Vivid Dreaming after Acetyl-L–Carnitine Administration: A Report of Two Cases

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Acetyl-L-carnitine (ALC), an acetylated form of L-carnitine, is able to influence the activity of cholinergic neurons, cell membrane stabilization and enhancing mitochondrial function. A 52-year-old woman was referred to neurology clinic for memory impairment within 1 year. She was administered ALC as dose of 1,500 mg per day for improving memory decline. After 14 days from administrating ALC, she complained vivid dreams at every night. Vivid dream was disappeared after ceasing ALC. Another patient, a 72-year-old man, visited neurology clinic for cognitive decline for 2 years. After 20 days from administrating ALC with dose of 1,500 mg per day, he also suffered from vivid dreams at every night. His previous stable sleep was also restored after ceasing ALC. ALC supplementation may present vivid dreams as a side effect. Possibility of vivid dream as a side effect should be considered during the management with oral ALC.

Key Words: Carnitine, Dreams

INTRODUCTION

Acetyl-L-carnitine (ALC) is an acetylated form of L-carnitine. ALC is able to influence the activity of cholinergic neurons, cell membrane stabilization and enhancing mitochondrial function. Previous studies have been showed beneficial outcomes of oral ALC supplementation in patients with cognitive decline in patients with dementia, such as Alzheimer disease and elderly patients with fatigue. However, there have been a little cautious observation for drug side effects, as compared with widely used drugs for patients with behavioral and cognitive deficits. Here, we report two patients who developed vivid dreaming after oral intake of ALC to improve cognitive function.

CASE REPORTS

CASE 1

A 52-year-old woman was referred to neurology outpatient clinic for memory impairment, developed within 1 year. She had no pertinent medical history including hypertension and diabetes. She had no drug history as well as nutritional supplement. She denied family history of dementia and neurodegenerative disease. Her Mini-mental Status Examination (MMSE) score was 27 out of a possible 30 points; she lost two points on the serial sevens from 100 and one point on the 3-stage command. Assessment of her activity of daily living (ADL) was normal. Brain magnetic resonance images (MRI) showed no significant findings, such as mild cerebral atrophic changes explainable to memory impairment (Fig. 1A). She was administered with ALC, dose of 1,500 mg per day, for improving memory decline. After 14 days from administrating ALC, she complained of vivid dreams at every night. She described that her dreams were too vivid and bothersome at night. The
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**CASE 2**

A 72-year-old man visited a neurology clinic due to cognitive decline for 2 years. He had no pertinent medical history, such as hypertension or diabetes mellitus. He denied any drug history as well as nutritional supplement. He also denied family history of dementia and neurodegenerative disease. His brain MRI showed mild ischemic changes in periventricular area and subcortical white matters and mild global atrophy (Fig. 1B). He performed neuropsychiatric function test. His MMSE was 25 points. He lost 2 points on memory recall and 2 points on the serial sevens from 100 and 1 point on the 3 stage command. However, his ADL was normal. He had no neurologic deficits at motor, sensory, coordination and gait tests. After 20 days from administering ALC with a dose of 1,500 mg per day for improving cognitive decline, he also suffered from vivid dreams at every night. He had 2 or 3 separate dreams that were graphic and vivid. The contents of his dream were standing surrounded colorful wall. And sometimes, his dream included familiar games called ‘go-stop’ with his friend. His dreams were too vivid to distress at night. In addition, his mornings were not fine and did not feel refreshed because of the dreams. His sleep was also restored after ceasing the ALC treatment.

**DISCUSSION**

The rapid eye movement (REM) sleep is distinct behavioral status, characterized by an activated cortical and hippocampal electroencephalogram and concurrent muscle atonia. Commonly, REM sleep is called as a dreaming sleep. Sublaterodorsal (SLD) nucleus which was located just ventral to caudal laterodorsal tegmental nucleus in the brainstem has been suggested as a “REM-on” cell based on several animal studies. The glutamate released from SLD nucleus is critical for regulating REM sleep phenomena.

Under the mechanism of ALC, the high level of serum...
carnitine is able to induce the glutamate releasing in SLD7).

As a result, it is able to enhance “REM-on” cell. Accenting projections of REM-on glutamate neurons might regulate the cortical features of REM sleep. We assumed that the vivid dream in our patients was a final consequence of these responses at high level of serum carnitine.

Dysfunction of fatty acid β-oxidation by abnormal low carnitine levels has been suggested in narcolepsy patient6). The deficiency of carnitine causes the fatty acid oxidation disturbances under fasting when fatty acids need to become a major energy source7). The orexin neuronal activity, which is an important role in the maintenance of wakefulness, is able to proportionally reduce with the level of serum carnitine7). Moreover, prolonged reduced orexin activity resultantly showed fragmentation of sleep and significant decreased amounts of REM sleep.

From all suggestive mechanisms, associated with carnitine metabolism, we suggested that overdose supplementation of carnitine could induce the dysfunction of fatty acid β-oxidation, increasing orexin neuronal activity and glutamate releasing from SLD8).

In addition, ageing is able to cause a number of changes in drug absorption, distribution and elimination. Therefore, the pharmacologic effect that is previously mentioned of ALC could be strengthened in elderly9). When we prescribe medicines to elderly with any purpose, we should have the pharmacologic changes of elderly in our mind.

However, there are some weak points to suggest vivid dreams after carnitine supplementation in our patients with cognitive declines.

First, we did not evaluate the level of serum carnitine before and after oral administration in each patient. Nevertheless, these two patients revealed suffering from vivid dreams within 2 weeks from carnitine administrations in same, and relieved after ceasing the ALC medications. We suggested that it could be an evidence for carnitine effects for vivid dreams.

Second, we felt the lack of this study for overnight sleep such as polysomnography, which is the best study for sleep quality and structure. However, polysomnography is not able to reveal whether the dream is vivid or not. It is only able to evaluate the amount of REM in proportion to the total sleep time. We thought the report about sleep problems by patients after nights were reliable to evaluate the dream contents.

Third, there were no exact pathomechanism for REM sleep disorder, such as nightmare, REM sleep behavior disorder and sleep related hallucination. The orexin neuronal activity, glutaminergic neuron in SLD have been just suggested a major neurotransmitters in REM sleep. In addition to these classical neurotransmitters, there are more considerable neurotransmitters and neuromodulators to influence the sleep regulation, including REM sleep.

In summary, oral ALC administration might present as vivid dream problems. This possibility should be considered during the management with oral carnitine, and should be checked for any possible sleep problems, including dream enhancing sleep disorders in patients treated with carnitine.

**REFERENCES**