ABSTRACT

The diagnosis and treatment of pituitary disease in pregnancy represents a special clinical challenge. Not least because there is very little data on the treatment of pregnant patients with pituitary disorders. A selective search of the literature was carried out with the aim of compiling evidence about the diagnosis and treatment of pituitary disease in pregnancy. The search covered the databases PubMed/MEDLINE including PubMed Central and also used the Livivo (ZB MED) search engine. Recent studies were evaluated for recommendations about the care of pregnant patients with hormone-inactive and hormone-active pituitary adenomas (prolactinoma, acromegaly and Cushing’s disease), pituitary insufficiency, pituitary apoplexy and hypophysitis. The most well-established forms of treatment are for prolactinoma, due to the incidence of this disease and its impact on fertility. When pregnancy has been confirmed, prolactinoma treatment with dopamine agonists should be paused. Although microprolactinomas rarely increase significantly in size after the administration of dopamine agonists is discontinued, symptomatic tumor growth of macroprolactinomas can occur. In such cases, treatment with dopamine agonists can be resumed. If the primary tumor is large and the risk that it will continue to grow is high, it may be necessary to continue medical treatment from the start of pregnancy. If one of the partners has a pituitary disorder, it is often still possible for many couples to achieve their wish of having children if they receive medical support to plan and the pregnancy is carefully monitored. Given the complexity of pituitary disease, pregnant patients with pituitary disorders
Introduction

During normal pregnancy, the endocrine system and metabolism of the pituitary gland undergoes significant changes as a result of placental hormone secretions. The pituitary gland increases in volume due to hyperplasia of the prolactin cells and changes in pituitary hormone secretions [1]. Prolactin levels rise throughout the entire pregnancy, with the placenta also taking over the synthesis of growth hormone (GH) and suppressing pituitary GH [2]. GH levels continue to rise until the 36th/37th week of pregnancy, achieving levels which are many times higher than before the start of pregnancy. Insulin-like growth factor 1 (IGF-1) levels may initially drop, but towards the end of pregnancy they will also be higher than at the start of pregnancy. The placenta additionally synthesizes corticotropin-releasing hormone (CRH), which stimulates the release of adrenocorticotropic hormone (ACTH) and cortisol. In the first trimester of pregnancy placental human chorionic gonadotropin (hCG) stimulates the thyroid, and the liver increases its synthesis of binding proteins [3]. The rise of binding globulins is followed by an increase in the concentrations of total thyroxine (T4) and cortisol until the 16th–20th week of pregnancy [1, 4].

Pituitary disorders are rare and may be accompanied by disorders of gonadal function and fertility. The diagnosis and treatment of pituitary disorders in pregnancy represents a special clinical challenge, not least because there is very little data on the treatment of pregnant patients with pituitary disorders. There are very few treatment recommendations, and they are often based on the approach used to treat non-pregnant women and attempt to take the specific physiology of pregnancy into consideration. Because the overall data is limited, this publication aims to provide a summary of the diagnosis and treatment of pituitary disorders in pregnancy. To this end, a search of the literature was carried out using the databases PubMed/MEDLINE together with PubMed Central and the Livivo (ZB MED) search engine. The search terms used were “pregnancy” or “Schwangerschaft” combined with the respective disease, e.g. “pituitary apoplexy” or “Hypophysenapoplex”. The publications found in the databases and the authors’ own experience were discussed by all of the authors at several meetings and the authors reached a consensus about the recommended treatment.

This review will start by discussing various aspects relating to tumors which occur in the region of the sella turcica in the order of their frequency of occurrence and then look at different types of pituitary axis insufficiency. The final part will consider issues of fertility in patients with pituitary disorders.

Review

Pituitary adenomas

Prolactinoma

Female patients with hyperprolactinemia almost always present with hypogonadotropic or normogonadotropic hypogonadism. This can usually be resolved by the administration of a dopamine agonist [5]. Regular ovulatory cycles usually return after therapy with a dopamine agonist is initiated, and spontaneous conception is possible in most cases. Dopamine agonist therapy should be discontinued as soon as the pregnancy has been confirmed [5]. Based on current data, taking the dopamine agonists cabergoline and bromocriptine during the period of conception and in early pregnancy does not pose a significant risk for mother or child [6, 7]. Microprolactinomas (< 1 cm) and small macroprolactinomas limited to the sella turcica (≥ 1 cm) rarely increase significantly in size even after the administration of dopamine agonists is discontinued; no special therapy is required. However, asymptomatic tumor growth may occur during pregnancy in 20 to 30% of cases with macroprolactinoma [6], although it is not possible to predict...
the growth behavior of the tumor. In principle, female patients with macroprolactinoma should only consider pregnancy after medical therapy has been shown to be effective or after surgery of the pituitary adenoma. The treatment options must be individually discussed with the patient before the start of pregnancy [6, 7] (Fig. 1). If the patient develops local symptoms such as headache or impaired vision during pregnancy due to growth of the tumor, then treatment with a dopamine agonist can be resumed; in cases with a large primary macroprolactinoma and a significant risk of tumor growth, treatment with a dopamine agonist may be continued from the start of pregnancy.

During pregnancy, patients with microprolactinoma should have a clinical examination every 3 months, with a particular focus on any headaches or visual impairments. It is recommended that patients with macroprolactinoma should have a clinical examination every month and visual field testing every three months [3, 5, 8]. The question whether prolactin values need to be measured regularly during pregnancy is still controversially discussed [9]; the authors are of the opinion that regular monitoring of prolactin levels is only useful for patients with macroprolactinoma. When measuring prolactin levels, the trimester-specific threshold values must also be taken into account [10] (Table 1). Exceeding the threshold values will not trigger any therapeutic consequences per se, but higher prolactin levels could be an indication that the prolactinoma is growing. In such cases, regular clinical examinations should be stepped up. If the prolactin levels rise to the level of the values measured at diagnosis, magnetic resonance imaging (MRI) may be indicated.

Regular clinical examinations and visual field testing is more important than measuring changes in prolactin levels [6, 11]. Magnetic resonance imaging without gadolinium is indicated for patients with macroprolactinoma and a pronounced rise in prolactin levels, symptoms such as persistent headache, visual impairment or reduced visual field [6, 11].

An asymptomatic prolactinoma is not a contra-indication for breastfeeding. Up to 35% of patients with microprolactinoma appear to be cured after pregnancy [12], probably either as a result of ischemia/necrosis of the adenoma or because of regressive transformation during the pregnancy. The administration of dopamine agonists should only be restarted after the patient has stopped breastfeeding [6].

Hormone-inactive pituitary adenoma/other tumors in the region of the sella turcica

If the patient has a macroadenoma which was detected prior to conception, the risk of developing Sheehan syndrome post par...

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**Table 1** Median prolactin concentrations in each trimester (10th–90th percentile) in uncomplicated pregnancies [10].

| Prolactin, ng/ml | All (n = 50) | Primipara (n = 25) | Multipara (n = 25) |
|------------------|-------------|-------------------|-------------------|
| 1st trimester    | 28.8 (16.3–57.6) | 32.6 (19.8–63.3) | 27.6 (10.6–47.0) |
| 2nd trimester    | 126 (54.9–206) | 105 (49.0–181)   | 139 (72–206)     |
| 3rd trimester    | 216 (124–318) | 177 (115–258)    | 225 (133–320)    |

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Acromegaly

Acromegaly is characterized by the increased secretion of circulating GH. The cause in more than 99% of cases is a benign GH-secreting tumor [21]. Other causes, for example a tumor secreting GH-releasing hormone (GHRH), are very rare [22]. GH and the target hormone IGF-1 which is synthesized in the liver do no cross the placental barrier [2], meaning that the infant is not affected by maternal acromegaly. According to retrospective case studies, active acromegaly in pregnancy is associated with a risk of the patient developing gestational diabetes and hypertension [23, 24]. Somatotrophic adenomas do not usually increase in size during pregnancy, with the possible exception of increased growth due to a “rebound” phenomenon after discontinuing therapy with somatostatin analogs.

Prior to conception, therapeutic options include surgical (usually transsphenoidal pituitary surgery), medical and radiotherapeutic procedures, which can be used as either primary or secondary treatment [25]. Medical therapy options include somatostatin analogs, dopamine agonists and the GH-receptor antagonist pegvisomant. It is recommended that women of child-bearing age treated with medication use effective contraceptive methods [26]. In case they have conceived, they should discontinue their acromegaly medication [24]. Therapy with a dopamine agonist may be considered for patients with tumors which extend to the chiasm or show invasive growth. The most data on the use of drugs in pregnancy is available for somatostatin analogs and the dopamine agonists bromocriptine and cabergoline [26]; continuation of therapy using somatostatin analogs during pregnancy has been described in case reports and small series [2, 23, 27]. There were no indications of teratogenicity or an increased rate of malformations. However, intrauterine growth retardation following treatment with somatostatin analogs was reported in individual cases [24]. The least data is currently available on the use of pegvisomant in pregnancy, and it should only be administered in exceptional cases during pregnancy [28, 29].

Large adenomas which extend to the chiasm should be evaluated clinically every three months with examination of the patient’s field of vision. Magnetic resonance imaging (MRI) may be indicated in patients who present with persistent headache, reduced vision or reduced field of vision, with MRI usually carried out without gadolinium, although contrast media may be used if necessary.

Patients with microadenoma who are not receiving medication and patients who have undergone surgical resection of a macroadenoma can breastfeed after an uncomplicated pregnancy [26]. Breastfeeding is contraindicated for patients receiving treatment with bromocriptine or cabergoline [26], and for somatostatin analogs or pegvisomant, as it is currently not known whether these substances pass into human breast milk [30 – 32].

Biochemical monitoring of acromegaly is difficult during pregnancy, as measured growth hormone concentrations increasingly mirror the production of placental GH over the course of the pregnancy and do not adequately reflect the activity of the somatotropic pituitary adenoma. Because of the high estrogen levels, IGF-1 levels usually drop at the start of pregnancy but increase again thereafter due to the effect of placental GH [2].
Cushing’s disease

Patients with florid Cushing’s disease rarely become pregnant due to the effect of hypercortisolism on the gonadal axis [33]. Fertility during remitting disease appears to correspond to that of the normal female population [34]. According to a systematic review of all 263 cases of pregnancy published between 1952 and 2015, pregnancy with active Cushing’s syndrome is often associated with diabetes mellitus, arterial hypertension, preeclampsia, maternal miscarriage as well as neonatal health problems [33]. Pregnancy during active disease is therefore inadvisable [33].

It is difficult to obtain an accurate diagnosis during pregnancy if there is a suspicion of newly occurring or recurrence of Cushing’s syndrome, not least because symptoms such as weight gain, glucose intolerance, hypertension, striae and mood swings can also be pregnancy-related. Physiologically, the excretion of free cortisol in urine (UFC) over a period of 24 hours doubles or triples in the first and second trimester [35], while the suppression of cortisol measured by the dexamethasone suppression test is reduced over the same period [36]. As the circadian cortisol rhythm remains the same in pregnancy (although with higher evening levels), alternative methods for cortisol determination include determining salivary cortisol levels or midnight serum cortisol test. When measuring UFC [37], only values which are more than triple the upper limit of standard levels are relevant as indications for disease. The treatment of Cushing’s disease during pregnancy depends on how active the disease is, as activity can range from mild hypercortisolism to aggressive disease. Whether transsphenoidal surgery should be considered during pregnancy depends on the MRI findings and the patient’s previous medical history/previous surgeries [38]. Individual reports have been published describing the use of drugs to treat Cushing’s disease during pregnancy, particularly the administration of metyrapone [39]. Considering its (limited) efficacy in non-pregnant patients [40, 41] treatment with the dopamine agonist cabergoline could be tried [42, 43]. Bilateral adrenalectomy may also represent an option, serving as a means of last resort to manage aggressive, uncontrolled progressive disease during pregnancy [38, 39, 44]. In a retrospective analysis of 136 pregnant women with Cushing’s syndrome, unilateral or bilateral adrenalectomy was carried out in 31 patients; the rate of live births in this cohort was 87% [38].

Lymphocytic hypophysitis

Lymphocytic hypophysitis is a rare autoimmune disease in which the pituitary gland is infiltrated by lymphocytes, plasma cells and macrophages [45]. In earlier publications, lymphocytic hypophysitis was predominantly reported in young women during or after pregnancy. More recent data suggest that this may have been a publication bias as an association between lymphocytic hypophysitis and pregnancy constitutes a distinctive constellation of symptoms [46]. It is often associated with other autoimmune diseases. In pregnancy, lymphocytic hypophysitis predominantly appears in the third trimester (cf. Honegger and Giese for an overview [46]). Early symptoms and imaging findings are often unspecific. Suggestive clinical symptoms are central diabetes insipidus and an unusual sequence of anterior pituitary lobe axis dysfunction (e.g. starting with the corticotropic hormone axis). Diagnostic pointers include thickened hypophyseal stalk, loss of the typically bright pituitary posterior lobe (“bright spot”) on T1-weighted MRI and a homogeneously tent-shaped pituitary gland with an intact, non-expanded sellar floor [46, 47]. Detection of hypophysial autoantibodies is not, as yet, diagnostically important. In addition to adequate hormone replacement therapy, treatment for patients experiencing visual deterioration and chiasma syndrome due to lesion expansion can consist of glucocorticoid therapy [48]. However, this is associated with a high rate of recurrence and, according to recent studies, its efficacy is limited [49]. Surgery during pregnancy may be considered in a few cases, particularly for patients with progressive deterioration of vision and reduced visual field acuity for whom glucocorticoid therapy was ineffective [38]. If symptoms are mild, treatment should be postponed until after the birth [46].

Pituitary apoplexy

Pituitary apoplexy is the term used to describe acute bleeding into the anterior pituitary or the infarction of a pre-existing pituitary adenoma [50]. A macroadenoma is present in more than 80% of cases. Typical symptoms are severe headache (“thunderclap headache”), visual deficits, double vision and altered mental status; they may be accompanied by symptoms of acute adrenocortical insufficiency. However, there may also be less clinically visible presenting symptoms. Risk factors for pituitary apoplexy are cardiovascular and other major surgical procedures, as well as hormone treatment, endocrine tests or taking antiocoagulants [51]. Because of the associated hypertrophy of lactotrophic cells and the increase in normal pituitary volume, pregnancy is also considered a risk factor for pituitary apoplexy [52]. Suggested causes for pituitary apoplexy include expansion of a tumor beyond the limits of its vascular supply and changes in the perfusion characteristics of the hormonally stimulated pituitary [50]. If the clinical symptoms only present in the form of headache and/or hormonal deficiencies, the recommended approach during pregnancy is usually surveillance combined with adequate substitution of the deficient hormone axis as well as the administration of a dopamine agonist if bleeding into a prolactinoma occurred [52]. Primary neurosurgical therapy should be discussed for patients with neuro-ophthalmological symptoms caused by a large symptomatic lesion. It is important to differentiate Sheehan syndrome (i.e., acute ischemic necrosis of the anterior pituitary due to extensive loss of blood and hypovolemic shock during childbirth) from pituitary apoplexy [53].

Pituitary insufficiency

Functional pituitary disorders can result from many different diseases occurring in the hypothalamic and pituitary region. In addition to hormone deficits due to pituitary adenoma, disorders may be caused by infections, radiotherapy, hypophysitis, pituitary apoplexy and metastasis [54]. Endocrine disorders are characterized by the partial loss of individual anterior pituitary functions (thyrotropic, gonadotrophic, somatotrophic and/or corticotrophic insufficiency), posterior pituitary function (diabetes insipidus) or complete pituitary insufficiency. Pituitary insufficiency is associated with an increased risk of pregnancy complications such as miscarriage, anemia, pregnancy-related hypertension, placental abruption, premature birth and postpartum bleeding [55–57].
The following sections provide information on the endocrine diagnosis of and therapy for specific functional losses of the anterior and posterior pituitary in pregnancy, including oxytocin deficiency. The management of pregnant women with complete pituitary insufficiency must be guided by the respective partial insufficiency. The management of pregnant women with complete pituitary in pregnancy, including oxytocin deficiency often require assisted pregnancy procedures [70]. There are some indications that growth hormone substitution can improve patients’ likelihood of conceiving. Once the pregnancy has been confirmed, growth hormone substitution can usually be terminated [70]. The results of a number of studies have been confirmed, growth hormone substitution can usually be terminated [70]. The results of a number of studies have been confirmed, growth hormone substitution can usually be terminated [70].

Secondary hypothyroidism

Because of the strong rise in human chorionic gonadotropin (hCG) during pregnancy with binding to the TSH receptor, a physiological decrease in TSH is often observed [58]. However, thyroid hormone substitution in patients with secondary hypothyroidism (thyrotropic insufficiency) must be guided by the levels of free thyroxine (fT4). Care must be taken when evaluating fT4 levels in pregnancy as the significant changes in protein and hormone levels that occur in pregnancy prevent accurate determination of fT4 levels using standard immunoassays [58]. During pregnancy, fT4 pregnancy-specific ranges apply [59] (Table 2). The aim should be to achieve values within the upper normal range. As the methods for measuring total thyroxine are less prone to interference, an alternative option could be to measure total thyroxine levels from the 16th week of gestation, with normal ranges increased by 50% to account for pregnancy changes [4].

During pregnancy, l-thyroxine dosages often have to be increased by up to 50% compared to pre-pregnancy dosages because of the increase in binding globulins [60,61]. But because it is not possible to estimate the quantitative impact of thyroid stimulation with hCG, in some cases the l-thyroxine dosage may even have to be reduced [60,61].

Up until at least the 12th week of gestation, the mother is the only source of thyroid hormones. Studies have suggested that maternal hypothyroxinemia may be associated with cognitive delays in early childhood [62]. The thyroid status of pregnant patients with secondary hypothyroidism should therefore be evaluated every 4 weeks.

Secondary adrenocortical insufficiency

Hypoglycemia, pituitary function failure or local symptoms caused by a pituitary lesion should also raise suspicion of secondary adrenocortical insufficiency (corticotropic insufficiency) [63]. Because of the massive increase in binding globulins, it is very difficult to interpret cortisol levels in serum correctly during pregnancy, and they are of only limited value for diagnosing adrenal gland insufficiency. However, in all three trimesters of pregnancy morning salivary cortisol levels do not differ significantly from those of healthy non-pregnant women [64]. The determination of morning salivary cortisol, preferably obtained during stress (e.g., “a cold shower”), is a useful first step to screen for adrenocortical insufficiency [65,66]. The most extensive data available on stimulation tests is for the 250 µg ACTH stimulation test. In this test, the cortisol serum threshold values in the second and third trimester are 60–80% higher than for non-pregnant women [67]. A peak cortisol value of > 30 µg/dl (828 nmol/l) is considered to exclude adrenocortical insufficiency [63]. Low dose (1 µg) ACTH stimulation tests provide a correct diagnosis in the majority of patients with threshold values of 18–20 µg/dl (497–553 nmol/l), while values of more than 30 µg/dl (828 nmol/l) in all probability exclude adrenocortical insufficiency [68]. Insulin hypoglycemia and metyrapone stimulation tests are contra-indicated in pregnancy [69].

Adequate glucocorticoid substitution is important, as is careful monitoring to prevent over-substitution. The standard substitution dosage usually does not have to be increased in the first and second trimester of pregnancy but sometimes in the third trimester due to the increase in binding globulins [63]. It is essential that the expectant mother is trained to recognize the signs of adrenal crisis, is prescribed emergency medication and given an emergency card and that information is sent in writing to the obstetrician who will attend the birth. During the birth, the patient should receive a 50 mg bolus of hydrocortisone, followed by continuous infusion of 100 to 200 mg/24 hours or 50 mg every 6 hours [63]. The dosage can then be reduced to the normal substitution dosage 1–2 days after giving birth. Medically, there is usually no reason for the birth to be by C-section rather than by vaginal delivery. Post partum the neonate must be monitored for signs of adrenocortical insufficiency.

Growth hormone deficiency

Patients with anterior pituitary insufficiency and growth hormone deficiency often require assisted pregnancy procedures [70]. There are some indications that growth hormone substitution can improve patients’ likelihood of conceiving. Once the pregnancy has been confirmed, growth hormone substitution can usually be terminated [70]. The results of a number of studies have been confirmed, growth hormone substitution can usually be terminated [70]. The results of a number of studies have been confirmed, growth hormone substitution can usually be terminated [70].
shown that growth hormone substitution is not required during pregnancy [71]. An alternative to immediately and completely ceasing substitution is to continue substitution, with the dosages adjusted to mimic physiological conditions. In such cases, substitution doses are reduced in the second trimester, and discontinued in the third trimester of pregnancy in parallel to the increases in placental growth hormone secretion [70].

Diabetes insipidus

Transient diabetes insipidus during pregnancy is primarily caused by increased placental vasopressinase activity and a higher rate of vasopressin degradation in the late second trimester and early third trimester of pregnancy [72]. Because of the greater placental mass, the risk is even higher in multiple pregnancies. Some studies have also reported an association with preeclampsia [73].

Exacerbation or demasking of pre-existing central diabetes insipidus is far rarer [74]. In such cases, symptoms occur even before peak vasopressinase activity is reached. Patients with pituitary disease who have an increased risk for diabetes insipidus and patients with pre-existing diabetes insipidus should be informed about the expected physiological changes and encouraged to regularly monitor how much they drink and how much they excrete. Diagnostic tests are primarily based on the measurement of serum/urine osmolality and the measurement of natrium levels. Pregnant women should not have a fluid deprivation test because of the risk of hypernatremia and dehydration. A lack of vasopressin is not necessarily associated with oxytocin deficiency [75, 76].

Therapy consists of the vasopressin V2 receptor agonist desmopressin (1-deamino-8-D-arginine vasopressin, DDAVP), usually administered intranasally in the evening. Dosages are usually higher than for non-pregnant patients [77]. After the patient has given birth, the administration of DDAVP to patients with transient diabetes insipidus can be discontinued after a few days or weeks; for patients with pre-existing disease, dosages should be reduced to the doses prescribed prior to pregnancy. A systematic review of 53 cases treated with DDAVP found no serious side effects [78]. The review also found no indications of adverse effects on mother or child when DDAVP was administered during pregnancy and the lactation period. The amount of DDAVP which can be transmitted to the infant through breastmilk is too low to affect the child’s diuresis.

Oxytocin deficiency

A recently published study suggested that patients with central diabetes insipidus have lower oxytocin levels than controls and that this could be associated with a reduced empathic ability [79]. But this was only investigated in a relatively small, mixed male and female population and never in pregnant patients.

The role of oxytocin in inducing birth in patients with pituitary insufficiency is controversial, and the literature and experience on this point are unfortunately very limited. Mice deficient in oxytocin had normal, spontaneous parturition [80]. In humans, most case reports and case series also report that normal spontaneous birth can occur even in mothers with complete hypophyseal insufficiency [55, 81] as it is possible that uterine oxytocin may be responsible for inducing parturition.

Fertility and pituitary disease

With careful planning and medical support prior to pregnancy, many couples with pituitary disease can fulfil their wish to have children [82]. Spontaneous pregnancy occurs in around half of affected couples.

Before starting assisted reproduction, affected couples should first wait to see whether spontaneous conception is possible. Men can still be fertile even if they present with small testicular volume (less than 3 ml) and a sperm count of less than 10 million/ml [83]. During fertility treatment, gonadotropins (i.e., hCG and recombinant follicle-stimulating hormone [rFSH]) are usually administered to men with hypogonadotropic hypogonadism [88, 89].

Pulsatile GnRH treatment is also a potential option to treat men with hypothalamic deficits. The first spermiogram should usually be done at 6 months after starting therapy, or even 4 months after starting therapy in patients who acquired their hypogonadism after puberty. A first stimulation carried out quite early after the end of puberty would probably lead to significantly faster sperm maturation at a time when the patient wishes to have a child.

The administration of gonadotropins achieves ovulation in most women with hypogonadotropic hypogonadism. Luteinizing hormone (LH) can be used as an alternative to hCG to support FSH-induced follicle maturation [84, 85]. Growth hormone deficiency can impair stimulation [1, 86]. If the patient also has partial impairment of somatotropic functions, growth hormone substitution can increase the probability of conception [87]. An important aspect of successful therapy is that men/women are cared for by

| Substance | Administration | Dosages |
|-----------|----------------|---------|
| Human chorionic gonadotropin (hCG), e.g. Brevactid® | SC or IM | 1500 IU, 2 to 3 times per week (prepubertal patients should receive higher doses in the first few months) After 2 months, titrate to achieve mean testosterone levels without a significant increase in estradiol. |
| Highly purified or recombinant FSH (e.g. Puregon®, Gonadotrope) | SC or IM | 150 IU, 3 times per week, usually only after 3 to 6 months of previous therapy with hCG |

FSH: follicle-stimulating hormone, hCG: human chorionic gonadotropin

Table 3 Gonadotropin substitution therapy to induce spermatogenesis and maintain androgenization in men [88, 89].
an experienced endocrinologist/andrologist/gynecologist who is familiar with such problems.

Conclusions

- In patients with small pituitary tumors, observant management during pregnancy with close monitoring is appropriate, although treatment with antiserotony drugs should be discontinued. Patients with large or symptomatically growing tumors and/or tumors with high disease activity require individually tailored therapies including neurosurgery and/or drugs.
- Prolactinoma therapy with dopamine agonists should be interrupted as soon as the pregnancy is confirmed. Microprolactinomas rarely increase significantly in size if dopamine agonist therapy is discontinued, although symptomatic tumor growth can occur with macroadenomas. In such cases, dopamine agonist administration can be resumed; patients with a large primary tumor and a significantly increased risk of tumor growth can be given a dopamine agonist from the start of pregnancy, if necessary.
- When treating pregnant women with secondary hypothyroidism, l-thyroxine substitution should be guided by the thyroid hormone changes (fT4) in normal pregnancies.
- In patients with adrenocortical insufficiency, cortisol substitution doses may sometimes need to be increased in the second half of pregnancy.
- Growth hormone substitution can usually be paused during pregnancy in patients with growth hormone deficits.
- If diabetes insipidus develops during pregnancy, treatment with DDAVP is indicated. Depending on the etiology of the diabetes insipidus, DDAVP may be gradually discontinued after the end of pregnancy or reduced to the dosage given to the patient prior to pregnancy.

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Conflict of Interest

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