Primary Hyperaldosteronism: Pitfalls in the Diagnosis and a not so Peculiar Evolution - Case Report

Joao Martin Martinsa,b, Rui Melo Cabrala, Sonia do Valea, Ana Filipa Martinsa, Ana Raquel Gomesa

Abstract

Primary hyperaldosteronism is now assumed to account for 10-15% of all cases of non selected high blood pressure. The differential diagnosis between bilateral hyperplasia and unilateral adenoma remains however a medical challenge, and the pathogenic stimulus for hyperplasia is still unknown. HS, a female patient aged 32, presented with High Blood Pressure and hypokalemia. An elevated aldosterone/renin ratio and non-suppressed aldosterone levels after saline infusion confirmed the diagnosis of primary hyperaldosteronism. Negative adrenal scintigraphic examination and a marked aldosterone response in the postural test supported the diagnosis of bilateral hyperplasia, despite the presence of left adrenal nodule in the CT-scan. Tetracosactide stimulation revealed a minor 21-hydroxilase defect and a marked ACTH-dependence of aldosterone secretion. Effective treatment with dexamethasone 1.0 mg every other night was maintained for 9 years, while in the meantime the left adrenal nodule apparently resolved and a right adrenal nodule become apparent. Finally worsening of blood pressure levels and increasing aldosterone concentrations with a positive scintigraphic labelling on the right adrenal gland led to right adrenalectomy that was so far curative. This case report demonstrates bilateral hyperplasia, despite the presence of left adrenal nodule in the CT-scan. Tetracosactide stimulation revealed a minor 21-hydroxilase defect and a marked ACTH-dependence of aldosterone secretion. Effective treatment with dexamethasone 1.0 mg every other night was maintained for 9 years, while in the meantime the left adrenal nodule apparently resolved and a right adrenal nodule become apparent. Finally worsening of blood pressure levels and increasing aldosterone concentrations with a positive scintigraphic labelling on the right adrenal gland led to right adrenalectomy that was so far curative. This case report demonstrates bilateral hyperplasia-adenoma evolution and suggests ACTH to be the relevant pathogenic stimulus, allowing for effective long-term treatment with dexamethasone.

Keywords: Primary hyperaldosteronism; Hyperplasia; Adenoma

Introduction

High blood pressure (HBP) occurs in 25% of the western developed world population [1]. It is now considered that up to 15% may correspond to secondary forms [2-4]. In particularly, recent studies suggest, primary hyperaldosteronism (PA), first described by Conn in 1955, is a rather common condition accounting for more than 10% of all cases [2-4]. This contrasts sharply with previous consensus, where essential or “idiopathic” hypertension was assumed to account for about 95% of all cases of non selected HBP [5].

Such a large difference may occur because a simple screening method is now available, namely the aldosterone/renin ratio and because we now believe many cases of PA occur without the classic hypokalemia, obviously without oedema, and even without HBP, so that in fact we may be diagnosing very early an asymptomatic condition [2-4, 6]. This occurs commonly in the context of the evaluation of adrenal incidentalomas [7].

HBP due to PA tends to be serious, with more marked evidence of target organ lesions and vascular events and the added risk of arrhythmias [8-10]. A postulated mechanism involving the effects of aldosterone on inflammation and tissue fibrosis and/or in the development of the cardiometabolic syndrome, has been invoked [11-13]. Besides either surgically curative or highly effective medical treatment is available highlighting the importance of this diagnosis.

After the condition is suspected confirmatory dynamic tests are necessary and the distinction between the different forms may be challenging since besides the classical unilateral adenoma (35%) and the bilateral hyperplasia (60%), a somehow paradoxical unilateral hyperplasia and a glucocorticoid remediable form have been described and treatment options are radically different [2-4]. Furthermore regarding the unilateral or bilateral hyperplasia, the stimulus for the persistent stimulation remains unidentified, and it seems possible that endocrine hyperplasia may evolve to autonomous adenomas [14]. In this regard, imaging methods, for example, adrenal computed tomography (CT) and nuclear magnetic resonance (NMR), may add to the confusion, given the high incidence of silent incidentalomas, the high probability of missing small (< 1 cm) adenomas, and the inability to definitively distinguish adenomas and uni- or bilateral hyperplasia [2-4, 15, 16].

Diagnostic work out generally includes screening tests in selected high risk groups, mainly ambulatory paired ran-
dom plasma aldosterone and renin, at specific timetables and postural conditions, optimally with an unrestricted salt diet and no anti-hypertensive drugs, but certainly without aldosterone receptor antagonists like spironolactone or eplerenone - sensitivity and specificity above 90% [2-4, 17-19]. Confirmatory tests must then be used, either the intravenous saline infusion test or an alternative like the oral sodium loading test or fludrocortisone suppression test - sensitivity and specificity again above 90% [2-4, 20]. Finally one must proceed to the differential diagnosis: 1) adrenal CT - only 25% sensitivity for small adenomas < 1 cm and an accuracy of 50% regarding the distinction between adenomas and bilateral hyperplasia [2-4, 21-24]; 2) adrenal venous sampling - now considered the reference test, it is not without risks, it is technically demanding and theoretically confusing with sequential instead of simultaneous sampling used, the need to correct for cortisol and tetracosactide stimulation - although a sensitivity and specificity above 95% have been reported [2-4, 21-24]. Regarding catecholamine secretion however, lateralization was found to be a normal finding [25]. Other historical tests seem to have lost favour like $^{131}$I-19-iodocholesterol or 6β-$^{131}$I iodomethyl-19-norcholesterol (NP-59) scintigraphy, the postural stimulation test and the measurement of 18-hydroxycorticosterone [2-4]. However regarding tests used in the differential diagnosis one must accept that they are either technically demanding and available only at selected sites or theoretically confusing [26].

We now report a case of PA where diagnostic workout proved confusing, but in the end probably very easy. This case demonstrates that bilateral hyperplasia - adenoma evolution can occur, and suggests ACTH to be the pathogenic stimulus, underlining the long term effectiveness of dexamethasone treatment.

## Case Report

HS, a Caucasian female patient, aged 32, unskilled worker, divorced, living in Lisbon, was referred to the Endocrine Out Patient Department of a public central hospital, on October

![Figure 1. Abdominal CT scan. (a) 1999 - a small nodular lesion around 1 cm is noted in the left adrenal gland; no abnormalities were found regarding the right adrenal gland. (b) 2001 - a nodular lesion hypodense with 15 mm maximal diameter, is noted in the right adrenal gland, which is not apparent in the previous scan; no apparent lesions were found regarding the left adrenal gland.](image)

### Table 1. Saline Suppression Test (NaCl 0.9%, 2,000 mL, ev, Over 4 h)

<table>
<thead>
<tr>
<th>Time (h)</th>
<th>Renin (μU/mL)</th>
<th>Aldosterone (pg/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1</td>
<td>310</td>
</tr>
<tr>
<td>2</td>
<td>0.8</td>
<td>150</td>
</tr>
<tr>
<td>4</td>
<td>0.7</td>
<td>226</td>
</tr>
<tr>
<td>RV</td>
<td>(D - 1 - 20 μU/mL)</td>
<td>(D - 10 - 160 pg/mL)</td>
</tr>
</tbody>
</table>

Renin and aldosterone were measured by radioimmunoassay (Diagnostic Products Corporation, Los Angeles); intra- and interassay coefficients of variation were less than 10%. Baseline decubitus (D) reference values (RV) are presented.
1999, because of HBP and hypokalemia. HBP was diagnosed 6 months before, on a routine medical examination and confirmed by several random measurements of 150 - 160/90 - 100 mmHg. The patient was asymptomatic with no headaches, no visual disturbances, no oedema, no dyspnea, no chest pain, no palpitations and no syncope. There were no episodes of sudden high blood pressure with or without anxiety and adrenergic symptoms, no weight loss, no renal disease, including renal stones, and no changes of body weight or menstrual cycles. At the time the patient was not using any medications, including contraceptive pills and she did not smoke nor did she regularly drink alcoholic beverages. She did not present any other known medical illness and was never submitted to a surgical intervention. Menarche occurred at age 12, followed by regular menstrual cycles. She was married at 20 years and used contraceptive pills for some time. She was pregnant once at 21 years, with an uneventful pregnancy not complicated by high blood pressure, gestational diabetes or pre-eclampsia. Both parents were still alive, aged 53 and 50 years, with no high blood pressure. There was no family record for other endocrine or renal diseases.

The patient presented several prior analytical reports with no relevant abnormalities, except for persistent hypokalemia (2.6 - 2.8 mEq/L (reference range 3.5 - 4.5 mEq/L)). A previous abdominal CT scan revealed a nodular lesion of 1 cm on the left adrenal gland with no other abnormalities (Fig. 1a). She was using Ramipril 5 mg, once daily with apparent success regarding the control of blood pressure levels and no significant side effects. Slow-releasing Nifedipine 30 mg was substituted for, and admission to the Endocrine In Patient department was arranged for two weeks later, after a weekend without any medication.

On admission, physical examination revealed an apparently healthy adult female subject, alert, oriented, with no cognitive disturbances and an adequate emotional behaviour. She looked her age, with slight excess weight, and moved normally with no apparent limitations. Her body temperature was normal with no respiratory distress, no cyanosis, and no peripheral oedema. Height: 1.54 m; Weight: 64 kg (without shoes or coats); Umbilical perimeter: 81 cm; Hip perimeter: 99 cm. Dorsal decubitus blood pressure was 190/110 mmHg in the right arm and 180/110 mmHg in the left arm and 190/110 mmHg in the right leg; standing right arm blood pressure was 170/100 mmHg. Heart beats varied between 82 - 86 bpm and respiratory excursions were large at 16 cpm. Head and neck examination did not reveal any relevant findings, namely regarding the thyroid or the neck vessels. The thorax was of normal shape with no skin or breast lesions and no apparent abnormalities on cardiopulmonary auscultation, except for a systolic murmur grade II/VI and a slightly increased S2 sound at the second right intercostal space. The abdomen was smooth, painless on palpation with no tenderness and moved freely with the respiratory excursion. There

### Table 2. Ambulatory Testing (4 h Brisk Walking)

<table>
<thead>
<tr>
<th></th>
<th>Before</th>
<th>After</th>
<th>RV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Renin (μU/mL)</td>
<td>1</td>
<td>2</td>
<td>(D -1 - 20 μU/mL)</td>
</tr>
<tr>
<td>Aldosterone (pg/mL)</td>
<td>280</td>
<td>367</td>
<td>(D - 10 - 160 pg/mL)</td>
</tr>
</tbody>
</table>

Reference values (RV) are present for the decubitus (D) and standing (S) position.

### Table 3. Tetracosactide Test (250 μg ACTH1-24, ev)

<table>
<thead>
<tr>
<th></th>
<th>0 min</th>
<th>30 min</th>
<th>60 min</th>
<th>RV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cortisol (μg/dL)</td>
<td>19</td>
<td>58</td>
<td>56</td>
<td>(8h am - 4 - 23 μg/dL)</td>
</tr>
<tr>
<td>17-hydroxiprogesterone (ng/mL)</td>
<td>3</td>
<td>10</td>
<td>7</td>
<td>(FF - 0.2 - 1.8 ng/mL)</td>
</tr>
<tr>
<td>11-deoxi-cortisol (ng/mL)</td>
<td>1</td>
<td>4</td>
<td>4</td>
<td>(&lt; 8 ng/mL)</td>
</tr>
<tr>
<td>Renin (μU/mL)</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>(1 - 20 μU/mL)</td>
</tr>
<tr>
<td>Aldosterone (pg/mL)</td>
<td>210</td>
<td>1,030</td>
<td>80</td>
<td>(10 - 160 pg/mL)</td>
</tr>
</tbody>
</table>

Cortisol was measured by chemiluminoscence immunoassay and 17-hydroxyprogesterone and 11-deoxi-cortisol were measured by radioimmunoassay (Diagnostic Products Corporation, Los Angeles). Reference values (RV) are presented at 8h am, or in the follicular phase (FF).
was no evidence of enlarged liver or spleen, and no bruits were found over the epigastric, peri umbilical or lumbar areas. The upper and lower members were apparently normal, without oedema and peripheral pulses were rhythmic and symmetric. Neurologic examination was negative.

During her 7-day hospital stay, repeated routine analytical evaluation failed to show any relevant finding except for persistent hypokalemia (2.4 - 2.8 mEq/L); in particular, there were no hematologic abnormalities, kidney and liver function tests were normal, as was acid-base status, mineral metabolism, glucose, proteins and lipids. Thorax x-ray was normal with no evidence of an increased or abnormal cardiac shadow and the EKG showed a normal sinus rhythm (75/min), with normal PR, QRS and QT intervals, an electric axis at +30º in the vertical plan and a transition between negative and positive complexes at V3-V4 in the horizontal plane. R was always less than 15 mm and R+S less than 35 mm; no other abnormalities could be found. Fundoscopic examination was negative for signs of hypertensive disease and 24 h urinary microalbumin was negative.

Twenty-four hours blood pressure measurements were obtained with mean values in the 24 h period of 123/80 mmHg, 134/89 mmHg during the day period (8 - 22 h) and 109/68 mmHg in the night period (22 - 8 h). Systolic blood pressure was greater than 140 mmHg at 30% of the measurements and diastolic blood pressure was greater than 90 mmHg at 31% of the time.

Routine endocrine evaluation was normal regarding thyroid, parathyroid, pituitary or gonadic function. ACTH and cortisol were also normal with the classical circadian rhythm; 24 h-urinary excretion of adrenaline, noradrenaline, metanephrine, normetanephrine and vanilmandelic acid were normal and so were androgen levels including total and free testosterone, androstenedione, dehydroepiandrosterone-sulphate, 17-hydroxyprogesterone and 11-deoxicortisol.

However analytical evaluation revealed repeated low renin levels: 0.5 - 0.8 μU/mL (reference range 1 - 20 μU/mL) with high aldosterone levels: 190 - 240 pg/mL (reference range 10 - 160 pg/mL) and an increased 24 h-urinary excretion of aldosterone: 48 - 56 μg/24 h (reference range 6 - 25 μg/24 h).

### Table 4. Saline Suppression Test (NaCl 0.9%, 2,000 mL, ev. Over 4 h)

<table>
<thead>
<tr>
<th></th>
<th>0 h</th>
<th>2 h</th>
<th>4 h</th>
<th>RV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Renin (μU/mL)</td>
<td>0.7</td>
<td>0.9</td>
<td>0.5</td>
<td>(1 - 20 μU/mL)</td>
</tr>
<tr>
<td>Aldosterone (pg/mL)</td>
<td>350</td>
<td>320</td>
<td>560</td>
<td>(10 - 160 pg/mL)</td>
</tr>
</tbody>
</table>
μg/24 h). These measurements were obtained in the supine position with a normal saline diet and no anti-hypertension drugs.

A saline suppression test was performed after an overnight fast while the patient remained in bed, with 2 L normal saline infused over 4 h. Results are presented in Table 1.

An ambulatory test was then performed again after an overnight fast, with blood samples obtained first while the patient was still in bed, and then after 4 h of brisk walking. The following results were obtained (Table 2).

A Synacthen test was finally performed again after an overnight fast. Tetracosactide (ACTH1-24) 250 μg was administered ev at time 0, and blood was then collected at 30 and 60 min. Results are presented in Table 3.

Nor-(125I)iodomethyl-colesterol isotope imaging under dexamethasone 0.5 mg, 6/6 h for 48 h, revealed “slight fixation at both adrenals, more so on the right, that is probably non relevant given the information regarding a nodular lesion on the left adrenal” (Fig. 2a).

A presumptive diagnosis of PA - bilateral hyperplasia was assumed given the ambulatory test, the tetracosactide test and non lateralization of scintigraphic labelling of the adrenals, while the nodular lesion at the left adrenal was not judged functionally relevant. Medical treatment was decided upon with a hyposaline diet and dexamethasone 1.0 mg at bedtime every other day. Under this treatment, and for 2 years blood pressure was normal (< 140/90 mmHg) with normal serum potassium and no relevant side-effects.

On January 2001, the patient now aged 33, was again admitted to the Endocrine Unit after a week end free of medication. The saline suppression test (Table 4) and the ambulatory test (Table 5) were again repeated. An adrenal CT scan was again obtained and a nodular lesion of 15 mm diameter, with low density and smooth contours was found in the right adrenal, with no apparent abnormalities on the left (Fig. 1b). Previous diagnosis and therapy were resumed after discharge.

On March 2002, a planned pregnancy occurred after dexamethasone substitution for alpha-methyldopa 500 mg tid. Pregnancy was uneventful with normal blood pressure levels and no pre-eclampsia or eclampsia. A boy was born with normal weight and normal vitality at 5 and 10 min. Fallopian tubes were ligated at the time and dexamethasone was again resumed 1.0 mg, at bedtime every other day.

On August 2005, the patient now aged 37 was again admitted to the Endocrine Department because of acute gastroenteritis with serious hypokalemia (1.8 mEq/L). The patient denied missing any drug schedule and she was not hypertensive. There was no fever and the stools were without blood, mucus or pus. C-Reactive protein was negative. Again high serum aldosterone levels (280 pg/mL) and low renin levels (0.7 μU/mL) were measured. Three days later after correction of the clinical condition and serum potassium the patient was discharged and previous medication was resumed.

Again for 2 years, evolution was uneventful with normal blood pressure levels and normal serum potassium, and no relevant side-effects.

Since 2008, blood pressure levels began to increase despite drug treatment and spironolactone 100 mg/day and later bisoprolol 5 mg/day were added. At this time there was still no evidence of target-organ lesions, namely regarding the heart or the kidneys. At the end of 2008 the patient was again admitted to the Endocrine Department because of resistant high blood pressure, although with no specific symptoms and still with a normal serum potassium levels. Dynamic tests were again repeated and are shown in Tables 6 and 7. A new scintigraphic examination with 19-noriodomethylcholesterol was performed that revealed persistent and

<table>
<thead>
<tr>
<th>Table 5. Ambulatory Testing (4 h Brisk Walking)</th>
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<tr>
<td>Before</td>
</tr>
<tr>
<td>--------</td>
</tr>
<tr>
<td>Renin (μU/mL)</td>
</tr>
<tr>
<td>Aldosterone (pg/mL)</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Table 6. Saline Suppression Test (2 L Normosaline Infused ev Over 4 h)</th>
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<tbody>
<tr>
<td>0 h</td>
</tr>
<tr>
<td>------</td>
</tr>
<tr>
<td>Renin (μU/mL)</td>
</tr>
<tr>
<td>Aldosterone (pg/mL)</td>
</tr>
</tbody>
</table>
abnormal fixation at the right adrenal gland (Fig. 2b). Surgery was decided and performed on March 2009; the right adrenal gland with 24 × 1 mm and 9 g was removed and an adenoma was identified. There was no reference to adrenal hyperplasia on the remaining gland.

Since the surgery the patient remains normotensive 130 - 140/80 - 85 mmHg without medication and a 24 h blood pressure ambulatory monitoring was again preformed with normal blood pressure levels along the 24 h period.

Discussion

This case begins with the classical presentation of hyperaldosteronism: a young woman presents with recent onset HBP and hypokalemia. Hypokalemia is indeed highly suggestive of hyperaldosteronism, even if it is present in only between 10-40% of the cases [2-4]. Furthermore the patient presented with 40% of apparent normal subjects present nodular lesions in the adrenal (incidentalomas); 2) while adrenal adenomas responsible for PA are generally small, less than 2 cm, uni- or bilateral hyperplasia may also present small nodules; 3) furthermore responsible or innocent lesions may be at the limit of detection and thus missed [15, 16, 21-24]. In fact CT scans have only a low and variable 45-87% sensitivity and a variable albeit higher specificity 71-92% [2-4]. CT scans results are discordant with adrenal vein sampling in up to 23% of the cases and using only the CT scan up to 34% of inappropriate surgeries would be preformed [2-4].

The adrenal vein sampling may be considered as the standard test with a reported sensitivity and specificity over 90% [2-4]. This procedure is however not without risks, technically demanding, requiring dedicated radiologist with acceptable personal experience and is theoretically confusing, since sequential instead of simultaneous sampling is generally used, the aldosterone/cortisol ratio must be used to account for differences regarding the activity level between the adrenals, and since bolus or continuous tetracosactide stimulation is necessary to correct for stress-induced fluctuations in adrenal function [2-4, 21-24]. In fact regarding pheochromocytoma and catecholamine secretion by the adrenals, a recent reported study found that in normal glands high lateralization ratios also occur [25]. This method was however not available at the time in our institution.

We performed an ambulatory test that revealed an aldosterone response suggesting at least partial angiotensin dependence and therefore favours adrenal hyperplasia. The ambulatory test by itself is not a definitive test but a response such as the one found in this case favours adrenal hyperplasia (88% of the cases) while a decrease suggesting partial ACTH sensitivity favours adenoma (83% of the cases). This test seems to have an overall accuracy of 85% [2-4].

Scintigraphic examination with nor-iodo-methylcholesterol was performed under dexamethasone and revealed “slight fixation on both adrenals, more so on the right.” This examination has an overall precision of 73%, but may miss small lesions. It suggested however the non functional nature of the lesion identified in the left adrenal gland by CT scan and again favours the diagnosis of bilateral hyperplasia [2-4].

Table 7. Ambulatory Test (4 h of Brisk Walking)

<table>
<thead>
<tr>
<th></th>
<th>0 h</th>
<th>4 h</th>
<th>RV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Renin (μU/ml)</td>
<td>0.5</td>
<td>0.4</td>
<td>(D -1 - 20 μU/mL)</td>
</tr>
<tr>
<td>Aldosterone (pg/mL)</td>
<td>631</td>
<td>1,000</td>
<td>(D - 10 - 160 pg/mL)</td>
</tr>
</tbody>
</table>

-nalectomy is in this setting a reasonable therapeutic option [2]. This method is however of little value at least for itself: 1) more than 10% of apparent normal subjects present nodular lesions in the adrenal (incidentalomas); 2) while adrenal adenomas responsible for PA are generally small, less than 2 cm, uni- or bilateral hyperplasia may also present small nodules; 3) furthermore responsible or innocent lesions may be at the limit of detection and thus missed [15, 16, 21-24]. In fact CT scans have only a low and variable 45-87% sensitivity and a variable albeit higher specificity 71-92% [2-4]. CT scans results are discordant with adrenal vein sampling in up to 23% of the cases and using only the CT scan up to 34% of inappropriate surgeries would be preformed [2-4].
We selected to perform a Synacthen test that is generally not used in this specific diagnostic work out. Based on our personal previous experience we wanted to exclude minor/late onset forms of congenital adrenal hyperplasia and also to evaluate the ACTH-dependence of aldosterone secretion [27]. The tetracosactide test revealed both a possible minor 21-hydroxylase defect [28] and a marked aldosterone response to synthetic ACTH. Several remarks may be made at this point: 1) ACTH dependence in the context of ambulatory testing is generally considered to suggest an adenoma [2-4]. 2) A glucocorticoid remediable hyperaldosteronism could have been suspected; however this is rare, less than 1% of all cases of primary aldosteronism, and clinical presentation is different with a positive family history and early onset of severe high blood pressure refractory to conventional treatment [2-4]; although genetic testing or the measurement of 18-hydroxycortisol or 18-oxyocortisol were not available and were not performed, we suppose this is not the case; in fact we have previously found this marked ACTH-dependence to be very common in patients with minor/late onset forms of congenital adrenal hyperplasia [27]; the fact remains however of a marked ACTH-dependence for aldosterone secretion.

A presumptive diagnosis of PA, probable bilateral adrenal hyperplasia was established, surgery was considered inappropriate and medical treatment was decided. In fact in favour of adrenal hyperplasia were the results of the ambulatory tests and negative scintigraphic examination, while hypokalemia and a lesion on the left adrenal gland favoured adenoma. Since a decision on medical treatment can always be later reversed while the opposite is not true and medical treatment is available although not curative, our decision seems justified.

Medical treatment was begun with dexamethasone 1.0 mg once in the evening, every other day and a sodium restricted diet. This treatment is unconventional and spironolactone is generally used instead; spironolactone in the female gender presents unpleasant side effects namely oligomenorhea [2-4]; dexamethasone was preferred instead because of the dramatic aldosterone response in the tetracosactide test, and was supported by the minor/late onset defect of 21-hydroxylase. In fact this therapy was very effective for 9 years, again suggesting a strong ACTH-dependence of aldosterone secretion that maybe in some cases, namely those with minor defects of adrenal steroidogenesis may be the relevant stimulus for the hyperplasia. Meanwhile an episode of acute gastroenteritis with hypokalemia occurred and a planned pregnancy was uneventful with dexamethasone that in this case at least, was long term effective.

Findings of the adrenal CT scan may be very misleading. Technical reasons are well known but it may be worth emphasising that at the detection limit of the method, adrenal nodules may “appear” and “disappear”, namely, may become apparent or inapparent along time in bilateral hyperplasia. Aldosterone secretion may at least in some cases, like this one, and more usual in the context of minor late onset defects of adrenal steroidogenesis be strongly ACTH-dependent. It is worth noting that aldosterone increase after ACTH was at least as high as the cortisol rise. ACTH may indeed be the stimulus leading to bilateral hyperplasia. At the same token this data suggests an alternative therapeutic option like dexamethasone that in this case at least, was long term effective.

An apparent evolution from bilateral hyperplasia to autonomous unilateral adenoma is strongly suggested in this case. Initial work out suggested adrenal hyperplasia, namely the ambulatory test and the scintigraphic examination. Follow up of the patient reinforces this evidence. A nodule was
first found on the left adrenal and later “resolved” while a nodule was found in the right adrenal. Strong ACTH dependence would lead first to a general hyperplasia not to a solitary nodule. Dexamethasone therapy would not be expected to be effective regarding an autonomous adenoma. In fact in the long term evolution dexamethasone lost efficacy while aldosterone levels markedly increased both baseline and post-stimulation. Adenoma autonomisation seems best fit in this change. Unfortunately we had no report regarding evidence of hyperplasia in removed gland, but probably one should look specifically for it. It is possible that in the long term other adenoma may appear and if so blood pressure will rise again. Follow up must therefore be maintained.

Consent

Written informed consent was obtained from the patient for publication of this case report and the accompanying images.

Competing Interests

All authors declare that they have no competing interests.

Authors’ Contributions

JMM assured the medical care of the patient, including the selection of appropriate diagnostic tests and therapeutic interventions. He also wrote the first and final version of this MS.

RMC wrote an intermediate version of the MS and collected the relevant bibliography. He also contributed deeply to the final version of the MS.

SV assisted JMM regarding the medical care of the patient, including the clinical discussion of relevant diagnostic tests and therapeutic measures. She contributed all along the writing of the first and final version of the MS.

AFM and ARG contributed fundamentally to the final version of the MS.

All authors give final approval of the version hereby submitted for publication.

References