The Beneficial Role of Thiamine in Parkinson’s Disease: Preliminary Report

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Abstract

Parkinson’s disease (PD) is the second most common form of neuro-degeneration in the elderly population. PD is clinically characterized by tremors, rigidity, slowness of movement and postural imbalance. A significant association has been demonstrated between PD and low levels of serum thiamine. Five PD patients presented with stone face, right-hand tremors, Parkinsonian gait and bradykinesia with occasional freezing. Two patients presented with sialorrhea and the plasma transketolase activity was low in one patient. All of the patients received 100 - 200 mg daily doses of parenteral thiamine. Within days of thiamine treatment, the patients had smiles on their faces, walked normally with longer steps, increased their arm swings, and experienced no tremors or sialorrhea. Three patients did not require carbidopa and levodopa without effects on their movements. Thiamine may benefit to PD. Further investigation of thiamine in PD patients is needed.

Keywords: Thiamine; Transketolase; Parkinson’s disease; Movement disorders

Introduction

Parkinson’s disease (PD) is a movement disorder characterized by tremor, rigidity, akinesia, and loss of posture reflexes, which leads to immobility and frequent falls. PD results from the selective loss of dopaminergic (DA) neurons in the substantia nigra (SN) of the brain. Recent studies highlight a possible relationship between thiamine and PD. Thiamine may be beneficial for PD patients. Lower central nervous system (CSF)-free thiamine levels were noted in PD patients compared with the controls [1]. In parkinsonism-dementia patients, thiamine-pyrophosphatase (TPP) activity was found to be significantly reduced in the frontal cortex [2]. In addition, Gold et al [3] reported that 70% and 33% of their PD patients had low plasma and red blood cell (RBC) thiamine levels, respectively. Starvation TD encephalopathy may also induce symmetrical lesions in the SN [4]. These findings suggest that thiamine might have a role in dopaminergic neuron activity. Interestingly, parenteral thiamine administration was used successfully in 9 non-alcoholic patients who presented with acute neurological disorders [5]. Administration of the combination of thiamine and acetazolamide reportedly reduced the Abnormal Involuntary Movement Scale (AIMS) and the Simpson-Angus Neurological Rating Scale (ANRS) scores of patients with tardive dyskinesia and parkinsonism symptoms [6]. In our previous publication, we discussed a number of genetic factors that link thiamine to PD pathology, including the DJ-1 gene, excitatory amino acid transporters (EAATs), the α-ketoglutarate dehydrogenase complex (KGDHC), co-enzyme Q10 (CoQ10 or ubiquinone), lipoamide dehydrogenase (LAD), chromosome 7, transcription factor p53, the renin-angiotensin system (RAS), heme oxygenase-1 (HO-1), and poly(ADP-ribose) polymerase-1 (PARP-1) gene [7]. In this paper, we report on the role of thiamine in PD patients.

Case Report

Case 1

The patient is a 76-year-old male, who has been diagnosed with PD for 8 years. He presented with rigidity, inability to close his mouth, sialorrhea, mask-like facies with infrequent blinking, and his eyelids dropped bilaterally. Additionally, he could not stand or walk alone. He is on carbidopa plus levodopa (25/100, one tablet twice a day) and benztropine mesylate (1 mg tablet twice a day). His blood test was remarkable for transketolase activity less than 2.0 nmol/L (normal, 8.0 - 30). A brain scan showed cortical atrophy and multiple nonspecific foci in the bilateral subcortical and deep white matter. After being given parenteral thiamine (100 mg daily) for 9 days, he could stand up unassisted and walks with...
normal associated movements in the arms. His eyes opened widely and smiling face. He experienced no sialorrhea. Unfortunately, he had an accident that resulted in a hip fracture, and the follow up ceased.

Case 2

The patient is a 74-year-old retired male physician, who has been diagnosed with PD for 7 years and presented with stone face with infrequent blinking, tremor of the right hand, loss of the normal associated movements in the arms during walking, walked-assisted Parkinsonian gait and bradykinesia with occasional freezing. He was on carbidopa plus levodopa (25/100, one tablet twice a day) and rasagiline 0.5 mg daily. He was prescribed with parenteral (200 mg daily). On the second day, he smiled and was able to walk longer steps without a walker and increased his arm swings. After 10 days of treatment, carbidopa plus levodopa and rasagiline were discontinued without any effect on his movement.

Case 3

The patient is a 68-year-old male who has been diagnosed with PD for 3 years and presented stone face with infrequent blinking, tremor of both hands, Parkinsonian gait and bradykinesia with occasional freezing. His memory had decreased over the last 2 years. He showed mild memory loss by the annotated mini mental state examination (AMMSE). His medications were carbidopa plus levodopa (25/100, one tablet twice a day). Computerized tomography (CT) of the brain revealed cortical atrophy. His plasma thiamine was 9.0 nmol/L (normal, 4.5 - 15.1). He was treated with parenteral thiamine 200 mg daily. On the fourth day, he smiled and was able to walk normally with longer steps, and increased his arm swings. His AMMSE score returned to normal. After 2 weeks of thiamine treatment, he was taken off carbidopa plus levodopa without any effect on his movement.

Case 4

The patient is a 65-year-old male, who has been diagnosed with PD for 8 years and presented stone face with infrequent blinking, tremor of both hands, difficulty pronouncing words, constant sialorrhea, Parkinsonian gait and bradykinesia with occasional freezing. His plasma thiamine was 6.0 nmol/L (normal, 4.5 - 15.1). His head CT scan revealed diffuse calcifications throughout the bilateral basal ganglia, caudate nucleus, bilateral occipital gyri, and bilateral cerebellum. He was treated with parenteral thiamine 200 mg daily. On the fourth day, he smiled and walked normally with longer steps, and increased his arm swings; no tremors or sialorrhea were reported. After 10 days of thiamine treatment, he was taken off carbidopa plus levodopa without any effect on his movement.

Case 5

The patient is an 82-year-old male who has been diagnosed with PD for 16 years and presented with stone face, tremor of the right hand, walked-assisted Parkinsonian gait and bradykinesia with occasional freezing. His medications were carbidopa plus levodopa (25/100, one tablet three times a day), stalevo 100 (carbidopa 25 mg, levodopa 100 mg, and entacapone 200 mg, one tablet a day), ropinirole extended release (12 mg a day), and rasagiline (1 mg daily). His whole-blood thiamine was 109 nmol/L (normal, 87 - 280). He was treated with parenteral 100 mg daily. On the fourth day, he smiled and walked with longer steps without a walker, increased his arm swing and no longer experienced hand tremor. Unfortunately, this patient was lost to follow-up.

Discussion

Note that our PD patients improved dramatically in a short time with thiamine supplements. Days after thiamine treatment, they smiled and walked normally with longer steps, and increased arm swings, and no tremors or sialorrhea was reported. Three patients did not receive carbidopa plus levodopa and cessation of those medications did not effect on their movements. The most effective treatment for PD is levodopa in combination with a peripheral decarboxylase inhibitor (carbidopa or benserazide). In a murine model, dopamine has been reported to suppress mouse-killing aggression (muricide) induced by a thiamine-deficient (TD) diet [8]. This suppressive effect can be potentiated with carbidopa [9]. Patients with PD who have undergone levodopa therapy have significantly higher cerebrospinal fluid (CSF) levels of thiamine diphosphate (TDP) and total thiamine than those patients who were not treated with this drug [1]. Moreover, thiamine deficiency can decrease the concentration of dopamine in the striatum, whereas animals fed on a diet that contained 5% ethanol exhibit increased dopamine turnover [10]. In an experimental TD study, a region-specific vesicular dysfunction (i.e., decreased levels of dopaminergic metabolites) was observed [11]. Dopamine release ii induced by intrastral administration of TPP or TDP (up to 1400% and 249% of the basal levels, respectively), reduced dopamine levels in the striatum may occur in cases of thiamine deficiency [12].

Sialorrhea is a disabling complication of advanced PD, and it interferes with PD patient’s abilities to speak, eat and socially interact with other people. Oral anticholinergic and botulinum toxin have been used as treatment, but the medications provided limited results that usually occurred weeks after treatment [13] and caused side effects. Interestingly, for two patients with sialorrhea, their symptoms disappeared after thiamine treatment.

Thiamine has also been implicated in PD via its effects
on L-type voltage-sensitive calcium channels (L-VSCC), matrix metalloproteinases (MMPs), prostaglandins (PGs), cyclooxygenase-2 (COX-2), reactive oxygen species (ROS), and nitric oxide synthase (NOS). These factors have been attributed to the pathogenesis of PD [7].

Gastrointestinal dysfunction is common in PD patients, and it potentially affects the therapeutic intervention [14]. Gastric emptying has been reported to be frequently delayed in PD patients [15]. Decreased non-mediated uptake across the enterocyte brush border membrane was demonstrated in PD patients [16]. In addition, the intestinal absorption of thiamine is sufficient in young people but may be reduced with age [17]. A single oral dose of thiamine above 2.5 mg is mostly unabsorbed in humans [18, 19]. Baker et al [20] demonstrated that only the intramuscular administration of thiamine was able to correct thiamine deficiencies in subjects over age 60. Sasaki et al [21] reported a case study of a patient with a thiamine deficiency and psychotic symptoms. Only repeated intravenous administration of thiamine ameliorated the patient’s condition. Furthermore, the patient responded rapidly to large doses of parental thiamine during the early stages of thiamine-deficient encephalopathy (namely Wernicke’s encephalopathy). The initial dose of thiamine is usually 100 mg two to three times daily for 1 to 2 weeks. Parental administration of thiamine was also used successfully in patients with general anxiety disorders [22].

Conclusions

Thiamine may have a beneficial role in PD. Further investigation of thiamine in PD patients is needed.

Conflict of Interest Statement

The authors, Dr. Khanh vinh quoc Luong and Dr. Lan Thi Hoang Nguyen, report no competing interests.

Ethical Approval

Not required.

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