Demonstration of the Anatomical Tachycardia Circuit in Sinoatrial Node Reentrant Tachycardia: Analysis Using the Entrainment Method

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Background—The anatomical tachycardia circuit of sinoatrial node reentrant tachycardia (SANRT) has not been well clarified. This study aimed to elucidate the tachycardia circuit of SANRT.

Methods and Results—Exit and entrance of the intranodal sinoatrial node conduction (I-SANC) of the reentry circuit were identified in 15 SANRT patients. After identifying the earliest atrial activation site (EAAS) during the tachycardia (EAAS-SANRT), rapid atrial pacing was delivered from multiple atrial sites to identify the entrainment pacing site where manifest entrainment and orthodromic capture of the EAAS-SANRT were demonstrated. Radiofrequency energy was then delivered starting at a site 2 cm proximal to the EAAS-SANRT in the direction of the entrainment pacing site and gradually advanced toward the EAAS-SANRT until tachycardia termination to localize the I-SANC entrance. The EAAS-SANRT was orthodromically captured by pacing delivered from the distal coronary sinus (n=7), high posteroseptal right atrium (n=2), low posteroseptal right atrium (n=2), low anterolateral right atrium (n=2), or coronary sinus ostium (n=2). Radiofrequency energy delivery to the entrance of the I-SANC, 10.4±2.8 mm away from the EAAS-SANRT, terminated tachycardia immediately after onset of energy delivery (3.4±2.3 seconds). The successful ablation site was located further from the EAAS during sinus rhythm (EAAS-sinus) than the EAAS-SANRT (12.8±4.5 versus 7.2±3.1 mm; P<0.0001).

Conclusions—The reentry circuit of SANRT was composed of the entrance and exit of the I-SANC being located at distinctly different anatomical sites. SANRT was eliminated by radiofrequency energy delivered to the I-SANC entrance, which was further from the EAAS-sinus than I-SANC exit. (*J Am Heart Assoc. 2020;9:e014472. DOI: 10.1161/JAHA.119.014472.*)

Key Words: atrial tachycardia • catheter ablation • mapping

Sinoatrial node reentrant tachycardia (SANRT) is an uncommon type of atrial tachycardia originating from the sinoatrial node region. Barker et al were the first to propose that reentry could occur within the sinus node.1 Han et al demonstrated the existence of sinoatrial reentrant echo beats in a superfused isolated rabbit right atrial preparation.2 Allessie and Bonke reported a study suggesting confinement of the reentry circuit to the sinus node in the rabbit heart.3 Glukhov et al examined the sinoatrial node reentry circuit in a canine model using high-resolution optical mapping.4 They revealed both a sinoatrial macro-reentry involving the atrial myocardium and a micro-reentry confined to within the sinoatrial node.4 In humans, Narura first described sustained SANRT, and his criteria to diagnose the SANRT are still valid.5 However, the details of the reentry circuit in patients with SANRT are still controversial to this day, because it is practically impossible to record the electrical activity of the human sinoatrial node directly by the currently used electrode catheters. Verapamil-sensitive atrial tachycardia arising from the atrioventricular annulus has electrophysiological features similar to SANRT, because a calcium-channel–dependent tissue forms the slow conduction zone of the reentry circuit.6–8 Recently, we have shown that the entrance of the slow conduction zone in verapamil-sensitive atrial tachycardia arising from the atrioventricular annulus can be identifiable and selectively ablated using the entrainment method.6–8 Therefore, we attempted to localize the entrance of the intranodal sinoatrial node conduction (I-SANC) in the SANRT reentry circuit using the same entrainment method and elucidated the anatomical reentry circuit of SANRT.

Methods

This study was approved by the hospital institutional review committee, and written informed consent was obtained from each patient. The data, analytical methods, and study
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Clinical Perspective

What Is New?

- We demonstrated manifest entrainment during the sinoatrial node reentrant tachycardia (SANRT) and showed the precise anatomical reentrant tachycardia circuit for the first time.
- SANRT was terminated by a radiofrequency energy application delivered to the entrance of the reentry circuit, distant from the exit of the reentry circuit.
- The reentry circuit was suggested to involve the perinodal atrium according to the findings of manifest entrainment and the results of the catheter ablation.

What Are the Clinical Implications?

- The entrance of the SANRT reentry circuit was located further from the earliest atrial activation site during sinus rhythm than from the exit of the SANRT reentry circuit.
- Thus, an entrance site ablation may be an alternative and safer therapeutic option than targeting the earliest atrial activation site during SANRT to avoid sinus node dysfunction.

Identification of the Entrance of the I-SANC

After the right atriography in the biplane view, the RA was mapped during sinus rhythm using a contact mapping system (EnSite NavX; St. Jude Medical), and the location of the earliest atrial activation site (EAAS) during sinus rhythm was identified. The RA was mapped again during the SANRT, to identify the location of the EAAS during the tachycardia. The location of the EAAS during SANRT was expressed relative to the EAAS during sinus rhythm. The proximity of the I-SANC in the reentry circuit of the SANRT was identified utilizing an entrainment technique. \(^\text{10}\) While recording the contact atrial electrogram at the EAAS during SANRT, rapid atrial pacing at a rate 5 beats/min faster than the tachycardia rate was delivered to demonstrate manifest entrainment and orthodromic capture of the EAAS. \(^\text{6–8,11}\) Rapid pacing was delivered from 9 sites in the RA and CS: high anterolateral RA, high posterolateral RA, high anteroseptal RA, high posteroseptal RA, low anterolateral RA, low posterolateral RA, low posteroseptal RA, CS ostium, and, distal CS. When the entrance site of the reentry circuit was located posterior to the sinoatrial node, demonstration of manifest entrainment by pacing from the RA was difficult because the entrance site was close to the left atrium. In those cases, we attempted to demonstrate the manifest entrainment by the pacing delivered from the distal CS, assuming that the orthodromic capture of the EAAS during SANRT could be obtained by the stimulated wave front propagating through Bachman’s bundle. When manifest entrainment with orthodromic capture of the electrogram at the EAAS was demonstrated, the pacing site was considered to be proximal to the I-SANC of the reentry circuit \(^\text{6–8,12}\) (Figure 1). Manifest entrainment was defined as when constant fusion of the surface P waves was demonstrated, and as when the orthodromic capture of the electrogram at the EAAS with a long conduction time and antidromic capture with direct activation by pacing showing different electrogram morphologies were both observed during 1 paced beat during pacing except for the last entrained beat \(^\text{6–8,10,12}\) (Figure 1). After identification of a site proximal to the I-SANC (entrainment pacing site, Figure 1), radiofrequency energy was delivered starting at a site 2 cm away from the EAAS in the direction of the entrainment pacing site because this site was considered to be proximally close to the entrance of the I-SANC (Figure 1). A current of

Study Population

Fifteen consecutive patients with SANRT of 359 patients with supraventricular tachycardia who were referred for radiofrequency catheter ablation from 2009 to 2019 were included in this study. There were 6 men and 9 women (mean age, 69 years; range, 59–76; Table).

Electrophysiological Study

Two 6-Fr quadripolar electrode catheters (St. Jude Medical, St. Paul, MN) were positioned in the His bundle region and right ventricular apex. A 6-Fr 20-pole electrode catheter (St. Jude Medical, St. Paul, MN) were positioned in the His bundle region and right ventricular apex. A 7-Fr 4 mm-tip, deflectable quadripolar electrode catheter with a 2-mm interelectrode distance (Japan Lifeline, Tokyo, Japan) was advanced into the right atrium (RA) for atrial mapping, pacing, and ablation. Bipolar electrograms were filtered between 50 and 600 Hz and recorded along with the surface ECG using a polygraph (EP-workmate; EP Med. Systems, Inc., Mt Arlington, NJ). Atrial and ventricular pacing was performed using a cardiac stimulator (SEC-4103; Nihon Kohden, Tokyo, Japan). The diagnosis of SANRT required the following criteria \(^\text{6,9}\):

1. (1) atrial activation sequence originating from the high lateral RA and similar to that during sinus rhythm; (2) P-wave configuration on the 12-lead ECG during the tachycardia similar to the P-wave configuration during sinus rhythm; (3) reproducible initiation and termination of the tachycardia by atrial burst and/or premature stimulation; and (4) termination with adenosine triphosphate or with maneuvers that increase the vagal tone.
15 to 20 W was delivered initially for 10 seconds with the temperature limit set to 55°C using a radiofrequency energy generator (CABL-IT; Central Inc., Ichikawa, Chiba). When the SANRT was not terminated within 10 seconds, the energy application site was advanced in a step-wise fashion by 2 to 3 mm toward the EAAS under the guidance of the EnSite anatomical map, until the tachycardia terminated in order to identify the entrance of the I-SANC of the reentry circuit (Figure 1). If the SANRT terminated within 10 seconds after onset of energy delivery, then we continued the energy delivery up to 30 to 60 seconds at that site.

Analysis of the Anatomical Reentry Circuit During the SANRT

To define the anatomical reentry circuit during the SANRT, the length of the I-SANC, which was expressed by the distance from the EAAS during SANRT (exit of I-SANC) to the successful ablation site (entrance of I-SANC), was measured. The atrial activation time between the EAAS during SANRT (exit of I-SANC) and the successful ablation site (entrance of I-SANC) during the tachycardia was also measured. To define the relative location of the SANRT reentry circuit in relation to the EAAS during sinus rhythm, the distance from the EAAS during sinus rhythm to the EAAS during the SANRT (exit of I-SANC) and that from the EAAS during sinus rhythm to the successful ablation site (entrance of I-SANC) were measured.
**Statistical Analysis**

Values are expressed as the mean±SD. Differences between clinical variables and electrophysiological parameters were analyzed using a paired Student t test for the quantitative data. A value of P<0.05 was considered significant.

**Statement of responsibility**

The authors had full access to the data and take responsibility for their integrity. All authors have read and agree with the manuscript for this article as written.

**Results**

In all patients, a tachycardia was induced and terminated by atrial rapid and extrastimulus pacing, and an inverse relationship between the A1A2 and A2Ae was observed during induction of the SANRT by atrial extrastimulus pacing. Initiation and termination of the tachycardia by atrial burst and/or premature stimulation were performed at least 3 times to confirm its reproducibility in all patients. Isoproterenol was used for induction of SANRT in 3 patients (patients 4, 5, and 10), but was not in the remaining 12. In those 3 patients, contact mapping during sinus rhythm and SANRT was performed during an isoproterenol administration. Intravenous administration of adenosine triphosphate (2.5–5.0 mg) terminated the tachycardia in all 15 patients. Vagal maneuvers were performed in 5 patients, and the SANRT was terminated in all 5 patients. Mean tachycardia cycle length was 477.0±67.4 ms (range, 370–600; Table).

**Location of the EAAS During SANRT Relative to the EAAS During Sinus Rhythm**

The EAAS during SANRT was observed in the vicinity of the EAAS during sinus rhythm in all patients (Table). The distance between the EAAS during sinus rhythm and EAAS during SANRT was 7.2±3.1 mm (range, 3–14). The EAAS during SANRT was located in the anterior (n=1), posterior (n=1), superior (n=3), inferior (n=4), anteroinferior (n=5), and posteroinferior (n=1) portion of the EAAS during sinus rhythm, respectively (Table).

**Manifest Entrainment and Catheter Ablation**

Manifest entrainment was demonstrated by rapid atrial pacing delivered from 1 specific site in each patient, being associated with the orthodromic capture of the earliest atrial electrogram (Table). Manifest entrainment was demonstrated by pacing from the distal CS (n=7), high posteroseptal RA (n=2), low posteroseptal RA (n=2), low anterolateral RA (n=2), and CS ostium (n=2), respectively (Table). In all patients, the SANRT was terminated by an application of radiofrequency energy, which was delivered to a site proximal to the EAAS during the SANRT in the direction of the manifest entrainment pacing site (Table). In 9 patients (patients 1–6, 10, 13, and 14), the successful ablation site (i.e., entrance of I-SANC) was located in the posterior portion of the EAAS during the SANRT (Table). The successful ablation site was located at the posteroinferior portion of the EAAS during the SANRT in 4 patients (patients 7–9 and 12; Table) and at the inferior portion of the EAAS during the SANRT in the remaining 2 patients (patients 11 and 15; Table). The distance between the EAAS during SANRT (exit of I-SANC) and the successful ablation site (entrance of I-SANC) was 10.4±2.8 mm (8–16; Table). Onset of the atrial electrogram at the successful ablation site occurred 14.5±3.4 ms later than that of the EAAS during the SANRT (Table). The SANRT was terminated immediately after onset of radiofrequency energy delivery (3.4±2.3 seconds; Table). The mean number of radiofrequency applications required for a successful ablation was 4±2. After termination of the tachycardia by the initial successful ablation, a reinduction of the tachycardia was attempted, but no reinduction of SANRT was observed in any of the patients. The ablation was not associated with any complications. During a mean follow-up period of 53±43 months (range, 12–122), neither tachycardia recurrence nor sinus node dysfunction were observed.

Figure 2 shows the tracing during manifest entrainment in patient 7. The EAAS during the SANRT was observed in the high right atrium (HRA; 7–8; Figure 2). During pacing from the low posteroseptal RA, the EAAS (HRA 7–8) and His bundle 5 to 6 recording sites were orthodromically captured by a long conduction interval (solid arrows). All of the electrograms at the EAAS (HRA 7–8) and His bundle 5 to 6 recording sites during pacing showed morphologies identical to those obtained during the SANRT, and the cycle lengths of those sites were all 425 ms, identical to the pacing interval, and shorter than the SANRT cycle length (440 ms). On the other hand, the atrial electrograms at CS 9 to 10 were captured antidromically during pacing (dashed red arrows). The atrial electrograms at CS 9 to 10 occurred 10.4±3.4 ms later than that of the EAAS during the SANRT (Table). The SANRT was terminated immediately after onset of radiofrequency energy delivery (3.4±2.3 seconds; Table). The mean number of radiofrequency applications required for a successful ablation was 4±2. After termination of the tachycardia by the initial successful ablation, a reinduction of the tachycardia was attempted, but no reinduction of SANRT was observed in any of the patients. The ablation was not associated with any complications. During a mean follow-up period of 53±43 months (range, 12–122), neither tachycardia recurrence nor sinus node dysfunction were observed.

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Figure 3 shows the isochronal map during the tachycardia showing the relative locations of the EAAS during the SANRT, the EAAS during sinus rhythm, successful ablation site, and entrainment pacing site (low posteroseptal RA) in patient 7. The EAAS during the tachycardia was observed at the posteroinferior portion of the EAAS during sinus rhythm. The SANRT was terminated by an energy application delivered at a site 12 mm proximal to the EAAS during the SANRT in the direction of the entrainment pacing site. Thus, the successful ablation site (entrance of I-SANC) was located at the posteroinferior portion of the EAAS during the SANRT.

Figure 4 shows the tracing during a radiofrequency energy application delivered at a site 12 mm proximal to the EAAS (successful ablation site) in patient 7. The atrial electrogram of the successful ablation site appeared 15 ms later than that of HRA 7 to 8 during the SANRT, and the unipolar electrogram exhibited an rS morphology (Figure 4A). However, the SANRT terminated immediately after onset of radiofrequency energy delivery (1.6 seconds; Figure 4B), which suggested that the radiofrequency energy application site was located at the entrance of the I-SANC.

Figure 5 shows the tracing during manifest entrainment in patient 13. The atrial electrogram at the EAAS during the SANRT preceded that at HRA 7 to 8 by 10 ms. During pacing from the distal CS (CS 1–2), the EAAS and HRA recording sites were orthodromically captured by a long conduction interval (solid arrows). All of the electrograms at the EAAS and HRA recording sites exhibited morphologies identical to those during the tachycardia, and the cycle lengths of those electrograms at the orthodromically captured sites were all 490 ms, identical to the pacing interval and shorter than the tachycardia cycle length (510 ms). On the other hand, the atrial electrograms at CS 5 to 6 were captured antidromically during pacing (dashed red arrows). The atrial electrograms at CS 5 to 6 occurred 25 ms later than those at HRA 7 to 8 during pacing, but they were observed 60 ms later during the tachycardia. Furthermore, the electrogram morphologies at CS 5 to 6 during pacing differed from those during the SANRT and the interval of the atrial electrogram at CS 5 to 6 just after pacing (535 ms) was longer than the pacing cycle length (490 ms), indicating the antidromic capture of the electrogram at CS 5 to 6 (dashed red arrows). In addition, the surface P-wave morphologies in lead II during pacing (open arrows) differed from those during the tachycardia (closed arrows), indicating fusion of the surface P waves during pacing except for the last captured beat.
Figure 6 shows the isochronal map during the tachycardia showing the relative locations of the earliest atrial activation site (EAAS) during the SANRT, the EAAS during sinus rhythm, successful ablation (ABL) site, and entrainment pacing site in patient 7. IVC indicates inferior vena cava; RA, right atrium; RAA, right atrial appendage; SANRT, sinoatrial node reentrant tachycardia; SVC, superior vena cava; TV, tricuspid valve.

Anatomical Location of the SANRT Reentry Circuit

The atrial electrogram at the successful ablation site was observed significantly later than that at the EAAS during the SANRT (14.5±3.4 versus 0±0 ms; P<0.0001; Figure 8A). The distance between the EAAS during sinus rhythm and successful ablation site was significantly longer than that between the EAAS during sinus rhythm and EAAS during the SANRT (12.8±4.5 versus 7.2±3.1 mm; P<0.0001; Figure 8B). That indicated that the successful ablation site (entrance of I-SANC) was located further from the EAAS during sinus rhythm than the EAAS during the SANRT (exit of I-SANC).

Discussion

This study revealed several new, important findings. First, we demonstrated manifest entrainment during the SANRT and, to the best of our knowledge, showed the precise anatomical
reentrant tachycardia circuit for the first time in humans. Second, the SANRT could be terminated by a radiofrequency energy application delivered to the entrance of the reentry circuit, distant from the exit of the circuit, under navigation by manifest entrainment. Third, it was suggested that the reentry circuit involved the perinodal atrium according to the findings of manifest entrainment and the results of the catheter ablation.

Tachycardia Circuit of the SANRT

In the present study, manifest entrainment associated with fusion of the surface P wave was demonstrated and the EAAS was orthodromically captured during entrainment in all patients. Orthodromic capture of the EAAS during manifest entrainment implies that there is an area of slow conduction within the reentry circuit between the entrance and exit sites.\(^6\)\(^-\)\(^8\),\(^10\),\(^11\) For fusion of surface P waves to occur, the tachycardia (ie, orthodromic) and stimulated (ie, antidromic) wavefronts must collide with each other in the atria after the tachycardia wavefront exits the I-SANC\(^6\)\(^-\)\(^8\),\(^10\),\(^11\) (Figure 1). This requires the stimulated wavefront to have access to an entrance site of the reentry circuit that is anatomically distinct from the exit site.\(^6\)\(^-\)\(^8\),\(^13\) Therefore, fusion of the surface P waves with orthodromic capture of the EAAS during manifest entrainment suggested the presence of an entrance of the I-SANC distinct from the exit of the I-SANC (EAAS during SANRT). Satoh et al showed orthodromic capture of the earliest atrial electrogram at the His bundle site by rapid pacing delivered from the coronary sinus during atrioventricular nodal reentrant tachycardia.\(^14\) They clearly demonstrated that orthodromic capture of the EAAS implies the presence of an entrance and an exit to the atrium located at distinctly different locations.\(^14\) In the present study, SANRT was successfully terminated by an energy delivery to a site proximal to the EAAS in the direction of the pacing site from where the manifest entrainment was demonstrated. These successful ablation sites were located 10.4±2.8 mm away from the EAAS during the SANRT, which was consistent with the previous findings of manifest entrainment.\(^6\)\(^-\)\(^8\),\(^10\)\(^-\)\(^14\)

Recent optical mapping of explanted human and canine hearts revealed that the sinoatrial node complex has at least 4
preferential sinoatrial conduction pathways, which are responsible for the transmission of electrical impulses to the atrial myocardium.\textsuperscript{15,16} These include the superior, lateral, inferior, and septal sinoatrial conduction pathways.\textsuperscript{15,16} The atrial breakthrough site is approximately 7.5 mm away from the leading pacemaker in the human sinoatrial node,\textsuperscript{17} but the distance between the sinoatrial node leading pacemaker and atrial breakthrough sites could vary from 3 to 25 mm because of multiple sinoatrial conduction pathways.\textsuperscript{15,16} It has been shown that these sinoatrial conduction pathways have different conduction properties and could lead to beat-to-beat variations in the atrial activation.\textsuperscript{15,16} In the present study, the EAAS during sinus rhythm slightly differed from that during SANRT. The difference in heart rate between sinus rhythm and SANRT might have resulted in a shift in the exit of the sinoatrial conduction pathways.\textsuperscript{15,16} These multiple active sinoatrial conduction pathways may form the macro-reentry circuit involving the atrial myocardium as shown in the canine model.\textsuperscript{4}

**Catheter Ablation of SANRT**

Catheter ablation of SANRT has been conducted targeting the EAAS during tachycardia.\textsuperscript{9,18,19} Kay et al reported that SANRT was terminated promptly upon targeting the EAAS; however, tachycardia could be reinduced in 2 of 4 patients and was associated with a slight shift in the EAAS.\textsuperscript{18} In those 2 patients, the EAAS shifted 2 to 3 mm laterally after the initial application of radiofrequency current. Therefore, successful ablation of the focus of the SANRT required the application of radiofrequency current within an arc spanning 3 to 5 mm in

Figure 5. Tracing during manifest entrainment by pacing from the distal coronary sinus (CS 1–2) in patient 13. The ECG leads I, II, and V1 and electrograms recorded from the high right atrium (HRA), coronary sinus (CS), mapping catheter (MAP), His bundle (HB) position, and right ventricle (RV) are shown. d indicates distal; EAAS, earliest atrial activation site; p, proximal; Stim, stimulation; uni, unipolar electrogram.
the region of the sinus node. Thus, they indicated that reentry occurs within the sinus node, sinoatrial atrial myocardium, or a combination of these regions with different exit sites into the surrounding atrial myocardium. Goya et al reported that ablation of SANRT targeting the EAAS was successful in all patients. However, the tachycardia was still inducible after the SANRT was terminated once by a radiofrequency energy delivery in 6 of 11 patients, associated with a shift in the EAAS from approximately 2 to 7 mm (mean, 4.3) laterally and inferiorly from the initial site of the current application. Therefore, they indicated that successful ablation of SANRT requires an energy application over a considerably large area, extending 2 to 7 mm in the region of the sinus node. Those previous findings indicated that the SANRT circuit can be changed accompanied by a shift in the exit sites, suggesting the presence of multiple sinoatrial conduction pathways, whereas in the present study neither reinduction nor a change in location of the EAAS during the SANRT were observed in any of the patients after the successful ablation of the SANRT. Thus, radiofrequency energy delivery to the entrance site of the reentry circuit might be more effective in preventing the occurrence of different SANRTs accompanied by a shift in the EAAS than targeting the exit of the reentry circuit.

It has been reported that a radiofrequency energy delivery to the sinus node region can produce sinus node dysfunction requiring a pacemaker implantation. In the present study, the EAAS during the SANRT was observed in the vicinity of the EAAS during sinus rhythm. However, the successful ablation site (entrance of the I-SANC) was located further from the EAAS during sinus rhythm than the EAAS during the SANRT. Therefore, application of radiofrequency energy to the entrance of the I-SANC may be an alternative and safer therapeutic option than targeting the EAAS during SANRT.
Study Limitations

Previous reports demonstrated a shift in the leading pacemaker and preferential sinoatrial conduction pathways in human hearts after exposure to adenosine or isoproterenol.\textsuperscript{4,21} Therefore, additional functional sinoatrial conduction pathways might be found if we compare the activation sequences between those with and without the administration of isoproterenol and adenosine triphosphate.

Although we regarded the last radiofrequency energy application site at where the tachycardia was terminated as the successful site, there was a possibility that the successful elimination of tachycardia occurred as a cumulative effect of the previous unsuccessful ablation lesions. Also, the successful ablation site might not be just at the entrance site, given that the successful ablation site was identified only by the findings from entrainment. It is possible that the successful ablation site was slightly different from the entrance of the SANRT circuit.

It may be possible that multiple locations can show manifest entrainment in cases in which multiple entrance sites contribute to the reentry circuit. In such cases, multiple radiofrequency energy deliveries to each entrance site may be necessary to terminate the tachycardia.

Although we used only a quadripolar catheter for mapping with the EnSite NavX system, the use of a high-density mapping catheter might have delineated the reentry circuit more precisely.

Conclusions

The reentry circuit of the SANRT was composed of the entrance and exit of the I-SANC located at distinctly different anatomical sites. A radiofrequency energy application to the entrance of the I-SANC, which was identified under navigation with entrainment, can eliminate the SANRT. The entrance of the I-
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SANC was located further from the EAAS during sinus rhythm than the exit site, suggesting that an entrance site ablation may be an alternative and safer therapeutic option than targeting the EAAS during SANRT to avoid sinus node dysfunction.

Disclosures
None.

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