

Immune Checkpoint Inhibitor-Induced Thyroidal, Pancreatic, and Pituitary Dysfunction: Diagnostic Challenges in a Cancer Patient

Disfunção Tiroideia, Pancreática e Hipofisária Secundária a Inibidores de Checkpoint Imunitário: Desafios Diagnósticos num Doente Oncológico

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ABSTRACT

Immune checkpoint inhibitors (ICPIs)-induced endocrine immune-related adverse events (irAEs) are common, can appear concurrently, and can be overlooked due to their nonspecific presentation overlapping with cancer-related symptoms. We describe the case of a 47-year-old woman with metastatic colorectal cancer treated with combined ICPI therapy. She presented thyrotoxicosis right after starting therapy, evolving into overt primary hypothyroidism, after two months, followed by abrupt-onset diabetes *mellitus* and hypophysitis-related secondary adrenal insufficiency and central hypothyroidism, six months after starting ICPIs. This case illustrates the complexity of diagnosing and managing overlapping endocrine irAEs, and the importance of high clinical suspicion. Clinical manifestations were attributed to the cancer and diabetes diagnosis, delaying recognition of adrenal insufficiency. Central hypothyroidism was initially interpreted as iatrogenic thyrotoxicosis. Glucocorticoid supplementation worsened diabetes management. Clinical and biochemical follow-up is essential in patients with ICPIs. Prompt recognition is essential to avoid life-threatening complications and ensure optimal long-term management.

Keywords: Endocrine System Diseases/chemically induced; Immune Checkpoint Inhibitors/adverse effects; Polyendocrinopathies, Autoimmune/chemically induced

RESUMO

Os efeitos adversos imunomediados (irAEs) endócrinos, secundários aos inibidores de *checkpoint* imunitário (ICPIs), são frequentes, podem surgir simultaneamente, e podem ser subdiagnosticados pela apresentação inespecífica, sobreponível à da doença oncológica. Descrevemos uma mulher de 47 anos com adenocarcinoma colorretal metastático, sob terapêutica combinada com ICPIs. A doente apresentou tirotoxicose transitória imediatamente após iniciar terapêutica, que evoluiu para hipotiroidismo primário, após dois meses, seguida de diabetes *mellitus*, e insuficiência adrenal secundária e hipotiroidismo central, secundários a hipofisite, seis meses após iniciar ICPIs. Este caso ilustra a complexidade do diagnóstico e gestão de irAEs endócrinos simultâneos, e a importância da elevada suspeição clínica. Os sintomas atribuídos à neoplasia e diabetes atrasaram o reconhecimento da insuficiência adrenal. O hipotiroidismo central foi interpretado como tirotoxicose iatrogénica. Os glucocorticóides prejudicaram o controlo glicémico. A vigilância clínica e laboratorial é fundamental, especialmente em doentes sob combinações de ICPIs. O reconhecimento precoce é crucial para prevenir complicações e garantir uma gestão eficaz a longo prazo.

Palavras-chave: Doenças do Sistema Endócrino/induzidas quimicamente; Inibidores de Checkpoint Imunológico/efeitos adversos; Poliendocrinopatias Autoimunes/induzidas quimicamente

INTRODUCTION

Immune checkpoints play a critical role in modulating and regulating the immune system's response, preventing overstimulation and autoimmunity.^{1,2} Immune checkpoint inhibitors (ICPIs) target these regulatory molecules, enabling immune recognition and subsequent destruction of cancer cells. However, their use can also lead to immune-related adverse events (irAEs) affecting any organ system.³ The prevalence and nature of each endocrine irAE vary depending on the ICPI used, and they are more common in case of combined therapy with multiple ICPIs.⁴

Endocrine toxicities are amongst the most common, accounting for 8% of ICPIs' irAEs.⁵ Unlike other irAEs, the inflammatory process is usually clinically silent and frequently results in permanent glandular damage. Clinical manifestations arise from the resulting hormone deficiency, often requiring lifelong hormone replacement therapy.⁶ Although most frequently observed during the early phases of treat-

ment, endocrine immune-related adverse events (irAEs) can arise at any time—even following discontinuation of ICPI therapy.³

Thyroid dysfunction is the most common endocrine irAE, followed by pituitary, pancreatic, and adrenal involvement and, less frequently, hypogonadism or hypoparathyroidism.⁵

We describe a case of multisystem endocrine dysfunction in a patient receiving combined therapy with ICPIs.

CASE REPORT

A 47-year-old-woman, receiving combination therapy with ipilimumab and nivolumab for progression of metastatic colon adenocarcinoma (CRC), after incomplete surgical resection (pT4bN1aM0) and palliative chemotherapy (FOLFIRI-bevacizumab), was referred to the endocrinology clinic for thyrotoxicosis (Table 1). She complained of a

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Diabetes *mellitus* is a rare but potentially life-threatening irAE of ICPIs, associated with diabetic ketoacidosis (DKA) in 70% of cases,³ primarily associated with PD-1/PD-L1 inhibitors or combination therapy.⁷ The sudden onset of hyperglycemia, low C peptide values and only slightly elevated HbA1c made the diagnosis of ICPI-induced diabetes *mellitus*, likely. The discrepant values reflect the rapid destruction of pancreatic beta cells, faster than what can be translated through the HbA1c value.^{3,7}

Immune checkpoint inhibitor-induced hypophysitis is most commonly seen after CTLA-4 inhibitors or combination therapy,⁵ usually resulting in panhypopituitarism. ACTH deficiency is the most common manifestation, followed by TSH and gonadotrophin deficiency.⁶

The clinical symptoms of cortisol deficiency were initially misattributed to the underlying cancer and newly diagnosed diabetes mellitus, leading to delayed recognition of adrenal insufficiency, which was ultimately identified through biochemical evaluation. This aligns with findings from several reviews and prospective studies, which emphasize that overlapping symptoms in cancer patients can hinder timely recognition of endocrinopathies—particularly adrenal insufficiency, which may rapidly progress to life-threatening states if not promptly diagnosed.⁸⁻¹¹

Central hypothyroidism emerged as a likely concurrent diagnosis. Retrospectively, the persistently low TSH levels, which prompted multiple levothyroxine dose adjustments, may have represented the early biochemical manifestation of central hypothyroidism, characterized by inappropriately low TSH and low thyroid hormone concentrations.

This report highlights the need for a high index of suspicion regarding the non-specific symptoms of endocrine irAEs. This is particularly relevant in patients in the oncology setting, in which cancer-related complaints can overlap and mask symptoms of ICPI-induced endocrinopathies, with life-threatening consequences. Regular clinical and biochemical evaluation is crucial in ensuring a prompt diagnosis and timely treatment.

This case also emphasizes the complexity of managing multiple, concurrent endocrine irAEs. Overlapping clinical features can delay diagnosis, and the management of one condition may directly complicate the diagnosis and treatment of another. In our patient, the diagnosis of central hypothyroidism was initially delayed due to the prior history of

primary hypothyroidism and ongoing levothyroxine therapy, which led to the assumption that the suppressed TSH levels reflected iatrogenic hyperthyroidism. Moreover, lifelong glucocorticoid replacement for adrenal insufficiency can interfere with glycemic management in the setting of ICPI-induced type 1 diabetes *mellitus*.

A multidisciplinary approach is essential for early detection, management and mitigation of long-term complications in patients with ICPIs' irAEs. Implementation of follow-up protocols may prove beneficial in these patients.

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AUTHOR CONTRIBUTIONS

SGS: Study design, data collection, analysis and interpretation, drafting and critical review of the manuscript.

RC, APS, II, JO: Study design, critical review of the manuscript.

All authors approved the final version to be published.

PROTECTION OF HUMANS AND ANIMALS

The authors declare that the procedures were followed according to the regulations established by the Clinical Research and Ethics Committee and to the Helsinki Declaration of the World Medical Association updated in October 2024.

DATA CONFIDENTIALITY

The authors declare having followed the protocols in use at their working center regarding patients' data publication.

PATIENT CONSENT

Obtained.

CONFLICTS OF INTEREST

The authors have no conflicts of interest to declare.

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REFERENCES

- Zhao Z, Wang X, Bao XQ, Ning J, Shang M, Zhang D. Autoimmune polyendocrine syndrome induced by immune checkpoint inhibitors: a systematic review. *Cancer Immunol Immunother.* 2021;70:1527-40.
- Iwama S, Kobayashi T, Arima H. Management, biomarkers and prognosis in people developing endocrinopathies associated with immune checkpoint inhibitors. *Nat Rev Endocrinol.* 2025;21:289-300.
- Husebye ES, Castinetti F, Criseno S, Curigliano G, Decallonne B, Fleseriu M, et al. Endocrine-related adverse conditions in patients receiving immune checkpoint inhibition: an ESE clinical practice guideline. *Eur J Endocrinol.* 2022;187:G1-21.
- Park R, Lopes L, Cristancho CR, Riano IM, Saeed A. Treatment-related adverse events of combination immune checkpoint inhibitors: systematic review and meta-analysis. *Front Oncol.* 2020;10:258.
- Cardona Z, Sosman JA, Chandra S, Huang W. Endocrine side effects of immune checkpoint inhibitors. *Front Endocrinol.* 2023;14:1157805.
- Percik R, Criseno S, Adam S, Young K, Morganstein DL. Diagnostic cri-

teria and proposed management of immune-related endocrinopathies following immune checkpoint inhibitor therapy for cancer. *Endocr Connect.* 2023;12:e220513.

7. de Filette JM, Pen JJ, Decoster L, Vissers T, Bravenboer B, Van der Auwera BJ, et al. Immune checkpoint inhibitors and type 1 diabetes mellitus: a case report and systematic review. *Eur J Endocrinol.* 2019;181:363-74.
8. Kotwal A, Perlman JE, Goldner WS, Marr A, Mammen JS. Endocrine dysfunction from immune checkpoint inhibitors: pearls and pitfalls in evaluation and management. *JCO Oncol Pract.* 2023;19:395-402.
9. Kotwal A, Kennedy R, Kikani N, Thosani S, Goldner W, Shariff A. Endocrinopathies associated with immune checkpoint inhibitor use. *Endocr Pract.* 2024;30:584-91.
10. Wright JJ, Johnson DB. Approach to the patient with immune checkpoint inhibitor-associated endocrine dysfunction. *J Clin Endocrinol Metab.* 2023;108:1514-25.
11. Wright JJ, Powers AC, Johnson DB. Endocrine toxicities of immune checkpoint inhibitors. *Nat Rev Endocrinol.* 2021;17:389-99.