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MP2006 - Pituitary Adenoma Order code: 4003.MP2006



Cena bez DPH Price with VAT 907,00 Eur 1.097,47 Eur

Parameters Quantitative unit

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Clinical History

A 29-year old male presented with a 22 month history of headaches and blurred vision. Examination revealed a bi-temporal hemianopia and a left 6th nerve palsy. Skull X-ray showed erosion of most of the sphenoid body with some dorsum sellae and anterior clinoid process intact. Carotid angiography showed upward and lateral displacement of the anterior and middle cerebral arteries. Pneumoencephalography (a common imaging procedure used until the 1970s in which CSF is drained and replaced by air, oxygen or helium that acted as a contrast medium in X-ray examinations) showed upward displacement of the lateral and third ventricles from below. A craniotomy was performed but the patient died immediately afterwards.

Pathology

The brain specimen is sliced in the sagittal plane to the right of the falx cerebri, which remains in-situ. The pituitary gland has been completely replaced by a round tumour 4cm in maximum diameter. The tumour cut surface is pale brown and homogenous (except for an area of haemorrhage superiorly, likely caused by surgical trauma). The tumour has resulted in upward displacement of the midbrain. Tumour erosion has destroyed the sphenoid bone; thus, the sella turcica is enlarged (arrow). The optic chiasma is compressed by the tumour. Histologically, this tumour was a chromophobe adenoma arising from the anterior pituitary.

Further Information

This specimen is from an old case and the investigations used would now be considered antiquated. Modern investigation would include an initial brain CT followed by an MRI of the brain to further visualise the pituitary lesion prior to any surgical intervention.

Pituitary adenomas are the most common pituitary tumour and are most commonly found in adults with peak incidence between 35-60 years. Primary carcinoma of the pituitary is very rare, and the pituitary is an uncommon site for metastases. Clinical manifestations of pituitary adenomas are related to local mass effect and tumour function. Local effects include increasing intracranial pressure (headache, nausea and vomiting), sellar expansion, bony erosion and compression of decussating nerve fibres in the optic chiasma, causing bitemporal hemianopia.

Pituitary adenomas can be functioning (i.e. associated with hormone excess) or non-functioning (i.e. without clinical symptoms of hormone excess). About 75% of adenomas are functional: usually secreting prolactin, growth hormone or ACTH. Secretion of TSH, LH and FSH from pituitary adenomas is very rare. Some adenomas can secrete two hormones: growth hormone and prolactin being the most common combination. Non-functional pituitary adenomas come to clinical attention later than those associated with endocrine abnormalities, and they may lead to hypopituitarism due to compression atrophy of the surrounding normal gland.