Fever of Unknown Origin: The Workup and Diagnosis of Pel-Ebstein Fever

Prachi Khanna 1, Natalie Malluru 1, Raaj Pyada 1, Mitul Gupta 1, Kartik Akkihal 1,2, Thomas C. Varkey 3,1

1. Internal Medicine, Dell Medical School, The University of Texas at Austin, Austin, USA 2. Kinesiology and Health Education, College of Education, University of Texas at Austin, Austin, USA 3. Business Management, Grand Canyon University, Phoenix, USA

Corresponding author: Prachi Khanna, prachikhanna@utexas.edu

Abstract

The clinical and diagnostic workup of fever of unknown origin (FUO) is key in the treatment of patients on the internal medicine service. In this article, the authors present a case of fever of unknown origin, walk through the differential diagnosis, explain the laboratory testing ordered in the workup of the patient as well as the resulting values of said testing, and discuss the pathophysiology and diagnostic criteria for the diagnosis of Pel-Ebstein fever. The authors also discuss a clinical pearl when working with electronic health records to ensure that the needs of the patient in question are met.

Categories: Internal Medicine, Oncology, Quality Improvement
Keywords: differential for fever of unknown origin, lymphoma, fever of unknown origin, hodgkin’s lymphoma, pel-ebstein

Introduction

Hodgkin’s lymphoma, a cancer of the lymphatic system, is characterized by genetic mutations leading to the proliferation of abnormally oversized lymphocytes [1,2]. Hodgkin’s is distinguished from non-Hodgkin’s by the presence of Reed-Sternberg cells, abnormal multinucleated lymphocytes [1,2]. Pel-Ebstein fever is a rare condition reported in patients with Hodgkin’s lymphoma, characterized by cyclic fevers that rise and fall every one or two weeks. Due to the rarity of disorder, many patients often suffer from lack of treatment or experience difficulties receiving sufficient care for their Pel-Ebstein fevers [1,2]. The authors provided a clinical case study, clinical workup, and discussed treatment for a patient suffering from Pel-Ebstein fevers secondary to Hodgkin’s lymphoma.

Case Presentation

The patient is a 53-year-old woman with a medical history of hypothyroidism, eczema, type II diabetes mellitus, and a reported history of non-Hodgkin’s lymphoma as indicated in the chart on previous hospital admission in March 2020, who presented with chief complaints of fever and a dry non-productive cough. She was sent to the emergency room directly from the infusion center where she was due to receive chemotherapy due to her low white blood cell count and increased temperature found at the clinic. The patient stated that for the last two and a half weeks she had been experiencing cough, intermittent fevers, diagnosed with pneumonia, and received treatment of a combination therapy consisting of 1 g of ceftriaxone twice daily and 750 mg of azithromycin once daily, for seven days at an outside hospital. The patient endorsed clear rhinorrhea but denied dysuria, diarrhea, shortness of breath, productive cough, constipation, or bleeding from any source. She denied any sick contacts stating that she lives alone but added that she does see her sister often. The patient reported that she had not yet received her coronavirus disease 2019 (COVID-19) vaccination due to her chemotherapy schedule. The patient had a past surgical history of cholecystectomy for gallstones, appendectomy for appendicitis, hysterectomy for uterine polyps, and port placement to facilitate chemotherapy infusions. The patient reported drug allergies to lisinopril, penicillin, and sulfa antibiotics which all resulted in whole body pruritus. The patient’s home medication regimen can be found in Table 1.
### Table 1: Patient’s home medication regimen.

| Medication   | Dosage and Frequency                                      | Indication                                                  |
|--------------|------------------------------------------------------------|-------------------------------------------------------------|
| Acyclovir    | 1200 mg taken twice daily by mouth                         | Viral illness prophylaxis                                   |
| Ondansetron  | 8 mg as needed every eight hours to be taken by mouth       | Chemotherapy associated nausea                              |
| Potassium chloride | 20 mEq taken twice daily by mouth          | To replace and maintain her potassium blood levels         |
| Naproxen     | 500 mg taken twice daily by mouth                         | For aches and pains                                         |
| Benzonatate  | 200 mg taken once daily by mouth                          | Cough suppressant                                           |
| Cyclobenzaprine | 5 mg by mouth as needed at bedtime                          | To help induce sleep                                        |
| Dupixent     | 300 mg by subcutaneous injection every two weeks           | To mitigate eczema symptoms                                 |
| Gabapentin   | 300 mg by mouth three times a day                          | For neuropathic pain symptoms                               |
| Glipizide    | 10 mg by mouth once a day                                  | For diabetes mellitus management                            |
| Hydroxyzine  | 10 mg by mouth at bedtime                                  | To help induce sleep                                        |
| Levothyroxine| 137 mcg by mouth in the morning                           | To manage her hypothyroid symptoms                          |
| Metformin    | 500 mg in the morning and 1000 mg at dinner by mouth        | To manage her diabetes mellitus                             |
| Guaifenesin  | 600 mg by mouth twice a day                                 | To manage mucus secretions                                  |

On physical examination, the patient was febrile to 100.7°F, but all other vital signs were within normal limits. She had some small strands of hair that were visible on the top of her head but was otherwise bald. There were no signs of erythema, swelling, induration, lymphadenopathy, or other signs of skin or tissue infection. Apart from her fever, the entirety of her physical examination was within normal limits.

### Differential diagnosis

The differential for fever of unknown origin (FUO) includes concurrent infection, malignancy, non-infectious inflammatory disease, as well as other lesser likely conditions such as drug-induced fever or factitious fever [3-9]. Please see Table 2 for an evidence-based differential of FUO. Infections are found to be the most common cause of FUO [3-9]. Bacterial and viral susceptibilities are potentially increased given the patient’s ongoing chemotherapeutic treatment [3-9]. While the patient’s reported history of non-Hodgkin’s lymphoma makes this diagnosis likely, other possible etiologies must be considered and eliminated before continuing immunocompromising treatment. Based on the patient’s medical history, care and attention should be drawn to her levothyroxine, a potential iatrogenic cause, which could lead to increased temperatures. However, without other signs and symptoms of systemic disease, this iatrogenic cause is considered less likely. Based on the patient’s objective data, it was unlikely that FUO was factitious or a secondary reaction to a drug.
**Subgroup** | **Cause**
---|---
**Infection (20-40%)** | Bacterial
| Abdominal or pelvic abscesses
| Dental abscesses
| Endocarditis
| Sinusitis
| Tuberculosis (especially extrapulmonary/disseminated)
| Urinary tract infection
| Viral
| Cytomegalovirus
| Epstein-Barr virus

**Malignancy (20-30%)**
| Colorectal cancer
| Leukemia
| Lymphoma (Hodgkin and non-Hodgkin)

**Non-infectious inflammatory disease (10-30%)**
| Connective tissue diseases
| Adult Still disease
| Rheumatoid arthritis
| Systemic lupus erythematosus
| Granulomatous disease
| Crohn disease
| Sarcoïdosis
| Vasculitis syndromes
| Giant cell arteritis
| Polymyalgia rheumatica/temporal arteritis

**Miscellaneous (10-20%)**
| Drug-induced
| Factitious fever
| Thromboembolic disease
| Thyroiditis

**TABLE 2: Common causes of fever of unknown origin.**

This information has been compiled from references [4-13] and is based on the currently available prevalence data.

**Laboratory testing**

Due to the patient’s recent history of chemotherapy treatment for cancer, the internal medicine team ordered several tests to rule out infection as the main cause of her fevers. The first set of tests included complete blood count (CBC) with differential, complete metabolic panel (CMP), chest x-ray, urinalysis with microscopy and culture, and three sets of blood cultures from two separate sites testing for bacterial, mycobacterial, and fungal infection. The bone marrow biopsy procedure was performed by the interventional radiology team. As per the pathologist, the patient’s biopsy was indicative of Hodgkin’s lymphoma - a surprise finding that led to an electronic medical record chart review. The pathologist sent her bone marrow for further evaluation of infection and found that there were no infectious materials. All laboratory results and their corresponding normal values can be found in Table 3.
Laboratory/Imaging Test | Subtest | Result | Normal Range
---|---|---|---
**Complete blood count** | White blood cell count | 2.2 thousand/mm³ | 4.5-11.0 thousand/mm³
| Hemoglobin | 8.6 g/dL | 12-14 g/dL
| Differential - lymphocytes | 6% of all white blood cells | 30-40% of all white blood cells
| Differential - neutrophils | 52.6% of all white blood cells | 50-65% of all white blood cells

**Blood culture** | N/A | No growth | No growth

**Complete metabolic panel** | Sodium | 127 mEq/L | 135-145 mEq/L

**Chest x-ray** | N/A | Normal chest anatomy | Normal chest anatomy

**Urinalysis** | Gross urinalysis | Normal light-yellow coloration | Normal light-yellow coloration
| Dipstick | Negative | Negative
| Microscopy | Negative | Negative
| Urine culture | No growth | No growth
| Spot urine sodium | 83 mmol/L | |
| Spot urine osmolality | 712 mOsm/kg | |

**Karius polymerase chain reaction (PCR) test** | N/A | Negative | Negative

**Bone marrow biopsy** | N/A | Hodgkin’s lymphoma | Normal hematopoietic stem cells

### TABLE 3: Pertinent laboratory and imaging results.

**Diagnosis and treatment**

Throughout the workup of her fevers, the patient had episodes where her fevers spiked to 104°F while her systolic blood pressure dropped into 90s and her diastolic blood pressure dropped into 50s. These factors necessitated transfer to the medical intensive care unit (MICU). Nevertheless, within hours the fever broke and her blood pressure normalized. These fever and shock episodes happened cyclically, with episodes happening every two to three days, raising the suspicion of a non-infectious etiology [4-13]. To ensure the patient would not be treated prematurely with immunocompromising drugs, the internal medicine team and the infectious disease team stated that it would be wise to await the results of the Karius polymerase chain reaction (PCR) test.

While waiting eight days for the Karius PCR test results, the team began treatment of the patient’s hyponatremia. The low salt was deemed to be the result of a syndrome of inappropriate antidiuretic hormone (SIADH) caused by her Hodgkin’s lymphoma. She was treated with 5 mg midodrine by mouth three times a day, 1.2 L fluid restriction daily, and 1000 mg sodium chloride tablets twice daily to bring her sodium levels within normal range.

Based on the patient’s medical history, negative laboratory results, and lack of infectious agent, the most likely etiology of her fevers was determined to be Pel-Ebstein fevers. Because of the lack of infection, the oncology team stated that they felt comfortable in moving forward with providing the patient with chemotherapy for her Hodgkin’s lymphoma. They started her on an intravenous infusion regimen of gemcitabine (1 g/m²) and oxaliplatin (75 mg/m²) for her Hodgkin’s lymphoma. This second-line treatment regimen was chosen because the patient had failed earlier treatment with the preferred first-line treatment of adriamycin, bleomycin, vinblastine, and dacarbazine.

The patient remained in the hospital for two additional days, after completing her chemotherapy regimen, to be monitored for unusual reactions to her chemotherapy. She was discharged when her absolute neutrophil count (ANC) was above 500 cells/mm³ and she was fever-free for 24 hours. Her total stay, including the workup, fever-shock episodes, chemotherapy, and recovery, was 14 days from initial admission.

**Discussion**

FUO is defined as the temperature of 38.3°C (100.9°F) or higher, which occurs for greater than three weeks with no known cause after investigation [3]. The most common etiologies are infection, malignancies, non-

---

2022 Khanna et al. Cureus 14(2): e21959. DOI 10.7759/cureus.21959

4 of 6
infectious inflammatory disease, and other miscellaneous causes [3]. The diagnostic workup for FUO centers around searching for potential diagnostic clues (PDCs), which can guide the team towards a potential etiology and treatment plan. The initial approach involves taking comprehensive history, physical examination, and appropriate lab testing [3-9]. The preliminary lab investigations should include complete blood count, liver function tests, erythrocyte sedimentation rate, urinalysis, and basic cultures. Specific techniques and imaging should be based on the PDCs, whereas more invasive techniques (i.e., lumbar puncture or bone marrow biopsy) should only be performed when there is a specific need or if an extensive investigation has failed to reveal a diagnosis [3-9].

While the patient’s complete history, physical examination, and laboratory results failed to yield an infectious agent, the bone marrow biopsy indicated that the patient had Hodgkin’s lymphoma. Therefore, the most likely etiology of her fevers was determined to be Pel-Ebstein fever, which is a rare but documented phenomenon associated with Hodgkin’s lymphoma [14-18]. Some estimates state that roughly 35% of those afflicted with Hodgkin’s lymphoma experience this phenomenon [14-18]. In this condition, patients experience cycles of high temperature that typically last one-to-two weeks, followed by febrile periods of similar duration. It is not unusual for patients with both Hodgkin’s and non-Hodgkin’s lymphoma to present with elevated daily temperatures, as it is theorized that in certain malignancies, such as lymphomas, there is a release of pyrogenic cytokines from tumor cells or macrophages, such as interleukin-1 (IL-1), interleukin-6 (IL-6), and tumor necrosis factor-alpha (TNF-α). These cytokines affect the thermostatic set point by inducing prostaglandin E2, which directly acts on the hypothalamus [17]. However, the exact mechanism behind the cyclic nature of Pel-Ebstein fever is still unknown and the phenomenon itself and its association with Hodgkin’s lymphoma is widely debated [18]. While more data and inquisitorial work must be done on the topic, FUO’s relatively consistent association with Hodgkin’s lymphoma leads the team to believe that Pel-Ebstein fevers are a true phenomenon that can therefore be treated.

One of the key pieces that became increasingly salient to the case was the patient’s misdiagnosis in the chart of “non-Hodgkin’s lymphoma.” When looking through the notes later, it was determined that the initial mistake had been entered into her chart during one of her previous admissions to the hospital. Following the biopsy which demonstrated Hodgkin’s lymphoma and the revelation of the error within the electronic medical record, the record was amended to ensure that the patient would be able to be treated appropriately and that the medical record was now clear as to her specific type of malignant disease. Herein, the authorial team wanted to stress the importance of verifying the information with the patient and with previous records to ensure accuracy. Nevertheless, human errors can (and often do) lead to the entry of “chart lore” [19,20]. Therefore, it is key for clinicians to be on the proverbial “lookout” for these errors and be ready to make amendments to ensure accuracy in the record [19,20].

Conclusions
This case study discussed the care and management of a patient with a fever of unknown origin, complicated by an error in the patient’s chart history which misreported the patient’s diagnosis and treatment of Hodgkin’s lymphoma as non-Hodgkin’s lymphoma. The patient was managed conservatively and thoroughly tested until the most likely cause of her symptoms was found to be Pel-Ebstein fever associated with Hodgkin’s lymphoma. A key clinical practice takeaway from this case is the importance of verifying the accuracy of patient charts with patients and previous records.

Additional Information
Disclosures
Human subjects: Consent was obtained or waived by all participants in this study. Conflicts of interest: In compliance with the ICMJE uniform disclosure form, all authors declare the following: Payment/services info: All authors have declared that no financial support was received from any organization for the submitted work. Financial relationships: All authors have declared that they have no financial relationships at present or within the previous three years with any organizations that might have an interest in the submitted work. Other relationships: Thomas is an Adjunct Professor at Grand Canyon University and receives payment for his teaching and grading, he is a faculty member with the National Multiple Sclerosis Society’s Monthly Fellows Difficult Case Discussion Webinar, and Thomas serves on the board of editors for ProClinS Cardiology.

Acknowledgements
Thomas C. Varkey and Kartik Akkihal both provided large contributions to the project and should be considered co-senior authors for this paper.

References
1. Racchi O, Raperzi D, Ferraris AM, Gaetani GF: Unusual bone marrow relapse of Hodgkin’s disease with typical Pel-Ebstein fever. Ann Hematol. 1996, 73:39-40. 10.1007/s002770050196
2. Reimann HA: Periodic (Pel-Ebstein) fever of lymphomas. Ann Clin Lab Sci. 1997, 7:1-5.
3. Bleeker-Rovers CP, van der Meer JW, Oyen WJ: Fever of unknown origin. Semin Nucl Med. 2009, 39:81-7.
4. Roth AR, Basello GM: Approach to the adult patient with fever of unknown origin. Am Fam Physician. 2003, 68:2223-8.
5. Hersch EC, Oh RC: Prolonged febrile illness and fever of unknown origin in adults. Am Fam Physician. 2014, 90:91-6.
6. Bleeker-Rovers CP, Vos FJ, de Kleijn EM, et al.: A prospective multicenter study on fever of unknown origin: the yield of a structured diagnostic protocol. Medicine (Baltimore). 2007, 86:26-38. 10.1097/MD.0b013e31802fe858
7. Varghese GM, Trowbridge P, Doherty T: Investigating and managing pyrexia of unknown origin in adults. BMJ. 2010, 341:i54:i5470
8. Hayakawa K, Ramasamy B, Chandrasekar PH: Fever of unknown origin: an evidence-based review. Am J Med Sci. 2012, 344:307-16. 10.1097/MAJ.0b013e31824ae504
9. Cunha BA: Fever of unknown origin: clinical overview of classic and current concepts. Infect Dis Clin North Am. 2007, 21:867-915. 10.1016/j.idc.2007.09.002
10. de Kleijn EM, Vandenbroucke JP, van der Meer JW: Fever of unknown origin (FUO). I. A prospective multicenter study of 167 patients with FUO, using fixed epidemiologic entry criteria. The Netherlands FUO Study Group. Medicine (Baltimore). 1997, 76:592-400. 10.1097/00005792-199711000-00002
11. Knockaert DC, Vandercruysen S, Blockmans D: Fever of unknown origin in adults: 40 years on. J Intern Med. 2005, 258:593-7. 10.1046/j.1365-2796.2005.01120.x
12. Kim SE, Kim UJ, Jang MO, et al.: Diagnostic use of serum ferritin levels to differentiate infectious and noninfectious diseases in patients with fever of unknown origin. Dis Markers. 2013, 34:211-8. 10.3233/DMA-130962
13. Zenone T: Fever of unknown origin in adults: evaluation of 144 cases in a non-university hospital. Scand J Infect Dis. 2006, 38:632-8. 10.1080/00365540600666564
14. Chung WJ, Howard MR: Pel-Ebstein fever with cyclical pancytopenia. J R Soc Med. 2001, 94:93-5. 10.1177/014107680109400210
15. Wilson AJ: Pel-Ebstein fever. N Engl J Med. 1995, 333:666-7. 10.1056/NEJM199507063330118
16. Talbot TR: Cases from the Osler Medical Service at Johns Hopkins University. Hodgkin’s disease with Pel-Ebstein fevers. Am J Med. 2002, 112:512-5. 10.1016/s0002-9545(02)01041-0
17. Fogg V, Cavenagh J: Malignant causes of fever of unknown origin. Clin Med (Lond). 2015, 15:292-4. 10.7861/clinmedicine.15-3-292
18. Schattner A, Krashet N: Pel-Ebstein cyclic fever: not just lymphoma. Am J Med. 2010, 123:10.1016/j.amjmed.2009.12.050
19. Bell SK, Delbanco T, Elmore JG, et al.: Frequency and types of patient-reported errors in electronic health record ambulatory care notes. JAMA Netw Open. 2020, 3:10.1001/jamanetworkopen.2020.5867
20. Gillum RF: From papyrus to the electronic tablet: a brief history of the clinical medical record with lessons for the digital age. Am J Med. 2013, 126:853-7. 10.1016/j.amjmed.2013.05.024