Markers of tumor invasion are major predictive factors for the long-term outcome of corticotroph microadenomas treated by transsphenoidal adenomectomy

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Abstract

Objective: To assess the postsurgical outcome of patients with corticotroph microadenomas and to define predictors of the long-term outcome, with special emphasis on markers of tumor extension.

Design: Prospective study of 53 corticotroph microadenomas treated by enlarged adenomectomy. Patients followed for at least 2 years were classified into two groups: those in long-term remission and uncured patients (immediate failures and recurrences). Pre-, per- and postoperative parameters were analyzed as predictors of the long-term outcome.

Methods: Baseline hormone assessments were performed preoperatively, 8 days after surgery and every 6–12 months thereafter. Pituitary magnetic resonance imaging (MRI) allowed analysis of possible tumor extension to adjacent structures. Apparent completeness of the surgical removal was determined, and fragments labeled either ‘tumor’ or ‘surrounding pituitary tissue’ were submitted to serial sectioning.

Results: Immediate control of hypercortisolism was achieved in 43/53 patients (81%). However, later recurrences were observed in five patients (9%). Preoperative MRI showed tumor extension into adjacent structures with good specificity (91%) for prediction of surgical failure. Evidence of local invasion at surgery was also significantly predictive of the long-term outcome. A corticotroph adenoma was found at histological examination in 96% of the patients, and 26% had irregular limits, a feature significantly correlated with a poor outcome. Immediate postoperative plasma cortisol did not allow discrimination between long-term remissions and recurrences.

Conclusion: Surgical failure was best predicted by signs of tumor ‘invasiveness’ at MRI, confirmed by peroperative examination and histology.

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Introduction

Cushing’s disease is the first cause of hypercortisolism, but remains a rare affliction. Its incidence in the general population is estimated to be between 0.7 and 2.4 cases/million per year (1, 2). Without treatment, hypercortisolism leads to severe complications and is associated with premature mortality. Most pituitary tumors in Cushing’s disease are microadenomas. Since the first removal of a corticotroph microadenoma by transsphenoidal surgery by Hardy in 1969 (3), pituitary microsurgery has become the treatment of choice for Cushing’s disease (4, 5). The immediate success rate ranges from 70 to 85% (1, 6–12). However, the recurrence rate after initially successful surgical treatment increases with the duration of the follow-up, to reach as much as 20% at 5–6 years (1, 9, 12–15).

It is widely accepted that measurement of postoperative cortisol concentrations can be used both to assess the immediate control of hypercortisolism and to provide a partial prediction of the likelihood of recurrence (9, 13, 16–18). In contrast, several attempts at using dynamic tests in this setting have provided controversial results (1, 12, 15–17, 19, 20). The principal aim of our study was thus to determine other prognostic factors that may be linked to either absence of immediate cure or later recurrence of Cushing’s disease after transsphenoidal surgery. In addition to the clinical and biological parameters usually monitored in these patients, we more particularly focused our attention on: 1) the evaluation of...
tumor invasion by preoperative magnetic resonance imaging (MRI) by the same neuroradiologist (N G) using three-dimensional sequences; 2) the systematic preoperative evaluation of the efficacy of the surgical removal by the neurosurgeon; 3) the careful histological analysis of the adenomas on serial sections of all removed fragments, in addition to the analysis of the juxtatumoral pituitary tissue. Identification of several possible risk factors might be helpful in improving the final success rate of surgery. In this respect, a surgical strategy characterized by a deliberate attempt at performing a curative removal of the tumor by the same experienced neurosurgeon (F G) was chosen. On the assumption that unsuccessful surgical removals were accounted for by insufficient adenomectomy because of the presence of tumoral cells at the periphery of the adenoma, an enlarged adenomectomy was carried out (21).

From 1989 to 1998, a cohort of 53 consecutive patients presenting with corticotroph microadenomas and treated by enlarged adenomectomy was prospectively studied using this approach and has been followed for a mean duration of 37.9 ± 22.1 months (range 24–120 months) after transsphenoidal surgery.

Patients and methods

Patients

In each patient, the diagnosis of corticotroph microadenoma was established on the basis of a combination of the usual criteria: clinical findings, baseline and pharmacodynamic hormone evaluations, radiological evidence of a microadenoma, pituitary–peripheral gradient at intrapetrosal sinus sampling, macroscopic and microscopic confirmation. During the study period, 53 consecutive patients with corticotroph microadenomas were operated on by enlarged adenomectomy. The population consisted of 49 women and four men, aged 39 ± 13 years (range 13–62 years). During the same period, five other patients with histologically proven small corticotroph microadenomas were treated by total hypophysectomy and four patients were treated by anticortisol therapy in the absence of radiological or peroperative visualization of a microadenoma.

Hormone evaluation

Nyctohemeral variations of plasma adrenocorticotrophic hormone (ACTH) and cortisol concentrations were measured every 4 h over 24 h in the preoperative period, 8 days after surgery, every 6 months for 18 months after surgery, and yearly thereafter. The plasma cortisol analysis was performed by a competition radioimmunoassay (Immunotech, Marseille, France), with normal values ranging from 200 to 700 nmol/l at 0800 h and 50 to 300 nmol/l at 2000 h, and an assay sensitivity of 20 nmol/l. Plasma ACTH concentrations were measured by an immunoradiometric assay (Allegro kit, Nichols Institute, San Juan, Capistrano, CA, USA), with normal values ranging from 10 to 55 ng/l at 0800 h and 5 to 20 ng/l at 2000 h, and an assay sensitivity of 5 ng/l. Urinary free cortisol (UFC) was measured after extraction, using a radioimmunoassay (Immunotech); normal values ranged from 80 to 300 nmol/d.

To confirm the diagnosis of Cushing’s disease, all patients underwent a low-dose dexamethasone suppression test (0.5 mg every 6 h during 48 h, absence of suppression of the plasma cortisol to 50 nmol/l or less confirming Cushing’s syndrome), followed by a high-dose dexamethasone suppression test (2 mg every 6 h during 48 h, suppression of plasma cortisol by 50% or more and UFC by 90% or more being consistent with Cushing’s disease) (9, 12, 22). Bilateral inferior petrosal sinus sampling for ACTH measurement before and after i.v. administration of 100 µg corticotropin-releasing hormone (CRH) was performed through a bilateral femoral vein approach in seven patients, to ascertain the pituitary origin of the ACTH hypersecretion. In all patients the central/peripheral gradient was >2 at baseline and >3 after CRH (9, 12, 22).

Other pituitary functions were assessed by determination of basal hormone concentrations, by thyrotrophin-releasing hormone (TRH) and gonadotropin-releasing hormone (GnRH) stimulation tests, and by measurements of urine volume and density. No substitutive cortisol replacement therapy was given before the hormone evaluation performed 8 days after surgery. All subsequent cortisol/ACTH measurements in the patients receiving postoperative cortisol replacement therapy were performed 48 h after discontinuation of their cortisol treatment.

Radiological investigations

Fifty-one patients underwent pituitary MRI performed by the same neuroradiologist (N G). Images were obtained on a 1.0 T system (Magnetom SP 40; Siemens, Erlangen, Germany) using precontrast 3-mm thick contiguous coronal spin echo T1-weighted images (repetition time (TR) 450 ms; echo time (TE) 22 ms; three excitations), followed by postgadolinium three-dimensional fourier transformed (FT) T1-weighted imaging using a fast, low-angle shot (flash) sequence (TR 40 ms; TE 5 ms; flip angle 40°) or turbo flash sequence (TR 10 ms; TE 4 ms; inversion time 200 ms; flip angle 12°). The three-dimensional FT T1-weighted sequences provided 1-mm thick contiguous sections and reformatted images in any desired plane. The acquisition was performed in the sagittal plane to avoid aliasing phenomena. Reformatted images were obtained in the coronal plane, the oblique plane oriented along the pituitary stalk, and the axial plane oriented along the sellar floor. Additional 3-mm thick coronal T2-weighted images (turbo spin echo sequence;
to classify each adenoma either as ‘invasive’ or ‘non-invasive’, cases without definite evidence of extension beyond the anterior pituitary being classified into the latter group. Cavernous sinus or posterior lobe invasion was suspected on the basis of an uninterrupted extension of the adenomatous signal into the corresponding structure as illustrated in Fig. 1b. For various reasons, the two remaining patients were investigated by computed tomography only.

**Surgery**

All patients underwent surgery via the transsphenoidal approach, performed by the same neurosurgeon (F G) using a microscopic technique. An enlarged adenomectomy, removing the adenoma and the surrounding parenchyma, was performed as described previously (21). Before the adenoma was sought or penetrated, the two lateral wings were dissected, the surgeon keeping the approach outside the capsule. If the adenoma did not cross the midline, a section from above downwards was made, starting from the edge of the stalk, allowing removal, in one block, of the adenoma, the adjacent pituitary gland and its capsule. If the adenoma crossed the midline, it was removed by cutting a slice of normal pituitary gland about 1 mm thick and finally removing the lateral hypophseal capsule in contact with the cavernous sinus. In each case, the neurosurgeon stated whether the removal was considered to be complete or partial on the basis of peroperative findings. Fragments labeled either ‘tumor’ or ‘surrounding pituitary tissue’ were placed into separate containers and sent to the pathologist.

**Histological methods**

All tumor specimens obtained at surgery were immediately fixed in Bouin-Holland and studied by the same pathologist (J T). When the tumor was not found at the first histological analysis ($n = 15$), the entire fragments were cut into serial sections. The presence of adenomatous tissue in surrounding pituitary samples was considered as indicative of microscopic tumor extension. The sections were stained by Herlant tetrachrome and PAS–Orange G and treated by the immunoperoxidase technique with streptavidin–biotin-complex. The following monoclonal (m) and polyclonal (p) antibodies were used at 1:200 to 1:4000 dilution: anti-p17-39-ACTH (donated by M P Dubois, Nouzilly, France), anti-pβendorphin and anti-pβlipotropin (LPH) (donated by B Clauserat Lyon, France), anti-m human prolactin (hPRL) and anti-m α-subunit (Immunotech), anti-p human growth hormone (donated by Dr Raiti, National Institute of Arthritis, Diabetes, Digestive and Kidney Diseases).
Surgical outcome

The patients were classified into two groups according to the findings of their individual latest evaluation performed 2–10 years after surgery (3.2 ± 1.8 years). Patients were considered to be in remission (n = 38) on the basis of clinical cure and a 0800 h plasma cortisol concentration <138 nmol/l at the day 8 evaluation (9), normalization of UFC and resumption of plasma ACTH and cortisol nyctohemeral variations at subsequent evaluations. Uncured patients (n = 15) included immediate failures (n = 10) characterized by 0800 h cortisol values >138 nmol/l at the day 8 evaluation and increased cortisol concentrations at subsequent evaluations that proved to be partially suppressible during the low-dose dexamethasone test, and recurrences of hypercortisolism (n = 5) defined as recurrence of clinical and biochemical markers (including partial suppression at the low-dose dexamethasone test) of hypercortisolism 12–84 months (mean 37.2 ± 27.6 months) after surgery in patients who had a morning cortisol concentration <138 nmol/l at the early postoperative evaluation.

Statistical analysis

Quantitative variables are expressed as mean ± S.E.M. and were analyzed with the Mann–Whitney test (non-parametrical study). Qualitative variables were analyzed using Fisher’s test. Statistical significance was achieved at a probability (P) value less than 0.05. All statistical tests were performed using the Stat Graph Plus statistical software package.

Results

Preoperative clinical and biological status

Age at diagnosis was not significantly different according to the outcome: 39 ± 12 years in patients in remission and 38 ± 15 years in those who were uncured. Preoperative UFC measurements averaged 1333 ± 985 nmol/day in the whole population and did not differ according to the outcome (1276 ± 969 nmol/day in cured and 1484 ± 1049 nmol/day in uncured patients), in keeping with ACTH and cortisol concentrations (data not shown). The other anterior pituitary functions were normal in all patients.

Preoperative MRI findings

Among the 51 patients examined by MRI, a focal lesion suggestive of an intrasellar adenoma was identified in 48 (sensitivity = 94%) in the left (n = 21) or right (n = 24) side or in the median part (n = 3) of the gland (Fig. 1a), and signs of tumor extension to adjacent structures were present in 10 (Fig. 1b). Taking into account the whole population of 62 patients with Cushing’s disease consecutively treated during the study period, MRI had a sensitivity of 48/60 (80%) in identifying a microadenoma, as nine patients not treated by enlarged adenomectomy (because of a lack of visualization of their adenoma) were not included in the present report. In this series, the pituitary gland appeared heterogeneous after gadolinium enhancement without a tumoral image in three patients (Fig. 1c). The largest diameter of the adenoma ranged from 2 to 10 mm, averaging 5.9 ± 2.6 mm, and was not significantly different in the two groups examined (Table 1). Signs of invasiveness were present in 10 of the individualized adenomas (21%): cavernous invasion was documented by the presence of tumoral tissue extending into the cavernous sinus (Fig. 1b) in a total of eight patients and posterior lobe or stalk invasion was demonstrated in four. Notably, two patients presented with both types of invasion. Irrespective of the structure invaded, signs of tumor invasion were found to be significantly associated with an increased risk of poor outcome, as such signs were observed in 9% of patients in remission but in 54% of those who remained uncured (Table 1). MRI had very good specificity (91%) for the prediction of surgery outcome (Table 2).

Neurosurgical evaluation

Macroscopic evidence of an intrasellar adenoma was found by the neurosurgeon in all the patients, enabling him to perform an enlarged adenomectomy. Taking into consideration all 58 patients operated on either by enlarged adenomectomy (n = 53) or by hypophysectomy

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**Table 1** Preoperative, peroperative and histological data in 53 patients with corticotroph microadenomas treated by enlarged adenomectomy.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Patients in remission (n = 38)</th>
<th>Uncured patients (n = 15)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Largest diameter at MRI (mm)</td>
<td>6.0 ± 2.7</td>
<td>5.6 ± 2.5</td>
</tr>
<tr>
<td>Cavernous or posterior lobe invasion at MRI</td>
<td>3/35</td>
<td>7/13†</td>
</tr>
<tr>
<td>Cavernous or posterior lobe invasion (evaluation by neurosurgeon)</td>
<td>4/35</td>
<td>11/14‡</td>
</tr>
<tr>
<td>Histological visualization of the adenoma</td>
<td>38/38</td>
<td>13/15</td>
</tr>
<tr>
<td>Presence of adenomatous tissue in surrounding tissue samples</td>
<td>5/26</td>
<td>3/5§</td>
</tr>
</tbody>
</table>

Significantly different from patients in remission: †P = 0.02, ‡P = 0.0001, §P = 0.03.
Sensitivity was calculated as the ratio of the number of true positives to true positives plus false negatives, and specificity as the ratio of the number of true negatives to true negatives plus false positives. Positive predictive value was calculated as the ratio of the number of true positives to true plus false positives, and negative predictive value as the ratio of the number of true negatives to true plus false negatives.

(n = 5) during the study period, an adenoma was identified by the surgeon in 91%. In 15 of those undergoing enlarged adenomectomies, the limits between the tumor and the surrounding normal pituitary were not clear. This macroscopic appearance of local invasiveness was a significantly discriminant parameter of the outcome, with good sensitivity (79%) and specificity (89%) (Tables 1 and 2).

**Histological findings**

An adenoma was identified by histological analysis in 51 of the 53 patients (96%). The two in whom no adenoma was found by the pathologist were classified as early failures and may be explained either by erroneous identification of the adenoma by the neurosurgeon or by aspiration of adenomatous tissue during surgical procedure. By immunohistochemistry, with the exception of two plurihormonal ACTH and PRL adenomas, all tumors were monohormonal corticotroph adenomas. The tumoral cells were positive for antibodies against pro-opiomelanocortin-derived peptides (anti-ACTH, anti-Bendorphin and anti-BlPH). Fragments of the non-tumoral pituitary (samples labeled as surrounding pituitary tissue) were studied in 31 patients. Although, in all of them, no capsule was observed (Fig. 2a), 74% of the tumors were well limited and no adenomatous tissue was present in the surrounding tissue samples from these patients (Fig. 2b). However, in eight, the limits were irregular and tumor cells invaded the juxtatumoral pituitary (Fig. 2c). This feature was significantly correlated with a poor outcome (Table 1), with a specificity of 81% (Table 2). In the juxtatumoral pituitary, numerous PRL cells were found in three patients. The corticotroph cells were either normal or displayed Crooke’s changes in 28 of the 31 patients. Their number varied from rare to numerous, but according to the mapping of human corticotroph cells (23), no ACTH cell hyperplasia was found, whether or not an adenoma was present.

**Postoperative results**

Irrespective of the eventual hormonal outcome, ACTH or cortisol circadian variations at day 8 after surgery were absent or moderate, and no statistically significant difference was found between outcome groups, leading us to compare groups on the basis of the 08:00 h cortisol or ACTH values. Patients in remission and those who remained uncurcured as defined in the Methods section had significantly (P = 0.0001) different morning plasma cortisol concentrations on day 8 (44 ± 43 nmol/l and 334 ± 421 nmol/l, respectively), but this value was not statistically different (P = 0.8) between patients in long-term remission and those who later proved to have recurrent disease (44 ± 43 nmol/l and 54 ± 28 nmol/l, respectively). Similarly, morning ACTH (11 ± 7 ng/l and 36 ± 39 ng/l, respectively) and UFC (24 ± 21 nmol/day and 365 ± 743 nmol/day, respectively) were significantly different (P = 0.03 and P = 0.0001, respectively) between patients in remission and uncurcured patients. Comparison of morning ACTH (11 ± 7 ng/l and 12 ± 7 ng/l, respectively) and UFC (24 ± 21 nmol/day and 36 ± 41 nmol/day, respectively) at day 8 showed no significant difference (P = 0.9 and P = 0.1, respectively) between those in long-term remission and those with later recurrent disease. The duration of postoperative cortisol replacement therapy was similar in all outcome groups: 17 ± 12 months (range 3–72 months) in patients in remission and 22 ± 10 months (range 12–36 months) in uncurcured patients. A transient diabetes insipidus was observed in the immediate postoperative period in nine patients who had normal posterior pituitary function when re-evaluated without treatment 3 months later. At the month 6 visit, urinary volumes and densities were normal in the entire population. Baseline PRL, free thyroxine, TSH, testosterone and estradiol were normal in all patients, as were TSH and gonadotropin responses to TRH or GnRH tests, respectively.

**Discussion**

Immediate control of hypercortisolism was achieved in 81% of our patients, thanks to a surgical strategy aimed at removing the adenoma whenever possible, using enlarged adenomectomy. However, 9% of our population experienced a late recurrence of disease.
As previously described (24), this study used three-dimensional FT MRI to achieve a more accurate determination of local invasiveness and, in particular, spread into the cavernous sinus and the neurohypophysis. Moreover, the peroperative macroscopic evaluation of the extent of tissue requiring removal by the neurosurgeon was closely related to the pathological diagnosis, including an immunocytochemical analysis of the juxtatumoral pituitary. These features, together with more commonly studied parameters were analyzed and confirmed that the frequency of recurrences in this disease is most probably linked to the local invasiveness of corticotroph microadenomas. Our series included 51 patients with histologically proven corticotroph microadenoma and two patients with Cushing's disease confirmed by standard endocrine findings and bilateral intrapetrosal sinus sampling. These patients were further analyzed in terms of prognostic factors to assess the results of transsphenoidal enlarged adenomectomy. It was noteworthy that, despite such an enlarged tumor removal, all pituitary functions – with the exception of the hypothalamo–pituitary–adrenal axis – were found to be normal 3 months after surgery in all patients. The minimum duration of follow-up selected in this study (24 months) does not permit exclusion of the possibility of later recurrences in some of the patients evaluated. However, according to published series (1, 25, 26) most recurrences occur within 2 years after surgery, and our mean duration of follow-up was 38 ± 22 months (range 24–120 months).

In this population, among hypo- or eucortisolic patients, early postoperative morning cortisol concentrations from patients in long-term remission were not

Figure 2  Histology of the corticotroph adenoma and the juxtatumoral pituitary. (a) Limits between a corticotroph adenoma (AD) and the juxtatumoral pituitary (JTP). Notice the absence of capsule (arrow) and the difference of cellular arrangement: diffuse in the tumor; in cords in the juxtatumoral pituitary. All the tumoral cells are PAS positive. The orangeophilic cells are normal somatotrophs (PAS-Orange G, ×1000). (b) This corticotroph adenoma (AD), positive for anti-ACTH antibodies, is round with regular limits (arrow) (×120). (c) This small adenoma (AD), less than 2 mm in diameter, has irregular limits, with cell cords invading (arrow) the juxtatumoral pituitary (JTP) (×250). Immunoperoxidase technique, with anti-17-39-ACTH at a dilution of 1:4000. (d) Diagram showing how tumoral cells may be left behind if a selective adenomectomy is performed.
significantly different from those of patients who later experienced recurrence of hypercortisolism, in contrast with other data reported in the literature (1, 16, 18, 25, 27–29). Several dynamic tests have been used in the assessment of the surgical outcome of Cushing’s disease. Patients with low serum cortisol concentrations that suppress with low-dose dexamethasone test may generally be cured (16, 17). In contrast, a restoration of cortisol suppression has been described after incomplete adenomectomy (19). Moreover, in a few cases, a partial cortisol suppression can exist before surgery for Cushing’s disease (16). Discrepant results were also described with CRH test as a predictor for surgical outcome (1, 12, 15, 20). Recently, desmopressin (ddAVP), a synthetic vasopressin analogue acting on pituitary V3 (or V1b) receptors, which mediates ACTH and cortisol secretion in patients with corticotroph adenomas, was used as a test for the diagnosis of Cushing’s disease (22, 30). However, preliminary postoperative results showed the persistence of ACTH hyperresponsiveness to ddAVP in patients with cured Cushing’s disease. Overall, no test has proved to be completely reliable in the etiological diagnosis of ACTH-dependent hypercortisolism or in the assessment of cure (17, 22).

In our series of patients, tumor visualization by MRI (94%), by the surgeon (100%) or by histological examination (96%) was not correlated with the outcome. This finding was expected because, during the study period, nine patients with corticotroph microadenomas were treated by either total hypophysectomy or anticortisolic therapy because no adenoma had been visualized, and were thus excluded from the present series that analyzed potential predictive criteria for cure after enlarged adenomectomy. Three-dimensional MRI appeared to be very helpful in the management of our patients. Indeed, it allowed visualization of an intrasellar adenoma in 80% of all patients, and in 94% of those treated by enlarged adenomectomy, with exact localization of the tumor in all the positive cases. In previously published series (1, 6, 31), a greater risk of immediate failure seemed to be associated with the absence of visualization of the tumor. Similarly, absence of visualization of the tumor by the surgeon or the pathologist seemed to be associated with a greater risk of surgical failure (1, 7, 12, 31). We paid particular attention to the local invasiveness of corticotroph microadenomas, by means of neuroradiology, surgery and histology. Previous studies had been unable to link the neuroradiological signs of invasiveness to the surgical outcome (6, 12). In our series, MRI revealed invasion of the surrounding structures in 10 patients. This parameter was significantly linked to an increased risk of poor outcome and was characterized by the best specificity score (91%). Combining both kinds of predictive values (positive and negative), signs of tumor invasiveness at MRI can be considered a reliable risk factor of surgical failure. Interestingly, macroscopic evaluation of local invasiveness, an unfrequently analyzed parameter in published series, was also indicative of the surgical result in our patients (sensitivity 79%; specificity 89%). Although such an assessment may have been influenced by the surgeon’s awareness of radiological signs of tumor extension, it was based on peroperative macroscopic observation. The histological analysis of the juxtatumoral pituitary performed on a blind basis showed that 26% of corticotroph microadenomas had irregular limits. This locoregional spreading, observed in our series in 60% of the recurrent tumors and in 19% of long-term remissions, is not related to malignancy or particular aggressiveness. The recurrence may be explained by tumor regrowth from tumoral cells lining the adjacent normal pituitary preserved by the surgeon, especially when tumor limits had not been clearly demarcated (Fig. 2d). Moreover, extension of tumoral tissue into the peritumoral rim in patients in remission suggests that removal of macroscopically normal pituitary tissue around the visible lesion might have been beneficial in these patients. Similar observations have been made in other types of pituitary adenomas (21). Unlike other authors (7), we found no evidence of corticotroph hyperplasia in the specimens analyzed. We thus believe that early surgical failure is not related to corticotroph hyperplasia, but to the fact that the microadenoma has been only partially removed by the surgeon.

The findings of the present study underline the high frequency of both macroscopic and microscopic invasion in corticotroph microadenomas. On a total of 10 parameters analyzed, signs of local tumor invasiveness were found to be good predictors of long-term outcome of patients with Cushing’s disease treated by transsphenoidal adenomectomy, at three-dimensional FT MRI examination, during the surgical procedure, or at histological analysis. Tumor extension thus appeared as the principal risk factor of these microadenomas and justified performance of enlarged adenomectomy. This surgical strategy should decrease the risk of immediate failure or of late recurrence. On an individual basis, however, a long-term clinical and biological follow-up is warranted in all patients, to ascertain a definitive cure.

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References


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