Cardioneuroablation in ictal asystole—New treatment method

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Introduction
We describe a case of successful parasympathetic denervation of the sinus node using cardioneuroablation in a patient with right temporal lobe epilepsy and prolonged ictal asystole. The procedure abolished seizure-induced bradyrhythmia occurrence and converted the patient’s dramatic seizures with severe cerebral hypoperfusion into short focal seizures with minimal motor signs. To the best of our knowledge, this is the first report of a successful cardioneuroablation procedure to potentially treat ictal asystole.

Case report
A 43-year-old right-handed male patient with pharmacoresistant focal epilepsy was admitted to the Department of Neurology for evaluation before potential epilepsy surgery. He had an extensive focal cortical dysplasia in the right temporal lobe and mild left-sided spastic hemiparesis. Seizures were resistant to several antiepileptic drugs, including levetiracetam and carbamazepine in maximal dosages he was taking before the admission. For the past 5 years, almost every seizure resulted in loss of consciousness, falls, and, on several occasions, traumatic injuries. During long-term video-electroencephalography (EEG) monitoring, the antiepileptic drugs were temporarily withdrawn, and 9 epileptic seizures were recorded. The seizures manifested with ictal pouting (“chapeau de gendarme”), right leg automatisms and subtle pelvic movements, and dystonic left hand posturing, followed by unresponsiveness, upward gaze deviation, and posturing resembling a decerebration pattern. In EEG, focal ictal activity started in the right temporal region, quickly evolving over both hemispheres, and was then followed by postictal EEG slowing over the right hemisphere. Unexpectedly, in the late postictal period we observed generalized EEG flattening and slowing, indicative of cerebral hypoperfusion. Simultaneously, on electrocardiography sinus heart rate slowing with asystole lasting up to 25 seconds was recorded, resulting in the patient’s syncope (Figure 1A, Supplemental Video).

Immediately surgical treatment of epilepsy was not feasible and also, considering the extensive nature of the presumed epileptogenic lesion, chances of favorable epilepsy surgery outcome were estimated to be low. The cardiology team was consulted. Since asystole after the seizure onset was suggestive of an increased direct vagal stimulation of the cardiac conduction system, we decided to try cardioneuroablation as an alternative to permanent pacemaker implantation.

After the patient’s consent, the cardioneuroablation procedure was performed using mild sedation with midazolam and additional fentanyl boluses. Mapping and ablations were performed using a 3.5-mm irrigated-tip catheter...

KEY TEACHING POINTS
- Seizure-induced (ictal) asystole is caused by direct vagal stimulation of the cardiac conduction system. It is estimated that only 0.27% of epileptic patients suffer from the condition, which is hypothesized as one of many potential mechanisms of sudden unexpected death in epilepsy.
- Several approaches are proposed to treat ictal asystole, including adjustment of antiepileptic drugs, epilepsy surgery, and, only recently, permanent pacemaker implantation.
- Cardioneuroablation—parasympathetic denervation of the sinus node—might represent a new treatment option in select patients with ictal asystole. In addition, with this treatment option permanent pacemaker implantation could be avoided and device-related complications prevented.

KEYWORDS Cardioneuroablation; Ictal asystole; Parasympathetic denervation; Syncope; Vagal denervation (Heart Rhythm Case Reports 2018;4:523–526)
Initially, 3-dimensional virtual anatomy of the right (RA) and left atrium (LA) was created using the CARTO 3 fast anatomic mapping system (Biosense Webster, Diamond Bar, CA), facilitated by intracardiac echocardiography (AcuNav, Siemens Medical Solutions, Mountain View, CA). Tagging the phrenic nerve capture points on the lateral RA allowed us to map the nerve course (Figure 2). Afterwards, the electrogram fractionations indicative of epicardial ganglia presence were mapped and tagged in the anatomic areas where epicardial parasympathetic ganglia for sinus and atrioventricular node innervation are located (anteriorly and superiorly to the right superior pulmonary vein [RSPV] and anteriorly to the right inferior pulmonary vein in LA). These locations correspond to locations on the posterior septal side of the RA, where fractionated electrograms were also mapped and tagged. Multiple ablations (power control, 25 W in RA, 30 W in LA, target contact force 10–30 g, duration up to 40 s, temperature limit 43°C, total radiofrequency time 1544 s) were performed in target areas from both the LA and RA, with care taken to avoid ablations in the proximity of the mapped course of the phrenic nerve. We aimed to achieve local electrogram attenuation and ablation index (Biosense Webster, Diamond Bar, CA) of 350 to 500. During initial ablations anteriorly to the RSPV, increase in heart rate was noticed, indicating parasympathetic denervation of the sinus node. Later ablations more inferiorly along the posterior interatrial septum from the LA and RA were anatomically guided. After no additional increase in heart rate was achieved with further ablations and target anatomic area was sufficiently densely ablated, the procedure was terminated.

Atropine test (3 mg of atropine sulfate given intravenously; dose calculated as 0.04 mg/kg body weight, maximal dose 3 mg) at the end of the procedure resulted in only 7% sinus rate increase (from 117 beats/min to 125 beats/min), which according to published data suggests successful denervation of the sinus node. The procedure duration was 200 minutes; fluoroscopy time was 6.8 minutes (dose-area product 560 μGy²).

To evaluate the impact of cardioneuroablation on bradyarrhythmia occurrence, an implantable loop recorder (ILR; Reveal Linq, Medtronic, Minneapolis, MN) was implanted. The patient resumed antiepileptic drugs and was released from the hospital.
Lesions on the interatrial septum.

Network.

Cingulate gyrus, and other parts of the central autonomic temporal and mesial frontal lobe epilepsy involving insula, a devastating condition is a feature of focal, most commonlylation of the cardiac conduction system. This rare but autonomous networks, which can result in direct vagal stimu-

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Discussion

Seizure-induced asystole, or ictal asystole, is caused by spread of ictal activity to loci where it intervenes with central autonomic networks, which can result in direct vagal stimulation of the cardiac conduction system. This rare but devastating condition is a feature of focal, most commonly temporal and mesial frontal lobe epilepsy involving insula, cingulate gyrus, and other parts of the central autonomic network.

It is estimated that only 0.27% of epileptic patients suffer from the condition, which is hypothesized as one of many potential mechanisms of sudden unexpected death in epilepsy.

Several approaches are proposed to treat ictal asystole, including adjustment of antiepileptic drugs, epilepsy surgery for medically refractory patients, and, only recently, permanent pacemaker implantation. However, despite the technological development of implantable electronic cardiac devices, complication incidence in modern pacing therapy is still substantial. Although most adverse events occur in the early postimplantation period (lead-related), complication rates during long-term follow-up are not scarce, ranging from 7.5% to almost 10% of the patients.

Pachon and colleagues first described cardioneuroablation in 2005 as a new treatment modality for neurocardiogenic syncope, functional sinus node dysfunction, and functional atrioventricular block. The technique is based on radiofrequency ablation of main epicardial parasympathetic ganglia in the heart. With this procedure partial parasympathetic denervation of the sinus and/or atrioventricular node is achieved and consequently the adverse parasympathetic influence on the heart diminished. Long-term follow-up of patients undergoing cardioneuroablation procedure for cardioinhibitory neurocardiogenic syncope showed very promising results, with only 3 patients out of 43 experiencing recurrent syncope. In addition, postprocedure 24-hour Holter electrocardiograms and stress tests showed no major abnormalities except mildly elevated basal heart rate.

Although the cardioneuroablation procedure considerably diminished the patient’s seizure presentation, longer follow-up and close monitoring of the patient are required to detect possible recurrence of bradyarrhythmias and worsening of seizures. Consequently, pharmacoresistant epilepsy merits constant evaluation for potential surgical treatment.

Conclusion

Cardioneuroablation might represent a new treatment modality in select pharmacoresistant patients with ictal asystole. In addition, with this treatment option permanent pacemaker implantation could be avoided and device-related complications prevented. Further research is warranted to evaluate this treatment option.

Appendix

Supplementary data

Supplementary data associated with this article can be found in the online version at https://doi.org/10.1016/j.hrcr.2018.07.018.

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