over 200,000 ICDs are implanted globally each year; however, the major cause of death in these patients (up to 70%) is not arrhythmic sudden cardiac death (SCD).1–3 Current risk stratification for ICD implantation and SCD include a New York Heart Association functional classification II–III, reduced left ventricular ejection fraction (LVEF <35%) with no improvement after 3 months of maximal therapy, QRS <120 μs, a predicted survival >1 year, age and comorbidity.1 2 4 5 Of all these risk factors, the single most important criterion is LVEF <35%.5 However, a substantial number of patients with HF at risk of SCD have LVEF >35% and would not qualify for an ICD.1 2 5 Previous studies in patients with HF with preserved ejection fraction showed that 20–40% of cardiovascular deaths were SCD, although the underlying mechanisms remain unknown.6 Clearly the current international guidelines for primary prophylaxis ICD in patients with HF are inadequate.

Fragmented QRS as a potential prognostic marker of arrhythmic sudden death
Additional prognostic criteria are urgently required to maximise the benefit of ICD therapy, and these include genetic testing, circulating biomarkers, improved knowledge of the myocardial scar and fibrosis and changes in the ECG waveform.2 7 In this viewpoint, we discuss the potential role of one or more fragmented QRS (fQRS) complexes to predict malignant ventricular arrhythmias and sudden death in primary prevention patients with HF. fQRS is defined as high-frequency notching and slurring within the QRS complex that reflects abnormal ventricular depolarisation and structural (anatomical) and functional (electrical) pathophysiology.1 7 The term fQRS was coined by Boineau and Cox in 1973, who reported the appearance of fragmented electrograms and re-entrant activity in dogs after coronary occlusion.1 7 Today, the most common fQRS forms include an additional R wave or notching in the lowest point of the S wave, or the presence of >1 R wave in the QRS complex measured in two contiguous leads, corresponding to a major coronary territory1 7 (figure 1).

The idea of linking fQRS to malignant arrhythmias is not new. It dates back to the early 1950s with Langner’s electrocardiographic investigations.8 Langner was among the first to show notching and slurring of an expanded ECG in patients following a ‘healed’ myocardial infarction.8 The association between fQRS and cardiomyopathy did not appear, however, until 1969 in a landmark study of Flowers and colleagues.9 This was followed 40 years later by another important study of Das and colleagues, which included primary and secondary prevention of patients with HF.7 The presence of fQRS...
has subsequently been shown to be an independent arrhythmic marker in patients with Brugada syndrome, right ventricular cardiomyopathy and other cardiac and non-cardiac pathologies. The next step was to identify and quantify the underlying causes of the fQRS complex, and how this zig-zag pathology leads to dyssynchrony of LV systolic function, and possibly ventricular fibrillation or ventricular tachycardia. Das and colleagues suspected that the primary substrates for fQRS complexes were myocardial scarring, fibrosis or inflammation, which has subsequently been confirmed. Spatial locations of scars and non-viable myocardium are predicted from a 12-lead ECG and confirmed using a variety of methods, including single photon emission tomography (SPECT), magneto-cardiography (MCG) and late gadolinium enhancement cardiac MRI (Ga-MRI).

**Paucity of high-quality experimental and clinical data**

Despite a growing interest in fQRS as a prognostic marker for ventricular arrhythmias, little progress has been made in assessing its clinical usefulness in patients with HF. This appears largely due to a paucity of high-quality experimental and clinical data. We recently performed a systematic review and meta-analysis of studies examining fQRS in patients with HF with or without an ICD, and with LVEF ≤ 40%. Outcome measures were ventricular arrhythmias and all-cause mortality. Using these search criteria, we analysed 10 studies involving 3885 patients and found that fQRS was statistically associated with ventricular arrhythmias and all-cause mortality. Furthermore, our study revealed a number of knowledge gaps in current reporting. None of the studies included all major forms of fQRS (narrow, wide, paced, premature ventricular contraction) (table 1), and only three reported on coronary artery territory. In addition, there were no uniform criteria on what constituted a ventricular arrhythmic event, which precluded post hoc assessment of what type of arrhythmia triggered ICD shocks. A further complication was the failure to separate primary and secondary prevention patients with HF as well as those with ischaemic (ICM) vs non-ICM (NICM) cardiomyopathies. There were also concerns on the reproducibility of ECG measurements, fQRS characterisation, counting of leads and intra and interobserver variability. Discounting leads and subjective visual yes/no diagnoses can make some studies inaccurate and non-reproducible. This lack of complete reporting illustrates the current gaps in knowledge when assessing fQRS as a potential predictor of ventricular arrhythmias and sudden death.

<table>
<thead>
<tr>
<th>Type</th>
<th>Definition</th>
<th>Morphology</th>
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<tr>
<td>Narrow</td>
<td>QRS duration &lt;120 ms with or without Q waves and has an additional R wave or notching at the lowest point of the S/R wave or the existence of &gt;1 R wave in 2 or more successive leads corresponding to a coronary artery (CA) territory.</td>
<td>Narrow</td>
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<tr>
<td>Wide</td>
<td>QRS duration ≥120 ms with or without Q waves and is explained with &gt;2 R waves or notching at the lowest point of the S/R wave or the existence of &gt;2 R waves in 2 or more successive leads corresponding to a CA territory.</td>
<td>Wide</td>
</tr>
<tr>
<td>Paced</td>
<td>QRS &gt;120 ms without any evidence of fusion when the beat is initiated by a pacing spike in pacemaker and ICD. Is explained with &gt;2 R waves or notching at the lowest point of the S/R wave or the existence of &gt;2 R waves in two or more successive leads corresponding to a CA territory.</td>
<td>Pacing</td>
</tr>
<tr>
<td>PVC</td>
<td>A premature ventricular contraction (PVC) with no evidence of supraventricular fusion and is explained with &gt;2 R waves or notching at the lowest point of the S/R wave or the existence of &gt;2 R waves in 2 or more successive leads corresponding to a CA territory, or if 2 notches in R wave are present and &gt; 40 ms apart.</td>
<td>PVC</td>
</tr>
<tr>
<td>Q wave</td>
<td>Notching in the Q wave can occur in any part of the Q wave. However, it must be negative (under the baseline). The exception is a Q wave occurring at the onset of the R wave (Q-R borderline-fQRS).</td>
<td>Descending, In-peak, Ascending, Q-R Border</td>
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<tr>
<td>R wave</td>
<td>Termed “notched R” classifies different R wave morphologies. A positive fragmentation must be present and can occur in any part of the ascent of the R wave. This is found at the R peak. However, the upper segment must be &gt;50% of the declining R wave. If occurring at the peak of the R wave the difference in amplitude cannot be &gt; 1.0 mm/0.1 mV. R-S borderline fQRS appears in borderline area of the R and the S wave and stretches across both sides of the baseline and can be both positive or negative.</td>
<td>Notched R Ascending in R peak Notched R in Upper Descending Notched R in Lower Descending</td>
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<tr>
<td>S wave</td>
<td>Notched S occurs in the descent of the S wave, at the top or in the ascending part of the S wave and is termed a “notched S” and must always have a negative QRS. The amplitude difference between the top or the peak of the S wave and the second QRS component cannot be &gt;1.0mm/0.1 mV. R-S borderline fQRS appears in borderline area of the R and the S wave and stretches across both sides of the baseline and can be both positive or negative.</td>
<td>Descending, In-S peak, Ascending, Q-R Border</td>
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Towards a functional linkage between fQRS, myocardial scar/fibrosis and arrhythmias

There is an urgent need for more complete and accurate reporting on the number, type, location and timing of formation of fQRS and its association with myocardial scar/fibrosis and arrhythmias during patient screening and after ICD implantation (table 1). Location and timing of formation of non-viable myocardium are important because 38–50% of patients with ICM have a myocardial scar (replacement fibrosis after necrosis), and up to 75% of patients with NICM have patchy or diffuse fibrosis from different secondary remodelling events. Triggering arrhythmias in patients with HF depends on many factors, including spatial variation, location and density of the scar and/or fibrotic architecture in the myocardium. For example, the onset of arrhythmias around a scar and/or ICM regions largely result from heterogeneous activation due to regional conduction slowing or block (figure 2), whereas arrhythmias triggered from a failing myocardium with more diffuse fibrosis involves a more spiral wave formation from myocyte–fibroblast interfaces, particularly where fibrosis encircles preserved myocytes (figure 2). Diffuse fibrosis also appears to be more strongly associated with arrhythmias than scar tissue secondary to myocardial infarction and may help explain why patients with NICM with fQRS have a significant 2- to 3-fold increased incidence of death compared with patients with ICM. The sensitivity and specificity of fQRS to predict a scar in patients with ICM or NICM is around 80–90% using SPECT, MCG or Ga-MRI, and future studies could benefit from including these quantitative assessments. Recently, late gadolinium enhancement was used in the prospective multicentre study of Klem and colleagues, and they showed in over 1000 patients with NICM that myocardial fibrosis was independently and strongly correlated to arrhythmic events and SCD. In this landmark study, no significant prognostic association was found between LVEF ≤ 35% and risk of sudden death, which provides further evidence why LVEF < 35% should not remain a major driver for determining a patient’s eligibility for an ICD. Unfortunately, fQRS, myocardial fibrosis and sudden death were not included in their study, and it again highlights the need for more studies in this area. Postimplant monitoring could include fQRS.

Table 1 Gaps in knowledge to assess fragmented QRS (fQRS) as a potential prognostic candidate for primary prevention ICD placement in heart failure (HF) patients

<table>
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<th>Number</th>
<th>Focus areas</th>
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<tbody>
<tr>
<td>1.</td>
<td>Identify the different forms of fQRS, that is, narrow, wide, paced and PVCs, as part of routine patient screening (see figure 1).</td>
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<td>2.</td>
<td>Specify location of the different fQRS forms according to coronary artery territory.</td>
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<tr>
<td>3.</td>
<td>Quantify proximity of fQRS to a myocardial scar/fibrosis region and functional status.</td>
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<tr>
<td>4.</td>
<td>Determine timing of fQRS formation during screening and post-implant follow-up.</td>
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<td>5.</td>
<td>Separate ischaemic from non-ischaemic cardiomyopathy patients and investigate sex differences.</td>
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<tr>
<td>6.</td>
<td>Provide a full evaluation of appropriate and inappropriate ICD shocks and unresponsive ICD therapy events.</td>
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<tr>
<td>7.</td>
<td>Quantify the nature and type of arrhythmias that triggered ICD shocks and rate of arrhythmia (VT, VF).</td>
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<tr>
<td>8.</td>
<td>Provide ICD type (dual chamber, single chamber, bi-ventricular or subcutaneous).</td>
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<tr>
<td>10.</td>
<td>Document status of co-morbidities (eg, obesity, diabetes, renal disease, liver disease, COPD, sleep apnoea and others).</td>
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COPD, chronic obstructive pulmonary disease; ICD, implantable cardioverter-defibrillator; PVC, premature ventricular contraction; VF, ventricular fibrillation; VT, ventricular tachycardia.

Figure 2 (A) In the healthy heart, the vector nature of the electric field provides visualisation of the timing of the QRS complex and helps to identify the extent and location of non-viable myocardium in diseased hearts, including those regions responsible for fQRS. (B) Differences in arrhythmia generation in ICM and NICM patients. ICM patients have more localised scarring compared with more diffuse fibrosis in NICM patients. These different pathologies generate different mechanisms to form different fQRSs and to initiate VT or VF (see text). Other mechanisms of VF and VT initiation include afterdepolarisations and enhanced automaticity (not shown). ATP, antitachycardia pacing; CAD, coronary artery disease; fQRS, fragmented QRS complexes; ICM, ischaemic cardiomyopathy; NICM, non-ischaemic cardiomyopathy; VF, ventricular fibrillation; VT, ventricular tachycardia.
and its proximity to the scar/fibrosis, fibrotic density, left ventricular geometry, nature and type of arrhythmias, the number of appropriate and inappropriate shocks, sex differences and disease progression (table 1). With this new information, the prognostic value of fQRS can be more fully assessed.

Molecular basis of fQRS, delayed Ca\textsuperscript{2+} cycling and arrhythmias

Alongside wider and more accurate reporting, there are other gaps in knowledge on the mechanisms responsible for fQRS notching and shuffling in the viable myocardium itself, beyond scar formation and/or fibrosis. Questions include examining whether the different types of zig-zag patterns in the QRS complex generated before contraction, alter Ca\textsuperscript{2+} cycling events during or following contraction? The answer appears to be yes. The QRS complex represents the depolarisation wave as it spreads through the heart prior to contraction (figure 3A).\textsuperscript{16} In the healthy heart, the three QRS waves last around 70 ms to 100 ms and comprise a rapid phase 0 upstroke and phase 1 transient repolarisation or notch of the ventricular action potential (figure 3B). Phase 0 is mediated by rapid activation and deactivation of voltage-gated Na\textsuperscript{+} fast channels (1–2 ms), which opens a tiny window for Na\textsuperscript{+} entry and depolarises the membrane from about −85 mV to +50 mV\textsuperscript{16} (figure 3B). This is followed by a brief period of repolarisation secondary to activation of a transient outward K’ current (I\textsubscript{o}) and the Na\textsuperscript{+}/Ca\textsuperscript{2+} exchanger (outward Na’ movement), which repolarises the membrane from +50 mV to around +30 mV or lower (phase 1). This phase ends the QRS complex (figure 3AB).

In the failing heart, changes in the ventricular action potential include a reduction in the rate of depolarisation and peak depolarisation during phase 0, and a decrease in the repolarisation (or even complete loss of the notch) in phase 1.\textsuperscript{17,18} The loss of the phase 1 notch is believed due to downregulation of the I\textsubscript{o}, Cooper and colleagues further showed that the loss of the notch in ventricular myocytes reduces the magnitude of Ca\textsuperscript{2+} transient from the sarcoplasmic reticulum, alters the time course of the Ca\textsuperscript{2+} transient and decreases the synchrony of later Ca\textsuperscript{2+} spark production with loss of excitation–contraction coupling.\textsuperscript{17,18} Thus, changes to the QRS complex that occur before contraction can alter intracellular Ca\textsuperscript{2+} cycling during contraction. New therapies targeting the restoration of phase 1 repolarisation and/or other defects in phase 0 may improve cardiac function in patients with HF and reduce arrhythmias by reducing early and delayed Ca\textsuperscript{2+}–triggered after depolarisations.\textsuperscript{18} These drugs should protect against arrhythmias without negative inotropic effects. However, a key question remains: How are the notches and shuffling in the QRS complex related to these aberrant Ca\textsuperscript{2+} releases, reduced late Ca\textsuperscript{2+} spark rates and/or increased arrhythmogenesis in primary prevention HF patients? Advances in the molecular basis of fQRS and aberrant Ca\textsuperscript{2+} cycling in the myocardium may also apply to other arrhythmogenic cardiomyopathies.

What are the effects of fQRS on arrhythmogenicity?

Figure 3  Schematic of the normal ECG (A), ventricular action potential (B) and changes in intracellular Ca\textsuperscript{2+} in normal and failing heart (C) over the duration of a single heartbeat. Modified after Cooper et al.\textsuperscript{11} Vertical lines separate the precontraction QRS complex from later contraction and relaxation phases. The molecular events underlying the different forms of fQRS, myocardial fibrosis and arrhythmias are not known (see text). I\textsubscript{o}, cardiac transient outward potassium current; I\textsubscript{Kr}, rapid delayed rectifier channel current for action potential repolarisation; I\textsubscript{to}, another major outward current responsible for repolarisation; I\textsubscript{t}, inwardly rectifying potassium current that stabilises the resting membrane potential and is responsible for shaping the initial depolarisation and final repolarisation of the ventricular action potential.

CONCLUSION

Before fQRS can become a potential prognostic candidate for primary prevention ICD placement in patients with HF, wider and more accurate reporting is urgently required into the different forms of fQRS, myocardial fibrosis and arrhythmias are not known (see text). I\textsubscript{o}, cardiac transient outward potassium current; I\textsubscript{Kr}, rapid delayed rectifier channel current for action potential repolarisation; I\textsubscript{to}, another major outward current responsible for repolarisation; I\textsubscript{t}, inwardly rectifying potassium current that stabilises the resting membrane potential and is responsible for shaping the initial depolarisation and final repolarisation of the ventricular action potential.

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