Pituitary macroadenomas are benign tumors. The most common subtype is clinically non-functioning pituitary adenomas, which are defined by an absence of clinical evidence of hormonal hypersecretion. Patients with pituitary macroadenomas may present to an eye clinic with symptoms related to mass effect on surrounding structures, which can include headache, bitemporal visual defects, and ocular motor deficits (see Key Ocular Findings in Pituitary Macroadenoma).

If you suspect that a patient has a nonfunctioning pituitary adenoma, the following steps are warranted: visual acuity testing, color plates, a cranial nerve examination, an assessment for a relative afferent pupillary defect, an evaluation of the optic discs’ appearance, visual field testing, and OCT imaging. In addition to narrowing the differential diagnosis, the information gathered by these assessments can guide treatment strategies, predict prognostic factors for recovery, inform providers’ and patients’ expectations, and document postoperative recovery after resection or radiotherapy.

### Key Ocular Findings in Pituitary Macroadenoma

- **Visual field defects**
  - Superior progressing to complete bitemporal hemianopsia
  - Junctional scotoma (postfixed chiasm or anterior tumor spread)
  - Rarely, macular bitemporal hemianopsia or homonymous visual field loss (prefixed chiasm or posterior tumor spread)
- **Pituitary apoplexy**
- **Diplopia**
  - Hemifield slide
  - Cranial nerve palsies, especially multiple
- **See-saw nystagmus**
  - Thinning of the binasal macular ganglion cell layer and temporal retinal nerve fiber layer on OCT
  - Temporal or bow-tie atrophy of the optic disc on examination

### Visual Field Defects

Visual field testing is central to the workup of pituitary adenomas. In a review of eight studies, visual field disturbances were the most common symptoms at presentation, found in approximately 46% of all patients. It is important to note, however, that slow tumor growth may delay patients’ presentation to an eye clinic, and some asymptomatic cases are detected incidentally on glaucoma screening.

Bitemporal hemianopsia can occur as a result of suprasellar growth of the pituitary lesion and direct inferior compression of the optic chiasm first.
where the axons that give rise to the superior temporal visual field cross (see Differential Diagnosis of Bitemporal Hemianopsia). Tumors may grow asymmetrically, and different configurations of the optic chiasm relative to the pituitary gland can contribute to the observed pattern of presentation. If the optic chiasm sits anterior to the pituitary gland within the tuberculum sellae, referred to as a prefixed chiasm, a pituitary adenoma is more likely to compress the posterior chiasm and optic tracts to produce a macular bitemporal hemianopsia or homonymous visual field loss. Conversely, postfixed chiasms overlying the dorsum sellae may produce patterns of visual field loss related to the optic nerve or a junctional scotoma due to compression at the confluence of the optic nerve and chiasm.

### Differential Diagnosis of Bitemporal Hemianopsia

- **Extrinsic = compression**
  - Pituitary adenoma
  - Parasellar meningioma
  - Craniopharyngioma
  - Parasellar internal carotid artery aneurysm
- **Intrinsic = chiasmal in origin**
  - Glioma/astrocytoma
  - Demyelinating lesion
  - Rarely, sarcoïd or inflammatory lesion
- **Not chiasmal = optic nerve anomalies**
  - (generally do not respect vertical midlines well)
  - Tilted nerve
  - Atypical glaucomatous optic neuropathy
- **Other**
  - Trauma with chiasmal contusion
  - Hydrocephalus

### Oculomotor Deficits

Cranial nerve (CN) palsies occur as a result of compression caused by the direct extension of a tumor out of the sella and into the adjacent cavernous sinuses. Multiple palsies of CN III, IV, V1, V2, and/or VI localize to this region. A patient with any of several CN palsies will complain of diplopia and, in severe cases, may present acutely with headache as a result of pituitary apoplexy. Acute-onset diplopia and headache in this scenario should prompt immediate neuroimaging (see Pituitary Apoplexy). Pituitary adenomas may also cause diplopia due to decompensation of a preexisting phoria by the lack of crossover of the remaining visual field in the setting of complete hemianopsia (see Hemifield Slide).

See-saw nystagmus (SSN) is a rare ocular motor syndrome characterized by a cycle of elevation and intorsion of one eye with synchronous depression and extorsion of the other, a scenario that is inverted in the next half-cycle. Two patterns of SSN exist. The pendular-type SSN is more typical of parasellar masses, and it is differentiated from jerk SSN by its smooth and rolling movements. Although the underlying pathophysiology of pendular SSN remains a subject of debate, a two-hit hypothesis has been proposed. First, compression of the chiasm may lead to a retinal calibration error of the vestibulo-ocular reflex, causing a disturbance in the roll plane (see Hemifield Slide). Second, compression of the interstitial nucleus of Cajal or the rostral interstitial nucleus of the medial longitudinal fasciculus can lead to nystagmus in the pitch and roll planes.

### Ocular Imaging in Diagnosis and Management

OCT imaging is useful for visualizing the retinal nerve fiber layer (RNFL) and macular ganglion cell layer (GCL) in patients with visual field loss from chiasmal compression (Figure 1). Pituitary adenomas cause compression of the axons from the nasal retinal fibers, which correspond to the temporal visual fields. This damage is reflected after weeks to months in binasal macular ganglion cell atrophy and...
PITUITARY APOPLEXY

Pituitary apoplexy is a rare vascular emergency caused by infarction or hemorrhage of the pituitary gland. Although eye care providers are unlikely to see a patient with apoplexy in the clinic, it is important to educate patients with a history of pituitary adenoma about the following red flags:

- Sudden changes in vision, especially peripheral visual field loss or diplopia
- Abrupt onset of severe thunderclap headache, usually retroorbital
- Altered mental status or personality changes from acute changes in cerebral blood flow
- Nausea, vomiting, or loss of appetite
- Low blood pressure
- Hormone insufficiency

Recent studies have shown that OCT imaging can provide useful guidance on treatment strategy. Because extensive RNFL thinning and macular GCL thinning indicate retrograde atrophy of fibers from chiasmal compression, patients with intact RNFL and GCL on OCT at diagnosis may have a greater potential for recovering optic nerve function and visual fields after surgical intervention.

MANAGEMENT

MRI with and without contrast of the brain and/or orbits with attention to the sella is the recommended medium for diagnosis (Figure 2). A patient with a confirmed pituitary macroadenoma should be referred to both neurosurgery and endocrinology. Asymptomatic patients with incidental macroadenomas should receive continued ophthalmologic surveillance because any new visual symptoms strongly indicate a need for surgical intervention.

HEMIFIELD SLIDE

Diplopia without oculomotor involvement, referred to as hemifield slide, can occur with bitemporal hemianopsia in patients who are experiencing compression of the optic chiasm. This syndrome arises when the eyes no longer share an area of visual field, leading to a loss of binocular fusion in a patient with preexisting phoria. When the eyes move apart from one another in decompensated exophoria, the patient may perceive the nasal hemifields overlapping centrally as a central strip. In esophoria, intermittent crossing of the eyes may be observed as a central strip of diplopia, causing visual confusion. In decompensated esophoria, the two nasal visual fields may appear to separate, leaving a linear central scotoma. Similarly, a vertical phoria may produce a superior or inferior segment of missing or jumbled vision (Figure).

Figure. Hemifield slide can arise from a loss of motor alignment in patients with pituitary adenoma by interfering with the binocular visual field. Exotropia creates an overlapping strip of nasal visual field that causes visual confusion, esotropia creates a central strip scotoma, and hypertropia results in a superior or inferior segment scotoma and central visual field misalignment.
established guidelines for the length of ophthalmologic follow-up, lesion size and a progression of visual field defects should inform decisions regarding timeline.

If visual impairment secondary to pituitary macroadenoma is confirmed or progressive, first-line treatment is surgical resection with endoscopic or microscopic transphenoidal surgery. Bromocriptine can be used for prolactin-secreting adenomas (usually small). Large masses may require superior craniotomy approaches, whereas gamma knife radiotherapy is reserved for large tumor remnants and the management of regrowth after surgery. In a review of 19 studies that describe visual outcomes after resection, approximately 40% of patients recovered completely, 80% showed an overall improvement, and 2% reported further deterioration in visual function. Preoperative visual field assessments are essential to estimate visual outcomes after resection and to set both patients’ and providers’ expectations.

**CONCLUSION**

By understanding the clinical signs associated with pituitary adenomas, you can ensure that patients with associated visual defects receive an early diagnosis and a thorough preoperative assessment. The information gathered can inform treatment plans and better predict ophthalmologic outcomes after treatment.


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