Optimal management of Cushing syndrome

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Abstract: Cushing syndrome (CS) caused by endogenous hypercortisolism is a diagnostic challenge. The most common cause is Cushing disease. Surgical treatment is the first-line therapy for Cushing disease. However, due to the often clinical instability of the patient’s condition, which needs acute treatment of hypercortisolism or inoperable tumors, initial surgery is often not possible. It is therefore important to provide appropriate initial medical treatment. Following surgery, the patient needs to be evaluated and confirmed for disease resolution based on standard criteria, and treated with appropriate supportive measures for the rest of life if necessary. This article reviews the current data and treatment options for Cushing syndrome and proposes a therapeutic algorithm for its optimal management.

Keywords: Cushing syndrome, Cushing disease, Hypercortisolism

Introduction

Cushing syndrome (CS) occurs as a result of long and chronic exposure to glucocorticoids (GCs). The most common cause is exogenous CS secondary to GC intake.1,2 Endogenous CS is more common in women. Its occurrence is from about 0.7 to 2.4 per 1 million population by year.3

GCs are synthetic steroids that are effective in the management of inflammatory, allergic, hematologic, and autoimmune processes, among others. Due to their remarkable anti-inflammatory effect and secondary pain reduction with rapid improvement of symptoms, the use of these drugs leads the patient to an indiscriminate intake and thus to side effects.4,5

Patients with noncontrolled CS have a five-fold increase in mortality.6 The screening studies performed in diabetic patients, especially in those diabetic patients with difficult control and with obesity and hypertension, suggest that it has a prevalence of 2%–5%.7

The present review aims to expose the “optimal” treatment options for CS, briefly stating the causes and offering a description of CS management induced by GCs.

Endogenous CS occurs in 80%–85% of the cases due to corticotropin (ACTH)-dependent causes: 80% is due to pituitary tumors (Cushing disease [CD]), and the remaining 20% is secondary to ectopic secretion of ACTH, which usually derives from carcinoid tumors or small-cell lung carcinoma.

ACTH-independent CS occurs in 15%–20% of cases, and most of these are caused by an adrenal tumor (adenoma or cancer, 75%) or, more rarely, macronodular adrenal hyperplasia and pigmented nodular adrenal disease.1,6,8–10 The diagnosis of endogenous
CS is a challenge; thus, this review does not cover such a target and is reserved only for treatment. For diagnosis it is worth consulting clinical guides.\textsuperscript{11}

**Exogenous CS and its treatment**

Exogenous CS occurs due to an excess of and chronic exposure to synthetic GCs, which suppress the synthesis and secretion of hypothalamic ACTH-releasing factor and therefore of ACTH, producing bilateral atrophy of adrenal glands and a decrease of endogenous cortisol.\textsuperscript{12} The use of prednisone at high doses over 5 days diminishes the weight of the adrenal gland, although it depends upon the dose and overall time in which it has been used. Prednisone administration from 20 mg to 30 mg over 7 days or more inhibits the hypothalamic-pituitary-adrenal axis (HPA). At a low dose (less than 7.5 mg/day), it may suppress the HPA axis in a 1-month interval.\textsuperscript{13,14}

Recovery of the HPA axis after suspension of the medicament may follow four stages: (1) stage I: a month after suspending GC, the ACTH is low and the plasmatic and urinary 17-hydroxycorticosteroids (17OHC) are low; (2) stage II: from 2 to 5 months after suspension of the medicament, the ACTH is normal but 17-OHC is low and without incurring response to ACTH stimulation; (3) stage III: from 6 to 9 months after suspension, the ACTH is normal and the plasmatic and urinary 17-OHC are normal but incurring an abnormal adrenal response to ACTH stimulation; and (4) stage IV: from 9 to 12 months after suspension, a complete recovery of the HPA axis is achieved.\textsuperscript{14,15} LaRochelle et al\textsuperscript{16} found in patients with rheumatologic disorders and receiving low doses of prednisone that lower doses of 5 mg per day had a normal HPA axis independently from evolution time, while those with a higher dose of 5 mg per day had a greater inhibition frequency of the HPA axis.

The diagnosis is established with the clinical manifestations, in addition to the background of GC therapy. Later on, suppressed endogenous cortisol levels and low ACTH levels are expected to be found.\textsuperscript{4,11} The specific clinical features of hypercortisolism may differ in terms of intensity depending upon the CS cause, either endogenous or due to GC intake.\textsuperscript{17}

The use of GC is very common; nonetheless, there is not enough evidence detailing suspension or withdrawal of steroids.\textsuperscript{18} The aim for decreasing or suspending GC is firstly based on achieving a GC decrease, which has been used in physiologic doses, and at the same time preventing the patient from onset of a suprarenal crisis. An option is decreasing the dose 10\%-20\% every 2 weeks. Therefore, as a first step after reduction of the steroid, we propose giving physiological doses (15–20 mg/day of hydrocortisone or 5–7.5 mg/day of prednisone).\textsuperscript{19} Afterwards, changing to an early morning dose of hydrocortisone or prednisone or even one dose every 48 hours allows a gradual recovery of HPA axis.\textsuperscript{20}

The greater risk of suspending GC consists of an adrenal crisis development, or by avoiding it by means of a gradual dose decrease of GC the risk will be the exacerbation of the underlying disease that in the first instance forced the prescription of GC.

The next stage is measuring the early morning serum cortisol levels (8:00 AM), wherein a serum level less than 3 mcg/dL would indicate cortisol deficiency rather than HPA axis recovery, being necessary to continue with replacement therapy. A serum cortisol level between 3 mcg/dL and 20 mcg/dL indicates that the patient has a basal cortisol secretion but lacks the capability of responding to physiological stress. When this occurs, an ACTH (Synacthen) stimulation test is recommended, or an insulin tolerance test for examining the capability response of adrenal glands and thus from the HPA axis. If serum cortisol levels are greater than 20 mcg/dL, then it can be assumed that the patient has recovered the HPA axis and therefore GC therapy can be withdrawn.\textsuperscript{19}

We will describe here the endogenous CS-specific treatment and its several causes, as well as the remission criteria for hypercortisolism.

**Surgical treatment of CS**

Because the main cause of endogenous hypercortisolism is ACTH-dependent (80\%-85\%), which is the most common cause of CD (hypophyseal hypersecretion by a corticotropic adenoma), we describe here some surgical treatment considerations.\textsuperscript{21,22}

**Transsphenoidal surgery**

For more than 3 decades transsphenoidal surgery (TSS) has been the best option for patients suffering from CD. Adenectomy was disclosed and performed by Hardy in the 1960s.\textsuperscript{23} Since then the technique has been improved, and selective adenomectomies have been possible. A problem we face is that many centers discuss their experiences in-house without satisfactorily documenting the healing and recurrence rates, although many of these reports show similar percentages.

Some classic studies that were performed in the 1960s report a remission rate within an interval from 50\%-90\%.\textsuperscript{25-27} After TSS, the overall recurrence rate
(micro- or macroadenoma) can be considered to be from about 3% to 17% and mortality up to 1.9%.21

Only 5%–20% of ACTH-producing adenomas are macroadenomas (>10 mm). Thus, the healing and recurrence may be due to this important factor. Most macroadenomas have a bad postsurgical prognosis. Some studies have disclosed remission ranging from 53% to 68%,28 and others point out lower rates of 33%.29 Fomekong et al28 report remission up to 92%, explaining that a noninvasive tumor of the cavernous sinuses with a size less than 15 mm may be a factor for an improved prognosis. Likewise, Rees et al30 indicate that tumor size has an influence as a healing predictor, independently of the ACTH serum levels. Another factor that appears to confer a favorable prognosis, in addition to adenoma characterization by means of image studies, is histologic identification by immuno-staining of ACTH.31 This makes sense because postsurgical identification of the said tumor increases remission probability.

Apparently, if we compare the type of surgical technique used, namely adenectomy (extraction of the adenoma only) versus hypophysectomy (resection of almost all the pituitary gland), there is no change regarding remission, but hypophyseal failure is more frequent in hypophysectomy (from 79% to 95%).21 The extent of hypopituitarism is 48%, excluding isolated ACTH deficiency, requiring up to 63% of the patients to temporarily receive hormone replacement therapy including GCs.32

According to different series, recurrence fluctuates between 3% and 17%, with an average of 9.4% and with variable intervals between 1.5 years and 8 years from the first surgery. Nevertheless, recurrences up to 12.5 years have been disclosed.31 In a recent report, overall recurrence rate was 12% at 4.2 ± 0.5 years after surgery.33 Remission rates after TSS are variable and depend principally upon tumor extent, the experience of the neurosurgeon, and the used biochemical criterion for its definition. Recently, it has been pointed out that postsurgical cortisol less than 2 µg/dL predicts long-term remission after TSS in CD.33 Remission in those with cortisol values 2–4.9 µg/dL at 3–5 days after surgery suggests that these patients do not require immediate surgical reintervention.33 However, all patients require long-term clinical follow-up because no cortisol limit value excludes patients from recurrence. With the aim of obtaining a more objective cortisol serum level and thus define remission of CD, we recommend the use of sensitive tests for postsurgical serum cortisol measurement.

Current standards of follow-up after hypophysis surgery in patients with CD consider obtaining a morning serum cortisol level < 5 µg/dL as the best criterion for defining remission.1,2,34 A lower level does not necessarily relate to lower rates of recurrence. We suggest using urinary free cortisol (UFC) or morning serum cortisol suppression (<2.4 µg/dL) after 1 mg of dexamethasone the previous night. UFC levels with a value less than 20 µg/24 hours (by means of ultrasensitive tests) predict remission, while normal or greater values are mistaken to suggest persistence. Following hypophysis surgery it will be necessary to regularly monitor all patients with CD, because there is no absolute cortisol value that predicts remission in 100% of cases.

Several methods have been used for evaluating prolonged remission of CD, namely overnight dexamethasone suppression test with a 1 mg dose and UFC and ACTH levels. Chen et al35 point out that less than 3 µg/dL of serum cortisol at day 3 after a 1 mg dose of dexamethasone overnight predicts 93% remission of the patients.

If there are no remission criteria and the surgical procedure was unsuccessful, treatment options include: (1) surgical reintervention, (2) radiotherapy (RT), (3) pharmacological treatment, (4) bilateral adrenalectomy (BA), and (5) a combination thereof. We recommend follow-up remission and regular base monitoring for life.

Mortality for TSS is 0.7%, with complications up to 42% disclosed. Thirty percent of the patients may present one or more postsurgical complications. Diabetes insipidus and hydroelectrolytic disorders (from 10%–15% and from 8.8%–12.5%, respectively) are the two most common transitory complications.36,37

Patients with corticotrophic adenomas appear to have more postsurgical complications of bleeding or neurological disorders compared with other hypophyseal tumors (5.6% vs 2.6%, respectively), showing greater risk within the first 3 days after surgery; thus, monitoring should be emphasized during this time by early identification of disturbances impacting final prognosis.36 Other reported complications are thromboembolic disease, infection, transient paralysis of cranial nerves III, IV, and VI, and epistaxis, all with a range less than 1%. A direct relationship between the number of annual procedures and the number of complications has been noted. Based on the aforementioned, an experienced surgeon has a greater rate of success and fewer complications.37

Following an unsuccessful first surgery, early reintervention within the first 60 days has a 38%–67% possibility of success.31 A second intervention increases the risk of complications, mainly cerebrospinal fluid fistula and hypopituitarism.31,38 Probability of remission after a second
intervention decreases by 50% if the remaining tumor invades the cavernous sinuses.\textsuperscript{39}

**Bilateral adrenalectomy**

BA for CD treatment began in the 1950s and is currently used in refractory cases to other treatments or when RT is not used, as in the case of young patients who wish to preserve fertility. A remission rate from 80%–100%\textsuperscript{23} is achieved by BA in CD, but at the same time it is a risk factor for the development of Nelson syndrome (NS), which occurs in 8%–38% of the cases.\textsuperscript{40} The development of NS is greater if BA is performed in young patients with imaged-confirmed adenomas before surgical treatment.\textsuperscript{41} Prophylactic RT decreases the risk of developing such syndromes; nevertheless, it is usually the chosen treatment for hypophyseal tumor remnant.\textsuperscript{42} It takes from about 5–10 years to develop NS after BA.\textsuperscript{43,44} Morbidity of 9.8% and mortality of 1.1% at 30 days after adrenalectomy are reported,\textsuperscript{45} although these may be greater in the presence of comorbidities that are present per se in patients with CS, such as obesity, susceptibility to infections, and increased risk of thromboembolism, as compared with adrenalectomy performed for other reasons.\textsuperscript{46}

It is possible that part of the adrenal function may be preserved following surgery, as adrenal remnants have been reported in 13%–27% of BAs carried out,\textsuperscript{47,48} this being a cause of disease persistence. Another noteworthy complication is deterioration of life quality due to depressive syndrome in varying degrees, fatigue being the most common symptom (85%).\textsuperscript{47} Undoubtedly, a high percentage of these patients will require chronic treatment with GCs and mineralocorticoids, thus partly improving alterations in life quality.

The treatment of choice for those patients diagnosed with CS independently from ACTH is adrenalectomy, and this can be unilateral in the case of patients with adenoma or carcinoma, or bilateral in hyperplasia cases. Adenomas have a healing rate of 100% with low morbidity and mortality when under expert management.\textsuperscript{49,50} Adrenal cancer prognosis is grim if there is recurrence of liver, thoracic, or loco-regional metastases.

**Radiotherapy**

RT was the first-line treatment used for CD from 1940 through to 1980, providing reduced healing rates lower than 50%;\textsuperscript{51} currently, it is considered as a second-line treatment after surgery failure. If used this way, the achieved remission is from 53% to 83% with a recurrence of up to 17%.\textsuperscript{52-54}

Different techniques have been applied. The use of a gamma knife has seen remissions reported from 63% to 73%; by using the lineal accelerator, data become less accurate but apparently with good results.\textsuperscript{55} If serious medical complications arise that increase anesthetic risk and therefore morbidity/mortality, then RT may be considered a first-line therapy. RT may be considered as the first-line therapy in children with remission rates similar to TSS. Nevertheless, the maximum effectiveness of RT may take as long as 1 year to be reached. Pharmacological treatment must be considered together with RT if hypercortisolism clinical manifestations are very pronounced. Growth hormone deficiency is the most common complication of RT with 50%, followed by the appearance of hypogonadism to a lesser extent; damage to the optical nerve is less than 1%.

The primary treatment of ectopic ACTH-dependent CS is the removal of the primary tumor, with which remission is achieved in 30%–47% of the cases.\textsuperscript{8} The probability of success depends upon tumor resectability and the expertise of the surgeon. However, if curative surgery fails or is not possible, then palliative treatment aims to reduce cortisol levels. BA is used in 56% of the hypercortisolemia cases that cannot be treated by other means.\textsuperscript{56} Recent advances in surgical techniques identify laparoscopic BA as an attractive alternative.\textsuperscript{57}

**Pharmacologic treatment for CS**

The principal treatment for CS’s most frequent causes is the excision of ACTH-producing tumors or adrenal adenomas; nevertheless, pharmacologic treatment has a well-established role. Indications of medical treatment for CD include preoperative treatment and persistent or recurrent hypercortisolism after hypophyseal surgery and for overcoming the period until RT becomes effective. Primary medical treatment may be considered in patients with no visible hypophyseal adenoma, in patients with high surgical risk and/or significant comorbidities, and in patients with low probabilities of resolution through surgery.\textsuperscript{58,59}

In the event of ectopic secretion of ACTH, patients may require medical treatment in order to lessen the impact of comorbidities and complications associated with hypercortisolism previous to a surgical intervention; furthermore, it can be used in such cases in which there has not been a conclusion on the source of the ectopic secretion and in patients with metastatic disease and ectopic production of ACTH without possible surgery resolution. Some patients may require continued therapy in the event of acute complications (psychotic break, hypertensive crisis).

Patients with unresectable adrenal carcinoma or metastasis of cortisol-producing adrenal carcinoma must be
treated with agents that decrease the cortisol production rate jointly with specific tumor therapy for treating symptoms and preventing complication from hypercortisolism.  

There are pharmacologic therapies that inhibit adrenal synthesis of GCs as well as therapies intended for decreasing ACTH secretion at a pituitary level or GC receptor antagonists. We will describe several drugs with pharmacological action at the aforementioned levels.

Adrenal steroid synthesis inhibitors

Ketoconazole

Since 1983, Engelhardt and Weber reported the usefulness of ketoconazole in a case of CS produced by an adrenocortical adenoma; afterwards, they recorded its inhibitory effect over adrenal enzymes 17, 20 desmolase (conversion of 17, 20 hydroxiprogesterone to androstendione), clarifying that the most powerful effect is here and to a lesser extent 17α-hydroxylase (conversion of progesterone to 17α-hydroxiprogesterone), 11β-hydroxylase (11-desoxicortisol to cortisol), and 16α- and 18-hydroxilase. Both in vivo and in vitro inhibition of androgens is greater than that of cortisol (30%–50% vs 19%, respectively). Ketoconazole also inhibits ACTH production and cellular growth, partly because of apoptosis induction. At a dose of 400–1200 mg/day it can decrease cortisol production in patients with CS from various etiologies. It can be administered two to three times a day, starting with 200 mg twice a day and being titrated up to a maximal dose of 1200 mg per day; therapy can be based on serum cortisol levels and UFC at 24 hours. Blood pressure in patients with CS may be better controlled if antihypertensive drugs are administered jointly with ketoconazole in comparison with antihypertensive drugs alone.

An adverse effect is the increment on serum concentration of liver enzymes in 5%–15% of the patients; additionally, it can cause male hypogonadism, gynecomastia (13%), gastrointestinal disorders (8%), edema (6%), and skin rash (2%).

Reports about paraneoplastic CS indicate complete hormonal response in up to 28% of the cases, leading to symptomatic hypoadrenalism in 12% of the cases. No prospective studies have been conducted based on ketoconazole monotherapy. The efficacy data have been drawn from retrospective studies, most of them with a small number of patients. Castinetti et al reported that ketoconazole induced biochemical remission in 50% of the patients with CD. Taking into account most of the trials as a whole, ketoconazole has an efficacy of 70%, and the remission rates in CD vary from 25% to 93%.

Ketoconazole has also been used in some cases of CS during pregnancy, having as side effects intrauterine growth retardation and potential antiandrogenic effects. Embryotoxic and teratogenic potential in animals has been reported. A patient with CD who refused surgical treatment had a normal birth through vaginal delivery at 37 weeks of gestation. We do not recommend the use of ketoconazole during pregnancy, as there is limited experience.

Metirapone

Metirapone it is a selective inhibitor of 11β-hydroxylase. It decreases cortisol and aldosterone synthesis, but the highest concentrations of 11-deoxycortisol preserve the mineralocorticoid functions. The maximal suppression of steroidogenesis requires the use of a 4 g/day dose. Electrolytic balance and blood pressure vary from patient to patient, depending upon the extent of aldosterone inhibition and stimulation of 11-deoxycorticosterone.

Cortisol suppression is dose-dependent within a few hours with a dose of 4.5 g/day. The dose varies from 500 mg/day to 6000 mg/day; patients must be carefully monitored due to potential development of hypercortisolism, being able to maintain chronic suppression with doses from 500 mg/day to 2000 mg/day. It may be used in patients with severe secondary hypercortisolism to ectopic cortical adrenocarcinoma or inoperable ectopic production of ACTH, as well as adjuvant in a case of surgical failure due to CD. In such a situation, a dose 500–750 mg three to four times daily is used.

The compensatory increase in ACTH production due to a decrease in negative feedback in hypophysial adenoma is a potential disadvantage in the treatment of CD. This in turn may lead to an increase in cortisol adrenal production, as well as androgens and mineralocorticoid precursors. Long-term administration creates hirsutism and hypertension. Other adverse effects include nausea, headache, sedation, and skin rash. A secondary increase of androgens is more likely in ACTH-dependent cases; meanwhile, hypertension and hypokalemia are less probable in ACTH-independent cases.

Etomidate

Etomidate it is a hypnotic anesthetic agent that suppresses steroidogenesis by inhibiting the cleavage of the side chain of cholesterol and 17-hydroxylase, 11-β hydroxylase, and 17–20 lyase enzymes. It has a rapid onset of action, and it is particularly useful in patients with acute complications that compromise life, such as psychosis and severe hypertension. Most of the time it is used in patients with ectopic production...
of ACTH and excessive production of cortisol. It can be administered parenterally in a dose between 0.03 mg/kg/hour and 0.3 mg/kg/hour in continuous infusions, achieving a cortisol decrease in 11–24 hours.\textsuperscript{74}

The use of etomidate has been reported via infusion at a dose of 3.0 mg/hour in pediatric patients presenting psychotic episodes, in addition to simultaneous intake of hydrocortisone, maintaining a stable serum cortisol and remitting the acute condition.\textsuperscript{75,76}

Adverse effects include sedation, hypotension, hypertension, and bradycardia.\textsuperscript{62} It has been associated with increased mortality in critically ill patients by causing acute adrenal insufficiency.\textsuperscript{74,77}

We suggest using it only in cases of complicated hypercorticism, in those who require short-term improvement, and in those who cannot use the oral route. Etomidate is recommended as a temporary treatment prior to definitive therapy.

Mitotane
Mitotane is a selective adrenolytic agent of adrenocortical cells, mainly used for adrenal carcinoma treatment. In addition to cellular growth suppression, mitotane inhibits the cleavage of the side chain of cholesterol and 11β-hydroxylase, leading to a decreased production of cortisol.\textsuperscript{78}

Treatment must continue for at least 3 months; if favorable effects are observed, the treatment continues in definite form. It should not be administered concomitantly with spironolactone, as it interferes with adrenal suppression produced by mitotane.\textsuperscript{78} At low doses (2–4 g/day), it has a lower adrenolytic effect within the glomerular zone, and it is possible to have less suppressed production of aldosterone.\textsuperscript{74}

Long-term remission rates with doses ≥ 4 g (adrenolytic doses) are about 30%.\textsuperscript{62}

Side effects are dose-dependent and usually intolerable with doses > 6 g/day. Patients must be monitored with UFC, as serum cortisol may be elevated even when free cortisol in circulation is not; the aforementioned is due to the fact that mitotane increases binding of cortisol into the corticosteroid-binding globulin. Anorexia, nausea, somnolence, lack of coordination, hepatotoxicity, and teratogenicity are some of its adverse effects.\textsuperscript{74} The adrenolytic effect involves oxidative damage by means of free-radical formation such as superoxide, which produces hydroxyl radicals and induces lipid peroxidation, leading to adrenal insufficiency and hypercholesterolemia.\textsuperscript{58}

Maintenance therapy using mitotane may be a reasonable option in the case of persistent and untreatable CD\textsuperscript{79} and should be used as a last resort in case of failure or intolerance to the other drugs.

In Table 1 we briefly describe the adrenal steroidogenesis inhibitors, as well as doses and collateral effects.

**Table 1** Steroidogenesis inhibitors used in patients with Cushing syndrome\textsuperscript{65,73}

<table>
<thead>
<tr>
<th>Drug</th>
<th>Initial dosage</th>
<th>Maximal dosage</th>
<th>Action mechanism</th>
<th>Side effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ketoconazole</td>
<td>200 mg bid</td>
<td>400 mg tid</td>
<td>Inhibits CYP11B1 and CYP11A1</td>
<td>Abnormal liver tests</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(15%), gynecomastia (13%), gastrointestinal effects (8%)</td>
</tr>
<tr>
<td>Metyrapone</td>
<td>250 mg qid</td>
<td>1500 mg qid</td>
<td>Inhibits CYP11B1</td>
<td>Dizziness and ataxia</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(15%), nausea (5%), skin rash (4%), edema (8%), and hirsutism (70%)</td>
</tr>
<tr>
<td>Mitotane</td>
<td>500 mg qid</td>
<td>3000 mg tid</td>
<td>Adrenolytic, inhibits CYP11B1 and</td>
<td>Abnormal liver tests,</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>CYP11A1</td>
<td>teratogen</td>
</tr>
<tr>
<td>Etomidate</td>
<td>Bolus 0.03 mg/kg intravenous followed by infusion 0.1 mg/kg/hour</td>
<td>0.3 mg/kg/hour</td>
<td>Inhibits CYP17, CYP17-20 and CYP11B1</td>
<td>Hypotension, sedation</td>
</tr>
</tbody>
</table>

Abbreviations: bid, twice daily; qid, four times daily; tid, three times daily.
Valproic acid
Valproic acid is a drug that can potentially suppress ACTH-releasing hormone. Different assays have not shown efficacy in CD. It has been used as a therapeutic adjuvant in NS cases. So far, there are not many data that can prove its usefulness. Other drugs like cyproheptadine have had the same results.

Somatostatin and pasireotide analogs
Somatostatin is a cyclopeptide that is present in two active forms of 14 and 28 amino acids widely distributed in the brain and peripheral tissues. It regulates the release of some hormones and inhibits endocrine and exocrine secretions. Biologic actions of somatostatin are mediated by 5 G-protein-coupled receptors, namely somatostatin receptor type 1 to 5 (sst<sub>1-5</sub>).<sup>89</sup> sst<sub>1</sub>, sst<sub>2</sub>, sst<sub>3</sub>, and sst<sub>4</sub> inhibit cell proliferation. sst<sub>5</sub> is cytotoxic and produces cell death or apoptosis.<sup>87</sup> The natural somatostatin is fixed with great affinity to all of the five receptors. However, utility of somatostatin is limited by its accelerated catabolism at serum level. Different metabolically stable analogs of somatostatin have been synthesized, like octreotide and lanreotide, which bind only to sst<sub>2</sub>, having a slight affinity for sst<sub>1</sub> and sst<sub>5</sub> and a very low or absent affinity for sst<sub>3</sub> and sst<sub>4</sub>. Somatostatin and octreotide have not been demonstrated to significantly inhibit ACTH secretion in patients with CD.<sup>90,91</sup>

Pasireotide is a recent analog of somatostatin with multireceptor activity. It is a cyclohexapeptide that bonds with great affinity to all sst receptors, except sst<sub>5</sub>.<sup>92</sup> In contrast with octreotide, pasireotide shows high subnanomolar affinity for sst<sub>1</sub> and an improved metabolic stability. Lesche et al<sup>93</sup> showed that pasireotide is more potent than octreotide for inducing internalization and signaling of sst<sub>1</sub> and sst<sub>5</sub> receptors, which are predominant in hypophyseal adenomas that produce ACTH, and at the same time their expression diminishes in adenomas with aggressive behavior.<sup>93</sup> The functional activity of pasireotide over sst<sub>1</sub>, sst<sub>5</sub>, and sst<sub>5</sub> is >30, eleven, and 158 times higher, respectively, and is seven times lower over sst<sub>1</sub>.<sup>92</sup> Pasireotide has shown inhibition of basal and stimulated release of ACTH from ACTH-producer hypophyseal adenomas.<sup>87</sup>

In a phase II trial<sup>94</sup> that included patients with recent CD diagnosis or with postsurgical relapse (excluding those receiving RT), a 600 g dose every 12 hours over 15 days was used, recording a UFC decrease in 76% of the patients. From these, 17% reached normal UFC concentrations; in addition, direct effects upon ACTH release were observed. Pasireotide safety has been assessed in healthy volunteers and in cases of acromegaly, CD, and carcinoid syndrome carriers; the most common side effects are the gastrointestinal ones and transient increases in fasting glucose.<sup>89</sup> In order to determine the role of pasireotide in patients with CD de novo, persistent or recurrent, the phase III, multicenter, randomized, double-blind PASPORT (Pasireotide Clinical Trial Portfolio) trial is being conducted, and we are still waiting for official results.<sup>95</sup> Because somatostatin receptors have been found in normal adrenal cortex and adrenocortical tumors, pasireotide may have a potential effect at this level.<sup>93</sup> An 80-day clinical trial has recently been performed in 17 patients with CD in order to examine the efficacy of the medical therapy based on treatment with pasireotide and sequentially extended with cabergoline and ketoconazole. Treatment began with 100 g subcutaneous pasireotide three times a day. If the UFC had not normalized at day 15, the pasireotide dose was increased up to 250 g three times a day. At day 28 and upon persistent elevation of UFC level, cabergoline was administered. If UFC was not normalized by day 60, a 200 mg dose of ketoconazole three times a day was added. Pasireotide induced normalization of UFC in four of 17 patients at day 60, while eight of 17 patients still had an elevated UFC, despite combination therapy; added ketoconazole induced complete response in 88% of patients, thus improving blood pressure and weight.<sup>96</sup>

Somatostatin and analogs are therefore capable of suppressing secretion of ACTH originating from ectopic neuroendocrine tumor; positive octreoscan gammagraphy of tumors (In-pentreotide) would be an indicator of somatostatin receptors, making them more likely to respond to this therapy with suppression of ACTH and cortisol. Nevertheless, expression or recording of such receptors is highly variable.<sup>74,97</sup> According to previous reports and probably due to the presence of high levels of cortisol that induce downregulation of sst<sub>2</sub>, octreotide (specifically) has proven to have limited usefulness.<sup>98</sup> A valuable tool in the future with the potential for optimizing drug therapy would be a targeted search of somatostatin tumor receptors using rabbit monoclonal antibodies UMB-4 and UMB-1 during routine histopathological studies, and thus the best drug selected according to the prevailing receptor.<sup>99</sup>

Dopaminergic agonists and cabergoline
Dopamine is the predominant catecholaminergic neurotransmitter in the central nervous system, acting by means of receptors (D<sub>1-5</sub>). The D<sub>2</sub> receptor is expressed in the anterior and intermediate lobe of the hypophysis, showing heterogeneous and variable expression in 89% of...
all hypophyseal tumor types. The expression of receptor D2 has been detected in 23%–80% of the ACTH-secreting hypophyseal adenomas, and it has been correlated with an inhibitory effect of dopaminergic agonists in the secretion of ACTH in vitro.87,93

Bromocriptine is a D2 agonist and D1 antagonist; in cell cultures of human hypophyseal tumors it suppresses the secretion of ACTH and induces apoptosis in AtT-20 cells.87 Moderate results have been reported with doses from 17.5 mg/day to 40 mg/day.108 In different studies, the effectiveness of bromocriptine varies between 0% and 50% and may be ≤10%,87,100 with few results on a long-term basis.1

Cabergoline is a dopaminergic agonist with greater selectivity and affinity for the D3 receptor than bromocriptine.100 Godbout et al101 performed a retrospective analysis of cabergoline monotherapy in 30 patients with CD, beginning with a dose of 0.5–1.0 mg/week and progressive titrations up to a maximum dose of 6 mg/week; taking UFC as a parameter, 36.6% of the patients showed a complete response within 3–6 months, 13.3% showed a partial response, and 30% maintained the response on a long-term basis (mean of 37 months) with a dose of 2.1 mg/week. Pivonello et al102 performed a prospective trial in 20 patients with surgical treatment and without disease remission with initial doses of 1 mg/week and monthly titrations until obtaining a response or until achieving a dose of 7 mg/week (mean 3.5 mg/week), reporting 3 months later with response to cabergoline in 15 patients (75%). In these patients it was maintained in ten. In only eight (40%) was sustained control at 24 months achieved; additionally, there was a tumor decrease in 20% of the patients and improvement of hypertension and glucose intolerance in most of them, independently from changes in UFC. In contrast, Lila et al103 conducted another prospective trial, which included 20 patients without surgery remission with or without RT adjuvant, with an initial dose of 1 mg/week and a maximal dose of 5 mg/week (mean 3.6 mg/week) and monitoring throughout the year, wherein serum markers were assessed and pharmacologic response reported of 28%, 25%, and 17%. Limitations of the study were the short follow-up and lack of UFC.

Vilar et al104 studied combination therapy by assessing the effectiveness of cabergoline (3 mg/week) alone or in combination with ketoconazole (<400 mg/day) in twelve patients with CD and unsatisfactorily treated with TSS. After a 6-month treatment with monodoses of cabergoline 2–3 mg/week, UFC was normalized in 25% of the patients, with UFC reductions from 15% to 48.8%; administration of ketoconazole to the remaining nine showed UFC normalization in six of them (66.7%), with doses from 200 mg/day to 400 mg/day.

It should be taken into account that there is no homology in the different biochemical criteria regarding complete or partial response and reduction percentage in several studies: some use UFC, others use a 1 mg dexamethasone test, and others refer to serum cortisol; likewise, maximum doses and follow-up times differ, therefore results are not totally equivalent. We need more prospective studies with better homogeneity regarding criteria, dose, and time in order to set up a clear position regarding the utility of dopaminergic agonists; for the time being, we could consider cabergoline as the best option due to its selectivity on D2, greater tolerability, and best evidence in the long-term follow-up. Variability on results also corresponds to the aforementioned variability in expression of dopamine receptors in hypophyseal adenomas.

Mifepristone is a synthetic steroid (derivative of the synthetic progestin norethindrone) acting as a potent antagonist of progesterone and cortisol.105,106 When administered in high doses, it is a potent antagonist of GC.107 It blocks the actions of hypercorticism on target organs by blocking the GC receptor type II.62 It can antagonize the acute effects of cortisol over the protein and carbohydrate metabolism and inhibit the cortisol-induced peripheral vasoconstriction. Its applications under hypercortisolism include CD, adrenal adenoma, adrenocortical carcinoma, and ectopic production of ACTH.108 Mifepristone and its metabolites have a greater affinity for the GC receptor (mifepristone, 100%; metabolites, 45%–61%) than dexamethasone (23%) and cortisol (9%).106 A problem with the drug is that ACTH and serum cortisol do not provide any relevant information regarding efficacy; it should be judged only upon a clinical basis.62 In fact, the increase of ACTH and cortisol by feedback is a potent adverse effect in patients with CD. Because no mineralocorticoid receptor is blocked, an increase in cortisol can lead to hypokalemia, which may require a mineralocorticoid antagonist such as spironolactone or eplerenone. Hypercortisolism symptoms, namely anorexia, nausea, arthralgia, and headache, may occur and require dose adjustments.74

Johanssen and Allovio108 conducted a retrospective analysis for hypercortisolism treatment with mifepristone in a total of 18 patients. Daily doses varied from 5 mg/kg to 30 mg/kg; the results suggest a significant improvement in patients in whom surgery and adrenal steroidogenesis inhibitors failed to control the overproduction of cortisol. A very positive feature is its rapid onset of action, making it particularly useful in acute cortisol-induced psychosis. Adrenal failure must be taken into account. There have been
two cases reported that point to the use of mifepristone allowing the “positivization” of two bronchial carcinoid tumors that initially did not capture $^{111}$In-pentetreotide (octreoscan). The findings support the theory of inhibition in expression of sst$_2$ in neuroendocrine tumor cells by hypercortisolism. This opens a new window in the potential indications of mifepristone, with “therapeutic-diagnostic” utility and even increasing diagnostic sensitivity previously reported for octreoscan ($\approx$50% in most of the series).

In February 2012, the Food and Drug Administration passed mifepristone use for treating chronic hyperglycemia in patients with uncontrolled CD. We suggest that it can be employed in cases of partial efficacy, intolerance, or lack of access to other drugs, either as a replacement of those or as adjuvant therapy, in a psychotic crisis case, or in those who require a rapid response because of the severity of the clinical symptoms. It must not be used during pregnancy.

As for future potential treatments, we can discuss dopastatin (BIM-23A760), which is a chimeric molecule with which we are seeking to exploit the theoretical potential of somatostatin and dopamine receptor interactions. This compound presents high activity on D$_2$ and sst$_2$ receptors and is moderate on sst$_5$. Its role in nonfunctional hypophysal adenomas is to exert a cytostatic and cytotoxic effect, mainly mediated by D$_2$. Regarding its role in ACTH-producing adenomas, in vitro preliminary results are encouraging because they show inhibition of dose-dependent cell replication in 60% of cases; however, there are still few data.

Continuation of hypercortisolism:
- Consider bilateral adrenalectomy
- Refuses surgical therapy
- Pharmacological treatment and RT, wants surgical therapy
- Refuses surgical therapy
- Nonresectable tumor, high surgical risk:
  - Treatment with mitotane
- Intensive care and management with mifepristone or etomidate infusion
- A stable and symptomatic patient with high surgical risk
- Unstable patient with uncontrollable symptoms and acute complications
- Presurgical management with ketoconazole or other drug
- ACTH-dependent Cushing: hypophysal tumor or other resectable tumor
- ACTH-independent Cushing: adrenal adenoma or cancer
- No remission criteria: pharmacological treatment (ketoconazole, cabergoline, pasireotide, or combination) with RT. Reconsider hypophysal surgery
- Continuation of hypercortisolism: consider bilateral adrenalectomy

**Figure 1** Treatment of Cushing syndrome.

**Abbreviations:** ACTH, corticotrophin; RT, radiotherapy.
Conclusion

We conclude by remarking that although diagnostic and therapeutic techniques have been substantially improved, CS is still a challenge regarding its treatment, particularly if the disease persists after first-line therapy. Finally, we suggest a simple therapeutic algorithm (Figure 1) for CS treatment.

Disclosure

The authors have no conflict of interest relevant to this manuscript to declare.

References


