INTRODUCTION

Patients generally have chemotherapy after surgery for cancer, regardless of its stage. Chemotherapy usually causes several side effects, such as nausea, hair loss, and fatigue. The central nervous system is relatively safe from such toxicity because the blood-brain barrier protects it, and it has a slower cell proliferation rate. Nevertheless, chemotherapy agents can cause neurological and psychiatric complications.

Introduced in 1958, 5-Fluorouracil (5-FU) is one of the most common chemotherapeutic agents. It has been widely used to treat solid organ tumors, such as tumors of the colon, rectum, breast, pancreas, and stomach. Its usual side effects are gastrointestinal symptoms, alopecia, cardiotoxicity, and neutropenia. Although neuropsychiatric complications have not been common, physicians have reported leukoencephalopathy, depression, organic mental disorder, psychosis, and mania as side effects.

We report the case of a woman who developed an acute manic episode after receiving 5-FU chemotherapy during treatment for rectal cancer.

CASE

In January 2010, a 77-year-old female underwent a colonoscopy due to an intermittent anal bleeding episode. The test found a rather freely-moveable, 1 cm ulcerofungating mass in the left anterior side of the anus, 1 cm above the anal verge. Colonoscopic biopsy confirmed an adenocarcinoma. There was no sign of metastasis in her abdominal computed tomography (CT) or whole body positron emission tomography scan. On January 18, 2010, she underwent a transanal local excision under general anesthesia. The final pathology revealed a submucosal invasion by the tumor and a lymphatic invasion by tumor cells. However, the patient and her family declined a further curative radical surgery. Therefore, we recommended adjuvant radiotherapy and chemotherapy as the alternative management. She received radiotherapy, scheduled as 28 sessions, whole-pelvis at 4,500 cGy and tumor bed boost at 540 cGy, from February 9th to March 22nd. During the same period, she received intravenous chemotherapy, in the form of 790 mg (500 mg/m²) fluorouracil (5-FU) and 32 mg (20 mg/m²) calcium folinate, for 5 consecutive days at 3-week intervals. Starting February 16th, she also received the antiemetic ondansetron.

After her 4th 5-FU and folinate calcium chemotherapy cycle on May 15th, her behavior changed rapidly. That evening, her mood was elated and talkative. She called a friend and asked to visit in the middle of night, around 3 am. Her voice became louder and her need to sleep decreased. The next day, she was still anxious and talkative as compared to her previous character. She became more impulsive and sometimes offered verbal assaults. Her mood was labile, and she wrote a will and
then insisted that she was superior to the president of the Republic of Korea. Her family took her to a local clinic, which gave her a sedative, but her mood and behavior displayed no remarkable improvement.

Finally, she came to the hospital, and the staff surgeon consulted the psychiatrist, who decided to admit her on May 25th. She had no family history of psychiatric illness or previous history of psychiatric treatment. On admission, her consciousness was clear, and orientation was intact. Her mood was elated, euphoric, and labile. Her pressure of speech and inattention made it difficult to carry out her interview. She was easily distracted by environmental stimuli and expressed energetic interest in the psychiatric ward. She had grandiose delusions, such as insisting she was stronger than the president of the Republic of Korea was. There was no evidence of auditory or visual hallucinations. Her neurological examination was normal. A brain magnetic resonance imaging (MRI) showed no evidence of metastasis or any signs of acute infarct or hemorrhage. There were chronic infarcts in the right frontal periventricular white matter and several small, chronic, ischemic foci in the subcortical white matter on both sides. These findings did not match her psychiatric symptoms. A routine blood test showed mild anemia (Hb=10.7 g/dL).

Our diagnostic impression was secondary manic episode due to substance (ICD-10 diagnosis, organic manic disorder F06.30). We prescribed olanzapine (5 mg/day), divalproex sodium (250 mg/day), and lorazepam (2 mg/day). Then, after she had been hospitalized 5 days, we increased the dose to 10 mg olanzapine and 1,000 mg divalproex sodium. We discontinued the lorazepam after the 2nd day. She was still in a manic state, with hyperactivity, grandiose ideation, insomnia, and labile mood. After her 10th day in the hospital, we increased her medication to 20 mg/day olanzapine and 1,000 mg/day divalproex sodium. After this 10th day, her mood gradually calmed down, and she started to sleep without interruption or extra medication. She could partially recall her symptomatic behaviors of 2 weeks ago and expressed feelings of shame. We decreased her medication to 15 mg/day olanzapine and 1,000 mg/day divalproex sodium. She gradually returned to a normal range of moods, sleep pattern, and psychomotor speed. We conducted psychological tests during her admission day. Her IQ was average (total score=98), and her attention span and executive functioning were within normal ranges. The Bender-Gestalt test revealed no sign of organic brain damage. Minnesota Multiphasic Personality Inventory profile showed hyperactivity and a higher energy level, as the increased 9th factor.

On June 19th, she left the hospital with the manic episode in full remission. She did not show any extraordinary behavior, flight of ideas, or cognitive deficits. Her final medication was olanzapine, 10 mg/day, and divalproate, 1,000 mg/day. Although she was scheduled to undergo more chemotherapy, she and her family declined this idea, and her physician respected their decision. She visited the outpatient clinic one week and again one month after discharge. Her mood remained stable and euthymic. She had been taking her medications, with olanzapine at 5 mg/day and divalproex sodium still 1,000 mg/day. Her manic episode was still in remission. We scheduled her to receive the medication for several months, to prevent a possible relapse.

DISCUSSION

Though the patient received additional folinate calcium and ondansetron, we did not assume they were directly associated with her development of manic symptoms. As radiotherapy was limited to the rectal lesion, it could not affect the central nervous system. Therefore, the chemotherapy series using 5-FU was associated with her abrupt manic episode.

Fluorouracil's mechanism of action blocks DNA synthesis by decreasing thymidine monophosphate formation through the inhibition of thymidylate synthetase. As 5-FU can easily penetrate the blood-brain barrier, it is reportedly associated with several neurological complications, including parkinsonian syndrome, oculomotor disturbance, and peripheral neuropathy.7 The ending metabolite of 5-FU is fluorocitrate; we regarded its inhibition of aconitase in the Krebs cycle, resulting in citrate accumulation in the neurons, as the neurotoxicity mechanism.7 Fluorouracil itself did not correlate with the neurotoxicity.7 Previous cases studies reported leukoencephalopathy in brain CTs. However, this case showed no evidence of leukoencephalopathy on a brain MRI. We attribute her vulnerability to psychiatric symptoms to her age, because 5-FU side effects are common in the elderly.8

Accumulation of 5-FU is one possible cause of such a secondary manic episode, via injury to several brain structures and neurotransmitter pathways. Secondary mania should be diagnosed in a case of no previous affective illness history and no confusional state. This patient's onset age was considerably older than the typical age for a primary manic episode.9

There are several possible mechanisms related to secondary mania. First, it can develop after a neuronal injury focuses on right side limbic area lesion. And the right basotemporal and orbitofrontal cortex area is more likely to correlate with a bipolar episode, whereas a right side cortical injury alone occurs in association with unipolar secondary mania.10 An orbitofrontal cortex lesion can cause development of irritability, labile mood, euphoria, and inappropriate social behavior.11 Additionally, dysfunction of the hypothalamic area can lead to insomnia and dysregulation of appetite and sleep.12 At the
neurotransmission level, the interactions of serotonin, dopamine, norepinephrine, and acetylcholine through the orbitofrontal-subcortical network are associated with development of manic symptoms.9

Based on these research findings, we thought that 5-FU’s neurotoxicity accumulated during the 4 chemotherapy cycles, then a subtle change in the orbitofrontal-subcortical network, correlated with the dopamine and serotonin system, and reversible damage to the right cortical and limbic area caused a rapid manic episode. The manic episode subsided after discontinuation of 5-FU and administration of antipsychotic drugs. Furthermore, we thought the additional factors, such as her lack of any record of previous mood episodes, her rapid remission, the reversibility of her symptoms, and lack of cognitive change showed the manic episode developed due to an exogenous substance. Moreover, 5-FU-related neurotoxicity is dose- and schedule-dependent and reversible upon discontinuation of the medication.1

We reported that the chemotherapeutic agent 5-FU can induce, not only neurological symptoms, but also psychiatric symptoms. Clinicians need to consider the possible development of psychosis or mood symptoms in cases of 5-FU usage.

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