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Published in:
American Heart Association. Journal. Cardiovascular and Cerebrovascular Disease

DOI:
10.1161/JAHA.113.000382

Publication date:
2014

Document Version
Publisher's PDF, also known as Version of record

Citation for published version (APA):
Bang, C. N., Gislason, G. H., Greve, A. M., Bang, C. A., Lilja, A., Torp-Pedersen, C., ... Wachtell, K. (2014). New Onset Atrial Fibrillation is Associated With Cardiovascular Events Leading to Death in a First Time Myocardial Infarction Population of 89 703 Patients With LongTerm FollowUp: A Nationwide Study. DOI: 10.1161/JAHA.113.000382
New-Onset Atrial Fibrillation is Associated With Cardiovascular Events Leading to Death in a First Time Myocardial Infarction Population of 89 703 Patients With Long-Term Follow-Up: A Nationwide Study

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Background—New-onset atrial fibrillation (AF) is reported to increase the risk of death in myocardial infarction (MI) patients. However, previous studies have reported conflicting results and no data exist to explain the underlying cause of higher death rates in these patients.

Methods and Results—All patients with first acute MI between 1997 and 2009 in Denmark, without prior AF, were identified from Danish nationwide administrative registers. The impact of new-onset AF on all-cause mortality, cardiovascular death, fatal/nonfatal stroke, fatal/nonfatal re-infarction and noncardiovascular death, were analyzed by multiple time-dependent Cox models and additionally in propensity score matched analysis. In 89 703 patients with an average follow-up of 5.0±3.5 years event rates were higher in patients developing AF (n=10 708) versus those staying in sinus-rhythm (n=78 992): all-cause mortality 173.9 versus 69.4 per 1000 person-years, cardiovascular death 137.2 versus 50.0 per 1000 person-years, fatal/nonfatal stroke 19.6/19.9 versus 6.2/5.6 per 1000 person-years, fatal/nonfatal re-infarction 29.0/60.7 versus 14.2/37.9 per 1000 person-years. In time-dependent multiple Cox analyses, new-onset AF remained predictive of increased all-cause mortality (HR: 1.9 [95% CI: 1.8 to 2.0]), cardiovascular death (HR: 2.1 [2.0 to 2.2]), fatal/nonfatal stroke (HR: 2.3 [2.1 to 2.6]/HR: 2.5 [2.2 to 2.7]), fatal/nonfatal re-infarction (HR: 1.7 [1.6 to 1.8]/HR: 1.8 [1.7 to 1.9]), and non-cardiovascular death (HR: 1.4 [1.3 to 1.5]) all P<0.001). Propensity-score matched analyses yielded nearly identical results (all P<0.001).

Conclusions—New-onset AF after first-time MI is associated with increased mortality, which is largely explained by more cardiovascular deaths. Focus on the prognostic impact of post-infarct AF is warranted.

Key Words: cardiovascular mortality • mortality • myocardial infarction • new-onset atrial fibrillation • re-myocardial infarction • stroke

New-onset atrial fibrillation (AF) is a common complication in post-myocardial infarction (MI) patients, which even in its paroxysmal stage independently predicts adverse outcome. This fact indicates that AF in post-MI patients is an ominous sign, which should be prevented and promptly treated if detected. However, the prediction of new-onset AF is difficult, because the mechanisms that promote the development of AF in the MI setting are complex and multifactorial, and our understanding of the pathophysiology remains incomplete. Similarly, and despite its frequent occurrence, very limited data exist on whether there are differences in competing causes of death in MI patients with AF versus those without AF, and very few have performed analysis with AF as a time-dependent variable on long-time follow-up. Moreover, only 1 trial has investigated the impact of AF on re-MI in post-MI patients.
Therefore, the aims of this study were first to examine whether new-onset AF was predictive of all-cause mortality in first-time MI patients and whether this could be explained by an increase in cardiovascular mortality; and second, to study the general causes of new-onset AF-related death, including risks of dying from stroke or re-infarction and noncardiovascular death during long-time follow-up in a large nationwide cohort of patients discharged after first-time MI in Denmark.

Methods

Since 1978, the Danish National Patient Registry has registered all hospital admissions as well as diagnoses obtained in outpatient clinics in Denmark. Each admission is registered with 1 primary diagnosis and 1 or more secondary diagnoses according to the International Classification of Diseases (ICD), until 1994 the ICD-8 and from 1994 the ICD-10. Since 1995, the Danish Registry of Medicinal Product Statistics (national prescription registry) has kept records of every drug prescription dispensed from pharmacies in Denmark. Each medication is classified by the international Anatomical Therapeutical Chemical system, and the registry also includes information about the date of dispensing, formulation, strength, and quantity dispensed. Information on patients' vital status (dead or alive) was obtained from the civil registration system through Statistics Denmark. In Denmark, every resident is provided with a permanent and unique civil registration number that enables linkage between these administrative registries.

Population

From the National Patient Registry we identified all patients, aged ≥30 years, hospitalized with acute MI (ICD-10 code I21 to I22) for the first time between 1997 and 2009. The diagnosis of acute MI and endpoints has been validated in the National Patient Registry, with a sensitivity of 91% and predictive value of 93%. A detailed description of patient selection has been published previously. All patients who were alive at discharge were included in the present study. The primary outcome was fatal and nonfatal stroke, fatal and nonfatal re-infarction or cardiovascular death, noncardiovascular death and finally all-cause mortality in respect to new-onset AF following first-time MI. Patients with a history of AF were excluded.

Atrial Fibrillation Ascertainment

Patients with atrial flutter were considered to have AF. From the National Patient Registry we identified all patients with new-onset AF and atrial flutter (ICD-10 code 427.93, 427.94, I48) between 1997 and 2009. The AF diagnosis in the registry was based on admission for AF or if AF was recorded during hospital admission for other disease. All AF episodes, whether permanent or paroxysmal, were verified on a 12-lead electrocardiogram. The diagnosis AF has previous been validated, with a specificity of 99%.

Drug Use

Since 1995, the Danish Registry of Medicinal Product Statistics has registered all prescriptions dispensed from Danish pharmacies. In Denmark the national health security system covers all inhabitants and partially reimburses drug expenses (with a maximum copayment for each individual of ≈US$ 500 a year). Therefore, pharmacies are required to register all prescriptions dispensed, which ensure complete registration, and the reimbursement results in minimal incentive for patients to obtain medication through other sources. Coding is done according to the Anatomic Therapeutical Chemical (ATC) system. The registry includes information about date of dispensing, strength and formulation, quantity dispensed, and the affiliation of the doctor who issues the prescription.

Comorbidity

Comorbidity was defined from diagnoses at discharge from index MI as specified in the Ontario acute MI mortality prediction rule. The comorbidity index was further enhanced by adding diagnoses from 1 year before the event, as done by Rasmussen et al. Diagnoses used in the comorbidity index are shown in Table 1.

Statistical Analysis

Descriptive data are reported as mean±standard deviation (SD) or frequencies expressed as percentages. The predictive value of new-onset AF was assessed using the Cox regression model with delayed entry (ie, counting the time from new-onset AF to censoring or event, provided that the analyzed end point had not occurred before new-onset AF) with adjustment for age, sex, calendar year, re-infarction, concomitant pharmacotherapy, and comorbidity. To deal with the time-to-event bias in the new-onset AF patients we made Landmark analysis with yearly landmarks including Cox multiple analysis on all the fatal end points. To analyze the impact of age on the outcome we made 2 additional multiple Cox models with yearly landmarks; first with fixed age adjustment, and second with increasing age adjustment. Proportional-hazard assumptions, linearity of continuous variables and lack of interactions were tested for each model. We calculated a propensity score for the likelihood of developing new-onset AF following first-time MI by a multiple Cox regression model using baseline covariates. Using the Greedy matching macro (http://mayoresearch.mayo.edu/mayo/
research/biostat/upload/gmatch.sas), we matched each new-onset AF case to 1 AF-free control from the risk set at the time of AF on the basis of the linear predictor from the propensity score Cox model and year of AF-onset. Matched pairs were then analyzed in a Cox regression model with delayed entry and with AF as the only covariate. Cumulative incidence of all-cause mortality as a function of incident new-onset AF from the Landmark analysis was plotted. The hazard ratios and confidence interval from each Landmark were plotted in a forest plot. Interaction was tested among the Landmarks to test whether there was a difference in outcome in subsequent years after discharge from MI. Finally, the cumulative incidence of AF was calculated accounting for the competing risk of death. A 2-tailed $P$ value $<0.05$ was regarded as statistically significant in all calculations. SAS statistical software package version 9.1 for UNIX servers (SAS Institute Inc) was used for statistical analysis.

**Ethics**

The Danish Data Protection Agency has approved this study (ref. 2007-58-0015, int. ref: GEH-2010-001), and data at the individual level were made available in a manner by which specific individuals could not be identified. Retrospective register studies do not require ethical approval in Denmark. The authors had full access to the data and take responsibility for its integrity. All authors have read and agree to the manuscript as written.

**Results**

In all, 89,703 post-MI patients with average follow-up of 5.0±3.5 years were identified. During the course of the study, 10,708 (11.9% or 23.9 per 1000 person-years) patients developed AF after discharge from the first-MI hospitalization and 78,992 (88.1%) remained in sinus rhythm (SR). The incidence of new-onset AF decreased each year after discharge from first time MI (Figure 1). A total of 29,695 (33.1%) patients received percutaneous coronary intervention. A detailed description of baseline characteristics of the study population and distributions are given in Table 1. Table 2 shows how the covariables affected the rate of AF.

**All-Cause Mortality After New-Onset Atrial Fibrillation**

All-cause mortality occurred in 5423 (50.6% or 173.9 per 1000 person-years) versus 26,885 (34.0% or 69.4 per 1000 person-years) among patients with and without new-onset AF, respectively (Figure 2). In a multiple Cox model, with new-onset AF inserted as a time-dependent variable, adjusted for age, gender, year, concomitant medical treatment, and comorbidity there was a increased all-cause mortality (HR: 1.89 [95% CI: 1.84 to 1.95], $P<0.001$) in patients with new-onset AF. The propensity-score matched analysis yielded nearly identical results (HR: 1.22 [95% CI: 1.17 to 1.27], $P<0.001$).

To address survival bias, Landmark analysis with yearly Landmarks was performed. This showed persistent results (Figures 3 and 4A). In addition, the Landmark analysis revealed an increasing risk of mortality during subsequent Landmarks ($P$ for interaction $<0.001$, Figure 4A). However, this association was no longer seen when adjusting each Landmark to increasing age (Figure 4B).

**Cardiovascular Mortality After New-Onset Atrial Fibrillation**

Cardiovascular mortality was registered in 4277 (50.6% or 137.2 per 1000 person-years) versus 26,885 (34.0% or 69.4 per 1000 person-years) among patients with and without new-onset AF, respectively (Figure 2). In multiple Cox models, with new-onset AF inserted as a time-dependent variable, adjusted for...
age, gender, year, concomitant medical treatment, and comorbidity there was a significant increased cardiovascular mortality (HR: 2.09 [95% CI: 2.01 to 2.16], \( P < 0.001 \)) in patients with new-onset AF. The propensity-score matched analysis yielded nearly identical results (HR: 1.24 [95% CI: 1.24 to 1.36], \( P < 0.001 \)).

## Fatal Stroke After New-Onset of Atrial Fibrillation

Fatal stroke was registered in 610 (5.7% or 19.6 per 1000 person-years) versus 2414 (3.1% or 6.2 per 1000 person-years) among patients with and without new-onset AF, respectively (Figure 2). Nonfatal stroke was registered in 606 (5.6% or 19.9 per 1000 person-years) versus 2172 (0.3% or 5.6 per 1000 person-years) among patients with and without new-onset AF, respectively (Figure 2). In multiple Cox model, with new-onset AF inserted as a time-dependent variable, adjusted for age, gender, year, concomitant medical treatment and comorbidity; there was a significant increased risk of both fatal and nonfatal stroke after new-onset AF (HR: 2.34 [95% CI: 2.12 to 2.57], \( P < 0.001 \)) and (HR: 2.47 [95% CI: 2.24 to 2.73], \( P < 0.001 \)) respectively. Propensity-score matched analysis on fatal-stroke yielded nearly identical results.

## Fatal Re-Infarction After New-Onset Atrial Fibrillation

Fatal re-infarction was registered in 905 (8.5% or 29.0 per 1000 person-years) versus 5502 (7.0% or 14.2 per 1000 person-years) among patients with and without new-onset AF, respectively (Figure 2). Nonfatal re-infarction was registered in 1521 (14.2% or 60.7 per 1000 person-years) versus 12 444 (15.8% or 37.9 per 1000 person-years) respectively (Figure 2). In multiple Cox model, with new-onset AF inserted as a time-dependent variable, adjusted for age, gender, year, concomitant medical treatment and comorbidity; there was a significant increased risk of both fatal and nonfatal stroke after new-onset AF (HR: 2.34 [95% CI: 2.12 to 2.57], \( P < 0.001 \)) and (HR: 2.47 [95% CI: 2.24 to 2.73], \( P < 0.001 \)) respectively. Propensity-score matched analysis on fatal-stroke yielded nearly identical results.

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### Table 2. Co-Variables Contribution to Atrial Fibrillation*

| Characteristic                  | Hazard Ratio | Confidence Interval   | P Value |
|--------------------------------|--------------|-----------------------|---------|
| Age                            | 1.041        | 1.039 to 1.043        | <0.001  |
| Male gender                    | 0.949        | 0.906 to 0.983        | <0.001  |
| Medical Condition              |              |                       |         |
| Cerebral vascular disease      | 0.930        | 0.846 to 1.022        | 0.196   |
| Pulmonary vascular disease     | 1.065        | 0.914 to 1.240        | 0.184   |
| Cancer                         | 0.975        | 0.836 to 1.137        | 0.855   |
| Cardiac arrhythmias            | 1.532        | 1.408 to 1.667        | <0.001  |
| Acute renal failure            | 1.291        | 1.054 to 1.582        | 0.006   |
| Chronic renal failure          | 1.282        | 1.027 to 1.512        | 0.002   |
| Diabetes with complications    | 0.971        | 0.873 to 1.080        | 0.759   |
| Pulmonary oedema               | 1.054        | 0.890 to 1.249        | 0.585   |
| Shock                          | 1.003        | 0.828 to 1.216        | 0.995   |
| Peptic ulcer                   | 1.102        | 0.949 to 1.280        | 0.199   |
| Prescribed Drugs               |              |                       |         |
| Beta blocker                   | 0.943        | 0.902 to 0.985        | 0.038   |
| Ace-inhibitor                  | 1.027        | 0.987 to 1.070        | 0.011   |
| Acetylsalicylic acid           | 0.925        | 0.888 to 0.964        | <0.001  |
| Clopidogrel                    | 0.889        | 0.844 to 0.936        | 0.001   |
| Antidiabetic drugs             | 1.082        | 1.015 to 1.154        | 0.003   |
| Loop diuretics                 | 1.470        | 1.408 to 1.535        | <0.001  |
| Digitalis                      | 2.867        | 0.851 to 0.935        | <0.001  |
| Statins                        | 0.892        | 0.851 to 0.935        | 0.002   |

*Based on results from propensity score from Cox multiple regression analysis.
among patients with and without new-onset AF, respectively (Figure 2). In multiple Cox models, with new-onset AF inserted as a time-dependent variable, adjusted for age, gender, year, concomitant medical treatment, and comorbidity there was a significant increase of both fatal and nonfatal re-infarction after new-onset AF (HR: 1.67 [95% CI: 1.55 to 1.80], \( P < 0.001 \) and HR: 1.79 [95% CI: 1.69 to 1.89], \( P < 0.001 \) respectively).

Non-Cardiovascular Mortality After New-Onset Atrial Fibrillation

Non-cardiovascular mortality was registered in 1146 (10.7% or 38.8 per 1000 person-years) versus 7502 (9.5% or 19.4 per 1000 person-years) among patients with and without new-onset AF, respectively (Figure 2). In multiple Cox models, with new-onset AF inserted as a time-dependent variable, adjusted for age, gender, year, concomitant medical treatment, and comorbidity there was an increased risk of noncardiovascular mortality (HR: 1.43 [95% CI: 1.34 to 1.53], \( P < 0.001 \)) in patients with new-onset AF. To address the cause of increased risk of noncardiovascular mortality, an interaction analysis with PCI was made. This showed a significant lower risk of noncardiovascular mortality in non-PCI treated patients with new-onset AF compared to PCI treated patients (HR: 1.42 [95% CI: 1.33 to 1.53]; (HR: 1.55 [95% CI: 1.30 to 1.86], \( P \) for interaction <0.001).

Discussion

This study is, to the best of our knowledge, the first study to examine the associations of new-onset AF with specific causes of death in a large contemporary post-MI population. Second,
this study adds information on the impact of new-onset AF on nonfatal cerebral stroke and nonfatal re-MI in a large cohort of first-time MI patients with long-term follow-up. The main results of the study were: (1) that new-onset AF is associated with an increased risk of all-cause mortality, fatal and nonfatal stroke, fatal and nonfatal re-infarction, cardiovascular mortality and non-cardiovascular mortality; and (2) rate of new-onset AF decreased each year after discharge from first-time MI, however the mortality increased with subsequent years.

New-Onset AF and Myocardial Infarction

In accordance with previous studies, incidence rate of patients admitted for AF was higher after MI discharge (23.9 per 1000 person-years) compared to the background population in Denmark of similar age (5 to 7 per 1000 person-years).11 Studies have suggested that increases in atrial pressure can occur in relation to MI either caused by atrial ischemia,5,6 increased LV filling pressure or LV pumping become affected.7,17 Moreover, AF in MI patients has been associated with a variety of clinical risk factors including increased age and history of hypertension and diabetes, kidney dysfunction, and, in particular, the presence of heart failure symptoms and left ventricular dysfunction.17–19 However, there is also evidence to suggest that AF in MI setting is seemingly independent of age, heart failure, and ventricular dysfunction.20

All-Cause Mortality After New-Onset AF

The observed rate of all-cause mortality among patients who developed new-onset AF in this study (173.9 per 1000 person-years) is higher than that reported in patients who developed new-onset AF in the Danish background population (2.84 to 3.54 per 1000 patient-years).21 Nevertheless, even after adjustment for several risk factors, new-onset AF was independently predictive of all-cause mortality in the present study. This is in accordance with the Global Utilization of Streptokinase and Tissue Plasminogen Activator for Occluded Coronary Arteries (GUSTO) I study22 that showed higher 1-year mortality in post-MI patients with AF compared to patients without AF. Increased long-term all-cause mortality in patients with AF in the setting of MI has also been verified in the Optimal Trial in Myocardial Infarction with the Angiotensin II Antagonist Losartan (OPTIMAAL),23 Trandolapril Cardiac Evaluation (TRACE)20 and VALsartan In Acute myocardial InFarcTion (VALIANT)24 studies.

The Cause of Death in MI Patients With and Without New-Onset AF

It has been demonstrated that AF can facilitate ventricular arrhythmias,25–27 which could explain the increased risk of all-cause mortality after new-onset AF in MI patients. However, to the best of our knowledge, no data exist on the cause of death in MI patients with and without new-onset AF. Pedersen et al28 reported increased risk of all-cause and cardiovascular mortality including sudden cardiac death in MI patients with AF. Unfortunately, they did not explore specific causes of death, did not exclude patients with previous AF and did not analyze AF as a time-dependent variable. Thus, the present study demonstrated for the first time that MI patients with new-onset AF have an increased risk of dying from stroke and re-infarction. Consistent with our data, the OPTIMAAL study23 showed an increased risk of stroke associated with AF in patients with MI and left ventricular dysfunction. Also the GUSTO I study22 showed an increased risk of stroke in patients with AF after MI. The observed increases in risks of fatal and nonfatal stroke are probably to a large extent explained by the well-established association of increased thromboembol in AF. This raises the question whether post-AMI patients with transient AF should be prescribed
anticoagulation similar to patients with primary AF and for how long?

Some studies have argued that AF is merely a risk marker of death, and not a causal mediator of MI. However, there is evidence to suggest AF is associated with impairment of endothelial function even in the absence of heart failure or hypertension which could explain the increased incidence of AF associated with ischemia-reperfusion injury after MI. Interestingly, inflammation has been demonstrated to be important not only in AF but also in MI pathophysiology. C-reactive protein (CRP) has been shown to reduce nitric oxide production in endothelial cells and increase endothelial expression of adhesion molecules. It also plays a crucial role in chemotaxis of monocytes and foam cell formation in atherosclerotic plaques and enhances vasoreactivity of unstable plaques. This could partly explain the associations among MI, new-onset AF, and re-MI, and support the idea that AF reflects an underlying disease in the heart.

Non-Cardiovascular Mortality After New-Onset AF

In our study we found that new-onset AF was associated with noncardiovascular mortality. To a large extent, this could be explained by the therapeutic challenge with the need of both vitamin K antagonists (VKAs) and 2 antiplatelet therapy to prevent stroke and further coronary events in the patients with new-onset AF, which we have shown to be associated with increased risk of fatal and nonfatal bleeding. This is supported by the interaction and lower risk of noncardiovascular mortality in non-PCI treated patients with new-onset AF in the present study. These results imply that new-onset AF in MI patients is a cardiovascular disease and the independent relation to adverse cardiovascular prognosis disproves that new-onset AF only reflects age, gender, an increased or general disease throughout the body such as cancer, chronic kidney, and/or lung diseases. In addition, you could imagine that AF makes physiologic stresses from other conditions harder to tolerate due to poor heart rate regulation and decrease in cardiac output in some patients, which would make new-onset AF patients more vulnerable to other diseases.

Increasing Mortality Despite Decreasing New-Onset AF After Discharge

This study revealed that there is a subsequently increasing risk of mortality in the later years after discharge from first-time MI, which was mainly due to increasing age. This is supported by earlier observations including addition of age in CHA2DS2-VASc score (Congestive heart failure, Hypertension, Age ≥75 years, Diabetes mellitus, Stroke/TIA, Vascular disease, Aged 65 to 74 years and female Sex [1 point for presence of each, except for 2 points for age ≥75 years and stroke/TIA]; score range from 0 to 6). The decreasing rate of new-onset AF after discharge could be caused by patients’ immediate atrial ischemia or backward failure from an affected LV pump, or simply that the sickest patient dies earlier, leaving increasingly healthy patients alive with lower risk of new-onset AF.

Strengths and Limitations

Our study suffers the limitations with any analysis of longitudinal registry data. However, the completeness of data, which enabled comprehensive follow-up for 10 years, adds strength to this study. The Danish National Patient Registry keeps records on discharge diagnosis from all hospitals in Denmark, and the present study thus avoids selection bias, which may be present in surveys that are based on selected centers. Furthermore, the Danish National Patient Registry as well as the national prescription registry has been shown to be accurate. The Danish healthcare system is government financed, which secures equal access to health care for all inhabitants free of charge, regardless of socioeconomic status. The present study based on registries did not include clinical data and some comorbid status, most importantly, hypertension, ejection fraction, MI severity, and heart failure, which are known to be associated with risk of developing AF, stroke, MI, and death. This, of course, was a limitation. In addition, we were not able to subdivide AF patients into groups according to the severity or persistence of the arrhythmia that would have allowed us to distinguish between patients with low risk and patients with even worse prognosis. We have addressed issues of potential bias and baseline imbalances between the groups, by performing 3 different sensitivity analyses; (1) a multiple model adjusting for potential confounders; (2) time-dependent variable analysis; and (3) a propensity score-matched analysis. New-onset AF, whether paroxysmal or persistent, was identified from the National Patient Registry. AF diagnosis is in this registry based on admission for AF and AF recorded in outpatient clinics or during hospital admission for other reasons. This method leaves episodes of paroxysmal AF without need of hospital admission undetected. Thus, our method may have underestimated AF occurrence and primarily detected symptomatic AF.

Conclusions

This study shows that MI patients with new-onset AF have an increased risk of all-cause mortality, which to a large extent is caused by cardiovascular mortality. This implies that new-onset AF after MI should be seen as an “adverse prognostic sign” which should remind the physician to monitor and treat underlying disease; in particular cardiovascular, as soon as possible after new-onset AF has occurred.
Sources of Funding

This work was supported by The Danish Heart Association. Grant number: [10-04-R78-A2962-22582], the A.P. Moller Foundation and Interreg Iva—European Regional Fund; and had no relationships with the industry. Dr Gislason is supported by an unrestricted research scholarship from the Novo Nordisk Foundation.

Disclosures

Drs Devereux and Wachtell received grant support from Merck & Co, Inc. Dr Torp-Pedersen is a consultant to Sanofi-Aventis, Merck, and Cardiome, and receives consulting fees from all 3 companies. Dr Devereux consults for Merck & Co, Inc and General Electric Medical Systems. The remaining authors report no conflicts.

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*J Am Heart Assoc.* 2014;3:e000382; originally published January 21, 2014;
doi: 10.1161/JAHA.113.000382

The *Journal of the American Heart Association* is published by the American Heart Association, 7272 Greenville Avenue, Dallas, TX 75231
Online ISSN: 2047-9980

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