Review

Perioperative Management of Endocrine Active Adrenal Tumors

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Key words
pheochromocytoma, paraganglioma, primary aldosteronism, Cushing’s syndrome, surgery, laparoscopy, perioperative management

received 13.05.2018
revised 22.06.2018
accepted 05.07.2018

Bibliography
DOI https://doi.org/10.1055/a-0654-5251
Published online: 14.8.2018
Exp Clin Endocrinol Diabetes 2019; 127: 137–146
© J. A. Barth Verlag in Georg Thieme Verlag KG Stuttgart · New York
ISSN 0947-7349

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ABSTRACT
Endocrine active adrenal tumors are associated with a relevant risk of complications during surgery, either due to direct hemodynamic effects of the hormone excess or due to hormone related comorbidities. Over the last decades, careful preoperative evaluation and improved peri-interventional medical management of affected patients has resulted in a significant reduction of perioperative complications. In addition, improvement in anesthesia and surgical techniques with the feasibility of laparoscopic adrenalectomy have contributed to reduce morbidity. Nevertheless, there are still several challenges to be considered in the perioperative care of these patients. Due to the rarity of functionally active adrenal tumors, there are no prospective data available to guide clinical management. Accordingly, most recommendations are based on retrospective data analysis, expert opinion or carry weak evidence based on small series or case reports. The aim of this review is to summarize the current knowledge and to suggest practical approaches to reduce perioperative complications in endocrine active adrenal tumors. This review exclusively deals with data from adult patients with functionally active adrenal tumors.

Abbreviations
PGL paraganglioma
PCC pheochromocytoma
PPG Pheochromocytoma and paraganglioma
PA primary aldosteronism
APA aldosterone producing adenoma
CS Cushing syndrome
BAH bilateral adrenal hyperplasia
HPA axis hypothalamic-pituitary-adrenal axis
ACC adrenocortical carcinoma

Pheochromocytoma and paraganglioma (PCC and PGL)

Background
Pheochromocytoma (PCC) are neuroendocrine tumors of chromaffin tissue mostly producing one or more catecholamines; epinephrine, norepinephrine and/or dopamine. Most PCC are intra-adrenal (90%). Rarely, extra-adrenal PCC can be found in the paraganglia cells of the sympathetic nervous system, and the organ of Zuckerkandl. [1] Most PCC are endocrine active. On the other hand, paraganglioma (PGL) are tumors arising from parasympathetic or sympathetic ganglia which depending upon location may or may not be function-
ally active. PGL arising from parasympathetic ganglia in the head and neck was previously thought to be non-functional. [2] However, the current literature has revealed that up to one third of head and neck PGL are, in fact hormonally active. [3] Sympathetic PGLs are commonly located around the inferior mesenteric artery or aortic bifurcation and are almost universally endocrine active.

Overall, PPGL represent tumors that carry a high risk of morbidity, even if benign. Further, these tumors (particularly PGL) carry a relevant risk of malignancy which requires long-term follow-up examinations. The major cause of morbidity results from hypersecretion of catecholamines, which may lead to hypertensive crisis, ventricular arrhythmia, myocardial infarction, stroke or other vascular manifestations. [2] In a retrospective study by Riester et al., patients with PCC were analyzed for incidence of life-threatening complications in three German centers from 2003 to 2012, whereby higher preoperative systolic blood pressure and the maximum tumor diameter were identified as predictors of life-threatening events. [4] The treatment of choice for PCC is laparoscopic adrenalectomy. [5] Surgical resection can be complicated by abrupt release of catecholamines during the induction of anesthesia or surgical manipulation of tumor leading to cardiovascular complications. From a physiological point-of-view, different adrenoreceptors are targeted differentially by catecholamines. Therapeutically, preoperative treatment with alpha-adrenergic receptor blockers to counteract their vasoconstrictive action has been shown to reduce the risk of perioperative cardiovascular complications in PPGL patients. Therefore, preoperative medical blockade has been recommended in current guidelines. [6]

Preoperative management

The catecholamine secretory profile depends on the localization of tumor (adrenal versus extra-adrenal) and underlying hereditary disease with a known germline mutation. Whereas PCC can produce any catecholamine, PGL mostly secrete norepinephrine or dopamine because of lack of expression of phenylethanolamin N-methyltransferase necessary to convert norepinephrine to epinephrine. [7–8]

Clinical presentation of the patient may vary depending on catecholamine secretory profile due to differential receptor activation by different catecholamines. [7, 8] For example, adrenalin-secreting tumors may not present with paroxysmal hypertension, but also hypotension whereas noradrenalin-secreting tumors more often result in persistent hypertension. Patients with dopamine-secreting tumors may have rather unstable blood pressure and may present with tachycardia. [9] Up to 50 % of PCC may even be clinically asymptomatic, particularly in case of an adrenal incidentaloma implying the need of biochemical testing even in the absence of hypertension in this subgroup of patients. [7–8]

The Endocrine Society guidelines do not provide detailed recommendations for the extent of preoperative cardiological assessment of PPGL patients. Some authors consider preoperative cardiological evaluation to screen for cardiomyopathy or coronary heart disease, [10] while others postulate an individual approach depending on the risk constellation. [11] Other authors suggest a routine preoperative echocardiography to exclude a rare case of primary cardiac PPGL. [12] We suggest preoperative echocardiography in symptomatic patients and in patients with long-standing arterial hypertension to evaluate the extent of left ventricular dysfunction or indirect evidence of coronary heart disease.

Preoperative medical treatment

**Alpha-adrenergic receptor antagonists**  In the current Clinical Practice Guideline of the Endocrine Society preoperative blockade with alpha-adrenergic receptor blockers is recommended in all patients with functionally active PPGL. The goal of preoperative medication is to normalize blood pressure and heart rate. After initiation of treatment with alpha-adrenergic receptor blockers, patients should be instructed to increase dietary sodium and fluid intake to reverse catecholamine-induced intravascular volume contraction. These measures can aid to decrease intraoperative hemodynamic instability and postoperative hypotension. [6]

The role of alpha-adrenergic receptor blockers in reducing perioperative cardiovascular complications has been suggested in many case series. Goldstein et al. observed a high perioperative complication rate of 69 % in patients who did not receive alpha-blockers in a case series of 104 patients from 1950 to 1998. [13] Livingstone and coworkers reported an improved postoperative outcome in a retrospective review of PCC resection from 1992 to 2013 in 88 patients. This impressive reduction of perioperative morbidity from 40 % to 7 % was evident over the last 10 years of the study period and was mainly attributed to preoperative treatment with alpha-adrenergic receptor blockers. In particular, higher dosages of preoperatively administered alpha-adrenergic receptor blockers correlated with a reduction in the risk of hemodynamic instability. [14]

There has been some controversy in the literature regarding the need to use preoperative alpha-blockade in patients with atypical presentation. These patients with atypical presentation are characterized by normal or even low blood pressure despite biochemical evidence of a functionally active PPGL and/or have a predominant dopamine-secreting profile. These patients may not tolerate preoperative alpha-blockade very well and may have a higher risk of postoperative hypotension. However, if untreated, these patients have an increased risk of abrupt and extensive release of catecholamines provoking a hypertensive crisis or ventricular arrhythmias during tumor manipulation or induction of anesthesia, as several case studies have shown. [15, 16] Therefore, the risk of possible side-effects of preoperative alpha-blockade needs to be carefully weighed against the risk of perioperative hypertensive derailing in untreated patients. However, Lafont et al. could not find any difference in intraoperative hemodynamic instability in patients with normotensive and hypertensive PCC treated with alpha-adrenergic receptor blockers in their retrospective study from 2004 to 2012. Endocrine Guidelines suggest use of preoperative alpha-adrenergic receptor blockers in all functionally active PPGL as the benefit of reducing perioperative risk seems to clearly outweigh the possible side effects of the therapy. However, it should be noted that the side effects of medication can indeed be more pronounced in normotensive patients implying the need of careful dose-titration. [17] The last dose of alpha-adrenergic receptor blocker (phenoxycobenzamine) is administered at our institution the evening before the operation and the morning dose is waived on the day of the operation to minimize postoperative hypotension.
Although there are no prospective studies on the treatment of choice in premedication of PPGL patients, the Endocrine Society suggests alpha-adrenergic receptor blocker as the first choice based on the available retrospective data. [6] There is no clear drug preference for nonselective versus selective alpha-adrenergic receptor blockers due to lack of randomized controlled studies directly comparing effectiveness and limited international availability of all drugs. Phenoxybenzamine is used most frequently as a long-lasting, nonselective, and irreversible inhibitor of alpha-1 and alpha-2 adrenoceptors. It may potentially lead to hypotension after tumor resection due to its long-lasting effect. [11] Current Endocrine Society guidelines suggest titration of phenoxybenzamine dose based on blood pressure. [6] In a retrospective study of 100 pheochromocytoma patients from 1992 to 2013, Livingstone reported median preoperative dose of phenoxybenzamine to be 119 mg with a trend to use higher dosages in later years of the study. In this study, higher preoperative phenoxybenzamine doses were a significant predictor of improved intraoperative hemodynamic stability. [14] At our institution we usually begin with a starting dose of 5 mg/day in the evening and increase it by 5–15 mg per day till the aim is achieved or intolerable side-effects occur. The aim is to achieve normal hemodynamic parameters (blood pressure and pulse) with tolerable side effects (nasal congestion, orthostatic hypotension, diarrhea, dizziness and reflex tachycardia).

Selective and competitive alpha-1 adrenoreceptor blockers such as prazosin, terazosin, and doxazosin are characterized by a shorter half-life. Some retrospective studies have reported fewer side effects, especially reflex tachycardia and postoperative hypotension, with selective alpha-1 adrenoreceptor blockers. [11, 18] However, a study by Kocak et al. could not find any significant difference in hemodynamic parameters intra- and postoperatively among phenoxybenzamine, doxazosin and prazosin. [19] In a further retrospective study on laparoscopic resection of PCCs, Weingarten and co-authors compared 50 Mayo Clinic patients (98% received nonselective alpha-1 adrenoreceptor blocker, phenoxybenzamine) with 37 Cleveland patients (65% received selective alpha-1 adrenoreceptor blocker). In this study, patients premedicated with selective alpha-1 adrenoreceptor blocker had significantly higher average blood pressure and required more volume (crystalloids and colloids) intraoperatively, whereas patients treated with nonselective alpha-1 adrenoreceptor blocker required more phenylephrine to manage hypotension during surgery. The authors postulated a more efficient volume repletion under phenoxybenzamine reducing the need of volume replacement. However, there were no significant differences in postoperative outcome or duration of hospitalization. [20]

Therapeutic goals According to the current guidelines of the Endocrine society a duration of 7 to 14 days of preoperative treatment with alpha-adrenoreceptor blocker is suggested. The therapeutic goal is to achieve blood pressure levels of < 130/80 mmHg in a seated position and a systolic blood pressure of > 90 mmHg upon standing. The optimal heart rate is regarded to be between 60–70 beats/minute when sitting and between 70–80/minute when standing. As mentioned above, salt-rich diet and sufficient fluid intake are recommended preoperatively to reverse catecholamine-induced blood volume contraction. [6]

Beta-adrenergic receptor antagonists Beta blockers can be recommended as add-on treatment to combat reflex tachycardia. It is important to note that beta-adrenoreceptor blockers are only indicated after treatment with alpha-adrenoreceptor blockers is already established because of the danger of inducing hypertensive crisis due to unopposed alpha-adrenoreceptor activation through catecholamine excess in untreated patients. [6] According to Pacak, preference should be given to cardioselective beta-adrenoreceptor blockers, whereas the Endocrine Society does not provide any preference for nonselective versus selective beta-adrenoreceptor blockers. There is, however, a consensus to avoid treatment with a combined alpha- and beta-blocker (labetalol) due to the risk of paradoxical hypertension or hypertensive crises owing to its weaker alpha-antagonist action and stronger beta-antagonist action. [6, 11]

Calcium-channel-blockers If the target blood pressure values are not reached with alpha-adrenoreceptor blocker, therapy can usually be expanded with calcium-channel-blockers. [6, 11] Some authors even propagate monotherapy with calcium-channel-blockers. [21–23] In our opinion, this should be considered as an option only in cases of mild hypertension or in patients suffering severe postural hypotension under treatment with alpha-adrenoreceptor blocker. [6]

Other agents Another therapeutic approach in preoperative medication is alpha-methyl-paratyrosine (metyrosine), a catecholamine synthesis inhibitor, in combination with alpha-adrenergic receptor blocker. There are few retrospective case studies suggesting a benefit in patients with more pronounced catecholamine release and a known metastatic disease. [24, 25] Wachtel et al. compared intraoperative hemodynamics in 174 PPGL patients (81.6% were treated with phenoxybenzamine in combination with metyrosine while the remaining were treated with phenoxybenzamine alone). The authors reported less intraoperative hemodynamic instability and less postoperative cardiovascular complications in the combined treatment group. [26] Nonetheless, limited availability and intolerable side effects of metyrosine (sedation, depression, galactorrhea and extrapyramidal symptoms) caused by central and peripheral inhibition of catecholamine synthesis prohibit wider use of this agent. [11]

If hypertensive derailment occurs despite preoperative medication, sodium nitroprusside, nitroglycerin, phentolamine or urapidil can be used intraoperatively to control blood pressure. Tachyarrhythmias can usually be treated with short-acting beta-blockers and lidocaine. [1] Some drugs such as dopamine D2 receptor antagonists, opioid analgesics, norepinephrine reuptake inhibitors, serotonin reuptake inhibitors, monoamine oxidase inhibitors, corticosteroids, peptides, neuromuscular blocking agents, and beta-blockers without prior alpha-blockade can predispose the patient to hypertensive crisis. Accordingly, these drugs should be avoided. [6]

Surgical management The standard treatment of PPGLs is surgical resection. Regarding surgical approach there are only retrospective analyses of patient data comparing laparoscopic with conventional transabdominal approach. However, in the recent literature the laparoscopic approach is clearly preferred over the transabdominal procedure due to many advantages such as reduced period of hospitalization,
shorter recovery time, less blood loss, less postoperative pain, and less surgical morbidity. [5, 27–29]

Fernandez-Cruz and coworkers compared laparoscopic and transabdominal adrenalectomy and reported more cardiovascular instability following the transabdominal approach despite similar increases in intraoperative catecholamine levels. [27] On the other hand, Weisman et al. found no significant difference in hemodynamic instability between minimally invasive and open access. [5]

Gumbs and Gagner compiled 2565 cases of laparoscopic adrenalectomy for adrenal tumors. The authors reported the lateral transabdominal adrenalectomy as the most commonly performed procedure, followed by posterior retroperitoneal endoscopic adrenalectomy and laparoscopic anterior trans-abdominal adrenalectomy. Overall, the most frequent complications were postoperative bleeding (40%), organ injuries (5%), and wound infections. Interestingly, cardiovascular complications resulting from hypertensive crises were documented in only 1% of cases. There were no differences between the different laparoscopic approaches in terms of duration of hospital stay or complications. The only limiting factor to retroperitoneal access was maximum tumor size of 6-9 cm. [30] According to a more recent retrospective study by Lee et al. comparing surgical access in 58 patients with adrenalectomy for adrenal tumors no relevant complications had occurred. [31] The Endocrine Society suggests to give priority to the retroperitoneal approach in patients who have previously undergone abdominal operations or those requiring a bilateral adrenalectomy. For patients with bilateral pheochromocytoma, Castillo et al. reported laparoscopic synchronous bilateral adrenalectomy to be a safe procedure. [32] Partial adrenalectomy is recommended in patients with bilateral disease or patients with hereditary PCC with a prior unilateral adrenalectomy to prevent postoperative adrenal insufficiency. [6] The negative corollary, of course, is the higher risk of recurrence in the adrenal-sparing approach. Asari and coworkers reviewed 348 patients with PCC due to MEN 2A. Following adrenal-sparing operation, 31% of 187 patients had a recurrent PCC (ipsi- or contralateral). In comparison, only 3% of 161 patients undergoing bilateral complete adrenalectomy had a recurrent PCC. [33]

In very large tumors and in tumors with apparent malignant potential, an open resection should be discussed to reduce the risk of intraoperative tumor rupture and to maximize the chances of complete resection. [30] According to the Endocrine Society guidelines, open surgery is recommended in tumor masses > 6 cm. However, in 2006 Gumbs and Gagner suggested that laparoscopic adrenalectomy may be considered for the resection of benign secreting and non-secreting tumors up to 12 cm. [30, 34]

Great care must be taken to avoid tumor rupture during removal of pheochromocytoma. Rafat et al. reported 5 cases of tumor recurrence or persistence which showed evidence of tumor capsule rupture upon macroscopic examination of the surgical specimen. Indeed, peritoneal tumor implantation due to tumor capsule rupture during the resection of an apparently benign primary tumor, is a rare, but potentially disastrous complication leading to tumor persistence or recurrence. [35]

We suggest discussion of PPGL cases in interdisciplinary tumor boards to decide the best surgical approach for the individual patient. Due to increased risk of hemodynamic instability in patients with PPGL und technically challenging surgical removal of these highly vascular tumors, all surgeries should be preferably done in a tertiary care center with an experienced team of surgeons, anesthesiologists and endocrinologists to maximize chances of a successful outcome.

For detailed information for the choice of anesthetic agents and anesthesia techniques, we refer the reader to consult the relevant literature in the field. [36]

**Postoperative management**

Postoperatively, the Endocrine Society recommends close monitoring of blood pressure, heart rate and plasma glucose during the first 24–48 h. There is a risk of postoperative hypotension due to long-lasting effect of alpha adrenergicceptor blockers after surgical removal of the source of catecholamine excess. [1] However, persistent hypertension can also occur due to various causes such as excessive intraoperative fluid administration, residual tumor or unwanted ligation of the renal artery. Therefore, adjustment of medical treatment in the immediate postoperative period is likely to be required based on blood pressure and general clinical parameter. [1]

There are only individual case reports of postoperative hypoglycemia after removal of a PPGL, so that no statement on its exact prevalence is possible. However, some case reports reported severe symptomatic hypoglycemia grade II–III, which led to delayed awakening from anesthesia or occurred 30 min to 2 h postoperatively. A possible cause for this seems to be rebound hyperinsulinemia after surgical correction of catecholamine excess. Another potential mechanism could be an increased glucose uptake in peripheral tissues. [37, 38] Following complete or partial bilateral adrenalectomy or unilateral partial adrenalectomy of a single remaining adrenal gland, attention should be paid to the possibility of adrenal insufficiency. [6]

**Postoperative follow-up**

According to the European Endocrine Society guidelines, biochemical screening should be undertaken by measuring either plasma metanephrines or urinary levels of metanephrines and 3-methoxytyramine in a properly collected 24-h urine sample ideally 2–6 weeks after surgery to confirm complete PPGL resection. Certain risk factors such as young age, underlying genetic disease, and larger tumor size may predict higher risk of recurrence. [39] These high-risk patients should undergo lifelong annual screening to enable early detection of recurrent disease (local or metastatic) as well as new tumors. In our opinion, the possibility of genetic screening should be discussed with all patients. Especially, patients with other symptoms of hereditary disease, positive family history or known genetic mutation, immunohistochemical evidence of an underlying SDH-mutation in surgical specimen, bilateral disease and young age should be strongly advised to undergo genetic testing. Patients with sporadic tumors should be checked annually for a duration of at least 10 years according to the recent European Endocrine Society guidelines. [39] In addition, we suggest to instruct patients to actively report specific symptoms (worsening arterial hypertension, palpitations, increased sweating), thus allowing for symptom-based detection of a recurrence which should be confirmed or refuted with a biochemical screening. Interfering medication should be suspended before undertaking biochemical screening. Patients
Primary Aldosteronism (PA)

Background and diagnosis

Primary aldosteronism (PA) is the most common form of secondary arterial hypertension, even though currently underdiagnosed due to lack of awareness to screen patients at risk. The cardiovascular sequelae occur more frequently than in patients with essential hypertension, which is why early targeted therapy is essential. The prevalence of PA in hypertensive patients is estimated between 5-10% [40-42] with a correlation of the prevalence with the severity of arterial hypertension. [41, 42] The most common causes of PA are adrenal adenoma (APA) and bilateral adrenal hyperplasia (BAH). Adrenal carcinoma or familial hyperaldosteronism are rare causes. [40, 43]

Therapeutic approach

Once the diagnosis of primary hyperaldosteronism is confirmed, the Endocrine Society guidelines recommend adrenal venous sampling in the majority of patients seeking surgical treatment options. However, patients < 35 years of age with pronounced PA (spontaneous hypokalemia, serum aldosterone > 30 ng/dl or 831 pmol/l) and a corresponding unilateral adenoma on imaging can forego adrenal vein sampling as the probability of a functionally inactive adenoma is believed to be low in young patients. [40] The choice of therapy depends on the subtype of aldosterone excess. In a unilateral source of aldosterone excess (adrenal adenoma or unilateral adrenal hyperplasia), laparoscopic adrenalectomy is the best treatment. Patients suffering from BAH or inoperable patients with serious comorbidities should be treated medically with mineralocorticoid receptor antagonists. Spironolactone is the first-line agent, while eplerenone can be administered as an alternative if side effects occur (especially gynecomastia and erectile dysfunction in men and menstrual disturbances in premenopausal women). [40]

In a recent prospective study by Rossi and colleagues, long-term cardiovascular effects such as blood pressure control, left ventricular changes and the incidence of cardiovascular events (including atrial fibrillation) were investigated in 323 patients with hypertension. 180 patients had PA, out of which 110 were adrenalectomized and 70 were treated with mineralocorticoid receptor antagonists, and compared to 143 patients with medically treated essential hypertension. Notably, patients with PA had significantly more left ventricular hypertrophy than patients with essential hypertension. After specific therapy (adrenalectomy or mineral corticoid receptor antagonists), patients with PA showed a regression of left ven-

with biochemical evidence of recurrence require imaging studies for further evaluation.

For an overview on perioperative management, please refer to Table 1.

Table 1: Perioperative management of secreting adrenal tumors, adapted according to the guidelines of the Endocrine Society.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Perioperative management of secreting adrenal tumors, adapted according to the guidelines of the Endocrine Society.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Preoperative evaluation</strong></td>
<td>ECG, physical examination Screening for metastatic tumors in patients with PPGL, patients with PCC and elevated 3-Methoxytyramine (3-MT) in plasma or urine and in patients with SDHB gene.</td>
</tr>
<tr>
<td><strong>Premedication and immediate postoperative care</strong></td>
<td>Alpha-adrenergic receptor antagonist 7-14d before surgery Thromboembolic prophylaxis</td>
</tr>
<tr>
<td><strong>Therapeutic goals</strong></td>
<td>BP &lt;130/80 mmHg Heart rate 60–70 bpm</td>
</tr>
<tr>
<td><strong>Postoperative period</strong></td>
<td>Monitoring of blood pressure, heart rate and blood sugar for 24-48h postoperatively</td>
</tr>
<tr>
<td><strong>Common complications</strong></td>
<td>Persistent hypertension Hypotension Hypoglycemia Adrenal insufficiency (rare)</td>
</tr>
<tr>
<td><strong>Follow-up</strong></td>
<td>Metanephrines in plasma or urine 2-6 weeks postoperatively, then annually for 10 years Lifelong screening in high-risk patients (young patients, genetic disease, large tumor and/or PPGL)</td>
</tr>
</tbody>
</table>
triculat changes comparable to patients with optimally treated es-
ential hypertension. [44] These findings argue for the necessity of
an early, specific therapy of PA.

**Preoperative management**

Preoperatively, blood pressure values and hypokalemia should be
optimally adjusted preferably with a mineralocorticoid receptor
antagonist according to the Endocrine Society Guideline. Mostly,
a starting dose of 12.5 mg–25 mg spironolactone is used with ti-
tration up to a maximum dose of 100 mg. [40] However, there are
no specific recommendations about the optimum duration of pre-
operative treatment. At our institution, we treat patients with
spironolactone for a duration of 4–6 weeks to achieve a target blood
pressure of < 140/90 mmHg and normokalemia without potassi-
um supplementation. Postoperatively, hypoaldosteronism may
occur due to chronic suppression of the contralateral adrenal gland.

Earlier, it was believed that preoperative spironolactone therapy
may antagonize renin suppression and thus may prevent postop-
ervative zona glomerulosa insufficiency, whereas Fischer et al. could
not support this hypothesis on the basis of a retrospective analysis
of 110 patients adrenalectomized for unilateral adenoma. [45]

Based on our case-based experience, we suggest to discontinue
spironolactone up to 1 week before surgery. The rationale for this
approach is long half-life of active metabolites, even though the
plasma half-life of spironolactone itself is short (1.3 h). Patients are
maintained on other antihypertensives and or potassium substitu-
tion to maintain normotension and normokalemia till the day of
surgery.

**Surgical management**

Laparoscopic adrenalectomy is the standard therapy for unilateral
disease due to shorter hospitalization time, less blood loss, fewer
complications and less need for analgesics. [46] In addition, the ad-
renal gland is not visibly enlarged in many cases of unilateral PA in
contrast to other adrenal tumors such as PCC or Cushing adenoma.

Mostly, total adrenalectomy is preferred over partial adrena-
ectomy to ensure remission. [40] Nanba et al. showed in a retro-
spective study of 55 patients following adrenalectomy in PA that
three of the patients (5.5 %) did not experience postoperative im-
provement despite preoperative lateralization by adrenal vein sam-
pling. All 3 patients had a partial adrenalectomy, whereas all pa-
ients with total adrenalectomy were cured postoperatively. [47]

In a study by Pautler et al. on 11 patients with laparoscopic partial
adrenalectomy, intraoperative ultrasound was used to improve the
differentiation between adrenal parenchyma and tumor tissue to
optimize the outcome of partial adrenalectomy. The authors were
able to detect an additional lesion in one patient that was not seen
in preoperative imaging. [48] However, whether these lesions cor-
respond to endocrine active nodules remains uncertain. In gener-
al, surgical treatment has been shown to be cost-effective as com-
pared to lifelong medical therapy in unilateral disease. [49–50]

**Postoperative management**

In a retrospective study, Kim et al. compared 26 PCC patients with
34 patients with PA who had laparoscopic adrenalectomy. As ex-
pected, tumor size, surgery duration and cardiovascular complica-
tion rates were higher for PCC than APA patients. [51] The possible
postoperative complications (postoperative bleeding, wound in-
fec tion, pain) were comparable among those groups.

A slight creatinine increase is not uncommon after unilateral ad-
renalectomy in PA patients. Vasodilatation of the arterioles and in-
creased sodium reabsorption via tubulo-glomerular feedback leads
to glomerular hyperfiltration and thus often masks a restricted kid-
ney function. [45] Furthermore, structural changes of the renal pa-
renchyma are observed in the context of PA, which could be ex-
plained by the long-standing arterial hypertension as well as by di-
rect toxic effects of aldosterone. [45, 52] A detailed analysis of the
German Conn’s Registry patients in 2009 revealed a moderate post-
operative decline in glomerular filtration rate, which stabilized in
the long-term follow-up. [53]

In severe cases, unilateral aldosterone excess can result in com-
plete suppression of the contralateral zona glomerulosa; as a re-

t, potassium substitution and mineralocorticoid receptor antag-
ons must be suspended postoperatively and electrolyte checks
should be carried out regularly. Fischer et al. showed in a retrospec-
tive chart review of 110 patients adrenalectomized for unilateral
adenoma that 16 % of patients had postoperative hyperkalemia due
to prolonged zona glomerulosa insufficiency. 5 % of patients even
required mineralocorticoid therapy for 11–46 months. Predictive
markers for development of postoperative hyperkalemia were re-
duced glomerular filtration rate before operation which worsened
further after operation and microalbuminuria. [45]

Recently, based on a multi-stroid analysis, it has been demon-
strated, that a subgroup of PA patients are characterized by a co-
secretion of glucocorticoids. Although, these patients did not dis-
play obvious clinical manifestations of Cushings’s syndrome within
this cohort, a number of metabolic parameters such as BMI, waist
circumference and and HOMA-IR were well correlated with total

glucocorticoid output. Following surgery for unilateral PA, elevated
cortisol levels returned to normal and 13 out of 45 patients
(29 %) failed a co-syntropin test indicating partial adrenal insuffi-
ciency. [54] In light of this recent evidence, we suggest to perform
low-dose overnight dexamethasone-suppression test to evaluate
cortisol co-secretion in PA patients. In case of a pathological result,
these patients should be carefully monitored during immediate
postoperative period to rule out postoperative adrenal insufficien-
cy and to provide hydrocortisone substitution as a stress dosage.

**Follow-up**

Blood pressure can be expected to improve or normalize within the
first 1–6 months after successful surgery. [40] Based on an interna-
tional retrospective analysis of 705 patients from 12 different cen-
ters (The Primary Aldosteronism Surgical Outcome study, PASO),
standardized criteria for postoperative outcome with regard to clin-
ical and biochemical response were defined. According to the PASO
consensus, blood pressure and serum potassium should be deter-
mined within the first 3 months postoperatively for optimal adap-
tation of antihypertensive therapy and any necessary potassium
substitution. Within the first 6–12 months after adrenalectomy,
blood pressure, serum potassium, aldosterone and renin should be
measured again for a final assessment of therapy success. Subse-
quently, follow-up should be carried out annually to evaluate long-
term blood pressure control and to assess general cardiovascular
risk profiles. Evidence on recurrence of primary aldosteronism fol-
lowing surgery is sparse but should be considered in patients with worsening of blood pressure control or development of hypokalemia. According to the PASO study, complete clinical remission of 37% and a partial clinical success rate of 47% after surgery can be expected. Biochemically, the success rate was significantly higher at 94%. Clinical outcome was significantly better in younger and female patients than in older and male patients or in patients with comorbidities especially OSAS and diabetes. [55]

For an overview on perioperative management, please refer to [Table 1](#).

**Cushing’s Syndrome (CS)**

**Background**

Cushing’s syndrome (CS) is a rare endocrine disorder characterized by either ACTH-dependent (caused by a pituitary tumor or ectopic ACTH-production) or ACTH-independent (adrenal tumor) hypercortisolism. The most common cause is pituitary adenoma, followed by adrenal adenoma. Adrenocortical carcinoma (ACC) and malignant ectopic ACTH overproduction are rare causes, but accompanied by a poor prognosis. [56] In a retrospective study by Lindholm et al., the incidence of pituitary Cushing’s disease was reported at 1.2–1.7/million/yr. For adrenal adenoma, an incidence of 0.6/million/yr. and for adrenal carcinoma 0.2/million/yr. has been estimated. [57]

Clinically, the patient with cortisol excess may present with a plethora of symptoms such as visceral obesity, proximal myopathy, alopecia, hirsutism, abdominal striae rubrae and easy bruising (echymosis). The predominant comorbidities are obesity, insulin resistance, arterial hypertension and dyslipidemia. Furthermore, steroid induced immunodeficiency can develop, which significantly increases the risk of opportunistic infections. In addition, osteoporosis and psychiatric diseases, in particular depression, restricted cognitive function and psychosis also occur more frequently. [58]

The treatment of choice is early removal of the lesion causing hormone excess. If this cannot be achieved (in case of extensive pituitary surgery or in occult ectopic ACTH production) bilateral adrenalectomy can be performed to normalize hypercortisolism. [56, 59] Overall, in patients with Cushing’s syndrome, a benign adrenal cortisol producing adenoma offers best chances of cure.

**Preoperative management**

The optimal treatment of comorbidities associated with cortisol excess is essential in the perioperative management and for the postoperative long-term course of CS. In particular, blood glucose control using antidiabetic medication and/or insulin therapy as well as blood pressure control should be optimized. Patients should be screened for dyslipidemia and treated accordingly. The treatment of possible secondary psychiatric diseases is unfortunately often overlooked, but it should not be neglected. Patients with electrolyte abnormalities, especially hypokalemia, need appropriate substitution before operation. Due to the increased risk of infection, vaccination for influenza, herpes zoster and pneumococcus are recommended before surgery. [56] Signs and symptoms of an underlying infection may be masked in a state of cortisol excess due to immunosuppressive effect of glucocorticoids. It is important to remain vigilant about the possibility of infection following adrenalectomy. There are some case reports of life-threatening postoperative infection (especially pneumocystis jiroveci). [60] The Endocrine Society does not make any precise recommendations, but points to the possibility of a prophylactic administration of trimethoprim-sulfamethoxazole to prevent Pneumocystis jiroveci infections. [56] Alternatively, an extended perioperative antibiotic prophylaxis may be evaluated.

Another potentially life-threatening complication of CS is venous thromboembolism, which makes perioperative thromboembolic prophylaxis mandatory. [56] This risk seems to be lower in patients with benign adrenal tumor as a cause of CS. In a retrospective multicenter cohort study of 473 patients, 360 of whom had ACTH-dependent pituitary CS, Stuijver et al. found a postoperative incidence rate of 3.4% in these patients within the first 3 months following surgery. However, in this study no patient with ACTH-independent hypercortisolism suffered a thromboembolic event within 3 months. [61] In an analysis of 8 studies on 476 patients with Cushing’s syndrome from Van Zaane et al., 78 patients with adrenal hypercortisolism and 398 patients with ACTH-dependent hypercortisolism had a postoperative risk of thromboembolism between 0.5 and 5.6%. [62] Koutroumpi et al. identified increased urinary cortisol excretion and increased levels of von Willebrand factor as predictive markers based on retrospective analysis of 108 patients with CS, 58 of whom suffered from thromboembolism. [63] There are no prospective data on the dosage and duration of thromboprophylaxis available in the literature. Nevertheless, preoperative and prolonged postoperative prophylaxis, at least during the 4-6 weeks after surgery, in addition to early mobilization are recommended. [64–65]

Another aspect that should not be neglected in the preoperative evaluation of Cushing’s patients is proximal myopathy, which significantly impair the quality of life. In a retrospective study by Berr et al. in which 47 patients with CS were compared with 149 in remission after CS and 93 control patients, it was shown that myopathy is relevant in both the florid stage and in long-term observation. [66] These patients may benefit from the ongoing intensive muscle training in the long-term postoperative course.

**Surgical management**

As already mentioned, the surgical procedure of choice is laparoscopic adrenalectomy in case of adrenal CS. There is also increasingly carried out in ACC, although many cases still require conventional transabdominal approach. [56] As the precise surgical strategies to approach ACC is beyond the scope of this overview we refer to a recent review by Rayes et al. on this topic. [67]

**Postoperative management**

Due to previous hypercortisolism, the hypothalamic-pituitary-adrenal axis is suppressed, which is why immediate postoperative glucocorticoid substitution is necessary. The Endocrine Society recommends a glucocorticoid substitution of 10–12 mg/m² per day in 2-3 daily doses. To assess the success of surgery serum cortisol and ACTH should be measured. According to the Endocrine Society, remission of hypercortisolism is defined as morning serum cortisol levels < 5 µg/dL ( < 138 nmol/L) or UFC < 28-56 nmol/d
(<10-20 μg/d) 7 days after surgery. [56] To avoid interference with substitution therapy, hydrocortisone needs to be paused for the evaluation. In patients with a high likelihood of adrenal insufficiency measurement of morning serum cortisol with the last dosage of hydrocortisone at the afternoon of the previous day seems more appropriate than longer discontinuation for urine sampling. Usually the function of the HPA axis recovers within the first 18 months after unilateral adrenalectomy. A retrospective analysis by Berr et al. [65] of 91 patients with 3 different subtypes of CS (Cushing’s disease, adrenal CS and ectopic CS) also showed that the recovery of adrenal function depends on the etiology of the cortisol excess. [68] After bilateral adrenalectomy, there is a lifelong need for glucocorticoid and mineralocorticoid replacement. All patients must be informed preoperatively about adrenal insufficiency and instructed postoperatively on the adequate adjustment of the glucocorticoid dose in case of stress or infections. The postoperative dose adjustment should be undertaken by an endocrinologist. Every patient should receive an internationally valid emergency card implying the need of glucocorticoid treatment in emergency situations. [56]

After surgical correction of hypercortisolism, patients often suffer from symptoms of hormone withdrawal. Within the first postoperative days, this may lead to fatigue, nausea, somnolence, fever and joint pain and are sometimes difficult to delineate from symptoms of an Addison crisis. Biochemically, hypercalcemia and hyperphosphatemia can often be detected. The administration of a supraphysiologic hydrocortisone doses may reduce the symptoms; however, the dose should be reduced as quickly as possible to avoid iatrogenic CS. The symptoms of withdrawal can persist for up to a year postoperatively. [69]

Due to the proximal myopathy as described above, postoperative muscle-training and physiotherapy should be recommended. [66]

**Follow-up**

Postoperatively, a regular endocrine check-up should be ensured to evaluate and adapt the dose of substitution therapy. Glucocorticoid - and in cases of bilateral adrenalectomy mineralocorticoid - doses should mainly be based on clinical evaluation, blood pressure as well as serum potassium and plasma renin concentration. [56] The Endocrine Society does not recommend a specific scheme, but in our institute, we measure morning cortisol between 6:00-8:00 a.m on the first postoperative day while the patient is placed on dexamethasone-treatment 0.5 mg every 6-8 h after adrenalectomy. This low-dose dexamethasone does not interfere with the analysis of early morning cortisol as this is expected to be low in case of successful surgery. A morning cortisol <50 nmol/l confirms successful removal of tumor. Then, we start treatment with hydrocortisone and monitor the patient clinically to adjust the hydrocortisone dose. The duration of hydrocortisone substitution can vary from patient to patient. We perform ACTH stimulation test (08.00 a.m) to assess recovery of HPA axis every 3 months until normalization (stimulated Cortisol > 441 nmol/l). The last dose of hydrocortisone should be taken at least 18–24 h. before blood examination. It should be mentioned that testing for adrenal insufficiency varies from institution to institution due to the lack of specific guidelines. [70]

Due to the comorbidities associated with hypercortisolism, mortality and morbidity in CS are significantly higher than in the normal population. Even 10 years after remission, overall mortality was significantly higher, especially due to cardiovascular diseases. [71] In a retrospective study of 253 patients with CS, Bolland and colleagues demonstrated that mortality and morbidity remained elevated despite biochemical remission of cortisol excess. After therapy, arterial hypertension, sexual dysfunction, myopathy and mild psychiatric diseases improved, whereas diabetes mellitus, severe psychiatric diseases and osteoporosis were still detectable years after treatment. [72] For this reason, regular postoperative follow-up and treatment of cardiovascular risk factors is indicated. [56] In addition, patients following resection of an adrenocortical carcinoma require long-term oncological/endocrine follow-up.

All patients with adrenal incidentaloma should be screened for autonomous cortisol secretion even though these patients mostly lack clinical signs or symptoms of Cushing’s syndrome. As described above, patients with aldosterone producing adenomas can co-secrete cortisol. Because of important implication of impending adrenal insufficiency upon removal of these tumors, we suggest performing overnight dexamethasone-suppression test to evaluate cortisol oversecretion.

For an overview on perioperative management, please refer to Table 1.

**Conflict of Interest**

No conflict of interest has been declared by the authors.

**References**


