Steroid-Induced Tumor Lysis Syndrome Accompanied by Diabetic Ketoacidosis and Acute Renal Failure in a Non-Hodgkin Lymphoma Patient

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Abstract

We present a case of a 66-year-old male with a past history of newly diagnosed non-Hodgkin lymphoma, diabetes, and recent surgical splenectomy secondary to splenic infarct who presented to the Emergency Department (ED) with several nonspecific symptoms that were consistent with tumor lysis syndrome. This case report discusses the clinical presentation, diagnosis, and management of spontaneous tumor lysis syndrome.

Introduction

Tumor lysis syndrome (TLS) is a rare emergent syndrome that results in the release of intracellular contents from cell lysis that occurs quickly. TLS is often triggered by chemotherapy but may also occur due to radiation, steroids, or spontaneously. Typically, tumor lysis occurs in patients with tumors with a significant proliferation rate, tumor burden, or sensitivity to cytotoxic therapy. TLS is commonly seen in those with hematological neoplasms such as acute lymphoblastic leukemia (ALL) and acute myeloid leukemia (AML). However, lysis may also happen in those with high-grade lymphomas such as Burkitt’s Lymphomas and other solid tumors [1, 2]. Intermediate grade non-Hodgkin lymphoma (NHL), such as diffuse large B cell lymphoma, can also be at high or intermediate risk for tumor lysis. Comparatively, slower dividing low-grade lymphoma or Hodgkin’s lymphoma are at lower risk of tumor lysis. Currently, there are only sparse case reports of TLS in low-grade and intermediate-grade NHL, Hodgkin’s disease, AML and ALL in blast crisis, and myeloproliferative disorders [3, 4].

Spontaneous tumor lysis syndrome (STLS) is an oncologic emergency initiated by releasing copious amounts of potassium, phosphate, nucleic acids, and uric acid into systemic circulation by extensive tumor breakdown. Lysis in STLS occurs without any antineoplastic agents or interventions such as chemotherapies [5]. As the tumors are catabolized, the tumor cells release electrolytes and nucleic acids into circulation. The nucleic acids are further broken down into uric acids, resulting in hyperuricemia. Uric acid can collect in renal tubules and cause an intrarenal obstruction due to the acidity [2]. The excessive level of these products may result in severe consequences such as acute renal injury, arrhythmias, altered mental status, seizures, and potentially death [1].

TLS can also be triggered by steroid treatment. There are a few cases in the literature documenting steroid-induced tumor lysis syndrome in lymphoma, causing acute renal failure [6, 7] and metabolic acidosis [8, 9]. However, cases that report steroid-induced TLS from lymphoma causing diabetic ketoacidosis (DKA) and acute renal failure are lacking.

Early recognition of metabolic derangements, rapid treatment, and early coordination of care with the oncology, nephrology, and intensive care team is critical for the care of a potentially lethal syndrome. Even in those with less prone to malignancies, including TLS as a differential diagnosis is imperative. Therefore, it is crucial for the emergency medicine physician to be aware of this life-threatening metabolic emergency and its associated complications and how to manage them.

Case Presentation

The patient is a 66-year-old male who presents to the ED for shortness of breath, nausea, generalized abdominal pain, bone pain, decreased appetite, oral intake, and decreased urine output for the past three days. He has a past medical history of diabetes, non-Hodgkin lymphoma that was newly diagnosed four weeks before presentation, and splenectomy secondary to splenic infarct of unclear etiology three months
prior to presentation. A month prior, the patient was seen for cervical lymphadenopathy and was subsequently diagnosed with non-Hodgkin lymphoma after cervical lymph node biopsy. The patient reported subjective fever and unintentional weight loss of approximately 30 pounds over the past three months. He denies coughing, vomiting, chest pain, or diarrhea. He endorses insomnia due to the symptoms for the past three days. During the interview with the patient, he stated that he had not started his chemotherapy for the lymphoma. He was instructed to take prednisone for 10 days but stop his metformin and insulin PM three days ago for mediport insertion today. However, this procedure was canceled due to his symptoms. On exam, the patient is ill-appearing with dry mucous membrane and epigastric and mid-abdominal tenderness. The patient was tachycardic, but otherwise, all other vital signs were stable. He was afebrile.

On initial workup, laboratory studies revealed that the patient was mildly hyponatremic, hyperkalemic, hypochloremic, with an elevated anion gap, elevated blood urea nitrogen (BUN) and creatinine, hypercalcemia, hyperphosphatemia, elevated aspartate aminotransferase (AST) and hypoalbuminemia, elevated beta-hydroxybutyrate, and hyperglycemia (Table 1).

| Arterial blood gas (left radial artery) |  |
|----------------------------------------|---|
| pH                                     | 7.35 - 7.45 | 7.44 |
| pCO2                                   | 35 - 48 mmHg | 19 L |
| pO2                                    | 83 - 108 mmHg | 87.1 |
| HCO3                                   | 21 - 28 mmol/L | 13.1 L |
| O2 Sat                                 | 94 - 98 % | 97.3 |
| Base Excess                            | -2 - 3 mmol/L | -8.9 L |
| Allen Test                             | Positive |
| Chemistry                              |  |
| Sodium                                 | (136 - 145 mmol/L) | 128 L |
| Potassium                              | (3.7 - 5.1 mmol/l) | 6.9 C |
| Chloride                               | (98 - 107 mmol/L) | 93 L |
| Carbon Dioxide                         | (21 - 32 mmol/l) | 11 L |
| Anion Gap                              | 30.9 |
| Blood Urea Nitrogen                    | (7 - 18 mg/dl) | 72 H |
| Creatinine                             | (0.55 – 1.3 mg/dl) | 2.34 H |
| Glucose                                | 74 - 106 mg/dl | 282 H |
| Calcium                                | 8.4 - 10.1 mg/dl | 8.5 |
| Phosphorus                             | 2.5 - 4.9 mg/dL | 7.6 H |
| Total Bilirubin                        | 0.2 - 1.5 mg/dl | 0.9 |
| Aspartate Aminotransferase             | 10 - 37 unit/L | 99 H |
| Alanine Aminotransferase               | 12 - 78 unit/L | 26 |
| Alkaline Phosphatase                   | 45 - 117 unit/L | 139 H |
| Creatinine Kinase                      | 35 - 232 unit/L | 111 |
| Troponin I                             | <0.05 ng/mL | 0.05 H |
| Total Protein                          | 6.4 - 8.2 g/dL | 6.5 |
| Albumin                                | 3.4 - 5.0 g/dl | 2.1 L |
| BHB                                    | 0.2 - 2.8 mg/dL | 21.4 H |
| Glucose                                | 74 - 106 mg/dL | 291 H |
| Lactic Acid                            | 0.4 - 2.0 mmol/L | 9.7 H |
**Complete blood count demonstrated leukocytosis and thrombocytopenia. Blood peripheral smear revealed immature granulocytes consisting of metamyelocytes, promyelocytes, and myelocytes. Additional laboratory studies also showed lactic Acidosis, hyperuricemia, and elevated lactate dehydrogenase. Arterial blood gas revealed a pH of 7.44, pCO2 of 19, pO2 of 87, and bicarbonate of 13. His COVID-19 test was negative. Presumptive diagnoses at this point were diabetic ketoacidosis due to steroids, tumor lysis syndrome complicated by acute renal failure and hyperkalemia, and sepsis. The patient was started on intravenous fluids for acute renal failure. The patient was then started on calcium gluconate, insulin infusion, and D5-NS to treat hyperkalemia and Acidosis. Ceftriaxone was initiated for presumptive sepsis. The patient was then admitted to the intensive care unit (ICU) for further management and oncology consultation, as his presentation was suggestive of STLS.**

**During hospital day one, the patient developed worsening hypocalcemia and Acidosis. It was determined that he would need to be stabilized prior to initiating mediport insertion and treatment of his non-Hodgkin lymphoma. On hospital day two, the patient became anuric, confused, and lethargic. Laboratory studies revealed improved hyperkalemia, a persistently high anion gap, and Acidosis despite insulin drip and IV...**
fluid. However, he then began to develop pulmonary edema. Nephrology was consulted, and the patient was started on hemodialysis due to worsening renal function and volume overload. Rasburicase was also started to manage the elevated uric acid level. On hospital day three, the patient had persistent worsening renal function, anuria, metabolic Acidosis, and volume overload despite hemodialysis. He later became hypotensive, requiring vasopressor support in the ICU. The patient had newly developed atrial fibrillation with rapid ventricular response. On hospital day four, due to his poor prognosis, the patient was made “do-not-resuscitate” and discharged to hospice for comfort care.

Discussion

TLS typically occurs after treatment of malignancy but may occur commonly after steroid treatment or spontaneously in rare cases prior to treatment. Non-Hodgkin lymphomas are a heterogeneous group of solid tumors. Caveats for lysis for solid tumors include bulky solid tumors or high sensitivity to cytotoxic therapies. Through a literature review, STLS in solid tumors is seen in a few cases, as demonstrated by its low likelihood for this syndrome [10]. Most cases of lysis seen in solid tumors were due to the initiation of radiation, chemotherapy, or steroids. Our patient had undergone a 10-day course of steroids, which is most likely the potential culprit as he has not started his chemotherapy. Current guidelines and literature reveal that most solid tumors are at low risk for spontaneous lysis. The occurrence of both TLS and STLS is typically rare for those with NHL, but certain features can make NHL at high risk for TLS. Certain tumor types and burden make NHL particularly high risk for TLS, such as advanced Burkitt’s lymphoma, early Burkitt’s lymphoma with high LDH, and diffuse large B-cell lymphoma with high LDH [11]. Pre-existing conditions such as renal dysfunction can also make an NHL patient at high risk for TLS. We do not know this patient’s NHL subtype, its tumor stage, disease burden, or baseline LDH, so it is difficult to risk stratifying this patient for TLS based on these tumor-specific features. However, we do know that this patient is at high risk for renal dysfunction given his diabetes history despite normal baseline creatinine. His steroid use likely exacerbated his hyperglycemia, subsequently leading to DKA and triggering TLS at the same time. The combination of these two conditions likely worsened his renal function, resulting in his renal failure and ultimate poor outcome.

STLS requires clinical correlation with the patient’s known malignancy, the likelihood for lysis, and laboratory studies. Current guidelines reflect that of the Cario-Bishop guidelines [12,13]. If the laboratory studies reveal at least two abnormalities, the patient may be diagnosed with TLS. Abnormalities noted by current Cario-Bishop guidelines classify patients with TLS or STLS have a serum uric acid level ≥8 mg/dL, or a 25% increase from baseline; a serum potassium level ≥7 mmol/L, or a 25% increase from baseline; serum phosphate levels ≥4.5 mg/dL in adults, or a 25% decrease from baseline; and a serum calcium level ≤7 mg/dL, or a 25% decrease from baseline [13-14]. In addition, these abnormalities must occur prior to or within seven days following the beginning of chemotherapy treatment to be considered for a diagnosis of TLS [12,13]. However, as this patient has not started any cytotoxic therapy, the evidence is clear that the patient was experiencing steroid-induced TLS given recent steroid use.

Aggressive IV fluid administration is the key to preventing TLS. The aim is to optimize renal perfusion with a target urine output of 2 ml/kg [15]. Adequate glomerular filtration and urinary output (UOP) are important in preventing precipitations of uric acid crystals or calcium phosphate deposits in the renal tubules, one of the main mechanisms behind acute renal failure in TLS patients. However, in vitro fertilization (IVF) administration also puts patients at risk for volume overload in the setting of preexisting acute kidney injury. In this case, reversing the underlying cause of acute kidney injury (AKI) should also be addressed. Our patient presented with TLS and AKI that gradually progressed to acute renal failure. IVF administration ultimately contributed to pulmonary edema in this patient due to poor renal function. Dialysis then had to be initiated due to volume overload, persistent electrolytes abnormalities, and acidosis. Diuretics such as furosemide could be a feasible therapeutic option for volume overload, but it is unlikely to work in patients that are already in renal failure and are anuric.

Uric acid-lowering agents, such as allopurinol (xanthine oxidase inhibitor) and rasburicase (urate oxidase), are additional key management strategies for preventing TLS. Allopurinol prevents the new uric acid formation and thus has the detrimental potential for obstructive AKI. However, allopurinol does not reduce pre-existing high uric acid levels, as seen in our patient. Therefore, it was not the best option in this case. In addition, allopurinol can also precipitate xanthine crystals in the renal tubules, worsening renal function. In patients at high risk for TLS with pre-existing high uric acid levels and those with impaired renal function, rasburicase is recommended. Rasburicase lowers the serum uric acid level by converting it to a more water-soluble compound, allantoin [15]. Two prospective trials have demonstrated the efficacy of rasburicase at reducing serum uric acid levels in patients at high risk for TLS [16-17]. Rasburicase should be started within the first 24 hours if possible [10,18]. In our patient, his TLS was further complicated by acute renal failure and DKA, making clinical management much more difficult. His clinical deterioration was progressing too quickly for rasburicase to have any meaningful clinical effect.

ED management includes IV hydration and stabilization of metabolic derangements. As the patient will have a variety of laboratory abnormalities, correction of elevated potassium, creatinine, and uric acid levels take precedence. Hyperkalemia is known to cause arrhythmias, as seen by the patient developing new-onset atrial fibrillation during his hospitalization. Uric acid levels further worsen the patient’s acute renal failure.
Acidosis may also be present. Nephrology consultation for hemodialysis may be required if renal function and acidosis do not improve. Oncology should also be consulted for further guidance, although limitations from TLS will further hold off treating the malignancy until the patient is stabilized.

Despite limited findings in the literature, research reveals that the prognosis of STLS and TLS is typically viewed as poor and remains guarded. Current research findings reveal that AKI is the strongest predictor of short-term and long-term mortality despite potentially confounding variables such as the malignancy itself and other pre-existing comorbidities [17]. Pre-existing conditions such as diabetes pose an additional layer of complexity in managing STLS and TLS. Cancer patients with diabetes tend to have higher mortality [20]. Patients with any malignancy and who are experiencing DKA will have poorer survival chances due to the sequela from the electrolyte imbalances during their hospitalization.

Conclusions
TLS is a condition that requires clinical acumen and immediate action. Although STLS is rare in those with non-Hodgkin lymphoma, this syndrome should still be considered as a differential for an acute oncologic emergency. Clinical correlation of laboratory studies with a thorough history is essential in elucidating the patient’s status and overall disposition. IV fluid administration, correcting electrolytes abnormalities, and lowering serum uric acid levels should be the cornerstone of TLS therapy. Dialysis may be required if electrolyte abnormalities, acidosis, volume overload, and uremia do not resolve with medical management. Oncology, nephrology, and intensive care specialists should be engaged early to assist with the management of TLS patients.

Additional Information
Disclosures
Human subjects: Consent was obtained or waived by all participants in this study. HCA Centralized Algorithms for Research Rules on IRB Exemptions (CARRIE) issued approval 2022-003. HCA Centralized Algorithms for Research Rules on IRB Exemptions (CARRIE)/IRB manager issued exemption 2022-005. Based on the information provided and attested as true, the research plan described does not require IRB oversight. This is because the investigators are either a) not engaging in research with human subjects as defined by federal regulations; b) engaging in research with human subjects deemed excluded from IRB oversight per 45CFR46.102(f) OR c) engaging in research with sufficient human subject protections in the design to meet one or more IRB exemption criteria set forth in 45CFR46.104. Conflicts of interest: In compliance with the ICMJE uniform disclosure form, all authors declare the following: Payment/services info: All authors have declared that there are no other relationships or activities that could appear to have influenced 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: All authors have declared that there are no other relationships or activities that could appear to have influenced the submitted work.

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References
1. Adeyinka A, Bashir K: Tumor Lysis Syndrome. 2022.
2. Spring I, Munshi L: Oncologic emergencies: traditional and contemporary. Crit Care Clin. 2021, 37:85-103. 10.1016/j.ccc.2020.08.004
3. Kekre N, Djordjevic B, Touchie C: Spontaneous tumour lysis syndrome. CMAJ. 2012, 184:913-6. 10.1503/cmaj.111251
4. Cairo MS, Coffler B, Reiter A, Younes A: Recommendations for the evaluation of risk and prophylaxis of tumour lysis syndrome (TLS) in adults and children with malignant diseases: an expert TLS panel consensus. Br J Haematol. 2010, 149:578-86. 10.1111/j.1365-2141.2010.08143.x
5. Opyrchal M, Figanbaum T, Ghosh A, Rajkumar V, Caples S: Spontaneous tumor lysis syndrome in the setting of B-cell lymphoma. Case Rep Med. 2010, 2010:619069. 10.1155/2010/619069
6. Lerza R, Botta M, Barsotti B, et al.: Dexamethasone-induced acute tumor lysis syndrome in a T-cell malignant lymphoma. Leuk Lymphoma. 2002, 43:1129-32. 10.1080/10428190290021452
7. Yang SS, Chau T, Dai MS, Lin SH: Steroid-induced tumor lysis syndrome in a patient with preleukemia. Clin Nephrol. 2005, 59:201-5. 5.10.5414/cnp59201
8. Griffin D, Myadam R, Patel P: Steroid-induced lactic acidosis in diffuse large B-cell lymphoma. Cureus. 2020, 12:e7446. 10.7759/cureus.7446
9. Malik IA, Abubakar S, Alam F, Khan A: Dexamethasone-induced tumor lysis syndrome in high-grade non-Hodgkin’s lymphoma. South Med J. 1994, 87:409-11. 10.1097/00007611-199403000-00024
10. Yaman S, Baço S, Turan G, et al.: Single-dose rasburicase might be adequate to overcome tumor lysis syndrome in hematological malignancies. Clin Lymphoma Myeloma Leuk. 2022, 22:271-6. 10.1016/j.clml.2021.08.009
11. Belay Y, Yirdaw K, Enawgaw B: Tumor lysis syndrome in patients with hematological malignancies. J Oncol. 2017, 2017:9684909. 10.1155/2017/9684909
12. Mirrakhimov AE, Voore P, Khan M, Ali AM: Tumor lysis syndrome: a clinical review. World J Crit Care Med. 2015, 4:130-8. 10.5492/wjccm.v4.i2.130
13. Cairo MS, Bishop M: Tumour lysis syndrome: new therapeutic strategies and classification. Br J Haematol. 2004, 127:3-11. 10.1111/j.1365-2141.2004.05094.x
14. Hande KR, Garrow GC: Acute tumor lysis syndrome in patients with high-grade non-Hodgkin’s lymphoma. Am J Med. 1995, 94:155-9. 10.1016/0002-9343(95)90174-n
15. Coiffier B, Altman A, Pui CH, Younes A, Cairo MS: Guidelines for the management of pediatric and adult tumor lysis syndrome: an evidence-based review. J Clin Oncol. 2008, 26:2767-78. 10.1200/JCO.2007.15.0177
16. Coiffier B, Mounier N, Bologna S, et al.: Efficacy and safety of rasburicase (recombinant urate oxidase) for the prevention and treatment of hyperuricemia during induction chemotherapy of aggressive non-Hodgkin’s lymphoma: results of the GRAAL1 (Groupe d’Etude des Lymphomes de l’Adulte Trial on Rasburicase Activity in Adult Lymphoma) study. J Clin Oncol. 2005, 23:4402-6. 10.1200/JCO.2005.04.115
17. Cortes J, Moore JO, Maziarz RT, et al.: Control of plasma uric acid in adults at risk for tumor Lysis syndrome: efficacy and safety of rasburicase alone and rasburicase followed by allopurinol compared with allopurinol alone—results of a multicenter phase III study. J Clin Oncol. 2010, 28:4207-15. 10.1200/JCO.2009.26.8896
18. Rahmani B, Patel S, Seyam O, Gandhi J, Reid I, Smith N, Khan SA: Current understanding of tumor lysis syndrome. Hematol Oncol. 2019, 37:537-47. 10.1002/hon.2668
19. Matuszkiewicz-Rowinska J, Malyszko J: Prevention and treatment of tumor lysis syndrome in the era of onco-nephrology progress. Kidney Blood Press Res. 2020, 45:645-60. 10.1159/000509934
20. Shahid RK, Ahmed S, Le D, Yadav S: Diabetes and cancer: risk, challenges, management and outcomes. Cancers (Basel). 2021, 13:10.3390/cancers13122735