Primary aldosteronism in normokalemic patients with adrenal incidentalomas

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Abstract

Objective: Since primary aldosteronism has been reported in asymptomatic incidental adrenal masses (adrenal incidentalomas, AI), the aim of our study was to detect primary aldosteronism in normokalemic patients with AI and to verify whether a raised plasma aldosterone (ALD)/plasma renin activity (PRA) ratio may be useful for diagnosis.

Design: One-hundred and twenty-five normokalemic patients with solid AI (90 hypertensives and 35 normotensives) and 82 essential hypertensives (EH) were studied. Upright ALD and PRA determination was performed in all cases while patients with abnormal ALD/PRA ratios were submitted to confirmatory tests (saline infusion and captopril tests) for diagnosis of primary aldosteronism.

Methods: ALD and PRA were measured by specific radioimmunoassays.

Results: PRA values in AI hypertensives were lower than in AI normotensives and in EH; P < 0.05 and in EH; P < 0.0001. The ALD/PRA ratio in AI hypertensives was higher than in AI normotensives and in EH; P < 0.03. Four patients with EH and 2 AI normotensive patients had elevated ALD/PRA ratios but normal responses to the suppressive tests, thus excluding diagnosis of primary aldosteronism. Eight patients with AI and hypertension had a high ALD/PRA ratio, and 7 of these were further studied: in 5 patients diagnosis of primary aldosteronism was well-established by dynamic tests, adrenal vein sampling or by surgery.

Conclusions: Primary aldosteronism in normokalemic patients with incidentally discovered adrenal masses was detected in 4 of all cases and in at least 5.5% of those with hypertension. Consequently, these patients, particularly if hypertensive, need to be routinely studied to exclude this hormonal disease. Evaluation of the ALD/PRA ratio seems to be a simple and reliable test for diagnosis.

Introduction

The incidental discovery of adrenal masses (incidentalomas) is becoming a common clinical problem for the physician, above all when detection occurs in hypertensive patients in whom a causal relationship between adrenal mass and hypertension may exist. Interestingly, the prevalence of hypertension in patients with adrenal incidentalomas (AI) is higher than in the general population. This finding has a possible explanation because the apparently nonfunctioning tumors may secrete catecholamines, cortisol and aldosterone (2, 4–9) causing diseases (pheochromocytoma, subclinical Cushing’s syndrome and primary aldosteronism) responsible for secondary hypertension.

Primary aldosteronism (PA) is an uncommon cause of endocrine hypertension. The prevalence of PA was previously thought to be less than 1%, above all in unselected hypertensive population (10–13). However, mainly as a result of improved diagnostic techniques, the prevalence of PA is now estimated to be more than 5% both in specialist referral centers (14–18) and also in a primary care setting (19, 20). PA in the classical form presents with aldosterone (ALD) excess, low plasma renin activity (PRA) and hypokalemia. However, several reports indicate that normokalemic PA constitutes the most common presentation of the disease, while the hypokalemic variant probably occurs only in the most severe cases (14, 15, 18, 20–25). For these reasons, detection of PA in patients with adrenal incidentalomas, who according to the criteria of the Italian Study Group on Adrenal Tumors are normokalemic (1, 2, 4), is not surprising.

It is well-known that in screening patients for primary aldosteronism, PRA determination is not specific, although very sensitive, because of the influence of...
medication and the high prevalence of low renin essential hypertension (26). On the other hand, measurement of plasma and urinary ALD alone may likewise be insufficient to diagnose hypokalemic patients with PA, because ALD values are sometimes in the normal range (25, 27, 28). To date, the best screening test for distinguishing patients with essential hypertension from those with possible PA is the ratio of plasma ALD level to PRA (12, 15, 28–31).

The aim of the present study was to detect primary aldosteronism in normokalemic patients with solid AI and to verify if a raised ALD/PRA ratio may be useful for diagnosis.

Materials and methods
Subjects
Among patients referred to our Department for radiological adrenal abnormalities, we studied only those in whom the discovery was incidental. In addition, patients found to have cystic adrenal lesions, hypokalemia (present or past), high catecholamine levels, abnormal pituitary–adrenal axis function or excess of sexual hormones and androgen precursors were not included in the study. We performed this selection because both in cystic tumors and in masses secreting catecholamines, cortisol or androgens, detection of PA is obviously unlikely. Thus, we consequently evaluated 125 normokalemic patients with solid AI (90 hypertensives and 35 normotensives). Detection of the mass, in the absence of signs or symptoms of adrenal dysfunction, had been obtained after ultrasonography performed for hepato-biliary (n = 58) and genito-urinary (n = 47) disorders or for aspecific abdominal (n = 20) pain. Computed tomography (CT) confirmed a solitary or clearly delineated nodule with a normal-appearing contralateral gland in 93 cases (right side n = 89, left side n = 34); maximum diameter 26.8±1.04 mm (mean±s.e.), range 10–80 mm). CT showed bilateral lesions in 32 cases: bilateral nodules (n = 8), enlargement (>15 mm) of both adrenals (n = 12) and monolateral enlargement associated with contralateral mass (n = 12).

Eighty-two essential hypertensives (EH) were also studied, chosen among patients referred to our Department. Diagnosis had been established after exclusion, by appropriate biochemical and morphological investigations, of all detectable causes of hypertension. Hypokalemia had never been documented in these patients and in no case had abdominal ultrasonography (n = 75) or CT (n = 7) shown adrenal masses. The patients had been followed-up for at least 1 year and their blood pressure was well controlled with conventional antihypertensive treatment.

Demographic and clinical data of our 207 patients are given in Table 1.

Experimental design
Patients taking drugs (anti-hypertensive or other drugs) suspended medication for at least 2 weeks prior to the study. All patients maintained their usual diet without salt restriction (31). Patients presented between 0800 and 0900 h in the morning, after overnight fasting. Twenty-four-h urine was collected for sodium, potassium and aldosterone determination. Blood samples for sodium, potassium, ALD and PRA measurements were obtained in patients who had been standing for at least 1 h (32).

The ALD/PRA ratio cut-off in our laboratory is 112. This value represents the 95% upper confidence limit of the 90th percentile of normal values obtained from patients with EH (mean±s.d. 33.2±3.5, median 19.72, range 3.72 –166.5). Therefore, patients with AI and an ALD/PRA ratio over 112 underwent confirmatory tests in order to demonstrate unsuppressibility of plasma ALD levels (33, 34). In this case, prior to testing the patients underwent controlled salt intake (80–100 mmol/day sodium and 60–80 mmol/day potassium), in addition to pharmacological wash-out. Subsequently, saline infusion (plasma ALD determination before and after 2 liters of normal saline i.v. over a 4-h period) and the captopril test (plasma ALD measurements before and 2 h after captopril 50 mg

Table 1 Demographic data of essential hypertensives (EH) and of patients with adrenal incidentaloma (AI) both hypertensive and normotensive. Mean values±s.e. and range (in parentheses) are reported.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Adrenal incidentalomas</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>EH (n = 82)</td>
</tr>
<tr>
<td>Sex (M/F)</td>
<td>48/34</td>
</tr>
<tr>
<td>Age (years)</td>
<td>47.0±1.4 (16–68)</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>28.5±0.5</td>
</tr>
<tr>
<td>SBP (mmHg)</td>
<td>153.8±1.7</td>
</tr>
<tr>
<td>DBP (mmHg)</td>
<td>97.6±0.9</td>
</tr>
</tbody>
</table>

BMI, body mass index; SBP, systolic blood pressure; DBP, diastolic blood pressure.
*P < 0.0004 and **P < 0.0001 vs EH; †P < 0.007 vs EH; ‡P < 0.0001 vs EH and AI hypertensives.
orally) were performed. ALD values above 15 ng/dl after captopril administration were considered a typical feature of PA (35, 36). As far as salt loading is concerned, it is well known that normal subjects and essential hypertensives suppress plasma ALD below 5 ng/dl (18, 33, 37, 38), while patients with PA fail to suppress plasma ALD below 10 ng/dl (20, 39, 40). Patients in whom plasma ALD ranges between 5 and 10 ng/dl after saline infusion may have PA (41–43). For this reason the patients in the present study who showed a highly suspicious ALD post-loading response were further studied by adrenal vein sampling (23, 44, 45).

The study was approved by the Local Ethical Committee and the patients gave their informed consent.

**Assays**

Plasma and urinary sodium and potassium were evaluated by photometry assay. Plasma samples for PRA and ALD determinations were collected in chilled anticoagulated glass tubes, immediately centrifuged and frozen at −20°C. Hormones were then assayed in duplicate and in the same run by specific RIAs. Intra-assay and interassay coefficients of variation were as follows: ALD (DiaSorin, Saluggia, Italy) 9.7% and 11.5%, PRA (DiaSorin) 7.6% and 9.1% respectively. Detection limits of the methods were below 2 ng/dl for ALD and below 0.2 ng/ml/h for PRA. Cortisol (Immunotech International, Marseille, France), expressed in ng/ml, had intra-assay and interassay coefficients of variation of 5.7% and 6.6% respectively, and the sensitivity was 13 nmol/l.

Blood pressure was recorded by mercury sphygmomanometer following recent guidelines (46).

**Statistical analysis**

PRA values and the ratio of plasma ALD to PRA were calculated with renin activity taken to be 0.2 ng/ml/h in those patients where this value was below the sensitivity limit of our assay. Variables are expressed as mean, standard error and range. Kruskal–Wallis and Wilcoxon tests were performed for each variable to compare groups. The unpaired t-test was also used, when appropriate. Finally, Spearman correlation coefficients were adopted as parameters of associations. *P* levels lower than 0.05 were considered statistically significant.

**Results**

Patients with adrenal incidentalomas, both normotensive and hypertensive, were older and leaner than EH. Systolic and diastolic blood pressure in AI normotensives was, by definition, lower than in AI hypertensives and in EH (Table 1).

As shown in Table 2, while plasma potassium did not significantly differ, patients with AI showed PRA values lower than EH. This finding was more evident in the AI hypertensive subgroup which, consequently, showed the highest ALD/PRA ratio.

In no group was a correlation found between plasma ALD/PRA ratio and age, body mass index, blood pressure, plasma potassium or tumor size (in the patients with AI).

Analytical data (Fig. 1) indicated that 4 patients with EH and 2 patients with AI and normal blood pressure had an ALD/PRA ratio that was borderline or over 112. When submitted to saline loading and the captopril test these patients (with urinary and upright plasma ALD lower than 30 μg/24 h and 30 ng/dl respectively) had normally suppressed plasma ALD levels. Thus they were false positives.

Eight patients with AI and hypertension (8.8%) were found to have an ALD/PRA ratio above the normal value. As expected, they had significantly lower PRA values and a significantly higher plasma ALD and ALD/PRA ratio than the remaining group (n = 82), without difference in age, blood pressure, body mass index, plasma potassium or mass size (Table 3). Among these eight patients with abnormal ALD/PRA ratios (Table 4), six were further investigated by the salt loading and captopril tests: two patients showed normal responses whereas ALD unsuppressibility or partial suppressibility was found in 4 cases. One patient (no. 3) with a clear monolateral nodule was surgically treated and cured. Three patients (nos 1, 2 and 4) with bilateral adrenal enlargement underwent successful (at

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Plasma humoral data of essential hypertensives (EH) and of patients with adrenal incidentaloma (AI) both hypertensive and normotensive. Mean values±S.E. are reported.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Adrenal incidentalomas</strong></td>
<td><strong>EH (n = 82)</strong></td>
</tr>
<tr>
<td>Plasma K (mEq/l)</td>
<td>3.97±0.02</td>
</tr>
<tr>
<td>ALD (ng/dl)</td>
<td>31.1±1.7</td>
</tr>
<tr>
<td>PRA (ng/ml/h)</td>
<td>1.68±0.15</td>
</tr>
<tr>
<td>ALD/PRA ratio</td>
<td>33.2±3.5</td>
</tr>
</tbody>
</table>

*P < 0.0005 vs EH; †P < 0.04 and ††P < 0.0001 vs EH; ‡P < 0.05 vs AI hypertensives; §P < 0.03 vs AI normotensives.
least 3-fold plasma cortisol elevation in the adrenal vein in comparison with the inferior vena cava) adrenal vein sampling (23, 44, 45) which revealed an ALD/cortisol index that was similar in both sides (23) and greater than in the inferior vena cava (Table 5). This procedure allowed diagnosis and etiology of PA, i.e. bilateral adrenal hyperplasia. The seventh patient, without testing, underwent surgery for radiological features of the mass suspected of malignancy: one year later blood pressure, plasma ALD and ALD/PRA ratios were maintained in the normal range. The eighth patient dropped out. Thus, in 5 out of 7 patients studied, the diagnosis of PA was well-established. Overall, primary aldosteronism was found in 4.0% of all AI and in 5.5% of those with hypertension. These results are shown in diagrammatic form in Fig. 2.

**Discussion**

Our results show that patients in whom an adrenal mass has been incidentally detected, above all if they are hypertensive, need to be screened for primary aldosteronism even in the presence of normal potassium levels. Thus, 8.8% of our patients with AI and hypertension showed an abnormal ALD/PRA ratio and at least 5.5% had proven hyperaldosteronism. Although apparently similar to that found in the general population, the diagnosis of PA was well-established in 5 out of 7 patients studied. Overall, primary aldosteronism was found in 4.0% of all AI and in 5.5% of those with hypertension. These results are shown in diagrammatic form in Fig. 2.

**Table 3** Demographic, humoral and clinical data of adrenal incidentaloma hypertensives with plasma ALD/PRA ratio lower ($n = 82$) and higher ($n = 8$) than 112. Mean values±s.e. are reported.

<table>
<thead>
<tr>
<th>ALD/PRA ratio</th>
<th>$&lt;112$</th>
<th>$&gt;112$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex (M/F)</td>
<td>28/54</td>
<td>2/6</td>
</tr>
<tr>
<td>Age (years)</td>
<td>59.9±1.1</td>
<td>57.1±2.8</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>26.1±0.3</td>
<td>27.8±1.2</td>
</tr>
<tr>
<td>SBP (mmHg)</td>
<td>159.0±1.4</td>
<td>155.7±4.3</td>
</tr>
<tr>
<td>DBP (mmHg)</td>
<td>97.4±0.9</td>
<td>95.7±3.4</td>
</tr>
<tr>
<td>Mass side: M/B</td>
<td>58/24</td>
<td>5/3</td>
</tr>
<tr>
<td>Mass size (mm)$^*$</td>
<td>26.8±1.3</td>
<td>29.5±7.4</td>
</tr>
<tr>
<td>Plasma K (mEq/l)</td>
<td>4.0±0.03</td>
<td>3.99±0.1</td>
</tr>
<tr>
<td>ALD (ng/dl)</td>
<td>21.5±1.4</td>
<td>38.2±4.1**</td>
</tr>
<tr>
<td>PRA (ng/ml/h)</td>
<td>1.13±0.14</td>
<td>0.21±0.01*</td>
</tr>
</tbody>
</table>

$^*$P<0.05; **P<0.0004

BMI, body mass index; SBP/DBP, systolic/diastolic blood pressure; $^*$Maximum diameter; M, monolateral; B, bilateral (bilateral nodules or bilateral enlargement of adrenals or increased thickness of one adrenal associated with a mass in the contralateral gland).

**Figure 1** Individual data of the plasma ALD/PRA ratio in essential hypertensives (EH) and in patients with adrenal incidentaloma (AI) both hypertensive and normotensive. The dashed line indicates the upper 95% limit of normal values (112).

**Figure 2** True positive and false positive results found in patients with adrenal incidentaloma.

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hypertensive population (29, 32, 47), this percentage is in effect higher since patients with hypokalemia are usually included among unselected hypertensives, while the present data were obtained in normokalemic subjects only.

Detection of primary aldosteronism in patients with AI has already been reported, with values ranging between 1.6% (2) and 3.0% (9). The number we found is even greater, probably because we selected only solid tumors after excluding glucocorticoid, sex steroid or catecholamine excess, while in the above reported series all AI (normo- and hyper-functioning, solid and cystic tumors) were studied.

As is known, the clinical and humoral features of PA are not always pathognomonic of the disease: plasma and urinary ALD may be in the normal range (25, 27, 28) and low PRA values are detected in patients with essential hypertension (26). In addition, serum potassium is frequently normal (14, 15, 18, 20–25) and even documented cases of normotensive patients with PA have been reported (48–50). On the basis of these findings, use of the plasma ALD/PRA ratio is believed, to date, to be the best screening test for primary aldosteronism (12, 15, 18, 28–31). This test is very accurate, simple, does not require the inconvenience of urine collection and does not seem to be influenced by variations in sodium intake, total body potassium deficit or by the drugs used to treat hypertension, with the exception of spironolactone (29, 31). Therefore, ALD/PRA ratio determination may be useful in the work-up of AI since the number of unexpectedly detected adrenal masses is constantly increasing, due to the growing spread of radiological procedures (51–54). Management of AI now includes not only a morphological study in order to rule out malignant tumors (52–54), but also hormonal screening because of the possible occurrence of abnormalities in glucocorticoid, catecholamine, sex steroid or mineralocorticoid secretion (2, 4–9). This hormonal evaluation is particularly indicated in hypertensive patients in whom high blood pressure values may be related to endocrine dysfunction of the adrenal glands. It is interesting to note that the prevalence of hypertension in patients with AI is higher than in the general population (1–3). This finding may suggest a possible pathogenetic link between hypertension and adrenal mass. However, in order to avoid the ‘cascade-effect’ in the clinical care of patients with AI, several restrictive diagnostic work-ups have been proposed to identify hormonal diseases. As far as primary aldosteronism is concerned, many authors claim that since the prevalence of the disease is low it may be counterproductive to extensively screen all patients with AI (even if they are hypertensive), while it is preferable to perform specific evaluations only in the presence of hypokalemia (52, 54–56). Our results show that high blood pressure values, more than plasma potassium levels, are crucial in alerting physicians to the suspicion of primary aldosteronism.

Table 4 Clinical data of adrenal incidentaloma hypertensives with probable primary aldosteronism (ALD/PRA ratio >112).

<table>
<thead>
<tr>
<th>Patient</th>
<th>BP (mmHg)</th>
<th>Plasma K (mEq/l)</th>
<th>Urinary Na (mEq/24 h)</th>
<th>Plasma ALD (ng/dl)</th>
<th>Urinary ALD (μg/24 h)</th>
<th>PRA (ng/ml/h)</th>
<th>ALD/PRA ratio</th>
<th>Mass size (mm)</th>
<th>Mass side (L/R/B)</th>
<th>Salt loading: ALD nadir vs basal (ng/dl)</th>
<th>Captopril test: ALD nadir vs basal (ng/dl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>160/90</td>
<td>4.0</td>
<td>119</td>
<td>30.0</td>
<td>3.2</td>
<td>0.2</td>
<td>150</td>
<td>24</td>
<td>B</td>
<td>8 vs 21</td>
<td>20 vs 31</td>
</tr>
<tr>
<td>2</td>
<td>152/82</td>
<td>4.0</td>
<td>119</td>
<td>26.8</td>
<td>3.6</td>
<td>0.2</td>
<td>154</td>
<td>25</td>
<td>B</td>
<td>8 vs 26</td>
<td>27 vs 32</td>
</tr>
<tr>
<td>3</td>
<td>176/108</td>
<td>4.1</td>
<td>24</td>
<td>33.6</td>
<td>35</td>
<td>0.2</td>
<td>176</td>
<td>24</td>
<td>L</td>
<td>16 vs 25</td>
<td>22 vs 29</td>
</tr>
<tr>
<td>4</td>
<td>160/108</td>
<td>3.9</td>
<td>196</td>
<td>41.7</td>
<td>31.7</td>
<td>0.2</td>
<td>196</td>
<td>10</td>
<td>B</td>
<td>11 vs 31</td>
<td>16 vs 30</td>
</tr>
<tr>
<td>5</td>
<td>140/90</td>
<td>4.7</td>
<td>104</td>
<td>39.2</td>
<td>35.8</td>
<td>0.2</td>
<td>198</td>
<td>28</td>
<td>R</td>
<td>4 vs 27</td>
<td>13 vs 26</td>
</tr>
<tr>
<td>6</td>
<td>160/104</td>
<td>3.8</td>
<td>86</td>
<td>39.7</td>
<td>29</td>
<td>0.2</td>
<td>198</td>
<td>20</td>
<td>R</td>
<td>1 vs 24</td>
<td>6 vs 27</td>
</tr>
<tr>
<td>7</td>
<td>138/94</td>
<td>3.7</td>
<td>70</td>
<td>63.9</td>
<td>29</td>
<td>0.2</td>
<td>194</td>
<td>25</td>
<td>L</td>
<td>nd</td>
<td>nd</td>
</tr>
<tr>
<td>8</td>
<td>160/90</td>
<td>3.8</td>
<td>90</td>
<td>52.8</td>
<td>52.3</td>
<td>0.2</td>
<td>155</td>
<td>80</td>
<td>nd</td>
<td>nd</td>
<td>nd</td>
</tr>
</tbody>
</table>

BP, blood pressure; L, left; R, right; B, bilateral; nd, not done.

*The patient underwent surgery. One year after the removal of adrenal mass, clinical and humoral data were as follows: plasma ALD 5.4 ng/dl, PRA 0.48 ng/ml/h, ALD/PRA ratio 11.2, urinary ALD 11.3 μg/24 h, urinary Na 7.5 mEq/24 h, plasma K 4.0 mEq/l, blood pressure 118/74 mmHg.

Table 5 Results of bilateral adrenal vein sampling in the three patients with bilateral adrenal enlargement, ALD/PRA ratio over 112 and partial ALD post loading response. Results are expressed as ALD in ng/dl/cortisol in ng/ml.

<table>
<thead>
<tr>
<th>Patient</th>
<th>Right side ALD/cortisol</th>
<th>Lift side ALD/cortisol</th>
<th>Peripheral* ALD/cortisol</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>380/680</td>
<td>290/587</td>
<td>30/130</td>
</tr>
<tr>
<td>2</td>
<td>193/391</td>
<td>230/724</td>
<td>26/92</td>
</tr>
<tr>
<td>4</td>
<td>272/1060</td>
<td>236/980</td>
<td>41/196</td>
</tr>
</tbody>
</table>

*Infrarenal portion of inferior vena cava.
aldosteronism in patients with AI. Significantly, no patient with PA was found in AI normotensives. We therefore suggest that in the diagnostic work-up of patients with AI and normal blood pressure, evaluation of the renin–angiotensin–aldosterone system may be avoided.

We are aware that the selection of our patients did not allow us to establish the prevalence of primary aldosteronism in AI. However, the study was not undertaken for this specific purpose, but rather to assess the presence of this form of endocrine hypertension and, therefore, to point out that in patients with solid AI normal potassium levels do not exclude the diagnosis of primary aldosteronism. Therefore, only focusing on primary aldosteronism in hypokalemic adrenal incidentaloma patients overlooks a significant subset of primary aldosteronism patients who could be cured of their hypertension if it were detected by a cost-effective screening tool such as the plasma ALD/PRA ratio.

In the choice of our ALD/PRA ratio cut-off we took many factors into account. First, the lower limit of PRA was moved to 0.2 ng/ml/h (the sensitivity limit of our assay) in those patients where this value was below. Secondly, since low renin levels are frequently observed in EH (26), the cut-off was obtained from a large number of patients with well-ascertained essential hypertension instead of from normotensive controls, in order to avoid the high number of false positive results in EH (13). Finally, the evaluation was made in conditions similar to those of patients with AI (pharmacological wash-out, free salt intake, upright position before testing). In these conditions our cut-off proved to be higher than that reported by other centers (12, 28, 29, 31). This variability is not surprising, due both to different assay techniques and laboratory conditions and the influence of several variables on PRA and ALD levels. Consequently, the importance of locally validated criteria for establishing the normal range must be stressed.

In conclusion, the present data show that in normokalemic patients with incidentally discovered adrenal masses primary aldosteronism is detectable. Therefore, at least the hypertensive patients with AI need to be routinely studied to exclude this hormonal disease. Evaluation of the ALD/PRA ratio seems to be a simple and reliable test for diagnosis.

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References


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