ACTH after 15 min distinguishes between Cushing’s disease and ectopic Cushing’s syndrome: a proposal for a short and simple CRH test

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Abstract

Objective: The aim of the present study was to validate criteria of corticotropin-releasing hormone (CRH) stimulation and 8 mg dexamethasone suppression (high-dose dexamethasone suppression, HDDS) to distinguish the etiology of ACTH-dependent Cushing’s syndrome.

Subjects and methods: We retrospectively analyzed cortisol and ACTH after the injection of 100 μg human CRH in confirmed Cushing’s disease (CD, n=78) and confirmed ectopic Cushing’s syndrome (ECS, n=18). Cortisol and ACTH increase (in percentage above basal (%B)) at each time point, maximal increase (Δmax %B), and area under the curve (AUC %B) were analyzed using receiver operator characteristics (ROC) curve analyses. Cortisol suppression (%B) after 8 mg of dexamethasone was evaluated as a supplementary criterion.

Results: An increase in ACTH of 43% at 15 min after CRH was the strongest predictor of CD, with a positive likelihood ratio of 14.0, a sensitivity of 83%, a specificity of 94%, a positive predictive value of 98% and a negative predictive value of 58%. All of the other criteria of stimulated ACTH and cortisol levels were not superior in predicting CD in response to CRH injection. The addition of cortisol suppression by dexamethasone did not increase the discriminatory power. However, the combination of a positive ACTH response at 15 min and a positive HDDS test excluded ECS in all cases.

Conclusion: The present findings support the use of plasma ACTH levels 15 min after the injection of human CRH as a response criterion for distinguishing between CD and ECS. The addition of the HDDS test is helpful for excluding ECS when both tests are positive.

Introduction

Adrenocorticotropic (ACTH)-dependent cortisol excess accounts for more than 80% of all cases of endogenous Cushing’s syndrome (CS) (1). Distinction between pituitary and ectopic ACTH production is crucial for further treatment decisions. Because imaging can be unsatisfactory, or even misleading, in the differential diagnosis of ACTH-dependent Cushing’s syndrome, a precise biochemical diagnosis is mandatory. Bilateral inferior petrosal sinus sampling (IPSS), the gold standard test, is invasive and requires technical experience. Additionally, although a positive result predicts Cushing’s disease (CD) with a positive predictive value (PPV) of about 99%, the negative predictive value (NPV) is only about 20% (2).
The corticotropin-releasing hormone (CRH) test is comparatively simple and therefore widely used for the evaluation of ACTH-dependent Cushing’s syndrome. The rationale behind the CRH test is that corticotroph adenomas usually exhibit a hyper-responsiveness to CRH, whereas the majority of ectopic ACTH-producing tumors do not respond to CRH. However, clinical experience and former research have demonstrated that a proportion of pituitary and ectopic tumors do not follow this functional pattern, which thus limits the reliability of the test (1). Ovine CRH was identified in 1981 and has since been utilized in a large number of studies. Human CRH, which is supposed to be less immunogenic, is primarily used in Europe. Human CRH has a shorter circulation half-life that seems to provoke a less pronounced and shorter ACTH response as compared to ovine CRH. Evoked increases in cortisol levels are not different because the adrenal cortex is maximally stimulated by endogenous ACTH in both forms of CRH (3, 4).

Although there is a body of literature that has examined the diagnostic accuracy of the CRH test alone or in combination with the high-dose dexamethasone suppression (HDDS) test, the results of these analyses are equivocal (5, 6). There is substantial disagreement about which response parameters and cutoff levels are most suitable for differential diagnosis of CS. As a result, protocols of the CRH test vary considerably at different centers with respect to time points and duration of sampling, the injection of human or ovine CRH, the dosage of CRH, and the criteria for interpreting the CRH test.

The aim of the present study was to reevaluate the criteria for ACTH and cortisol responses to 100 µg human CRH in a large series of patients with ACTH-dependent Cushing’s syndrome. Furthermore, we evaluated the additional value of the HDDS test in order to distinguish the etiology of ACTH-dependent CS.

**Subjects and methods**

We retrospectively analyzed serum cortisol and plasma ACTH concentrations of CRH tests in patients with confirmed CD and confirmed ECS. A cohort of 87 of the 152 (57%) patients treated at our institution for ACTH-dependent CS from 1994 until 2014 was included in the analyses. The remaining patients were excluded because no CRH test was performed at our institution (n=56), because ACTH-dependent CS was of undefined origin, or because the patient was lost to follow-up (n=9). The pituitary origin of ACTH secretion was established by biochemical remission following transsphenoidal surgery (TSS), histopathology, and/or temporary adrenal insufficiency after surgery. Ectopic ACTH production was confirmed by histopathology and/or cure after surgery. In nine of the patients with CD, a second CRH test was performed because of recurrent disease. Altogether 96 CRH tests were analyzed. In 81% of the tests, the final diagnosis was CD (n=78), and in 19% of the tests the patients were diagnosed with ECS (n=18). Additionally, in 74 of 96 cases (77%), data from overnight 8 mg dexamethasone suppression testing were available.

The CRH test was performed in the morning after an overnight fast. Blood samples for serum cortisol and plasma ACTH were drawn at −15 and 0 min before and at 15, 30, 45, 60, 90, and 120 min after the injection of 100 µg human CRH (Ferring Arzneimittel GmbH).

The following criteria for differentiating between CD and ECS were tested: i) the percentage increases above basal (%B) for ACTH and cortisol at each time point between 15 and 120 min; ii) the sum of ACTH and cortisol increases (%B) at various time points; iii) the maximal increase in ACTH and cortisol above basal (Δmax %B) across the investigation interval; iv) the area under the curve of ACTH and cortisol responses (AUC %B) as calculated by trapezoidal approximation; and v) the mean cortisol increase 15 and 30 min after CRH, which was a criterion proposed in a previous study (7).

In addition to the CRH test, a subgroup of the patients underwent HDDS testing, with 8 mg of dexamethasone (Jenapharm GmbH, Jena, Germany) being administered at 2300 h and a cortisol measurement at 0800 h the next morning. The suppression of cortisol secretion as a percentage of the basal cortisol level measured on a different day between 0800 and 1000 h was calculated. Additionally, the sum of ACTH secretion following CRH stimulation and cortisol suppression following dexamethasone administration was examined.

Data are presented as means and s.e.m. The optimal cutoff level and associated sensitivity and specificity were evaluated by receiver operator characteristics (ROC) curve analyses based on the highest positive likelihood ratio (+LR) using GraphPad Prism 5 software (GraphPad, San Diego, CA, USA). PPV and NPV were calculated with Bayes’s formula. Post-test probability was calculated using the following formulas: post-test probability = post-test odds/(1 + post-test odds); post-test odds = pre-test odds × LR, pre-test odds = pre-test probability/(1 − pre-test probability); pre-test probability = prevalence. Finally, we estimated the number of patients that were either correctly diagnosed or misdiagnosed when the strongest
criteria of the CRH test and the dexamethasone test were applied.

ACTH and cortisol were measured using an automated chemiluminescence assay (Liaison, Diasorin, Italy) as per the manufacturer’s instructions. In our hands, for ACTH, within-assay coefficient of variations (CV values) were 6.7% at 5.8 pg/ml and 1.2% at 355.4 pg/ml, and between-assay CV values were 12.7% at 13.9 pg/ml and 9.4% at 354.7 pg/ml. For cortisol, within-assay CV values were 2.2% at 0.5 µg/dl and 3.9% at 4.5 µg/dl, and between-assay CV values were 9.8% at 0.2 µg/dl and 9.6% at 4.4 µg/dl.

Data for the present analysis were obtained from the Munich center of the German Cushing Registry. All of the patients included in the present analysis gave written informed consent for the use of their data. The German Cushing Registry was approved by the ethics committee of the University Hospital of Munich and the ethics committees of the participating centers.

**Results**

The patient’s characteristics are reported in Table 1. Patients with confirmed ECS suffered from pulmonary carcinoid in 14 cases, from small-cell lung cancer in three cases, and from an ACTH-producing paraganglioma in one case.

In patients with CD, ACTH concentrations significantly increased from 71 (±8) to a maximum of 191 (±20) pg/ml (P<0.001) at 15 min and reached basal values 90 min after CRH injection. In patients with ECS, there was no significant rise in ACTH values (Fig. 1A). Cortisol concentration after CRH increased from 23 (±1) to a maximum of 36 (±4) µg/dl after 45 min (P<0.001) in patients with CD. Again, there was no significant increase in serum cortisol from the basal value (31 ±5 µg) after CRH in patients with ECS (Fig. 1B). There was considerable overlap of ACTH and cortisol responses between patients with pituitary and ectopic Cushing’s syndrome (Fig. 2). The results of the ROC curve analysis of different test variables along with the corresponding cutoff level and statistic characteristics are provided in Table 2.

An ACTH increase of ≥43% at 15 min after CRH administration had the highest +LR, sensitivity, and NPV along with high specificity and high PPV. ACTH values obtained after 15 min after CRH injection had lower sensitivity, and those obtained after 60 min also had lower specificity as compared to the early increase. Incremental ACTH in percentage above basal over 120 min had lower +LR, sensitivity, and NPV as compared to ACTH increase after 15 min as well. Maximal ACTH increase over 120 min had less favorable characteristics than ACTH measurement after 15 min did.

For cortisol, the most favorable ROC curve statistic was achieved after 30 min, with a cutoff of ≥32% increase. All of the other time points were associated with lower sensitivity and a lower LR. Likewise, incremental cortisol over 120 min and maximal cortisol rise were less sensitive for discriminating between CD and ECS.

The combination of cortisol and ACTH increase in percentage above basal at various time points showed lower sensitivity than ACTH increase at 15 min alone, along with the same specificity and slightly lower LR.

In clinical practice, commonly used cutoff values of the CRH test are a 30% rise in cortisol and a 50% rise in ACTH after human CRH injection. Applying these criteria, ROC curve analysis showed a sensitivity of 78%, a specificity of 78%, and an LR of 3.5 for cortisol and a sensitivity of 83%, a specificity of 89% and an LR of 7.5 for ACTH.

When ROC analysis was performed using mean cortisol increase after 15 and 30 min, as previously published (7), a maximal LR of 10.5 at a cutoff level of 24% with a sensitivity of 59% and a specificity of 94% was achieved. A cutoff value of 14%, as proposed by Newell-Price et al., had higher sensitivity (73%) at the cost of lower specificity (78%), which resulted in a low LR of 3.3. Mean ACTH increase at 15 and 30 min after CRH injection resulted in a maximal LR of 13.1 at a cutoff level of 77%, with 73% sensitivity and 94% specificity. When a previously hypothesized cutoff level of 9% was applied (8), specificity dropped to 50%.

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**Table 1 Patient’s characteristics.**

<table>
<thead>
<tr>
<th></th>
<th>n (%)</th>
<th>Age (years)</th>
<th>Females, n (%)</th>
<th>TSS, n (%)</th>
<th>BADx, n (%)</th>
<th>ACTH&lt;sub&gt;A&lt;/sub&gt; (pg/ml)</th>
<th>Cortisol&lt;sub&gt;B&lt;/sub&gt; (µg/dl)</th>
<th>HDSS, n (%)</th>
<th>i Cortisol (%&lt;sub&gt;B&lt;/sub&gt;)</th>
<th>IPSS, n (%)</th>
<th>Positive IPSS (%)</th>
</tr>
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<tbody>
<tr>
<td>CD</td>
<td>78 (81)</td>
<td>43 ±2</td>
<td>61 (78)</td>
<td>78 (100)</td>
<td>11 (14)</td>
<td>71 ±8</td>
<td>23 ±1</td>
<td>64 (82)</td>
<td>71 ±5</td>
<td>23 (30)</td>
<td>19 (83)</td>
</tr>
<tr>
<td>ECS</td>
<td>18 (19)</td>
<td>51 ±4</td>
<td>11 (61)</td>
<td>2 (11)</td>
<td>7 (39)</td>
<td>106 ±19</td>
<td>31 ±5</td>
<td>14 (78)</td>
<td>31 ±7</td>
<td>5 (28)</td>
<td>0 (0)</td>
</tr>
</tbody>
</table>

CD, Cushing’s disease; ECS, ectopic Cushing’s syndrome; TSS, transphenoidal surgery; BADx, bilateral adrenalectomy; HDSS: high dose (8 mg) dexamethasone suppression; i Cortisol %<sub>B</sub>, percentage suppression of cortisol after HDSS above basal; IPSS, inferior petrosal sinus sampling; Positive IPSS, central to peripheral gradient after CRH >3.
ROC curve analysis of the HDDS test showed the highest LR (8.9), with a cutoff value of 71% suppression of basal cortisol and a corresponding sensitivity of 64% and specificity of 93%. The clinically recommended cutoff for 50% cortisol suppression had an LR of only 3.0, and even though sensitivity was strong (86%), specificity reached only 71%. The combination of both tests calculated the sum of cortisol suppression and ACTH increase at 15 min after CRH injection and had an LR of 11.0, a cutoff value of 119%, a sensitivity of 80%, a specificity of 93%, a PPV of 98%, and an NPV of 53%. When the strongest criteria of the CRH test (43% increase in ACTH 15 min after CRH injection) and the results of the HDDS test were concurrently applied (n = 74), 68% of the patients were correctly diagnosed as CD, and no patient with ECS was misdiagnosed (PPV: 100%, NPV: 43%). Both tests were negative in only 2% of the patients with CD, whereas 64% of the patients with ECS had negative results when both tests were applied together (Fig. 3).

The results of the ROC curve analysis separated for female and male patients (data not shown). The cortisol and ACTH excursions after CRH were not statistically different between female and male patients (data not shown). The results of the ROC curve analysis of the female cohort were similar to the results obtained from the entire cohort. An increase in ACTH after 15 min did not differ between the female cohort and the entire cohort. Values of post-test probability calculated directly from the female cohort did not differ much from the post-test probability of the entire cohort (data not shown). ROC curve characteristics of the male cohort differed from the female cohort as well as from the entire cohort.

**Discussion**

Although the CRH test for the differential diagnosis of ACTH-dependent CS has been evaluated in the past, the present analysis reveals some new aspects. The most interesting outcome is that ACTH increase after 15 min was the strongest predictor of CD and was superior to all of the additional parameters evaluated. The amplitude of fast ACTH rise after 15 min in patients with CD yielded the highest discriminatory power as compared to the flat curve of ACTH in patients with ECS (Figs 1 and 2). In addition, ACTH increase after 15 min was a stable criterion in the sex-specific analysis, with almost identical results for the female and for the total cohort. Results from ROC curve analysis of the male cohort did not allow firm conclusions to be drawn because of the low number of subjects in the male cohort (<20 cases). ROC curve analysis is suggested to allow valid conclusions only with more than 90 observations.

The present analysis was performed with human CRH. Former kinetic analyses have demonstrated that in comparison to ovine CRH, the human peptide has a faster metabolic clearance rate, which results in a less pronounced and shorter ACTH peak (3, 9). This might explain why in previous studies with ovine CRH, later time points across the time series showed better diagnostic abilities as compared to ACTH increase after 15 min in the present series (10). Thus, the present results are specific for human and not ovine CRH tests.

A large European series examined 171 CRH tests using human and ovine CRH (11) and reported a sensitivity of 85% and a specificity of 100% at a maximum increase in
ACTH of $50\%$ after CRH. An increase in ACTH of $50\%$ is also often used in clinical practice. We applied this criterion to the present series, and it resulted in similar sensitivity but lower specificity. In the present series, there were two patients with ECS who responded with a distinct rise in ACTH and cortisol to CRH ($90\%$ and $50\%$ respectively) despite a histology-proven ACTH-producing carcinoid. These patients had particularly low basal ACTH and cortisol levels, which possibly contributed to a diminished negative feedback inhibition that is otherwise believed to be one mechanism for negative CRH response in patients with ECS (12). Thus, mild or cyclic hypercortisolism could be a cause for false-positive ACTH response to CRH in ECS, and the timing of functional testing might impact the outcome.

In the present analysis, an ACTH increase at 15 min had a better discriminative power, with a higher number of correctly identified patients with ECS as compared to maximal ACTH increase after CRH injection. A possible explanation for this observation might be that a longer sampling period increases the likelihood of spontaneous bursts of ACTH secretion in ectopic tumors that resemble CRH-stimulated peaks in CD.

Newell-Price et al. (7) investigated a large series of human CRH tests and found a maximal ACTH response of $105\%$ above basal to have $100\%$ specificity but a rather low sensitivity. Because of the low sensitivity, these authors postulated the superiority of cortisol as the best criterion for differentiating between CD and ECS. In contrast, the present data showed a sluggish and quantitatively small cortisol increase (Fig. 1) and less favorable results in ROC analysis as compared to ACTH (Table 2). The major methodological difference between their work and the present study is that we used ROC curve analysis based on the highest LR to define cutoff values. LRs express the clinical usefulness of a test by measuring the change in the certainty of diagnosis. An LR greater than 10 is considered to be a strong indicator that the test will predict the presence of the disease. In contrast, Newell-Price et al. estimated their cutoff levels at a preset

**Figure 2**

Distribution of different response variables of the corticotropin-releasing hormone (CRH) test and the high-dose dexamethasone suppression (HDDS) test in a series of 78 patients with Cushing’s disease (CD, closed symbols) and 18 patients with ectopic Cushing’s syndrome (ECS, open symbols). The bar indicates the cutoff level obtained by ROC curve analysis based on the highest likelihood ratio. The ROC curve of each analysis is inserted in the right corner of the graph.
specificity of 100%. We decided not to follow this approach, because we believe that 100% specificity is difficult to obtain with the CRH test. It has been well documented that a small number of patients with ECS show a clear cortisol and ACTH response to CRH, especially patients with carcinoids (13, 14, 15, 16, 17, 18, 19). Presetting specificity to 100% will always lead to a pronounced reduction in sensitivity. Because the pretest probability of CD is more than four times higher than that of ECS, a significant reduction in sensitivity can cause a greater proportional increase in false-negative results.

In contrast to the results of the present analysis, several other reports have found serum cortisol to be a stronger predictor of CD than ACTH is (8, 20, 21). Those earlier analyses reported a cortisol increase of 20% or even less as the most suitable cutoff. In the present series, a cutoff for cortisol of less than 20% would indeed be associated with a higher sensitivity, but it would simultaneously result in a low specificity of less than 75%, because more than one-quarter of the patients with ECS exhibited maximal cortisol excursions of more than 20% above basal. In general, using a threshold that is less than four times the intra-assay CV (5–8%) is controversial, seeing as test results become more vulnerable to other effects than they do to the CRH stimulus. Therefore, we believe that the ACTH increment in response to CRH is a more robust criterion than a modest cortisol rise is.

In the present series, ROC curve analysis proved the HDDS test to be inferior to the CRH test in distinguishing CD from ECS. The combined analysis of the CRH and the HDDS test did not increase diagnostic precision in the ROC analysis. These results are in accordance with previous data that showed that the HDDS test in combination with the CRH test was not superior to the CRH test alone (13, 19, 22) or the CRH test in combination with the 48 h low-dose dexamethasone suppression test (16). Nevertheless we believe that the HDDS test can still have some impact in the differential diagnosis of

<table>
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<tr>
<th>Test variable</th>
<th>Cutoff value (%)</th>
<th>ROC AUC (S.E.M.)</th>
<th>Positive LR</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
<th>PPV (%)</th>
<th>NPV (%)</th>
<th>Post-test probability (%)</th>
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<tbody>
<tr>
<td>ACTH (15 min)</td>
<td>≥43</td>
<td>0.89 (0.03)</td>
<td>14.9</td>
<td>83</td>
<td>94</td>
<td>98</td>
<td>58</td>
<td>98</td>
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<tr>
<td>ACTH (30 min)</td>
<td>≥62</td>
<td>0.91 (0.03)</td>
<td>13.9</td>
<td>77</td>
<td>94</td>
<td>98</td>
<td>50</td>
<td>98</td>
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<tr>
<td>ACTH (45 min)</td>
<td>≥51</td>
<td>0.89 (0.04)</td>
<td>12.5</td>
<td>69</td>
<td>94</td>
<td>98</td>
<td>40</td>
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<tr>
<td>ACTH (60 min)</td>
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<td>0.85 (0.05)</td>
<td>8.0</td>
<td>42</td>
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<td>97</td>
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<tr>
<td>Cortisol (15 min)</td>
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<td>ACTH (15 min) + Cortisol (30 min)</td>
<td>≥110</td>
<td>0.90 (0.03)</td>
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<td>74</td>
<td>94</td>
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<td>ACTH (30 min) + Cortisol (30 min)</td>
<td>≥108</td>
<td>0.90 (0.03)</td>
<td>12.8</td>
<td>71</td>
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<td>45</td>
<td>98</td>
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<tr>
<td>Δmax ACTH (120 min)</td>
<td>≥95</td>
<td>0.90 (0.04)</td>
<td>12.9</td>
<td>71</td>
<td>94</td>
<td>98</td>
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<td>98</td>
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<tr>
<td>Δmax Cortisol (120 min)</td>
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<td>0.88 (0.04)</td>
<td>10.5</td>
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<tr>
<td>AUC ACTH (%B)</td>
<td>7375</td>
<td>0.89 (0.04)</td>
<td>9.0</td>
<td>52</td>
<td>94</td>
<td>97</td>
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<tr>
<td>AUC Cortisol (%B)</td>
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<td>0.86 (0.05)</td>
<td>10.4</td>
<td>67</td>
<td>94</td>
<td>98</td>
<td>38</td>
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<tr>
<td>HDDS + ACTHCRH 15 min</td>
<td>≥119</td>
<td>0.94 (0.03)</td>
<td>11.2</td>
<td>72</td>
<td>93</td>
<td>98</td>
<td>53</td>
<td>98</td>
</tr>
</tbody>
</table>

AUC, area under the curve; +LR, positive likelihood ratio; PPV, positive predictive value; NPV, negative predictive value.

Figure 3
Combined test response to the corticotropin-releasing hormone (CRH) test (response criterion: ACTH rise of ≥43%B 15 min after the injection of 100 μg human CRH) and the high-dose dexamethasone suppression (HDDS) test (response criterion: cortisol suppression of ≥50%B after 8 mg of dexamethasone) in patients with Cushing’s disease (CD) and patients with ectopic Cushing’s syndrome (ECS).
ACTH-dependent CS. We found that not one single patient with CD was misdiagnosed as having ECS when the combination of a positive CRH test and a positive HDDS test were present (PPV 100%). Therefore, a positive CRH test with an ACTH of >43% after 15 min and a positive HDDS test of more than 50% cortisol substantiated the diagnosis of pituitary CS, which led to TSS without further biochemical tests. Still, 30% of the patients with CD and 36% of the patients with ECS had a mixed response. In these situations, the next logical step for differentiating CD from ECS would be IPSS. Imaging of the pituitary region is generally performed before IPSS. Pituitary MRI might reveal a larger pituitary tumor, which would indicate an ACTH-producing pituitary adenoma. On the other hand, in up to 50% of patients with proven CD, the MRI fails to detect the source of ACTH secretion, and more than 15% of the patients with ECS carry incidental pituitary tumors. Thus, from a clinical point of view, imaging is required before IPSS, whereas from a scientific point of view, the diagnostic abilities of imaging are limited and are inferior to the discriminatory abilities of dynamic biochemical tests. IPSS has been suggested to be the most accurate test for differentiating CD from ECS. The predictive value of a positive IPSS with CRH stimulation is high. False-positive IPSS are either a result of ectopic CRH secretion or of the insufficient suppression of regular corticotropin feedback by only mild or periodic hypercortisolism. The rate of false-negative IPSS is, to some extent, influenced by the investigators’ experience, and it ranges from 80 to 99%. In cases of negative IPSS despite normal venous anatomy:draining, prolactin normalized IPS:P ratios might decrease the number of false-negative results. Because neither biochemical functional testing nor imaging and IPSS have 100% sensitivity and specificity in the differential diagnosis of ACTH-dependent CS, only a differentiated combination of these tests can minimize the number of patients with uncertain diagnoses.

In summary, the present study has two main findings: i) an early ACTH increase after CRH stimulation identifies patients with CD; and ii) ECS is excluded when the CRH test plus the HDDS test are positive. However, like all previous series, the present work has important limitations. First, the present study was designed as retrospective analysis that is susceptible to selection bias, especially seeing as not all of the patients who underwent CRH testing were available for follow-up. Second, the overall number of patients included in the present analysis and, in particular, the number of patients with ECS was small, which limited the study power. If these results can be confirmed in larger prospective series and be implemented in clinical practice, the diagnostic work-up in patients with ACTH-dependent CS could be simplified and standardized, and this is urgently needed.

Conclusion

The present analysis supports the use of plasma ACTH increase 15 min after CRH injection as a powerful response criterion for distinguishing between CD and ECS. If these results can be confirmed in a larger prospective series, a single sample after a CRH injection could replace the time-consuming sampling of time series. In addition, a positive CRH test using ACTH response after 15 min in combination with a positive HDDS test allowed excluding ECS in patients with ACTH-dependent CS in the present series.

Declaration of interest

M Reincke has served on the advisory boards of Novartis and has received lecture fees and grants from Novartis, Ipsen, and Pfizer. F Beuschlein has received grant support from Novartis. H Schneider has received lecture fees from Novartis and Pfizer and research grants from Pfizer. J Schopohl has received lecture fees from Novartis, Ipsen, and Pfizer. All the other 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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