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Karavitaki, Niki

DOI: 10.1016/j.mpmed.2017.05.008

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Document Version Peer reviewed version

Citation for published version (Harvard): Karavitaki, N 2017, 'Diagnosis of Pituitary Disease', *Medicine*, vol. 45, no. 8, pp. 464-469. https://doi.org/10.1016/j.mpmed.2017.05.008

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Diagnosis of pituitary disease

Athanasios Fountas Niki Karavitaki

Athanasios Fountas MD, MSc is an Honorary Research Associate at the College of Medical and Dental Sciences. - of University of Birmingham, UK. Competing interests: none declared.

Niki Karavitaki MD, MSc, PhD, FRCP is a Senior Clinical Lecturer in Endocrinology at the College of Medical and Dental Sciences, of University of Birmingham and an Honorary Consultant Endocrinologist at Queen Elizabeth Hospital, UK. Niki-Karavitaki has received fees for speaking, funds for research and fees for consulting from Novartis, Pfizer, Merck Serono and NovoNordisk in the last 5 years.

Abstract

The prevalence of pituitary disease is increasing mainly due to thbecause of advances in modern imaging techniques and to the an increased awareness amongst the medical community. Pituitary tumours constitute 10–15% of all diagnosed intracranial neoplasms, and their clinical manifestations result from local mass effects (mostly neurological, visual, hypopituitarism) and/or hypersecretion. Pituitary adenomas are the most common pituitary tumours and <u>are</u>clinically <u>are</u>classified as functioning or non-functioning. The majority isMost are sporadic, but₇ in rare cases, they can be related with to hereditary syndromes. Other lesions involving the (para)sellar region include inflammatory and infiltrative diseases, cysts, primary or metastatic neoplasms, abscesses and internal carotid artery aneurysms. The clinical manifestations of hypopituitarism depend mainly on the type, number and severity of hormonal deficits. The Establishingment of the diagnosis requires hormonal measurements (basal or after dynamic tests), and the management includes relevant hormonal replacement and life-long monitoring.

Keywords

Craniopharyngioma; hypophysitis; hypopituitarism; pituitary adenoma; pituitary apoplexy; pituitary incidentaloma; pituitary stalk lesions; Rathke's cleft cyst

Key points

- Pituitary adenomas comprise the majority of pituitary tumours, and can be functioning or non-functioning.
- Clinical features of pituitary masses maycan result from local mass effects and/or hypersecretion.
- Sellar or parasellar masses mainly include adenomatous and non-adenomatous tumours, inflammatory and infiltrative diseases, cysts, primary or metastatic malignancies, pituitary infections and internal carotid artery aneurysms-
- All patients with pituitary masses should undergo testing for hypopituitarism, and for hormonal hypersection (in cases of pituitary adenomas), radiological assessment and neuro-ophthalmological evaluation.

Introduction-(A)

The pituitary gland, or 'hypophysis cerebri', is considered to be the 'master gland' of the endocrine system, integrating, together with the hypothalamus, hormonal signals that control a plethora of endocrine and metabolic functions.

The prevalence of pituitary disease is-has increaseding in over the last 10 years duetobecause of the advances in modern imaging techniques and hormonal measurements, as well as due toan the increased awareness and rate of suspicion rate for these disorders amongston the part of the medical community. Fortunately, the improvements in pituitary Comment [CMW1]: AQ: should this be 'hypersecretion'? Comment [NK2]: Yes, please change it

to hypersecretion

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surgery and radiotherapy techniques, combined with the development of medical treatments for pituitary tumours and the advances in pituitary hormone replacement therapy, have led to more optimal outcomes.

Pituitary anatomy (A)

The pituitary gland consists of the an_anterior lobe (adenohypophysis), the posterior lobe (neurohypophysis), and the vestigial intermediate lobe. It lies at the base of the brain in the sella turcica, within the sphenoid bone, and is overlain by the dural sellar diaphragm through which the pituitary stalk connects to the median eminence of the hypothalamus. The sellar diaphragm also protects the pituitary from compression by the cerebrospinal fluid (CSF). In both sides of the sella turcica, and lateral and superior to the sphenoid sinus, are the cavernous sinuses, in which; the cavernous segments of the internal carotid arteries and the cranial nerves III, IV_7 and VI are located in these. The optic chiasm is anterior to the pituitary stalk, and typically sits 5--10 mm above the sellar diaphragm.

The pituitary measures approximately 13 mm transversely, 9 mm antero-posteriorly, and 6–9 mm vertically, and in adults it-weighs around 600 mg (range 400–900 mg). However, the size and the volume of the gland change in different situations; the pituitary increases during pregnancy to almost twice its normal size, whilst whereas it decreases in older people.

| The posterior pituitary lobe is comprisesd of the distal axons of the magnocellular neurosecretory cells extending from the supraoptic and paraventricular nuclei of the hypothalamus. These cells synthesize the neurohypophysial neurohypophyseal hormones oxytocin and vasopressin and store them into neurosecretory granules at their axon terminals; from where they are released from here into the neurohypohyseal capillaries and the systemic circulation. Blood supply(B) The anterior pituitary receives most of its blood supply from the hypothalamo-hypophyseal arteries. Through this system, which originates from the capillary plexus of the median eminence and superior stalk, derived from the superior hypophyseal arteries. Through this system, the hypophysiotrophic hormones are delivered to the hormone-producing cells of the adenohypophysis. The remainder of the blood supply is through via the pituitary capsular vessels, which that also originate from the superior hypophyseal arteries. The posterior lobe | | obe (adenohypophysis) (B) | Formatted: Font: Bold, Not Italic |
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| and the stalk are directly supplied with blood from the hypophyseal arteries. | | alk are directly supplied with blood from the hypophyseal arteries. | |
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-The venous drainage from both lobes is through the cavernous sinuses into the petrosal sinuses and the internal jugular veins.

Pituitary tumours (A)

Pituitary tumours constitute 10–15% of intracranial neoplasms and are often discovered incidentally on imaging performed for an unrelated reason (pituitary incidentaloma). Their clinical features maycan result from local mass effects and/or hypersecretion.¹

The local mass effects depend on the size of the tumour and its anatomical position <u>/ and</u> extensions. Headache is usually the consequence of dural stretching. The neuroophthalmological effects include visual field defects (usually bitemporal hemianopia) from compression of the optic pathways, and ocular nerve palsies caused by lateral extension to the cavernous sinuses. Erosion of the sellar floor <u>maycan</u> result in sinusitis, CSF rhinorrhoea, and meningitis. The anterior pituitary hormone deficits tend to occur in a specific order, with GH and gonadotrophins affected first, followed by ACTH and TSH. PRL secretion is the most resistant, and decreased <u>levelconcentration</u>s indicate severe pituitary damage.

All patients with a pituitary mass should undergo testing for hypopituitarism and neuroophthalmological evaluation. In cases of <u>With</u> pituitary adenomas, hormonal hypersecretion needs to be assessed. Careful neuroradiology review aiming to identify imaging features helpful for the differential diagnosis is also mandatory.

| Pituitary adenomas-(B) | Formatted: Font: Bold, Not Italic |
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| Pituitary adenomas account for 90% of pituitary tumours and have a prevalence of 77.6+ | Formatted: Indent: First line: 0 cm |
| cases <mark>/per</mark> 100,000 inhabitants in the UK. They are benign lesions arising from | |
| adenohypophyseal cells and, based on their size, are classified as microadenomas (<10 mm | |
| in diameter) or macroadenomas (≥10 mm in diameter). They may hypersecrete hypophyseal | |
| hormones (functioning) or maycan be clinically non-functioning. Whilst-Although mostthe | |
| majority is are sporadic, <u>, they are in rare cases</u> , they may be related with to hereditary | |
| syndromes, like-such as multiple endocrine neoplasia type 1, Carney complex or familial# | |
| isolated pituitary adenomas. | |
| | |
| Non-functioning pituitary adenomas (C) | Formatted: Font: Bold, No underline |
| Non functioning pituitary adenomas (NFA): these comprise 15–37% of all pituitary | Formatted: Font: Bold |
| adenomas and have a prevalence of 722/ per 100,000 inhabitants. As they are not | |
| associated with hormonal hypersecretion, they usually escape early diagnosis, and are | |
| mostly recognized when they are large enough to exert pressure effects to on surrounding | |
| tissues; thus, at the time of detection, 67–90% of them are macroadenomas. Additionally, at | |
| diagnosis, 60- <u></u> 85% of the patients have at least one pituitary hormone deficiency. | |

First-line treatment for the macroadenomas is surgery, usually with theusing a trans_sphenoidal approach; this, which aims to improve or resolve the mass effects on adjacent structures, and especially the optic pathways. Radiotherapy is <u>maycan be</u> offered as adjuvant treatment after surgery, aiming to prevent tumour regrowth. The management of regrown <u>non-functioning pituitary adenomas NFAs</u>-includes observation, surgery, radiotherapy or <u>a</u> combination of surgery and radiotherapy.

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Functioning pituitary adenomasthese release excessive amounts of active hypophyseal hormones into the systemic circulation, resulting in multiple clinical manifestations. Prolactinomas are the most prevalent hormone-secreting adenomas followed by GH-producing, corticotroph₇ and thyrotroph adenomas. The clinical presentation, diagnosis and treatment of functioning adenomas depend on the type of hormone(s) secreted (Table 2).

Functioning pituitary adenomas (C):

| Other sellar or parasellar masses | (| Formatted: Font: Bold, Not Italic |
|---|---------------|------------------------------------|
| ,(8) | (| Formatted: Font: Bold, Not Italic |
| Rathke's cleft cysts- (C) | | Formatted: Font: Bold, No underlin |
| thesey are benign sellar and/or suprasellar lesions that arise from remnants of Rathke's | - (| |
| pouch. ² Their size varies, as well asdoes their content (ranging from a clear CSF-CSF-like | | |
| liquid to a thick mucoid material made up of cholesterol and protein). Patients with a | | |
| Rathke's cleft cyst usually present with symptoms of compression to of adjacent structures, | | |
| although incidentally detected cases are also reported. Surgery is the treatment of choice in | | |
| patients with symptomatic cysts. | | |
| Other cystic lesions usually <u>found</u> in the suprasellar region include arachnoid, | | |
| epidermoid, and dermoid cysts. | | |
| Craniopharyngiomas: - (C) | (| Formatted: Font: Bold, No underlin |
| these are sellar/parasellar tumours that arise from embryonic remnants of Rathke's pouch. | \rightarrow | Formatted: Font: Bold, No underlin |
| They are commonly found during childhood and adolescence ₂ - ³ However, theybut can be | C | |
| also-diagnosed at any age. They are usually large masses with suprasellar extension and can | | |
| invade the third ventricle and other brain structures. On imaging, craniopharyngiomas are | | |
| mostly often predominantly cystic and usually filled with a cholesterol-rich fluid; purely or | | |
| predominantly solid or purely cystic masses can beare also detected. Another common | | |
| finding The presence of is calcifications inside the tumour is also another common finding. | | |
| Patients with craniopharyngioma demonstrate many clinical features due toresulting | | |
| from local tumour effects (headache, vomiting, papilloedema, visual field deficits) and due | | |
| to-dysfunction of the hypothalamus and pituitary (obesity, problems with appetite, satiety, | | |
| temperature control, hypopituitarism). Treatment of these tumours involves hypothalamus- | | |
| sparing surgery followed by local radiation therapy. The long-term outcomes are not optimal | | |
| due to the<u>because of</u> significant morbidities. | | |
| Hypophysitis- (C): <u>this</u> | (| Formatted: Font: Bold, No underlin |
| Hypophysitis is an inflammatory disease that can affect both lobes of the pituitary , as well as $$ | | Formatted: No underline |
| the stalk. The Diagnosis of this condition can be difficult given that 50% of the cases are | | |
| misdiagnosed as pituitary adenomas (Table 3). | | |
| Pituitary stalk lesions- (C) : | (| Formatted: Font: Bold, No underli |
| the spectrum of pituitary stalk is extensive, but can be considered and is divided in three | - (| |
| main categoriesunder three main headings: | | |
| neoplastic — thesey account for the majority of most pituitary stalk lesions, with | | |
| metastases (mainly from lung and breast cancer) and lymphoma being the most | | |
| frequent, followed by germ cell tumors and astrocytomas- | | |
| inflammatory and infiltrative diseases_; hypophysitis is the most common cause, | (| Formatted: Font: Bold |
| followed by neurosarcoidosis and Langerhanhs' cell histiocytosis- | | |
| congenital conditions_: thesey constitute the minority of stalk lesions; pituitary | | |
| hypoplasia and Rathke's cleft cyst are the most frequent causes. | | |
| Central diabetes insipidus and hyperprolactinaemia (absence of normal hypothalamic | | |
| dopamine suppression of prolactin release due tocaused by stalk interruption) are the most | | |
| common est hormonal findings among st patients with pituitary stalk lesions. Anterior | | |
| hypopituitarism can also be observed. All patients should undergo clinical, biochemical, and | | |
| imaging investigations, and if diagnosis remains unclear, then a pituitary stalk biopsy maycan | | |
| be considered <u>if the diagnosis remains unclear</u> . ⁴ | | |
| Other lesions (C) | کسہ | Formatted: Font: Bold, No underli |
| | | TOTHELEO, FUIL BUIL, NU UNDER |

Other less frequent sellar and parasellar lesions include:

- non-adenomatous tumours<u>—</u>meningiomas (comprising <u>the majoritymost</u> of this group), chordomas, gliomas and pituicytomas.
- pituitary infections÷ <u>–</u> haematogenous or local spread of infectious agents can result in pituitary abscess and perisellar arachnoiditis.
- vascular lesions: ____internal carotid artery aneurysms, which can manifest as parasellar lesions.

| parasellar lesions. | |
|---|--|
| Hypopituitarism-(A) | Formatted: Font color: Blue |
| Hypopituitarism is the result of conditions that reduce or destroy the pituitary function or interfere with the hypothalamic secretion of pituitary-releasing hormones, leading to <u>a</u> complete or partial deficiency in <u>of</u> pituitary hormones. ⁵ | Formatted: Indent: First line: 0 cm |
| <u>Etiology Actiology(B)</u> | Formatted: Font: Bold, Not Italic |
| Apart from the space-occupying lesions of the pituitary, other conditions resulting in hypopituitarism-include: | Formatted: Font: Bold, Not Italic |
| • Vascular | Formatted: Font: Bold |
| frequent vascular cause of hypopituitarism. It can be a life-threatening condition that requires acute management (Table 4). On the other handln contrast, | |
| postpartum isch <u>a</u> emic pituitary necrosis (Sheehan's syndrome) is <u>now</u> relatively rare due tobecause of the advances in obstetric care. | |
| • Traumatic — | Formatted: Font: Bold |
| <u>cause in</u> hypopituitarism. | |
| Jatrogenic:surgery and irradiation therapy for sellar/extasellar masses or brain tumours maycan compromise pituitary function. In addition, partial hypopituitarism | Formatted: Font: Bold |
| may be seen as a result of <u>can be caused</u> various medications (glucocorticoids, opiates, etc.). | |
| Congenital → They can <u>maythese can manifest as be</u> isolated <u>deficiencies</u> due | Formatted: Font: Bold |
| tocaused by mutations in the genes coding for a specific hormone, or multiple deficiencies resulting from abnormal pituitary development (e.g. PROP1, HESX1, | Formattad, Fost, Nat Italia |
| and POU1F1 gene mutations). | - Formatted: Font: Not Italic |
| Clinical manifestation and diagnosis-(B) | Formatted: Font: Bold, Not Italic |
| The clinical features of hypopituitarism vary and depend on the rapidity of onset, the severity of the hormonal defect(s), and the number and type of hormones affected. The Establishingment of the diagnosis requires hormonal measurements (basal or after dynamic tests) (Table 5). | |
| Treatment (B) | Formatted: Font: Bold, Not Italic |
| Hydrocortisone is the treatment of choice in central hypoadrenalism (usual total daily dosage of 15–20 mg divided into two or three doses). Patients should take the highest dose | Comment [CMW3]: AQ: please confirm |
| in the morning <u>on at a</u> wakening and the second in the afternoon (two-dose regimen), or the second and third at lunch-time and <u>in the</u> late afternoon, respectively (three-dose regimen). | that these doses match those recommended by the BNF. |
| Central hypothyroidism is managed with levothyroxine in doses sufficient to achieve serum free thyroxine levelconcentrations in the mid to upper half of the reference range, | Comment [NK4]: These are the doses mostly used in clinical practice. |
| <u>but</u> and only after adequate hydrocortisone initiation (because thyroid hormone replacement maycan aggravate adrenal insufficiency in patients with untreated | Formatted: Indent: First line: 1.25 cm |
| steroid <u>corticosteroid</u> deficiency). | Comment [CMW5]: AQ: can 'females' |
| Males Men and premenopausal females with central hypogonadism should be | include children, or is it just adult women? |
| offered sex-steroid replacement therapy (provided there are no contraindications). Diabetes insipidus is managed with desmopressin, and GH deficiency with recombinant GH. | Comment [NK6]: This is just premenopausal women (the statement does not apply to children) |

Key references

- 1. Freda PU, Beckers AM, Katznelson L, et al. Pituitary incidentaloma: an endocrine society clinical practice guideline. *J Clin Endocrinol Metab* 2011; **96**: 894–904.
- 2. Trifanescu R, Ansorge O, Wass JA, Grossman AB, Karavitaki N. Rathke's cleft cysts. *Clin Endocrinol (Oxf)* 2012; **76**: 151–60.
- Karavitaki N, Cudlip S, Adams CB, Wass JA. Craniopharyngiomas. *Endocr Rev* 2006; 27: 371–97.
- 4. Catford S, Wang YY, Wong R. Pituitary stalk lesions: systematic review and clinical guidance. *Clin Endocrinol (Oxf)* 2016; **85**: 507–21.
- 5. Fleseriu M, Hashim IA, Karavitaki N, et al. Hormonal replacement in hypopituitarism in adults: an Endocrine Society Clinical Practice Guideline. *J Clin Endocrinol Metab* 2016; **101**: 3888–2921.

 Table 1: Hormone-producing cells in the anterior pituitary gland

| Type of cell | Percentage of cells in anterior pituitary lobe | Distribution of cells |
|---------------|--|--|
| Somatotrophs | 4050% | Lateral wings of adenohypophysis |
| Lactotrophs | 15 <mark></mark> 20% | Dispersed populations throughout the anterior lobe; mainly in the posterior part of the lateral wings |
| Corticotrophs | 15_20% | Middle and posterior portion of anterior lobe |
| Gonadotrophs | 1015% | Distributed through anterior lobe |
| Thyrotrophs | 510% | Anterior medial part of adenohypophysis |

 Table 2: Clinical features, diagnosis and treatment of functioning pituitary adenomas

| Brolactinama | Comment [MC7], p. J. H. D. H. |
|--|---|
| Prolactinoma Clinical features | Comment [MG7]: Production Dept to cross-reference to Prolactinoma chapter |
| Females:galactorrhoea, hypogonadism (oligo/amenorrhoea, and infertility) | from Niamh Martin |
| Maleshypogonadism (impotence, decreased libido), galactorrhoea (very rare) | Comment [CMW8]: AQ: throughout the table: does 'Females' and 'Males' refer |
| Diagnosis | to all ages or just adult patients? |
| Hyperprolactinaemia (after excludingsion of macroprolactinaemia and of other causes of increased PRL levelconcentrations) | Comment [NK9]: These apply to adults and wherever the statement involves children, we have mentioned this. |
| Immunoradiometric PRL measurement at a serum dilution of 1:100 in cases offor | |
| pituitary macroadenomas with normal or mildly elevated PRL values when the so- called 'hook effect' is suspected (high <u>levelconcentration</u> s of circulating PRL causes antibody saturation in the immunoradiometric assay, leading to artifactually | |
| artefactually low results) | - |
| Treatment Medical therapy with dopamine agonists (cabergoline as firstline treatment, <u>alternatively</u> bromocriptine alternatively) Surgery if <u>there is</u> resistance or intolerance to medical treatment | |
| Radiotherapy in resistant or aggressive prolactinomas, usually after surgery | |
| GH-secreting adenomas | Comment [MG10]: Production Dept to |
| Clinical features | cross-reference to Acromegaly chapter |
| Adults: <u>–</u> acromegaly (acral enlargement, prognathism, frontal bossing, soft tissue overgrowth, hyperhidrosis, arthralgias, fatigue) | from Will Drake |
| Children and adolescenceadolescents:- – gigantism | |
| Diagnosis | |
| Serum IGF-1 level<u>concentration</u>s above the age-and sex-adjusted reference range Lack of suppression of GH level<u>concentration</u>s (<0.4 μg/4/litre) during a 75 g⁺ oral glucose load | |
| Treatment | |
| Surgery | |
| Medical therapy÷<u></u>somatostatin analog<u>ue</u>s, dopamine agonists, pegvisomant Radiotherapy if no control of the disease is not controlled by surgery and medical treatment | |
| ACTH-secreting adenomas (Cushing's disease) | Comment [MG11]: Production Dept to |
| Clinical features | cross-reference to Cushing's chapter from John Newell-Price |
| Cushing's syndrome phenotype (weight gain and central obesity, skin thinning, purple striae, moon face, buffalo hump, proximal muscle weakness, spontaneous ecchymosis, increased supraclavicular fullness) | |
| Diagnosis | |
| Endogenous hypercortisolism (festablished by increased 24-hour urinarye free cortisol levelconcentrations, loss of diurnal rhythm of cortisol secretion (serum, or salivary), lack of serum cortisol suppression on overnight or low-low-dose dexamethasone suppression test]. | |
| Non-suppressed plasma morning–ACTH levelconcentrations | |

| | • | CRH stimulation test, high-high-dose dexamethasone suppression test, bilateral | | |
|---|----------------------------------|--|--|--|
| • | l | inferior petrosal sinus sampling | | |
| | Treatment | | | |
| | • | Surgery | | |
| | • | Radiotherapy | | |
| I | • | Medical therapy÷ <u>–</u> most commonly <u>used-with</u> steroidogenesis inhibitors | | |
| | • | Bilateral adrenalectomy | | |
| | TSH-se | creting adenomas | | |
| | Clinical | features | | |
| | • | Hyperthyroidism | | |
| | Diagno | sis | | |
| Ī | • | Non-suppressed TSH levelconcentrations in the presence of high FT4free T4 and- | | |
| | l | Ffree T3 concentrations | | |
| | • | TRH stimulation test, T3 suppression test | | |
| | Treatm | | | |
| | • | Surgery | | |
| l | • | Medical therapy with somatostatin analogues (usually after non-curative surgery) | | |
| | • | Radiotherapy if no control of the disease is not controlled by surgery and medical | | |
| ļ | l | treatment | | |
| | Functioning gonadotroph adenomas | | | |
| | Clinical features | | | |
| I | • | Females <u>- – menstrual irregularities (oligo/amenorrhoea, spotting, menorrhagia)</u> , | | |
| 1 | l | infertility, ovarian hyperstimulation syndrome (premenopausal women); no clinical | | |
| | l | syndrome in postmenopausal women | | |
| I | • | Males : <u>–</u> testicular enlargement, hypogonadism | | |
| ļ | Diagno | | | |
| I | • | Females:- <u>-</u> hyperoestrogenism (occasionally normal or fluctuating oestrogen | | |
| | l | levelconcentrations); serum FSH levelconcentrations mildly elevated or within | | |
| • | I | reference range; serum LH suppressed or less often within reference range | | |
| | • | Males: <u>–</u> serum FSH elevated; varying serum LH and testosterone | | |
| | I | level <u>concentration</u> s (slightly below the reference range, normal or elevated); | | |
| • | I | increased sperm count may be seen | | |
| | Treatm | | | |
| | • | Surgery combined or not with radiotherapy | | |
| ľ | In all ca | ises, pressure effects of the adenoma to on surrounding structures from the | | |
| | - | na-may be seen. | | |
| I | CRH <mark>÷</mark> | corticotrop <u>h</u> in-releasing hormone , ; (PRL: prolactin, GH: growth hormone, IGF-1 : | | |
| | | like growth factor 1, ACTH: adrenocorticotropic hormone, TSH: thyroid-stimulating | | |
| I | | ne, FFT3, free triiodothyronine; T4, free thyroxine, FSH: follicle stimulating | | |
| I | | ne_LH: luteinizing hormone). | | |

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Table 3: Types, clinical features, imaging characteristics and treatment of hypophysitis

Types of hypophysitis

- Lymphocytic hypophysitis-__classically in women during or after parturition
- Granulomatous hypophysitis
- Xanthomatous hypophysitis
- IgG-4 hypophysitis _ in patients with IgG-4-4-related disease
- Drug-induced hypophysitis (ipilimumab, nivolumab, pembrolizumab)

Clinical features

- Local tumour effects (headache, visual deterioration)
- Anterior hypopituitarism
- Diabetes insipidus

Imaging characteristics

- Symmetrical enlargement of the pituitary; stalk <u>maycan</u> be thickened, and suprasellar extension <u>maycan</u> be seen
- A highly cystic lesion is often found in xanthomatous hypophysitis

Treatment

- High doses of glucocorticoids (although potential side eadverse effects should be carefully considered on an individual case basis)
- Hormone replacement therapy in cases of hypopituitarism
- Trans_sphenoidal surgery if visual deterioration and for histological confirmation of the diagnosis

IgG, immunoglobulin G.

Comment [CMW12]: AQ: please confirm definition is correct.

Comment [NK13]: This is correct.

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 Table 4: Pituitary apoplexy: pathophysiology, clinical presentation, differential diagnosis and management

| Patho | physiology |
|---------|---|
| ٠ | Haemorrhage or infarction, usually on a background of a pituitary tumour |
| ٠ | Precipitating Causative factors: — hypertension, major surgery, coagulopathies, |
| | anticoagulation therapy, pregnancy, dynamic pituitary function testing, and head |
| | trauma |
| Clinica | l presentation |
| ٠ | Acute severe headache that maycan be accompanied by nausea and vomiting |
| ٠ | Ocular nerve palsies, reduced visual acuity, visual field defects (usually bitemporal |
| | hemianopia) |
| ٠ | Meningism (fever, neck stiffness, photophobia) |
| ٠ | Altered consciousness |
| Differe | ntial diagnosis |
| • | Subarachnoid haemorrhage |
| • | Meningitis (bacterial or viral) |
| ٠ | Brainstem infarction |
| ٠ | Cavernous sinus thrombosis |
| Manag | jement |
| • | Supportive measures to ensure haemodynamic stability and careful monitoring of |
| | fluid and electrolyte balance |
| • | Immediate administration of high-dose glucocorticoids, especially in |
| | haemodynamically unstable cases or with severe neurological or neuro- |
| | op <u>h</u> thalmological signs; thisit covers the increased risk of hypoadrenalism and has |
| | significant anti-inflammatory and anti-oedematous effects |
| • | Surgery is offered in the presence of if there is severe visual acuity and visual field |
| | impairment (not isolated ophthalmoplegia), altered consciousness, deteriorating |
| | visual or neurological signs or of further enlargement of the sellar mass on serial |
| | imaging |

 Table 5:
 Diagnosis of pituitary hormone deficits

| GH deficiency | |
|--|---|
| • GH stimulation testing is mandatory (insulin tolerance test, GHRH and arginine | |
| stimulation test, glucagon stimulation test) | |
| Normal IGF-1 levelconcentrations does not exclude the diagnosis | |
| FSH/LH deficiency | |
| • Males | Comment [CMW14]: AQ: Does 'Males |
| and ideally corrected for SHBG) and low or normal gonadotrop <u>h</u> ins | and 'Females' refer to all ages? |
| Females:low serum oestradiol levelconcentrations and low or normal | Comment [NK15]: This section refers to post-pubertal subjects. I would suggest |
| gonadotrop <u>h</u> ins in the presence of oligomenorrhoea or amenorrh <u>o</u> ea | we leave it the way it is. |
| (premenopausal women); absence of high serum FSH and LH (postmenopausal | |
| women) | |
| ACTH deficiency | |
| • Serum cortisol levelconcentrations at 08:00–09:00 hours <100 nmol/L/litre (in the | |
| absence of <mark>steroid</mark> corticosteroid administration) are indicative of adrenal | |
| insufficiency | |
| If morning cortisol values are between 100 - and 400 nmol/4/litre, a dynamic test | |
| (e.g. insulin tolerance test, glucagon stimulation test) is required to establish the | |
| diagnosis- (eg. insulin tolerance test, glucagon stimulation test) . The Cut-offs for | |
| serum cortisol need to<u>should</u> be defined by each lab<u>oratory,</u> but overall peak serum | |
| cortisol >500 nmol <mark>/4/litre</mark> at 30 or 60 minutes excludes the diagnosis | |
| Central hypothyroidism | |
| • Low free thyroxine levelconcentrations in conjunction with a-low, normal, or mildly | |
| elevated TSH in the setting of pituitary disease | |
| Central diabetes inspidus | |
| • Confirm hypotonic polyuria - simultaneous measurement of serum/plasma and | |
| urine osmolarity in the presence of polyuria (>50 mLml/kg of body weight/24 | |
| hours *), <u>a</u> | Formatted: Superscript |
| • Urine osmolarity >600 mOsmol/kg) effectively excludes the diagnosis of diabetes | |
| insipidus (urine osmolarity/plasma osmolarity ratio should be ≥ 2 during urine | |
| concentration); urine dipstick should be negative for glucose | |
| Water deprivation test may also be needed | |
| *-Clinical correlation is important in this context (for example <u>e.g.</u> hypotonic polyuria | Formatted: Font: Not Italic |
| naycan also result following from the infusion of large volumes of intravenous fluids). | Formatted: Font: Not Italic |
| GH: growth hormone, GHRH: growth hormone-releasing hormone, i IGF-1: j insulin-like | Formatted: Font: Not Italic |
| growth factor 1, -: FSH: follicle stimulating hormone, LH: luteinizing hormone, ACTH: | Formatted: Font: Not Italic |
| adrenocorticotropic hormone, TSH: thyroid-stimulating hormone, SHBG:sex hormone | Formatted: Font: Not Italic |
| binding globulin 🚽 | Formatted: Font: Not Italic |
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TEST YOURSELF

To test your knowledge based on the article you have just read, please complete the questions below. The answers can be found at the end of the issue or online here.

Question 1

A 30-year-old woman was referred to Endocrinology_the endocrine clinic withfor a one1year history of headaches, amenorrhoea and decreased libido. She had no visual deterioration. She hadThere was no significant past medical-history, and she was not on taking any medication. Hormonal workupInitial i

Investigations

-<u>erevealed</u> Hypogonadotrop<u>h</u>ic hypogonadism<mark>, as well as increased</mark>

Serum prolactin levels (15,000 mU/L/litre, reference range (60-_620).
 The Insulin-like growth factor 1IGF-1 and adrenocorticotrophic hormoneACTH reserve were normal

•Thyroid function was normal and there was no hypothyroidism. She had no visual deterioration.

•MR scan of the pituitary Pituitary MRI revealed a macroadenoma abutting, but not compressing, the optic chiasm.

What is the best choice of treatment?

- A. Dopamine agonist
- B. Levothyroxine
- C. Pituitary radiotherapy
- D. Somatostatin analogue
- E. Trans-sphenoidal adenomeatectomy

Correct answer: A.

This patient has a macroprolactinoma. Dopamine agonists are the first-line treatment for this tumour, as they can <u>lead tocorrect</u> prolactin <u>normalizationabnormalities</u>, restor<u>eation of</u> gonadal function and <u>lead to</u> tumour shrinkage. Surgery (E) is an alternative option for patients who show resistance or intolerance to medical treatment. Radiation therapy (C) is used only <u>in for</u> resistant or *f*aggressive prolactinomas, and usually after surgery. Somatostatin analogues (D) are not effective in the treatment of prolactinomas, and levothyroxine (B) has no place in the management of this patient as there is no hypothyroidism.

Question 2

A 45-year-old man was referred to <u>the endocrine clinic for further assessment.</u> <u>ofEndocrinology due to a 3 cm pituitary mass, which had been</u> found on <u>limaging</u>, performed for headaches, had shown a 3 cm pituitary mass, likely to be a pituitary adenoma, occupying the sella, with suprasellar extension and invasion of the cavernous sinuses. <u>The</u> <u>patient had</u> reported tiredness and low libido <u>but no polyuria or polydipsia</u>. The <u>Nneuroradiologist reported that gy review suggested that the mass wasis most likely to be a pituitary adenoma, occupying the sella with suprasellar extension and invasion of the cavernous sinuses. There were no clinical manifestations of acromegaly, Cushing's <u>syndrome</u> or hyperthyroidism.</u>

Investigations

<u>Hormonal tests revealed IGF-1Insulin-like growth factor 1</u> below the reference range,
<u>Hypogonadotrophic hypogonadism</u>,

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Comment [JC16]: Authors: Could we have some figures and ranges for this to simulate clinical practice and so they have to do some interpretation?

Comment [NK17]: I would suggest we leave it like this because the values of the FSH, LH and oestradiol depend on the day of the cycle. Here by mentioning amenorrhoea and hypogonadotrophic hypogonadism, we pass the message we would like to. Happy to follow your advice though if you felt this is necessary.

Comment [CMW18]: AQ: do UK report sheets use mU/L for prolactin, rather than SI units of pmol/litre?

Comment [NK19]: Yes, UK uses mU/L.

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Comment [JC20]: Author: Could we have some figures for the first two points, with reference ranges and check that the next two are realistic

Comment [NK21]: The reference range for IGF-I ranges varies between labs and assays (also affected by age) and this is why we left it like this. For the hypogonadotrophic hypogonadism, we would suggest adding 9.00 am serum testosterone and gonadotrophins below the reference range. •ACTH Plasma adrenocorticotropic hormone 1.8 pmol/L/litre (3.3–15.4) and •TSH-Serum thyroid-stimulating hormone 0.2 mU/L/litre (0.4–5.0) deficiency. There was no evidence of diabetes insipidus.

What should be the first step in the patient's management?

- A. <u>Administer Dd</u>esmopressin
- B. Growth hormone replacement therapy
- C. Hydrocortisone replacement therapy
- D. Levothyroxine replacement therapy
- E. Testosterone replacement therapy

Correct answer: C.

In patients with anterior hypopituitarism, hydrocortisone therapy should be initiated before any other hormonal replacement. Levothyroxine (D) should be offered after adequate hydrocortisone replacement is-has been established, as otherwise, levothyroxine maycan otherwise aggravate adrenal insufficiency and lead to adrenal crisis. Growth hormone (B) and sex-steroid replacement therapy (E) can be offered later (provided there are no contraindications). -Treatment with desmopressin (A) would be indicated if there was-were diabetes insipidus.

Question 3

A 52-year-old man<u>presented acutely with</u> was admitted to the Emergency Department with acute-headache, fever, confusion<u>-and complaining</u>ts that light hurt his eyes., photophobia, and hypotension. His wife <u>confirmed_reported</u> a <u>12–12-</u>month history of tiredness and episodes of feeling lightheaded. <u>-There was no history of polydipsia or polyuria</u>. He had a history of is medical history included hypertension.

On clinical examination, he was confused, with a temperature of \mp 38.0°⁶C, HRheart rate 115 beats/minminute, and BPblood pressure 102/68 mmHg. There was nNeck stiffness was identified <u>noted on clinical examination</u>. There were but no signs of ocular nerve palsies. The

Investigations

•Cerebrospinal fluidSF examination showed<u>A lumbar puncture report revealed</u> normal level<u>concentration</u>s of <u>CSF</u>-protein, <u>normal CSF:a</u>_plasma:-glucose ratio and no-<u>presence of</u> cells. <u>CSF cC</u>ulture and Gram stain_, <u>as and well as _</u>analysis for xanthochromia were negative.

Hormonal evaluation identified hypogonadotropic hypogonadotrophic hypogonadism₇
 Plasma adrenocorticotrophic hormone 1.6 pmol/L/litre (3.3–15.4)

<u>Serum thyroid-stimulating hormone 0.3 mU/L/litre (0.4–5.0)</u> ACTH and TSH deficiency.
 There was no evidence of diabetes insipidus.

<u>•MR scan of the brain</u><u>Brain</u> MRI showed a large pituitary tumour with signs of haemorrhage, but no other abnormalities-

What is the most likely final diagnosis?

- A. Brainstem infarction
- B. Cavernous sinus thrombosis
- C. Meningitis
- D. Pituitary apoplexy
- E. Subarachnoid haemorrhage

Correct answer: D.

Comment [CMW22]: AQ: does this imply 'should only be offered after'?

Comment [NK23]: This is correct, please change the statement as you suggest.

Comment [JC24]: Author: Or an investigation which excludes DI

Comment [NK25]: I would not use this phrase – investigations for DI include also blood and urine tests. Here we just present the history.

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Comment [CMW26]: AQ: please clarify 'a plasma:glucose ratio'. Is some text missing?

Comment [NK27]: Please change to glucose > 60% of serum glucose

Comment [JC28]: Could there be some figures for this with normal ranges?

Comment [NK29]: As in the previous case, I would suggest we add we would suggest adding 9.00 am serum testosterone and gonadotrophins below the reference range.

This patient <u>has developedhad a</u> pituitary apoplexy on a background of a pituitary tumour. The normal <u>results on CSF-cerebrospinal fluid</u> analysis <u>results</u> exclude <u>the a</u> diagnosis of meningitis (<u>C</u>) and subarachnoid haemorrhage (<u>E</u>). The fact that no cranial nerve palsies were noted, the absence of a history of central face or paranasal sinus<u>es</u> infection, and the negative findings <u>o</u>in <u>the</u> MRI eliminate the possibility of cavernous sinus thrombosis (<u>B</u>). Finally, there were no signs <u>o</u>in the MR <u>scan</u>¹ to indicateing brainstem infarction (<u>A</u>).

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