MINI-REVIEW

Endocrine dysfunctions during treatment of immune-checkpoint inhibitors

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ABSTRACT

Immune-checkpoint inhibitors (ICIs) are novel agents directed to various malignant tumors. During ICI therapy, however, immune related adverse effects (irAEs) including endocrine dysfunctions have been reported. Dysfunctions in the pituitary gland and the thyroid gland by ICI are often observed, and those in the adrenal glands and the pancreas are less frequent. Positive correlation of the prevalence of endocrine irAEs to clinical antitumor effectiveness during ICI therapy has been reported. The mechanisms of endocrine irAEs by ICI, however, remain unclear, and optimal prevention, prediction, and treatment of the irAEs are still uncertain. This review describes possible mechanisms involved in ICI-related immunity, and discusses clinical management of endocrine irAEs during ICI therapy.

Keywords: PD-1; CTLA-4; immune-checkpoint inhibitors; endocrine organs; irAEs

Introduction

Immune-checkpoints play an indispensable role in anti-tumor immunity, anti-infection and autoimmunity⁴. Their major components include cytotoxic T-lymphocyte associated antigen 4 (CTLA-4), programmed death protein 1 (PD-1), and ligand for PD-1 (PD-L1) (Figure 1, 2). CTLA-4 is located on the surface of activated T-cells, therefore inhibiting binding of CD28 to B7 molecule on antigen presenting cells (APC)¹. CTLA-4 pathway predominantly acts in lymph nodes. PD-1 is majorly expressed on T-cells². PD-L1 is usually expressed on tumor cells, T-cells, B-cells, dendritic cells². PD-1 pathway is involved with tumor microenvironment. APC or tumor cells present cancer-related antigen with major histocompatibility (MHC)-class I to T-cell receptor (TCR) on cytotoxic CD8⁺ T-cell (Figure 1). Additionally, APC presents cancer-related antigen with MHC-class II to TCR on helper CD4⁺ T-cell (Figure 2). CD4⁺Th1-T-cells activate CD8⁺ T-cells, and CD4⁺Th2-T-cells stimulate antibody production. In antitumor immunity, CD4⁺ T-cell and CD8⁺ T-cell cooperate in the network.

Monoclonal antibodies to immune-checkpoints are referred to as immune-checkpoint inhibitors (ICIs), and are currently considered to be novel promising agents for treatment of malignant tumors⁵. ICI promotes T-cell mediated cytotoxicity directed to cancer cell antigens. It has been reported to improve prognosis of patients with malignant melanoma, renal cell cancer, non-small cell lung cancer, Hodgkin’s lymphoma, and Merkel cell carcinoma. Approximately 20%-30% of patients with malignant melanoma, renal cell cancer, and non-small cell lung cancer were found to be responders of ICI. More than 50% of patients showed objective responses in the patients with refractory Hodgkin’s lymphoma or Merkel cell carcinoma. On the other hand, various adverse events were reported as immune-related adverse
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IrAEs include dermatological, gastrointestinal, hepatic, neurological, and endocrine disorders\(^3,4\) (Table 1).

In endocrine organs, irAEs in the pituitary gland, the thyroid gland, the parathyroid glands, the adrenal glands, the pancreas (type 1 diabetes mellitus) were reported\(^5,6\). Although ICI activates T-cells through inhibition of immune-checkpoints molecules, the etiology of endocrine irAEs remains unclear. Endocrine dysfunctions are sometimes crucial, and diagnostic procedures in endocrine dysfunctions are often complicated. Routine follow-up including physical examination with endocrinological investigation, as well as image testing is important. This review describes endocrine dysfunction during ICI therapy, and raises possible mechanisms involved with ICI-related immunity, highlighting diagnostic approaches and appropriate clinical management during ICI therapy.

**Hypophysitis related to irAE**

The pituitary gland is an endocrine organ central to regulate peripheral endocrine glands, by secreting several pituitary hormones; adrenocorticotrophic hormone (ACTH), thyrotropin (TSH), gonadotropin, growth hormone (GH), and prolactin, and antidiuretic hormone (ADH). The estimated prevalence of irAE related pituitary dysfunction is 9.1%\(^3\), in contrast to idiopathic autoimmune hypophysitis which is only observed in 1 in 9 million individuals\(^14\). Incidence of irAE-related hypophysitis varies (0.4%-17.2%) in previous reports, the wide range associated with differences in surveillance\(^4,7,15,16\). Hypophysitis related to irAE occurs mostly within 6 months. In pituitary dysfunction, the incidence of secondary adrenal insufficiency: ‘ACTH insufficiency’, secondary hypothyroidism; ‘TSH insufficiency’, and secondary hypogonadism were 6.1%, 7.6%, and 7.5%, respectively\(^15\). The prevalence of GH insufficiency is not clear. Although rare, central diabetes insipidus due to pituitary posterior lobe dysfunction; ‘ADH insufficiency’ during ICI therapy was reported recently\(^17\).

Most of hypophysitis related to irAE is induced by anti-CTLA-4 antibody, and anti-PD-1 antibody-induced hypophysitis is less common (<1.0%)\(^4,18\). Hypophysitis related to irAE occurs more often in men than women\(^15,16\). In contrast, female predominance is seen in idiopathic autoimmune hypophysitis\(^14\). Common clinical manifestations in secondary adrenal insufficiency include headaches, fatigue, muscle weakness, and nausea, anorexia, weight loss, and hypotension are often reported\(^7,15,16,17\). In secondary hypothyroidism, constipation, fatigue, edema, and bradycardia can be seen. Manifestations in secondary hypogonadism and GH insufficiency can be not clear, but may be general fatigue and appetite loss. Notably, ACTH or TSH insufficiency are the most common pituitary hormone abnormalities in hypophysitis related to irAE\(^15,16\).

Secondary adrenal insufficiency is diagnosed with decreased ACTH and cortisol, and severe cases show hypoglycemia, hyponatremia, and hyperkalemia. Secondary hypothyroidism is diagnosed with decreased FT3, FT4, TSH, and sometimes hyponatremia, and elevated CK or cholesterol are seen. Secondary hypogonadism was diagnosed with decreased gonadotropin with decreased sex hormones (estradiol or testosterone). Low levels of GH and insulin-like growth factor-1 (IGF-1) are seen in GH insufficiency. In imaging tests, pituitary enlargement is sometimes seen in hypophysitis related to irAE in pituitary MRI\(^15,16\). Incomplete secretion of ADH and disappearance of T1-weighed high signal in posterior lobe in pituitary MRI is observed in central diabetes insipidus\(^15\). In order to precisely diagnose, pituitary provocation tests is often recommended. ACTH insufficiency is usually permanent, on the other
| Affected organ                        | Diagnosis                                   | Prevalence | Onset                  | Major symptoms                      | Laboratory data | Imaging tests                     | Recommended managements                                      | Additional requirements                                              |
|--------------------------------------|---------------------------------------------|------------|------------------------|-------------------------------------|-----------------|-----------------------------------|---------------------------------------------------------------|---------------------------------------------------------------------|
| Pituitary gland                      | Secondary adrenal insufficiency             | 6.1%       | Mostly within 6 months | Fatigue, muscle weakness, nausea, anorexia | ACTH↑ cortisol↓ | With/without pituitary swelling in pituitary MRI | Hydrocortisone replacement                                      | 1) Pituitary provocation tests, and ACTH infusion test is often recommended. 3) Adrenal insufficient should be firstly treated if hypothyroidism is concomitant. |
|                                     | Secondary hypothyroidism                    | 7.6%       | Depend on each report (0.4-17.2%) | Constipation, fatigue, edema, bradycardia | FT3↓FT4↓TSH↓   | L-T4 replacement                   |                                   | 2) Higher dosage of prednisolone is used for severe persistent headache or visual symptoms. |
|                                     | Secondary hypogonadism                      | 7.5%       | Mostly with anti-CTLA-4 Ab, and uncommon with anti-PD-1 Ab (<1%) | General fatigue appetite loss       | Gonadotropin↓   | Gonadotropin replacement if needed | GH replacement, if needed | 3) Adrenal insufficient should be firstly treated if hypothyroidism is concomitant. |
|                                     | GH insufficiency                            | Not determined | Not determined | Polydipsia and poluria              | GH↓IGF-1↓      | Disappearance of T1-weighed high signal in posterior lobe in pituitary MRI | Desmopressin acetate (DDAVP) |  |
|                                     | Central diabetes insipidus                  | Not determined | Not clear | Serum osmolality↑ADH↓              |                 |                                   |  |
| Thyroid gland                        | Thyrotoxicosis (destructive thyroiditis)    | 9%         | 2-6 weeks, mostly within 6 months | Palpitation, weight loss, tremor, fatigue, diarrhea | FT3↑FT4↑TSH↓   | Color Doppler flow↓¹²³I/¹³¹I Tc uptake↓ | Beta-blockers may be used |  |
|                                     | (often)                                     | 0-4%       | Depend on each report (0-39.1%) | Constipation, fatigue, edema, bradycardia | FT3↑FT4↑TSH↓ TRAb(+) | Color Doppler flow↑¹²³I/¹³¹I Tc uptake↑ | Anti-thyroid drug,¹³¹I treatment, or thyroid surgery |  |
|                                     | Thyrotoxicosis (hyperthyroidism)            | Rare       | 2-6 weeks, mostly within 6 months | Palpitation, weight loss, tremor, fatigue, diarrhea | FT3↑FT4↑TSH↑   | Color Doppler flow↑¹²³I/¹³¹I Tc uptake↑ | L-T4 replacement Adrenal insufficient should be firstly treated if hypothyroidism is concomitant. |  |
|                                     | Hypothyroidism                              | Not determined | Not determined | Polydipsia and poluria              | FT3↓FT4↓TSH↑   | Color Doppler flow↑¹²³I/¹³¹I Tc uptake↑ | L-T4 replacement Adrenal insufficient should be firstly treated if hypothyroidism is concomitant. |  |
| Adrenal gland                        | Primary adrenal insufficiency               | 1%         | Mostly within 1-6 months | Fatigue, muscle weakness, nausea, anorexia | ACTH↑ cortisol↓ | With/without adrenal swelling in abdominal CT | Hydrocortisone replacement                                      |  |
| Pancreas                             | Type 1 diabetes mellitus                    | 0.6%       | Mostly within 1-6 months | Thirst, polydipsia, and polyuria     | plasma glucose↑HbA1c levels are not always elevated. | With/without pancreas swelling in abdominal CT | Insulin therapy                                                   |  |
hand, secondary hypothyroidism/hypogonadism may recover. Positivity of anti-pituitary antibody may be helpful for diagnosis of hypophysitis related to irAE\(^{[18,19]}\). As predictive factors for hypophysitis related to irAE, eosinophilia, elevation of ESR/CRP/LDH, and hyponatremia can be useful\(^{[19]}\).

In monitoring and treatments, baseline and follow-up thyroid function (serum levels of FT3, FT4, and TSH) and morning ACTH and cortisol are crucial, because imbalance of hypothalamus-pituitary-thyroid (H-P-T) axis or hypothalamus-pituitary-adrenal (H-P-A) axis is life-threatening. A cosyntropin (; synthetic ACTH), infusion test is helpful to distinguish primary or secondary (pituitary-originated) adrenal insufficiency. Levels of GH-IGF-1, gonadotropin-sex hormones, and prolactin, should be assessed in patients with hypophysitis or hypopituitarism. High-dose steroids (e.g. 1-2 mg/kg/day of oral prednisolone) is used as treatment for those patients with severe headache, or visual abnormalities\(^{[15,16]}\). In cases of critical hypophysitis or hypopituitarism, intravenous glucocorticoid infusion is considered. Otherwise, oral hydrocortisone (15-30 mg/day) is used for treatment of adrenal insufficiency. In cases of adrenal crisis, higher dose of hydrocortisone (300 mg/day) is administrated intravenously. Levothyroxine (L-T4) (typically starting from 25 μg/day with increasing dosage) is used for secondary hypothyroidism; but adrenal insufficiency should be treated first to avoid adrenal crisis induced by increasing thyroid hormone effect. In cases of secondary hypoponadism or GH insufficiency, gonadotropin replacement or GH replacement therapy can be considered, respectively. In cases of central diabetes insipidus, Desmopressin acetate (DDAVP) is used\(^{[17]}\).

**Thyroiditis related to irAE**

The thyroid gland is an organ to produce thyroid hormone (T3, T4), and controls systemic metabolism. Major autoimmune thyroid diseases (AITD)s are Graves’ disease (GD) in which TSH receptor is autoantigen (20), and Hashimoto’s thyroiditis (HT) in which TPO and Tg are autoantigen\(^{[23]}\). Cell-mediated immunity and antibody-dependent cellular cytotoxicity (ADCC) are involved in AITD. An interim analysis of Japanese patients with malignant melanoma undergone ICI therapy showed high prevalence of thyroid disorders (any grade; 23.53%, and ≥Grade 3; 0.44%)\(^{[29]}\). In contrast to pituitary irAEs, the thyroid gland is preferentially affected by anti-PD-1 antibody rather than anti-CTLA-4 antibody. Thyroiditis related to irAE occurs in 0-4% patients treated with anti-CTLA-4 antibody\(^{[7]}\). Prevalence of thyroiditis by anti-PD-1 antibodies is estimated to be 9%\(^{[8]}\). IrAE-associated thyrotoxicosis and hypothyroidism similarly prevalent\(^{[6,23,24]}\). Transient thyrotoxicosis may precede hypothyroidism\(^{[23,24]}\). Dual treatment of nivolumab and ipilimumab induces thyroiditis in 22% of patients\(^{[25]}\). More recent reports showed increased incidence of anti-PD-1 antibody-induced thyroid dysfunction in 17.1% of malignant melanoma patients\(^{[24]}\) and 21% of non-small cell lung cell cancer patients, respectively\(^{[26]}\). Morganstein et al reported that ipilimumab-induced thyroid dysfunction occurred in 23% of patients, anti-PD-1-induced thyroid dysfunction in 39.1% of those, and dual treatment induced thyroid dysfunction in 50% of those\(^{[27]}\). Anti-PD-L1 antibody treatment was reported to induce 2-4% of thyroid dysfunction\(^{[28]}\). Patients who have anti-TPO-antibody (TPOAb) or anti-Tg-antibody (TgAb) are reported to be prone for development of thyroiditis related to irAE\(^{[29]}\). Thyroiditis related to irAE can occur early; between 2 and 6 weeks and mostly within 6 months after treatment. Notably, a proportion of the ICI-treated patients exhibit ‘non-thyroidal illness syndrome’ (normal FT4, normal TSH, and decreased FT3), reflecting lower nutrition status. Thyroid dysfunction during pembrolizumab treatment was reported to have possible association with feasible outcome of lung cancer\(^{[29]}\).

Thyrotoxicosis: thyrotoxicosis related to irAE is classified into two subtypes; 1) destructive thyroiditis, and 2) hyperthyroidism manifested as GD. Palpitation, weight loss, tremor, fatigue, and diarrhea are typically seen as thyrotoxic symptoms.

1) Destructive thyroiditis: destructive thyroiditis is often induced by ICI, transient increase of FT3 and FT4 are seen, concomitant with suppressed TSH levels. Anti-TSH receptor antibody (TRAb) is negative in most of the cases. TPOAb or TgAb are often positive at diagnosis\(^{[23,24]}\). Elevation of serum Tg levels may help the diagnosis of thyroiditis\(^{[30]}\). In thyroid ultrasonography, Doppler blood flow is not increased, thyroid uptake in \(^{123}\)I or \(^{99m}\)Tc scintigram is usually decreased, and FDG-PET is positively accumulated in the thyroid\(^{[24]}\). Diffuse thyroid enlargement can be seen\(^{[30]}\). In treatment, β-blocker is effective for relieving thyrotoxic symptoms. Effectivity of steroid treatment is not clear.

2) GD induced by ICI: Rare cases of thyrotoxicosis showed elevated FT3 and FT4 concomitant with suppressed TSH, and positivity of TRAb, suggesting GD during ICI treatment\(^{[31]}\). In those cases, increased color Doppler flow and increased uptake in \(^{123}\)I or \(^{99m}\)Tc scintigram were seen.
and treatment options are anti-thyroid drug, ¹³¹I treatment, or thyroid surgery.

Hypothyroidism: Hypothyroidism related to irAE is classified into two subtypes; 1) transient thyrotoxicosis followed by hypothyroidism, and 2) a single phase of hypothyroidism. Hypothyroidism is due to thyroiditis by ICI. Patients who have TgAb or TPOAb are prone to be hypothyroidism. Primary hypothyroidism is diagnosed by decreased FT3 and FT4, and increased TSH. Primary hypothyroid symptoms are similar to those in secondary hypothyroidism mentioned above. In thyroid ultrasonography, Doppler blood flow may be decreased, and thyroid uptake in ¹²³I or ⁹⁹ᵐTc scintigram is decreased. Diffuse thyroid swelling is seen in acute phase of thyroiditis, and atrophy can be seen in subsequent hypothyroid phase. L-T4 (typically starting from 25 μg/day with increasing dosage) is used to treat hypothyroidism. Cases of hypothyroidism often progress to permanent hypothyroidism. In monitoring, baseline measurement of thyroid autoantibodies (TRAb, TgAb, and TPOAb), and routine checkup of serum FT3, FT4 and TSH levels are recommended. Eosinophilia, thrombocytopenia, ESR/CRP/LDH elevation, and liver dysfunction might be important for earlier detection of thyrotoxicosis.

Adrenalitis related to irAE

Adrenal glands secrete adrenal hormones to maintain systemic activity. Cortisol is one of the critical adrenal hormones, regulating glucose metabolism, blood pressure, and other homeostasis. Cases of autoimmune adrenalitis were reported, with presence of autoantibodies to P450c21 or P450c17; adrenal enzyme involved with adrenal hormone synthesis. Adrenal failure is due to incomplete secretion of adrenal hormones including cortisol. IrAE-associated adrenal hypofunction is classified to primary or secondary adrenocortical insufficiency. Primary adrenocortical insufficiency is due to adrenalitis, and secondary adrenocortical insufficiency is due to pituitary damage described above. Prevalence of adrenalitis related to irAE is reported as approximately 1%. Adrenal dysfunction usually occurs within 1-6 months after initiation of ICI. Either anti-PD-1 antibody or anti-CTLA-4 antibody induces adrenalitis.

Symptoms in primary adrenal insufficiency are almost the same as those seen in secondary adrenal insufficiency. Decreased serum cortisol, increased plasma ACTH, and decreased urine cortisol levels indicate diagnosis of adrenalitis. In an ACTH stimulation test, response of cortisol is lost in cases of primary adrenal insufficiency. Presence of autoantibodies to P450c21 or P450c17 may be useful for diagnosis of adrenalitis. As a treatment, oral hydrocortisone replacement therapy (15-30 mg/day) is used. In cases of adrenocortical crisis, higher dose of hydrocortisone (300 mg/day) are administrated intravenously.

Type 1 diabetes mellitus and other endocrine dysfunctions related to irAE

Pancreatic beta cells produce and secrete insulin. Insulin controls blood glucose levels. In patients with type 1 diabetes (T1DM), pancreatic beta cells are destroyed, and insulin secretion is attenuated. Insulin is considered to be an autoantigen in T1DM. As a pancreatic irAE, patients with T1DM during ICI therapy were reported. Prevalence of T1DM associated with ICI therapy was reported to be 0.6%. Anti-PD-1 antibodies more often induce T1DM than anti-CTLA-4 antibodies. As a severe subtype of T1DM, fulminant type 1 diabetes (F1DM) was also reported. T1DM related to irAE occurs mostly within 1-6 months after ICI treatment, and often rapidly progressing with ketoacidosis. T1DM related to irAE is thus one of the urgent complications of irAE.

Thirst, polydipsia, and polyuria are seen as hyperglycemic symptoms, and elevated plasma glucose is seen. HbA1c levels are not always elevated, reflecting acute development of hyperglycemia. Sometimes swelling of pancreas is seen in abdominal CT. Anti-GAD, anti-IA2, and anti-insulin autoantibodies are sometimes positive. Immediate insulin therapy is required in patients with T1DM related to irAE. Monitoring of plasma glucose and HbA1c are useful for detection of T1DM. Measurement of anti-GAD antibody is also helpful to find individuals who may be susceptible to T1DM. Although rare, hypoparathyroidism related to irAE was reported as an endocrine irAE.

Possible mechanisms in endocrine irAEs

Multiple factors influence in the development of endocrine-related irAEs. In certain genetic and environment backgrounds, ICI may activate immune systems especially autoreactive cytotoxic CD8+ T-cells directed to endocrine organs.
Genetic factors of immune-checkpoints in irAEs

Altered function or genetic mutation of immune-checkpoint molecules may contribute to the development of endocrine irAEs. In AITD, genetic mutations in CLTA-4 in GD[38] and HT[39] were reported. An association of a SNP in PD-L1 with GD was shown[40]. SNPs in CLTA-4 and PD-L1 also contribute to the development of autoimmune adrenitis[41]. PD-L1 and PD-1 genetic mutation are reported to be involved in T1DM development[42,43].

Molecular mimicry of tumor antigen and auto-antigen in endocrine organs

One of the tumor-associated antigen: NY-ESO-1 possess common amino acid sequence with thyroid autoantigens (TSH receptor, Tg, and TPO), and administration of NY-ESO-1 was reported to induce GD or HT in individuals who have risk allele of MHC (class I or class II)[44]. In a case of malignant melanoma with F1DM during ICI treatment, HLA-DR4 restricted insulin autoantigen with positive conversion of anti-insulin antibody was suggested[45]. Therefore, upon both activation of anti-pancreatic beta cells and anti-melanoma immunity, exacerbation of pancreatic beta cell function and improvement of malignant melanoma can be seen in patients with F1DM related to irAE. Distinct manifestations of irAE inpatients with various malignant diseases were reported (e.g. malignant melanoma patients had a higher frequency of gastrointestinal/skin irAE, and a lower frequency of pneumonitis related to irAE)[45]. Thus, various tumor epitope may possess common amino acid sequences with endocrine epitope (e.g. TSH receptor, Tg, TPO, insulin). And cross-presentation of those on HLA molecules may be related to endocrine irAEs (Figure 1,2).

Inhibition the role of immune-checkpoints in endocrine irAEs

Anti-CTLA-4 antibodies often induce autoimmunity, by inhibiting regulatory T-cells function, and by activation of immunogenic CD4+ T-cells and CD8 T-cells[46,47]. In animal models, anti-CTLA-4 antibodies were reported to induce thyroiditis[48], adrenitis[49], and T1DM[50]. Anti-PD-1 antibodies blocks PD-1 pathway which suppress autoimmunity. Subsequently, immunogenic CD4+ T-cells and CD8 T-cells can be activated. PD-1 knockout mice showed lupus-like autoimmune disorders[2], and PD-1 blockade was reported to induce T1DM in mice[51].

Discussion

To date, most authors have recommended both discontinuation of ICIs and high dose glucocorticoid for treatment of irAEs. However, this has been challenged by recently, particularly if the endocrine irAEs can be managed, and the anti-tumor therapy is effective[53]. In addition, correlation of prevalence of irAEs to anti-tumor immunity was recently shown[54]. It is of interest to examine whether biomarkers which have been reported in malignant melanoma can be applied to the patients with endocrine irAEs[55]. Further investigations with larger number of cases over longer periods are warranted to establish diagnostic and therapeutic approaches to endocrine irAEs. In the future, identification of individuals (e.g. genetic predispositions such as HLA type, age, gender) who are susceptible to irAEs are necessary[55].
Conflict of interest

The authors declare no potential conflict of interest with respect to the research, authorship, and/or publication of their article.

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