Proton pump inhibitor–induced hypomagnesaemia leading to recurrent falls and delirium: a case report

Tanya Joy Zapata Quicho, MD, GDGRM, Christine Yuanxin Chen, MBBS, MRCP

ABSTRACT
Older adults are at increased risk of developing hypomagnesaemia secondary to chronic use of proton pump inhibitors. We report a 73-year-old woman who presented with acute functional decline, recurrent falls, and cognitive impairment likely owing to unrecognised persistent hypomagnesaemia secondary to use of proton pump inhibitors.

Key words: Aged; Proton pump inhibitors

INTRODUCTION
Proton pump inhibitors have been reported to cause hypomagnesaemia. Manifestations of hypomagnesaemia include muscle weakness, tremors, delirium, coma and even death. We present a 73-year-old woman who had progressive muscle weakness, recurrent falls, arrhythmias, and delirium secondary to hypomagnesaemia.

CASE PRESENTATION
In February 2020, a 73-year-old Chinese woman with comorbidities of hypertension, type II diabetes mellitus, hyperlipidaemia, and ischaemic heart disease was admitted to the acute geriatric ward with sudden unresponsiveness during a meal.

Before October 2019, the patient could walk independently. However, she started to have recurrent falls and was hospitalised in late October 2019. Since then, she had an acute decline in function and cognition. She was unable to walk and needed moderate to maximum assistance in transfers. She also became less conversant and had a decline in cognitive function. It was noted that her magnesium levels were persistently low at 0.4 to 0.5 mmol/L.

Upon this current admission, she had irregular heart sounds and weakness of all four limbs. She was not able to stand independently. She could not complete the abbreviated mental test, as she was not responding consistently. She was found to have a stage 3 sacral pressure injury.

Magnetic resonance imaging result of the brain was unremarkable. Blood test results showed hypernatraemia (150 mmol/L), hypokaelaemia (2.5 mmol/L), and hypomagnaesemia (0.19 mmol/L). She had no hypoglycaemia or anaemia and her blood pressure was normal. Inflammatory markers were normal. Electrocardiography showed frequent premature ventricular complexes. A 24-hour Holter showed some atrial ectopies and infrequent ventricular ectopies. Transthoracic echocardiography showed an ejection fraction of 55% with no wall motion abnormalities or valvular disease.

She had dysphagia and was placed on a modified diet. On transfer to the subacute ward, she was drowsy. She could open her eyes to calling but was not orientated and could not sustain attention. The power of all four limbs was 2/5. Her sitting balance was very poor and there was no neck control.

Repeat blood tests revealed that her magnesium levels dropped to 0.33 mmol/L. This was replenished intravenously. A drug review showed that one of her
chronic medications was omeprazole, which was stopped while magnesium was being replenished. Her thyroid and parathyroid hormone levels and serum calcium and potassium levels were normal. The magnesium levels recurrently dropped and several courses of magnesium replenishment were given. After intravenous and oral replenishment and maintenance magnesium tablets, her serum magnesium level normalised to around 0.7 mmol/L, and her attention span improved. She scored 5/10 on the abbreviated mental test. She recovered some sitting balance, and the power in all limbs improved to at least 3/5.

The renal specialist concurred that the hypomagnesaemia was most likely caused by omeprazole. Omeprazole was stopped permanently, and she was maintained on oral magnesium. At the home visit 1 month post discharge, her magnesium level was normal. She was followed up for 3 months and had no further improvement in cognition. She was subsequently diagnosed with dementia.

DISCUSSION

Proton pump inhibitor–induced hypomagnesaemia may result from impaired intestinal absorption owing to reduced active transport of the Mg$^{2+}$ in the reduced intraluminal pH.1 The reported time to development of hypomagnesaemia ranges from 14 days to 13 years.2 Older adults with multiple comorbidities (diabetes mellitus, polypharmacy, and poor dietary intake) are at higher risk of developing hypomagnesaemia.

When older adults present with falls and loss of consciousness, initial management is to rule out cardiac and neurologic pathologies. Hypomagnesaemia is usually not considered a cause of cognitive impairment or delirium. Our case illustrates the importance of including testing of serum levels of magnesium in patients with falls and functional decline. Serum levels of magnesium should be monitored frequently during treatment of hypomagnesaemia, as magnesium is an intracellular ion, and its repletion may not be reflected correctly on the first normal level result. Repeat testing is thus important.

It is imperative to evaluate the underlying cause of hypomagnesaemia to prevent recurrence.3 Proton pump inhibitor use of more than a year significantly increases the risk of hypomagnesaemia.4,5 As a large population of older patients are prescribed with proton pump inhibitors indefinitely, medication review for appropriateness of long-term proton pump inhibitor use should be routinely performed. Serum magnesium levels should be checked before and during proton pump inhibitor therapy, or when there are acute changes in function or cognition. This may prevent prolonged state of hypomagnesaemia and its complications.

REFERENCES