Catheter-based renal denervation for treatment of patients with treatment-resistant hypertension: 36 month results from the SYMPLICITY HTN-2 randomized clinical trial

Murray D. Esler1*, Michael Böhm2, Horst Sievert3, Christian L. Rump4, Roland E. Schmieder5, Henry Krum6, Felix Mahfoud2, and Markus P. Schlaich1

1Baker IDI Heart and Diabetes Institute, Monash University, 75 Commercial Road, Melbourne, Victoria 3004, Australia; 2Universitätsklinikum des Saarlandes, Homburg/Saar, Germany; 3CardioVaskulares Zentrum, Frankfurt, Germany; 4Department of Nephrology, Heinrich-Heine-University, Medical Faculty, Düsseldorf, Germany; 5Department of Nephrology, Hypertension of the University Hospital, Clinical Research Competence Center of Hypertension and Vascular Medicine, Erlangen, Germany; and 6Monash Centre of Cardiovascular Research and Education in Therapeutics, School of Public Health and Preventive Medicine, Monash University, Melbourne, Australia

Received 3 March 2014; revised 28 April 2014; accepted 30 April 2014; online publish-ahead-of-print 4 June 2014

Aim

The aim of this study was to determine long-term results of renal artery denervation for treatment of treatment-resistant hypertension in the SYMPLICITY HTN-2 study.

Methods

SYMPLICITY HTN-2 randomized 106 subjects with treatment-resistant hypertension to renal denervation or medical therapy alone. At 6 months, 37 control subjects crossed over to renal denervation. Office blood pressure measurements, antihypertensive medication use, and safety events were followed every 6 months through 3 years.

Results

Follow-up was available at 36 months in 40 of 52 subjects in the initial renal denervation group and at 30 months in 30 of 37 subjects who crossed over and received renal denervation at 6 months. Baseline blood pressure was 184 ± 19/99 ± 16 mmHg in all treated subjects. At 30-month post-procedure, systolic blood pressure decreased 34 mmHg (95% CI: −40, −27, P < 0.01) and diastolic blood pressure decreased 13 mmHg (95% CI: −16, −10, P < 0.01). The systolic and diastolic blood pressure reduction at 36 months for the initial renal denervation group was −33 mmHg (95% CI: −40, −25, P < 0.01) and −14 mmHg (95% CI: −17, −10, P < 0.01), respectively. Procedural complications included one haematoma, and one renal artery dissection before energy delivery that was treated successfully. Later complications included two cases of acute renal failure, which fully resolved, 15 hypertensive events requiring hospitalization, and three deaths.

Conclusion

Renal denervation resulted in sustained lowering of blood pressure at 3 years in a selected population of subjects with severe, treatment-resistant hypertension without serious safety concerns.

Clinical trial registration

NCT00888433.

Keywords

Resistant hypertension • Renal denervation • Symplicity

Introduction

The prevalence of hypertension is increasing worldwide at alarming rates.1–3 Furthermore, the prevalence of uncontrolled hypertension is also increasing, in spite of advances in pharmacotherapy.4,5 Patients with treatment-resistant hypertension (blood pressure ≥140/90 mmHg despite ≥3 antihypertensive medications, including a diuretic)5,6 often have other risk factors for cardiovascular disease and consequently are at even greater risk for end-organ damage and cardiovascular morbidities.5 When confounding causes, such as white-coat hypertension, non-adherence to drug therapy, and inappropriate drug selection or dosing, are eliminated, the estimated proportion of patients with treatment-resistant hypertension ranges from 5 to 16%.7–11

* Corresponding author. Tel: +61 385321393, Fax: +61 385321100, Email: murray.esler@bakeridi.edu.au

Published on behalf of the European Society of Cardiology. All rights reserved. © The Author 2014. For permissions please email: journals.permissions@oup.com.
Renal sympathetic outflow is commonly overactive in patients with essential hypertension; this has been demonstrated by measurements of norepinephrine spillover from renal sympathetic nerves. Pre-clinical work revealed that surgical sectioning of the renal nerves in experimental animal models of hypertension lowers blood pressure or can prevent development of hypertension.\(^2\)\(^3\)

Percutaneous catheter-based renal denervation (RDN) using the Symplicity\(^TM\) Renal Denervation System (Medtronic, Inc., Santa Rosa, CA, USA) involves the application of low-power (\(\approx 8\) W) radiofrequency energy to the main renal arteries in a helical pattern of ablations (four to six per artery). The renal artery nerves passing to and from the kidneys transit in the adventitia of the renal artery or just beyond and are therefore within reach of energy delivery. The Symplicity generator uses a proprietary algorithm to monitor and control temperature, impedance, and power output to assure safe delivery of energy to each site. The SYMPLICITY HTN-1 proof-of-principle trial demonstrated the feasibility of this procedure and showed that subjects with severe, treatment-resistant hypertension experienced a significant reduction in blood pressure that was sustained over at least 3 years (\(-32/-14\) mmHg at 36 months, \(P < 0.01\)).\(^18\)\(^-20\)

The SYMPLICITY HTN-2 randomized clinical trial compared the safety and effectiveness of RDN plus medical management to medical management alone (control group) in subjects with severe treatment-resistant hypertension.\(^21\)\(^22\) At 6-month post-randomization, there was a significant decrease from baseline blood pressure for the initial RDN group (\(n = 49; -32/-12\) mmHg, \(P < 0.0001\)) and no change in blood pressure from baseline in the control group (\(n = 51\)).\(^21\) No serious device or procedure-related adverse events were reported at the 6-month primary endpoint.

The SYMPLICITY HTN-2 trial allowed control subjects to crossover to receive RDN after the initial 6-month primary endpoint evaluation. We now report 36-month follow-up of the initially denervated subjects and 30-month follow-up of the control subjects who crossed over to RDN at 6 months.

**Methods**

**Study population and protocol**

SYMPLICITY HTN-2 enrolled subjects from June 2009 to January 2010 at 24 clinical sites in Europe, Australia, and New Zealand. The trial was approved by the local Ethics Committees in accordance with the Declaration of Helsinki and all subjects provided written informed consent.

Details of the trial methods have been previously reported.\(^21\) In summary, hypertensive adult subjects with a systolic blood pressure (SBP) \(\geq 160\) mmHg (\(\geq 150\) mmHg if they had type 2 diabetes mellitus) while receiving \(\geq 3\) antihypertensive medications were eligible for randomization in the trial. Exclusion criteria included a history of a prior renal artery intervention, main renal arteries \(<4\) mm in diameter, or \(<20\) mm in length and haemodynamically or anatomically significant renal artery abnormalities. The baseline estimated glomerular filtration rate (eGFR) was required to be \(>45\) mL/min/1.73 m\(^2\) as calculated by the Modification of Diet in Renal Disease formula.\(^23\) Subjects with type 1 diabetes mellitus and stenotic valvular heart disease, or myocardial infarction, unstable angina, or cerebrovascular accident within 6 months prior to enrolment were also excluded.

All the subjects underwent a 2-week screening period both before randomization and before their 6-month follow-up to establish adherence to a stable pharmacological treatment regimen and to confirm that SBP was \(\geq 160\) mmHg. Subjects were asked to record medications in a diary for 2 weeks at baseline and again at 6-month post-randomization. Control subjects, however, were not required to keep a diary again before the 6-month measurement post-RDN treatment. The study did not include any dietary recommendations. Physicians were encouraged to maintain subjects on a stable drug regimen throughout the study period, although clinically necessary modifications of drug choice or dose were allowed. All the subjects underwent renal artery imaging (renal duplex, computed tomography, MRI, or angiogram) before randomization to establish anatomical eligibility. Among the 190 subjects assessed for eligibility, 84 subjects were excluded from eligibility, and 106 were randomly allocated 1:1 to the RDN group (\(n = 52\)) or control group (\(n = 54\)).\(^21\) (Appendix Figure 1). The RDN procedure has been previously described.\(^21\)\(^-24\) Neither study personnel nor subjects were blinded to the study group allocation. At the 6-month follow-up visit, control group subjects were eligible to crossover to receive RDN treatment. The treated crossover group reported here had a SBP \(\geq 160\) mmHg (\(\geq 150\) mmHg in subjects with type II diabetes mellitus) at the 6-month visit. Control group subjects (\(n = 9\)) with a SBP \(<160\) mmHg (\(<150\) mmHg in subjects with type II diabetes mellitus) at the 6-month visit who chose to proceed to RDN on compassionate grounds are excluded from the present blood pressure analysis but are included in the safety analysis. Participating study sites, where subjects were treated, were designated European hypertension centres of excellence in 16 of 24 instances.

**Statistical analysis**

Descriptive statistics were used to present results for the initial RDN and crossover groups. Comparisons of blood pressure and heart rate measurements at trial milestones were compared with pre-procedure measurements using the paired t-test. A change was considered significant if the two-side alpha level was \(\leq 0.05\). Variability of office blood pressure in all follow-up visits was calculated as the standard deviation (SD) or coefficient of variation (\%, 100 * SD/mean). Between group differences were compared using confidence intervals and tested using unpaired t-tests. Statistical analyses were performed using SAS version 9.2.

**Results**

**Subject characteristics**

A total of 70 subjects were available for long-term follow-up; 40 subjects in the original RDN treatment group had follow-up to 36-month post-randomization and 30 control subjects who crossed over to RDN treatment at 6-month post-randomization had follow-up to 30 months following denervation (Figure 1). Baseline characteristics are shown in Table 1. Baseline SBP was 178 ± 18 mmHg in the RDN group and 191 ± 20 mmHg in the crossover group prior to the procedure. The baseline BP for all treated subjects combined was 184 ± 19/99 ± 16 mmHg. The crossover group had more women, a higher baseline eGFR, and a higher baseline office SBP. Most subjects received a diuretic, beta-adrenergic blocker, calcium channel blocker, and an angiotensin receptor blocker or angiotensin-converting enzyme (ACE) inhibitor (Table 2).

**Blood pressure and heart rate measurements**

Systolic blood pressure and diastolic blood pressure (DBP) measurements were significantly lower than the pre-procedure measurements at all-time points (6, 12, 24, 30, and 36 months, Figure 2A...
and B); blood pressure reduction did not diminish during extended follow-up. For the 69 subjects (1 initial RDN subject missed the 30-month visit) with 30-month post-procedure follow-up, the change in SBP was −34 mmHg (95% CI: −40, −27, P < 0.01) and the change in DBP was −13 mmHg (95% CI: −16, −10, P < 0.01). The SBP and DBP reduction at 36 months in the initial RDN

---

Table 1 Baseline demographics stratified by original renal denervation and crossover groups

<table>
<thead>
<tr>
<th></th>
<th>Original RDN (n = 52)</th>
<th>Crossover (n = 52)</th>
<th>P-value</th>
<th>Original RDN (n = 40)</th>
<th>Crossover (n = 30)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Systolic pressure, mmHg systolic diastolic</td>
<td>178 ± 18</td>
<td>191 ± 20</td>
<td>0.002</td>
<td>178 ± 17</td>
<td>191 ± 21</td>
<td>0.008</td>
</tr>
<tr>
<td>Age, years</td>
<td>58 ± 12</td>
<td>57 ± 13</td>
<td>0.76</td>
<td>59 ± 11</td>
<td>58 ± 13</td>
<td>0.69</td>
</tr>
<tr>
<td>Women, n (%)</td>
<td>18 (35)</td>
<td>22 (60)</td>
<td>0.03</td>
<td>12 (30)</td>
<td>19 (63)</td>
<td>0.008</td>
</tr>
<tr>
<td>Caucasian race, n (%)</td>
<td>51 (98)</td>
<td>36 (97)</td>
<td>0.84</td>
<td>39 (98)</td>
<td>29 (97)</td>
<td>0.50</td>
</tr>
<tr>
<td>Body mass index, mean (SD)</td>
<td>31 ± 5</td>
<td>31 ± 5</td>
<td>0.06</td>
<td>31 ± 5</td>
<td>32 ± 5</td>
<td>0.01</td>
</tr>
<tr>
<td>Type II diabetes mellitus, n (%)</td>
<td>21 (40)</td>
<td>11 (30)</td>
<td>0.37</td>
<td>19 (48)</td>
<td>9 (30)</td>
<td>0.22</td>
</tr>
<tr>
<td>Coronary artery disease, n (%)</td>
<td>10 (19)</td>
<td>2 (5)</td>
<td>0.11</td>
<td>7 (18)</td>
<td>1 (3)</td>
<td>0.13</td>
</tr>
<tr>
<td>eGFR, mL/min/1.73 m²</td>
<td>77 ± 19</td>
<td>89 ± 20</td>
<td>0.005</td>
<td>77 ± 18</td>
<td>88 ± 20</td>
<td>0.02</td>
</tr>
<tr>
<td>Heart rate, b.p.m.</td>
<td>75 ± 15</td>
<td>73 ± 15</td>
<td>0.65</td>
<td>73 ± 15</td>
<td>72 ± 14</td>
<td>0.63</td>
</tr>
</tbody>
</table>

---

eGFR, estimated glomerular filtration rate.

*Initial RDN subjects followed to 3-year post-RDN therapy; crossover subjects followed to 30-month post-RDN therapy.

*Among the 51 control subjects at 6 months, 46 chose to crossover, of whom 37 met the inclusion/exclusion criteria and are included in the efficacy analysis.

*Body mass index is calculated as weight in kilograms and height in meters squared.
group was 33 mmHg (95% CI: −40, −25, P < 0.01) and 14 mmHg (95% CI: −17, −10, P < 0.01), respectively. Both the original RDN group and the crossover subjects had a similar reduction in SBP at 30-month post-RDN (−34 ± 23 vs. −33 ± 33, respectively). The coefficient of variation of systolic office blood pressure measurements (based on all subjects with at least two measurements) was 9.5 ± 4.7 (n = 80).

At 30 months after RDN, 83% of the subjects had achieved a reduction in SBP of ≥10 mmHg, and among the 40 RDN subjects with 36-month follow-up, 85% of the subjects had achieved a reduction in SBP of ≥10 mmHg (Figure 3). This response rate was similar in both the original RDN subjects and in crossover subjects: at 30 months after RDN 72% of original RDN subjects and 70% of crossover subjects had ≥20 mm reduction in SBP.

Heart rate was slowed following RDN and this decrease persisted to 36 months, with a mean decrease of 4 b.p.m. (n = 36, 95% CI: −8, −0.1, P = 0.04) (Figure 2A).

### Medications and body weight

Changes in medication class and/or prescribed dose were permitted beyond 6 months. Comparison of the number of antihypertensive medications at 36-month follow-up showed a significant reduction vs. baseline (4.6 ± 1.6 vs. 5.1 ± 1.5, P = 0.02) due in part to significantly lower usage of ACE inhibitors and centrally acting sympatholytics at 3 years (Table 2). At 30 months in all 69 subjects, the mean change in antihypertensive medications was 5.1 ± 1.4 at baseline vs. 4.8 ± 1.5 at 6 months, P = 0.06, due in part to a significant reduction in ACE inhibitors, diuretics, centrally acting sympatholytics, and direct renin inhibitors, and a significant increase in aldosterone antagonists (Table 2).

Weight was measured at all-time periods. Among subjects with 30-month follow-up, mean body weight was 91.9 ± 19.4 kg at baseline (n = 68) vs. 93.0 ± 19.9 kg at 30-month post-RDN (n = 65, P = 0.17).

### Safety

Previous safety results to 12 months on the RDN and crossover subjects have been published. Peri-procedural complications included one haematoma at the femoral access site in the original RDN therapy group and one renal artery dissection before energy delivery that required renal artery stenting in the crossover group. Clinical events to 1 year included nine hypertensive events requiring hospitalization and two hypotensive events requiring hospitalization.

Between 12 and 36 months, there were five hypertensive events requiring hospitalization as well as one case of mild transient acute renal failure due to dehydration that responded to volume expansion and temporary discontinuation of antihypertensive medications. There was one report of acute renal failure due to acute interstitial nephritis that resolved with conservative treatment and was deemed unrelated to the RDN treatment. Two subjects were hospitalized with atrial fibrillation, one of whom underwent cardioversion. There were three deaths (two occurred at home of unknown causes and one occurred due to cardiogenic shock following aortic valve replacement) that were adjudicated to be unrelated to the device or therapy by the independent safety and monitoring board. There was no change in the mean eGFR, and no renal vascular events were reported.

### Discussion

SYMPLICITY HTN-2, the first randomized controlled RDN trial, now with data out to 3 years, adds additional long-term data to the body of evidence concerning the safety and efficacy of RDN first reported in SYMPLICITY HTN-1. Our similar results confirm the substantial magnitude and long-term durability and safety of the...
RDN antihypertensive effects. Observations are available at 36 months in the group initially randomized to RDN, and at 30 months following denervation in the crossover group, the group initially randomized to control who subsequently received RDN after the 6-month primary endpoint assessment. At 36 months SBP was reduced by 33 mmHg (95% CI: 24, 22, \( P, 0.01 \)). This consistent long-term effect was achieved without increasing medication or serious long-term safety concerns.

While results in SYMPLICITY HTN-2 were consistent with results in SYMPLICITY HTN-1 and other reports with the Symplicity catheter, results in SYMPLICITY HTN-3 have been inconsistent to previous studies. SYMPLICITY HTN-3 was a rigorously designed study that required screening with 24-h ambulatory blood pressure monitoring to exclude subjects with white-coat hypertension and randomized 535 subjects in a 2:1 fashion to active denervation or a sham procedure. Both subjects and the involved staff (outside the catheterization laboratory) were blinded to treatment. But there is an Achilles heel with most clinical trials of RDN for hypertension, including this one. Typically no validated test is applied during the procedure, or subsequently to document that adequate renal denervation is performed. The 6-month reduction in office SBP was \(-14 \pm 24 \text{ mmHg}\) in the denervation group compared with \(-12 \pm 26 \text{ mmHg}\) in the sham-procedure group (\( P = 0.26 \) for prespecified superiority with a margin of 5 mmHg). The lack of significant improvement in SBP over a control arm may be related to study design, patient selection criteria and procedural details that may have caused failure to consistently achieve renal denervation. The level of procedural training for the 100+ operators in SYMPLICITY HTN-3

Figure 2 (A) Office systolic blood pressure (all \( P < 0.01 \) by 6 months after renal denervation and after crossover) and heart rate (\( P < 0.01 \) at 6, 12, 18 months, \( P = 0.06 \) at 24 months, \( P = 0.16 \) at 30 months, and \( P = 0.04 \) at 36 months and for difference in the mean heart rate in the pooled population from baseline to follow-up). RDN, renal denervation. Solid bar: original renal denervation treatment group. Hashed bar: crossover to renal denervation treatment from control group. (B) Post-procedure change in office blood pressure. Systolic blood pressure, SBP; diastolic blood pressure, DBP. \( P < 0.01 \) at all-time periods for blood pressure difference from baseline. White bar: SBP. Black bar: DBP.
at 88 centres, all new to renal denervation, was far less than that in SYMPLICITY HTN-2, which was conducted at only 24 centres with hands-on renal denervation experience for all operators prior to their performing their first denervation in the trial. This flaw in the design of SYMPLICITY HTN-3, the use of renal denervation novices, with minimal training, was likely based on the notion that renal denervation is easily achieved, which it is not. Additionally, participating SYMPLICITY HTN-2 study sites were designated European hypertension centres of excellence in 16 of 24 instances, whereas subjects in SYMPLICITY HTN-3 were often referred to the study from outside institutions and may not have exhausted all treatment options.

Despite rational multidrug antihypertensive prescribing in severe essential hypertension and patient adherence to both lifestyle measures and to medication dosing, blood pressure targets are not met in a significant proportion of patients. In the early phases of SYMPLICITY HTN-1 and SYMPLICITY HTN-2 trials, antihypertensive medications were not changed unless clinically required prior to assessing the primary BP endpoint at 6 months following RDN to allow the evaluation of the antihypertensive effect of RDN free from any confounding by dosing or medication changes. Subsequently, dosing or medication changes could be initiated by treating physicians when clinically required. On average, at the current limits of follow-up, there was a significant decrease in the number of medication classes. Thus, the sustained decrease in blood pressure beyond 6 months and out to at least 3 years cannot be readily explained by increased drug regimens. Indeed, the intent of the trial was to lower blood pressure and thereby reduces cardiovascular risk. The aim was not to achieve medication-free control of hypertension; therefore down-titration of antihypertensive drug treatment after the procedure was not part of the study protocol.

The antihypertensive response rate in this trial was 85% for reduction in office systolic BP of ≥ 10 mmHg, and 68% for ≥ 20 mmHg fall at 3 years. The introduction of RDN into clinical patient care may be aided by tests to identify good responders. A higher baseline SBP has been consistently associated with greater blood pressure reduction but other consistent predictors of response have not been identified to date. Lower-response to RDN could be due to multiple possible causes including technical failure of the RDN procedure, variable patient drug adherence, or non-neurally driven pathogenesis of the hypertension in an individual subject.

Renal denervation as a treatment option for severe uncontrolled hypertension should only be considered after careful clinical evaluation to eliminate both white-coat hypertension (using 24-h ambulatory blood pressure monitoring) and secondary causes of hypertension (primary aldosteronism being most likely to remain unidentified), and after pharmacological and other non-pharmacological treatment options have been optimally applied. The results of our trial should be interpreted in the context of some limitations. First, the number of subjects enrolled was relatively small, although significant changes in BP were observed. Secondly, our findings should not be extrapolated to patients who do not fulfil the inclusion and exclusion criteria of the SYMPLICITY HTN-2 study. Thirdly, there were losses to clinical follow-up. Fourthly, after the 6-month primary endpoint, control subjects had the option to crossover and therefore form a registry. There may have been a selection bias in terms of control subjects who chose to crossover and still met the inclusion and exclusion criteria of the SYMPLICITY HTN-2 study. Fifthly, neither study personnel nor subjects were blinded to the study group allocation. Sixthly, ambulatory blood pressure measurement (ABPM) was commonly measured at baseline in order to avoid white-coat hypertension, but this was not part of the inclusion and exclusion criteria. The study previously reported 6-month change in ABPM in initially enrolled subjects. Owing to subject non-adherence and incomplete records, an overall change in ABPM is unavailable. After the 6-month end point, 24-h blood pressure records were not collected. Seventhly, crossover subjects began with a higher BP at the time of RDN treatment when compared with the initial RDN group. The crossover subjects represent a subset of the control group; there may have been a selection bias of subjects.
who continued to have a SBP $\geq 160$ mmHg and who chose to cross-over, and this increase in blood pressure could represent a natural tendency for blood pressure to increase in subjects with treatment-resistant hypertension. However, at 30-month post-RDN, the original RDN group and the crossover group had a similar reduction in SBP and similar response rates regardless of a different baseline measurement at time of treatment.

In conclusion, we have demonstrated that renal artery nerve ablation results in sustained lowering of blood pressure out to at least 3 years (30 months for crossover subjects) in a selected population of subjects with severe, treatment-resistant hypertension. This long-term benefit occurred with minimal procedural or late safety concerns and was not associated with increased drug therapy.

Acknowledgements
The sponsor designed the study in collaboration with the study investigators and was responsible for data collection and data analysis. The authors are responsible for data interpretation and writing of the report. The corresponding author, M.D.E., had full access to all the study data and had final responsibility for the decision to submit.

Clinical Investigators: Michael Bo¨hm and Felix Mahfoud (Universita¨tsklinikum des Saarlandes, Homburg, Germany), Horst Sievert and Nina Wunderlich (CardioVascular Center Frankfurt, Frankfurt, Germany), Lars Christian Rump and Oliver Vonend (Universita¨tsklinikum Dusseldorf, Dusseldorf, Germany), Roland E Schmieder and Michael Uder (Universita¨t Erlangen-Nu¨rnberg, Erlangen, Germany), Mel Lobo and Mark Caulfield (William Harvey Research Institute, Barts and The London NIHR Cardiovascular Biomedical Research unit, Queen Mary University of London and Barts and the London NHS Trust, London, UK), Andrejs Erglis (Pauls Stradins Clinical University Hospital, Riga, Latvia), Michel Azizi and Marc Sapoval (Assistance Publique des Hôpitaux de Paris, Hôpital Européen Georges Pompidou, Paris, France), Suku Thambar (John Hunter Hospital, Newcastle, NSW, Australia), Alexandre Persu and Jean Renkin (Cliniques Universitaires Saint-Luc, Brussels, Belgium), Henibert Schunkert and Joachim Weil (Universitaetsklinikum Schleswig-Holstein, Luebeck, Germany), Hannes Reuter and Uta C Hoppe (Universita¨t zu Ko¨ln, Ko¨ln, Germany), Henry Krum and Antony Walton (The Alfred Hospital, Melbourne, VIC, Australia), Markus P Schlaich and Murray D Esler (Baker IDI Heart and Diabetes Institute, Melbourne, VIC, Australia), Dierk Scheinert (Universita¨t Leipzig—Herzzentrum, Leipzig, Germany), Thomas Binder (Allgemeines Krankenhaus der Stadt Wien, Vienna, Austria), Andrzei Januszewicz and Adam Witkowski (Samodzielnia Pracownia Hemodynamiczna, Warsaw, Poland), Luis M Ruilope (Hospital 12 de Octubre, Madrid, Spain), Robert Whitbourn (St Vincent’s Hospital, Melbourne, VIC, Australia), Heike Bruck (Universitaetsklinikum Essen, Essen, Germany), Mark Downes and Robert Kaikini (Kent and Canterbury Hospital, Canterbury, UK), Thomas F Luscher (University Hospital Zurich, Zurich, Switzerland), Alan G Jardine (University of Glasgow, Glasgow, UK), Mark W Webster (Auckland City Hospital, Auckland, New Zealand), Thomas Zeller (Herz-Zentrum Bad Krozingen, Bad Krozingen, Germany), Jerzy Sadowski and Krzysztof Bartus (The John Paul II Hospital, Jagiellonian University, Krakow, Poland).

Funding
This work was supported by Medtronic, Inc.

Conflict of interest: Nicole Brilakis, MS, MBA, Colleen Gilbert, PharmD, and Sidney A. Cohen, MD, PhD from Medtronic, Inc. provided editorial support. Martin Fahy, MS and Minglei Liu, PhD from Medtronic, Inc. provided statistical support.

Appendix

Figure A1 Study randomization, 24-h ambulatory blood pressure measurement, ABPM; month, mo; week, wk; systolic blood pressure, SBP; classes of antihypertensive medications, meds; magnetic resonance angiography, MRA; computed tomography angiography, CTA; duplex ultrasonography, duplex.
References


