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Opioids and endocrine dysfunction

Oliver Seyfried and Joan Hester

Abstract
The endocrine effects of opioids used for the management of persistent pain are poorly understood by clinicians and patients, and hormone levels are rarely measured. It is recognized that opioids exert this effect via the hypothalamic-pituitary-gonadal axis. Additional effects on adrenal hormones, weight, blood pressure and bone density may also occur. Symptoms and signs of sex hormone deficiency occur in both men and women but are under-reported and are often clinically unrecognized. The potential effects of long term opioid therapy on the endocrine system should be explained to patients before opioid therapy is commenced. Monitoring of sex hormones is recommended; if there are deficiencies opioids should be tapered and withdrawn, if this is clinically acceptable. If opioid therapy has to continue, hormone replacement therapy should be initiated and monitored by an endocrinologist.

Keywords
Opioids, endocrine, sex hormones, hypogonadism, testosterone, hormone replacement
It is imperative that all those managing and prescribing opioids for persistent pain are aware of the long-term effects of opioids on endocrine function and are able to diagnose deficiencies, monitor hormone levels, understand and explain to the patient the possible consequences of these deficiencies, strive to taper and withdraw opioids when appropriate and liaise with other clinicians to replace hormone deficiencies.

The endocrine effects of opioids

Hypogonadism

In 1895 Reverend RH Graves noted how ‘opium ate out the virility of the individual’ and in 1925 Surgeon General HS Cumming stated ‘opium makes a man effeminate’. Katz has previously quoted Charles Bruce, who, in 1839, referred to the opium addicts in Assam as ‘more effeminate than women’.

It has been reported on many occasions that opioids, given by any route, suppress the hypothalamic–pituitary–gonadal axis and have a measurable impact on gonadal function.

Physiology of opioid–endocrine interactions

The hypothalamus is central to the regulation of sex hormones. It exerts its control via the secretion of gonadotrophin-releasing hormone (GnRH) from the preoptic area into the hypophyseal portal circulation at the median eminence. GnRH stimulates the release of follicle-stimulating hormone (FSH) and luteinising hormone (LH) from the anterior pituitary by activating its own GnRH receptor, which, via increased levels of calcium and protein kinase C, leads to the formation and secretion of FSH and LH (Figure 1).

GnRH is normally released in bursts in all studied vertebrates. These pulses produce circadian peaks which are essential for the correct functioning of the reproductive system by determining when each hormone is released. Non-pulsatile secretion of GnRH results in downregulation of the pituitary and leads to failed LH secretion.

FSH is responsible for the early growth of ovarian follicles in females and maintenance of the spermatogenic epithelium in males. LH is responsible for the final maturation of follicles and their oestrogen secretion, as well as ovulation and the initial formation of the corpus luteum in the female. In the male it stimulates Leydig cells to secrete testosterone. These hormones are also required for the appropriate development of the human – physiologically, physically and socially.

The hypothalamic–pituitary axis is constantly under the effect of multiple substances including neurotransmitters, steroid hormones and endogenous opioids. Exogenous opioids exert an effect on the same receptors as endogenous opioids and have been shown to interfere with the release (including its pulsatile nature) of GnRH. Naloxone showed increased GnRH levels, and thereby increased LH concentration and pulse frequency, which suggested a basal level of opioid-based inhibition of LH secretion. It has been suggested that morphine inhibits the biosynthesis of GnRH. Opioids also decrease the negative feedback

Figure 1. Representation of the normal female and male hypothalamic–pituitary–gonadal axis. (Reproduced with permission from NIAAA).
of sex steroids on the anterior pituitary, as well as its response to GnRH. In contrast, FSH secretion is not, or only minimally, affected.

With the reduction in LH levels, testosterone and oestradiol are commensurately lowered. Li Shizhen wrote of opium in his *Compendium of Materia Medica* (1578) that ‘lay people use it for the art of sex, particularly to “arrest seminal emission”’11. Animal studies have added credence to Shizhen’s observation, demonstrating inhibited sexual receptivity in both sexes of rats with opioids and the converse with naloxone.12 This appears to be due to decreased arousal rather than impotence. Alarmingly, related studies showed that prepubertal opioid exposure inhibited sexual maturation.13 In humans, menstrual irregularities, including amenorrhoea, occurred following treatment with opioids via the oral and intrathecal routes. Daniell14 noted that there was also a decrease in adrenal androgens, and thereby explaining the decreased libido and sexual performance so often encountered in those exposed to long-term opioids. It appears that decreased sexual behaviour may also be due to the direct action of opioids on μ and δ receptors in the hypothalamus.15,16

The term ‘sex hormones’ encompasses the sex steroids (testosterone and oestradiol) and the non-steroid hormones (GnRH, FSH and LH).

Testosterone is the principal hormone of the testes, synthesised from cholesterol in Leydig cells and from androstenedione in the adrenal cortex. Its secretion is controlled by LH at 4–9 mg/day. Small amounts are secreted by the ovaries and possibly the adrenal cortex in women. It is 98% protein bound (gonadal steroid-binding globulin and albumin) in the plasma. Only the free and weakly albumin-bound testosterone is available to act on androgen receptors. Some target cells convert testosterone to dihydrotestosterone, which forms more stable hormone–receptor complexes. In males testosterone has a circadian rhythm, with the highest levels in the morning. Maximal variation may be in the order of 35%.

Testosterone imparts a negative feedback on LH release from the pituitary. It develops and maintains the male secondary sex characteristics and encourages male sexual behaviours. It is anabolic, increases rate of growth and, along with FSH, promotes spermatogenesis. It does all of this by binding to intracellular receptors and forming complexes that bind to DNA, and thus facilitating certain gene expression.

Oestrogens (17ß-oestradiol, oestrene and oestriol) are the primary female sex steroids and are biosynthesised from testosterone and androstenedione. They are secreted by the granulosa cells in ovarian follicles. Ninety-eight per cent of oestrogens are protein bound. They are metabolised by the liver and excreted in the urine. Oestrogens facilitate the development of ovarian follicles, aid in the regulation of the menstrual cycle and the requisite changes in female internal anatomy, and have anabolic effects on the uterus and fallopian tubes. They decrease FSH levels and alter LH levels. They are responsible for female sexual behaviours and libido and are largely responsible for breast development. These effects, combined with the absence of androgens, lead to female secondary sexual characteristics. Oestrogen, being a steroid hormone, exerts its effects via two principal intracellular receptors.

In summary, opioids lead to decreased secretion of GnRH, which in turn leads to reduced levels of LH. This results in decreased testosterone and oestradiol secretion, which leads to the signs and symptoms listed

### Table 1. Clinical effects of reduced sex hormone secretion.

<table>
<thead>
<tr>
<th>Male</th>
<th>Female</th>
<th>Males and females</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reduced testicular volume</td>
<td>Menstrual irregularities</td>
<td>Infertility</td>
</tr>
<tr>
<td>Erectile dysfunction</td>
<td>Reduced breast size</td>
<td>Reduced libido</td>
</tr>
<tr>
<td>Loss of muscle mass</td>
<td>Hot flushes</td>
<td>Depression and anxiety</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Glucose intolerance</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Hypercholesteremia</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Gynaecomastia</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Polyuria</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Diarrhoea</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Night sweats</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Fatigue</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Osteoporosis</td>
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<td></td>
<td></td>
<td>Anosmia</td>
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<td></td>
<td></td>
<td>Myalgia</td>
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<tr>
<td></td>
<td></td>
<td>Galactorrhoea</td>
</tr>
</tbody>
</table>

Glucose intolerance; Hypercholesteremia; Osteoporosis; Anosmia; Myalgia; Galactorrhoea.
Diagnosis of hypogonadism in the male

The diagnosis of hypogonadism is dependent on history and examination in conjunction with laboratory testing. In the postpubertal male the potential causes of primary hypogonadism mentioned in Table 3 can cause diagnostic difficulty, especially with regard to their gradual onset. If hypogonadism is suspected, it is therefore important to differentiate primary gonadal failure from that of the hypothalamic–pituitary axis.

According to the World Health Organization’s Second International Consultation on Erectile Dysfunction, which considered the circadian rhythm and pulsatile nature of testosterone secretion, two blood samples should be taken between 8.00 a.m. and 11.00 a.m. (when testosterone levels are presumed to be at their highest, although the circadian rhythm can diminish with increasing age). Samples should be sent for measurement of serum testosterone, sex hormone-binding globulin (SHBG), prolactin, FSH and LH levels.

Normal to high FSH/LH levels might indicate primary hypogonadism (see Table 3). It is important to measure the level of FSH as it has a longer half-life and demonstrates less variability than LH. Secondary hypogonadism (Table 4) is indicated by low testosterone and normal to low FSH/LH levels (Box 1).
Box 1: According to the Endocrine Society of Australia Consensus Guidelines\textsuperscript{17}.

<table>
<thead>
<tr>
<th>Condition</th>
<th>Testosterone Levels</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary hypogonadism</td>
<td>$&lt; 231 \text{ng/dl with LH} &gt; 1.5 \text{ times the upper limit of normal (1.5 x ULN)}$</td>
</tr>
<tr>
<td>Secondary hypogonadism</td>
<td>$&lt; 231 \text{ng/dl without elevation of LH}$</td>
</tr>
</tbody>
</table>

With ageing comes diminished diurnal fluctuation in serum testosterone. The level falls markedly throughout the day, reinforcing the importance of early morning sampling. In hypogonadism in this age group total testosterone levels may be normal if sex hormone-binding globulin (SHBG) levels are raised. SHBG levels increase with age, and thereby decrease the bioavailability of testosterone.

Recent data from the Massachusetts Male Ageing Study (MMAS) provide perspectives on normal androgen ranges, as shown in Table 1.

**Diagnosis of hypogonadism in the female**

Hypogonadism in females can similarly be due to a hypothalamic–pituitary axis or primary gonadal defect. The signs and symptoms are closely linked to the menstrual cycle in the postmenarche and premenopausal years. Common signs include oligomenorrhea, amenorrhoea and failure to conceive. More subtle symptoms such as hot flushes and anxiety may occur, rarely with changes in pubic hair distribution and breast size\textsuperscript{21}. Although the clinical effects can be as severe for women as for men, the outward or visible changes are not so obvious and have not been studied in any detail, especially in postmenopausal women.

Although androgen deficiency has been shown to be symptomatic in females using intrathecal opioid therapy and those on maintenance methadone, testosterone levels have not been routinely measured. Dihydroepiandrosterone (DHEA) is thought to be lowered and it is known that LH levels are markedly reduced. DHEA is a marker of adrenal androgen production, and approximately 50% of androgens produced in the female are adrenal in origin. There is a paucity of data with regards to hormone levels in females, and further studies are needed to quantify the clinical significance of this potentially very interesting aspect of opioid-induced androgen deficiency (OPIAD).

**Effect of opioids on adrenal hormones**

Acute and chronic pain, as physiological responses, result in increased secretion of adrenocorticotropic hormone (ACTH) and cortisol. However, the chronic use of exogenous opioids has been found in several studies to decrease ACTH and cortisol levels and cortisol responses to adrenocorticotropic challenges.\textsuperscript{19} Opioids also affect the circadian rhythms of cortisol secretion, resulting in persistently raised levels of ACTH and cortisol and eventually blunting the stress response.\textsuperscript{20} Levels of dehydroepiandrosterone sulphate (DHEAS), a precursor of adrenal androgens, have also been markedly reduced in both male and female chronic opioid users.\textsuperscript{14,18} Occasionally, opioid administration may cause frank adrenal insufficiency, but the risk factors for this are unknown at present.\textsuperscript{15}

**Prolactin**

Acute administration of opioids stimulates prolactin release from the anterior pituitary through an effect at the hypothalamus.\textsuperscript{15} This effect can be blocked by metoclopramide, suggesting that it is mediated through dopaminergic mechanisms. The effect of long-term administration of opioids on prolactin is less clear. There is an occasional increase in prolactin release, possibly dependent on the type of opioid. The clinical significance of this is unknown, but it can cause galactorrhoea.

**Thyroid hormones**

In general opioids do not appear to alter thyroid function in any meaningful way,\textsuperscript{19} though opioids can stimulate thyroid-stimulating hormone (TSH) via the hypothalamus.\textsuperscript{15} This is unimportant in individuals with normal free thyroxine, but individuals with hypothyroidism may have prolonged and exaggerated responses to opioids.\textsuperscript{22}

**Growth hormone**

Acute administration of opioids leads to increased growth hormone (GH) secretion, through mechanisms involving opioid receptors, feedback levels and gene transcription. The minimum dose required is approximately 15 mg morphine.\textsuperscript{15} Abs et al.\textsuperscript{23} found GH deficiency in approximately 15% of patients receiving long-term intrathecal opioids, but not in all patients. Naloxone has been shown to inhibit GH in healthy subjects, but to increase it in obese women. The effect of chronic opioid administration on GH is complex and currently poorly understood, but it appears to be related to sex hormones, body composition and degree of insulin resistance.\textsuperscript{15}

**Vasopressin**

Tramadol has been found to cause hyponatraemia through serotonin-induced release of vasopressin.\textsuperscript{24}
effect of opioids on the posterior pituitary is unclear: both increased and decreased levels of vasopressin have been found, depending on hydration status.22

Oxytocin
Studies on pregnant women have shown that morphine inhibits oxytocin production in the early stages of labour and during breastfeeding post partum.25

Obesity and diabetes
An increasing number of data suggests that opioids play a role in regulating food intake and food choice, and perhaps the reward associated with good-tasting foods, via central mechanisms.15 Chronic opioid use is associated with weight gain, hyperglycemia and worsening diabetes. This may be a central action via the sympathetic nervous system and impaired insulin secretion.26 Hypogonadism is associated with increase insulin resistance and risk for diabetes mellitus,15 a risk that is improved by testosterone replacement.27

Catecholamine metabolism
Opioid therapy increases catecholamine secretion via the hypothalamus and brain stem. Patients on long-term opioids should be screened for hypertension.22

Bone metabolism
There are many risk factors for decreased bone mineral density and osteoporosis in patients treated with opioids, including possible poor nutritional status, hypogonadism, inhibition of osteoblasts, decreased osteocalcin synthesis, abnormal calcium and parathyroid hormone and increased bone resorption, mediated by interleukin 1. There is an increased risk of bone fracture in patients using opioids.3

Route of administration
Opioids for chronic use are administered orally or intrathecally. Other routes, for example repeated injection, may occasionally be used, though this practice is not recommended.28

Hormonal changes occurring after intrathecal administration were reported by several authors.23,29,30 It is estimated that 90% of patients receiving intrathecal opioids will develop hypogonadism. Oral opioids have the same effect, though the onset of action may be slower.

Type and dose of opioid
Classes of opioids differ in their effect on gonadal suppression. Tramadol and buprenorphine do not significantly alter testosterone levels in animals and humans, and buprenorphine does not suppress serum cortisol.

The incidence of hypogonadism was higher in cancer survivors receiving a dose equivalent to or more than 200 mg morphine per day for at least 1 year compared with age-matched survivors not on opioid therapy, suggesting that the effects are dose related.31 Duration of opioid therapy also seems to increase the chance of hormonal suppression, though this needs to be studied further.

There is a possible gender difference in endocrine effects, biased towards more symptomatology in women, but further studies are needed.22

When and what to measure
Before commencing chronic opioid therapy it is recommended that the following are measured:

- blood pressure;
- electrolytes (especially if tramadol is used);
- fasting glucose levels;
- thyroid function (to exclude hypothyroidism);
- testosterone, sex-binding globulin, LH/FSH and oestradiol levels; and
- bone density (in an ‘at-risk’ group).

Monitoring
There are no accepted standards, but it seems reasonable to repeat the above tests every 6 months.

- Consider morning fasting blood cortisol, DHEA, ACTH and GH levels. (Remember that an abnormally high fasting blood cortisol level can represent loss of diurnal variation, and advice should be sought.)
- Repeat bone density yearly in ‘at-risk’ group.
- Measure prolactin levels if there is galactorrhoea.

Replacement therapy
There are no accepted standards for the management of opioid-induced endocrine dysfunction.

The best option is to taper and withdraw opioids and monitor the response over a period of a few months if this is appropriate. It is not known if switching opioids is of any benefit. Buprenorphine seems to have less effect on adrenal hormones but has a greater effect on TSH than morphine. Response to different opioids is largely unknown at the time of writing.

If opioid withdrawal cannot be achieved, and the patient has definite symptoms of endocrine dysfunction,
hormone replacement is recommended, with monitoring by an endocrinologist.

Testosterone can be replaced, in both men and women, as a transdermal patch or gel or by injection. Careful monitoring is required as side-effects include site reactions, polycythemia and increased risk of prostate cancer in men and menstrual irregularities and hirsutism in women.

Oestrogen replacement therapy is best monitored by a gynaecologist.

Conclusion

The use of opioids for the management of pain continues to increase and therapy that involves these drugs has significant adverse effects in both the short and long term. Both patients and prescribing physicians should be aware, or made aware, of them. Careful risk–benefit analysis should be applied and all possible opioid-sparing manoeuvres considered.

The effects of opioids on the endocrine system are becoming increasingly apparent, although more research is required to further elucidate the mechanisms by which these happen. It affects both sexes, but the clinical signs are more obvious in males. Direct action of opioids on the hypothalamus reduces the release of GnRH, and thereby adversely affects LH levels, and subsequently testosterone synthesis and secretion.

Symptoms include infertility, decreased sexual function, loss of muscle mass and anxiety and/or depression. Osteoporosis may result from long-term opioid therapy, leading to fractures and their associated morbidity and cost.

More research is required to elucidate the aetiology of opioid-induced endocrine dysfunction. Differences between methods of opioid delivery, the effects of OPIAD in both sexes and the potential value of opioid rotation need further investigation.

The potential long-term effects of opioids on the endocrine system should be explained to the patient before therapy is commenced and regular monitoring should be performed.

The use of opioid therapy should be minimised by implementing a biopsychosocial approach to pain management.

The benefits of opioid therapy should be carefully assessed; if therapeutic goals have not been achieved, opioids should be tapered and withdrawn.

If it is necessary to continue opioids, hormone replacement therapy should be initiated and monitored by an endocrinologist.

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Conflict of interest

The authors declare that they do not have any conflict of interest.

References


**Multiple choice questions (more than one answer may be correct for each question)**

1. Which serum levels of the following hormones may be reduced by taking oral opioids for more than one year?
   a) adrenocorticotropic hormone
   b) testosterone
   c) growth hormone
   d) luteinising hormone
   e) thyroid-stimulating hormone

2. Which of the following possible side-effects of taking opioids for persistent pain should be explained to women of childbearing age?
   a) greater risk of pregnancy
   b) weight gain
   c) osteoporosis
   d) amenorrhoea
   e) fatigue

3. High fasting blood cortisol level may indicate which of the following:
   a) loss of diurnal variation
   b) opioid overuse
   c) adrenal tumour
   d) stress
   e) overexercising

4. Testosterone replacement therapy:
   a) can cause secondary polycythaemia
   b) should be offered to women who have low levels of free testosterone
   c) leads to an increase in muscle mass
   d) increases the risk of prostate cancer
   e) can be safely prescribed by any doctor

5. Long-term use of opioids for non-cancer pain:
   a) is usually effective in relieving pain
   b) carries an increased risk of accidental overdose
   c) requires no monitoring
   d) may cause hypertension
   e) is known to impair cognitive function

**Answers**

1. a) True; b) True; c) False; d) True; e) False.
2. a) False; b) True; c) True; d) True; e) True.
3. a) True; b) True; c) True; d) False; e) False.
4. a) True; b) True; c) False; d) True; e) False.
5. a) False; b) True; c) False; d) True; e) True.