Surgical Management of Giant Pituitary Adenomas

Gérard Mohr, Jules Hardy, Ronald Comtois and Hughes Beauregard

ABSTRACT: During the past 25 years, 77 giant pituitary adenomas have been treated surgically, including suprasellar extensions of type C in 66 cases and of type D in 11 cases. Non-secreting adenomas were present in 53 and secreting adenomas in 24 cases. All patients except 3 presented with significant visual field defects; including bitemporal hemianopia, superior quadranopia or unilateral temporal hemianopia, contralateral blindness in 73% of the cases, and one case with sudden bilateral blindness due to acute pituitary apoplexy. A single transsphenoidal procedure was carried out in 74% of the patients while 11 patients (7%) required reoperations for recurrent or residual tumor. Only 3 patients required a subsequent transcranial procedure. Complications included 1 CSF-leak, 1 empty-sella syndrome and 4 fatal post-operative hematomas. We prefer the transsphenoidal route even in very large or giant pituitary adenomas, since it allows rapid and adequate decompression of the optic nerves and chiasm, avoids major pituitary insufficiency in 60% of the cases and is associated with low morbidity-mortality rates.

RESUME: Traitement chirurgical des adénomes pituitaires géants. Au cours des 25 dernières années, 77 tumeurs hypophysaires géantes ont été traitées chirurgicalement, incluant 66 cas d’extensions suprasellaires de type C et 11 cas de type D. Il s’agissait d’adénomes non-secrétants dans 53 et d’adénomes sécrétants dans 24 cas. Presque tous les malades sauf 3 présentaient des altérations importantes des champs visuels, dont des hémianopsies ou quadransopsies bitemporales, des hémianopsies unilatérales avec cécité contralatérale dans 73% des cas et un patient avec cécité bilatérale brutale par apoplexie pituitaire aiguë. Une seule intervention transphénoidale a été effectuée chez 74% des malades, tandis que 11 (7%) subirent des réinterventions pour tumeur résiduelle ou récidivante. Seulement 3 malades nécessitèrent ultérieurement une chirurgie par voie transcrânienne. Parmi les complications, on observa 1 cas de fistule de LCR, 1 cas de selle vide et 4 hématomes postopératoires mortels. La voie transphénoidale est l’abord de choix même en cas d’adénomes très volumineux ou géants, permettant une décompression rapide et efficace des structures optiques, tout en évitant une insuffisance hypophysaire majeure dans 60% des cas et en assurant des taux de morbidité-létalité très réduits.


Since Jefferson's description of unusually large suprasellar extensions of pituitary adenomas almost fifty years ago,1,2 these giant lesions remain a significant challenge for neurosurgeons.3,4 In spite of the development of transsphenoidal microsurgical techniques,5 some authors still consider the subfrontal transcranial approach preferable for very large tumors.6

The introduction of dopamine agonistic agents (bromocriptine) has opened new perspectives in the management of prolactinomas since significant tumor shrinkage and reduction of serum-prolactin levels can be obtained thus preparing the patient's tissues for an easier surgical procedure.7,8

Although the term macroadroma of the pituitary seems to be generally accepted as defining a tumor measuring 2 cm or more in at least one diameter,9,7 some ambiguity remains as to the definition of giant,4 massive, very large and huge10 adenomas. According to Symon et al.11 giant adenomas should have an extension of more than 40 mm from the midline of the jugum sphenoidale in any direction or within 6 mm of the foramina of Monro. On the other hand, the classification used by Guist and coworkers11,12 implies that the largest suprasellar extensions (grade 3) reach or exceed the foramen of Monro, which is usually located 30 mm above the tuberculum sellae.

Our initial classification of pituitary tumors13 has recently been revised in the light of the newer imaging techniques, CT-Scanning and more particularly MRI-Scanning. Accordingly, microadenomas are now classified as intrasellar tumors of less...
than 10 mm in diameter, macroadenomas with suprasellar extensions (SSE) include grade A (moderate SSE within 10 mm above the jugum sphenoidale, filling the chiasmatic cistern), grade B (large SSE, up to 20 mm, elevating the anterior recess of the third ventricle), grade C (very large SSE, up to 30 mm, filling the anterior third ventricle) and grade D (huge SSE, in excess of 30 mm, above the level of foramina of Monro, or grade C with asymmetrical lateral or multiple expansions). We therefore consider “giant” those adenomas with SSE of types C and D, the critical threshold being a superior margin more than 20 mm above the jugum sphenoidale, regardless of the volume of the intrasellar tumor portion (Figure 1).

**MATERIAL AND METHODS**

During the twenty-five years from 1962 to October 1987, 77 patients with “giant” macroadenomas (SSE types C and D) have been treated surgically at Notre-Dame Hospital, comprising 9.6% of the 800 pituitary adenomas treated by one of us (J.H.) over the same period of time in this hospital, the total series includes 400 additional cases operated on at the Montreal General Hospital (Table 1).

**Pathologic classification**

Non-secreting adenomas were present in 53 cases (69% of 77); these represented 31% of the 168 non-secreting adenomas treated. Patients were males in 29 cases (mean age 47.8 years) and females in 24 cases (mean age 53.8 years).

Secreting adenomas were present in 24 cases (31% of 77) with 16 males (mean age 40.3 years) and 8 females (mean age 28.2 years), accounting for only 3.8% of the total of 632 secreting adenomas treated during the same time-lapse. These 24 adenomas included 13 prolactinomas, 7 human growth hormone (HGH) adenomas, 2 gonadotrophic adenomas, 1 adenocorticotropic adenoma and 1 thyreotropic adenoma.

**Radiologic classification**

With respect to the bony structures, the sella turcica was mildly enlarged and asymmetrical (grade I) in only 1 case, significantly enlarged, including local destruction of the sellar floor (grades II and III) in 62 cases (80.5%) and diffusely destroyed (grade IV, “phantom sella”) in 14 cases (18.1%). The suprasellar extensions were of type C (more than 20 mm above the jugum) in 66 cases (85.7%), 49 of which were non-secreting and 17 secreting adenomas, and SSE type D were found in 11 cases (14.3% of 77) including 7 secreting and 4 non-secreting.

**Visual field disturbances**

Visual field defects were demonstrated preoperatively in 74 of 77 cases (96%) and included complete bitemporal hemianopia in 41 cases (53.2%), bilateral blindness in one patient with acute pituitary apoplexy, bilateral temporal hemianopia and contralateral blindness in 8 cases (10%), unilateral quadranopia and contralateral hemianopia in 8 cases (10%), unilateral hemianopia or quadranopia in 6 cases (7.6%), bitemporal quadranopia in 8 cases (10%). One patient with lateral temporal expansion showed a lateral homonymous hemianopia and 3 patients with presumably postfixed chiasm had normal visual fields.

**Endocrinologic disturbances**

The endocrinologic deficits were classified as major deficits (ACTH-insufficiency), or more minor deficits (FSH, LH, HGH or TSH insufficiency). Adequate endocrinologic preoperative data were available in 70 patients. Twenty-seven patients had major endocrine deficits (38.5%), 20 of which had non-secreting and 7 secreting adenomas. Twenty-three patients (33% from 70) had minor endocrine deficits, and 20 had no deficit at all (28.5%).

Serum prolactin levels were available in 9 from 13 giant prolactinomas and ranged from 900 ng/ml to 30,900 ng/ml, the mean PRL-level being 7,464 ng/ml. The highest PRL-levels were found in invasive tumors of type D.15,16,17,18,19

**Surgical Treatment and Operative Modalities**

The general surgical aspects of the transsphenoidal approach for large pituitary adenomas with suprasellar extension have been described in detail elsewhere.20,21,22
Table 2: Type of 99 Surgical Procedures Performed in 77 Patients with Giant Pituitary Adenomas

<table>
<thead>
<tr>
<th>Procedure Description</th>
<th>Number</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>SINGLE TRANSSPHENOIDAL PROCEDURE</td>
<td>57</td>
<td>57.5%</td>
</tr>
<tr>
<td>(6 pat. previously op. elsewhere)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(T.S.: 3, T.C.: 3)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TWO T.S. (&lt;2 MONTHS)</td>
<td>8</td>
<td>8%</td>
</tr>
<tr>
<td>(two stages: 2, post-op. hemat.: 4)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(CSF-leak: 1, chiasmapexy: 1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TWO T.S. (LONG TERM)</td>
<td>9</td>
<td>9%</td>
</tr>
<tr>
<td>(residual tumor or recurrence)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ONE OR TWO T.S. PLUS CRANIOTOMY</td>
<td>3</td>
<td>3%</td>
</tr>
</tbody>
</table>

Ninety-nine surgical procedures have been carried out in these 77 patients with giant adenomas: 57 patients, 6 of which had previously undergone surgery elsewhere for the same condition, 3 transsphenoidally and 3 transcranially, had a single transsphenoidal procedure performed; 8 patients underwent two transsphenoidal procedures within 2 months from the first operation (2 cases for deliberate two-staged removal, 4 because of post-operative hematoma, 1 for post-operative CSF-leak, and 1 for chiasmapexy related to post-operative empty sella); two patients had two transsphenoidal procedures after prolonged intervals for residual or recurrent tumor; 3 patients had a transcranial approach after one or two transsphenoidal procedures (Table 2).

Operative Technique for Large and Giant Pituitary Adenomas

After exposure of the sellar floor using the conventional transrhinoseptal approach, the sella floor is widely opened to allow a maximal access to the lateral portions of the tumor. After removal of the intrasellar portion of the adenoma, which is usually relatively easy, attention is directed toward the suprasellar extension. Intra-operative pneumoencephalograms using air administered via a lumbar catheter and in some instances before the era of CT-Scanning, immediate preoperative cisternographies were performed in 45 cases to obtain an outline of the

Figure 2 — A. Voluminous pituitary adenoma filling the sphenoid sinus and expanding upwards as outlined by air cisternography via lumbar injection (A). B. Partial debulking of tumor resulting in collapse of capsule as outlined by further air injection. C. Detachment of tumor fragments from underneath the diaphragm. D. Collapsed diaphragm outlined by air above in chiasmatic cistern and air below in the emptied sellar cavity.
suprasellar extension and to follow the downward migration of the diaphragm on radiofluoroscopy (Figure 2). Angled curettes are used to scrape the residual suprasellar tumor remnants adherent to the diaphragm, and Valsalva-maneuvers are performed to help push the SSE down. Intratumoral contrast injections have been used in 3 instances, but this method has been abandoned since two patients demonstrated penetration of contrast into the intracranial subarachnoid space. Before the era of CT-Scanning metallic marker clips were applied in 21 cases on the diaphragma sellae to assess post-operatively the presence of early hematomas or later recurrent tumor growth. In 11 instances, intraoperative CSF-leak required extensive duroplasty using either fascia lata or more recently several layers of folded lypocephalic homologous dura.

Significant amounts of recent liquefied blood were found in 9 tumors (11.6%) and cystic degeneration with yellowish fluid were found in 11 cases (14.2%).

The normal pituitary tissue was identified and preserved in 62 cases (82%). The degree of tumor removal at first operation judged in 75 from 77 cases was: total selective removal in 49 cases (65.3%), total non-selective (normal pituitary not identified) in 4 cases (5.3%), subtotal selective in 13 cases (17.3%) and subtotal non-selective in 9 cases (12%).

Operative complications

From 99 procedures performed in 77 patients, the most significant complication was the development of postoperative hematomas which required operative decompression in 4 cases (4.1%), all of whom died from related causes (one postoperative hematoma with ventricular hemorrhage in a 70-year-old patient with a large invasive prolactinoma, one postoperative hematoma after a second transsphenoidal procedure in a 25-year-old patient with a prolactinoma who experienced cardiac arrest immediately after surgical decompression, one postoperative hematoma in a 72-year-old patient with a large invasive prolactinoma who developed bilateral carotid occlusions and cavernous sinus thrombosis, and one acute myocardial infarction 6 weeks after a postoperative hematoma and meningitis developed in a 52-year-old patient with a non-secreting adenoma). The overall procedure mortality is 4.1% of the 99 procedures and the management mortality is 5.2% (4 of 77 patients). One patient developed an empty sella syndrome with visual field deficit attributed to traction on the optic chiasm which required chiasmopexy: the transphenoidal filling of the sella turcica with muscle and fascia lata resulted in reversal of symptoms. Two patients developed early postoperative Korsakoff’s syndrome attributed to the compressive effect on the hypothalamus and possible intrasellar hemorrhation of the diencephalon. Five patients required permanent therapy because of diabetes insipidus (6.5%).

Operative Results

The overall results of the surgical management were estimated from 48 patients only, since adequate follow-up data were unavailable in 29 cases, mainly from overseas.

Residual or recurrent tumor

Altogether, 19 patients (39.5%) were shown to have residual or recurrent tumor after the first operation on CT-scanning: of these, 12 underwent repeat surgery (9 transphenoidally, and 3 transcranially after one or two repeat transphenoidal procedures).

Radiotherapy was administered in 23 cases (47.9%) after one or two operations.

Bromocriptine was administered postoperatively in 6 of 14 prolactin adenomas because of residual hyperprolactinemia.

Postoperative endocrinologic deficits

Sixty-one patients had adequate early postoperative biological endocrine data. A major deficit (ACTH or more) was present in 26 cases (42.6%), a minor deficit (TSH, HGH, LH, FSH but normal ACTH) was present in 15 cases (24.6%) and no deficit was found in 20 cases (32.7%); in total almost 60% of cases did not get major adrenocortical insufficiency.

Postoperative visual function

Sixty-three patients were evaluated postoperatively as to visual function (3 cases without visual symptoms pre-operatively and 11 cases without postoperative data): of these, 22 patients (31.7%) showed total recovery and 36 (57.1%) showed partial recovery. Five patients remained unchanged (7.9%) and one patient (1.5%) became completely blind.

Discussion and Conclusions

Based on the assessment of a large series of 77 giant adenomas (suprasellar extensions of more than 20 mm above the diaphragm sellae) we conclude that the transsphenoidal route is the preferred surgical approach. It allows an adequate removal of the tumor in most cases, either after one or two procedures. The deliberate two-staged removal has been shown to be particularly useful. Thus, transcranial procedures are rarely needed in our experience.23,24 We have not found any indication for simultaneous combined intra and extracranial approach which seems to be an excessively traumatic procedure with increased risk of infectious complications.

Complications of this transsphenoidal approach include postoperative hematomas which should be dealt with surgically very promptly to avoid hypothalamic damage: these hematomas account for the entire mortality in this series.

Efficient decompression of the optic structures can be obtained by the transsphenoidal approach and usually results in rapid improvement of visual acuity and visual fields.26,27,28

The recent introduction of the dopamine agonist bromocriptine in the preoperative management of large and giant secreting adenomas has shown that significant tumor reduction can be obtained,29,30 but its long-term effect needs to be demonstrated.

The transsphenoidal route allows, even in giant tumors, the identification of the normal pituitary tissue in the majority of the cases, so that major pituitary insufficiency can be avoided in about 60% of the cases.

Acknowledgements

This work was supported by the Pituitary Research Foundation (Canada). It was presented on June 16th 1988 at the XXIIrd Canadian Congress of Neurological Sciences, Quebec City. The manuscript was typed by Miss Helène Lefrançois, scientific secretary.

References


