Intestinal Pseudoobstruction Caused by Chronic Lyme Neuroborreliosis. A Case Report

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Chronic intestinal pseudoobstruction is often classified as idiopathic. This condition is associated with poor quality of life and high morbidity, and treatment options are often unsatisfactory. A case of chronic intestinal pseudoobstruction in a 66-year-old woman, presenting with back and abdominal pain, urinary retention and severe constipation is described. The patient lived in an area in which Lyme disease is endemic and had been bitten by ixodes ticks. Intrathecal synthesis of anti-borrelia IgM and IgG and lymphocytosis in the cerebrospinal fluid was found, consistent with chronic Lyme neuroborreliosis since symptoms had lasted for more than six months. The patient’s gastrointestinal function recovered and the pain subsided significantly following treatment with antibiotics. Lyme neuroborreliosis (LNB) often results in palsy, but rarely affects the autonomic nervous system. Three patients have been described with intestinal pseudoobstruction due to acute LNB. However, this is the first described case of intestinal pseudoobstruction due to chronic Lyme neuroborreliosis. LNB must be suspected in patients with intestinal pseudoobstruction, in particular in patients who have been bitten by an ixodes tick and in patients living in an endemic area. (J Neurogastroenterol Motil 2015;21:440-442)

Key Words
Constipation; Intestinal pseudoobstruction; Lyme neuroborreliosis

Introduction

Chronic constipation is a common disease in the Western world, especially among the elderly population. This condition can usually be treated conservatively with lifestyle changes and laxatives. Nonetheless, occasionally patients with no previous history of gastrointestinal disease develop severe chronic constipation, often refractory to conservative treatment. This condition is also called chronic intestinal pseudoobstruction (CIP) and is often classified as idiopathic. Since the pathogenesis in some of these cases is due to a neuropathy involving the enteric nervous system, diseases that involve autonomic neuropathy, eg, diabetes mellitus, alcohol abuse, and paraneoplastic syndrome, are known causes of CIP.

In this case report we describe a case of sudden onset CIP due to chronic Lyme neuroborreliosis (LNB), which is known to cause neurological disorders, but has never before been described as the cause of CIP.
Case Report

A 66-year-old woman, with no major medical history was hospitalized due to urinary retention and lower back-pain radiating to the right lower extremity and lower abdomen. The patient lived in an area where Lyme disease is endemic and had been bitten by ixodes ticks previously. Physical examination on admission, including clinical neurological examination, was normal. Blood tests, including white blood cell count and C-reactive protein, were normal.

Spinal MRI revealed a suspected right-sided paramedian disc prolapse at L4/L5 with impingement of the L5 nerve root and a small disc prolapse at L5/S1. This was treated conservatively with physiotherapy and analgesia, including opioids. While the urinary retention resolved spontaneously, the pain intensified and the patient developed severe constipation, unresponsive to laxatives. Abdominal X-ray and CT scan suspected colonic ileus with some dilation of the colon (Figure). Absence of intestinal obstruction was confirmed by barium X-ray and the condition was interpreted as intestinal pseudoobstruction resulting from opioid therapy. Treatment with an acetylcholinesterase inhibitor resulted in defecation and flatulence, but the effects were only temporary. Colonoscopy revealed a flaccid, poorly-prepared bowel, but no other pathology was found. Colonic transit time investigation showed significantly delayed colonic emptying. Treatment with laxatives was intensified and transanal irrigation initiated.

More than 6 months after onset of symptoms the patient only defecated sporadically, even with twice daily transanal irrigation, supplemented with high dose osmotic and peristaltic stimulation. By this time she had lost 15 kg.

As a result of persistent back pain a repeat clinical neurological assessment was performed, which remained normal. Lumbar puncture was performed and investigation of the cerebrospinal fluid (CSF) showed CSF lymphocytosis with spinal leucocyte count on $103 \times 10^6/L$, 100% mononuclear, elevated IgG-index, elevated total spinal protein content, positive oligoclonal bands and intrathecal borrelia antibody synthesis, IgG as well as IgM. Since the symptoms had lasted for more than 6 months this was consistent with chronic LNB. No other bacteria or malignant cells were found in the CSF. Anti-Bb IgM and IgG was found in the serum.

The patient received a course of intravenous ceftriaxone 2 g administered once daily for 3 weeks. The frequency of transanal irrigation was reduced 4 days after onset of antibiotic therapy and 2 weeks later she had daily defecation without transanal irrigation, with decreased use of laxatives. Following improvement in gastrointestinal function the patient experienced a decrease in pain. After 3 weeks of antibiotic therapy, intensity of the pain slowly subsided with subsequently reduced need for analgesia. Three months after antibiotic treatment the patient was having a daily bowel movement with no need for laxatives. The pain was now of low intensity (visual analog scale 1-2) when treated with secondary analgesics. In a repeated lumbar puncture, the spinal leucocyte count had decreased to $9 \times 10^6/L$, spinal protein content had decreased and there was a dramatic decrease in intrathecal synthesis of anti-Bb IgM and IgG.

Subsequent nerve conduction studies showed no certain signs of myelopathy. The sensory evoked potentials showed bilateral peripheral affection of the upper extremities and the motor evoked potentials showed bilateral peripheral affection to musculus abductor pollicis brevis.

Discussion

This case report describes a case of CIP due to chronic LNB. LNB is a nervous system disorder caused by *Borrelia burgdorferi* (Bb), a tick-borne spirochete. Borreliosis is highly endemic in southern Scandinavia and central Europe. In North America, borreliosis occurs mainly in the North Eastern states but chronic
LN B is almost solely reported in Europe.4

LN B is divided into early and chronic disease. Chronic LNB is defined as disease duration of more than six months, CSF inflammation and Bb specific intrathecal IgG synthesis. Less than 1-2% of patients with LNB develop chronic LNB.4

Borrelia often presents as erythema migrans and if untreated approximately 5% will develop LNB in 2-6 weeks. The most common manifestation of early LNB is a subacute painful meningoarachiditis called Bannwarth’s syndrome and consists of CSF inflammation, painful radiculitis and palsies. Bannwarth’s syndrome comprises more than 80% of LNB cases but approximately 30% of these patients present without motor signs as seen in the present case. Characteristically, the pain is migrating, burning, often located in the back and radiating to the upper and lower extremities, abdomen and/or chest. Approximately 50% have cranial nerve involvement, predominantly a peripheral facial nerve palsy.

In the chronic stage, patients typically present with headache, malaise, significant weight loss, sensorineural hearing loss or a progressive spastic-ataxic gait disturbance.4

Autonomic dysfunction is rarely seen in LNB.4 If LNB is suspected, investigation of anti-Bb in the serum must be made. In the first stage of Lyme borreliosis, 20-50% of the patients will be anti-Bb IgM positive. Twelve weeks after onset of neurological symptoms, all patients will have detectable anti-Bb IgG in the serum.4 If the blood sample is positive for anti-Bb the CSF must be examined. Specific CSF antibodies appear during the second week after onset of neurological symptoms and are detectable in 100% of patients 8 weeks after onset. All patients have pronounced CSF inflammation with lymphocytosis and high protein content. Disease duration for more than 6 months, CSF inflammation and Bb specific intrathecal IgG synthesis is diagnostic of chronic LNB.4

Chronic LNB seems not to be self-limiting, but antibiotic therapy arrests disease progression and patients recover significantly. However, sequelae may occur. There is no evidence of improved outcome when treating LNB patients with antibiotics for more than 2 weeks.4

The patient in the present case, initially experienced radicular pain, but she also developed urinary retention and severe intestinal pseudoobstruction as manifestations of autonomic dysfunction.

The results of the nerve conduction studies were regarded as insignificant since the patient had no neurological symptoms of the upper extremities during the course of disease.

Autonomic dysfunction in LNB has been reported before. Urinary retention in LNB is described previously.5-7 Intestinal pseudoobstruction in patients with acute LNB has been described in three patients, two of whom had symptomatology very similar to the patient in the present case, with urinary retention in addition to the intestinal pseudoobstruction.5,8

To our knowledge, the present case report is the first reported case of CIP due to chronic LNB.

CIP is associated with poor quality of life and high morbidity.1 Pathophysiological aspects of the condition remain obscure and treatment options are limited and often unsatisfactory.9 Consequently, it is crucial that all physicians and gastroenterologists in particular, are aware of the non-idiopathic causes, and especially those that can be treated successfully. Infection with Bb can manifest as CIP and all of the 3 patients described, with acute LNB and intestinal pseudoobstruction, experienced complete remission after 2-3 weeks of antibiotic therapy. In the present case, 3 weeks of antibiotic treatment was effective even though the patient had developed chronic LNB.

In conclusion, Bb infection must be suspected in patients with sudden onset, severe constipation or intestinal pseudoobstruction, in particular in patients with simultaneous emergence of symptoms such as palsies, radicular pain, arthritis, and myocarditis, and in patients living in an endemic area.

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