BACKGROUND AND EPIDEMIOLOGICAL CONSIDERATIONS

Pituitary tumor apoplexy (the second syllable of the suffix “-plexy,” originating from the Greek term meaning to strike or to have a stroke) refers to the abrupt onset of a severe headache frequently coupled with clinical indications of decreased endocrine function. Hemorrhage into or necrosis of a preexisting sellar mass, usually a pituitary macroadenoma, produces an expansion of sellar contents. Compression of adjacent structures elicits the variable expression of symptoms referable to displacement of the optic nerves and chiasm and impingement of the third, fourth, and sixth cranial nerves. Damage to or destruction of the anterior pituitary leads to multiple acute and/or chronic hormone deficiencies in many patients. Medical management may be used in rare cases in which the signs and symptoms are mild and restricted to meningismus or ophthalmoplegia deemed to be stable. In patients with visual or ocular motility lability or an altered level of consciousness, expeditious surgical decompression, accomplished most commonly through a transphenoidal approach, should be performed to save life and vision and to optimize the chance of regaining or maintaining pituitary function.

KEY WORDS • pituitary adenoma • pituitary tumor apoplexy • pituitary apoplexy • pituitary hemorrhage • pituitary infarction • pituitary necrosis

PRESENTATION, DIAGNOSIS, AND PATHOPHYSIOLOGY

Clinical unawareness of the presence of a preexisting syndrome is associated with a pituitary adenoma in the overwhelming number of cases and is linked with healthy glands in only a few isolated instances.

Pituitary adenomas comprise approximately 10% of intracranial tumors;66-72 the reported incidence of apoplexy in these lesions ranges from 0.6% to 27.7%.49 Values quoted at the high end of this range may reflect the inclusion of cases in which blood discovered during surgical or histopathological examination was obtained from specimens of asymptomatic patients.8,24,47,81 Most reports of series indicate that the incidence of apoplexy among pituitary adenomas is 2 to 7% when clinical signs coupled with surgical or histopathological evidence are considered.1,8,21,46,47,52,81 Recurrence following apoplexy is rare and has only been documented in a few cases.1,7,8,15,37,59 The clinical syndrome usually evolves from within a few hours to 3 days.19,42

All types of tumors are at similar risk for developing apoplexy. Nearly 50% of apoplectic events occur in patients who were not previously known to harbor a pituitary lesion.14,24,41,64 Men are affected more commonly than women and in some series the ratio of male to female patients is as divergent as 2:1.8,21,23,36,59,79,83 The age range of affected individuals stretches from the first through the eighth decade, with most cases presenting among individuals in the fifth or sixth decade.1,12,15,19,24,48,50,65

Abbreviations used in this paper: CSF = cerebrospinal fluid; CT = computerized tomography; DI = diabetes insipidus; IHA = inferior hypophysial artery; MR = magnetic resonance; SAH = subarachnoid hemorrhage; SHA = superior hypophysial artery.
pituitary mass often renders apoplexy an overlooked possibility; consequently, the diagnosis is often delayed and the clinical signs are mistaken as produced by a different cause.\textsuperscript{1,13,15,19,21,24,34,36,42,56,60,80} The nonspecific nature and variability of signs and symptoms further confound expeditious establishment of an appropriate diagnosis.\textsuperscript{1, 24,25,40,48}

Headache is the most commonly reported symptom and is present in up to 100\% of affected patients; it is almost universally described as severe and abrupt, and located retroorbitally or bifrontally,\textsuperscript{8,9,15,21,24,35,59,65,80} A deterioration in visual acuity leading to blindness, visual field defects, and ophthalmoplegia are frequently associated with the ictus, as are nausea, vomiting, lethargy, meningismus, and a decreased level of consciousness.\textsuperscript{3,10,19,36,38,76,81} Altered facial sensation or facial pain, hemiparesis, seizures, or miosis and ptosis associated with Horner syndrome are less frequently encountered.\textsuperscript{8,21,24,26,41,42,47,49,58,62,63,65,67,74} Several authors describe compression of the third nerve against the interclinoid ligament as reason for the familiar inclusion of pupillary dilation, diabetic third nerve palsies, and the clinical signs are mistaken as produced by a different cause.\textsuperscript{1,13,15,19,21,24,34,39,42,53,60,80} The nonspecific nature and variability of signs and symptoms further confound expeditious establishment of an appropriate diagnosis.\textsuperscript{1, 24,25,40,48}

Distension of the optic nerves and chiasm account for restriction of visual fields, specifically bitemporal hemianopia.\textsuperscript{60} Lateral extension of hemorrhage and necrosis leads to defects in the third, fourth, fifth (first and second divisions), and sixth cranial nerves.\textsuperscript{60} The third cranial nerve is most commonly affected in isolation or combined with other nerve deficits.\textsuperscript{21,26,41,42,47,49,58,62,63,65,67,74} Several authors describe compression of the third nerve against the interclinoid ligament as reason for the familiar inclusion of this nerve in the grouping of symptoms or in a singular presentation.\textsuperscript{35,58,67,70} The position of the third nerve parallel to the pituitary gland in the lateral wall of the sinus may make this nerve more directly susceptible to pressure transmitted via rapid lateral expansion.\textsuperscript{58,68}

Medially located within the sinus, the sixth cranial nerve is the next most frequently affected.\textsuperscript{47,56,66} The fourth cranial nerve is the least commonly involved in plural defects and it is rare for this nerve to be affected separately. Facial pain, altered facial sensation, hypesthesia, and/or loss of corneal reflex are occasionally mentioned as associated symptoms.\textsuperscript{13,20,58,65}

Endocrine abnormalities associated with hypopituitarism, pressure transmitted to the brainstem, or hypothalamic compression may account for a diminished level of consciousness.\textsuperscript{21,70,80} Vasospasm due to SAH following the apoplectic event may provoke hemispheric signs.\textsuperscript{27,25,65,80} Acute diminishment of the caliber of the intracavernous carotid artery and decreased blood flow secondary to constriction from pressure of edematous, hemorrhagic material or from being wedged against the anterior clinoid process have also been postulated as accounting for episodes of lethargy or hemiplegia.\textsuperscript{9,25,58,60,64,65,79,80} Cortical irritation from extension of hemorrhage into the brain par enchyma may elicit seizures.\textsuperscript{59} Encroachment on the sympathetic plexus can lead to the Horner sign. Attenuation of venous channels draining into the sinus may trigger proptosis and swelling of the eyelid.

Signs of mass effect on the optic nerves and chiasm are commonly noted at presentation in patients with pituitary adenosomas (243 of 247 patients in a series conducted by Cushing),\textsuperscript{67} yet only 5 to 15\% of patients with adenosomas are noted to have optomotor defects.\textsuperscript{3,5,20,67,68} In contrast, although chiasmal abnormalities continue to be a prominent sign in patients with necrosis or hemorrhage, the incidence of optomotor palsies increases to 60 to 100\% in these patients.\textsuperscript{37,70,80}

The reason for the disparity in the frequency or pattern of occurrence between chiasmal and optokinetic signs associated with adenosomas, as opposed to pituitary tumor apoplexy, has been attributed to the vulnerability of the optic nerves and chiasm to even gradual stretching, which occurs with an adrenoma, compared with the relative tolerance and greater mobility of the third, fourth, and sixth cranial nerves, which are comparatively undisturbed by the more insidious effects of an adenoma. These more durable nerves frequently do not show signs of compromise unless or until they are abruptly compressed.\textsuperscript{18,34,67,80}

The diagnosis of pituitary tumor apoplexy is frequently complicated by signs and symptoms that may resemble those of other intracranial pathological entities (Table 2). Headache, photophobia, stiff neck, and oculomotor palsy may mimic a ruptured intracranial aneurysm,\textsuperscript{24} whereas fever, meningismus, and lethargy can appear to indicate bacterial or viral meningitis.\textsuperscript{14,20,24,66} Periorbital edema from compression of the venous plexus contained in the cavernous sinus paired with defects resulting from encroachment on adjacent cranial nerves can emulate cavernous sinus thrombosis.\textsuperscript{49,70} Unilateral ptosis and mydriasis, in addition to restriction of adduction, elevation, and downward gaze, may raise consideration of the presence of a posterior communicating artery aneurysm or, if lacking the inclusion of pupillary dilation, diabetic third nerve palsy.\textsuperscript{52}

Hypopituitarism, with variably decreased levels of all or multiple pituitary hormones, is evident on presentation in the majority of patients who present with apoplexy.\textsuperscript{3,19,31,59,79,80} This attenuation in hormone levels may

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<th>TABLE 1</th>
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<td><strong>Signs and symptoms of pituitary tumor apoplexy</strong></td>
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<tr>
<td>headache (frontal or retroorbital)</td>
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<tr>
<td>restriction of visual fields</td>
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<tr>
<td>decrease in visual acuity</td>
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<tr>
<td>ophthalmoplegia</td>
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<tr>
<td>nausea</td>
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<td>vomiting</td>
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<td>vertigo</td>
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<tr>
<td>meningismus</td>
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<tr>
<td>decreased level of consciousness</td>
</tr>
<tr>
<td>facial pain or altered or impaired facial sensation</td>
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<tr>
<td>epilepsy</td>
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<tr>
<td>fever</td>
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<tr>
<td>hemiparesis</td>
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<td>Horner syndrome</td>
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Differential diagnosis of pituitary tumor apoplexy

<table>
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<th>Diagnosis</th>
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<tr>
<td>SAH from aneurysmal rupture</td>
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<tr>
<td>viral or bacterial meningitis</td>
</tr>
<tr>
<td>temporal arteritis</td>
</tr>
<tr>
<td>migraine</td>
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<tr>
<td>spontaneous hemorrhage (hypertensive</td>
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<tr>
<td>or amyloid bleeding or from a</td>
</tr>
<tr>
<td>metastatic tumor or primary lesion)</td>
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<tr>
<td>diabetic oculomotor palsy</td>
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<tr>
<td>optic neuritis</td>
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<tr>
<td>vertebrobasilar insufficiency</td>
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<tr>
<td>hypertensive encephalopathy</td>
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<tr>
<td>transtentorial herniation</td>
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<tr>
<td>cavernous sinus thrombosis</td>
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<td>carotid–cavernous fistula</td>
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Result from increased intrasellar pressure, preexisting deficits from an adenoma, or destruction of the gland. Elevated intrasellar pressure can compress the pituitary, instigating dysfunction and producing subsequent endocrine abnormalities. Lack of recognition of endocrine perturbations can significantly increase the incidence and extent of morbidity.

Diabetes insipidus is an uncommon sequela to pituitary apoplexy, this may reflect the differential vascularization between the pituitary gland and the hypothalamus. The anterior communicating artery vascularizes the preoptic nucleus of the hypothalamus, whereas the A1 segment of the anterior cerebral artery supplies the anterior hypothalamic and the posterior communicating artery sustains the posterior region. Conversely, the SHA supplies the anterior lobe of the pituitary gland and the infundibulum as the IHA furnishes blood to the posterior lobe. The bilateral SHAs divide into the hypophysial portal system at the infundibulum and are joined by the ipsilateral and contralateral branches of the IHA. The hypophysial portal system supplies the anterior lobe of the pituitary.

Diabetes insipidus has been postulated to occur as a result of impingement on the intracavernous portion of the IHA, causing diminished perfusion to the posterior lobe. Alternatively, kinking or pressure on the infundibulum by edematous, hemorrhagic material, which impedes transit of antidiuretic hormone from the preoptic and paraventricular nuclei of the hypothalamus, may prove to be the originating cause of DI.

Computed tomography scans may reveal the hyperdensity of acute hemorrhage if obtained within 3 or 4 days after the event or demonstrate the mixed density of acute blood and hypodense necrotic tissue (Fig. 1). Subarachnoid hemorrhage may be evident if blood has invaded the basal cisterns. The greater sensitivity, precision, and tissue definition of MR imaging usually reveal the heterogeneous intensity of hemorrhage, edematous pituitary gland, and necrotic tumor, leading more readily to a diagnosis. The multiplanar capability of this type of imaging allows meticulous characterization of the extent of necrosis and/or hemorrhage in addition to an evaluation of the relationship of hemorrhage and denuded tissue to neurovascular structures (Fig. 2).

In their study of 37 patients with a diagnosis of pituitary tumor apoplexy, Bills and colleagues reported a 46% detection rate of apoplexy on CT scans compared with hemorrhage or infarction ascertained using MR images in all patients thus studied. Randeva, et al., reported that CT scans revealed 21% of the cases of pituitary tumor apoplexy in their series of 35 patients, whereas MR images demonstrated 100% of the cases. Although MR imaging has emerged as the most exact and definitive imaging modality in this circumstance, it remains important to recognize apoplexy on CT scans if hemorrhage or necrosis is apparent, as CT scans are often obtained initially when patients present to emergency departments with nonspecific neurological complaints. Considering the low sensitivity of CT scanning in these cases, it is important not to ignore or reject the possibility of pituitary tumor apoplexy if clinical assessment indicates this diagnosis.

Cerebrospinal fluid analysis is rarely helpful in establishing a diagnosis because the fluid may be clear if blood has not broached the subarachnoid space. Alternatively, CSF may be bloody or xanthochromic, depending on the time interval from the ictus. Yellow-tinged CSF will be present in 90% of specimens at > 12 hours from onset. Increased protein levels and pleocytosis noted in the CSF of some patients are similarly nondiagnostic.

**PRECIPITATING FACTORS AND TUMOR CHARACTERISTICS**

Predisposing factors have been identified in approximately 50% of cases of pituitary tumor apoplexy. Head trauma, bromocriptine administration or withdrawal, anticoagulation, pregnancy, cardiac bypass, bowel or other general surgery, atherosclerosis, diabetic ketoacidosis, estrogen therapy, and radio-
tion therapy, hypertension, and hypotension have been cited as inciting apoplectic episodes. Pituitary apoplexy has also been instigated by dynamic testing with thyrotropin-releasing hormone, gonadotropin-releasing hormone, and corticotropin-releasing factor. Biousse and associates reported increased demand on the pituitary coupled with blood pressure fluctuations as an inciting factor in the generation of pituitary tumor apoplexy. Fluctuations in blood pressure associated with cardiac bypass and surgical interventions may provoke necrosis. Fragmentation of atherosclerotic plaques with lodging of particles within the pituitary vasculature may potentiate gland infarction. The association of pituitary tumor apoplexy with surgery, childbirth, and infection may reflect the greater activity of the pituitary gland in response to the increased need for stress hormones in these circumstances. Additionally, estrogen therapy and pregnancy stimulate and enhance the demand on the pituitary. Sneezing, coughing, and severe acidosis elicit alterations in blood flow or an abnormal osmotic composition of blood delivered to the pituitary. Administration of releasing hormones during stimulation tests may increase intratumoral blood flow and provoke bleeding in friable, immature vessels.

Numerous investigators have reported a greater propensity toward necrosis and hemorrhage in larger tumors due to the discrepancy between the rate of neoplastic progression and the availability of circulatory input. Nevertheless, small tumors also hemorrhage. The inherent fragility of the vascular infrastructure of tumors may provide an explanation for hemorrhage in these instances. Predisposing factors may also prove more important in the genesis of necrosis and/or hemorrhage in smaller lesions. Ebersold, et al., have emphasized vascular stasis and subsequent thrombosis as the likely mechanism eliciting necrosis and subsequent bleeding.

Impingement of the SHA from compression of the pituitary stalk and superior hypophysial vessels against a dilated or more closely maintained diaphragma sella is cited as placing pituitary adenomas at high risk for ischemia and ensuing hemorrhage. Yet it has been suggested that tumor vessels originate from the IHA. Attenuation in the caliber of the lumen of feeding vessels from the SHA would thus place the basis of the apoplectic event within the anterior lobe of the pituitary, rather than within the tumor itself.

There is no preponderance of tumor type in the genesis or evolution of pituitary tumor apoplexy. Although adenomas are most commonly implicated in apoplectic events, necrosis and/or hemorrhage has been noted to occur in Rathke cleft cysts, in nontumorous pituitary glands, and in association with an empty sella turcica. Although clinically evident apoplexy has commonly been deemed to occur in less than 10% of pituitary adenomas, the incidence of subclinical hemorrhage has been identified in as many as 28% of histologically scrutinized tumors.

In their retrospective review of 560 cases of pituitary adenoma, Wakai, et al., found 93 tumors (16.6%) with evidence of degenerated blood or intratumoral hematoma, yet only 38 patients (6.8%) with pituitary lesions experienced an apoplectic event. Similarly, in his examination of specimens obtained from 280 patients, Deb identified 48 patients (17.1%) in whom hemorrhage was discovered at surgery, whereas only 15 patients (5.4%) demonstrated clinically evident symptoms or signs.

Comprising only 10% of intracranial neoplasms, pituitary adenomas constitute a greater proportion of tumor-related hemorrhages. In considering intracranial hemorrhage derived from tumors, Glass and Abbott reported that 25% of intracranial hemorrhages originated secondary to adenomas compared with 50% instigated by gliomas, which are five times more common. Based on their...
Pituitary tumor apoplexy

examination of 1861 hemorrhagic intracranial tumors, Wakai, et al., 52 discovered that adenomas are 5.4 times more likely to bleed.

TREATMENT

The pituitary gland remains capable of secreting adequate amounts of hormones when as little as 10% of residual tissue remains; however, a dearth or absence of sufficient hormone can lead to adrenal crisis. 5, 15, 19, 25, 43, 56, 58, 60. The prompt recognition of apoplexy permits the immediate initiation of steroid replacement. 5, 53 One hundred milligrams of hydrocortisone should be administered intravenously on establishment of the correct diagnosis. 40, 70

The definitive treatment for pituitary tumor apoplexy is surgery for decompression of constricted cavernous and/or suprasellar structures, especially in cases in which visual acuity or field defects, decreased level of consciousness, or progressive deterioration of visual or oculomotor abilities are present. 1,5, 15, 19, 25, 39, 43, 56, 60. The definitive treatment for pituitary tumor apoplexy is surgery for decompression of constricted cavernous and/or suprasellar structures, especially in cases in which visual acuity or field defects, decreased level of consciousness, or progressive deterioration of visual or oculomotor abilities are present. 1, 5, 15, 19, 25, 39, 43, 56, 60. Use of the intracranial route to reach suprasellar sites of hemorrhagic and necrotic material places the delicate perforating vessels extending from the anterior communicating artery and the SHA at risk.

A significant visual compromise, diminished level of consciousness, and declining visual status are clear indications for operative intervention. 19, 56, 79 Surgery offers a means for direct and immediate decompression and optimizes the chance for restitution of endocrine function and recovery of neural elements. 5, 59 In addition, surgery allows tumor resection, decreasing the chance of continued neoplastic viability. 53 Extensive intracavernous extension or invasion limit the opportunity for complete tumor removal.

In a minority of published papers, cases of isolated and stable meningismus and/or ophthalmoplegia have been shown to be managed medically, because meningismus will resolve once blood no longer irritates the dura mater and ophthalmoplegia can resolve spontaneously. 22, 43, 56, 60, 76 Medical management includes close monitoring of endocrine, neurological, and ophthalmological function, hormone administration, and support with intravenous fluids and electrolytes.

In our own experience with 15 patients who presented with signs and symptoms indicative of pituitary tumor apoplexy, surgical decompression within 2.2 days after symptom onset led to the postoperative return of endocrine function in 11 patients. This recovery leads us to believe that the source of pituitary compromise in the wake of apoplexy, in some cases, is compression rather than destruction, and that in certain patients there remains the chance for improvement of function. 3, 5 Of course, if the apoplectic event involves overt gland necrosis or ischemia from attenuation of the caliber of the SHA, return of function would be less likely. Still, in light of our data, the potential for restitution of pituitary activity is another reason for advocating timely surgical intervention.

The unpredictable clinical course of pituitary tumor apoplexy provides a strong rationale for prompt decompression. 3, 5, 15, 19, 22, 43, 56, 60. The capricious nature of pituitary tumor apoplexy makes it difficult or impossible to foretell precisely in which patients the physical conditions will remain stable and in which it will deteriorate. It is impossible comfortably to exclude patients from surgical intervention by speculating that their conditions will remain stable.

PATIENT OUTCOMES

Among visual perturbations, abnormalities in optic motility generally rebound quickly once the third, fourth, and sixth cranial nerves are relieved of compression. 3, 5, 15, 20, 21, 43, 47, 56, 59, 60 Congruous eye movements are more readily restored than the chiasmal and optic nerve defects of decreased visual acuity and restriction of visual fields. 15, 23, 56, 70, 80 In their series of 37 patients, Bills and colleagues 8 reported that ocular paresis was resolved in 100%, visual field deficits in 95%, and deficits in acuity in 88% of patients. This pattern of resolution or improvement is reflected in numerous other studies. 59, 60 Similarly, patients in whom decompression is accomplished expeditiously generally experience greater improvements in both cranial nerve and chiasmal deficits. 14, 58, 59, 68 Bills and colleagues also described the largest recovery in patients who underwent surgery within 1 week after inception of signs and symptoms.

More than half of affected patients require hormone replacement following an episode of pituitary tumor apoplexy. 3, 8, 15, 22, 43, 47, 56, 59, 79 In their study of 37 patients, Bills and colleagues reported that administration of steroid, thyroxine, and testosterone was essential postoperatively in 82, 89, and 64% of patients, respectively. From their review of the literature, Veldhuis and Hammond 79 ascertained that, following an episode of pituitary tumor apoplexy, 88% of patients lacked sufficient growth hormone, 76% adequate amounts of lutenizing hormone, 67% sufficient prolactin, and 66% enough adrenocorticotropic hormone, and that 33% of patients experienced deficiencies in estradiol. Diabetes insipidus occurred in only 2 to 3% of patients examined for this disorder. 79

Despite these outcomes, as mentioned previously, by performing surgical decompression within 3 days after symptom onset, we were able to produce a return of endocrine function in 11 of 15 patients. Restitution of pituitary function was recognized within 3 days after surgery by the return of endogenous adrenocorticotropic hormone secretion.

CONCLUSIONS

Pituitary tumor apoplexy is most commonly a discrete event marked by onset of the constellation of severe, sudden headache, restriction of visual fields, and paresis of ocular muscles. Signs and symptoms generally arise consequent to the following: 1) endocrine abnormalities from acute pituitary dysfunction; 2) subarachnoid extravasation of blood and dural irritation; and 3) cranial nerve or hemi-
spheric compression from lateral or superior extension of necrotic and/or hemorrhagic material. Clinical indications associated with neuroimaging and surgical or pathological findings constitute the syndrome of pituitary apoplexy. The variable presentation of nonspecific signs and symptoms can mimic numerous other causes, including meningitis, midbrain infarction, and cavernous sinus thrombosis; the dramatic onset may resemble that of a ruptured intracranial aneurysm.

Pituitary macroadenomas are most commonly implicated as the entity underlying the ictus. Although pituitary tumor apoplexy most commonly occurs spontaneously, numerous precipitating associations have been described, including bromocriptine therapy, pregnancy, and hypertension. Abrupt expansion of a pituitary lesion causes inexcipient pressure on intrasellar and parasellar structures, producing clinical signs referable to the size and contour of necrotic and/or hemorrhagic material. Magnetic resonance imaging most accurately identifies this pathological condition, precisely defines its extent, and establishes its proximity to neurovascular elements.

Swift recognition of an accurate diagnosis allows prompt steroid administration and surgical decompression, which is most commonly accomplished via the transsphenoidal approach. Although hypopituitarism is a prominent finding preoperatively among patients following an apoplectic episode, an improvement in endocrine function has been attained in some patients. Long-term hormone replacement in some axes is required in a major proportion of cases in which surgery is performed within 1 week after clinical onset.

References

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