CLINICAL STUDY

Confirmatory testing in primary aldosteronism: extensive medication switching is not needed in all patients

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Abstract

Objective: Confirmatory testing of suspected primary aldosteronism (PA) requires an extensive medication switch that can be difficult for patients with severe complicated hypertension and/or refractory hypokalemia. For this reason, we investigated the effect of chronic antihypertensive medication on confirmatory testing results. To allow the results to be interpreted, the reproducibility of confirmatory testing was also evaluated.

Design and methods: The study enrolled 114 individuals with suspected PA who underwent two confirmatory tests. The patients were divided into two groups. In Group A, both tests were performed on the guidelines-recommended therapy, i.e. not interfering with the renin–angiotensin–aldosterone system. In Group B, the first test was performed on chronic therapy with the exclusion of thiazides, loop diuretics, and aldosterone antagonists; and the second test was performed on guidelines-recommended therapy. Saline infusion, preceded by oral sodium loading, was used to suppress aldosterone secretion.

Results: Agreement in the interpretation of the two confirmatory tests was observed in 84 and 66% of patients in Groups A and B respectively. For all 20 individuals in Group A who ever had end-test serum aldosterone levels ≥240 pmol/l, aldosterone was concordantly nonsuppressible during the other test. Similarly, for all 16 individuals in Group B who had end-test serum aldosterone levels ≥240 pmol/l on modified chronic therapy, aldosterone remained nonsuppressible with guidelines-recommended therapy.

Conclusion: Confirmatory testing performed while the patient is on chronic therapy without diuretics and aldosterone antagonists can confirm the diagnosis of PA, provided serum aldosterone remains markedly elevated at the end of saline infusion.

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Introduction

Primary aldosteronism (PA) is not an infrequent form of secondary arterial hypertension. The cardiovascular consequences of this disorder go beyond blood pressure elevation (1, 2). It is important to identify PA cases as affected individuals can benefit from specific medical or surgical treatment. The PA diagnostic algorithm consists of screening, confirmation, and subtype differentiation.

The main obstacle when diagnosing PA is the need for a medication switch before confirmatory testing. According to the current guidelines (3), only drugs with minimal or no effects on the renin–angiotensin–aldosterone system (RAAS) can be administered before and during the confirmatory testing.

However, the medication change as suggested by the guidelines carries a risk of uncontrolled hypertension and severe hypokalemia. For this reason, confirmatory testing is often contraindicated in individuals with severe complicated hypertension or a history of significant hypokalemia.

Conversely, the effects of chronic antihypertensive therapy on confirmatory testing are not completely understood. and it is not known whether an extensive medication change is necessary in all patients referred for confirmatory testing.

These facts inspired us to investigate this aspect of the PA diagnostic process. To interpret the effects of chronic therapy on confirmatory testing, we also evaluated the reproducibility of confirmatory testing performed under the recommended conditions.

Oral sodium loading with subsequent saline infusion was used to suppress aldosterone secretion. This simple procedure can be performed on an outpatient basis.

Subjects and methods

Patients with suspected PA who were referred for confirmatory testing on the basis of screening results were prospectively enrolled. The study was approved by the Local Ethics Committee, and all participants...
provided written informed consent. All enrolled individuals were examined according to the protocol described below.

PA screening
The aldosterone-to-renin ratio (ARR) was used to screen patients with possible PA. Before ARR testing, no changes in the patients’ medications were required except for withdrawal from any aldosterone antagonist for at least 2 months. These drugs were not allowed until the PA diagnostic procedures were completed.

Blood sampling for ARR was always performed in the morning (between 0700 and 1000 h) when the patients had been in the upright position for at least 30 min. To calculate ARR, both serum aldosterone and plasma renin concentrations were expressed in nanograms per liter. On the basis of our observations in healthy volunteers, ARRs > 20 were considered abnormal (4).

Referral for confirmatory testing resulted mainly from repeatedly elevated ARR in screened patients. However, testing was also indicated for selected hypertensive individuals who presented with otherwise unexplained hypokalemia or an adrenal adenoma.

Confirmatory testing
Two confirmatory tests were performed for all enrolled patients. The patients were divided into two groups (A and B) that differed in the medication the patients were given before testing. The patients were allocated into the study groups consecutively, i.e. after enrolment in one group was completed, enrolment in the other group was initiated. The investigators had no influence on the allocation of the patients into the respective groups.

Group A
For patients in Group A, both confirmatory tests were performed under identical RAAS-neutral conditions, i.e. excluding any antihypertensive therapy interfering with RAAS. These drugs had to be stopped at least 2 weeks before the first test. When the risks of uncontrolled blood pressure were considered high, the patients were given doxazosin and/or verapamil. The suggested period between the two tests was 2 weeks. No changes in the antihypertensive medications were allowed during the period between tests. All patients who required any antihypertensive therapy adjustment before the second suppression test were excluded from the final analysis.

Group B
For patients in Group B, the first confirmatory test was performed under minimally modified chronic antihypertensive therapy. Excluding aldosterone antagonists that were not allowed at any point in the diagnostic process, thiazides and loop diuretics had to be stopped at least 2 weeks before the test. In patients with a history of severe hypokalemia, amiloride was permitted at a dose not exceeding 15 mg/day.

The second confirmatory test was performed with the exclusion of any antihypertensive therapy potentially interfering with the RAAS. These drugs had to be discontinued at least 2 weeks before testing. The only antihypertensive medications allowed were doxazosin and/or verapamil. The results of the second confirmatory test were used as the reference for clinical decision-making.

In both study groups, oral potassium supplementation was administered as needed at any time during the confirmatory testing.

Figure 1 Study protocol. Flowchart describes the study protocol in brief, including the study procedures and the patients excluded from the analysis. *Denotes a therapy with no or minimal effect on the RAAS, only doxazosin and/or verapamil were allowed. †Only amiloride was permitted at a dose not exceeding 15 mg/day.
All confirmatory tests in the study were performed according to the protocol that is routinely used in our center.

Oral sodium loading preceded the i.v. administration of normal saline, as a short-lasting infusion itself might not be sufficient to suppress renin and aldosterone secretion.

The patients were advised to enhance their oral salt intake after adjusting their medication before testing. Three days before the saline infusion was administered, oral salt intake was further increased by at least 6 g/day. The day before the saline infusion, a 24-h urine sample was collected to measure urinary sodium excretion.

Saline was always infused in the morning. After 2 h of bed rest, 2 l of normal saline was infused over a 4-h period. The patients remained in the supine position until the end of the infusion, when blood samples were taken for the estimations of plasma renin and serum aldosterone.

The criteria for interpreting confirmatory tests were based on our observations in healthy volunteers (4) as follows: patients who had plasma active renin levels ≥7 ng/l at the end of any confirmatory test performed during RAAS-neutral therapy were excluded from the final analysis. The exclusion limit reflects the 97.5 percentile of the values observed in healthy volunteers who underwent the same diagnostic protocol. Serum aldosterone was considered suppressible at levels ≤100 pmol/l at the end of the saline infusion. This cutoff is routinely used in our center. It was set arbitrarily between the 97.5 percentile of the values of healthy volunteers (60 pmol/l) and the lowest cutoff mentioned in the current guidelines (139 pmol/l).

Adrenal venous sampling

Adrenal venous sampling (AVS) was offered to all patients who had nonsuppressible serum aldosterone at the end of any suppression test performed during RAAS-neutral therapy.

The AVS protocol used in our center has been described in detail elsewhere (5). In brief, a continuous cosyntropin infusion was given during the procedure. The adrenal veins were sequentially cannulated. Adrenal samples were considered adequate if the cortisol concentration was at least fivefold greater than that of the inferior vena cava. Adrenalectomy was recommended to patients whose cortisol-corrected aldosterone concentration from the affected gland was at least fourfold higher than that of the contralateral side.

Laboratory measurement

For all collected blood samples, the material was processed and frozen within 1 h after the blood was taken. Plasma renin measurements were performed using the Renin III Generation radioimmunometric assay (CIS Bio International, Gif sur Yvette Cedex, France). The method’s detection range was 2.7–340 ng/l. When the estimated renin concentration was below the assay’s lower limit of detection, the renin concentration was considered the lowest detectable value, i.e. 2.7 ng/l. The observed intra-assay coefficients of variation (CV values) were 9.5 and 3.2% for renin concentrations of 7.6 and 347.3 ng/l respectively. The observed interassay CV values were 11.9 and 5.2% for renin concentrations of 24.6 and 281 ng/l respectively.

Serum aldosterone concentrations were measured using a commercially available RIA (Coat-A-Count aldosterone; Siemens/DPC, Los Angeles, CA, USA). The assay’s range was 30–3880 pmol/l. When the estimated aldosterone concentrations were below the lower detection limit, the samples were considered to have the lowest detectable serum aldosterone concentration, i.e. 30 pmol/l. The observed intra-assay CV values were 10.2 and 4.31% for aldosterone concentrations of 160 and 550 pmol/l respectively. The observed interassay CV values were 10.3 and 7.6% for aldosterone concentrations of 155 and 1560 pmol/l respectively.

To calculate ARRs, both aldosterone and renin concentrations were expressed in nanograms per liter. To convert aldosterone concentration to mass units, a coefficient of 0.361 ng/pmol was used.

Statistical analysis

MedCalc Software (version 11.6) was used for the statistical analysis of the acquired data.

<table>
<thead>
<tr>
<th>Table 1 Patients’ principle characteristics. The table shows the data of patients who were enrolled in the analysis. Unless otherwise stated, the data are expressed as medians (ranges). Group A patients underwent two identical confirmatory tests under RAAS-neutral therapy, only doxazosin and/or verapamil were allowed. Group B patients underwent two confirmatory tests: the first during modified chronic antihypertensive therapy and the second during RAAS-neutral therapy, only doxazosin and/or verapamil were allowed.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
<tr>
<td>No. of patients (women)</td>
</tr>
<tr>
<td>Age (years)</td>
</tr>
<tr>
<td>Systolic BP (mmHg)a</td>
</tr>
<tr>
<td>Diastolic BP (mmHg)a</td>
</tr>
<tr>
<td>No. of antihypertensive drugsa</td>
</tr>
<tr>
<td>Aldosterone-to-renin ratioa,b</td>
</tr>
</tbody>
</table>

aData obtained during the screening for primary aldosteronism.

bTo calculate the aldosterone-to-renin ratio, both aldosterone and renin were expressed in nanograms per liter.
The study consecutively enrolled 114 patients with suspected PA. The patients were divided into two groups: Groups A and B. All of the patients were examined according to the study protocol (Fig. 1).

**Group A**

Group A included 57 patients, all of whom underwent two identical confirmatory tests under RAAS-neutral therapy conditions, only doxazosin and/or verapamil were allowed. Group B patients underwent two confirmatory tests: the first with modified chronic antihypertensive therapy and the second with RAAS-neutral therapy, only doxazosin and/or verapamil were allowed.

**Group B**

Fifty-seven patients with suspected PA were enrolled in Group B. All were examined according to the study protocol. Four individuals were excluded because of high plasma renin at the end of the second confirmatory test performed during RAAS-neutral therapy. The data of 53 patients were included in the analysis. The principal characteristics of these patients are shown in Table 1. The antihypertensive medications given before both tests are summarized in Tables 2 and 6.

Figure 2A compares the serum aldosterone concentrations at the end of both confirmatory tests performed in Group A patients. The difference in aldosterone concentrations ranged from 0 to 350 pmol/l (median: 50 pmol/l).

Detailed results of both confirmatory tests in Group A patients are displayed in Tables 3 and 4. Overall, agreement in the interpretation of the two tests was noted in 43 (84%) patients. The data from the eight patients who exhibited discordant results are shown in more detail in Table 5.

All 20 patients who had serum aldosterone concentrations ≥ 240 pmol/l at the end of either confirmatory test exhibited aldosterone nonsuppressibility at the end of the other test.

AVS was performed in 31 patients in Group A. Aldosterone secretion lateralization was observed in 15 individuals. All patients with proven aldosterone secretion lateralization had consistently nonsuppressible aldosterone at the end of both tests. None of the five patients who had discordant confirmatory test results and underwent AVS exhibited evidence of aldosterone secretion lateralization.

**Figure 2** Comparison of serum aldosterone at the end of the confirmatory tests. This graph compares the end-test serum aldosterone (open circle) in Groups A (A) and B (B). Passing–Bablok regression analysis was used to analyze the differences between the compared confirmatory tests. In both study groups, wide 95% confidence intervals (CI) describing the regression line indicate only moderate agreement in the main results of the compared confirmatory tests. Bold lines indicate regression lines; grey lines indicate 95% CI; and dotted lines indicate identity lines.
The detailed data from both tests administered to Group B patients are summarized in Tables 3 and 7. Consonant interpretations of confirmatory tests were noted in 35 (66%) patients, and discordant results were found in 18 (34%) patients.

We have not identified any specific type of chronic antihypertensive medication associated more frequently with discordant results in the compared confirmatory tests.

Concerning amiloride administration, discordant results were observed in four (20%) of 20 patients who were given this drug as part of the modified chronic therapy provided during the first test.

All 16 individuals who had serum aldosterone levels ≥240 pmol/l on modified chronic therapy end of testing exhibited consonant results when the test was repeated under RAAS-neutral therapy.

On the basis of the results of the second confirmatory test, 39 individuals underwent AVS. Aldosterone secretion lateralization was observed in 14 patients, three of these patients exhibited aldosterone suppression during the confirmatory test performed under modified chronic antihypertensive therapy conditions.

Table 3 Confirmatory testing results. The data are expressed as medians and ranges. Group A patients underwent two identical confirmatory tests under RAAS-neutral therapy conditions. Group B patients underwent two confirmatory tests: the first with modified chronic antihypertensive therapy and the second with RAAS-neutral therapy.

<table>
<thead>
<tr>
<th></th>
<th>Group A (51 patients)</th>
<th>Group B (53 patients)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Test 1&lt;sup&gt;a,b&lt;/sup&gt;</td>
<td>Test 2&lt;sup&gt;a,b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Test 1&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Test 2&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Aldosterone-to-renin ratio&lt;sup&gt;d,e&lt;/sup&gt;</strong></td>
<td>45.7 (0.7–204.5)</td>
<td>47.0 (1.4–160.4)</td>
</tr>
<tr>
<td><strong>Serum potassium level (mmol/l)&lt;sup&gt;f&lt;/sup&gt;</strong></td>
<td>3.7 (2.7–4.6)</td>
<td>3.7 (2.9–4.7)</td>
</tr>
<tr>
<td><strong>Urinary sodium excretion (mmol/24 h)&lt;sup&gt;f&lt;/sup&gt;</strong></td>
<td>368 (124–814)</td>
<td>358 (142–721)</td>
</tr>
<tr>
<td><strong>Diastolic blood pressure (mmHg)&lt;sup&gt;g&lt;/sup&gt;</strong></td>
<td>92 (70–118)</td>
<td>94 (70–122)</td>
</tr>
<tr>
<td><strong>Systolic blood pressure (mmHg)&lt;sup&gt;g&lt;/sup&gt;</strong></td>
<td>150 (120–192)</td>
<td>158 (116–220)</td>
</tr>
<tr>
<td><strong>Serum aldosterone at the end of saline infusion (pmol/l)</strong></td>
<td>140 (30–2400)</td>
<td>160 (118–220)</td>
</tr>
<tr>
<td><strong>Plasma active renin at the end of saline infusion (ng/l)</strong></td>
<td>2.7 (2.7–6.9)</td>
<td>2.7 (2.7–5.4)</td>
</tr>
<tr>
<td><strong>Diastolic blood pressure (mmHg)&lt;sup&gt;g&lt;/sup&gt;</strong></td>
<td>92 (70–118)</td>
<td>94 (70–122)</td>
</tr>
</tbody>
</table>

<sup>a</sup>Confirmatory tests performed under RAAS-neutral therapy, only doxazosin and/or verapamil were allowed.
<sup>b</sup>Confirmatory tests performed with identical dosing of doxazosin and/or verapamil.
<sup>c</sup>Confirmatory tests performed under modified chronic antihypertensive therapy conditions, aldosterone antagonists and diuretics were withdrawn before testing.
<sup>d</sup>Blood sampling was performed after oral sodium loading before the saline infusion was given.
<sup>e</sup>To calculate the aldosterone-to-renin ratio, both aldosterone and renin were expressed in nanograms per liter.
<sup>f</sup>A 24-h urine sample was collected the day before the saline infusion was administered.
<sup>g</sup>Blood pressure measurement after saline infusion.

**Hypokalemia**

Serum potassium was measured after oral sodium loading before the saline infusion was given. Hypokalemia, defined as a potassium level <3.5 mmol/l, was observed in 24 (23%) patients. All but one of these patients received potassium supplementation, with a mean dose of 46 (±18) mmol/day. No patient with hypokalemia achieved aldosterone suppression at the end of saline infusion.

**Side effects**

We did not encounter any major complications that required acute hospital admission in any of the enrolled patients.

Peripheral edema was observed in seven patients and was more commonly associated with therapy limited to verapamil and/or doxazosin. An episode of atrial fibrillation developed in one patient on doxazosin and verapamil; hypokalemia was not detected. Dyspnea of effort was reported by three patients, and two patients reported headaches; neither complaint was related to any specific therapy.

**Discussion**

Our study assessed the clinical value of confirmatory testing performed under modified chronic antihypertensive therapy with the exclusion of thiazides, loop diuretics, and aldosterone antagonists. To interpret the results, the study protocol also evaluated the reproducibility of confirmatory testing performed under RAAS-neutral antihypertensive therapy conditions.
The reproducibility of the confirmatory testing performed with RAAS-neutral antihypertensive therapy

Consonant results for two identical confirmatory tests were observed in a majority of examined patients; however, the number of patients with discordant results was not negligible. The only definite predictor of concordantly positive results was the marked elevation of serum aldosterone at the end of saline infusion.

We cannot provide any definite explanation for the observed discrepancies. Some of the discordant findings may result from imperfect renin suppression (in patients 2 and 7, Table 5) or insufficient oral sodium loading (patient 4, Table 5).

We suppose that autonomous aldosterone secretion may be excluded in all patients who ever had serum aldosterone suppression levels < 60 pmol/l, i.e. below the 97.5 percentile of values obtained from healthy volunteers who underwent identical confirmatory testing (4).

The discrepant interpretations of the compared tests cannot be attributed solely to the inherent imprecision of laboratory methods. As is apparent from the data presented in Table 5, the expected inaccuracies of hormonal estimations were usually below the observed differences in aldosterone concentrations.

We do not assume that previous chronic medical therapy influenced the results. In all patients whose data are presented in Table 5, the periods during which they were not taking chronic medication were sufficient for the complete elimination of previously administered antihypertensive medications.

We did not encounter hypokalemia ‘masking’ an autonomous aldosterone secretion; the instances of hypokalemia in our study resulted from excessive aldosterone overproduction.

The effect of chronic antihypertensive medication on the results of confirmatory testing

In one-third of Group B patients, there were discrepancies in the interpretation of the two confirmatory tests. Modified chronic medication was associated with aldosterone suppressibility, even in a few of the patients who were eventually found to have aldosterone secretion lateralization.

Conversely, the highest end of the serum aldosterone test with modified chronic therapy uniformly predicted aldosterone nonsuppressibility when the test was repeated under RAAS-neutral therapy. It appears that severely excessive aldosterone overproduction may not be effectively suppressed by modified chronic antihypertensive medication.

It is also important to note that amiloride administration did not influence the confirmatory test results in the majority of patients who received this drug as a part of their treatment.

Table 5 The analysis of data from Group A patients whose confirmatory test results were discordant. Both tests were performed under the same RAAS-neutral therapy conditions.

<table>
<thead>
<tr>
<th>n</th>
<th>Days off chronic therapy (mmol/day)</th>
<th>Urinary sodium (mmol/day)</th>
<th>Serum potassium (mmol/l)</th>
<th>Plasma renin (ng/l)</th>
<th>Serum aldosterone (pmol/l)</th>
<th>AVS/lateralization</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>43</td>
<td>506</td>
<td>3.7</td>
<td>2.7</td>
<td>30</td>
<td>503</td>
</tr>
<tr>
<td>2</td>
<td>18</td>
<td>457</td>
<td>3.8</td>
<td>5.9</td>
<td>210</td>
<td>636</td>
</tr>
<tr>
<td>3</td>
<td>22</td>
<td>161</td>
<td>3.7</td>
<td>3</td>
<td>190</td>
<td>142</td>
</tr>
<tr>
<td>4</td>
<td>15</td>
<td>376</td>
<td>3.9</td>
<td>2.7</td>
<td>40</td>
<td>153</td>
</tr>
<tr>
<td>5</td>
<td>48</td>
<td>814</td>
<td>4.5</td>
<td>2.7</td>
<td>80</td>
<td>436</td>
</tr>
<tr>
<td>6</td>
<td>19</td>
<td>359</td>
<td>4.2</td>
<td>2.7</td>
<td>60</td>
<td>300</td>
</tr>
<tr>
<td>7</td>
<td>31</td>
<td>586</td>
<td>4.2</td>
<td>4.6</td>
<td>140</td>
<td>502</td>
</tr>
<tr>
<td>8</td>
<td>132</td>
<td>651</td>
<td>4.5</td>
<td>2.7</td>
<td>90</td>
<td>719</td>
</tr>
</tbody>
</table>

* n, number of patients; -, indicates not applicable.

aDays of days that chronic antihypertensive therapy with a potential influence on RAAS had been stopped before the first confirmatory test was performed.

bA 24-h urine sample was collected after oral sodium loading the day before the saline infusion was given.

cBlood sampling was performed after 3 days of oral sodium loading before the saline infusion was given.

dBlood samples were taken at the end of saline infusion. AVS denotes successful adrenal venous sampling. Lateralization denotes the lateralization of aldosterone secretion from one adrenal gland.

Table 6 The modified chronic antihypertensive therapy administered to Group B patients during the first confirmatory test. The data are expressed as numbers of patients (percentages).

<table>
<thead>
<tr>
<th>Beta blockers</th>
<th>45 (85%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Angiotensin-converting enzyme inhibitors</td>
<td>35 (66%)</td>
</tr>
<tr>
<td>Angiotensin receptor blockers</td>
<td>8 (15%)</td>
</tr>
<tr>
<td>Calcium channel blockers</td>
<td>41 (77%)</td>
</tr>
<tr>
<td>Alpha blockers</td>
<td>31 (58%)</td>
</tr>
<tr>
<td>Centrally acting drugs</td>
<td>8 (15%)</td>
</tr>
<tr>
<td>Amiloride</td>
<td>20 (38%)</td>
</tr>
</tbody>
</table>

* The mean dose of amiloride was 9 ± 3.8 mg.
of their modified chronic therapy. This observation may be important for patients with severe hypokalemia, for whom there are concerns about the safety of performing confirmatory testing according to the therapy recommended by the guidelines.

**Study limitations**

The main limitation of our study is the unclear definition of aldosterone suppressibility. There is no gold standard for reliably confirming autonomous aldosterone production. Our study, as well as other reports (6, 7, 8, 9, 10), used an arbitrary criterion to define aldosterone suppressibility. Despite the fact that our cutoff for defining PA was rather low, we cannot be sure that it was sufficient to identify all individuals with autonomous aldosterone secretion.

We are aware that the addition of one more confirmatory test to the study protocol would further improve our understanding of the clinical question studied. However, the extension of the study procedures was considered highly impractical from the patients’ point of view.

Finally, our observations cannot be automatically extrapolated as the reported data are closely related to the protocol and the hormonal assays used. Significant differences among kits from different manufacturers exist (11).

**Conclusion**

Our data suggest that an extensive medication switch is not necessary for all patients who are referred for confirmatory testing for suspected PA. Chronic therapy with the exclusion of aldosterone antagonists, thiazides, and loop diuretics may offer an alternative to RAAS-neutral therapy in patients with severe, complicated hypertension and/or refractory hypokalemia.

The results of confirmatory testing performed with modified chronic therapy are valid only when the serum aldosterone is markedly elevated at the end of saline infusion. In the remaining cases, tests conducted under modified chronic therapy conditions must be considered inconclusive.

Another important finding of our study is the limited reproducibility of confirmatory testing under RAAS-neutral therapy conditions. The confirmatory test is worth repeating unless the serum aldosterone concentration is markedly high or markedly low at the end of the test. The aim of repeated testing is to find patients who are able to suppress serum aldosterone to very low levels, which makes a PA diagnosis improbable. The need for AVS can be decreased by this approach.

**Declaration of interest**

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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**References**


**Table 7** The interpretation of confirmatory tests in Group B patients. The table compares the interpretation of two confirmatory tests in Group B patients. The data are expressed as the numbers of patients.

<table>
<thead>
<tr>
<th>Serum aldosterone at the end of saline infusion</th>
<th>Test during modified chronic antihypertensive therapy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Suppressible (≤ 100 pmol/l)</td>
<td>Nonsuppressible (&gt; 100 pmol/l)</td>
</tr>
<tr>
<td><strong>Test during RAAS-neutral therapy</strong></td>
<td></td>
</tr>
<tr>
<td>Suppressible</td>
<td>5</td>
</tr>
<tr>
<td>Nonsuppressible</td>
<td>7</td>
</tr>
</tbody>
</table>


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