A case of ‘tachy-brady syndrome’: What is the mechanism?

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A young male presented with incessant narrow QRS tachycardia and left ventricular dysfunction. 24-Holter monitoring revealed multiple episodes of sustained and non-sustained episodes of tachycardia with prolonged sinus pauses at termination. The analysis of the electrocardiogram, followed by an invasive electrophysiological study, suggested an unusual mechanism for this tachy-brady syndrome.

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1. Case presentation

An 18-year old male presented with recurrent palpitations associated with occasional presyncope for the preceding 3 months. Nonsustained episodes of narrow QRS tachycardia with sinus pauses at termination were noted on the electrocardiogram (ECG) (Fig. 1A & B). Holter monitoring showed sustained and nonsustained episodes of tachycardia, with a cumulative duration of 14 h in a 24-h recording, and intervening sinus pauses (Fig. 1C) lasting up to 4.5 s. His left ventricular ejection fraction was 35% at presentation.

During electrophysiological study, sinus cycle length was 734 ms with normal atrio-hisian (AH) and His-ventricular (HV) intervals. Atrial pacing till atrioventricular (AV) weneckbeaching didn’t reveal any preexcitation. No dual AV nodal physiology was seen. The clinical tachycardia was easily induced on ventricular pacing. The termination and re-initiation of the tachycardia are shown in Fig. 2.

2. Discussion

The ECG in Fig. 1A and B show a narrow QRS long RP tachycardia with negative P waves in inferior leads. The episodes of this nonsustained tachycardia are initiated with sinus beats, and the second QRS complex of the tachyarrhythmia shows right bundle branch block morphology. The morphology and axis of the subsequent P waves (P’) are different from the sinus beats. The short tachycardia episodes terminate with P’ wave and are followed by sinus pauses. Occasionally, the isolated sinus beat is followed by a single P’ wave, without continuing as tachycardia. The differential diagnosis of this long RP tachycardia includes atrial tachycardia (AT) originating from inferior
atrium, AV nodal reentrant tachycardia (AVNRT) of fast-slow sub-form and persistent form of junctional reciprocating tachycardia (PJRT). During AT, the P waves of the initial and subsequent beats tend to be similar, unlike what is observed in this case. The fast-slow type of AVNRT is usually initiated following ventricular premature beat and the incessant nature is unusual. Orthodromic reentrant tachycardia using an accessory pathway (AP) with decremental retrograde conduction properties, i.e., PJRT can be initiated by sinus beat, and this tachyarrhythmia tends to be incessant. Termination on either antegrade or retrograde limb can occur in both reentrant mechanisms, but to terminate with a P’ wave is unusual for AT.

The intracardiac tracing in Fig. 2 helps to confirm PJRT as the mechanism in this case. The initial 2 cycles of tachycardia in Fig. 1 have cycle length of 360 ms with the earliest atrial activity recorded with ‘bracketing’ at proximal coronary sinus (CS) bipolar (CS 7-8). The tachycardia terminates following a paced beat from the right ventricular apex, timed 30 ms later than the His signal. The morphology of this paced complex suggests QRS fusion between the paced and tachycardia beat. Termination following a paced ventricular beat not conducted to atrium and resulting in QRS fusion suggests orthodromic AV reentry as the tachycardia mechanism. The prolonged local ventriculoatrial (VA) interval of 210 ms even at the site of earliest retrograde atrial activation during the tachycardia suggests slow conduction through the AP. Immediately following its termination, the next sinus beat re-initiates the tachycardia. Notably, the first local V–V interval at CS 7-8 after the re-initiation is 370 ms, as compared to 360 ms during its stable phase. The increment of 10 ms is contributed by a delay in the AH interval by 60 ms (40–100 ms) despite the shortening of the conduction time across the AP by 40 ms (210–170 ms). Despite a delay in conduction down the His bundle, the anterograde impulse finds the right bundle in refractoriness, and conducts down the left bundle with an HV interval similar to that in other tachycardia beats. This functional block and its mitigation in the subsequent beats follows that the recovery period in the bundle branch is directly related to the previous RR interval.

The conduction times through the pathway, as measured by the local VA interval in CS 7-8 bipolar, showed progressive prolongation (170, 190 and 210 ms respectively), in response to a shortening of the preceding local V–V intervals (720, 370 and 330 ms respectively). This shows a decremental nature of the conduction through the AP. The progressive reduction in the time interval of the retrograde impulses at the ventricular end of the AP during the re-initiation, related to the conduction delay in the AV node in the second beat, unmasked.

![Intracardiac tracing](image_url)

**Fig. 1** – A) 12-lead ECG showing the long RP tachycardia initiated by a sinus beat. The first sinus beat is followed by a non-conducted atrial echo beat. B) Rhythm strips (lead II) showing the onset and termination of the nonsustained episodes of tachycardia. C) Holter showing episodes of tachycardia interspersed with sinus pauses.
the decremental properties of the AP in this case. Similarly, the decremental nature of the conduction in the AV node accounted for the minimal changes in AV interval in the initial cycles during the re-initiation. Frequently, the sinus beats were followed by single atrial echo beats, non-conducted down the AV node (Fig. 1B). However, if this echo beat could find AV node during its relative refractory period and conduct with a prolonged AH interval and bundle branch aberrancy, the reentry could be initiated. During the following cycles, the minimal but incremental prolongation in the conduction time through the AP due to its decremental nature helped the retrograde impulse find a non-refractory atrium and AV node to maintain the reentry.

The tachycardia was incessant during the study and could be terminated only with ventricular pacing. The AP was successfully ablated at CS, within 1 cm of its ostium. Sinus nodal function tests done immediately following the ablation showed normal values. Holter monitoring done 3 months after the study showed no pauses or tachycardia and the ventricular function recovered completely to suggest tachycardiomopathy at presentation.

Symptomatic sinus pauses at the termination of the tachycardia episodes, mimicking a ‘tachy-brady syndrome’ is unusual in PJRT. The patient was not any negative chronotropic agents to account for the prolonged pauses. Suppression of the sinus node related to chronic atrial arrhythmias and its reverse remodeling over weeks to months following successful ablation of the culprit arrhythmia has been described [1,2]. However, the complete recovery of sinus node function immediately after the ablation makes this mechanism unlikely. A compensatory vagotonia in response to the initial hypotension and resultant sympathetic activity has been proposed as the dominant mechanism in spontaneous and early termination of supraventricular tachyarrhythmias [3]. The tachycardia was incessant during the electrophysiological study with infrequent spontaneous termination. However, no significant sinus pauses occurred at the termination of tachycardia episodes on pacing protocols or successful ablation. This also would suggest that a common denominator like vagotonia accounted for the spontaneous termination of tachycardia and the sinus pauses followed. Like the AV node, the AP mediating PJRT is also known to be vagal sensitive. Interestingly, the termination with a block on the antegrade limb was more commonly noted in this case presumably related to a higher sensitivity of the AV node to vagotonia compared to the AP.

Fig. 2 – Intracardiac electrogram during the tachycardia. The tracings from the top to bottom are leads V1, I and II, followed by intracardiac electrograms from high right atrium (HRA), distal His bundle (HBD), coronary sinus proximal to distal (CS), and the right ventricular apex (RVA). A paced beat from the RVA during His refractoriness terminates the tachycardia and the next sinus beat reinitiates the tachycardia. The QRS morphology of the paced beat suggests ventricular fusion.
Disclosures

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References

[1] Elvan A, Wylie K, Zipes DP. Pacing-induced chronic atrial fibrillation impairs sinus node function in dogs: electrophysiological remodeling. Circulation 1996;94:2953–60.
[2] Sparks PB, Jayaprakash S, Vohra JK, Kalman JM. Electrical remodeling of the atria associated with paroxysmal and chronic atrial flutter. Circulation 2000;102:1807–13.
[3] Waxman MB, Sharma AD, Cameron DA, Huerta F, Wald RW. Reflex mechanisms responsible for early spontaneous termination of paroxysmal supraventricular tachycardia. Am J Cardiol 1982;49:259–72.