ABSTRACT

Introduction. Camptocormia is an extreme flexion of the toracolumbar spine that exacerbates by walking and is relieved in the supine position. It is described with increasing frequency in association with Parkinson disease (PD). Its pathogenesis remains unclear. One of the hypotheses proposed defends that when associated with PD probably it is a focal action dystonia of the paraspinal muscles. Dystonia may be accompanied of a "geste antagoniste". It is a kind of trick that enables the patient to alleviate dystonic posture. We present two cases in which the patients use sensory tricks to alleviate their posture during gait. Case Reports. Patient 1: A 74-year–old woman, with a 2-year of PD. Clinical examination disclosed parkinsonian signs and camptocormia. She was able to partially overcome it by pushing her hands against her thighs. Patient 2: A 80-year-old woman, with a 1-year probable PD. Neurological examination revealed postural instability, freezing gait and camptocormia that partially remits when she pushes a bar at the level of her waist. Conclusions. The particularity of these cases is that patients are able to improve camptocormia during gait by specific tricks. This clinical improvement supports the hypothesis that camptocormia in PD is a focal dystonia of the paraspinal muscles.

Keywords. Parkinson disease, Posture, Distonic Disorders.

Citation. Semedo C, Calado A, Dias M, Almeida M, Pedrosa R. Tricks that relieve camptocormia during gait in Parkinson’s disease patients.
INTRODUCTION

Camptocormia is defined as an abnormal flexion of the trunk that appears when standing or walking and disappears when in the supine position.

The condition was described by Earle in 1815 and by Brodie in 1837, but the term itself was introduced by Souques to describe the forward flexion posture of some soldiers in World War I who had to bend to move through the trenches to avoid injury\(^4\)-\(^3\). The word is derived from the Greek words “kamptos” to bend and “kormos” trunk.

Despite considered evidence of a conversion disorder in earlier cases, camptocormia has been linked to a variety of organic diseases in the past decade, including parkinsonism, primary and secondary dystonias, spinal deformities, neuromuscular disorders, psychogenic causes, and others\(^4\).

Initially considered a rare feature of Parkinson’s disease (PD), a prevalence of 6.9% was reported by Tiple and co-workers\(^5\). It is associated with greater discomfort and disability\(^6\).

The mechanism of pathogenesis in PD remains unclear and several hypothesis have been put forward. One of them postulates that it is probably a central disorder, considering camptocormia as a focal action dystonia of the paraspinal muscles resulting from an excessive activation of the abdominal wall muscles\(^4\).

Dystonia is defined by involuntary, sustained contractions of muscles, leading to sustained abnormal postures or movements\(^7\). One of the most distinctive features of focal dystonias is their frequent improvement with sensory tricks (or \textit{geste antagoniste}). It is a kind of trick or manoeuvre that enables the patient to alleviate the dystonic posture. These often involve tactile stimulation of a particular body part, most often with mild force. The pathophysiology of this phenomenon has not been elucidated so far. One of the most postulated hypotheses suggests that it influences proprioceptive “input”\(^8\).

We present two case reports of patients with PD according to the UK Brain Bank Criteria that use a “\textit{geste antagoniste}” to improve posture during gait\(^9\).

Patients’ reports
Ref: 16/2011

Case 1: This 74-year-old woman presented a 2-year history of progressive gait difficulty and stooped posture. Upon examination she had hypomimia, bradykinesia and rigidity more pronounced on the right side of the body. Her gait was notable for its slow shuffling step and anteroflexion of the trunk with backward extension of the head (Fig. 1). The patient’s arm swing was diminished bilaterally and more pronounced on the right side. There was complete resolution of her camptocormia in supine.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure1.png}
\caption{Case 1. Camptocormia in a 74-year-old patient with Parkinson's disease (A). Her posture is markedly improved when she pushes her hands against her thighs (B). Patient consent has been received to publish this figure.}
\end{figure}
Non-demented, without frontal release signs, extra ocular movements were normal. No weakness or wasting of paraspinal muscles. Neuroimaging revealed chronic microvascular leukoencephalopathy. DaTSCAN showed a symmetric reduction of tracer uptake with more marked loss in putamen. At the time of presentation the patient was on Carbidopa-Levodopa (CD-LD) 25-100mg 3 times a day and selegiline 5mg 2 times a day, without significant improvement. It was supplemented with entacapone and selegiline. On her own initiative, she interrupted medication for two weeks resulting in worsening camptocormia. Higher dose of dopaminergic therapy was not tolerated. She was able to partially overcome the camptocormia during gait by pushing her hands against her thighs. This manoeuvre allowed her to straighten her back while walking (Fig. 2).

Case 2: This 80-year-old woman presented a 1-year postural instability, freezing gait and incapability to walk straight. She reported lower back pain. Upon examination she exhibited bradykinesia, marked flexion of the spine and lateral side deviation to her left side. She had no difficulty in lying down flat in supine position. No paraspinal muscle paresis or wasting. She was able to partially overcome her camptocormia by holding a bar at waist level, like when pushing a shopping trolley. She was progressively placed on a high dose of levodopa (CB-LD 25/100 up to 6 times a day plus CB-LD 50-200 sustained-release) which resulted in a marked improvement of parkinsonian symptoms but no change in her posture. Neuroimaging revealed a small cortical, left temporal infarction, and chronic microvascular leukoencephalopathy. Cervical MRI demonstrated multilevel osteoarticular degenerative disease.

**DISCUSSION**

Although camptocormia is receiving more attention, its pathogenesis and consequently its treatment still need to be determined. Three mechanisms have been considered as probable aetiology for camptocormia: dystonia, rigidity, and myopathy.

The more attractive hypothesis to consider for our two PD patients (case 1 and 2) is to consider camptocormia as a segmental dystonia of the trunk. The abnormal flexion of the trunk that appears when standing or walking and disappears in supine position could be caused by an excessive activation of the abdominal wall muscles. The improvement of flexed posture the patients achieved with the sensory tricks previously described also supports camptocormia as a form of dystonia. Our patients could even improve posture during gait (when it usually gets worse) using a sensory trick. The fact that patients use sensory tricks to improve their posture has already been

*Figure 2A*  
Figure 2. Case 2. Photographs of an 80-year-old woman with camptocormia (A) that is markedly improved when she to partially overcome her camptocormia by holding a bar at waist level (B).  
*Patient consent has been received to publish this figure*
reported in literature. The past history of spondyloarthrotic vertebral changes could be considered as a peripheral trauma acting as a trigger for dystonia.

It is theoretically possible that camptocormia in PD is the result of specific nigrostriatal dopaminergic projections, but the lack of correlation between the degree of camptocormia and clinical and treatment-related variables could imply that the pathophysiology of camptocormia involves additional non-dopaminergic mechanisms. The role of dopaminergic therapy on this disorder is complex. There are a few anecdotal reports of camptocormia that respond to dopaminergic treatment, botulinum toxin and deep brain stimulation.

Furthermore, another hypothesis for explaining parkinsonian camptocormia is the axial rigidity of flexion muscles, with weakness of the erector spinal muscles pointed out as a possible cause of camptocormia in PD patients. This would be a consequence of dysfunction of the basal ganglia controlling the reticulospinal pathway. It is unlikely that camptocormia could be a result of paraspinal rigidity in our patients otherwise it would not remit in supine position.

Currently, the discussion remains, especially since Margraf and co-workers published their study where they concluded that the most likely parkinsonian camptocormia is a focal myopathy of the paravertebral muscles. On the other hand, the possibility of an over-use myopathy has been mentioned by some researchers. We did not find any axial weakness in our patients and, for this reason, no further studies were performed to evaluate an eventual myopathic disorder.

**CONCLUSIONS**

In summary, we support the theory that considers camptocormia in PD patients a form of focal action dystonia, considering that the most finding of our two cases was the clinical demonstration of the use of a sensory trick to improve patients’ posture during gait. The relevance of determining the mechanism behind camptocormia consists in its implication on the therapeutic approach of this disabling condition.

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