Detection of atrial high-rate events by continuous Home Monitoring: clinical significance in the heart failure–cardiac resynchronization therapy population

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Received 18 March 2011; accepted after revision 5 August 2011; online publish-ahead-of-print 20 September 2011

Aims
Uncertainty exists over the importance of device-detected short-duration atrial arrhythmias. Continuous atrial diagnostics, through home monitoring (HM) technology (BIOTRONIK, Berlin, Germany), provides a unique opportunity to assess frequency and quantity of atrial fibrillation (AF) episodes defined as atrial high-rate events (AHRE).

Methods and results
Prospective data from 560 heart failure (HF) patients (age 67 ± 10 years, median ejection fraction 27%) patients with a cardiac resynchronization therapy (CRT) device capable of HM from two multi-centre studies were analysed. Atrial high-rate events burden was defined as the duration of mode switch in a 24-h period with atrial rates of > 180 beats for at least 1% or total of 14 min per day. The primary endpoint was incidence of a thromboembolic (TE) event. Secondary endpoints were cardiovascular death, hospitalization because of AF, or worsening HF. Over a median 370-day follow-up AHRE occurred in 40% of patients with 11 (2%) patients developing TE complications and mortality rate of 4.3% (24 deaths, 16 with cardiovascular aetiology). Compared with patients without detected AHRE, patients with detected AHRE > 3.8 h over a day were nine times more likely to develop TE complications (P = 0.006). The majority of patients (73%) did not show a temporal association with the detected atrial episode and their adverse event, with a mean interval of 46.7 ± 71.9 days (range 0–194) before the TE complication.

Conclusion
In a high-risk cohort of HF patients, device-detected atrial arrhythmias are associated with an increased incidence of TE events. A cut-off point of 3.8 h over 24 h was associated with significant increase in the event rate. Routine assessment of AHRE should be considered with other data when assessing stroke risk and considering anti-coagulation initiation and should also prompt the optimization of cardioprotective HF therapy in CRT patients.

Keywords
Heart failure • Atrial fibrillation • Home monitoring • Thromboembolism • Anticoagulation • Cardiac resynchronization therapy

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Introduction

The morbidity and mortality of atrial fibrillation (AF) have a major impact on health-care expenditure, largely as a consequence of thromboembolic (TE) complications.\(^1\)

A significant proportion of AF is asymptomatic but carries important prognostic implications, as emphasized in the AFFIRM study.\(^2\)

Patients with undetected, ‘unprotected’ paroxysms of AF remain at a continued risk of TE. Conventional detection methods, such as Holter monitoring, are ineffective at detecting asymptomatic paroxysmal atrial fibrillation (PAF).\(^3\) Although data monitoring of modern implantable pacemakers has enhanced diagnostic capability,\(^4\) recognition of relevant data generally remains unchecked until the scheduled device review with consequent delay in therapeutic intervention. Implantable devices with remote monitoring offer a solution to this problem. Continuous atrial diagnostic data provide a unique opportunity to assess the frequency and quantity of AF episodes defined as atrial high-rate events (AHRE). The home monitoring (HM) technology (BIOTRONIK, Berlin, Germany), provides both daily and instantaneous transmission on event detection. This does not require patient involvement, is validated, and is a safe and reliable automatic remote system.\(^5\)

However, interpretation of the comprehensive data provided by these devices increasingly falls outside of conventional guidelines. There are limited data available as to what level of atrial high-rate burden is clinically relevant and when thromboprophylaxis should be employed.

Therefore, we proceeded to evaluate the clinical impact of high atrial arrhythmic burden in a heart failure (HF) population receiving cardiac resynchronization therapy (CRT) with remote monitoring technology. Data were pooled from two international prospective multi-centre studies.

Methods

Patient population

Data were considered from all patients enrolled in the international prospective multi-centre observational Home CARE (clinicaltrials.gov code NCT00376116) and everesT trials (Evaluation of the new Biotronik Resynchronization+ICD System) (Appendix 1). All patients had HF and a CRT device capable of continuous heart rhythm monitoring via HM. Patients in sinus rhythm (including patients with a prior history of AF) with >70% HM transmissions in the follow-up period and >3 months of HM follow-up were included in the evaluation. All trials complied with the Declaration of Helsinki and were approved by the locally appointed ethics committee of each participating centre. Informed consent had been obtained from all the patients.

Device and Home Monitoring characteristics

Patients received CRT devices with or without a defibrillator (Stratos LV-T, Kronos LV-T, or Lumax HF-T from BIOTRONIK) with HM turned on. This wireless, mobile remote monitoring system continuously records details of relevant clinical events (e.g., rhythm disturbances and delivered therapies). However, as this was an observational study, no interventions were undertaken when high atrial arrhythmia burden was detected. Also for the purposes of study analysis we were unable to verify the detected atrial arrhythmias with the available intracardiac electrograms (IEGM Online HD).\(^6\)

Detection of atrial burden

We defined AHRE as the duration of mode-switch in a 24 h period. Mode-switching in Stratos and Kronos devices occurs when five of eight consecutive atrial beats are >180 bpm and continues until five of eight are below 180. In Lumax devices the criteria for onset are 36 out of 48 atrial cycles with a rate >180 bpm and termination occurs when 20 out of 24 atrial beats are at a slower rate than a programmed value. Prior studies using similar detection algorithms have shown >95% sensitivity and specificity for the detection of atrial tachycardia/atrial fibrillation (AT/AF) episodes and measurements of AT/AF burden.\(^6\) Patients were included in the analysis if the total detected AHRE burden in a 24 h period exceeded an accumulative total of 14 min. This 14 min limit is due to the definition of AF burden in the HM system in per cent per day. So 1% (=14.4 min) is the minimal amount detected per day.

Outcome events

The primary study endpoint was the incidence of (TE) event, including stroke, transient ischaemic attack (TIA), and peripheral arterial embolism (PAE). The secondary endpoints were cardiovascular death, hospitalization because of AF, or worsening of HF. Ischaemic stroke was defined as an acute onset neurological deficit due to focal cerebrovascular inhibition of flow, persisting for >24 h.

Data analysis

Baseline demographic characteristics were collated including individual CHADS2 score.\(^7\) The patients analysed in this study were divided into the following groups: (Figure 1):

- **Group 1A.** Sinus rhythm at enrollment, no prior history of AF but HM detected AHRE over the follow-up period.
- **Group 1B.** Sinus rhythm at enrollment and no prior history of AF with no detected AHRE.
- **Group 2A.** Prior history of AF with detected AHRE over the follow-up period.
- **Group 2B.** Prior history of AF with no detected AHRE over the follow-up period.

Patients were included in Group 1A or 2A, if they experienced ≥1 AHRE event during the follow-up period. A prior history of AF was defined as attacks of AF lasting from 2 min to ≤7 days (PAF) or AF >7 days but <1 year (persistent AF).\(^8\) A diagnosis of persistent AF required documentation by EKG or Holter monitoring. All patients were in SR at enrollment.

![Figure 1](https://example.com/figure1.png)

**Figure 1** Study flow chart. AHRE, atrial high rate events; SR, sinus rhythm.
Statistical analysis

Normally distributed continuous variables were calculated using mean and standard deviation. Categorical data are presented as number and percentage. Distributions were tested for normality using Shapiro–Wilk statistics. Non-normally distributed variables were represented as median with 25th–75th interquartile range (IQR).

Comparison of normally distributed continuous variables was performed using Student’s t-test for paired and unpaired data. Non-normally distributed variables were compared using Mann–Whitney Rank Sum tests and Kruskal–Wallis tests. Comparison of categorical data was performed using χ² and Fisher’s Exact tests where appropriate. A global P value was calculated when comparing all four groups. Statistical significance was established as P < 0.05.

Differences in event rates over time were analysed by the Kaplan–Meier analysis with Gehan–Breslow test for each outcome in the four study groups. Multiple comparisons for the four different groups were performed with the Holm–Sidak method. The Cox regression model was used to analyse the effect of daily AHRE on TE and other composite endpoints (TEE/cardiovascular death) and (AF and HF admissions/TE/cardiovascular death). Thereby, the categorization of no AF, daily AHRE, > median (3.8 h), and daily AHRE ≥ median was used. SPSS software (version 16 statistical package, SPSS Inc., Chicago, IL, USA) was used for all the statistical analyses.

Results

Study population

Between August 2004 to August 2008, 936 HF/CRT patients were enrolled from 77 European cardiac centres in the two trials. Of the 376 patients excluded from analysis, pre-enrollment AF status was missing in 87 (9%) patients, 75 (8%) had <70% HM data available for analysis, 56 (6%) had <90 days surveillance via HM, and 158 (17%) dropped out of the study. Analysis was undertaken on prospective data from 560 patients, over a median follow-up period of 370 days (IQR from 253 to 390 days). See Table 1 for study population demographics.

All four groups had similar baseline characteristic, although there were significantly more patients with ischaemic HF in Group 1A compared with the other three groups. Accordingly, the uptake of cardioprotective medication including beta-blockers,

### Table 1 Patient Characteristics for all 560 patients including subgroup demographics

| Variable baseline patient characteristics | Total study population (n=560) | AHRE Group 1A (n=126) | No AHRE Group 1B (n=256) | Prior history of AF pre entry Group 2A AHRE (n=97) | Prior history of AF pre entry Group 2B no AHRE (n=81) | Global P value |
|------------------------------------------|-------------------------------|-----------------------|---------------------------|---------------------------------|---------------------------------|----------------|
| Male                                     | 434 (77.4%)                   | 107 (84.9%)           | 189 (73.8%)               | 73 (75.3%)                      | 65 (80.2%)                      | 0.086          |
| Age (years)                              | 66 ± 10                       | 65 ± 10               | 67 ± 10                   | 68 ± 10                         | 69 ± 9                          | 0.07           |
| QRS (ms)                                 | 160 (140–178)                 | 160 (140–180)         | 160 (132–173)             | 160 (140–180)                   | 160 (140–173)                   | 0.14           |
| Non-isaemic                              | 258 (46.1%)                   | 48 (38.1%)            | 126 (49.2%)               | 46 (47.4%)                      | 38 (46.1%)                      | 0.23           |
| NYHA I                                   | 7 (1.25%)                     | 1 (0.8%)              | 4 (1.6%)                  | 0 (%)                           | 2 (2.5%)                        | 0.31           |
| NYHA II                                  | 108 (19.3%)                   | 24 (19.0%)            | 51 (19.9%)                | 17 (17.5%)                      | 16 (19.8%)                      | 0.81           |
| NYHA III                                 | 401 (71.6%)                   | 94 (74.6%)            | 178 (69.5%)               | 74 (76.3%)                      | 55 (67.9%)                      | 0.21           |
| NYHA IV                                  | 36 (6.43%)                    | 5 (4.0%)              | 19 (7.4%)                 | 6 (6.2%)                        | 6 (7.4%)                        | 0.71           |
| EF (%)                                   | 25 (20–30)                    | 27 (20–30)            | 25 (20–30)                | 28 (20–32)                      | 25 (20–28)                      | 0.14           |
| Diabetes                                 | 179 (32%)                     | 37 (29.4%)            | 82 (32.0%)                | 31 (32%)                        | 29 (35.8%)                      | 0.82           |
| Hypertension                             | 210 (37.1%)                   | 45 (35.7%)            | 95 (37.1%)                | 35 (36.1%)                      | 35 (43.2%)                      | 0.71           |
| Pre-implant Stroke                       | 1 (0.18%)                     | 1 (0.79%)             | 0 (0.00%)                 | 0 (0.00%)                       | 0 (0.00%)                       | 0.33           |
| CHADS2 Score 1                           | 202 (36.1%)                   | 49 (38.9%)            | 95 (37.1%)                | 34 (35.1%)                      | 24 (29.6%)                      | 0.47           |
| CHADS2 Score 2                           | 216 (38.6%)                   | 48 (38.1%)            | 95 (37.1%)                | 36 (37.1%)                      | 37 (45.7%)                      | 0.47           |
| CHADS2 Score 3                           | 128 (22.9%)                   | 29 (23.0%)            | 56 (21.9%)                | 24 (24.7%)                      | 19 (23.5%)                      | 0.31           |
| CHADS2 Score 4                           | 14 (2.5%)                     | 0 (0.0%)              | 10 (3.9%)                 | 3 (3.1%)                        | 1 (1.2%)                        | 0.07           |
| ACEI                                     | 354 (63.2%)                   | 89 (70.6%)            | 165 (64.5%)               | 55 (56.7%)                      | 45 (55.6%)                      | 0.07           |
| Beta-blockers                            | 353 (63%)                     | 91 (72.2%)            | 166 (64.8%)               | 50 (51.5%)                      | 46 (56.8%)                      | 0.008          |
| Amiodarone                               | 103 (18.4%)                   | 17 (13.5%)            | 36 (14.1%)                | 22 (22.7%)                      | 28 (34.6%)                      | 0.0001         |
| Other anti-arythmics                     | 110 (19.6%)                   | 18 (14.3%)            | 38 (14.8%)                | 24 (24.7%)                      | 30 (37.0%)                      | 0.0004         |
| Anti-platelets                           | 239 (42.7%)                   | 64 (50.8%)            | 119 (46.5%)               | 28 (28.9%)                      | 28 (34.6%)                      | 0.002          |
| Warfarin                                 | 67 (12.0%)                    | 9 (7.14%)             | 22 (8.59%)                | 19 (19.6%)                      | 17 (21.0%)                      | 0.0007         |
| CRT (%) bivent pacing                    | 98 (95–99)                    | 98 (95–99)            | 98 (95–99)                | 98 (95–99)                      | 99 (95–99)                      | 0.07           |
| HM performance %                         | 93 (87–97)                    | 94 (89–97)            | 93 (87–97)                | 92 (84–96)                      | 94 (86–97)                      | 0.31           |

Data are presented as the mean value ± SD or median value (25th–75th percentile) for continuous variables and number and percentage of patients for categorical data.

ACEI: angiotensin-converting enzyme inhibitor; ARB: angiotensin receptor blocker; AHRE: atrial high-rate events; BiV biventricular pacing; EF: ejection fraction; HM: Home Monitoring; NYHA, New York Heart Association; other anti-arythmics include Sotolol and Class 1 anti-arythmics.
angiotensin-converting enzyme inhibitor (ACEI)/angiotensin receptor blocker (ARBs) and anti-platelets was increased in this group. Anticoagulation use among patients with known prior AF (Group 2A 19.6% and Group 2B 21%) was significantly higher than in Group 1A (7%) or Group 1B (8%). The zero atrial burden noted in Group 2B may have been explained by the significantly increased usage of anti-arrhythmic medication, including amiodarone, observed in this group. Of further relevance to our CRT cohort, the median frequency of biventricular pacing over the total follow-up period was >98% (IQR 95–99) for all four groups.

Quantification of atrial tachyarrhythmic burden

At least, 40% of the study population had at least 1 day where they had atrial tachyarrhythmias totalling >14 min (1% of day). Furthermore, in the sinus rhythm group with no prior history of AF, the annual incidence of a device detected atrial episode was 33%. Despite the prior history of AF only 64% of patients in Group 2 demonstrated a monitored atrial event. However, the median daily AHRE was substantially greater in Group 2A (647 min IQR 84–700) compared with Group 1A (44 min IQR 11–179; P = 0.001).

Clinical events

See Table 2, for summary of clinical event rates in all four groups.

Arterial thromboembolic complications

Over a median follow-up period of 370 days TE complications occurred in 11 patients (2.0%). In particular, five patients suffered a PAE, four ischaemic strokes, and two TIA. There were no associated TE events in those patients with no prior history of AF and zero atrial burden over the follow-up period. However, two PAE events (18.1%) did occur in Group 2B. Consequently, if a patient developed new atrial episodes or had a prior history of AF, they were more likely to develop a TE complication compared with patients in Group 1B. Furthermore, there was no statistical difference in the cumulative probability of developing a TE complication between Group 1A and 2A (P = 0.124). Despite there being no detected atrial arrhythmia in Group 2B, the TE outcomes were similar to those patients experiencing AHRE (Group 2B vs. 1A P = 0.15 and Group 2B vs. Group 2A, P = 0.5) (Figure 2).

Mortality

There were 24 (4.3%) deaths in total, with 16 (67%) due to a cardiovascular aetiology. There was no difference in all-cause mortality between the four study groups.

Heart failure admissions

A total of 52 (9.3%) admissions for HF were observed in the study population. Patients with a prior history of AF, whether or not they experienced a period of atrial burden, were more likely to be admitted with decompensated HF [7.9% (Group 1A), 5.1% (Group 1B) vs. 16.5% in Group 2A and 16% in Group 2B; P = 0.001] (Figure 3).

The temporal association between atrial arrhythmias and clinical events

Arterial thromboembolic complications

The last detected episode of AHRE was 46.7 ± 71.9 days (range 0–194) before the TE event. Of those with detected atrial burden, who suffered a TE complication, three (27.3%) were in an atrial arrhythmia at the time of diagnosis. Interestingly, four patients (36.4%) who developed TE events had no prior atrial arrhythmia recorded, including two patients from Group 2B.

Heart failure admissions

As for TE complications, there was a similar period of delay between the admission for HF and the detected atrial event (mean 47.4 ± 88.6 days, range 0–339). Moreover, 63.5% (n = 33/52) of patients had no atrial trigger recorded prior to their admission. Only seven (13.4%) patients were in an AHRE at the time of admission.

An atrial burden cut-off

Based on published analytical strategy,9 a cut-off was defined before data analysis as the observed median value of daily AHRE burden. Among those with detected AHRE the median value was 16% per day or 3.8 h and the study population was divided into three subsets: zero AHRE, low AHRE burden if below the median and high AHRE burden if above.

After adjusting for TE risk factors, Cox multivariate analysis demonstrated that a maximum AHRE longer than the median 3.8 h over a 24 h period was independently associated with TE events compared with patients with zero AHRE (HR 9.4; CI 1.8–47.0 P = 0.006). However, when comparing patients with a high AHRE burden (≥3.8 h) to those with a low burden

Table 2 Adverse event rates for all four study groups

| Clinical event | Total events | Group 1A n = 126 | Group 1B n = 256 | Group 2A n = 97 | Group 2B n = 81 | Global P value |
|----------------|-------------|-----------------|-----------------|----------------|----------------|---------------|
| TE             | 11 (2.0%)   | 5 (4.0%)        | 0 (0%)          | 4 (4.1%)       | 2 (2.5%)       | 0.02          |
| All-cause mortality | 24 (4.3%)   | 7 (5.6%)        | 8 (3.1%)        | 4 (4.1%)       | 5 (6.2%)       | 0.56          |
| CVS mortality  | 16 (2.9%)   | 4 (3.2%)        | 4 (1.6%)        | 4 (4.1%)       | 4 (4.9%)       | 0.33          |
| HF admission   | 52 (9.3%)   | 10 (7.9%)       | 13 (5.1%)       | 16 (16.5%)     | 13 (16%)       | 0.001         |

Data are presented as number and percentage.
(<3.8 h), a composite endpoint of admissions for AF or HF, or TE event or cardiovascular death was highly predictive for an AHRE burden cut-off of 3.8 h (HR 3.9 CI 1.9–7.9, P < 0.0001). (Table 3).

### Discussion

Using data from two international multi-centre studies, this study has shown that the detection of atrial events by remote device monitoring is associated with TE complications in sinus rhythm patients with HF and CRT; moreover, they also have a similar TE event rate to those patients with a prior history of AF.

Compared with patients with no detected AHRE, patients with device-detected AHRE burden >3.8 h in 1 day were nine times more likely to develop TE complications and four times more likely to die from a cardiovascular cause.

In addition, those patients with 3.8 h or more of AHRE were at significantly higher risk of being admitted with AF or decompen-sated HF, to develop a TE complication or die from a cardiac cause when compared with patients with an AHRE burden <3.8 h. Interestingly, the majority of patients (73%) did not show a temporal association with the detected atrial episode and their adverse event, with a mean lag period of 46.7 ± 71.9 days before the TE complication.

### Atrial burden studies to date

In the few studies that have assessed the clinical impact of device-detected AHRE only three have been in patients with HF and CRT, although none assessed the impact on TE event rates. Two of the studies relied on routine device interrogation. In a retrospective analysis of their single-centre experience of 161 patients with CRT devices, Caldwell et al. did not demonstrate significant difference in outcomes between patients in sinus rhythm or newly detected PAF. In a subsequent study by Borleff et al. there was a significant difference in outcomes between the AHRE group and the sinus rhythm group, using a composite endpoint of all cardiac hospitalizations and (appropriate or inappropriate) shock therapy.

However, Santini et al. have recently demonstrated similar adverse outcome data to our study with the use of continuous device diagnostics in 1193 patients in sinus rhythm with CRT. They clearly showed a higher incidence of the composite endpoint of death and HF hospitalizations among patients with AT/AF during follow-up when compared with patients without AT/AF. However, no data were reported on the TE event rates.

The Mode Selection Trial (MOST) substudy showed that almost 50% of patients with bradycardia pacing devices and sinus node disease had at least one atrial high-rate event lasting a minimum of 5 min, and this was an independent predictor for the combined endpoint of death or non-fatal stroke. Subsequently, Glotzer et al. attempted to define a threshold for stroke risk in device-detected AF. They demonstrated that patients, with moderate stroke-risk factors, evidence of atrial arrhythmias >5.5 h/day in a 30-day period had a TE event rate of 2.4% per year. Although the hazard ratio of 2.2 suggested that this cut-off doubled the risk of TE event compared to the low-burden group, the confidence intervals were wide (0.96–5.05). Capucci et al. reported that patients with bradycardia pacing indications and atrial events >24 h were three times more likely, after adjustment for risk factors, to have TE complications than those who had atrial arrhythmias <24 h. Similarly, in the recent ASSERT Trial (Asymptomatic AF and Stroke...
Evaluation in Pacemaker Patients and AF Reduction Atrial Pacing Trial), Healey et al. were able to demonstrate, in hypertensive patients receiving a dual-chamber pacemaker with no prior history of PAF, more than a doubling of embolic risk with the presence of atrial arrhythmias.  

### Clinical relevance of device-detected atrial burden

Although it is accepted that the stroke risk in PAF, persistent and permanent AF is similar, uncertainty exists over the importance of short-duration atrial arrhythmias. While prothrombogenic electrical remodelling of the human atria may occur with AF durations >20 min, few would consider anti-coagulation at this level. Our study supports the findings in the MOST study suggesting that device-detected atrial events of short duration may be as relevant as having a clinical history of PAF. Certainly, our study shows that the incidence of TE complications was similar in the group with a pre-existing AF compared with the sinus rhythm group with newly detected atrial arrhythmias.

However, the difficulty encountered for predicting a clinically relevant AHRE burden threshold, is the low TE event rate in the atrial burden studies. In a total of five atrial burden studies (including our data) totalling 4651 followed up over a median period of 18 months, there were only 51 TE events (1.1% incidence). This low event rate may be a consequence of a short follow-up period or reflect under-reporting of embolic events. Interestingly, in the relatively longer ASSERT study (mean 3-year follow-up) the overall rate of the TE complications was 0.72% per year.

**The presence of device-detected AHRE: is it a surrogate marker of disease severity**

Although, a causal link between the onset of AHRE and TE complications can be inferred in (3 of 11) 27% of patients in our study, the majority did not demonstrate a direct temporal link with both TE complications and HF admissions. Our findings correlate with the recently reported 40-patient subgroup analysis of patients with TE complications from the TRENDS study. Daoud et al. were able to show that in their 40 patients, who demonstrated atrial episodes prior to a TE event, 29 (73%) did not have an atrial arrhythmia in the 30 days prior to a TE event; a similar incidence as shown in our study. They go on to postulate that there may be other mechanisms, such as vascular disease risk factors, other than cardioembolism, which may come into play. Others have suggested that patients with device-detected atrial arrhythmias have more ‘severe underlying heart disease’ and hence higher mortality, while it is also well known that the development of AF is a marker of increased mortality in patients with underlying heart disease.

**Heart failure admissions**

The significantly increased admissions for decompensated HF observed in Groups 1A and 2 compared with those with no prior history of AF who remained in sinus rhythm throughout the follow-up period (Group 1B) may be explained by Borleffs et al. observation that patients with newly detected AF post-CRT-D, had less reverse LV remodelling and no improvement in MR severity or LA size. Although our results do not demonstrate a causal relationship between the development of AF and worsening HF, our study highlights that in CRT HF patients the device detection of AHRE may, in fact, represent a marker for worsening HF or HF hospitalization. Furthermore, the excellent biventricular pacing rate, observed in all four study groups, would further reiterate that the presence of AHRE may represent a marker of increased disease state. This would counter the findings shown by Santini et al. who had linked the deterioration in HF with the impact of reduced biventricular pacing, due to an overriding atrial rate.

### The role of anti-coagulation for device-detected atrial arrhythmias

Although our study does not define a level of atrial burden at which anti-coagulation is required it does support prior studies

### Table 3 Results of hazard ratios comparing atrial high-rate events variables for each clinical outcome based on Cox regression model

| Clinical outcome | AHRE variable | Low AHRE vs. zero AHRE | High AHRE vs. zero AHRE | High AHRE vs. low AHRE |
|------------------|---------------|-------------------------|-------------------------|------------------------|
| TE               | HR 4.3 (CI 0.73–26.2) | P = 0.11 | HR 9.4 (CI 1.8–47.0) | P = 0.006 |
| TE + cardiovascular death | HR 2.1 (CI 0.72–6.0) | P = 0.17 | HR 4.0 (CI 1.5–10.1) | P = 0.004 |
| TE + AF + HF + cardiovascular death | HR 1.0 (CI 0.49–2.1) | P = 1.0 | HR 3.8 (CI 2.3–6.3) | P < 0.0001 |

Low AHRE corresponds to a burden of 14 min to <3.8 h in 24 h monitoring period; high AHRE corresponds to a burden >3.8 h in a 24 h monitoring period.

aAF admissions.
bHF admissions.

**Clinical relevance of device-detected atrial burden**

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in considering lower levels of atrial fibrillation to be clinically significant.9–15 Following an interesting study correlating stroke risk and device-detected AHRE, Botto et al.,16 recommended anti-coagulation if patients developed 5 min or more of AF and had a CHADS2 score of >2. They also suggested that if patients experienced 24 h or more of AF despite a CHADS2 score of <1, they also should be considered for therapy. We eagerly await the results of the ongoing interventional prospective multi-centre study of remote monitoring of AF, which should provide the strongest evidence for the role of anticoagulation.21 In the worldwide survey, Lazarus retrospectively analysed 3 million transmissions in over 11 000 patients with brady pacing, implantable cardioverter defibrillator (ICD) and CRT-D devices. He reported that events were detected 2–5 months earlier than allowed by standard care and was associated with expedited anti-coagulation in 30% of patients and modification of anti-arrhythmic therapy in 42%.22

Study limitations
The major limitations were that atrial events were not verified by evaluating the electrograms and that two different AF burden-detection methods were employed. Although, all atrial rates <180 beats per minute were excluded to minimize the false-positive detection of arrhythmias, patients were included if they had an accumulative total of (1%) or 14 min of data detected per day, which would allow shorter episodes (<5 min) to be included and increase the chance of oversensing. Previously published data suggest that excluding episodes <5 min reduces the chance of oversensing.23 Also, the inclusion requirement of >70% HM data transmission and the minimum of 90 days of follow-up over the total study period may have contributed to some patients having limited amount of data for analysis.

The low frequency of TE complications introduces room for statistical error, especially when assessing for predictors of TE events and obtaining an AHRE burden cut-off. Another limitation is that we are unable to truly verify whether patients in Group 1 did in fact have previous asymptomatic AF. Furthermore, our study was not designed to assess the clinical response to CRT and this could have had an impact on the incidence of atrial arrhythmia burden. However, in the larger randomized trials CRT has not been shown to decrease the incidence of AF.24,25

Conclusions
Remote monitoring of implantable cardiac devices is a feasible and effective means of arrhythmia surveillance in HF patients. We have demonstrated that atrial arrhythmias detected by implanted devices are associated with an increased incidence of TE events, with a cut-off point of 3.8 h being associated with a significant increase in event rate. Device-detected atrial episodes may also represent a marker of disease severity or risk and hence the routine assessment of AHRE data, in patients with HF, should be considered with other diagnostic information when assessing stroke risk and considering anti-coagulation initiation. Furthermore, it should also prompt the optimization of cardioprotective HF therapy and arrhythmia management in CRT patients.

Acknowledgements
The authors thank Bernd Brüsehaber for his expert assistance in statistical analysis.

Conflict of interest: N.S. has received grant support from Medtronic and St Jude Medical. A.B. and J.P. are employees of Biotronik. P.O. and O.V. have no conflicts of interest. S.K.G.M. has received speaker honoraria/grant support from Biotronik, Medtronic and St Jude Medical. W.R.B. is an advisor for Biotronik. V.P. has received speaker fees and fees for proctoring from Biotronik. S.S. receives grant support and is an advisor for Biotronik.

Funding
Home CARE and everesT trials were supported by BIOTRONIK SE & Co. KG (Woermannkehre 1, D-12359 Berlin, Germany), who also provided funding for the Open Access publication charges.

Appendix 1
everesT Study
Evaluation of the new BIOTRONIK Resynchronization + ICD System
Objective
The international multi-centre prospective study ‘everesT’ is designed to evaluate safety and efficacy of BIOTRONIK’s CRT-D device: Lumax HF-T 300 and 340, and BIOTRONIK’s bipolar LV lead: Corox OTW BP Steroid and Corox OTW-S BP Steroid. This study documents first clinical experience with the handling and function of these products during the intra- and post-operative course.

Patient selection
Inclusion criteria
• All the following criteria must be fulfilled to include the patient in this study:
  • Patient is willing and able to comply with the protocol and has provided written informed consent.
  • Indication for CRT
  • Indication for implantation of an ICD
  • Stable residence anticipated for 6 months after enrollment

Exclusion criteria
• The patient is not eligible to enter this study, if one of the below listed exclusion criteria is fulfilled:
  • Planned cardiac surgical procedures within 6 months after enrollment
  • Life expectancy ≤ 6 months
  • Pregnant and breast-feeding women
  • Age < 18 years or otherwise missing complete contractually capability
  • Participation in another clinical study
  • For part B only: patient inclusion stopped by sponsor
  • For part B only: Corox OTW-(S) BP not yet available on the market.
For part B only: LV lead implanted or any failed attempt to implant an LV lead prior to enrolment

Primary endpoints
(1) Safety and efficacy of Lumax 300/340 HF-T - Part A
(2) Safety and efficacy of Corox OTW(-S) BP - Part B

Secondary endpoints
(1) Observations and complications related to Lumax 300/340 HF-T (A)
(2) Rate of inappropriate ICD therapies (A)
(3) Shock impedance (A)
(4) VF detection time (A)
(5) Observations and complications related to Corox OTW(-S) BP (B)
(6) Lead parameters (B)
(7) Patient deaths (A/B)

Publication
Results of the study have not been published yet. But the data received and analysed during the everesT clinical study demonstrated the safety and efficacy of the Lumax HF-T CRT-D device and Corox OTW(-S) BP Steroid LV leads and supported the FDA approval of Corox BP lead.

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