Hyponatremia and Cerebral Edema Associated with Radioiodine Therapy for Papillary Thyroid Carcinoma

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Abstract

Among the serious complications associated with radioiodine therapy (RAI) for thyroid cancer, cerebral edema is uncommon and has been reported previously in cases of swelling of brain metastases. This case is of a patient with papillary thyroid carcinoma who complained of nausea and vomiting after RAI and was then found unconscious the next day. Laboratory results showed electrolyte imbalances including hyponatremia and cranial imaging only revealed cerebral edema, and she regained consciousness after sodium correction. The etiology of the cerebral edema here is likely multifactorial - due to hyponatremia from hypothyroidism, aggravated by vomiting, copious water intake and a low sodium diet.

Keywords: hyponatremia, hypothyroidism, radioiodine

Background

In the treatment of well-differentiated thyroid cancer, radioiodine (I-131) therapy (RAI) for thyroid remnant ablation is safe and complications are usually mild and tolerable. More serious complications after RAI are rare but they can cause significant morbidity and mortality. The most critical acute complications are cerebral edema or hemorrhage due to swelling of brain metastases. However in this case, alteration of sensorium and cerebral edema occurred in the absence of brain lesions, and other factors may have been contributory. This is important to consider when faced with a patient who loses consciousness after RAI.

Case report

A 30-year-old female presented with a slowly growing anterior neck mass over the course of 2 years. She was diagnosed with nodular non-toxic goiter and subsequently had a subtotal thyroidectomy. Final histopathologic examination however revealed papillary thyroid carcinoma and a completion thyroidectomy with selective node dissection was then performed (Stage I, T3N0M0, <45 years old). After a few months the patient was admitted for RAI after cessation of levothyroxine (LT4) for 3 weeks and being advised to limit iodine intake in accordance with local guidelines.1 On the day of admission she complained of some light-headedness, which she attributed to ongoing menstruation. Pregnancy test was negative while vital signs were stable and physical examination unremarkable and 131I 3.7 GBq (100 mCi) was administered. Six hours later, she complained of nausea and had 2 episodes of vomiting. On the second hospital day, nausea and vomiting persisted, although some relief was obtained with anti-emetics. She reported tolerance of small feedings and compliance with liberal fluid intake. She was normotensive and did not appear to be dehydrated. Two hours after last being seen however, she was found unconscious and could only be minimally roused with noxious stimuli. Vital signs were within normal limits and physical examination was unrevealing, with no evidence of trauma. Neurologic examination did not reveal any lateralizing signs. Pupils were equal and briskly reactive, reflexes were symmetric and extensor toe sign was negative. There was no note of abnormal posturing, involuntary movements or withdrawal to painful stimuli. Dose rate at a distance of 1 meter from the patient was 10 mR/hr.

Differential diagnosis

Differential diagnoses for her altered sensorium at this time included hypotension due to dehydration from vomiting and possible head trauma. Metabolic causes could have been either from hypoglycemia due to poor intake and excessive vomiting or hyponatremia secondary...
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Vol. 27 No. 1 May 2012

www.asean-endocrinejournal.org

Among the serious complications associated with radioiodine therapy (RAI) for thyroid cancer, cerebral edema is likely multifactorial—due to hyponatremia from hypothyroidism, aggravated by vomiting, copious water intake and a low sodium correction. If they do occur it is after large single administrations of cumulative amounts of RAI have been given.2,3 In this case, this was the patient’s first, relatively low dose of RAI. Based on these findings and the patient’s response to sodium correction, it was concluded that the cerebral edema was secondary to hyponatremia probably due to hypothyroidism and exacerbated by a low iodine and sodium diet, gastrointestinal losses due to acute radiation sickness and excessive fluid intake.

Discussion

Serious complications associated with RAI are unusual, but if they do occur it is after large single administrations of RAI at doses above 131I 7.4 GBq (200 mCi) or when large cumulative amounts of RAI have been given.2,3 In this case, this was the patient’s first, relatively low dose of RAI.

Treatment

Given the possible etiologies, the most critical was that of cerebral edema due to brain metastases and intravenous (IV) dexamethasone was started as soon as the patient was found unconscious. The Neurology service was also consulted, and once the CT scan was obtained, furosemide and mannitol were given for the cerebral edema. Intravenous correction was started for the electrolyte deficiencies.

Outcome and follow-up

The patient regained consciousness the next day, 16 hours after the being found unresponsive. She could not recall the previous events, except for the sensation of dizziness and episodes of vomiting. She admitted to drinking numerous glasses of water but could not recall the exact amount. The patient was found to have 2 almost-empty 5-gallon containers of drinking water in her hospital room which were full on admission, with no spillage was observed. Physical and neurologic examination did not reveal any lateralizing signs suggestive of an acute cerebrovascular event. Serum sodium was corrected to 128 mmol/L and subsequently to 130 mmol/L by the fourth hospital day, with the other electrolytes corrected to within normal range. A follow-up plain cranial CT scan showed no evidence of the previously seen frontal infarcts (Figures 1-B and 1-C) with resolution of the pontine and cerebral edema (Figures 2-C and 2-D). A cranial CT scan with contrast was also done and this did not reveal any other abnormalities. A post-ablative whole body scan done 7 days after RAI showed only functioning residual thyroid tissue in the anterior neck and no evidence of metastases to the brain (Figure 3).

Figure 1. Plain cranial CT scan showing acute bilateral frontal infarcts (A) and subsequent resolution (B and C).

Figure 2. Plain cranial CT scan showing mild pontine and cerebral edema (A and B) and resolution after electrolyte correction (C and D).

Initial capillary blood glucose was 4.38 mmol/L and after administration of 1 vial of intravenous 50% dextrose, her blood glucose was rechecked and noted to be 12.21 mmol/L. The patient did not regain consciousness. Multiple electrolyte abnormalities were found, including hyponatremia (sodium:118 mmol/L, NV:136-145 mmol/L), hypokalemia (potassium:2.5 mmol/L, NV:3.5-5.1 mmol/L), hypomagnesemia (magnesium:0.57 mmol/L, NV:0.7-1.0 mmol/L) and hypocalcemia (calcium:1.71 mmol/L, NV:2.12-2.52 mmol/L). Serum osmolality was low at 251 mmol/kg. Urine specimens for electrolytes and osmolality were not obtained due to the patient’s elevated radiation levels. A 12-lead electrocardiogram showed occasional premature ventricular contractions (PVCs) with no other specific changes.

While obtaining a computed tomographic (CT) scan, a lead gown and thyroid shield were placed over the patient. This was in accordance with radiation safety practices, providing healthcare providers with adequate shielding from radiation as the patient had just received RAI the day before. Plain cranial CT scan showed hypodensities in the frontal subcortical regions that were interpreted as acute bilateral frontal infarcts. (Figure 1-A). Mild pontine and cerebral edema were also noted. (Figures 2-A and 2-B).

In this case, alteration of sensorium and cerebral or hemorrhage due to swelling of brain metastases. The most critical acute complications are cerebral edema but they can cause significant morbidity and mortality. Ablation is safe and complications are usually mild and years old). After a few months the patient was admitted for RAI after cessation of levothyroxine (LT4) for 3 weeks. Examination however revealed papillary thyroid hypokalemia (potassium:2.5 mmol/L, NV:3.5-5.1 mmol/L), hypomagnesemia (magnesium:0.57 mmol/L, NV:0.7-1.0 mmol/L) and hypocalcemia (calcium:1.71 mmol/L, NV:2.12-2.52 mmol/L). Serum osmolality was low at 251 mmol/kg. Urine specimens for electrolytes and osmolality were not obtained due to the patient’s elevated radiation levels. A 12-lead electrocardiogram showed occasional premature ventricular contractions (PVCs) with no other specific changes.

Investigations

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Early complications
Gastrointestinal symptoms be due to acute radiation sickness may occur as early as 2-12 hours and up to 36 hours after RAI and typically lasts 1-2 days. Symptoms include fatigue, headache, nausea and vomiting which are controlled with anti-emetics. The reported incidence varies from 36-73%, but vomiting occurs in no more than 5%.3,4

Central nervous system complications
Radiation-induced swelling may also affect metastatic tissue. Aside from bony and soft tissue, this may occur in the spinal cord or brain, causing pain, edema, hemorrhage and compressive symptoms.3,5,6 The brain accounts for only 1% of the 10% of distant metastases that occur in patients with well-differentiated thyroid carcinoma,7-9 but the effects of radiation-induced swelling can be devastating. There are a few reports on serious central nervous system complications after RAI, including a case of hemorrhage of brain metastases and 2 of cerebral edema, with 1 being fatal.3,5,6 It has been recommended pre-treatment prior to RAI with corticosteroids and mannitol may reduce the risk for this complication.2,10,11

Hyponatremia and hypothyroidism
Symptomatic hyponatremia has been reported to complicate RAI, although usually in the period prior to treatment during withdrawal of LT4. Reported cases complained of dizziness and confusion just before RAI, and all were found to have low sodium levels which were either treated with saline, fluid restriction or liothyronine (LT3) prior to RAI.12,13 In another reported case, the symptoms of hyponatremia manifested almost 2 weeks after RAI with LT4 replacement already resumed.14 In all of these cases the patients were over 60 years old and had been on LT4 withdrawal and low iodine diets in preparation for RAI.

For patients undergoing RAI, current guidelines recommend LT4 withdrawal for at least 2-3 weeks or LT3 treatment for 2-4 weeks and LT3 withdrawal for 2 weeks to achieve hypothyroidism with a TSH level of >30 mIU/L.11 Hypothyroidism has been noted to induce hyponatremia, although the mechanism by which this occurs is incompletely understood. Hyponatremia itself is the most commonly encountered electrolyte disorder in hospitalized patients, and is more of a disorder in water rather than sodium balance.15 In hypothyroidism, the principal abnormality in patients with normal fluid intake appears to be an inability to maximally suppress antidiuretic hormone (ADH).16 There is also a decrease in renal blood flow and glomerular filtration and this can directly diminish free water excretion by diminishing water delivery to the diluting segments.17,18 These however do not lead to significant hyponatremia unless there are other contributory factors such as diuretic use, excessive extra-renal losses as in diarrhea and vomiting or large volumes of fluid intake,18 as in our case. Reduced cardiac output in hypothyroidism may also play a role, stimulating volume-mediated ADH release via the carotid sinus baroreceptors.17,19

It is unclear if there truly is a relationship between hypothyroidism and hyponatremia. Studies in hypothyroid children and adults have shown that low serum sodium concentrations were not altered despite LT4 correction.20-22 In one report of almost 1,000 hypothyroid patients, none were found to have serum sodium concentrations <120 mmol/L although this was present in 2 controls.23 Given this data it would appear that the presence of a low serum sodium in a hypothyroid patient would require further investigation into other possible causes for hyponatremia.

Hyponatremia and low iodine diets
Iodine depletion has been found to allow for increased delivery of the radiation dose in thyroid cancer ablative therapy.24 This is achieved by a low iodine diet (<50 mg/d of dietary iodine) for 1-2 weeks prior to RAI, especially in patients with high iodine intake. Patient should also avoid any iodine contamination (e.g., intravenous contrast, amiodarone use) to increase the effective radiation dose.11

Unfortunately, many patients equate a low iodine diet with a low salt diet.25 In our setting where it is mandated by law that all salt be iodized,25 it is difficult to obtain non-iodized salt and this leads to the avoidance of salt altogether. In states of low solute intake, the ability to generate and therefore excrete free water becomes limited if solute excretion is markedly decreased, which can further worsen hyponatremia, if present.26 In susceptible patients with hypothyroidism, such as the elderly, serum
sodium levels should be measured in the event that they exhibit symptoms of hyponatremia.13

**Hyponatremia and polydipsia**

It is possible that excessive fluid intake may have contributed to the condition of the patient in this case. Patients are advised to drink water liberally and urinate frequently to avoid unnecessary radiation to the bladder and pelvic area after RAI therapy.22 The patient admitted to drinking large amounts, but the actual volume of water intake was not documented. Hyponatremia related to polydipsia is generally considered to be an acute dilutional state in which water intake overwhelms renal excretory capacity. It is also said to be a mixed disorder of increased intake of water and loss of solute.23 In normal subjects, 10 to 15 liters of urine may be excreted per day, a response that is mediated by suppression of ADH secretion and the subsequent formation of a dilute urine. Thus, if ADH regulation were intact, intake of about 10 to 12 liters of water should not lead to clinically important disturbances in the plasma sodium concentration.24 Water intoxication and hyponatremia could occur however if intake were massively increased,25 or if there was volume depletion, use of diuretics, or the presence of co-existing disturbances in renal, endocrine, cardiac and hepatic function. In these situations, amounts of fluid that are below the water excretory capacity of the kidneys can already cause hyponatremia.25

**Conclusion**

Loss of consciousness and the development of cerebral edema after RAI is uncommon and its occurrence should alert physicians to the possibility of swelling of undetected brain metastases. However in hypothyroid patients, this may also occur in the setting of hyponatremia, and in this case, was exacerbated by vomiting, a low sodium diet and excessive fluid intake.

This was a young patient who presented with symptomatic hyponatremia, which is unusual in her age group, as this has only been observed in elderly patients placed on low iodine diets who may equate this with a low sodium diet. Proper instruction on dietary restriction should be discussed with patients prior to RAI, and monitoring of serum sodium levels should be considered for symptomatic or elderly patients. Once RAI has been administered, patients are also advised liberal fluid intake during their hospital stay but they should avoid excessive amounts.

**Acknowledgements**

Thank you to Dr. Ruben Ogbac, Dr. Lorraine Almelor and the nurses at the Cancer Institute of the Philippine General Hospital for their assistance in the management of this case.

**References**