Atrial fibrillation in heart failure
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Heart failure (HF) and atrial fibrillation (AF) were predicted to become epidemics of the 21st century, in part due to increased longevity and the success in reducing overall cardiovascular (CV) mortality. Both conditions are increasingly prevalent, with spiralling cost to healthcare services globally. The incidence of AF is also predicted to double over the next 20 years, with expectations of 120–215 000 new cases per year by 2030 in Europe alone. Rates of HF in a global AF registry were 33% in paroxysmal, 44% in persistent and 56% in permanent AF. Thus, the combination of these two conditions will have a dramatic impact on healthcare and require a refocusing of CV care.

The pathophysiology and risk factors for HF and AF are closely aligned, and affected patients are usually elderly with a significant burden of comorbidity. Atrial fibrillation is both a cause and consequence of HF, with complex interactions leading to impairment of systolic and diastolic function not present in sinus rhythm. Atrial fibrillation is associated with a three-fold increased risk of incident HF. Vice versa, the structural and neurohormonal changes in HF make the development and progression of AF much more likely, both in heart failure with reduced ejection fraction (HFrEF) and preserved ejection fraction (HFpEF). Regardless of which comes first, patients with concomitant HF and AF have significantly worse prognosis. Given the poor outcomes associated with HF and AF, finding effective therapies for these patients is of paramount importance but also challenging—treatments shown to be effective in one or other of these conditions alone have also been observed to have efficacy or safety concerns in patients with HF and AF combined. In this review, we summarize the mechanisms and inter-relationship of HF and AF, and provide a state-of-the-art synopsis on optimal management, considering how best to combine therapies and the evidence-base supporting their use.
Mechanisms and pathophysiology of atrial fibrillation in heart failure

Heart failure and AF share risk factors and common pathophysiologic processes (see Figure 1). Hypertension, smoking, obesity, diabetes, renal impairment, sleep apnoea, and coronary artery disease are all associated with an increased risk of developing both HF and AF. In HF, neurohormonal imbalance and activation of the renin–angiotensin–aldosterone system (RAAS) leads to maladaptive physiological changes including increased filling pressures and afterload. These can lead to increased left atrial stretch and fibrosis, contributing to the development of conduction abnormalities and facilitating the initiation and maintenance of AF. The renin–angiotensin–aldosterone system also directly contributes to proarrhythmic remodelling, with angiotensin II causing atrial fibrosis and anisotropic conduction. Patients with HF also demonstrate altered calcium handling and calcium overload, which can lead to afterdepolarizations and arrhythmia.

Atrial fibrillation can promote the development of HF by a number of established mechanisms. Loss of atrial systole in AF impairs LV filling and can decrease cardiac output by up to 25%, particularly in patients with diastolic dysfunction. Irregular and/or rapid ventricular conduction in AF can lead to LV dysfunction and in some patients, a tachycardia-induced cardiomyopathy. Restoration of sinus rhythm increases stroke volume and LV emptying even before contractility improves, explaining why some patients with HF gain rapid haemodynamic improvement with cardioversion.

Prevention of atrial fibrillation in heart failure (and heart failure in atrial fibrillation)

In the Framingham study, 41% of patients with AF and HF developed HF first, 38% developed AF first, and in the remaining 21% AF and HF occurred at the same time. While there are no therapies proven to prevent the risk of incident HF in patients with established AF, the treatment of modifiable CV risk factors (especially hypertension), effective rate control and the diagnosis and treatment of associated comorbidities (e.g. sleep apnoea) would seem to be sensible interventions.

What about preventing AF in patients with known HF? Meta-analysis of RCTs suggests that angiotensin converting enzyme inhibitors (ACEi) and angiotensin receptor blockers (ARBs) reduce the risk of incident AF, with RR of 0.79 (95% CI 0.62–1.00) and 0.78 (95% CI 0.66–0.92), respectively. Data from the Candesartan in Heart Failure-Assessment of Reduction in Mortality and morbidity program (CHARM) show that ARBs can decrease the risk of new-onset AF in patients with HFrEF and HFpEF. In the β-blocker vs. placebo trials in HFrEF with baseline sinus rhythm, allocation to β-blockers was associated with a significant reduction in the adjusted odds of incident AF (odds ratio 0.67; 95% CI 0.57–0.79). In a small analysis pointing towards the potential benefits of personalized therapy in patients with HF and AF, HF patients who were homozygotes for β1 adrenergic receptor 389 Arginine had a 74% reduction in new-onset AF (95% CI 43–88%) when treated with bucindolol vs. placebo.

Management of concomitant heart failure and reduced ejection fraction and atrial fibrillation

Currently, clinicians often manage patients with combined HFrEF and AF by focusing on particular therapeutic aspects that have an evidence-base in one or other of these conditions (see Figure 2). Researchers have started to investigate if treatment efficacy differs in patients with concomitant disease, but at present these data are limited. In this section, we summarize the evidence-base for common...
treatment modalities and suggest a simple clinical mnemonic for the initial management of newly diagnosed concomitant HF and AF. The CAN-TREAT HFrEF + AF algorithm (see Figure 3) distinguishes the management of these patients from those with sinus rhythm. The presence of haemodynamic instability should be treated with urgent cardioversion (C). Anticoagulation (A) should be instituted to prevent thromboembolism, and diuretic therapy to normalize (N) fluid balance and reduce symptoms of HF. Subsequent therapy should target (T) an initial heart rate < 110 b.p.m. and initiate RAAS antagonism (R), though with limited data on efficacy (see details below). Early (E) rhythm control in patients with symptoms refractory to rate control, and consideration of advanced (A) HF therapies should follow (e.g. cardiac resynchronization therapy), with aggressive treatment (T) of other concomitant CV disease, particularly ischaemia and hypertension.

**Anticoagulation**

Stroke is the most feared complication of AF, most commonly due to embolization of thrombus from the left atrial appendage. Thrombus formation in AF fulfils Virchow’s triad of a prothrombotic milieu, stagnant blood flow and endothelial dysfunction. Anticoagulation with vitamin K antagonists (VKA; e.g. warfarin) or non-VKA anticoagulants (NOACs) prevent ~2/3 of ischaemic strokes in AF patients. Reduced left ventricular ejection fraction (LVEF) is independently associated with stroke, and the combination of HFrEF with AF doubles the risk of stroke compared with AF alone.

Although no trials have investigated this specific population, indirect sub-group data from the NOAC RCTs suggest the effect of anticoagulation for AF is similar in patients with concomitant HF. With the combination of higher stroke risk and effective therapy, anticoagulation is essential in all patients with HF and AF that do not have an

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**Figure 2** Major priorities of management in patients with heart failure and reduced ejection fraction and those with atrial fibrillation. HFrEF, heart failure with reduced ejection fraction; AF, atrial fibrillation; CV, cardiovascular; HF, heart failure; LV, left ventricle; RV, right ventricle.

**Figure 3** CAN-TREAT HFrEF + AF Management of newly diagnosed concomitant heart failure with reduced ejection fraction and atrial fibrillation.

- **C**: Cardioversion if signs of haemodynamic compromise
- **A**: Anticoagulation unless absolute contraindication
- **N**: Normalise fluid balance diuretics to control signs and symptoms of failure
- **T**: Target initial heart rate < 110 bpm consider stricter control if persistent symptoms
- **R**: Renin-angiotensin-aldosterone system ACEI/ARB/mineralocorticoid receptor antagonists
- **E**: Early consideration of rhythm control amiodarone/cardioreversion and ablation
- **A**: Advanced heart failure therapies resynchronization/defibrillator/mechanical support
- **T**: Treatment of other CV disease control of ischaemia and hypertension

**Diagnosis/management of non-CV comorbidities, including diabetes, renal dysfunction, anaemia and airways disease**

**Patient-centred approach**

**Education and support**

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absolute contraindication, and the NOACs are particularly attractive due to lower rates of intracranial haemorrhage compared to VKA therapy.33

Guideline-recommended heart failure and reduced ejection fraction therapy

Achieving euvolaemia and the resolution of HF symptoms using loop and thiazide diuretics are an important first step in the management of all HF patients, regardless of heart rhythm. Activation of neurohormonal pathways and RAAS are well described in HF, and the majority of evidence-based therapies target these compensatory mechanisms.41 Angiotensin converting enzyme inhibitors have proven efficacy in HFrEF for significant reduction in mortality, sudden cardiac death, and HF hospitalization, but no trials have examined their benefit in concomitant AF. Angiotensin receptor blockers are recommended as alternatives to ACEi in cases of intolerance, and there are numerous RCTs supporting their use in HFrEF.41 In CHARM, randomization to candesartan significantly reduced CV death or HF hospitalization in HFrEF patients with concomitant AF (HR 0.83; 95% CI 0.69—0.99), similar to that observed in patients without AF at baseline (HR 0.84; 95% 0.77—0.92). In contrast, irbesartan did not reduce the composite outcome of hospitalization due to HF, stroke, myocardial infarction, or death from vascular causes in AF patients enrolled in the Atrial Fibrillation Clopidogrel Trial with Irbesartan for Prevention of Vascular Events A or W trials. For those with a history of HF, the HR was 0.90 comparing irbesartan to placebo (95% CI 0.75—1.08). It should be pointed out that both of these results were post hoc defined sub-group analyses.

β-Blockers are now a standardized part of treatment in HFrEF following numerous RCTs describing a substantial reduction in all-cause mortality, CV death and hospitalization compared with placebo. In these trials, between 8 and 23% of enrolled participants were in AF at baseline.34 Pooling individual patient data from 11 RCTs (with 96% of recruited participants ever enrolled in such trials), the adjusted HR for all-cause mortality for β-blockers vs. placebo was 0.73 (95% CI 0.67—0.80) in sinus rhythm. In patients with AF the HR was 0.97 (95% CI 0.83—1.14), with the interaction P-value for baseline rhythm highly significant at 0.002 with no heterogeneity (see Figure 4).14 The lack of benefit in AF was consistent across all sub-groups, including age categories, gender, NYHA class, and baseline heart rate. There were also no significant reductions in a range of secondary outcomes despite a sample size of over 3000 patients, including CV hospitalization and composite clinical outcomes.41 Again, these data are based on sub-group analysis and the patients with AF differed to those in sinus rhythm. However, pooling individual patient data allows more robust handling and statistical analysis, as well as sufficient power.45 Hence clinicians should not expect a prognostic benefit from β-blockers in HFrEF patients with concomitant AF; however, there was no apparent harm and patients may have other indications, such as symptom or heart rate control. We are currently exploring whether this variance in efficacy is due to heart rate, LVEF, or other fundamental differences in how AF patients respond to β-blockers.46

Mineralocorticoid receptor antagonists (MRAs), such as spironolactone and eplerenone, are recommended in all HFrEF patients with persisting symptoms (NYHA classes II—IV) after treatment with ACEi and β-blockers.41 Although the majority of data on MRA are positive, in a post hoc analysis of the Atrial Fibrillation and Congestive Heart Failure (AF-CHF) trial evaluating rate and rhythm-control strategies, spironolactone was associated with increased mortality (HR 1.4, 95% CI 1.1—1.8).47 Despite a propensity-matched statistical model, it is not possible to exclude residual confounding as an explanation for this unexpected finding (i.e. sicker patients with AF differed to those in sinus rhythm).48 Hence clinicians should not expect a prognostic benefit from β-blockers in HFrEF patients with concomitant AF; however, there was no apparent harm and patients may have other indications, such as symptom or heart rate control. We are currently exploring whether this variance in efficacy is due to heart rate, LVEF, or other fundamental differences in how AF patients respond to β-blockers.46

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Figure 4 β-Blockers in heart failure and reduced ejection fraction with sinus rhythm and atrial fibrillation. Kaplan–Meier survival curves for β-blocker vs. placebo in heart failure patients with (A) sinus rhythm and (B) atrial fibrillation. Data are unadjusted survival curves for all reported deaths. Hazard ratios are derived from an adjusted one-stage Cox regression model, stratified by study and censored at 1200 days (3.3 years). Reproduced from Kotecha et al.14

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patients receiving MRA). Baseline AF was not reported in the Randomized Aldactone Evaluation Study of spironolactone vs. placebo.48 In the Eplerenone in Mild Patients Hospitalization and Survival Study in Heart Failure trial, the reduction in CV death or HF hospitalization was similar for HFrEF patients with or without a history of AF (P for interaction 0.59).49

To summarize, there are scarce data on the efficacy of ACEi, ARBs, or MRA in HFrEF with concomitant AF to decrease morbidity or mortality; however, their use is still recommended to reduce adverse remodelling in HF. The totality of RCT data on β-blockers in HFrEF patients with AF have now been analysed, and suggest that β-blockers have a neutral effect on death and hospitalization in these patients.

Rate vs. rhythm control of atrial fibrillation

Although sub-group data suggest that sinus rhythm is associated with improved outcomes in patients with AF (including all-cause survival),50 clinical trials have failed to demonstrate superiority of either a rate or rhythm-control strategy. For example in the AF-CHF trial, there was no difference in CV death when comparing a rate vs. rhythm-control strategy in patients with HFrEF and NYHA classes II–IV (HR 1.06, 95% CI 0.86–1.30, P = 0.59), with similar findings for all-cause mortality and worsening HF.51 There are several reasons that rhythm control has failed to prove survival in clinical trials, including limited efficacy and adverse effects of available treatments, or delayed intervention such that the cumulative effects of AF are already unable to be reversed. Sinus rhythm can be difficult to achieve and maintain, particularly in patients with HF. In the rhythm control arm of AF-CHF, 21% crossed over to rate control, 82% were taking amiodarone, 27% were in AF at 4-year follow-up, and 58% had at least one episode of AF during the trial.51 On the other hand, in studies of catheter ablation of AF, restoration of sinus rhythm is associated with significant improvement in left ventricular function (11% increase in LVEF on average).52 While there are no clear differences in CV outcomes, patients with AF and HF who spend a higher proportion of time in sinus rhythm suffer less severe functional impairment (NYHA class III symptoms in 27 vs. 35%, P < 0.0001).53 Based on these and other data, current guidelines reserve rhythm-control therapy for those patients who experience AF-related symptoms or worsening HF despite adequate rate control.54

Specific rate-control therapies

The three available therapies for rate control of AF in the context of HFrEF are discussed below and summarized in Table 1.

β-Blockers

As previously discussed, β-blockers in HFrEF patients with AF do not appear to improve mortality or reduce hospital admissions.14 However, their use is widespread for control of heart rate in AF, both acutely and for long-term management. In the acute setting of HF with rapid AF, β-blockers are useful for rate-reduction and preferred to digoxin due to their effectiveness at high sympathetic tone.59 As β-blockers can initially be negatively inotropic, initiation of β-blockers requires a measured approach using incremental dosage to achieve a heart rate that balances the need for rate control with other haemodynamic parameters. For long-term control of heart rate, β-blockers are traditionally the first-line choice for clinicians.60

Digoxin

Cardiac glycosides, such as digoxin and digitoxin, have seen recent declines in use after the publication of the DIG trial which showed no mortality benefit from digoxin in HF patients with sinus rhythm.61,62 Importantly, patients randomized to digoxin suffered less hospitalizations. In observational studies and post hoc analysis of RCTs, there have been concerns about increased mortality with digoxin,63 but equally a number of studies have found no association.64–67 As clearly demonstrated in a systematic review of all digoxin vs. control studies, the main problem with non-randomized assessment is that clinicians are more likely to prescribe digoxin to the sickest patients with HF and/or AF, which results in bias that cannot be adjusted for, even with complex statistical modelling.55 Unfortunately, there are currently no direct RCT comparisons of digoxin use in patients with AF. Until further evidence becomes available, digoxin should be used cautiously in appropriate patients, with no expectation of any effect on mortality.55 In a crossover mechanistic RCT of 47 patients with HFrEF and AF, there were no differences in heart rate, blood pressure, walk distance, or LVEF comparing carvedilol and digoxin, although β-blockers did result in higher BNP levels (183 pg/mL with carvedilol vs. 79.5 with digoxin; P = 0.03). There was a small and marginally significant improvement in LVEF with combination β-blocker/ digoxin compared with placebo/digoxin after 4 months of treatment (30.6 ± 9.6% vs. 26.0 ± 12.4%; P = 0.048).56

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<th>Guidelines</th>
<th>Agent</th>
<th>Safety</th>
<th>Efficacy</th>
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<tbody>
<tr>
<td>Recommended</td>
<td>β-Blockers</td>
<td>Individual patient data sub-group meta-analysis of RCTs suggests no safety concerns.54</td>
<td>Individual patient data sub-group meta-analysis of RCTs shows no impact on mortality or hospitalization in concomitant HFrEF and AF.54</td>
</tr>
<tr>
<td>Recommended as</td>
<td>Digoxin</td>
<td>Systematic review suggests no increase in mortality in concomitant HF and AF; higher mortality in AF patients in observational studies is likely due to residual confounding.55</td>
<td>No RCTs vs. placebo in AF patients; combined therapy with β-blockers improves LVEF.56</td>
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<tr>
<td>Avoid/use caution</td>
<td>Non-dihydropyridine calcium channel blockers</td>
<td>Limited sub-group data in post-MI patients only; suggestive of increased death, re-infarction, and HF.57</td>
<td>None demonstrated.58</td>
</tr>
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Table 1 Rate control of atrial fibrillation in heart failure with reduced ejection fraction
Calcium channel blockers
Non-dihydropyridine calcium channel blockers (verapamil or diltiazem) are not recommended in patients with significantly impaired left ventricular function due to their negative inotropic effects, although specific data are limited. In the Multicenter Diltiazem Postinfarction Trial, patients were randomized to diltiazem or placebo 3–15 days after the onset of myocardial infarction. HF patients, including 490 with evidence of pulmonary congestion, had an increase in the composite of cardiac death or non-fatal re-infarction (HR 1.41, 95% CI 1.01–1.96). In subsequent analysis, diltiazem was found to increase late-onset HF in those with LVEF <40%. Verapamil did not improve outcomes after myocardial infarction in patients who developed HF in the Danish Verapamil Infarction Trials.

Heart rate targets for atrial fibrillation in the context of heart failure and reduced ejection fraction
Following the publication of the Rate Control Efficacy in Permanent Atrial Fibrillation (RACE II) trial, patients with AF can initially be treated to a more lenient heart rate regime (<110 beats/min resting heart rate). To summarize, 614 patients with permanent AF were randomized to a heart rate <80 b.p.m. at rest and <110 b.p.m. during moderate exercise or lenient control, with results showing a similar rate of composite clinical events in each arm. There were also no differences in functional outcomes, hospital admissions, or symptoms. Fifteen percent of the population had LVEF <40% but it remains unclear if these targets apply to AF patients with HFrEF. However, these findings are consistent with other results and would suggest that heart rate in AF is more a marker of disease than a therapeutic target and that a lenient heart rate may be acceptable if symptoms are controlled and tachycardia is avoided. It is worth noting that other guideline recommendations continue to advocate a resting heart rate <80 b.p.m. in patients with symptomatic AF and left ventricular dysfunction.

Specific rhythm-control strategies
Cardioversion
The first step in rhythm control is the restoration of sinus rhythm, which often requires cardioversion. Urgent cardioversion is recommended in any patient with significant haemodynamic impairment secondary to AF. Elective cardioversion is indicated in individuals with symptomatic persistent AF. Unfortunately, recurrence of AF after successful cardioversion is a frequent problem (~50% at 6 months), particularly in patients with HF. Over half of inpatients undergoing cardioversion for atrial arrhythmias have HF.

Antiarrhythmic drugs
Antiarrhythmic drugs (AAD) for the maintenance of sinus rhythm in patients with AF and HFrEF are limited to dofetilide or amiodarone; however, dofetilide is not approved in Europe. Both drugs have associated safety concerns (see Table 2). Other AAD are not recommended for general use in HFrEF. When antiarrhythmic medications are used to treat AF in patients with HF, every effort should be made to avoid toxicity.

Catheter ablation
Catheter ablation has been shown to significantly improve freedom from AF in patients who have failed AAD, and avoids their toxicity. Accordingly, the use of catheter ablation has increased in clinical practice. Observational data suggest that in patients with AF and HF, LVEF improves by 11.1% after ablation (95% CI 7.1–15.2). Higher recurrence rates of AF are seen after ablation in patients with HF, leading to a need for additional ablation procedures. The PABA-CHF pilot study compared a rate-control approach with atrioventricular (AV) node ablation and cardiac resynchronization therapy (CRT), vs. catheter ablation in 81 patients with drug-refractory AF and mild-to-moderate HF. In the ablation group, 71% maintained sinus rhythm without AAD. The patients randomized to ablation had better HF-related quality of life, better

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<tr>
<td>Recommended</td>
<td>Amiodarone</td>
<td>Mixed channel blockade</td>
<td>Risks of toxicity, including thyroid, hepatic, pulmonary, and neurological.</td>
<td>Superior efficacy for maintenance of sinus rhythm vs. placebo: odds ratio 0.15 (95% CI 0.10–0.22).</td>
</tr>
<tr>
<td>Dofetilide</td>
<td>III</td>
<td>Requires inpatient stay for loading. Risk of torsades 0.8–3.3%. Not approved in EU.</td>
<td>Lower risk of all-cause rehospitalization in patients with AF at baseline vs. placebo: relative risk 0.70 (95% CI 0.56–0.89).</td>
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<tr>
<td>Caution required</td>
<td>Dronedarone</td>
<td>Mixed channel blockade</td>
<td>Increased mortality in patients with HF and permanent AF.</td>
<td>Decreased risk of CV hospitalization or death in patients with AF and no recent HF decompensation vs. placebo: 0.76 (95% CI 0.69–0.84).</td>
</tr>
<tr>
<td>Sotalol</td>
<td>III</td>
<td>Concern for excess proarrhythmia in patients with acute myocardial infarction or LVEF &lt;40%; relative risk 1.65 (95% CI 1.15–2.36) for all-cause mortality.</td>
<td>Sotalol was inferior to amiodarone in patients with AF (28% had NYHA class III HF).</td>
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<tr>
<td>Contraindicated</td>
<td>Flecainide and Propafenone</td>
<td>I</td>
<td>Flecainide, encainide and morazaine increased mortality in patients with myocardial infarction. Propafenone can precipitate decompensated HF, particularly in CYP 2D6 slow-metabolizers.</td>
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*SWORD evaluated D-sotalol rather than D,L-sotalol.*

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6-min walk times, and greater improvement in LVEF. Similar findings were reported in the ARC-HF trial of 52 HFrEF patients randomized to ablation or rate control with β-blockers/digoxin. More recently, pre-publication results from the Ablation vs. Amiodarone for Treatment of Atrial Fibrillation in Patients with Congestive Heart Failure and an Implanted ICD/CRTD trial suggest promising results for catheter ablation in this patient group, including greater maintenance of sinus rhythm. While there is debate as to which ablative approach is most effective for restoring sinus rhythm in patients with persistent forms of AF, the Substrate and Trigger Ablation for Reduction of Atrial Fibrillation Trial Part II demonstrated that neither additional linear ablation nor ablation of complex fractionated electrograms improves efficacy. Larger, more definitive trials are underway to help clarify whether ablation leads to improved CV outcomes in patients with AF and HF.

Recognizing the limitations of percutaneous ablation in patients with more advanced forms of AF, there are emerging alternative approaches to catheter ablation, including both surgical ablation and hybrid ablation. In the ‘convergent’ procedure, a surgical transdiaphragmatic endoscopic approach is used to make non-contiguous epicardial ablation lesions. These lesions are later completed with catheter ablation to produce complete pulmonary vein isolation. In one study of 101 patients, in which 30% had symptomatic HF, freedom from AF was 66% at 1 year with a major complication rate of 6%. More traditional surgical ablation, like the Cox-Maze procedure, also has a role, particularly when HF patients with symptomatic AF are undergoing surgery for valvular disease or revascularization. Preliminary data in patients with concomitant HF suggests that Cox-Maze procedures may be effective and safe in those with LVEF <40% and symptomatic HF, with sinus rhythm maintained in >80% at 6-month follow-up.

**Device therapy and management of advanced heart failure**

Cardiac resynchronization therapy decreases mortality and prevents hospitalizations in patients with symptomatic HF, LVEF ≤35%, and QRS duration ≥120 ms. Between 25 and 50% of patients eligible for CRT have AF, although patients with AF are not well represented in randomized trials of CRT. At present, CRT is a class Ila recommended therapy in patients with HF and concomitant AF. Loss of AV synchrony and rapid ventricular rates in AF may impair benefit from CRT and small studies have suggested that the beneficial effect is rather due to AV node ablation. A meta-analysis of 23 observational studies including 7495 recipients suggests that patients with AF have a higher rate of non-response to CRT (35 vs. 28%, P = 0.001). Recent data from 8951 patients with AF and HF in the US NCDB registry demonstrated that CRT-D therapy was associated with lower mortality, all-cause, and HF readmissions compared with ICD therapy alone. Thus while response rates are lower in patients with AF, CRT should still be pursued in appropriate patients and every attempt should be made to ensure aggressive rate control. This is important to ensure biventricular pacing is as close to 100% as possible and avoid inappropriate shocks. AV node ablation should be considered in cases of tachycardia refractory to medical therapy.

Patients with HFrEF and AF often present unique challenges when implanted pump-support is required. Increased hospitalization and mortality have been observed in patients with the HeartMate II left ventricle assist device (adjusted HR for persistent AF 3.54; 95% CI 1.52–8.25; P < 0.01), with more frequent thromboembolic events in AF despite higher INR.

**Tachycardiomyopathy**

Tachycardia-induced cardiomyopathy is a long-recognized complication of AF, affecting as few as 3% and as many as 25% of patients with atrial tachyarrhythmias. Several mechanisms have been proposed to contribute to tachycardia-induced cardiomyopathy, including decreased density of L-type calcium channels and β-adrenergic receptors, increased intracellular calcium and diastolic contracture, impaired myocardial blood flow due to raised left ventricular diastolic pressure, oxidative stress, and even deleterious polymorphisms in angiotensin converting enzyme.

The diagnosis should be considered in a patient with no prior CV history who presents with new-onset HF in the setting of AF with rapid ventricular conduction. When evaluating patients with AF and left ventricular dysfunction, it is paramount to exclude other underlying causes of ventricular dysfunction, including ischaemia. Once ischaemia has been ruled out, other indicators of non-ischaemic cardiomyopathy should also assessed (e.g. left ventricular hypertrophy, alcohol/drug use, infiltrative disorders, etc.). It is important to emphasize that there are no established diagnostic criteria for tachycardia-induced cardiomyopathy and the diagnosis can be elusive in the majority of patients with established CV disease or when the initial presentation is missed.

Once anticoagulation has been initiated and the risk of thrombus has been addressed, sinus rhythm should be restored with cardioversion. Several methods can be used to maintain sinus rhythm; short-term amiodarone (3 months) is often helpful and allows for recovery before deploying a more durable treatment modality such as catheter ablation. Recovery of ventricular function confirms the diagnosis and may take up to 6 weeks. Patients with tachycardia-induced cardiomyopathy have similar outcomes following catheter ablation compared with patients without structural heart disease.

**Heart failure with preserved ejection fraction and atrial fibrillation**

Heart failure with preserved ejection fraction is common, responsible for over half of prevalent HF, yet no therapies have yet been shown to reduce mortality or morbidity. Atrial fibrillation and HFrEF are closely linked, with similar risks and mechanisms as discussed above. Current management is no different to that in sinus rhythm; diuretics to reduce signs and symptoms of fluid overload, and optimisation of hypertension and other comorbidities. Whether MRA have a specific role in improving exercise capacity and diastolic function by reducing fibrosis is currently under investigation. The risk of stroke in AF with HFrEF is similar to HfRHF, and therefore all suitable patients require anticoagulation.
Future directions

Given the limited treatment options for patients with HF and AF, there is a clear unmet need in this important patient population. Future investigation is particularly required for rate control, optimal methods of rhythm control, and prevention. There is also a need to stratify the use of various treatments to improve efficacy and safety, and a number of studies are addressing whether genetic profiling can help to personalize our therapeutic approach.

Despite advances in many areas of AF, it seems that the evidence to guide best practices for rate control is more uncertain than ever. Adequately powered prospective clinical trials are needed to clarify optimal rate-control targets and the best pharmacologic treatments to achieve rate control in patients with HF and AF. While ablation represents a promising alternative to AAD, clinicians require more information on the balance between effectiveness, complications, and potential prognostic benefits. Finally, given the poor outcomes observed in patients with HF and AF, perhaps the best treatment strategy is to prevent AF from occurring in the first place. Clinical trials of interventions targeted at left atrial substrate and preventing disease progression may have an important role in this regard. Table 3 presents some of the main studies currently recruiting that will assess patients with HF and AF. It is clear that the combination of these two common CV conditions will continue to challenge physicians both in CV and general medicine for many years to come.

Acknowledgements

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References


Table 3  Upcoming trials relating to heart failure and atrial fibrillation

<table>
<thead>
<tr>
<th>Trial</th>
<th>Objective</th>
<th>Status</th>
<th>Further information</th>
</tr>
</thead>
<tbody>
<tr>
<td>CASTLE-AF</td>
<td>Catheter ablation for AF and HFrEF</td>
<td>Funded, recruiting</td>
<td><a href="https://clinicaltrials.gov/ct2/show/NCT00643188">https://clinicaltrials.gov/ct2/show/NCT00643188</a></td>
</tr>
<tr>
<td>RAFT AF</td>
<td>Rate vs. rhythm control for AF and HFrEF</td>
<td>Funded, recruiting</td>
<td><a href="https://clinicaltrials.gov/ct2/show/NCT01420393">https://clinicaltrials.gov/ct2/show/NCT01420393</a></td>
</tr>
<tr>
<td>GENETIC AF</td>
<td>Genetically targeted AF therapy in HF</td>
<td>Funded, recruiting</td>
<td><a href="https://clinicaltrials.gov/ct2/show/NCT01970501">https://clinicaltrials.gov/ct2/show/NCT01970501</a></td>
</tr>
<tr>
<td>IMPRESS-AF</td>
<td>Spironolactone in AF with HFrEF</td>
<td>Funded, recruiting</td>
<td><a href="http://www.isrctn.com/ISRCTN10259346">http://www.isrctn.com/ISRCTN10259346</a></td>
</tr>
<tr>
<td>EAST</td>
<td>Early rhythm control for AF</td>
<td>Funded, recruiting</td>
<td><a href="https://clinicaltrials.gov/ct2/show/NCT01288352">https://clinicaltrials.gov/ct2/show/NCT01288352</a></td>
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<tr>
<td>CAVANA</td>
<td>Early rhythm control for AF</td>
<td>Funded, recruiting</td>
<td><a href="https://clinicaltrials.gov/ct2/show/NCT00911508">https://clinicaltrials.gov/ct2/show/NCT00911508</a></td>
</tr>
<tr>
<td>CATCH ME</td>
<td>AF genetics and tissue profiling</td>
<td>Funded, recruiting</td>
<td><a href="http://www.catch-me.info">http://www.catch-me.info</a></td>
</tr>
<tr>
<td>DIGIT-HF</td>
<td>Digitoxin vs. placebo in HFrEF</td>
<td>Funded, recruiting</td>
<td><a href="https://clinicaltrials.gov/ct2/show/NCT00783576">https://clinicaltrials.gov/ct2/show/NCT00783576</a></td>
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<tr>
<td>RATE-AF</td>
<td>Digoxin vs. β-blockers in AF</td>
<td>Funded, not started</td>
<td><a href="https://clinicaltrials.gov/ct2/show/NCT02391337">https://clinicaltrials.gov/ct2/show/NCT02391337</a></td>
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100. Yancy CW, Jessup M, Bozkurt B, Butler J, Casey DE Jr, Drazner MH, Fonarow GC, Heart failure & atrial fibrillation
