Case Report

The Challenge of Diagnosing Heparin-Induced Thrombocytopenia with Negative Immunologic and Functional Assays

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Abstract

Heparin-Induced Thrombocytopenia (HIT) is a prothrombotic and potentially fatal immune complication of heparin therapy. HIT is challenging to diagnose, particularly in critically ill patients where multiple causes of thrombocytopenia must be considered. Diagnostic algorithms for HIT begin with a clinical assessment, followed by laboratory testing when indicated. If Platelet Factor-4 (PF4)/heparin immunoassay and Serotonin Release Assays (SRA) are negative, HIT is deemed unlikely and heparin therapy may be resumed. Current recommendations have excluded the next step in work up for thrombocytopenia after immunoassay and functional assays result negative despite worsening thrombocytopenia following heparin re-initiation. We present the case of an 85-year-old male with multiple comorbidities, found to have a clinical course consistent with HIT despite negative serologic and functional assay results. Our case highlights the challenge in diagnosing heparin-induced thrombocytopenia in a medically complex patient and demonstrates the need for standardized recommendations following negative laboratory results despite high clinical suspicion.

Keywords: Heparin-induced thrombocytopenia; Platelet Factor 4; Serotonin Release Assay; Immunoassay

Introduction

Heparin-Induced Thrombocytopenia (HIT) is a prothrombotic immune complication that occurs in up to 5% of patients treated with heparin therapy [1]. HIT arises when IgG antibodies target complexes of Platelet Factor 4 (PF4) and heparin, which subsequently induce widespread endothelial and platelet dysfunction [1]. HIT should be suspected in patients with declining platelet counts within 5-14 days of initiating Unfractionated (UFH) or Low-Molecular Weight Heparin (LMWH) [2]. When HIT is suspected, a clinical assessment such as the 4Ts scoring system is performed to determine the pretest probability [3]. If the pretest probability is intermediate or high, patients are switched to a non-heparin anticoagulant and an immunoassay is obtained [4]. Immunoassays are sensitive for detection of HIT and thus a negative result warrants heparin resumption [4]. However, false-negative results have been reported, especially early in the disease course. There is no standardized workup in patients with negative immunologic and functional assays in the context of high clinical suspicion. We present a patient with a clinical picture of heparin-induced thrombocytopenia despite negative serological results, highlighting the need to expand on current diagnostic guidelines.

Case Presentation

An 85 year-old male with a past medical history significant for transcatheater aortic valve replacement, congestive heart failure (CHF), chronic kidney disease, and two prior episodes of deep venous thrombosis presented to our hospital with a four day history of worsening lower extremity edema. He was admitted for CHF exacerbation complicated by cardiorenal syndrome and managed with bumetanide diuresis. Upon admission, his anticoagulation regimen was switched from his home dose of Apixaban (2.5mg twice daily) to a therapeutic heparin drip. His platelet count was 168. After 2 days of heparin therapy, his platelet count declined to 148. On day 6 of hospitalization, he was started on bridge to warfarin. On day 9 his BUN peaked at 146mg/dl, thus a peripherally inserted central catheter was placed for hemodialysis. On day 12, his platelet count also dropped to 84 from 118 the day prior. His 4Ts score was calculated to be 4 (+1 platelet count fall by 30-50%, +2 clear onset between 5-10 days after heparin, +1 for other possible causes of thrombocytopenia) indicating a moderate pretest probability for HIT at 14% [3]. Following the HIT 4Ts score guidelines, heparin was discontinued and switched to argatroban. PF4/heparin antibody and serotonin release assays were ordered.

PF4/heparin antibody results returned negative on day 15, measuring 0.37 (ref range <0.4 optical density). With a platelet count of 85, anticoagulation was switched back to a heparin drip with

Abbreviations

HIT: Heparin-Induced Thrombocytopenia; PF4: Platelet Factor 4; SRA: Serotonin Release Assay; LMWH: Low Molecular Weight Heparin; UFH: Unfractionated Heparin

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bridge to warfarin, as argatroban was not preferred in the setting of acute kidney injury. Due to the potential of ceftriaxone-mediated thrombocytopenia, cephalaxin was given on Day 16 for the last antibiotic dose treatment of UTI. Two days following resumption of heparin, his platelet count dropped to 60. During the next 5 days of heparin bridge to warfarin, the platelet count ranged from 60-81 (Figure 1). On day 22, the bridge was completed and heparin was discontinued. Over the next 5 days after heparin discontinuation, his platelet count rose to 117 and the patient was discharged. The serotonin release assay later returned negative. Alternative etiologies of his thrombocytopenia were explored, and further workup for HIV, Covid-19, and Hepatitis B/C were negative.

Discussion

Current diagnostic guidelines for heparin-induced thrombocytopenia recommend immunoassay in patients with a 4Ts score of 4 and above [4]. The Platelet Factor-4 (PF4)/heparin immunoassay is reported to have a sensitivity of >99%, with a specificity of 30-70% [5]. If immunoassay is positive, a functional assay such as the SRA is ordered for confirmation [4]. Due to interlaboratory variability and lack of standardization, sensitivity of the SRA is suboptimal at 62-100% with a specificity of 89-100% [5]. The SRA further requires specialized laboratory settings not widely available, and results can take up to 1-2 weeks [6].

Limited data is available to support further workup for HIT in the context of negative laboratory results. According to the American Society of Hematology 2018 guidelines, a negative immunoassay in patients with low or intermediate pretest probability excludes HIT and warrants resumption of heparin [4]. Follow-up with repeat immunoassay, a different immunoassay, or functional assay is recommended only in patients with high pretest probability [4]. If further follow-up remains negative, there are no further diagnostic recommendations available.

There are several factors that impact the reliability of testing for HIT including time to testing, transfusion-related dilution of serum, and technical limitations. False-negative PF4/heparin antibodies and SRA testing have been reported early in disease course, followed by positive results on repeat testing [7-12]. Another case report demonstrated negative enzyme immunoassay and SRA results following massive blood transfusion, believed to be caused by dilution of HIT antibodies and should be considered with fluid administration [11]. Chan et al analyzed the utility of repeat HIT immunoassay testing in patients with initially negative GTI-PF4 Enzyme-Linked Immunosorbent Assay (ELISA) [12]. The study found that 80% of patients with an initial high Optical Density (OD) reading of 0.268-0.399 had a positive ELISA on repeat testing [12]. Our patients OD reading of 0.37; unfortunately, repeat ELISA testing was not performed.

Our patient had several risk factors for HIT including unfractionated heparin use, duration of treatment, therapeutic dosing, and increased age [2,13]. This patient’s platelets rose from 63 to 85 after the heparin was initially stopped for 3 days but dropped to 60 following re-initiation of heparin. 4 days after completing the final heparin dose, the platelets rose to 128. No other cause of thrombocytopenia was established. Given the timing of heparin administration and subsequent platelet decline, our patient’s clinical picture was congruent with HIT. The negative PF4/heparin and SRA may have been due to low levels of antibodies early in the disease course or laboratory error. Repeat testing was not ordered, but should be considered when the clinical picture conflicts with laboratory results.

Conclusion

Diagnosis of heparin-induced thrombocytopenia poses many challenges. Diagnostic guidelines do not account for clinical variability, technical limitations, and lack of laboratory standardization. Our case represents the need to account for false negative results and delineate when repeat testing is recommended. By fulfilling this need, future patients may benefit from avoidance of unnecessary medical interventions and associated costs.
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