Introduction

Several neurological complications associated with metronidazole therapy have been reported, including sensory polyneuropathy, cerebellar dysfunction and encephalopathy. Proposed risk factors for toxicity include liver dysfunction [1], duration of treatment and total cumulative dose [2], but this is less clear for Central Nervous System (CNS) involvement, as has been recently suggested [3].

Case Report

A 55 years old man was admitted to the Neurology department with a three days history of fluctuating walking instability, and dysarthria. There was no history of alcoholism or drug addiction. One week after admission, cerebellar dysfunction progressively worsened, becoming permanent, but fluctuating, depending on the day. The pattern of the fluctuation was neither predictive nor related with the moment of administration of metronidazole. Neurological examination revealed gaze-evoked nystagmus, slurred speech, mild limb dysmetria, truncal ataxia and unstable gait. The patient had been under treatment with metronidazole for vascular necrosis and infection of the first finger of the right toe. After 2 months of treatment, the symptoms began (total duration of treatment: 2 months and 3 weeks; dose: 500mg/8h). There was no liver or renal dysfunction, and vitamin levels were normal (B1: 4,5mcg/dl and B12: 188pmol/L). CSF was also normal. Brain MRI at admission showed bilateral and symmetric hyperintensities in the cerebellar White Matter (cWM) and Dentate Nuclei (DN), Superior Cerebellar Peduncles (SCP), medulla, medullary Pyramid (P), pons, midbrain, Splenium of the Corpus Callosum (SCC) and subcortical White Matter (WM). Diffusion-Weighted Imaging (DWI) demonstrated high signal intensity within the SCC and low apparent diffusion coefficients, suggesting cytotoxic edema, whereas lesions involving the brainstem showed faint hyperintensity on DW and increased ADCs, consistent with T2 shine-through effect and/or vasogenic edema (Figure 1).

MRI findings suggested metronidazole neurotoxicity, and recognition of this entity was crucial for the patient’s management. Interruption of metronidazole treatment quickly induced clinical improvement, and the patient was almost asymptomatic after fifteen days. Two months later, a second brain MRI demonstrated resolution of most lesions, but the appearance of bilateral inferior olivary hypertrophy (Figures 1g and h).

Discussion

Metronidazole is a five-nitro-imidazole derivative antibiotic commonly prescribed for the treatment of anaerobic and protozoal infections of the gastrointestinal and genitourinary tracts.
Sensory-motor neuropathy is occasionally seen in patients treated with metronidazole, but other rare neurological complications including cerebellar dysfunction, seizures or encephalopathy have been also reported [4]. Neurological features usually become apparent when the drug is used with a dose exceeding 2g/day for prolonged periods. The cumulative dose implicated ranges from 13.2 to 228g, with neuropathy developing 2 weeks to 6 months following initiation [4]. Quick recognition of the clinical and radiological picture is crucial to avoid persistent disability in patients suspected of having metronidazole-related complications [5,6].

Here, we report a patient with highly suggestive MRI findings of neurotoxicity, who presented with cerebellar and brainstem dysfunction after metronidazole treatment. Previously reported cases of metronidazole neurotoxicity include similar findings to those of our patient, with bilateral, symmetric T2/FLAIR-hyperintensities in the cWM and DN, midbrain, dorsal pons, and SCC. On DWI, most of the lesions probably reflect underlying vasogenic edema, whereas involvement of the corpus callosum rather suggests underlying cytotoxic edema [7]. The lesion distribution in the brainstem is heterogeneous, as many structures can be affected. Lesions in the medulla may involve the vestibular and inferior olivary nuclei [5,6].

Restricted ADC values in the corpus callosum could indicate irreversible damage. Corpus callosum degeneration may produce memory impairment and slowdown retaining information [8]. Clinical follow-up of our patient after 4 years, showed a minor cognitive impairment, and neuropsychology demonstrated impairment of verbal memory, and difficulties in retaining new information. Therefore, we suggest clinical follow-up of cases having metronidazole neurotoxicity, as there may be long impairment, similarly to our case.

Inferior olivary hypertrophy in metronidazole toxicity has only been described previously in 2 cases [9,10]. In these, inferior olivary hypertrophy is suggested to be secondary to lesions involving the midbrain and cerebellum (interruption of the triangle of Guillain Mollaret), rather resulting from direct metronidazole neurotoxicity. In our case, both the central tegmental tract and the superior cerebellar peduncles were involved, and olivary hypertrophy occurred rapidly. Follow-up MRI two months after metronidazole withdrawal showed improvement of all lesions, except for the inferior olivary nucleus, which became hypertrophic. The findings in our case thus seem to support the view that olivary hypertrophy is secondary to disruption of the Guillain Mollaret triangle caused by metronidazole neurotoxicity, even if this is reversible [11].

The Guillain Mollaret triangle connects the red nucleus, via the central tegmental tract, to the inferior olivary nucleus, via the inferior cerebellar peduncle, to the contralateral DN, back to the red nucleus, via the superior cerebellar peduncle. When both the central tegmental tract and superior cerebellar peduncle are involved, bilateral hypertrophic olivary degeneration can occur. On pathology, gliosis appears 15-20 months after the primary lesion. Clinically, olivary hypertrophic degeneration may produce palatal myoclonus [10], which was nevertheless absent in our patient Hypertrophic olivary degeneration may last for almost three years after the initial presentation [12].

In the case described here, the combination of clinical and radiological signs reported were suggestive of metronidazole neurotoxicity, but the differential diagnosis should include other neurological conditions, especially Wernicke's encephalopathy [13].

Classical MRI features in Wernicke's encephalopathy consist on bilateral, symmetric T2 hyperintensities involving the mammillary bodies, and other structures, such as the medial thalamus, the floor of the third and fourth ventricles, the periaqueductal grey matter, midbrain tectum, and less commonly the pericentral cortex and superior cerebellar vermis [14]. The DN can also be affected in cases of malnourished Wernicke's encephalopathy, thus suggesting that vitamin B1 antagonism may play a role in the neurotoxicity associated to metronidazole [13]. Moreover, bilateral DN involvement has also been reported in cases of methyl bromide intoxication [15], metabolic conditions such as maple syrup urine disease [16] and enteroviral and other viral encephalomyelitis [17-19].

The pathophysiology of the brain changes related to imidazole toxicity is unclear, but similar clinical and radiological findings as metronidazole have been described in others 5-nitroimidazole drugs, such as tinidazole, secnidazole and ornidazole [20,21].

Conclusion

Differential diagnosis of a subacute progressive ataxia in adults should include drug intoxications/toxicity, among others. Metronidazole neurotoxicity should be considered in patients undergoing metronidazole treatment presenting characteristic MRI findings, especially in the DN and SCC, similarly to our patient. Rapid olivary hypertrophy may appear secondary to metronidazole neurotoxicity. Even though this entity is reversible, minor cognitive impairment may occur, therefore follow-up of these patients is recommended.

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