Diabetic ketoacidosis caused by fulminant type 1 diabetes during adjuvant chemotherapy for colon cancer: A case report

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Abstract. Development of diabetic ketoacidosis (DKA) caused by fulminant type 1 diabetes (FT1D) during administration of uracil-tegafur (UFT) with leucovorin (LV) as adjuvant chemotherapy is extremely rare. Here, we report a case of DKA caused by FT1D during administration of UFT with LV as adjuvant chemotherapy for colon cancer. A woman in her 60s was transferred to the emergency medical center of our hospital with complaints of impaired consciousness and vomiting. She had undergone left hemicolectomy and D3 lymph node dissection for transverse colon cancer 8 months earlier. She was provided UFT with LV as adjuvant chemotherapy. Laboratory analysis revealed hyperglycemia, high anion gap metabolic acidosis and urinary ketones. She was diagnosed with DKA and was started on intravenous infusion of fluid and continuous subcutaneous insulin injections. Following admission, she was examined and diagnosed with FT1D. The present case describes an extremely rare case of DKA caused by FT1D during adjuvant chemotherapy with UFT + LV for colon cancer.

Introduction

Most colorectal cancers are adenocarcinomas. The primary treatment method is surgery, which may result in cure depending on the disease stage at diagnosis. We usually perform adjuvant chemotherapy for high risk of recurrence cases. The regimens of adjuvant chemotherapy are various such as 5-fluorouracil (5-FU) + leucovorin (LV), oral uracil-tegafur (UFT) + LV, oral capecitabine, and FOLFOX (oxaliplatin, 5-fluorouracil, and leucovorin). UFT often causes neutropenia, stomatitis, hand-foot syndrome, and alopecia; however, fulminant type 1 diabetes (FT1D) is a rare complication.

Diabetic ketoacidosis (DKA) is a life-threatening complication of type 1 diabetes (T1D), which leads to death if not treated. FT1D is a subtype of T1D characterized by noticeably rapid and almost complete destruction of pancreatic B cells, with acute onset leading to severe metabolic disorders. Several studies have reported that interferon alpha or immune checkpoint inhibitors induce FT1D; however, its mechanism of development remains unclear. Here, we report a case of DKA caused by FT1D during administration of UFT with LV as adjuvant chemotherapy for colon cancer.

Case report

A woman in her 60s was transferred to the emergency medical center of our hospital with complaints of impaired consciousness and vomiting. Eight months earlier, she underwent left hemicolectomy and D3 lymph node dissection for transverse colon cancer with perforation. The final pathological diagnosis was Stage II colon cancer (TNM classification: T4bN0M0). The patient had been taking adjuvant chemotherapy with UFT + LV; the dose for UFT and LV was 500 and 75 mg/day, respectively. The patient had been medicated for 28 days with a 14-day washout period; she successfully underwent 4 courses of chemotherapy without any major side effects. During the chemotherapy, she didn’t have neither upper respiratory tract nor abdominal symptoms. In the fifth course, she developed impaired consciousness and vomiting on the last day of medication.

Her surgical history included appendectomy. Neither she nor her family had a history of diabetes mellitus. She had no history of drinking or smoking and no known allergies. Upon admission, the Glasgow Coma Scale revealed a score of 11 (E3V3M5). Her abdomen was flat and soft. Laboratory data upon admission are presented in Table I. Arterial blood gas analysis revealed high anion gap metabolic acidosis. Urinalysis was positive for glucose and ketone bodies, and biochemical analyses revealed hyperglycemia. Abdominal computed tomography revealed fatty liver and pancreatic swelling. Based on these findings, the patient was diagnosed with DKA and was started with intravenous fluid infusion and continuous subcutaneous insulin injections. Her blood glucose
level gradually improved, and feeding and intensive insulin therapy was initiated on day three after admission. Laboratory findings on day four are presented in Table II. Each exocrine pancreatic enzyme was remarkably elevated. Her urinary C-peptide level after admission was 0.1 µg/day, whereas fasting serum C-peptide level was 0.01 ng/ml. C-peptide level remained stable at 0.01 ng/ml for six minutes after glucagon loading. Both anti-glutamic acid decarboxylase antibody and anti-insulin antibody were negative. We determined that the DKA was caused by FT1D. The patient was discharged on day 22 after admission, and no recurrence has been observed for five years since the radical surgical procedure.

Discussion

Our patient was provided adjuvant chemotherapy with UFT (500 mg/day) + LV (75 mg/day). The patient did not have a history of alcohol or drugs and had undergone four courses of chemotherapy without any major side effects. She developed impaired consciousness and vomiting on the last day of medication during the fifth course of chemotherapy. Because of the marked insulin deficiency and increased levels of the insulin counter-regulatory hormones cortisol and adrenaline, high levels of plasma glucose (≥300 mg/dl), hyperketonemia (increase of β-hydroxybutyric acid), and acidosis (pH, <7.3) occur together; this condition is known as DKA (Treatment Guide for Diabetes, 2014-2015). DKA is often considered the initial symptom of T1D, including FT1D. FT1D is a novel subtype of T1D that has been reported in Japan since the end of the 20th century (1). FT1D is characterized by rapid-onset DKA, low HbA1c levels, undetectable serum C-peptide concentrations, and negative islet-related autoantibodies (2). Although the pathogenesis of FT1D remains unknown, recent studies indicate its cause as either viral infection or genetic factors, such as HLA-II or CTLA4 CT60 (3-5). In this case, we didn't examine the antibodies against virus and genotyping for HLA-II or CTLA4.

Some studies have reported that interferon alpha induces autoimmune-mediated TID (6,7). The precise role of interferon alpha in the onset of TID in humans is unclear; however, interferon alpha is known to promote maturation of dendritic cells and activation of B cells in mice, thereby leading to autoimmune diabetes. Interferon alpha also directly influences pancreatic beta cells by inducing cytokines and enhancing their susceptibility to invasion by diabetogenic T cells (8). In addition, immune checkpoint inhibitors are known to cause endocrine-related adverse events. Although the evidence is limited, some reports have described new-onset T1D after anti-programed cell death-1 therapy (9-13). Their common point is that both interferon alpha and immune checkpoint inhibitors stimulate the immune system.

UFT is an anti-cancer agent (14). Tegafur produces 5-FU by hepatic metabolism. 5-FU acts as an anti-cancer agent by inhibiting RNA synthesis through active metabolites and
thymidylate synthase activity (15). Co-administration with uracil enhances the inhibition of 5-FU degradation, thereby increasing the concentration of 5-FU (16). UFT is widely used as adjuvant chemotherapy for Stage III colorectal cancer (17); its efficacy has been applied to high-risk Stage II colorectal cancer, such as in cases of perforation (18,19). Although several patients receive UFT, the development of FT1D during chemotherapy is extremely rare. To the best of our knowledge, only one case with FT1D during chemotherapy with UFT has been reported to date. Adachi et al (20) suggested two possible mechanisms for this development. The first is via immune suppression or immunological reaction, and the second is via the effects of thymidine phosphorylation. With regard to the first opinion, UFT have the common feature with interferon alpha and immune checkpoint inhibitors in the point of effect on the immune system, but the reason why other anti-cancer agents induce FT1D is remained. With regard to the second opinion, no reports of FT1D caused by 5-FU, TS-1, or capecitabine exist, although these belong to the same family of fluoropyrimidine anti-cancer agents. The mechanistic differences between UFT and other fluoropyrimidine anti-cancer agents might participate in development of FT1D. More cases and further studies are warranted to investigate the mechanisms of FT1D pathogenesis.

We reported an extremely rare case of DKA caused by FT1D during adjuvant chemotherapy with UFT + LV for colon cancer. The difference in the mechanisms between UFT and other fluoropyrimidine anti-cancer agents might contribute to the development of FT1D. More cases and further studies are warranted to investigate the mechanism of FT1D pathogenesis.

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Availability of data and materials

The datasets used and analyzed during the current study are available from the corresponding author on reasonable request.

Authors’ contributions

Y1, NM and KYo participated in the conception and design of the case report, analyzed and interpreted the data and wrote the manuscript. TS, MF, IY, TI, TTan, SM, HI, YT, KYa, NM and TTak evaluated the patient and participated in the therapy. NM, TTak and KYo revised the manuscript for intellectual content. All authors read and approved the final manuscript.

Ethics approval and consent to participate

Not applicable.

Patient consent for publication

Written informed consent was obtained by the patient.

Competing interests

The authors declare that they have no competing interests.

References