

Medtronic Plans for Renal Denervation Pivotal Trial After New Study Shows Significant Blood Pressure Reductions

Published Simultaneously in The Lancet, the Late-Breaking SPYRAL HTN-OFF MED Study at ESC Successfully Isolates RDN Treatment Effect to Show Compelling Efficacy and Safety of Hypertension Procedure

DUBLIN and BARCELONA - August 28, 2017 - Medtronic plc (NYSE:MDT) today announced its intent to move forward with a new renal denervation pivotal trial following positive first results from a sham-controlled study in patients with high blood pressure. Investigators of the SPYRAL HTN-OFF MED Study found statistically significant and clinically important blood pressure reductions in the patients treated with renal denervation (RDN) across both office and ambulatory systolic and diastolic measurements. The data in the first 80 patients enrolled in the study at three months were presented in a late-breaking clinical trial session at the European Society of Cardiology (ESC) meeting in Barcelona, Spain and published simultaneously in [The Lancet](#).

"Based on the clinical results presented at ESC, we are confident that the SPYRAL HTN-OFF MED Study has successfully demonstrated the treatment effect of RDN," said co-principal investigator (PI) and presenter of the data at ESC, Professor Michael Böhm, M.D., Ph.D., chairman, department of internal medicine, University of Saarland in Homburg/Saar, Germany. "With consistent and significant drops across ambulatory and office-based blood pressure-including both systolic and diastolic measurements-we are confident to be moving forward with a pivotal trial so that doctors and patients may have an alternative approach to lowering blood pressure."

The initial three-month data found that patients who underwent renal denervation had their office systolic blood pressure (OSBP) decline an average 10 mm Hg, resulting in a 7.7 mm Hg difference with the patients in the sham control arm ($p=0.016$), and their diastolic blood pressure declined an average 5.3 mm Hg, resulting in a 4.9 mm Hg difference with the sham control arm ($p=0.008$). With a systolic blood pressure difference of 7.7 mm Hg between the two groups, a recent meta-analysis published in [The Lancet](#) associated that decrease with a relative risk reduction of approximately 20 percent for major cardiovascular events, including heart attack and stroke.

In SPYRAL HTN OFF-MED renal denervation patients, the 24-hour systolic ambulatory blood pressure (ABPM) declined 5.5 mm Hg, resulting in a 5 mm Hg difference from the sham control arm ($p=0.041$), and the diastolic AMBP declined 4.8 mm Hg, resulting in a 4.4 mm Hg difference from the sham control arm ($p=0.002$). Additionally, there were no reported cases of new or worsening renal failure, vascular complications, or other major adverse events, confirming the safety of the RDN procedure - even with a revised procedural approach that extended ablations into the kidney's branch arteries and an increase in total number of ablations compared with previous RDN trials.

"After SYMPPLICITY HTN-3, we learned a lot about the procedure itself, medication adherence and which patients may have less response to the RDN procedure-these insights have been incorporated into the revised clinical approach in the SPYRAL HTN program," said co-PI David Kandzari, M.D., director of interventional cardiology and chief scientific officer, Piedmont Heart Institute in Atlanta, Ga. "With this new approach and protocol design, coupled with new technology that allows more consistent circumferential treatment and easier access into the distal anatomy, based on these compelling results we are confident that we've appropriately addressed the issues in previous renal denervation trials."

The SPYRAL HTN-OFF MED Study evaluated RDN in the absence of any antihypertensive medications compared

to a sham-controlled population to isolate its effect on blood pressure reduction. The Study enrolled 353 patients who were either drug naïve or willing to discontinue medication for four months, with mild-to-moderate hypertension (OSBP equal to or more than 150 mm Hg and less than 180 mm Hg) at 24 centers in the U.S., Europe, Japan, and Australia. Prior to randomization, patients were required to stop taking their antihypertensive medication during a period of three to four weeks, known as a medication washout. During the washout period and the three months following randomization, patients were monitored to ensure their blood pressure would not exceed 180mm Hg.

"These results are extremely encouraging for the more than 50 percent of hypertension patients who become non-adherent to their medications within one year of initiating therapy," said co-PI Raymond Townsend, M.D., director of the hypertension program, University of Pennsylvania. "The societal implications are potentially huge as well, as high blood pressure contributes more than 500 billion dollars in direct costs to our healthcare systems globally."

Unlike SYMPLICITY HTN-3, where patients self-reported adherence, and change in medications occurred among 40 percent of patients, the SPYRAL HTN-OFF MED Study limited that variability, as patients remained off their medication throughout the duration of the study as confirmed by serum and urine testing for antihypertensive medication. Additionally, the aftermath of SYMPLICITY HTN-3 revealed new anatomical insights showing that renal nerves are easier to access when the catheter treats the renal artery segments located close to the kidney. While SYMPLICITY HTN-3 only averaged five ablations in the main renal artery and no ablations in the branch arteries, the unique, flexible design and low profile of the investigational Symplicity Spyral(TM) system achieved an average of 10 main renal artery and 13 branch artery ablations per kidney.

"These data confirm our long-held belief that the underlying science behind renal denervation is strong and we are committed to continuing our mission to realize the full potential of this procedure to help address an unmet need for the more than one billion people worldwide living with high blood pressure," said Sean Salmon, senior vice president and president of the Coronary and Structural Heart division, which is part of the Cardiac and Vascular Group at Medtronic. "We intend to continue consulting with our physician advisors and global regulatory authorities about these results so that we can appropriately move forward with a pivotal trial design to ultimately support regulatory approval in the U.S., Japan and other countries where the technology is not currently available."

The SPYRAL HTN-OFF MED study is part of the SPYRAL HTN Global Clinical Trial Program and accompanies the SPYRAL HTN-ON MED study as two global, prospective, randomized, sham-controlled trials conducted simultaneously by hypertension experts and experienced proceduralists to investigate the impact of renal denervation both in the absence of and in the presence of antihypertensive medications. The clinical program utilizes Medtronic's next-generation renal denervation technology, composed of the flexible, 6 Fr guide catheter compatible, Symplicity Spyral multi-electrode renal denervation catheter and Symplicity G3(TM) renal denervation RF (radiofrequency) generator. The Symplicity Spyral system is investigational in the United States and Japan.

About Medtronic

Medtronic plc (www.medtronic.com), headquartered in Dublin, Ireland, is among the world's largest medical technology, services and solutions companies - alleviating pain, restoring health and extending life for millions of people around the world. Medtronic employs more than 84,000 people worldwide, serving physicians, hospitals and patients in approximately 160 countries. The company is focused on collaborating with

stakeholders around the world to take healthcare Further, Together.

Any forward-looking statements are subject to risks and uncertainties such as those described in Medtronic's periodic reports on file with the Securities and Exchange Commission. Actual results may differ materially from anticipated results.

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Contacts:

Wendy Dougherty
Public Relations
+1-763-381-1204

Ryan Weispfenning
Investor Relations
+1-763-505-4626

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