GUIDELINES AND GUIDANCE

SOCiETY FOR ENDOCRINOLOGY

ENDOCRINE EMERGENCY GUIDANCE

Acute management of the endocrine complications of checkpoint inhibitor therapy

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Abstract

Immunotherapy treatment with checkpoint inhibitors (CPI) (CTLA-4 and PD-1 inhibitors) significantly improves survival in a number of cancers. Treatment can be limited by immune-mediated adverse effects including endocrinopathies such as hypophysitis, adrenalitis, thyroiditis and diabetes mellitus. If endocrinopathies (particularly hypocortisolemia) are not recognized early, they can be fatal. The diagnosis and management of endocrinopathies can be complicated by simultaneous multi-organ immune adverse effects. Here, we present Endocrine Emergency Guidance for the acute management of the endocrine complications of checkpoint inhibitor therapy, the first specialty-specific guidance with Endocrinology, Oncology and Acute Medicine input and endorsed by the Society for Endocrinology Clinical Committee. We present algorithms for management: endocrine assessment and management of patients in the first 24 hours who present life-threateningly unwell (CTCAE grade 3–4) and the appropriate management of mild-moderately unwell patients (CTCAE grade 1–2) presenting with features compatible with an endocrinopathy. Other important considerations in relation to hypophysitis and the maintenance of glucocorticoid therapy are discussed.

Introduction

Immunotherapy treatment with checkpoint inhibitors (CPI) such as ipilimumab (CTLA-4 inhibitor), nivolumab and pembrolizumab (PD-1 inhibitors) significantly improves prognosis in a number of cancers (1, 2, 3). Combination therapy with ipilimumab and nivolumab is approved in the United Kingdom for the treatment of advanced melanoma but indications for immunotherapy, the cancers that benefit and the
number of agents available are increasing. However, treatment can be limited by immune-mediated adverse effects particularly with combination treatment (3, 4, 5, 6).

Immune-mediated endocrinopathies as a consequence of treatment with checkpoint inhibitors include hypophysitis, adrenalitis, thyroiditis and diabetes mellitus (7, 8, 9, 10, 11, 12, 13, 14, 15). These can be life-threatening if not recognised and treated appropriately; deaths have been reported.

Diagnosis and management in this group can be complicated by simultaneous multi-organ immune adverse effects, e.g. presentation with colitis and hypophysitis.

Early recognition and appropriate management of these endocrinopathies is essential. Multiple, informative review articles have been published with regards to the mechanisms, incidence and screening strategies. While endocrinologists and oncologists may be familiar with the complications of CPI treatment, these patients frequently present as emergencies to those unfamiliar with these agents. This guidance has been developed as an expert consensus between endocrinologists, oncologists and an acute physician and is designed to aid the early phase of care.

This document therefore covers:

- Endocrine assessment (first 24 h) of patients treated with CPI’s who present life-threateningly unwell (CTCAE (Common Terminology Criteria for Adverse Events) grade 3–4: Algorithm 1).
- Appropriate management of a mild-to-moderately unwell patient presenting with clinical features compatible with an endocrinopathy (CTCAE grade 1–2: Algorithms 2 and 3).
- Other important considerations; hypophysitis and maintenance glucocorticoid therapy.

Management of a life-threateningly unwell patient (CTCAE grade 3–4)

Cortisol

- Features of acute cortisol deficiency may be non-specific. Any patient receiving a CPI who presents severely unwell should be assumed to have acute cortisol deficiency until proven otherwise and treated with glucocorticoids until serum cortisol result available (20; https://doi.org/10.1530/EC-16-0054) (Algorithm 1).

- In the acute setting, primary (e.g. caused by adrenalitis) and secondary (e.g. caused by hypophysitis) cortisol deficiency are treated identically.
- A baseline (pre-glucocorticoid treatment) serum cortisol of >450 nmol/L excludes cortisol deficiency (for exceptions see clinical considerations in Algorithm 1), and glucocorticoid treatment can be discontinued at this point if this is the only indication. If there is any doubt about the presence of cortisol deficiency glucocorticoids should be continued and an endocrine opinion sought. It is crucial to obtain a good drug history with regards to recent glucocorticoid use to enable correct interpretation of results.
- Methylprednisolone is not an appropriate treatment for acute cortisol deficiency secondary to hypophysitis or adrenalitis (16). Methylprednisolone may be beneficial for pressure effects such as optic chiasm compromise, visual field defects, cranial nerve palsies and in some cases, intractable headache. If methylprednisolone or other pharmacological dose glucocorticoids are administered for this or other non-endocrine immune complications, additional hydrocortisone is not required.
- If significant polyuria, polydipsia and/or hypernatremia occurs following glucocorticoid replacement; consider the possibility of diabetes insipidus. Seek urgent specialist/endocrine input.
- In view of the multiplicity of immune adverse events seen with CPI’s if there is not a significant improvement once cortisol deficiency has been corrected over the first 24 h, then additional diagnoses must also be explored.

Thyroid dysfunction

It is rare for acute CPI thyroiditis to cause a patient to be life-threateningly unwell although one potential case of ‘thyroid storm’ and one of ‘myxedema’ have been reported (17, 18) (Algorithm 3).

- If severe thyrotoxicosis or thyroid storm features are present, we recommend supportive management in a critical care setting and endocrine input (19).
- If myxedema secondary to hypothyroidism is suspected, specialist endocrine input should be sought. Thyroxine should never be instigated unless cortisol deficiency is excluded as it can trigger an adrenal crisis. If in doubt, treat for cortisol deficiency first.
Management of a life-threateningly unwell (CTCAE grade 3–4) patient

Assess for the following signs/symptoms:
- hypotension: (systolic BP <90 mmHg)
- postural hypotension (>20 mmHg drop in BP from standing to sitting)
- dizziness / collapse
- hypovolemic shock
- abdominal pain, tenderness and guarding
- nausea and vomiting

• tachycardia +/- cardiac arrhythmias
• fever
• confusion/delirium
• coma
• hyponatraemia/hyperkalaemia/hypoglycaemia
• pre-renal/renal failure

Severe, potentially life threatening and possibility of hypoadrenalism: needs urgent management

Measure (alongside other acute assessment measures as indicated e.g. blood cultures):
- random serum cortisol and plasma ACTH (footnote 1)
- U+E/LFTs/CRP/FBC/SH/Hb/glucose (footnote 2)
- Prolactin, testosterone/oestradiol, LH/FSH (footnote 3)

Treat as adrenal insufficiency as per Society for Endocrinology Emergency Endocrine Guidance:
Hydrocortisone (immediate bolus injection of 100 mg hydrocortisone i.v. or i.m., followed by continuous intravenous infusion of 200 mg hydrocortisone per 24 h (alternatively 50 mg hydrocortisone per i.v. or i.m. injection every 6 h)

Random serum cortisol <450 nmol/l (footnotes 1 & 5) - stop adrenal insufficiency management
- reassess cause of signs and symptoms (footnote 6)

Random serum cortisol >450 nmol/l (footnotes 1 & 5)
- once clinically stable: convert to oral hydrocortisone (initially 20/10/10 mg to reduce to maintenance of 10/5/5 mg) or oral prednisolone (maintenance 3–5 mg per day)
- consider primary adrenal failure: assess renin/aldosterone (particularly if ACTH elevated/normal and hypotension present) (footnote 8)
- continue immunotherapy if no other contraindications

Rehydration with rapid intravenous infusion of 1000 mL of isotonic saline infusion within the first hour, followed by further intravenous rehydration as required (usually 4–6 L in 24 h; monitor for fluid overload in case of renal impairment and in elderly patients)

Footnotes:

Footnote 1: Important clinical considerations
- Review patient information for evidence of recent steroid use:
  - any supraphysiological dose of glucocorticoids can suppress the adrenal axis.
  - patients receiving doses of dexamethasone >0.75 mg or prednisolone >3 mg daily will likely have a supressed endogenous HPA axis and may have a serum cortisol measurement of <50 nmol/l. If the glucocorticoid treatment is ongoing they are not adrenally insufficient but may need higher doses of glucocorticoids when clinically unwell. Seek specialist advice from endocrinology.
  - prednisolone can cross-react in cortisol immunoassays causing spuriously elevated values. Seek specialist advice from local laboratory/endocrinologist.
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  - steroid inhalers, nasal sprays, creams and intra-articular injections can all suppress the HPA axis. Seek specialist endocrine advice if patient has been administered any of these.
  - women on oral oestrogen (hormone replacement therapy or oral contraceptive pill) will have elevated serum total cortisol levels as a result of oestrogen effect on cortisol binding globulin levels. A ‘normal’ serum cortisol level under these circumstances may be falsely reassuring. If patients are taking these preparations seek specialist advice from an endocrinologist.

Footnote 2: If patient has significant hyponatraemia but with a normal serum cortisol, consult the Society for Endocrinology Emergency Endocrine Guidance: Emergency management of severe symptomatic hyponatraemia in adult patients (21).

Footnote 3: Acute and subacute illness can lead to hypothalamic suppression of gonadotrophin and thyroid axis that recovers. This can be misleading in the diagnosis of hypothyroidism.

Footnote 4: Society for Endocrinology Emergency Endocrine Guidance: Emergency management of acute adrenal insufficiency (adrenal crisis) in adult patients (20).

Footnote 5: Absolute cut-offs for hyponatraemia are local assay and reference range dependent. We suggest amending these cut-offs in consultation with local laboratory/biochemistry advice.

Footnote 6: Immunosuppression can induce a number of toxicities which can mimic, precipitate or coexist with adrenal insufficiency. These must all be taken into consideration clinically.

Footnote 7: MRI of the pituitary can assist with a diagnosis of hypophysitis secondary to hypophysitis and exclude other causes (e.g. pituitary metastases, coincidental macroadenoma). There is no indication for i.v. methylprednisolone therapy in the absence of neurological features.

Footnote 8: Immunosuppression can induce a number of toxicities which can mimic, precipitate or coexist with adrenal insufficiency. These must all be taken into consideration clinically.

Footnote 9: Adrenal insufficiency may mask Diabetes Insipidus, which manifests once hydrocortisone is started.

Algorithm 1
Management of a life-threateningly unwell (CTCAE grade 3–4) patient.

http://www.endocrineconnections.org
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Hyperglycaemia

CPIs can induce pancreatic failure and an insulin-deficient diabetes mellitus.

- Plasma glucose should be measured in all unwell patients treated with a CPI.
- Hyperglycaemia should be managed as per local guidance for diabetic keto-acidosis or hyperglycemic hyperosmolar state as appropriate. Pancreatic antibodies (e.g. GAD 65) and C-peptide should be measured.
- Supraphysiological doses of glucocorticoids used as therapy for immune-mediated complications can induce diabetes, potentially causing or worsening hyperglycaemia and therefore monitoring of CBGs (capillary blood glucose) is essential in all patients.

If important to the oncology outcome, CPI therapy could be recommenced early after effective treatment for life-threatening endocrinopathy, with close observation.

Management of patient with mild/moderate symptoms (CTCAE grade 1–2) compatible with cortisol deficiency

<table>
<thead>
<tr>
<th>9 am cortisol &lt;200 nmol/l</th>
<th>9 am cortisol 200–450 nmol/l</th>
<th>9 am or random cortisol &gt;450 nmol/l</th>
</tr>
</thead>
<tbody>
<tr>
<td>adrenal insufficiency likely</td>
<td>adrenal insufficiency possible</td>
<td>adrenal insufficiency unlikely</td>
</tr>
</tbody>
</table>

- refer to Endocrinology
  - measure remainder of pituitary profile
  - IGF-1/TSH/T4/LH+FSH/TorE2/prolactin
  - For TFT abnormalities see Algorithm 3
  - consider SST (but interpret with caution if ACTH low as may be falsely reassuring in recent onset pituitary disease – discuss with Endocrinology)
  - continue immunotherapy if no other contraindications

- consider other causes of symptoms
- continue immunotherapy if no other contraindications

- if delay in Endocrine referral anticipated start oral hydrocortisone (10, 5, 5 mg) or prednisolone (3–5mg)

- consider other causes of symptoms
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Footnotes:

Footnote 1 Review patient information for evidence of recent steroid use:
- any supraphysiological dose of glucocorticoid can suppress the adrenal axis.
- patients receiving doses of dexamethasone >0.75 mg or prednisolone >3mg daily likely have a suppressed endogenous HPA axis and may have a serum cortisol measurement of <50 nmol/l. If the glucocorticoid treatment is ongoing they are not adrenally insufficient but may need higher doses of glucocorticoids when clinically unwell. Seek specialist advice from endocrinology.
Patients should be provided with a Steroid Emergency Card, education with regards to ‘sick day rules’, and an hydrocortisone emergency injection kit as per Society for Endocrinology guidance (20).

Immunotherapy can be continued once patient is clinically stable on appropriate endocrine replacement therapy (Algorithms 2 and 3).

Thyroid dysfunction

The most common CPI endocrinopathy is thyroiditis. In general, it starts with a transient hyperthyroid phase, which may not be symptomatic, followed by permanent subclinical or overt hypothyroidism. There have been reports of Graves’ hyperthyroidism and ophthalmopathy, thyroid storm and myxedema. Thyroid function tests should be performed as per Algorithm 3.

Thyrotoxic phase of thyroiditis should be managed symptomatically with beta blockers and regular monitoring of thyroid function tests. Subsequent hypothyroidism is likely to occur and requires treatment with thyroxine (Algorithm 3 footnote2). Pharmacological glucocorticoid therapy is not required and thioamide treatment (carbimazole/propylthiouracil) is of no value.

If there is clinical uncertainty, an iodine uptake scan should be considered (uptake reduced in thyroiditis and increased in Graves’ thyrotoxicosis). Iodinated contrast imaging may result in false-negative (sodium iodide) thyroid uptake scans. Institution-specific advice should be sought if thyroid nuclear medicine imaging is being performed after contrast administration. If thyrotoxic phase is persistent or there is evidence of thyroid eye disease, TSHRAbs should be measured and treatment with thioamides should be considered. Severe thyroid eye disease (CAS (Clinical Activity Score) >3 or evidence of optic nerve compression) should be referred urgently to ophthalmology/local thyroid eye service.

Levothyroxine should never be instigated until co-existent cortisol deficiency is excluded or treated as this can trigger an adrenal crisis.

Immunotherapy can be continued once patient is clinically stable on appropriate endocrine therapy.

Hypophysitis

Hypophysitis can present with either hormone defects or mass effect. It is an immune-mediated inflammation of the pituitary gland, the exact mechanism of this in
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If evidence of primary adrenal insufficiency, the serum cortisol cut-offs described in these algorithms are designed to ensure patient safety and err on initiating hydrocortisone therapy if there is any possibility of cortisol deficiency. Initiating glucocorticoid therapy on the basis of this guidance should not be taken as a definitive diagnosis of adrenal insufficiency. Referral to endocrinology services is advised in all cases. Patients should be re-evaluated once stable to confirm diagnosis and at intermittent intervals to assess for HPA axis recovery.

Immunotherapy can be continued once patient clinically stable on appropriate endocrine replacement therapy.

Maintenance steroid therapy following acute cortisol deficiency

Tapering of treatment dose glucocorticoids (hydrocortisone or prednisolone) can be started after clinical recovery (usually 48–72h). Aim for a daily replacement dose of hydrocortisone 10mg mane, 5mg lunchtime and 5mg early evening or prednisolone 3–5mg once daily. Referral to endocrinology is recommended. Maintenance glucocorticoid will usually be hydrocortisone but prednisolone is an acceptable alternative. If evidence of primary adrenal insufficiency, mineralocorticoid replacement (fludrocortisone)

• If hypophysitis is suspected clinically then a contrast-enhanced MRI pituitary scan should be performed as soon as possible.
• Headache, diplopia and cranial nerve palsies (CTCAE grade 3–4) should trigger an urgent MRI scan of the pituitary; to assess the potential need for treatment with IV methylprednisolone and to distinguish intracranial metastases as a differential diagnosis.

The value of IV methylprednisolone in the treatment of autoimmune hypophysitis, of any cause, is controversial, and there is minimal evidence of reversal of hormone deficiencies in CPI-induced hypophysitis but it may be of benefit for pressure effects such as optic chiasm compromise causing visual field defects, cranial nerve palsies and in some cases, headache.

Growth hormone deficiency does not need diagnosis or treatment in the acute phase. TSH and gonadotrophin deficiencies are also potential consequences of hypophysitis (frequency may differ with different CPIs) and should be treated if persistent. Acutely unwell patients of any etiology can have a suppressed TSH (sick euthyroid syndrome) and gonadotrophin axis but acute initiation of replacement is not indicated. Referral to an endocrine service is recommended.

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