Paradigm shifts in pathophysiology and management of atrial fibrillation—a tale of the RACE trials in the Netherlands

H. J. G. M. Crijns · I. C. Van Gelder

Abstract In the past 20 years the Netherlands-based RACE trials have investigated important concepts in clinical atrial fibrillation (AF). Their scope ranged from rhythm versus rate control to early or delayed cardioversion and also included early comprehensive management of AF in two trials, one focusing on early ‘upstream therapy’ and risk factor management and the other on integrated chronic nurse-led care. Studies were mostly triggered by simple clinical observations including futility of electrical cardioversion in persistent AF; many patients with permanent AF tolerating day-after-day ‘uncontrolled’ resting heart rates of up till 110 beats/min; patients being threatened more by vascular risks than AF itself; and insufficient guideline-based treatments for AF. Also the observation that recent-onset atrial fibrillation generally converts spontaneously, obviating cardioversion, triggered one of the studies. The RACE trials shifted a number of paradigms and by that could change the AF guidelines. The initial ‘shock-and-forget’ attitude made place for increased attention for anticoagulation, and in turn, broader vascular risks were recognised. In a nutshell, the adage eventually became: ‘look beyond the ECG, treat the patient’.

Keywords Atrial fibrillation · Rate control · Rhythm control · Heart failure · Quality of life · Randomised clinical trial

Introduction

The first RACE study [1] originated from the Academic Hospital Groningen and spread rapidly across the Netherlands. All studies from the RACE consortium were strongly supported by the Working Group Cardiology Centres in the Netherlands (WCN), several non-WCN cardiology centres as well as the academic centres in Maastricht, both centres in Amsterdam, and Nijmegen. Per RACE study the participation of centres varied but the group remained cohesive. The Netherlands Heart Institute (the former Interuniversity Cardiology Institute in the Netherlands—ICIN) formed the scientific basis and a meeting place for all investigators and research nurses. The initial acronym RACE stands for 'RAte Control versus Electrical cardioversion for persistent atrial fibrillation'. We kept the acronym for the subsequent studies since the consortium partners and theme of study remained connected and constant through time. Most RACE trials were advised by a data safety monitoring committee of which Hein Wellens—now deceased—was a prominent member. Many medical scientists served on the respective endpoint event committees (Electronic Supplementary Material). A very important scientist for all RACE studies was Jan Tijssen, at present emeritus professor at the University of Amsterdam. His cardiovascular biostatistical insights and excellent knowledge of the field yielded robust biostatistics and by that a solid basis for high ranking publications of the results.

The RACE trials investigated and challenged important concepts in clinical atrial fibrillation (Tab. 1). In the early years of our careers, cardiologists looked...
upon atrial fibrillation (AF) as an arrhythmia for which rhythm control, especially antiarrhythmic drugs and electrical cardioversion (ECV), were most important ('shock-and-forget'). Subsequently, the focus shifted massively towards anticoagulation, fuelled by observations that patients had spontaneous or cardioversion-provoked cardioembolic strokes. The initial ‘big 6’ stroke trials [2] were—much later—followed by industry-academy alliances propelling the trials on non-vitamin K dependent anticoagulants. Thereafter, driven by observations from the EuroHeartSurvey on AF [3–5] as well as the two initial RACE studies [1, 6], we began to focus on comprehensive management of AF, i.e. treatments in addition to rate and rhythm control, and anticoagulation ('look beyond the ECG, treat the patient') [7]. For us a very important trigger was the observation that rates of mortality and heart failure were significantly higher [1, 4, 6] than the rate of stroke, and that patients suffer stroke despite anticoagulation. Then, comprehensive risk factor management—including rehabilitation and lifestyle—was studied in RACE3 [7, 8], in parallel with the study of integrated chronic care for AF (RACE4) [9]. RACE V, still ongoing, concerns a registry with the concept of hypercoagulation as a central mechanism for atrial remodelling and AF progression. RACE6-CV@H (cardioversion at home) is a proof-of-concept study into ECV performed at patients’ homes for their convenience and to test reallocation of hospital care. Conceived well before the 2020 COVID-19 pandemic, its concept perfectly fits the current hospital-at-home care strategies. Also in the early versus delayed cardioversion trial (RACE7-ACWAS) [10, 11] avoidable hospital procedures were the focus. As we became senior scientists, our successors Kevin Vernooy, Michiel Rienstra and Dominik Linz recently started two RACE studies which focus on ablation in heart failure (RACE8) and tele-checked rate control of recent-onset AF (RACE9). The present paper gives a perspective of all RACE studies and their contribution to the field.

The first RACE study: setting the stage, execution and aftermath

The first RACE trial, evaluating whether simple rate control was non-inferior to complex rhythm control, was stimulated by several clinical observations. First and foremost, long-term maintenance of sinus rhythm after ECV was disappointingly low, with 90% of patients having a recurrence of persistent AF after 4 years [12]. With that, cost-effectiveness as well as the effect on quality of life of cardioversion were questioned. In fact it was Henk Lie, former head of the Department of Cardiology in Groningen and our well-respected PhD and cardiology residency supervisor, who challenged us by asking the simple question ‘Cardioversion . . ., does it help?’ During the preparation of the RACE trial, friend and colleague scientist Maurits Allessie, Department of Physiology in Maastricht, cautioned us that the trial might come too early since the therapies to maintain sinus rhythm were still unsatisfactory. At that time, the AFFIRM trial [13] was about to start and therefore we were eager to go through with RACE. On the basis of Allessie’s advice we optimised the rhythm control treatment requirements for the investigators. ECV was performed in a serial manner with repeat cardioversion as needed and a change of prophylactic antiarrhythmic if the recurrence was early, i.e. within 6 months (Fig. 1; [1, 12, 14]). If, on the other hand, a recurrence happened beyond 6 months of sinus rhythm, the drug in use was left unchanged. We asked investigators to maximise efforts to keep patients in sinus rhythm by performing repeat cardioversion as quickly as possible after a recurrence and tailor antiarrhythmic drug treatment to the patient, avoiding both underdosing as well as side effects. During those years catheter ablation was not widely used for AF. Rate control was performed with a lenient target of 100 beats per minute (bpm). A non-electrical primary endpoint was chosen in RACE since it is not the eventual rhythm during follow-up that counts but rather longevity of the patient and avoiding severe adverse events including heart failure, stroke, bleed-

Dutch contribution to the field

- The revolutionary notion of ‘electrical remodelling’ inspired many investigators to develop new clinical concepts including ‘the second factor’ to complement electrical remodelling, AF progression as an endpoint in clinical trials, and early AF management and early comprehensive upstream therapy to improve prognosis.
- Several significant paradigm shifts in AF treatment happened: rhythm control was offset by rate control in persistent AF; aggressive by lenient rate control in permanent AF; and acute restoration of sinus rhythm by the wait-and-see approach in recent-onset AF. Lately the concept that electrical cardioversion should be considered a diagnostic rather than a therapeutic procedure emerged. The RACE studies also fed the notion that besides stroke, AF patients are even more threatened by heart failure and cardiovascular death.
- Cardiovascular risk scores to steer AF management were developed and swept the world, among which CHA2DS2-VASC, HAS-BLED and HATCH scores.
- The RACE consortium strongly advocated nurse-led integrated chronic care for atrial fibrillation and demonstrated its overall effectiveness; nurses steering integrated care perform better than stand-alone doctors.
| Concept/Hypothesis                                      | Result/Change                                                                 | References |
|--------------------------------------------------------|-------------------------------------------------------------------------------|------------|
| **RACE**                                               |                                                                               |            |
| Sinus rhythm better than AF                            | Rate control not inferior to rhythm control                                   | [1, 21]    |
| Mending the rhythm improves prognosis                  | No change                                                                     | [19]       |
| Rhythm control affects sudden death?                   | No impact                                                                      | [42]       |
| Sex differences may exist in rate and rhythm control   | Females suffer excess cardiovascular events under rhythm control              | [26]       |
| outcomes                                               |                                                                               |            |
| Rhythm control gives better QoL                        | No difference with RC                                                          | [25]       |
| Costs lower with RC                                    | Costs proven lower with RC                                                     | [24]       |
| RC may be deleterious in patients with CHF             | In patients with mild to moderate CHF, RC is not inferior to rhythm control   | [43]       |
| Clinical lone AF is not associated with cardiovascular  | Clinical lone AF is associated with bleeding and thromboembolism              | [44]       |
| events                                                 |                                                                               |            |
| Underlying comorbidities may affect outcome differen-  | In hypertensives, pharmacological rhythm control is associated with           | [45]       |
| tently between rate and rhythm control                 | cardiovascular morbidity/mortality; consider default RC                        |            |
| Anticoagulation should be bridged around surgery       | Extremely low perioperative thromboembolism risk; interruption of warfarin    | [22]       |
|                                                         | less dangerous than previously thought                                         |            |
| **RACE-II**                                            |                                                                               |            |
| Strict RC is standard of care (comparison of RC in     | Strict RC causes CV events, including excess artificial pacemaker implanta-    | [28]       |
| RACE (lenient) and AFFIRM (strict))                    | tions                                                                         |            |
| Strict rate control with resting heart rate in AF     | Lenient RC not inferior to strict RC                                           | [6, 46, 47]|
| recommended as <80 bpm                                 |                                                                               |            |
| Strict RC in AF and HF improves symptoms, CV prog-    | No beneficial effect of strict RC in permanent AF patients                    | [48]       |
| nosis and QoL                                          |                                                                               |            |
| Strict RC improves QoL                                  | Stringency of RC does not affect QoL; symptoms, sex, age, underlying          | [32]       |
| disease affect QoL                                      |                                                                               |            |
| Strict RC may fail, which predisposes to events        | Strict RC fails in 33% of patients but is not associated with events;          | [30]       |
|                                                         | lenient RC is preferred                                                       |            |
| Digoxin affects morbidity and mortality                | The use of digoxin was not associated with increased morbidity and            | [31]       |
|                                                         | mortality                                                                      |            |
| **RACE3**                                              |                                                                               |            |
| Targeted ‘upstream therapy’ for secondary AF preven-    | First study to show improved rhythm outcome with upstream therapy             | [7, 8, 49] |
| tion unproven                                          |                                                                               |            |
| Optimal upstream therapy may not be feasible in all    | Upstream therapy feasible in 57% of patients; it is associated with            | [37]       |
| patients                                               | enhanced rhythm outcome                                                        |            |
| QoL change uncertain                                   | Targeted therapy improves QoL, not necessarily through obtaining sinus         | [33, 38]   |
|                                                         | rhythm                                                                         |            |
| **RACE4**                                              |                                                                               |            |
| Doctors manage AF better than nurses                    | (Experienced) nurses manage better                                            | [9, 39, 41]|
| **RACE-V**                                             |                                                                               |            |
| AF progression is driven by hypercoagulation           | Expected change: anticoagulation prevents AF progressions, not only stroke    |            |
| Uncertain role for ILR                                 | Expected: ILR detects temporal types of AF                                    |            |
| **RACE6-CV@H**                                         |                                                                               |            |
| Electrical cardioversion must be done in-hospital      | Expected change: cardioversion can be safely performed at home                |            |
| **RACE7-AFCWS**                                        |                                                                               |            |
| Early cardioversion better than delayed cardioversion  | Delayed cardioversion not inferior to early cardioversion                     | [10, 11, 50]|
| for recent-onset AF                                    |                                                                               |            |
| **RACE8-HF**                                           |                                                                               |            |
| Cryoballoon PVI improves prognosis in persistent AF    | Expected change: uncertain, remains to be seen                                 |            |
| and heart failure                                      |                                                                               |            |
| **RACE9**                                              |                                                                               |            |
| Cardioversion (early or delayed) remains a key proce-   | Expected change: interventional rhythm control has no significant role in     |            |
| dure in recent-onset AF                                | stable recent-onset AF                                                        |            |
| Telemonitoring in management of recent-onset AF has-   | Expected change: telemonitoring prevents needless interventions and keeps     |            |
| —asyet—no place!                                       | patients safely out-of-hospital                                                |            |

**AF** atrial fibrillation, bpm beats per minute, **CHF** congestive heart failure, **ILR** implantable loop recorder, **LV** left ventricle, **PVI** pulmonary vein isolation, ablation therapy, **QoL** quality of life, **RC** rate control

Paradigm shifts in pathophysiology and management of atrial fibrillation
an association between sinus rhythm and survival [17]. In contrast, ‘mending the rhythm’ was not associated with event-free survival in RACE [18]. What critics did not understand was that staying in sinus rhythm is not only by virtue of the intervention but is heavily influenced by the underlying cardiovascular condition at the outset of the procedure, in particular the state of atrial remodelling. They also misunderstood the fact that RACE compared two strategies rather than the very acts of rate control (should yield appropriate rate) and rhythm control (should give permanent sinus rhythm). Like AFFIRM (and many later trials in this area, including CASTLE-AF) [19], RACE was a strategy evaluation and not an evaluation of the efficacy of cardioversion in maintaining sinus rhythm or of rate control to obtain an acceptable heart frequency. Also it was not an exercise in keeping the rate control arm patients away from cardioversion if that was deemed clinically indicated nor an exercise in obstructing channelling a rhythm control patient to rate control after repeated failure of cardioversion. In this respect, concepts like ‘per protocol analysis’ and ‘cross-over’ are pointless since failing rate or rhythm control may all be the outcome of an otherwise perfectly executed strategy. Post-hoc per-protocol analyses as performed in CABANA [20] are misleading and unfortunately they feed endless and perfectly fruitless debates. The bottom-line here is that sinus rhythm is frequently a marker of survival, and not the mechanism of survival!

Numerous sub-analyses were performed in the RACE population (Tab. 1). One of the main lessons learned from RACE is that anticoagulation must be continued if stroke risk factors are present even if patients maintain sinus rhythm [21]. RACE provided an excellent opportunity to check thromboembolism in almost 94 patients undergoing 121 non-cardiac surgeries. It was found that perioperative interruption of anticoagulation is far less dangerous than previously believed [22]. The latter findings were in line with the subsequent study by Douketis et al. [23] which led to widespread abandoning bridging of anticoagulation. A cost analysis indicated that costs were lower in the rate control arm compared with the rhythm control arm [24]. Over 2.3 years of follow-up, the mean costs per patient were €7386 under rate control and €8284 under rhythm control. Under rhythm control, more costs were generated due to electrical cardioversions, hospital admissions and antiarrhythmic medication. Quality of life appeared unaffected by strategy [25]. A sex-related sub-study showed that female patients with persistent AF had significantly higher cardiovascular morbidity and mortality under rhythm control compared with rate control [26]. Events were mainly heart failure, thromboembolism and adverse effects of antiarrhythmic drugs. Presumed mechanisms include higher prevalence of diastolic heart failure not amenable to rhythm control and heart failure associated with AF recurrence, lower rate of adequate anticoagulation in females, bradycardias and higher pacemaker implantation rates due to unmasking of
Fig. 2  Main results from 5 RACE trials.  

- **a** RACE: rate control is non-inferior to rhythm control [1];  
- **b** RACE-II: lenient rate control is non-inferior to strict rate control [6];  
- **c** RACE3: upstream therapy was associated with a modest improvement of rhythm control [8];  
- **d** RACE4: nurse-led care appeared not superior to usual-care provided by a cardiologist, although in proficient centres, nurse-led care was associated with significantly fewer events [9];  
- **e** RACE7-ACWAS: sinus rhythm at 4 weeks after delayed cardioversion and standard-of-care early cardioversion did not differ;  
- **f** RACE7-ACWAS: sinus rhythm during index visit, according to type of cardioversion – in 69% of patients in the delayed cardioversion strategy an intervention may be avoided. In contrast, an early intervention allows only for 16% spontaneous conversion with 80% obligatory cardioversions [10].

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chronotropic and dromotropic incompetence, mainly in females. Taken together, rhythm control should be applied judiciously in females, and when in doubt be avoided.

The birth of RACE-II

RACE-II was ushered in by a criticism on RACE indicating the rate control arm was too lenient allowing an upper limit for resting rate in AF of 100 beats per minute (bpm). The AF guidelines at that time recommended an upper limit of 80 bpm [27]. Subsequently we constructed a lenient rate control arm in RACE-II with a limit at 110 bpm. The latter was based on the observation that most of our patients did well using a target resting heart rate below 100–110 bpm. We also reasoned that an at-all-costs strict rate control would be associated with iatrogenic bradycardia and excess pacemaker implants. A post-hoc comparison of the rate control arms of RACE (lenient control, below 100 bpm) and AFFIRM (strict rate control, resting heart rate below 80 bpm) showed a significantly lower heart rate but a higher composite of cardiovascular death, hospitalisation and myocardial infarction in AFFIRM compared with RACE (34 vs 25%) and conspicuously more patients in need of pacemaker therapy (11 vs 1% over 3 years, \( p < 0.009 \)) [28]. A few years later, in a randomised controlled comparison, RACE-II showed that lenient rate control is non-inferior to strict rate control (Fig. 2; [6]), which led to a change in the AF guidelines [29]. RACE-II has remained unique: unfortunately no further randomised clinical trials have been performed in this area. Note that patients with severe heart failure were not included.

Sub-analyses of RACE-II are shown in Tab. 1. An interesting finding was that strict rate control fails in up to one third of patients, i.e. a resting heart rate below 80 bpm could not be achieved. In line with the main results of RACE-II, failure of strict rate control was not associated with excess events compared with successful strict or lenient rate control [30]. Another interesting finding is that in stable permanent AF digoxin may be used safely to control heart rate [31]. This is in contrast to several post-hoc analyses from large studies suggesting that digoxin increases mortality, but all of these studies on AF and digoxin are post-hoc and many suffer from extensive selection biases or bias by indication. Note that both in RACE and RACE-II there was no difference in quality of life between the intervention and control groups [25, 32]. In contrast, RACE3 (below) showed that quality of life improved under upstream therapy compared with control, an effect which was independent of whether patients maintained sinus rhythm during follow-up or not [33].

Upstream to RACE3

An unpublished comparison of RACE and RACE-II (Fig. 3) showed that over the 10 years between those two studies sustained rather than interrupted anticoagulation as well as management of high blood pressure and heart failure with renin-angiotensin system inhibition had improved prognosis: less stroke and bleeding, fewer admissions for myocardial infarction or heart failure, and fewer severe side effects of drugs. The scientific community started to see AF as a vascular disease rather than an arrhythmia. The latter held for the majority of patients while, in only relatively few patients, AF is an exclusively electrical disease in which the electrical abnormality precedes onset of AF. In vascular AF, however, the underlying heart disease precedes AF by many years, usually more than a decade or two, and that mostly concerns hypertension, atherosclerotic heart disease, obesity or heart failure. The term vascular AF refers not only to underlying vascular disease but also to the notion that vascular remodelling (in particular left atrial dilation and fibrosis) precedes the onset of AF by decades, setting the electrophysiological stage for AF. Cosio and Crijns, supported by a group of experts in the field, were among the first to recognise that in clinical practice vascular disease precedes AF in many instances (their Fig. 3; [34]). Once AF emerges, most patients already suffer from vascular remodelling. These notions were fed by remarkable findings from studies such as the LIFE trial [35] indicating that at similar blood pressure reduction in hypertensive patients also suffering from left ventricular hypertrophy, the angiotensin receptor blocker (ARB) losartan halved the incidence of AF compared with the beta-
blocker atenolol (primary prevention). Of even greater significance, those investigators showed that in patients with incident AF during the study, strokes were halved by losartan compared with atenolol, strongly suggesting that ARBs, as non-antithrombotic drugs, may help to prevent stroke through their vascular protective effects. In 2007, Savelieva and Camm introduced the notion of ‘upstream therapy’, indicating that through primary prevention, AF and its cardiovascular sequels can be effectively reduced. Typically, upstream therapy and risk factor management would target remodelling processes through reduction of inflammation, oxidative stress and extracellular matrix remodelling driven atrial fibrosis, using ARBs, mineralocorticoid receptor antagonists (MRAs) and statins [36]. In addition, cardiovascular risk factors become reduced. In RACE3 we reasoned that single-element upstream therapy would leave other remodelling pathways and other risk factors open [7]. Therefore, we hypothesised that a combination of different classes of upstream therapies would have synergistic effects on the atrial substrate and thereby decrease AF. In addition to anti-remodelling drugs and optimised risk factor management, we introduced a lifestyle intervention with cardiac rehabilitation since regular exercise may reduce AF. We tested our hypothesis in patients with early persistent AF and early heart failure (predominantly heart failure with preserved ejection fraction, HFpEF) since in advanced stages of these diseases AF would no longer be amenable to upstream therapy. Note that the primary aim was to reduce recurrent AF after cardioversion. RACE3 showed that upstream therapy significantly reduces risk factors (blood pressure, cholesterol) as well as recurrent AF (Fig. 2). Therefore, lifestyle intervention, ARBs, MRAs and statins all should be considered in persistent AF and stable heart failure. Obviously, this composite of upstream therapies yielded only a moderate rhythm control effect and cannot replace antiarrhythmic drug therapy and catheter ablation. It, however, fits perfectly into a comprehensive therapeutic strategy of treating not only AF but also underlying comorbidities and risk factors. It needs to be seen whether better maintenance of sinus rhythm using upstream therapy will result in improved survival. This holds especially since a post-hoc sub-analysis showed that only 57% of all patients in the interventional group reached their upstream targets (i.e. had optimal therapy) [37]. Another sub-analysis [33, 38] showed that quality of life improves significantly with the multi-faceted upstream therapy compared with control treatment. This was seen independent from rhythm outcome, meaning that other pathways than rhythm control seem active in maintaining quality of life under targeted upstream therapy (Tab. 1).

### The Hendriks study and RACE4

In 2012—under the supervision of Robert Tieleman—Jeroen Hendriks reported that guideline-based, ICT-supported, physician-supervised, nurse-driven care for AF was superior to usual-care provided by a cardiologist [39]. That then unique study was criticised for being mono-centre. In RACE4 we adopted the approach of integrated chronic care in a multi-centre trial. RACE4 showed that among patients recently referred for management of first-detected AF, nurse-led care did not significantly reduce cardiovascular death or hospitalisation compared with usual-care (Fig. 2). Remarkably, nurse-led care did not enhance patient knowledge on AF or quality of life. Nevertheless, a predefined exploratory analysis showed that centres with higher proficiency and experience in nurse-led care performed significantly better concerning cardiovascular endpoints than less experienced centres. On the basis of the similar event rates between both approaches, one may conclude that nurse-led care is a safe and effective way of providing care for patients with AF. Therefore, continued education and sharing of knowledge between centres are key to increasing the impact of nurse-led integrated care in AF clinics [40]. Patient numbers are growing and the average age of AF patients is increasing. Therefore, nurse-led integrated chronic care will become inevitable to enable cost-effective and widely accessible care for AF in the future [41].

### RACE7-ACWAS—toward cardiovert now or later?

Patients reporting to the emergency department (ED) frequently convert spontaneously under the eyes of
the attending physician. Also, in the outpatient setting, many patients report self-termination of AF for which they do not even bother to come to the hospital. Obviously, mainly patients with prominent symptoms present to the ED and cardioversion is performed almost automatically by a willing team eager to clear the department. However, the opportunity to observe spontaneous conversion is wasted, which in itself may reveal important information for chronic management. Acute conversion also distracts from what really matters: the need for anticoagulation and how to manage recurrent episodes. These two approaches were evaluated in the RACE7-ACWAS trial [11]. Fig. 2 shows the main result. Wait-and-see with delayed cardioversion as needed appeared non-inferior to early or acute cardioversion in terms of sinus rhythm at one month after the index visit to the ED. The study provided insight into the advantages and disadvantages of both approaches (Tab. 2). One important aspect is that patient and physician can take an informed shared decision on the management of preference in the ED. In addition, patients having experienced a spontaneous conversion will more likely stay at home in case of a new episode. All medical information on episodes emerging over time may feed into a decision for interventional therapy or not. An ongoing cost-effectiveness analysis will provide insight into reduction of costs of the wait-and-see strategy.

The RACE trials—a clinical perspective of cardioversion

Rhythm control by ECV is still seen as a therapeutic procedure although several RACE trials have shown that it is often therapeutically futile. Nevertheless, ECV may have an important application as a diagnostic procedure. Firstly, it may help to establish whether the arrhythmia causes symptoms, e.g. by assessing the symptom/rhythm correlation in the work-up towards an ablation. In persistent AF it is frequently difficult to establish whether eliminating AF (e.g. by ablation with its intrinsic risks) will reduce symptoms since patients’ complaints may be related to other mechanisms, conspicuously to concomitant HFpEF. HFpEF is very frequently associated with AF. Secondly, a diagnostic ECV may underpin the diagnosis of tachycardiomyopathy due to AF in patients suffering from AF and heart failure with a reduced left ventricular ejection fraction (HFrEF). If, after ECV, the left ventricular ejection fraction improves, tachycardiomyopathy with HFrEF is a most probable diagnosis. To prevent future recurrences of tachycardiomyopathy, ablation therapy may then be warranted or at least stringent rate control is needed if ablation fails or is not considered. Although this all sounds great, its wider application needs a change of attitude among attending physicians. Up till now, the greatest challenge is to bridge the disconnect between the world of arrhythmologists and heart failure cardiologists, with only very few heart failure specialists offering their HFrEF or HFpEF patients also suffering from AF a way out of their electrical heart failure. Vice versa, electrophysiologists should look beyond the ECG and provide heart failure and other risk mitigating treatments more attentively.

Conclusion

The Netherlands RACE trials were a concerted action of many centres in the Netherlands and challenged established clinical concepts. Starting from simple clinical observations with the perspective of improving care, robust clinical trials could be constructed. As they moved forward they helped to change guidelines and improve care for AF patients.

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Conflict of interest H.J.G.M. Crijns and I.C. Van Gelder declare that they have no competing interests.

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References

1. Van Gelder IC, Hagens VE, Bosker HA, et al. A comparison of rate control and rhythm control in patients with recurrent persistent atrial fibrillation. N Engl J Med. 2002;347:1834–40.
2. Hart RG, Pearce LA, Aguilar MI. Meta-analysis: antithrombotic therapy to prevent stroke in patients who have nonvalvular atrial fibrillation. Ann Intern Med. 2007;146:857–67.
3. Nieuwlaat R, Capucci A, Camm AJ, et al. Atrial fibrillation management: a prospective survey in ESC member countries: the Euro heart survey on atrial fibrillation. Eur Heart J. 2005;26:2422–34.
4. Nieuwlaat R, Prins MH, Le Heuzey JY, et al. Prognosis, disease progression, and treatment of atrial fibrillation patients during 1 year: follow-up of the Euro heart survey on atrial fibrillation. Eur Heart J. 2008;29:1181–9.
5. Nieuwlaat R, Olsson SB, Lip GY, et al. Guideline-adherent antithrombotic treatment is associated with improved outcomes compared with undertreatment in high-risk patients with atrial fibrillation. The Euro heart survey on atrial fibrillation. Am Heart J. 2007;153:1006–12.
6. Van Gelder IC, Groenveld HF, Crijns HJ, et al. Lenient versus strict rate control in patients with atrial fibrillation. N Engl J Med. 2010;362:1363–73.
7. Alings M, Smit MD, Moes ML, et al. Routine versus aggressive upstream rhythm control for prevention of early
atrial fibrillation in heart failure: background, aims and design of the RACE3 study. Neth Heart J. 2013;21:354–63.

8. Rienstra M, Hobbelt AH, Alings M, et al. Targeted therapy of underlying conditions improves sinus rhythm maintenance in patients with persistent atrial fibrillation: results of the RACE3 trial. Eur Heart J. 2018;39:2987–96.

9. Wijtvliet E, Tielemans RG, van Gelder IC, et al. Nurse-led vs. usual-care for atrial fibrillation. Eur Heart J. 2020;41:634–41.

10. Pluymaekers N, Dudink E, Luermans J, et al. Early or delayed cardioversion in recent-onset atrial fibrillation. N Engl J Med. 2019;380:1499–508.

11. Dudink E, Essers B, Holvoet W, et al. Acute cardioversion vs a wait-and-see approach for recent-onset symptomatic atrial fibrillation in the emergency department: rationale and design of the randomized ACWAS trial. Am Heart J. 2017;183:49–53.

12. Van Gelder IC, Crijns HJ, Tielemans RG, et al. Chronic atrial fibrillation. Success of serial cardiotherapy and safety of oral anticoagulation. Arch Intern Med. 1996;156:2585–92.

13. [This study has no authors listed] Atrial fibrillation follow-up investigation of rhythm management—The AFFIRM study design. The Planning and Steering Committees of the AFFIRM study for the NHLBI AFFIRM Investigators. Am J Cardiol. 1997;79:1198–202.

14. Crijns HJ, Van Gelder IC, Van Gilst WH, Hillege H, Gosselink AM, Lie KL. Serial antiarrhythmic drug treatment to maintain sinus rhythm after electrical cardioversion for chronic atrial fibrillation or atrial flutter. Am J Cardiol. 1991;68:335–41.

15. Wyse DG. Selection of endpoints in atrial fibrillation studies. J Cardiovasc Electrophysiol. 2002;13(1 Suppl):S47–S52.

16. Wyse DG, Waldo AL, DiMarco JP, et al. A comparison of rate control and rhythm control in patients with atrial fibrillation. N Engl J Med. 2002;347:1825–33.

17. Corley SD, Epstein AE, DiMarco JP, et al. Relationships between sinus rhythm, treatment, and survival in the Atrial Fibrillation Follow-Up Investigation of Rhythm Management (AFFIRM) study. Circulation. 2004;109:1509–13.

18. Rienstra M, Van Gelder IC, Hagens VE, Veezer NJ, Van Veldhuisen DJ, Crijns HJ. Mending the rhythm does not improve prognosis in patients with persistent atrial fibrillation: a subanalysis of the RACE3 study. Eur Heart J. 2006;27:357–64.

19. Marrouche NE, Brachmann J, Andresen D, et al. Catheter ablation for atrial fibrillation with heart failure. N Engl J Med. 2018;378:417–27.

20. Packer DL, Mark DB, Robb RA, et al. Effect of catheter ablation vs antiarrhythmic drug therapy on mortality, stroke, bleeding, and cardiac arrest among patients with atrial fibrillation: the CABANA randomized clinical trial. JAMA. 2019;321:1261–74.

21. Hagens VE, Van Gelder IC, Crijns HJ. The RACE study in perspective of randomized studies on management of persistent atrial fibrillation. Card Electrophysiol Rev. 2003;7:118–21.

22. Vink R, Rienstra M, van Dongen CJ, et al. Risk of thromboembolism and bleeding after general surgery in patients with atrial fibrillation. Am J Cardiol. 2005;96:822–4.

23. Douketis JD, Spyropoulos AC, Kaastra S, et al. Perioperative bridging anticoagulation in patients with atrial fibrillation. N Engl J Med. 2015;373:823–33.

24. Hagens VE, Vermeulen KM, Ten Vergert EM, et al. Rate control is more cost-effective than rhythm control for patients with persistent atrial fibrillation—results from the RACE Control versus Electrical cardioversion (RACE) study. Eur Heart J. 2004;25:1542–9.

25. Hagens VE, Ranchor AV, Van Sonderen E, et al. Effect of rate or rhythm control on quality of life in persistent atrial fibrillation. Results from the rate control versus electrical cardioversion (RACE) study. J Am Coll Cardiol. 2004;43:241–7.

26. Rienstra M, Van Veldhuisen DJ, Hagens VE, et al. Gender-related differences in rhythm control treatment in persistent atrial fibrillation: data of the rate control versus electrical cardioversion (RACE) study. J Am Coll Cardiol. 2005;46:1299–306.

27. Fuster V, Ryden LE, Asinger RW, et al. ACC/AHA/ESC guidelines for the management of patients with atrial fibrillation. A report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines and the European Society of Cardiology Committee for Practice Guidelines and Policy Conferences (Committee to develop guidelines for the management of patients with atrial fibrillation) developed in collaboration with the North American Society of Pacing and Electrophysiology. Eur Heart J. 2001;22:1852–923.

28. Van Gelder IC, Wyse DG, Chandler ML, et al. Does intensity of rate-control influence outcome in atrial fibrillation? An analysis of pooled data from the RACE and AFFIRM studies. Europace. 2006;8:935–42.

29. Camm AJ, Kirchhof P, Lip GY, et al. Guidelines for the management of atrial fibrillation: the Task Force for the Management of Atrial Fibrillation of the European Society of Cardiology (ESC). Europace. 2010;12:1360–420.

30. Groenveld HF, Tijssen JG, Crijns HJ, et al. Rate control efficacy in permanent atrial fibrillation: successful and failed strict rate control against a background of lenient rate control: data from RACE II (rate control efficacy in permanent atrial fibrillation). J Am Coll Cardiol. 2013;61(7):741–5.

31. Mulder BA, Van Veldhuisen DJ, Crijns HJ, et al. Digoxin in patients with permanent atrial fibrillation: data from the RACE II study. Heart Rhythm. 2014;11:1543–50.

32. Groenveld HF, Crijns HJ, Van den Berg MP, et al. The effect of rate control on quality of life in patients with permanent atrial fibrillation: data from the RACE II (rate control efficacy in permanent atrial fibrillation II) study. J Am Coll Cardiol. 2011;58:1795–803.

33. De With RR, Rienstra M, Smit MD, et al. Targeted therapy of underlying conditions improves quality of life in patients with persistent atrial fibrillation: results of the RACE3 study. Europace. 2019;21:563–71.

34. Cosio FG, Aliot E, Botto GL, et al. Delayed rhythm control of atrial fibrillation may be a cause of failure to prevent recurrences: reasons for change to active antiarrhythmic treatment at the time of the first detected episode. Europace. 2008;10:21–7.

35. Wachtell K, Lehto M, Gerdts E, et al. Angiotensin II receptor blockade reduces new-onset atrial fibrillation and subsequent stroke compared to atenolol: the Losartan intervention for end point reduction in hypertension (LIFE) study. J Am Coll Cardiol. 2005;45:712–9.

36. Savelieva I, Camm J. Is there any hope for angiotensin-converting enzyme inhibitors in atrial fibrillation? Am Heart J. 2007;154:403–6.

37. Nguyen BO, Rienstra M, Hobbelt AH, et al. Optimal treatment of underlying conditions improves rhythm control outcome in atrial fibrillation—Data from RACE 3. Am Heart J. 2020;https://doi.org/10.1016/j.ahj.2019.12.005.

38. Chen LY, Chung MK. Risk factor modification: another win for our fight against atrial fibrillation. Europace. 2019;21(4):527–8.

39. Hendriks JM, de Wit R, Crijns HJ, et al. Nurse-led care vs. usual care for patients with atrial fibrillation: results of
a randomized trial of integrated chronic care vs. routine clinical care in ambulatory patients with atrial fibrillation.
Eur Heart J. 2012;33(21):2692–9.

40. Bunting KV, Van Gelder IC, Kotecha D. STEEER-AF: a cluster-randomized education trial from the ESC: the STEEER-AF trial is designed by the European Society of Cardiology (ESC) to see if better education for healthcare professionals can improve how patients are treated and how AF is managed.
Eur Heart J. 2020;41:1952–4.

41. Crijns H, Wijtvliet EJP, Pluymaekers N, Van Gelder IC. Newly discovered atrial fibrillation: who(see) care(s)? Europace. 2020;22:677–8.

42. Hagens VE, Rienstra M, Van Veldhuisen DJ, Crijns HJ, Van Gelder IC. Determinants of sudden cardiac death in patients with persistent atrial fibrillation in the rate control versus electrical cardioversion (RACE) study. Am J Cardiol. 2006;98:929–32.

43. Hagens VE, Rienstra M, Van Veldhuisen DJ, Crijns HJ, Van Gelder IC, Investigators R. Enhanced cardiovascular morbidity and mortality during rhythm control treatment in persistent atrial fibrillation in hypertensives: data of the RACE study. Eur Heart J. 2007;28:741–51.

44. Van Gelder IC, Van Veldhuisen DJ, Crijns HJ, et al. RAte Control Efficacy in permanent atrial fibrillation: a comparison between lenient versus strict rate control in patients with and without heart failure. Background, aims, and design of RACE II. Am Heart J. 2006;152:420–6.

45. Groenveld HF, Crijns HJ, Tijssen JG, et al. Rate control in atrial fibrillation, insight into the RACE II study. Neth Heart J. 2013;21:199–204.

46. Van Gelder IC, Van Veldhuisen DJ, Crijns HJ, et al. Lenient vs. strict rate control in patients with atrial fibrillation and heart failure: a post-hoc analysis of the RACE II study. Eur J Heart Fail. 2013;15:1311–8.

47. Groenveld HF, Crijns HJ, Tijssen JG, et al. Rate control in atrial fibrillation, insight into the RACE II study. Neth Heart J. 2013;21:199–204.

48. Mulder BA, Van Veldhuisen DJ, Crijns HJ, et al. Lenient vs. strict rate control in patients with atrial fibrillation and heart failure: a post-hoc analysis of the RACE II study. Eur J Heart Fail. 2013;15:1311–8.

49. Rienstra M, Van Veldhuisen DJ, Crijns HJ, et al. RAte Control Efficacy in permanent atrial fibrillation: a comparison between lenient versus strict rate control in patients with and without heart failure. Background, aims, and design of RACE II. Am Heart J. 2006;152:420–6.

50. Healey JS, McIntyre WF. The RAte Control Efficacy in permanent atrial fibrillation in hypertensives: data of the RACE study. Eur Heart J. 2007;28:741–51.

51. Groenveld HF, Crijns HJ, Tijssen JG, et al. Rate control in atrial fibrillation, insight into the RACE II study. Neth Heart J. 2013;21:199–204.