Resistant Hypertension: A Clinical Case

Hipertensión resistente: un caso clínico

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Abstract

Introduction: Secondary hypertension corresponds to 15% of the causes of arterial hypertension, and among them, primary hyperaldosteronism presents a variable incidence of about 3% in hypertensive patients. It has a slightly higher prevalence in women, between 30 and 60 years, and is usually unilateral. Case presentation: The authors describe the clinical case of a patient, followed by a severe hypertension medicated with four antihypertensive drugs for tension stabilization, maintaining systolic arterial tensions superior to 170 mmHg. In the aetiological study of hypertension, analytical alterations suggested hyperaldosteronism and a nodular lesion was detected in the left adrenal gland. The patient was submitted to surgery and excision of the lesion was done with histological confirmation of the diagnosis of corticomedullary adenoma of the adrenal gland. The patient presented improvement of the tension profile, with need to suspend two of the four antihypertensive drugs and to reduce the dose of the remaining ones. Conclusion: A tumor of the adrenal cortex producing aldosterone is the main cause of primary hyperaldosteronism and should always be excluded when the presence of difficult to control, severe hypertension is detected, since the standard treatment is surgical, leading to a stabilization of the tension pattern after a few months.

Keywords: Conn’s adenoma, adrenal adenoma, arterial hypertension, secondary hypertension, primary aldosteronism, resistant hypertension.

Resumen

Introducción: la hipertensión secundaria corresponde al 15% de las causas de hipertensión arterial, y entre ellas, el hiperaldosteronismo primario presenta una incidencia variable de sobre 3% en pacientes hipertensos. Tiene una prevalencia ligeramente mayor en mujeres, entre 30 y 60 años, y generalmente es unilateral. Presentación del caso: los autores describen el caso clínico de un paciente, seguido por una hipertensión resistente medicada con cuatro fármacos antihipertensivos para la estabilización de...
la tensión, con mantenimiento de las tensiones arteriales sistólicas superiores a 170 mmHg. Las alteraciones analíticas en el estudio etiológico de la hipertensión sugirieron hiperaldosteronismo y una lesión nodular en la glándula suprarrenal izquierda. El paciente fue sometido a cirugía y se realizó la escisión de la lesión con confirmación histológica del diagnóstico de adenoma cortical medular de la glándula suprarrenal. El paciente presentó una mejora en el perfil de tensión, con la necesidad de suspender dos de los cuatro fármacos antihipertensivos y reducir la dosis de los restantes. Discusión: un tumor de la corteza suprarrenal que produce la aldosterona es la principal causa de hiperaldosteronismo primario y siempre debe excluirse cuando se presenta hipertensión grave, difícil de controlar, ya que el tratamiento estándar es quirúrgico y conduce a una estabilización del patrón de tensión después de unos meses.

Palabras clave: adenoma de Conn, adenoma adrenal, hipertensión arterial, hipertensión secundaria, aldosteronismo primario, hipertensión resistente.

Resumo
Introdução: a hipertensão secundária corresponde ao 15% das causas de hipertensão arterial, e entre elas, o hiperaldosteronismo primário apresenta uma incidência variável de sobre 3% em pacientes hipertensos. Tem uma prevalência ligeiramente maior em mulheres, entre 30-60 anos, e geralmente é unilateral. Apresentação do caso: os autores descrevem o caso clínico de um paciente, seguido por uma hipertensão resistente medicada com quatro fármacos anti-hipertensivos para a estabilização da tensão, com manutenção das tensões arteriais sistólicas a 170 mmHg. As alterações analíticas no estudo etiológico da hipertensão sugeriram hiperaldosteronismo e uma lesão nodular na glândula suprarrenal esquerda. O paciente foi submetido a cirurgia e se realizou a incisão da lesão com confirmação histológica do diagnóstico de adenoma cortical-medular da glândula suprarrenal. O paciente apresentou uma melhora no perfil de tensão, com a necessidade de suspender dois dos quatro fármacos anti-hipertensivos e reduzir a dose dos restantes. Discussão: um tumor do córtex suprarrenal que produz a aldosterona é a principal causa de hiperaldosteronismo primário e sempre deve excluir-se quando se apresenta hipertensão grave, difícil de controlar, pois o tratamento standard é cirúrgico e conduz a uma estabilização do patrão de tensão depois de uns meses.

Palavras-chave: Conn’s adenoma, adrenal adenoma, hipertensão arterial, hipertensão secundária, aldosteronismo primário, hipertensão resistente.

Introduction
Secondary hypertension corresponds to 15% of the causes of arterial hypertension, and among them, primary hyperaldosteronism is the most common cause of reversible hypertension, with a very characteristic clinical frame. It has a slightly higher prevalence in women, between 30 and 60 years, and is usually unilateral.

An aldosterone producing tumor of the adrenal cortex is the main cause of primary hyperaldosteronism and should always be excluded when in the presence of severe hypertension, since the standard treatment is surgical and leads to a stabilization of the tension pattern after a few months.
Case presentation

The authors describe the clinical case of a 63-year-old male patient who presented with a severe hypertension with five years of evolution, treated with four antihypertensive agents (nifedipine 60mg daily, losartan 100mg daily, nebivolol 5 mg daily, and rilmenidine 1mg daily) introduced sequentially, maintaining systolic blood pressure [BP] greater than 170 mmHg, with progressive worsening in the last two years. He is an autonomous patient, with history of non-insulin treated type 2 diabetes mellitus and a stroke, four years previously, without chronic sequelae. In the etiological study of hypertension, several analytical alterations were identified, and they are explained in table 1.

Table 1. Representation of the analytical alterations

<table>
<thead>
<tr>
<th></th>
<th>Patient values</th>
<th>Reference values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potassium</td>
<td>3.19 mmol/L</td>
<td>3.5-5.1 mmol/L</td>
</tr>
<tr>
<td>Aldosterone</td>
<td>21.9 ng/mL</td>
<td>&gt; 20 ng/mL</td>
</tr>
<tr>
<td>Active renin</td>
<td>0.8 uU/mL</td>
<td>7-76 uU/mL</td>
</tr>
<tr>
<td>Aldosterone / active renin index</td>
<td>27.4 ng/mL/h</td>
<td>&gt; 25 ng/mL/h</td>
</tr>
<tr>
<td>Vanilmandelic acid (Urinary)</td>
<td>7.46 mg/24h</td>
<td>2.0-7.0 mg/24h</td>
</tr>
<tr>
<td>Metanephrines and normetanephrines (Urinary)</td>
<td>114 mcg/24h</td>
<td>&lt;350 mcg/24h</td>
</tr>
<tr>
<td></td>
<td>141 mcg/24h</td>
<td>&lt;600 mcg/24h</td>
</tr>
</tbody>
</table>

By ultrasonography, a nodular lesion was detected on the left adrenal gland (figure 1) which was subsequently confirmed by abdominal CT, showing a solid lesion of about 2.5 cm on the left adrenal gland. The patient was submitted to surgery and the lesion was excised with histological confirmation of a corticomедullary adenoma of the adrenal gland.

Figure 1. Echographic image of Conn’s adenoma
The patient presented improvement in his ambulatory blood pressure profile (with systolic BP around 130mmHg), and confirmed in 24 h ambulatory blood pressure monitoring (systolic BP 113-138 mm Hg and diastolic BP 66-78 mm Hg) seven months after the procedure, despite suspending two of the four antihypertensive drugs (nebivolol and rilmenidine, the latter still during the admission) and reducing the dose of the remaining ones (nifedipine to 30mg daily, losartan to 50mg daily). For the time being the patient maintains normalized aldosterone/active renin index and no hypokalemia.

Discussion

Hypertension causes significant morbidity and mortality worldwide, having a prevalence of about 30% of the adult world population, and around 85% of them above 55 years-old (1, 2). Hypertension has the largest prevalence of all cardiovascular risk factors and, due to its deleterious effects on the cardiovascular and renal systems, it is considered responsible for 45% of the heart disease related deaths and 51% of them due to stroke (1, 3).

The prevalence of hypertension increases with age and most individuals with hypertension are diagnosed with primary hypertension. Primary hypertension accounts for approximately 85% of the diagnosed cases. It is estimated that approximately 15% of hypertensive patients have identifiable conditions that result in blood pressure elevation (secondary hypertension) (4).

Common causes of secondary hypertension include obstructive sleep apnea, renal artery stenosis, chronic kidney disease, and endocrine alterations (3, 4, 5). Endocrine hypertension accounts for approximately 3% of the secondary forms of hypertension and is a term assigned to states in which hormonal derangements result in clinically significant hypertension (4). The most common causes of endocrine hypertension are excess production of mineralocorticoids (primary hyperaldosteronism), catecholamines (pheochromocytoma), thyroid hormone, and glucocorticoids (Cushing syndrome) (4).

Primary hyperaldosteronism is the most common cause of reversible hypertension, with a reported prevalence of 5% to 20%, and presents in a myriad of clinical scenarios (1, 4, 6). Adrenocortical adenoma is the most common adrenal tumor. Because the majority of adenomas are non-functioning, most of these lesions are detected incidentally on routine imaging performed for unrelated reasons (7).

Primary aldosteronism is characterized by autonomous production of aldosterone by the adrenal glands along with suppressed renin production by the renal juxtaglomerular apparatus. This leads to volume expansion, severe hypertension, metabolic alkalosis, and hypokalemia (1, 5, 6, 8). Classically, excessive aldosterone secretion in primary hyperaldosteronism not only results in difficult to manage hypertension in the majority of
patients, but also produces biochemical effects of hypokalemia in 10% to 30% of patients (1). In many occasions, primary hyperaldosteronism is completely asymptomatic. When symptoms occur, they are usually secondary to hypertension or the resulting hypokalemia: cramps and fatigue; cardiac arrhythmias, manifested by palpitations; progressive weakness, even reporting cases of generalized paralysis; polydipsia and polyuria due to diabetes insipidus induced by hypokalemia (8). It is important to identify the existence of primary aldosteronism among resistant hypertensive patients since it is frequently curable, with clear reduction in disease burden (9).

Mineralocorticoid receptor antagonists absolutely interfere with the interpretation of the aldosterone/renin plasmatic index and should be discontinued at least six weeks before testing (8, 10). Angiotensin converting enzyme inhibitors and angiotensin receptor blockers falsely increase plasma renin activity; therefore, a useful clinical point is that when plasma renin activity is undetectably low in a patient taking this type of medication, primary aldosteronism is suspected (10). Although it is the most used screening method in patients with suspected primary hyperaldosteronemia, given the low prevalence of this condition, the screening tests are reserved for those patients with resistant hypertension or hypertension associated with hypokalemia (even when hypokalemia is suspected to be secondary to the use of diuretics) (8).

The elevation of the aldosterone/renin index is a sensitive but not specific test, so that confirmatory tests are necessary in patients with elevated indexes (10).

There are four suppression tests used to confirm the alterations: oral sodium load, oral fludrocortisone load, oral captopril load and saline infusion to suppress aldosterone secretion; with the first and last being the most commonly used (10).

There is also an adrenal venous sampling, which is considered the gold-standard test to distinguish between unilateral or bilateral disease in patients with primary aldosteronism who desire surgical treatment of their hypertension. It is done by catheterization of femoral venous access, sampling blood from both adrenal veins and from the inferior vena cava below the renal veins and measurement of aldosterone and cortisol levels. The aldosterone/cortisol index should be elevated in secretory tumor (10).

Adrenalectomy is the procedure of choice for unilateral aldosterone-secreting adenomas, while medical therapy is best for bilateral adrenal hyperplasia (1). Biochemical cure following adrenalectomy occurs in 99% of patients, and hemodynamic improvement is seen in over 90%, prompting a reduction in quantity and dosing of anti-hypertensive medications in most patients (1). End-organ damage secondary to hypertension and excess aldosterone is significantly improved by both surgical and medical treatment (1, 4). Resistant hypertension is an increasingly common clinical problem that is often heterogeneous in etiology, risk factors and comorbidities. In light of diagnostic advances, evidence points to higher prevalence of secondary causes of hypertension, about 10 to
35% (5). For diagnosing, patient's history and physical examination are important, as are assessing compliance, regular blood pressure measurement, biochemical evaluation and noninvasive imaging (3).

Although the most common causes of therapeutic failure are undiscovered secondary causes of hypertension and lack of patient compliance, other causes should be suspected, tested and, if confirmed, curable forms of arterial hypertension should be treated (3). Given the large impact on global health, controlling hypertension is of extreme importance (1).

**Declaration of Interest**

The authors report no conflicts of interest.

**References**
