Rapid control of ectopic Cushing’s syndrome during the COVID-19 pandemic in a patient with chronic hypokalaemia

Ziadoon Faisal¹ and Miguel Debono ²

¹Department of General Medicine, Royal Victoria Infirmary, Newcastle Upon Tyne Hospitals NHS Foundation Trust, Newcastle Upon Tyne, UK and ²Department of Diabetes and Endocrinology, Royal Hallamshire Hospital, Sheffield Teaching Hospitals NHS Foundation Trust, Sheffield, UK

Summary

In this case report, we describe the management of a patient who was admitted with an ectopic ACTH syndrome during the COVID pandemic with new-onset type 2 diabetes, neutrophilia and unexplained hypokalaemia. These three findings when combined should alert physicians to the potential presence of Cushing’s syndrome (CS). On admission, a quick diagnosis of CS was made based on clinical and biochemical features and the patient was treated urgently using high dose oral metyrapone thus allowing delays in surgery and rapidly improving the patient’s clinical condition. This resulted in the treatment of hyperglycaemia, hypokalaemia and hypertension reducing cardiovascular risk and likely risk for infection. Observing COVID-19 pandemic international guidelines to treat patients with CS has shown to be effective and offers endocrinologists an option to manage these patients adequately in difficult times.

Learning points:

• This case report highlights the importance of having a low threshold for suspicion and investigation for Cushing’s syndrome in a patient with neutrophilia and hypokalaemia, recently diagnosed with type 2 diabetes especially in someone with catabolic features of the disease irrespective of losing weight.

• It also supports the use of alternative methods of approaching the diagnosis and treatment of Cushing’s syndrome during a pandemic as indicated by international protocols designed specifically for managing this condition during Covid-19.

Background

The COVID-19 pandemic has hugely influenced medical practice. To maintain social isolation and lockdown restrictions, authorities have modified standard protocols to reduce COVID-19 spread. A typical complex condition influenced by these changes has been Cushing’s syndrome (CS). We present a case of a patient with persisting neutrophilia and hypokalaemia diagnosed with ectopic ACTH syndrome from small cell lung cancer with sacral metastases. The patient had recently been diagnosed with type 2 diabetes; irrespective of clinical and biochemical features of CS, the diagnosis was delayed.

Supported by COVID-19 international protocol modifications, routine diagnostic testing to confirm ectopic CS was unnecessary and the patient was started on second-line treatment metyrapone immediately when serum cortisol was found to be 3333 nmol/L. This intervention achieved a rapid control of diabetes, normalization of neutrophils and potassium levels and a
reduction in cortisol levels with significant improvement in symptoms allowing delays in further Cushing’s treatment during the COVID pandemic.

Case presentation
A 65-year-old retired tarmac layer man presented to the emergency department with worsening mobility over the last 6 weeks. He reported unintentional weight loss of 8 kg over 8 weeks. He was independently mobile and lived with his wife prior to this. He was an ex-smoker (stopped 3 months previously). He had a background medical history of T2DM which had been diagnosed 4 months previously when he was referred to his GP. He also had a history of hypertension which had been difficult to control and a previous history of atrial fibrillation and ischaemic heart disease. His medications were atorvastatin, ramipril, doxazosin, gliclazide, metformin, bisoprolol and sublingual glyceryl trinitrate.

On examination, his temperature was 36.8°C, BP 178/115 mmHg, pulse rate 122 bpm, weight 89.6Kg and BMI 28.2kg/m². The patient had Cushingoid features, including a round face, interscapular adiposity, wasted upper and lower limbs proximally, thin skin, purple stria and central obesity. He had clinical evidence of bilateral proximal myopathy and multiple bruises were noted on his forearm and legs. Lower limb examination showed wasting of the quadriceps muscles with no fasciculation and knee flexion of 3/5 bilaterally. Sensory examination was normal. Rectal examination was normal as well.

Investigation
Initial investigations showed his WCC was slightly elevated at 11.8 × 10⁹ and neutrophils 10.4 × 10⁹ and potassium 3.2 mmol/L. Two months previously during admission, potassium levels varied between 2.2 to 3.4 mmol/L and neutrophils were 12.4 × 10⁹. COVID swabs were repetitively negative during the admission.

In view of the lower limb symptoms and signs, an urgent MRI of spine was requested on admission and this showed involvement of the sacrum by extensive tumour with the involvement of the exiting right and, to a lesser extent left, S2 nerve roots (Fig. 1).

Based on all these findings, CS was suspected. Baseline cortisol was 3333 nmol/L and ACTH level was 96 ng/L. Routine diagnostic testing, overnight dexamethasone suppression test and 24-h urine cortisol were unnecessary (1). CXR showed a suspicious lesion in the right lung and CT of thorax, abdomen and pelvis showed two irregular nodules in the lung’s right lower lobe which is highly suspicious for primary lung malignancy, with a pleural-based mass invading the chest wall consistent with metastasis. There was a 67 mm left adrenal gland metastases. Right adrenal gland was hyperplastic. An ultrasound-guided biopsy from the pleural mass showed an ACTH secreting metastatic small cell lung cancer confirmed by histology.

Treatment
Oral metyrapone 1000 mg three times per day was started, and within 24 h potassium levels went back to the normal range and blood glucose levels dropped significantly to an extent that gliclazide had to be stopped to avoid hypoglycaemia (Fig. 2). Blood pressure also improved to readings within normal (Table 1). Neutrophils normalized.

Cortisol levels improved significantly varying between 200 and 500 nmol/L initially on 1000 mg 8 h, with metyrapone then titrated up to 1000 mg 6 h followed by a block replacement regime adding on hydrocortisone 10 mg three times per day. He was also put on prophylaxis for Pneumocystis jirovecii with oral co-trimoxazole 960 mg/day in view of immunosuppression risk and a low-molecular-weight heparin dalteparin 5000 units subcutaneously daily in view of thromboembolism risk.

![Figure 1](https://edm.bioscientifica.com/)

**Figure 1**
Small cell lung metastasis to sacrum by extensive tumor, principally involving the vertebral bodies, but also extending into the pedicle and lamina of S2 (right), with expansion along the anterior margin of the sacrum into the presacral space, particularly on the right (left).
Outcome and follow-up

For the first few weeks, the patient felt better and was transferred to oncological care. Potassium and neutrophil counts remained within the normal range whilst glucose and BP were controlled. Unfortunately, due to the advanced metastatic small cell lung cancer, the patient passed away weeks later.

Discussion

The COVID-19 pandemic has hugely influenced medical practice. Following formal guidelines to attain perfection is not generally possible so authorities have designed more appropriate protocols to be used during the pandemic. These recommendations aim to maintain social isolation and lockdown restrictions by reducing out-patient visits, imaging, and biochemical testing. A typical condition influenced by these changes has been Cushing’s syndrome (CS). Patients with suspected or confirmed CS usually undergo complex investigations and management, therefore, active management has been recommended for only patients with clear discriminating features of Cushing’s and those with moderate to severe features. Importantly, those with diabetes and high blood pressure should be treated actively (1). In addition, in patients with severe clinical signs of CS, who are not stressed for example, septic, as opposed to carry out dexamethasone suppression tests and 24-h urine cortisol, a serum cortisol which is very high >1000 nmol/L is sufficient to make a diagnosis (1).

Treatment of CS is aimed to reduce its complications. As per Endocrine Society guidelines, the first line treatment of CS is surgical but, in some circumstances, if a patient is not fit for surgery or as part of preoperative preparation, medical treatment can be used first line. In view of COVID restrictions, surgery as a first-line therapeutic strategy as expected is not viable (2). Medical therapy, usually a second line option, could temporarily be used to provide rapid reduction in cortisol levels and allow a safe delay in surgery and possibly achieve good control of complications.

This case report highlights the importance of having a low threshold for suspicion and investigation for CS in a patient with chronic neutrophilia and hypokalaemia, recently diagnosed with type 2 diabetes especially in someone with catabolic features of the disease irrespective of losing weight. It also supports the use of alternative methods of approaching the diagnosis and treatment of CS during a pandemic.

A common cause for ectopic CS is small cell lung cancer (SCLC). CS is a clinical disorder resulting from high levels of circulating glucocorticoids, with concomitant loss of the negative feedback mechanism of cortisol on the hypothalamic-pituitary-adrenal axis (HPA) (3). Ectopic ACTH syndrome is responsible for one-fifth of the ACTH dependent CS, most arise from SCLC or bronchial carcinoid, though any endocrine tumor can cause ectopic ACTH CS (4).

CS can present clinically with a wide range of signs and symptoms including weight gain, lethargy, hirsutism, acne, purple abdominal stria, depression, and muscle weakness. Though the presentation can be nonspecific, certain clinical features are highly suggestive of CS such as proximal myopathy, easy bruising, facial plethora, and abdominal striae. In patients who are not phenotypically cushingoid (e.g. slim rather than overweight), the suspicion should be high for a malignancy-secreting ACTH, especially in the presence of pronounced myopathy, neutrophilia, and profound hypokalaemia (5).

Our patient presented with weight loss. Weight loss is atypical in CS with obesity present in around 85% of patients in Cushing’s disease (6), and the rest is usually overweight. Obesity is lower in ectopic ACTH syndrome occurring in 66% but only 10% lose weight so high
clinical suspicion is necessary (7). Importantly, one should recognize the effect of advanced malignancy as a cause of weight loss in some patients with ectopic ACTH syndrome and this confounds the classical clinical features of CS.

Two-thirds of patients with CS have diabetes and 50% have impaired glucose metabolism (8). In ectopic CS, diabetes is present in 59% (7). Hypertension is also common in CS. Hypertension is present in 68% of patients with Cushing’s disease (6), and 59% of patients with ectopic disease (7).

Both surgery and excess glucocorticoids are known to cause immunosuppression and increase the risk of thromboembolic events (1). So making a quick diagnosis and controlling glucocorticoid excess rapidly reduce the risks of such complications and prevents long periods of admission; this is ideal during the COVID-19 pandemic in a patient with CS.

Our patient presented with new-onset diabetes and this together with neutrophilia and hypokalaemia should alert physicians to the possible presence of hypercortisolism. Chronic neutrophilia with hypokalaemia is highly suggestive of hypercortisolism and this pair of findings with new-onset diabetes should alert the general physicians of the possibility of Cushing’s syndrome. When the patient presented to us with a cortisol level >1000 nmol/L and a high ACTH level with clear features of CS, we were able to make a rapid diagnosis supported by new guidelines "Endocrinology in the time of Covid-19: management of Cushing’s syndrome" (1) and avoid extra dexamethasone suppression testing and 24-h urine cortisol tests which would have delayed diagnosis confirmation and treatment.

Treatment options for CS as per Endocrine Society Guidelines recommendations include surgery as a first-line treatment for pituitary lesions responsible for Cushing’s disease, for ectopic ACTH-secreting tumors or for unilateral or bilateral adrenal tumors. During COVID, in view of social isolation restrictions to try and reduce exposure time to medical services especially in conditions where patients are immunosuppressed, surgery might not always be the best option. Medical treatments are possible though to try and achieve rapid control of the disease and its complications.

Metyrapone is one of the first-line options for medical treatment. Metyrapone works by reducing cortisol synthesis through inhibition of 11β hydroxylase, it also blocks 18 hydroxylase. Metyrapone is given orally and has a short half-life, so it should be given 3–4 times a day, and it is well tolerated (9). In one study looking at the short- and long-term response to metyrapone in patients with CS, 75% showed biochemical normalization in less than 4 months and 70–83% had long-term response (10). Metyrapone, in our case, helped to control hypokalaemia, hyperglycaemia and hypertension rapidly during the COVID-19 pandemic. This highlights the effectiveness of this drug in treating CS at least temporarily.

In this case report, we describe the management of a patient who was admitted with ectopic ACTH syndrome during the COVID pandemic with new-onset type 2 diabetes, neutrophilia and unexplained hypokalaemia. These three findings when combined should alert physicians to the potential presence of Cushing’s syndrome. On admission, a quick diagnosis of CS was made based on clinical and biochemical features and patient was treated urgently using high dose metyrapone thus allowing delays in surgery and rapidly improving the patient’s clinical condition. This resulted in the treatment of hyperglycaemia, hypokalaemia and hypertension reducing cardiovascular risk and likely risk for infection. Following guidelines to treat patients with CS have shown to be effective and offer endocrinologists an option to manage these patients adequately in difficult times.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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Patient consent
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M D and Z F managed patient’s clinical care, obtained consent and designed the manuscript.

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