Raised parathyroid hormone levels in the milk alkali syndrome: an appropriate response?

David W. Bullimore and Krzysztof J. Miloszewski

St. James's Hospital, Leeds LS9 7TF, UK.

Summary: A case of the 'milk alkali syndrome' associated with grossly elevated levels of amino terminal parathyroid hormone is described. The hypercalcaemia (calcium 4.09 mmol/l) and hyperparathyroidism settled on conservative measures. Factors in the milk alkali syndrome which might stimulate the release of parathyroid hormone include parathyroid gland hyperplasia secondary to suppression of ionized calcium, alteration in sensitivity of calcium receptors on the cells of the parathyroid glands, the stimulation of an intermittent alkaline tide in the blood and the high intake of phosphate and bicarbonate.

We suggest that high levels of parathyroid hormone in the milk alkali syndrome may be appropriate rather than paradoxical.

Introduction

The Sippy diet for the management of peptic ulceration involved the consumption of large amounts of calcium and absorbable alkali.1 Soon after its introduction toxic complications were noted.2 In 1949 Burnett3 described the chronic form of toxicity characterized by hypercalcaemia, alkalosis and renal dysfunction in subjects with a prolonged and excessive intake of milk and absorbable alkali. The condition has become known as the milk alkali syndrome or Burnett's syndrome.

Several cases have been reported in association with primary hyperparathyroidism.4-6 The association has been considered to be either coincidental or representing cases of primary hyperparathyroidism in subjects taking large quantities of calcium and absorbable alkali for relief of dyspeptic symptoms, peptic ulceration occurring more commonly in subjects with hyperparathyroidism.7

The case described is of a woman with the milk alkali syndrome and raised levels of parathyroid hormone, in the presence of marked hypercalcaemia, suggestive of hyperparathyroidism. The hypercalcaemia resolved rapidly (days) and the hyperparathyroidism gradually (months) on rehydration and cessation of excessive intake of calcium and alkali. The possible causes of the apparently paradoxical elevation of parathyroid hormone are discussed.

Case report

A 35 year old woman presented with chronic constipation, anorexia and a 6 week history of nausea and vomiting. Following a Heller's operation for achalasia at age 13 and two subsequent antireflux procedures she had persistent symptoms of oesophageal reflux. Her typical daily calcium intake was 6.5 g (normal 0.6 g) in the form of 4 pints of milk, 12 'Rennies', a proprietary antacid containing calcium and magnesium carbonate, and 'Complan', a food supplement, plus large amounts of absorbable alkali. Investigations showed a hypochloroaemic alkalosis, marked hypercalcaemia and evidence of renal dysfunction. Calcium 4.09 mmol/l (normal 2.20-2.60); calcium (albumin adjusted) 3.84 mmol/l; chloride 85 mmol/l (normal 98-109); bicarbonate 38 mmol/l (normal 20-28); urea 24 mmol/l (normal 2.2-7.7); creatinine 290 μmol/l (normal 45-120); phosphate 1.07 mmol/l (normal 0.8-1.3). Blood pH was 7.55 (7.36-7.44) and urinary pH was 8. A diagnosis of the 'milk alkali syndrome' was made.

With rehydration and a low calcium diet the urea and bicarbonate became normal within a few days and by nine days the serum calcium, corrected for albumin, fell to within the normal range. With a corrected calcium of 2.59 mmol/l, phosphate of 1.17 mmol/l and creatinine of 137 μmol/l urinary excretion studies were undertaken. Results (normal range) were as follows: calcium/creatinine ratio 1.17 (0.10-0.45), calcium excretion 0.16 mmol/l of glomerular filtrate (0.01-0.05), phosphate excretion 0.18 mmol/l of glomerular filtrate (0.05-0.22), TmP/GFR 0.99 (0.71-1.36).8 These results suggest a degree of renal resistance to the action of parathyroid hormone. Other studies...
included normal thyroid function tests, serum cortisol, serum gastrin, serum magnesium and ultrafiltrable magnesium. Endoscopy with oesophageal biopsy and 24-hour oesophageal pH monitoring confirmed the presence of reflux oesophagitis and the absence of duodenal and gastric ulceration. A skeletal survey showed no evidence of hyperparathyroidism or metastatic calcification and thallium scintigraphy and computerized axial tomography showed no parathyroid adenoma. However, amino terminal parathyroid hormone was initially elevated at 1100 pg/ml (normal <120) when the calcium was 3.18 mmol/l (albumin 38 g/l). This level of parathyroid hormone is within the range (100–1350 pg/ml) normally seen in primary hyperparathyroidism.9

At 20 weeks the corrected calcium was 2.44 mmol/l and, using a calcium ion specific electrode, the ionized calcium was 1.27 mmol/l (normal range 1.15–1.27 mmol/l). The amino terminal parathyroid hormone remained elevated at 1000 pg/ml.

Alkaline phosphatase levels varied between 9 and 16 KA units/dl (normal range 3–13 KA units/dl) suggesting a degree of increased bone turnover in response to the high levels of parathyroid hormone.

By 38 weeks the calcium had fallen to the mid-normal range at 2.35 mmol/l as had the corrected calcium (2.29 mmol/l) and the ionized calcium (1.22 mmol/l). The amino terminal parathyroid hormone also fell to the mid-normal range. TmP/GFR (0.91), calcium excretion (0.02 mmol/l of glomerular filtrate) and phosphate excretion (0.13 mmol/l of glomerular filtrate) were normal.

Discussion

Hyperparathyroidism has been reported in association with the milk alkali syndrome and exploration of the neck undertaken with removal of an enlarged parathyroid gland.46 The differentiation between the two conditions has sometimes proven difficult.10 11 It would be expected that the development of assays for parathyroid hormone would allow a clear separation of cases of milk alkali syndrome from primary hyperparathyroidism. In the milk alkali syndrome the elevated calcium should be associated with suppression of parathyroid hormone. In hyperparathyroidism the parathyroid hormone should be elevated or inappropriately high in the face of the hypercalcaemia. However, the separation has not proven to be so clear cut. Elevated parathyroid hormone levels have occasionally been noted in the milk alkali syndrome10 12 and have led to confusion with primary hyperparathyroidism. In some cases surgical exploration of the parathyroid glands has been undertaken and no abnormality found.13 In other cases the elevated parathyroid hormone levels have resolved over a number of months.10–12 Unfortunately the assay used has sometimes been a C-terminal parathyroid hormone assay which can give falsely elevated values in renal failure as the biologically inactive C-terminal fragments are excreted via the kidney.13

There are a number of factors in the milk alkali syndrome which might stimulate parathyroid hormone release: (1) The milk alkali syndrome is a chronic condition and the severe hypercalcaemia and marked renal dysfunction seen at presentation in this case reflect an acute deterioration of a chronic state. At presentation the ionized calcium, given the extreme elevation in the total calcium, would not be depressed despite the marked alkalosis.14 However, for much of the course of the milk alkali syndrome a chronic alkalosis could produce a low ionized calcium even though the total calcium might be normal or slightly increased. Thus tetany is a recognized feature of the excessive oral intake of absorbable alkali.2 A chronic low ionized calcium would result in parathyroid gland stimulation and resultant hyperplasia analogous to the secondary hyperparathyroidism seen as a response to a low serum calcium in renal failure.15 Following correction of the alkalosis the hyperplastic parathyroid glands could continue to secrete an inappropriate amount of parathyroid hormone as parathyroid cells are not totally suppressible by serum calcium levels within the normal range.16 This would explain the delay in normalization of the parathyroid hormone seen in this case. An additional factor which could stimulate parathyroid gland hyperplasia is that the interaction of calcium with protein receptors on the parathyroid cells is likely to be a pH dependent process, as is the binding of calcium to other proteins. A chronic alkalosis might prevent the parathyroid cell from detecting the ionized calcium as efficiently as normal and adapting its level of secretion appropriately.

(2) Intake of calcium-containing absorbable alkalis is followed by neutralization of gastric acid and systemic alkalosis. This systemic alkalosis can produce a decrease of ionized calcium which, while still within the normal range, produces a doubling of parathyroid hormone secretion.17 18 Such swings in ionized calcium taking place several times per day in response to frequent self-medication with absorbable alkali could produce marked parathyroid gland stimulation without ionized calcium or total serum calcium being outside the normal range. Parathyroid gland hyperplasia is commonly seen in Wermer’s syndrome (multiple endocrine adenomatosis type 1).19 20 The increased gastric acid secretion secondary to the presence of non-beta cell tumours of the pancreas in this syndrome can also produce an alkaline tide in the blood.18 A similar systemic alkalosis is seen in the Zollinger-Ellison syndrome. Elevated parathyroid hormone levels have been found in the Zollinger-Ellison syn-
drome and these elevated levels have returned to normal following resection of the non-beta cell gastrin secreting pancreatic tumours.

(3) Other recognized effects of parathyroid hormone include promotion of bicarbonaturia, which would tend to limit any alkalosis present, and phosphate excretion. Phosphate intake is frequently high in these subjects due to their high milk intake although the syndrome may occur even when excessive amounts of milk are not consumed. The phosphaturic effect of parathyroid hormone promotes excretion of excess phosphate. Thus, while parathyroid gland stimulation in the milk alkali syndrome at first appears paradoxical there are a number of factors present which make it likely to occur.

It is interesting to note that in several of the reported cases of hyperparathyroidism and milk alkali syndrome occurring together the findings at surgery were of hyperplasia of several parathyroid glands or adenoma formation plus hyperplasia rather than solitary adenoma formation with suppression of the other parathyroid glands. We suggest that these cases may develop as a result of chronic parathyroid stimulation as discussed above and that cases of hyperplasia alone may resolve spontaneously over a number of months without recourse to surgery.

The original paper by Burnett over 35 years ago noted in Case 4 that there was 'parathyroid hyperplasia of the chief and oxyphil cell variety thought to be characteristic of secondary hyperparathyroidism'. We suggest that this description may well have been entirely appropriate.

Acknowledgements

We thank Dr B. Payne and Dr E. Will for helpful discussion, Dr I. Gunn for measurements of ionized calcium and the Supra-Regional Assay Service at the Middlesex Hospital, London for assays of amino terminal parathyroid hormone.

References

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