Case Report

Hypothalamic-Pituitary-Adrenal Suppression and Iatrogenic Cushing’s Syndrome as a Complication of Epidural Steroid Injections

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Epidural steroid injections are well accepted as a treatment for radicular low back pain. Nearly 50 million in Medicare dollars went to this treatment in 1999 [1], and from 1994 to 2001, there was a 271% increase in Medicare funded lumbar epidural steroid injections [2]. A typical epidural steroid is triamcinolone in 40 mg or 80 mg doses. Some expert recommendations promote giving up to 3 injections within a year with a minimum of 30 days between injections. Further or more frequent injections are not recommended due to concerns about hypothalamic-pituitary-adrenal suppression [3]. Other practice guidelines provide no recommendations on maximum injection number or dosing interval [4]. The American Society of Interventional Pain Physicians-Interventional Pain Management comments in their 2009 guidelines that there is no “basis for reported assumptions and limitations” on epidural steroid injection doses or frequency and furthermore, that “administration must be based solely on patients’ response, safety profile of the drug, experience of the patient, and pharmacologic and chemical properties such as duration of action and suppression of adrenals [5].” However, they go on to recommend epidural steroid injections be given no more frequently than every 1-2 weeks during the diagnostic phase and thereafter no more frequently than every 2 months.

2. Case

The patient is a 53-year-old Caucasian woman of Northern European ancestry referred to Endocrinology by a specialty Pain Clinic for concerns of possible hypercortisolism. She had been seen by the pain specialist for chronic radicular back pain and was given 3 doses of fluoroscopically guided epidural triamcinolone 80 mg via the caudal approach over the course of 4 months. Four weeks elapsed between her first and second doses, 10 weeks between her second and third, her last dose having been given 7 weeks prior to our initial evaluation. She noticed bloating and a buffalo hump between her second and third doses. She then noted progressive muscle weakness, difficulty rising from sitting to standing, worsening in her baseline insomnia, loss of hair on her head,
Table 1: Laboratory evaluation pertinent to our initial evaluation (normal values for our clinical lab in parentheses).

<table>
<thead>
<tr>
<th>Time following 3rd epidural injection (time of day for lab draw)</th>
<th>3 wks</th>
<th>7 wks</th>
<th>11 wks</th>
<th>15 wks</th>
</tr>
</thead>
<tbody>
<tr>
<td>AM serum cortisol</td>
<td>1505</td>
<td>63.5 nmol/L (2.3 mcg/dL)</td>
<td>80.0 nmol/L (2.9 mcg/dL)</td>
<td>63.5 nmol/L (2.3 mcg/dL)</td>
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<tr>
<td>118.6–618 nmol/L (4.3–22.4 mcg/dL)</td>
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<tr>
<td>PM serum cortisol</td>
<td>16.6 nmol/L (0.6 mcg/dL)</td>
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<tr>
<td>8.53–460.8 nmol/L (3.1–16.7 mcg/dL)</td>
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<tr>
<td>DHEA-S</td>
<td>Undetectable</td>
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<tr>
<td>90.2–6940 nmol/L (2.6–200 mcg/dL)</td>
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<tr>
<td>Prolactin</td>
<td>208.7 pmol/L (4.8 ng/mL)</td>
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<tr>
<td>121.7–1269.6 pmol/L (2.8–29.2 ng/mL)</td>
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<tr>
<td>Bioavailable testosterone</td>
<td>0.045 nmol/L (1.3 ng/dL)</td>
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<tr>
<td>0.052–0.326 nmol/L (1.5–9.4 ng/dL)</td>
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<tr>
<td>Free testosterone</td>
<td>0.0017 nmol/L (0.5 pg/mL)</td>
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<tr>
<td>0.0021–0.0132 nmol/L (0.6–3.8 pg/mL)</td>
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<tr>
<td>Total testosterone</td>
<td>0.07 nmol/L (2 ng/dL)</td>
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<tr>
<td>0.21–0.87 nmol/L (6–25 ng/dL)</td>
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</table>

redness of her chest, and acne. Furthermore, she stated her face had rounded and her arms and legs looked thin while her abdomen seemed to be expanding. She denied easy bruising, newly visible striae, increased tan or scar pigmentation, or other skin changes, but complained of increased hair growth on her face including thickening of her sideburns. While there was a family history of hirsutism, there was no family history of endocrinopathies. Her past medical history included chronic back pain, three uncomplicated pregnancies, regular menses until menopause at the age of 45, and long-standing hypertension. Notably, she had been told that she had “borderline diabetes” within the six months prior to her presentation. Medications included naproxen sodium, gabapentin, nortriptyline, simvastatin, hydrochlorothiazide, benazepril, calcium carbonate, multivitamin, and omega-3 fish oil.

On exam, she was obese (103.8 kg, BMI 35.9) and hypertensive (175/88 mm Hg). She was plethoric along the neck, face, and upper back. She had increased vellus facial hair. There was fat accumulation at the base of the neck. There were no striae or bruises.

Given the rapid onset of her symptoms and the complaint of hirsutism, she was evaluated for endogenous hypercortisolism and nonclassic congenital adrenal hyperplasia. A random afternoon cortisol level drawn three weeks after her last epidural injection (four weeks prior to our initial evaluation) was 16.6 nmol/L (0.6 mcg/dL. normal morning value 118.6–618 nmol/L (4.3–22.4 mcg/dL)), and repeat at 10 am four weeks later was 63.5 nmol/L (2.3 mcg/dL). Bioavailable testosterone level was low at 0.045 nmol/L (1.3 ng/dL. normal: 0.052–0.326 nmol/L (1.5–1.9 ng/dL)), and DHEA-S level was undetectable. A midnight salivary cortisol level three weeks after our initial visit was only 0.77 nmol/L (0.028 mcg/dL) (normal < 3.09 nmol/L (0.112 mcg/dL)). Prolactin level and thyroid function tests were normal. Chemistries were consistently normal without aberration in potassium, sodium, or creatinine. Laboratory data pertinent to our initial evaluation are summarized in Table 1.

Although no adrenocorticotropic hormone (ACTH) level was obtained, laboratory investigations were felt to be consistent with hypothalamic-pituitary-adrenal (HPA) suppression from exogenous steroids. The patient was empirically placed on prednisone 5 mg daily, was given instructions on stress-dosing of steroids, and was provided with a dose of injectable hydrocortisone to be used in case of emergency. At one and two months after our initial visit (2-3 months after her last epidural injection), the prednisone was held for 24 hours, and repeat morning cortisol levels remained low at 80.0 and 63.5 nmol/L (2.9 and 2.3 mcg/dL), respectively (Table 1). The patient underwent a cosyntropin stimulation test four months after her last epidural injection, which showed a suboptimal rise from 209.7 to 389 nmol/L at 60 minutes (7.6 to 14.1 mcg/dL). A random 17-hydroxyprogesterone was not elevated at 0.21 nmol/L (7 ng/dL). Shortly thereafter, the patient underwent an outpatient rotator cuff repair. She was instructed to increase her prednisone to 15 mg the day of her surgery and to return to 5 mg daily afterwards. Her surgery and postoperative period were uneventful.

Given her morning cortisol level remained suppressed, the patient’s prednisone dose was tapered slightly to 4 mg daily. One month later (six months after her last epidural injection), her cortisol level was normal at 328.3 nmol/L (11.9 mcg/dL) at 0935. The same day, she also demonstrated an appropriate response to cosyntropin with cortisol levels rising to 485.6 and 554.6 nmol/L (17.6 and 20.1 mcg/dL) at 30 and 60 minutes after stimulation, respectively. During this
3. Discussion

Case reports and some experimental data suggest that adrenal disorders can result from epidural steroid injection. Patients can develop isolated HPA suppression, or HPA axis suppression can be accompanied by overt iatrogenic Cushing’s. The potential for HPA suppression has been clearly documented and is referenced briefly by expert recommendations [3]. Meanwhile, the potential for accompanying iatrogenic Cushing’s Syndrome is less well recognized and goes unmentioned in any guidelines. Moreover, Jacobs et al. assert that the lack of a detectable plasma methylprednisolone level after epidural injection in 12 patients indicated that systemic absorption does not occur, and therefore, patients are unlikely to have systemic side effects [6]. Previously published case reports of patients presenting with systemic symptoms, including the present case, do not support this assertion. In addition to the present case, five other case reports of biochemically proven adrenal suppression in patients with Cushing’s Syndrome have been reported [7–11], and another four cases of patients with Cushing’s features were reported in 1980 by Knight and Burnell, but no biochemical data was provided [12]. Case reports by Stambough et al. [7] and Horani and Silverberg [8] described patients presenting with moon facies, supraclavicular fat pad, buffalo hump, bruising, peripheral edema, abdominal striae, and hypertension. In addition to similar common manifestations of hypercortisolism, the patient presented by Boonen et al. suffered from significant steroid myopathy [11]. Finally, Lansang et al. performed urine screens for synthetic glucocorticoids in their patient who had received epidural triamcinolone injections, and were able to detect triamcinolone in the patient’s urine sixty-two days after her last injection, establishing biochemical evidence of systemic absorption [10].

The risk of iatrogenic Cushing’s with HPA axis suppression in an individual patient receiving epidural steroid injections is unknown. The occurrence of this serious side effect has been reported in only a small number of cases, and a detailed review of the literature by Chou et al. found that many of the randomized placebo-controlled trials of epidural steroids did not report harm overall, and certainly not for rarer side effects [13]. Furthermore, only 5 randomized trials included more than 100 patients limiting ability to capture rare events. One could deduce that the risk for iatrogenic Cushing’s is relatively low given the small number of case reports and the widespread use of epidural steroid injections, but the true incidence is unknown.

Meanwhile, the risk of adrenal suppression without overt iatrogenic Cushing’s has been more clearly established. As early as 1974 [14], Burn and Langdon showed plasma cortisol levels in 50 of 56 patients to be depressed for 1-2 weeks after an epidural injection of 80 mg methylprednisolone. In 1983, Jacobs et al. [6] further studied the potential for one extradural injection of 80 mg prednisolone acetate, approximately 1mg/kg body weight, to affect the HPA axis. After extradural drug administration, they found that patients’ adrenals had a diminished ability to respond to ACTH compared to pretreatment responses despite minimally detectable drug levels in the systemic circulation. They also found evidence of central suppression in morning plasma cortisol and ACTH levels that did not resolve to pretreatment levels at 3 weeks after one injection. Similar to the Burn and Langdon study, suppression of the axis was most marked one week after the extradural steroid injection, with gradual recovery thereafter. In the case presented by Horani and Silverberg [8], the patient showed evidence of adrenal suppression over two months after her one and only epidural glucocorticoid injection. The patient was given replacement hydrocortisone 20 mg orally once daily and without dose adjustments or tapering of this dose, her HPA axis was still abnormal at eight months, but recovered by ACTH stimulation testing at twelve months.

The risk for adrenal suppression is likely higher in patients who have received serial epidural steroid injections as in our case and that of Lansang et al. [10]. This scenario was evaluated in a series by Kay et al. [15] wherein they showed that 5 of 14 patients who received 80 mg triamcinolone epidural injections weekly for 3 weeks had a subnormal response to cosyntropin one month after the last epidural injection.

4. Conclusion

Hypothalamic-pituitary-adrenal axis suppression due to epidural steroid administration has been established as a potential complication of treatment. Concomitant iatrogenic Cushing’s Syndrome is less common, but suggestive physical symptoms or signs should prompt referral to an endocrinologist for further testing and management. Epidural steroid injections should be held in the setting of HPA suppression even if overt Cushing’s is not present.

Given the regular use of epidural steroid treatment for chronic lumbar radicular pain, physicians administering the treatment must be aware of the potential for suppression of the HPA axis that could indicate an increased risk for adrenal crisis in the setting of stress. Some have advocated evaluation of the HPA axis via baseline cortisol levels and stimulation testing or empiric stress-dosed steroids for any patient undergoing a surgical procedure within three weeks of an epidural steroid injection [14, 15]. The case report evidence shows that some patients have suppression far beyond three weeks, supporting the use of formal testing of HPA status prior to surgery as the safest route. Conversely, it is notable that despite evidence of routine HPA suppression, there are no case reports of adrenal crisis occurring in patients with recent epidural steroid exposure. Perhaps a prudent approach would be to perform routine screening for symptoms of adrenal insufficiency in all patients who have received epidural steroids in the preceding year and to perform preoperative testing in all patients less than three weeks out from an epidural injection, those who have symptoms suspicious of...
adrenal insufficiency, and in those with a history of multiple injections in the preceding 4–6 months.

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References


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