Complete heart block in cardiac sarcoidosis reversed by corticosteroid therapy: time course of resolution

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SUMMARY
A 53-year-old man was admitted for recurrent syncope and found to have complete heart block (CHB). Cardiac magnetic resonance imaging (MRI) showed extensive patchy late gadolinium enhancement in the apical and lateral walls, consistent with cardiac sarcoidosis (CS) but no scar in the septum. A fluorodeoxyglucose (FDG)–positron emission tomography showed FDG uptake in the septum and basal lateral walls. Imaging suggested active inflammation in the septum affecting atrioventricular (AV) conduction but no irreversible fibrosis. Diagnosis of isolated CS requires a high level of suspicion and multidisciplinary teamwork involving heart failure specialists, electrophysiologists and rheumatologists. After specialist and patient discussion, treatment of the disease was initiated with prednisone 40 mg daily, 11 months after presenting with CHB. Three weeks later, ECG with pacing inhibited showed second-degree AV block Mobitz type II and 4 weeks later, AV conduction recovery. This highlights the importance of immediate therapy in reversing AV conduction abnormalities in CS.

BACKGROUND
Isolated cardiac sarcoidosis (CS) often escapes detection given no clinically apparent disease in other organs. Diagnosis requires high index of suspicion based on cardiac manifestations and advanced imaging modalities. Its presentation is usually that of an idiopathic atrioventricular block (AVB). Imaging may show active disease versus myocardial fibrosis which determines the benefit of immunosuppressant therapy. A patient with complete heart block (CHB) received prednisone for suspected CS 11 months after presentation. Three weeks later, the underlying rhythm was second-degree AVB Mobitz II and 4 weeks after therapy initiation, he had atrioventricular (AV) conduction recovery. We report a case of improved AV conduction after a delayed diagnosis of CS as well as collaborative work between electrophysiologists, heart failure specialists and rheumatologists in the management of isolated CS.

CASE PRESENTATION
A 53-year-old Caucasian man with a history of obesity, fatigue for 7 years and right bundle branch block for 10 years was admitted at an outside facility for recurrent syncope. At the time of his presentation, he was found to have CHB on ECG. On physical examination, his blood pressure was 97/54 mm Hg with a heart rate of 32 beats per minute. The rest of his examination was otherwise unremarkable.

INVESTIGATIONS
Prior to developing syncope with CHB and to having his care transferred to our facility, he underwent extensive evaluation of other symptoms which included dyspnoea and fatigue. A sleep study revealed severe sleep apnea for which he was placed on continuous positive airway pressure. Two trans thoracic echocardiograms (TTEs) 1 year apart showed an ejection fraction (EF) of 50%–55%, with no valvular disease and normal atrial sizes. However, it revealed inferior wall hypokinesis and grade I diastolic dysfunction. An exercise stress echocardiogram revealed mid-inferior and interseptal infarction without ischaemia. Over the years, he had coronary angiograms due to continued concerns of dyspnoea which were negative for obstructive coronary artery disease. Multiple chest X-rays were negative for any cardiopulmonary processes. A chest CT revealed hepatic steatosis but no pulmonary process or lymphadenopathy. He then developed light-headedness and eventually, syncopal episodes. For the newly discovered CHB on ECG, he had a dual-chamber pacemaker implanted. During his follow-up clinic visits, he continued to have dyspnoea prompting another evaluation of his coronary arteries with a coronary angiogram which was again negative for obstructive disease. Seven months after his last TTE, he had cardiac magnetic resonance imaging (MRI) which showed an EF of 21%. On pacemaker interrogation, he was noted to have episodes of sustained ventricular tachycardia (VT). His device was therefore upgraded to a cardiac resynchronisation therapy defibrillator. He did not have any physical signs of volume overload. At the time of the cardiac MRI, he was noted to be ventricular paced 86% of the time. Given the findings on imaging, he was referred to our facility for further evaluation and management of his heart failure.

On further evaluation at our institution, Lyme antibody titres were negative. During a 6-minute walk test, he ambulated 1480 ft with no hypoxia. A repeat cardiac MRI showed extensive patchy late gadolinium enhancement (LGE) accounting for 16% of the left ventricle in the lateral and apical walls, concerning for CS (figure 1). EF was 21% and the septum appeared hypokinetic. Due to concerns of CS, his care was collaborated between the advanced heart failure and sarcoidosis clinics in the electrophysiology department at our facility.

In order to determine whether he was having active inflammation due to sarcoidosis, he had a fluorodeoxyglucose–positron emission tomography (FDG-PET). This showed a large, severe perfusion defect in the mid-inferior, interseptal and basal...
anteroseptal walls with FDG uptake in the septum and basal lateral walls (figure 2). There was no FDG uptake of lymph nodes in the axilla, mediastinum or hila. Endobronchial ultrasound–guided lymph node biopsy was negative for inflammation or granuloma. Troponin levels were negative on two occasions. ACE levels were not obtained. An evaluation by a hematologist was negative for any skin manifestations of sarcoidosis. There was no concern for eye inflammation. Spirometry revealed mild restrictive lung disease with mildly reduced diffusing capacity of the lungs for carbon monoxide. Endomyocardial biopsy was offered to the patient; however, he declined.

TREATMENT
Given imaging findings on cardiac MRI and FDG-PET were suggestive of presumptive CS, prednisone 40mg daily was initiated 11 months after presentation with syncope and CHB. Systolic congestive heart failure was treated with carvedilol, eplerenone and sacubitril/valsartan. Due to hepatic steatosis, methotrexate was not initiated. However, azathioprine was initiated 2 months after prednisone, after obtaining thiopurine methyltransferase levels. TTE 3 months later showed an EF of 40%–45%. At follow-up, ECG with pacemaker inhibited showed sinus rhythm with AV conduction recovery in an additional 1 week.

OUTCOME AND FOLLOW-UP
Four weeks after starting steroid therapy, the patient showed improvement in fatigue. Steroid tapering was initiated at that time. Three months after initiating heart failure treatment, he showed near-complete resolution of fatigue and dyspnoea. He has been able to return to work as a security officer with no limitations. He remains on guideline-directed medical therapy for heart failure and on azathioprine for management of CS.

DISCUSSION
Sarcoidosis is a multi-system granulomatous disorder that exhibits non-caseating granulomas in involved organs. This case presents a patient with CS whose CHB is reversed to normal AV conduction, nearly 1 year after initial diagnosis of CHB and 4 weeks after steroid therapy. Unfortunately, cardiac findings are discovered later in the disease even though CS often presents as an idiopathic AVB in young and middle-aged individuals.1 As recommended by the 2014 Heart Rhythm Society (HRS) Expert Consensus Statement, all patients younger than 60 years with unexplained Mobitz II or third-degree AVB should be screened for CS with high-resolution chest CT or advanced cardiac imaging with MRI or FDG-PET.2 In a study of 32 patients aged 18–60 years presenting with high-degree AVB with no history of sarcoidosis, 34% had evidence of CS on FDG-PET.1 Identifying CS as the aetiology of high-degree AVB is critical. Nordenswan et al remarkably showed that the 5-year incidence of sudden cardiac death (SCD) was 9% in patients with CS and AVB and normal EF and the 5-year incidence of SCD or VT was 24%.1

No prospective randomised trials have evaluated the benefit of corticosteroid therapy in patients with CS and CHB; however, small uncontrolled trials have shown the benefit of glucocorticoid therapy in this patient population. In a retrospective study of 41 patients, 63% had clinical manifestations and 22% had electrical findings of sarcoidosis. Nine patients had nodal, supraventricular or infraventricular block identified via an EP study, and 75% of the patients had improvement in conduction after steroid therapy. In another study evaluating the rate of undiagnosed CS and giant cell myocarditis, there was a tendency toward less pacing on follow-up at 1 year related to early start of immunosuppression therapy after pacemaker implantation. AVB was reversible in 2 (13%) of 16 patients after the institution of corticosteroid therapy. In a meta-analysis, 27 (47.4%) of 57 patients with CS presenting with AVB who received steroid therapy showed improvement in AV conduction, compared with recovery in none of the 16 patients who did not receive steroids.5 The 2014 HRS Expert Consensus Statement gave a class IIa recommendation stating that immunosuppression can be useful in patients with CS with second-degree (Mobitz II) or third-degree AVB.2 While corticosteroids can reverse high-degree AVB in patients with CS, the time course of resolution is not well defined. Our patient showed improvement from third degree to Mobitz II within 3 weeks and further improvement to sinus rhythm with AV conduction recovery in an additional 1 week.

In the case reported here, imaging revealed FDG uptake in the basal septum without evidence of LGE demarcating fibrosis on cardiac MRI. These findings suggest that this patient had ongoing granulomatous inflammation in the septum affecting AV conduction, but had not yet developed irreversible fibrosis in this region. Patients with fibrosis affecting the electrical
conduction system are unlikely to have AV conduction recovery with steroids, while patients with earlier disease, who have inflammation but no or minimal fibrosis, are more likely to have conduction system improvement with treatment. The timing of progression from inflammation to fibrosis in patients with CS is not known and may differ from patient to patient. However, early diagnosis and treatment are critical to maximize the chance of improvement with therapy. How long into the disease process AVB may reverse is not known. Our patient had AV recovery with steroid initiation 11 months after presentation with AVB. In some patients with CS, fibrosis formation may be delayed and a late response to steroids may be achieved. Our patient had suspected isolated CS based on imaging but did not have tissue biopsy confirming the diagnosis. Isolated CS can be particularly challenging to diagnose. In fact, the HRS Expert Consensus Statement requires endomyocardial biopsy to confirm isolated CS. The revised 2017 Japanese Ministry of Health and Welfare Diagnostic Guidelines allow for a diagnosis of isolated CS based on a combination of clinical and imaging findings without tissue diagnosis, and our patient had isolated CS based on these criteria. While the incidence of isolated CS is not known, with strict diagnostic criteria, it is likely quite rare. In one study using whole body FDG-PET to look for extra-cardiac inflammation, isolated CS was found in only 1 of 31 patients (3%).

CONCLUSION
In this case study, we reported a patient with suspected isolated CS who reverted from CHB to Mobitz II heart block to normal AV conduction 4 weeks after starting corticosteroids, 11 months after development of CHB. This case highlights the importance of screening young patients with AVB for CS and the possibility of AV conduction recovery if irreversible fibrosis has not developed.

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Patient’s perspective
I had been so tired for so long and could not quite understand why that was the case. I saw my primary care doctor a lot of times and had a lot of tests but nobody could explain why I was feeling the way I was. I did everything I was told to do. I used my CPAP machine but I was still tired. I tried to lose some weight but I did not feel better. I did a lot of other tests and everything always came back normal. I saw cardiologists and they could not find anything. Then, I started passing out. At first, I thought maybe I was working a lot or I was dehydrated but it happened again. That’s when I went to the hospital and they told me my heart rate was very slow. That is why they put the pacemaker in my chest. I initially felt a little better but was not back to normal. I had to follow-up with a cardiologist. He then did the MRI of my heart which showed that my heart was not functioning normally. That is when he sent me to the heart failure specialist for congestive heart failure.

I did a lot of other scans. The MRI and the PET scan showed that I probably have sarcoidosis so he sent me to the electrophysiologist and the rheumatologist. It required a lot of drives back and forth to the hospital because I had so many tests and so many appointments scheduled but it was all worth it. I was very scared because I was not sure of what was going to happen. Initially, like I said, I was not back to normal and then my device had to be changed because of ventricular tachycardia. Those were very uncertain and scary times. It was a disease I had never heard of before and have since read a lot about it. I am glad I made it through and did what I was advised to do. I definitely feel a lot better now.

What is interesting is that nobody in my family has this but I am glad they found it. I can work, walk and I don’t get as tired or short of breath as I did before. I can do more activities with my family. My heart is functioning better and I will continue to see all the doctors at your hospital. There are three I see there and one closer to my house because of how far I live. I tell my story all the doctors at your hospital. There are three I see there and if it helps even just one person, I am happy. This has all made a big difference and I am recovering well.

Learning points
► Isolated cardiac sarcoidosis (CS) is difficult to diagnose and often presents as conduction disease including complete heart block (CHB), or ventricular arrhythmia.
► The differential diagnosis for unexplained CHB in a young patient includes Lyme disease, congenital CHB, cardiac amyloid, Lamin A/C-mutation-related cardiomyopathy, giant cell myocarditis and CS.
► Early diagnosis and treatment of CS with corticosteroids, in the absence of myocardial fibrosis, could potentially revert conduction disease.
► In this case, multidisciplinary teamwork was crucial and necessary for accurate diagnosis and management.
Case report

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