Foot drop - an uncommon presentation of FOLFOX toxicity

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Chemotherapy based on FOLFOX (oxaliplatin, leucovorin, and 5-fluorouracil) regimen is frequently used in colorectal cancer patients. Oxaliplatin and other platinum agents are known to be a class of chemotherapy drugs that commonly induce peripheral neurotoxicity. The most frequent oxaliplatin related neurotoxicity is sensitive symptoms. Here, we present two cases of patients with colon adenocarcinoma, both undergoing chemotherapy with FOLFOX4, who developed uncommon neurotoxicity, presenting with foot drop after the third treatment cycle. Foot drop may be explained by axonal damage of peripheral motor neurons of the common peroneal (fibular) nerve, which provides motor innervation to the foot muscles. Peroneal nerve palsy causes sudden weakness in the muscles of the foot that seems to be temporary. Both patients completely recovered from the event. There was no need for treatment adjustments, neither introduction of different drugs. Foot drop as chemotherapy toxicity is still poorly understood. The reported cases show foot drop as a severe and uncommon manifestation of FOLFOX-induced neuropathy, that might be transitory, and does not necessarily requires specific intervention.

Keywords: Foot Drop; Oxaliplatin; Neurotoxicity.
INTRODUCTION

The FOLFOX regimen (based on oxaliplatin, leucovorin, and 5-fluorouracil) is frequently used as treatment of colorectal cancer. Oxaliplatin and other platinum agents are a drug class in which peripheral neuropathy is a common adverse effect. Many patients have neurological toxicity during and after treatment with these protocols. The most frequent oxaliplatin related neurotoxicity is sensitive symptoms. Peripheral motor neuropathy and autonomic symptoms are less common. The severity of neurological adverse effects varies from mild paresthesia to severe pain with limitation in quality of life. This often leads to changes in chemotherapy protocols such as dose reduction, delay or treatment suspension.

In this study, we report uncommon form of oxaliplatin-related neurotoxicity, in two cases of colon cancer patients, during their chemotherapy with FOLFOX regimen.

CASE REPORT

CASE 1

A 59 years old female presented with constipation, abdominal pain and asthenia, performance status 1 and a positive fecal occult blood test. The colonoscopy was performed, showing a stenosing tumor at 50cm from anal margin, which pathological report was compatible with differentiated colon adenocarcinoma. Promptly, the patient underwent left hemicolectomy. The final pathological staging was pT3dpN1 cM1 (liver), RAS wildtype. Palliative chemotherapy with FOLFOX and panitumumab was initiated. She complained about transient CTCAE grade 1-2 paresthesia and sensibility changes in the oral cavity. After 3 cycles, the patient sought medical care due to frequent stumbles and falls, few days of its onset. She presented with a right sided foot drop, with a motor strength grade 3 in the scale ranging 0 to 5. The remaining physical exam was unremarkable. She denied traumatic injury. Radicular compression was the main suspicion. Further investigation was performed. Blood tests and Brain CT scan (Figure 1) were normal. MRI revealed bone metastasis in lumbar and sacral spine without pathologic fractures, medullar or radicular compression (Figure 2). A neurosurgeon and a rehabilitation doctor excluded spine or central causes. The patient got spontaneously better after one month. Treatment with FOLFOX was never changed or suspended and foot drop did not recur.

CASE 2

A 54 years old man presented with weight loss, performance status 0 and abnormal finding on screening colonoscopy. Biopsy revealed the diagnosis of well-differentiated colon adenocarcinoma and patient had sigmoidectomy surgery. Final pathological stage was pT3b N2a cM0, RAS wildtype. Palliative chemotherapy with FOLFOX and panitumumab was initiated. She complained about transient CTCAE grade 1-2 paresthesia and sensibility changes in the oral cavity. After 3 cycles, the patient sought medical care due to frequent stumbles and falls, few days of its onset. She presented with a left sided foot drop, with no traumatic history. The patient was admitted at the emergency room, where a full neurologic evaluation was performed by the neurologist, who confirmed the peripheral and focal weakness of the left foot with any other abnormalities. Blood tests were normal. He was discharged with recommendation of physiotherapy program of rehabilitation. All the clinical manifestations disappeared spontaneously two weeks later. Treatment with FOLFOX was never changed or suspended during this time.

Figure 1. Brain CT scan images without abnormalities (case 1).

Figure 2. MRI of lumbar and sacral spine without pathologic fractures or spinal cord compression.
DISCUSSION

Usually, chemotherapy neurotoxicity is dose dependent and cumulative\(^2\). Oxaliplatin is frequently associated with peripheral neurotoxicity, specially as with paresthesia/dysesthesia of the extremities and perioral region\(^3\). However, this drug may also be related to neuropathy by a non-dose dependent process. As described in MOSAIC trial, 92% of patients treated with a FOLFOX scheme experienced neuropathy of any degree during treatment, and in up to 30% of these patients, the symptoms remained even after the conclusion of the treatment\(^1,4\).

Our patients developed foot drop after the third treatment cycle (total cumulative dose of oxaliplatin of 255mg/m\(^2\) - 85mg/m\(^2\)/cycle). We assume that chemotherapy was responsible for this manifestation and interpreted as peripheral motor neuropathy caused by oxaliplatin toxicity. Foot drop can be caused also by central nervous system pathology, stroke or tumor, spine lesion or mononeuropathy/vasculitis. Extensive evaluation and complementary exams were performed to rule out these differential diagnosis.

There is still a lot to know about the mechanism of how chemotherapy agents provoke neuropathy. In the literature, it is hypothesized that platinum drugs target the dorsal root ganglion, with platinum DNA-binding resulting in neuropathy. Both of our patients were in treatment with oxaliplatin and had associated neuropathy. The mechanism of pathogenesis was not clarified.

Foot drop can be explained by axonal damage of peripheral motor neurons of the common peroneal (fibular) nerve that provides motor innervation to the foot muscles. Peroneal nerve palsy causes sudden weakness in the muscles of the foot that seems to be temporary. Both patients completely recovered from the event and there was no need for treatment adjustments, neither addition of different drugs.

The incidence of peroneal nerve palsy related with chemotherapy is unknown\(^5\).

Among the recommended treatment, it is included dose decreasing, interruption and discontinuation of chemotherapy agent. Physiotherapy might be helpful to stabilize the ankle and maintaining the range of movements\(^4\). Duloxetine is recommended as pharmacological approach, but strong evidences of its benefit is still lacking\(^4\).

In both cases reported in this paper, the recovery was prompt and spontaneous, and no dose modification were necessary. It differs from other publications, such as Dharmapuri and Pintova (2019)\(^3\), in which the patients took longer to recover and required physical therapy.

CONCLUSION

The foot drop as a chemotherapy toxicity entity is still poorly understood. Physicians of different specialties should be aware that oxaliplatin neurotoxicity pattern includes this type of manifestations. Further studies are needed to better describe this clinical finding in order to prevent it and to avoid potential delays/suspensions in chemotherapy treatments.

CONFLICTS OF INTEREST

The authors do not have any conflicts of interest to declare.

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