Cyclic Cushing’s disease with misleading inferior petrosal sinus sampling results during a trough phase

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Diagnosing Cushing’s syndrome is challenging and is further hampered when investigations are performed in a patient with cyclic Cushing’s syndrome. A subset of patients with Cushing’s syndrome exhibit periods of abnormal cortisol secretion with interspersed normal secretion. Patients can have periods of clinical improvement during these quiescent phases or remain symptomatic. Initial diagnostic testing can be challenging because of the unpredictable durations of the peak and trough phases, and it is especially challenging when the diagnosis of cyclic Cushing’s syndrome has not yet been determined. Here, the authors present the case of a patient with Cushing’s disease with a pathology-proven adrenocorticotropic hormone (ACTH)–secreting pituitary adenoma and whose initial inferior petrosal sinus sampling (IPSS) results were deemed indeterminate; further studies elucidated the diagnosis of cyclic Cushing’s syndrome. Repeat IPSS was diagnostic of a central source for ACTH secretion, and the patient was treated successfully with transsphenoidal resection. Literature concerning the diagnosis and management of cyclic Cushing’s syndrome is also reviewed.


KEY WORDS cyclic Cushing’s disease; diagnosis; pitfalls

Case Report

A 63-year-old woman with a medical history of resistant hypertension presented with complaints of intermittent weight gain, bruising, progressive fatigue, and emotional lability. On examination, she was noted to have classic cushingoid features with moon facies, plethora, central obesity, upper arm and leg muscle atrophy, and bruises. Her blood pressure was 110/74 mm Hg, her weight was 180 lbs, her height was 61 inches, and her body mass index was 34 kg/m². The neurological examination revealed that she had an inability to rise from a squatting position.

Investigations

Biochemistry. Two 24-hour urinary free cortisol (UFC) tests were completed and revealed elevated levels of 225 and 175 μg/24 hours (reference range < 50 μg/24
hours). Her plasma ACTH level was 26 pg/ml (reference range 5–27 pg/ml). A 1-mg overnight dexamethasone suppression test revealed a morning serum cortisol concentration of 22 µg/dl (reference range < 1.8 µg/dl), and a low-dose 2-day (2 mg/day) dexamethasone suppression test revealed 24-hour UFC levels that were suppressed from 48 to 38 µg/24 hours (reference range > 50% suppression). A high-dose (8-mg) overnight dexamethasone suppression test revealed a pretreatment serum cortisol level of 21 µg/dl, which was suppressed to 5.4 µg/dl after dexamethasone administration. Her insulin-like growth factor–1 level was 76 ng/ml, her prolactin level was 8.3 ng/ml, her free thyroxine level was 1.3 ng/dl, her thyroid-stimulating hormone level was 0.7 mIU/L, her luteinizing hormone level was < 0.2 IU, and her follicle-stimulating hormone level was 0.7 mIU/L, her luteinizing hormone level was < 0.2 IU, and her follicle-stimulating hormone level was < 0.7 IU (inappropriately suppressed for a postmenopausal woman). The results of an oral glucose-tolerance test were consistent with impaired glucose tolerance.

Imaging. Pituitary MRI was suggestive of a small central hypointense lesion. A dual-energy x-ray absorptiometry scan confirmed osteoporosis.

Follow-Up

With a diagnosis of ACTH-dependent CS, the patient underwent the initial inferior petrosal sinus sampling (IPSS) (Table 1), 7 months after her presentation and initial workup from her referring endocrinologist. The IPSS results revealed a < 2:1 central-to-peripheral gradient at baseline, with a < 3:1 gradient of central-to-peripheral ACTH in the petrosal sinus samples after corticotropin-releasing hormone (CRH) administration. An isolated central-to-peripheral gradient was seen (> 3:1) in samples obtained from the jugular vein, but it was believed to be inconsistent with petrosal samples and nondiagnostic. Without a clear central-to-peripheral gradient between the petrosal samples and the peripheral measurements, the IPSS results were considered indeterminate and not supportive of a diagnosis of central ACTH-dependent Cushing’s disease.

No additional samples for 24-hour UFC testing had been collected since the time of the patient’s initial presentation. At that time, she reported that her symptoms associated with CS had resolved in the weeks leading up to the IPSS procedure. Her facial plethora and weakness had improved dramatically. It was decided to reevaluate the patient with further testing before proceeding with further investigation of the source of the ACTH. Serial samples for 24-hour UFC measurements (Fig. 1) were then collected for 11 months to document cortisol hypersecretion. A second IPSS procedure was scheduled during a cortisol hypersecretory phase (Table 2). The second IPSS procedure was diagnostic of a central source for ACTH, which supported a diagnosis of Cushing’s disease. The peaks and troughs revealed by serial measurements of 24-hour UFC levels suggested a diagnosis of cyclic Cushing’s disease.

Treatment and Posttreatment Course

The patient underwent transsphenoidal surgery, with removal of a microadenoma and the inferior central portion of the pituitary gland. Pathology confirmed a sparsely granulated corticotroph adenoma (Ki 67 < 1%) that was p53 negative. Postoperatively, her serum cortisol concentration reached a nadir of 1.1 µg/dl on Postoperative Day 2, falling from 9.6 µg/dl on Postoperative Day 1. She was discharged from the hospital after being placed on a hydrocortisone replacement regimen and reported symptoms of steroid withdrawal in the perioperative time period. She lost 8 lbs, complained of joint aches, and experienced resolution of her hypertension without taking her antihypertensive medication. Her recovery was otherwise uneventful, and she returned to her referring endocrinologist for long-term follow-up.

Discussion

This case clearly documents several peaks and troughs of cortisol secretion as measured by frequent 24-hour sample collections for UFC testing. The results of IPSS performed during a trough phase were more consistent with a peripheral source of ACTH, given the lack of gradient between the petrosal and peripheral samples at baseline and after CRH administration, but were indeterminate at the time and required further investigation. Although it was not recognized at the time, one potential explanation for the results was that the patient had cyclic CS and was studied during a trough phase. Another potential source for error was improper cannulation of the petrosal sinuses leading to the errant measurement of ACTH levels. Both of these situations necessitate repeat performance of IPSS. The correct diagnosis was made by restudying the patient after demonstrating periodicity of cortisol secretion and ensuring that the IPSS was performed during a phase of active cortisol secretion.

CS is an important clinical entity with several deleterious effects on health, including dyslipidemia, impaired glucose tolerance, hypertension, bone loss, proximal muscle wasting, weight gain, and mood disturbances. In classic CS, cortisol levels are elevated without diurnal varia-

### Table 1. Initial IPSS

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Petrosal ACTH (pg/ml)</th>
<th>Peripheral ACTH (pg/ml)</th>
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<tr>
<td>Time (mins)</td>
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<td>Rt</td>
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<tr>
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<td></td>
</tr>
<tr>
<td>Lt</td>
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<td></td>
</tr>
<tr>
<td>15</td>
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<tr>
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<tr>
<td>19</td>
<td>112</td>
<td></td>
</tr>
<tr>
<td>Inferior Vena Cava Infra-adrenal (pg/ml)</td>
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<td></td>
</tr>
<tr>
<td>20</td>
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</table>
Cyclic Cushing's disease with misleading IPSS results

Cyclic CS is a rare subtype of CS in which periods of biochemical and/or clinical hypercortisolism alternate with periods of eucortisolism. A variety of subcategories have been described.10 Cyclic patterns of biochemical hypercortisolism may be regular or irregular and have variable or constant clinical manifestations. Cycle durations have also been noted to vary between weeks and years.8 Although it is rarely diagnosed, cyclic CS may in fact be more frequent than commonly thought. Indeed, a retrospective study of 201 patients with CS found cyclic variability in 30 (15%) of them.1

Difficulties in diagnosis underlie the underreporting of this phenomenon. As demonstrated in our case report, capturing a period of cortisol excess and confirming the diagnosis during this time period may be challenging. A generally accepted diagnostic criterion for cyclic CS is the recognition of 2 cycles (3 peaks and 2 troughs) of cortisol production,3 although others consider 1 complete cycle (2 peaks and 1 trough) sufficient for making the diagnosis.1

Any etiology of CS can produce cyclic patterns of hypercortisolism. In a review of 65 published cases of cyclic CS, 54% were caused by ACTH-producing pituitary adenomas, 26% were caused by ectopic ACTH production, and 11% were a result of ACTH-independent causes.8 The pathophysiology has yet to be defined clearly, although some theories have been proposed. A histological finding of necrosis within tumors led some investigators to propose that episodic hemorrhage and necrosis of cortisol-producing cells causes a cyclical release of cortisol,7 although others have identified cases of cyclical CS without necrosis in the tumor specimen.2 Another theory posulates that tumor calcifications lead to infarction with subsequent eucortisolemia;4 however, this theory does not explain the periodicity. A third theory involves potential hypothalamic causes as mediators of pulsatile ACTH hypersecretion. Neurotransmitters thought to be involved include CRH, noradrenaline, dopamine, acetylcholine, and γ-aminobutyric acid.8

Diagnosis of typical CS is known to be challenging because no single test has ideal diagnostic accuracy, which results in the need for multiple tests to confirm a diagnosis. According to Endocrine Society practice guidelines, screening tests include 24-hour UFC measurement, midnight salivary cortisol measurement, and the 1-mg dexamethasone suppression test, with a note that urinary and salivary levels should be measured at least twice given the
variability in results. A diagnosis of cyclical CS is complicated further by the relapse and remission of biochemical and clinical findings and the need for close monitoring and coordination of confirmatory tests. Furthermore, an increased duration of testing is required given the varying periods of remission, which are known to last up to 5 years. This poses a major obstacle in the ability to administer appropriate management in a timely fashion.

Novel diagnostic methods are being explored to circumvent this pitfall. Hair cortisol levels are known to be significantly elevated in patients with CS over those in the general population (sensitivity 86%, specificity 98%). The advantage of using hair cortisol levels is that they can provide data over the previous several months in one analytical setting. In one study, hair samples were obtained from one lock of hair each from healthy controls, patients with CS, and patients with cyclic CS. Cortisol measurements at 1-cm intervals correspond to monthly time frames. In patients with cyclic CS, hair cortisol-level fluctuations were found to correlate with the clinical course of CS. In another report, a desmopressin acetate (DDAVP) stimulation test was used for a case in which biochemical evidence was not confirmed, resulting in a 2-year delay of the eventual discovery of an ACTH-producing pituitary adenoma. Thus, both the analysis of hair cortisol and the DDAVP stimulation test may improve on the diagnostic approach by significantly shortening the time to diagnosis and enabling more timely management.

As with classic CS, the treatment of cyclical CS generally involves surgical removal of the tumor or, for patients who are not candidates for surgery, medical management. However, posttreatment management poses a unique set of challenges in these patients. Patients with cyclic CS caused by pituitary disease have a significantly higher recurrence rate postoperatively (63%) and lower cure rates (25%) than those of patients with noncyclical CS (approximately 80% cure rate). Moreover, postoperative management is complicated by the fact that normal cortisol levels in these patients may not be indicative of cure but instead correspond to a period of quiescence. In this situation, there is a need to closely monitor for longer periods of time, and perhaps more frequently than usual, to ensure the capture of a relapse if it occurs. It was fortunate that, in our case, the postoperative symptoms of cortisol withdrawal and the need for steroid replacement were strongly indicative of remission in this patient.

We report here a case that clearly documents multiple peaks and troughs of cortisol secretion, as measured by frequent sampling of 24-hour UFC levels. The IPSS performed during a trough phase provided misleading results because of the lack of gradient at baseline and after CRH administration. The correct diagnosis was made by repeating the IPSS and ensuring that it was performed during an active phase of cortisol secretion. It is important to recognize cyclic CS as a clinical entity and to note that its prevalence may be higher than currently believed given the challenges in establishing a diagnosis. A diagnosis of cyclic CS is made by showing peaks and troughs of cortisol secretion either by frequent measurement of cortisol levels in urine or saliva or perhaps by using novel techniques such as the analysis of hair cortisol levels or DDAVP stimulation testing. As illustrated by this case, the timing of IPSS (i.e., during a period of quiescence) may lead to an incorrect diagnosis and further delay appropriate management. It is important that testing for the differential diagnosis be done only during periods of biochemical or clinical hypercortisolism. Management of these patients is complicated by high recurrence rates and low cure rates, which necessitate the need for continued closer monitoring for longer durations of time than for patients with noncyclical CS.

References


Author Contributions

Conception and design: Carmichael. Acquisition of data: Carmichael, Bonert. Analysis and interpretation of data: Carmichael, Bonert. Drafting the article: all authors. Critically revising the article: all authors. Reviewed submitted version of manuscript: all authors. Approved the final version of the manuscript on behalf of all authors: Carmichael.

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