Immune checkpoint inhibitor-induced takotsubo syndrome and diabetic ketoacidosis: rare reactions

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SUMMARY
Immune checkpoint inhibitors (ICIs) are increasingly used to treat certain malignancies due to their higher efficacy compared with conventional chemotherapy. As familiarity with these agents increases, it is becoming apparent that a significant number of patients treated with ICIs experience adverse events. With time, more immune-related adverse events (IRAEs) are being recognised. It is important to be vigilant for IRAEs and recognise that a patient may have multiple IRAEs affecting multiple organ systems. Common cardiovascular adverse events associated with ICIs include myocarditis, arrhythmias and pericarditis. This case report identifies a patient presenting with takotsubo syndrome followed by ketoacidosis (associated with sodium-glucose transport protein 2 (SGLT2) inhibitor) in the setting of combination ipilimumab and nivolumab therapy for metastatic melanoma.

BACKGROUND
Immune checkpoint inhibitors (ICIs) are increasingly used to treat certain malignancies due to their higher efficacy compared with conventional chemotherapy. Emergence of ICI is a turning point in the field of immuno-oncology. Tumour cells escape immunosurveillance by activation of immune checkpoint pathways that inhibits anti-tumour immune responses. ICIs reactivate antitumour immune responses by blocking co-inhibitory signalling pathways and promote immune-mediated destruction of tumour cells.1

As familiarity with these agents increases, it is becoming apparent that a significant number of patients treated with ICIs experience adverse events. Literature suggests that 75%–90% of patients on a cytotoxic T lymphocyte antigen-4 (CTLA-4) inhibitor and 30%–70% of patients on an anti-programmed cell death protein-1 (PD-1)-blocking and/or anti-programmed cell death protein-1 ligand (PD-L1)-blocking monoclonal antibody experience an immune-related adverse event (IRAE).2,3 Combination therapy has up to a 40% higher rate of grade 3 or 4 adverse events.4 Common cardiovascular adverse events associated with ICIs include myocarditis, arrhythmias and pericarditis. There are published case reports of ICI-triggered takotsubo syndrome (TS).3-5 ICI-induced endocrinopathies are well described in the literature and these occur at a higher frequency with combination therapy.5 This case report identifies a patient presenting with TS followed by ketoacidosis (associated with SGLT2 inhibitor) in the setting of combination ipilimumab and nivolumab (Ipi/Nivo) therapy for metastatic melanoma.

CASE PRESENTATION
A 76-year-old man presented to the emergency department with central crushing chest pain and diaphoresis. This was on a background of metastatic melanoma with intracranial metastases, and a transfusion of Ipi/Nivo therapy in the preceding weeks. Two months prior to his presentation, he had had a craniotomy and debulking of his intracranial metastases. His other past comorbidities included type 2 diabetes mellitus (for which he took empagliflozin, an SGLT2 inhibitor), dyslipidaemia and hypertension. An ECG performed in the emergency department at presentation revealed 1–2 mm ST elevation in leads V2–V6 (figure 1). The initial management included sublingual glyceryl trinitrate, anticoagulation with enoxaparin, aspirin and fentanyl, and this resulted in the resolution of the ST segments elevation on subsequent ECG. He was taken to the Cath lab and as his ST segments had resolved the on-call interventionalist deemed it prudent to seek an oncology and neurosurgical opinion about the safety of heparin, dual antiplatelets and a staged angiogram.

INVESTIGATIONS
His initial troponin I was 938 ng/L, which subsequently peaked at 2679 ng/L with the reference range being < 20 ng/L. He was cleared by neurosurgery and subsequently underwent coronary angiography. This revealed non-obstructive coronary artery disease (CAD) (figure 2) and the left-ventriculogram showed reduced left ventricular ejection fraction (LVEF) of about 40% with apical ballooning. Echocardiogram showed an LVEF of 50% and apical akinesis with ballooning and hypokinetic basal and mid segments (video 1). This picture was consistent with TS. A detailed history did not reveal any recent emotional or physical stress. He made an uneventful recovery and was discharged from the hospital. On discharge, his troponin had already started to decrease, falling to 1428 ng/L indicating no ongoing insult to his myocardium.

Six days later, the patient received his second course of Ipi/Nivo. Four days after the ICI treatment, he re-presented with recurrent chest pain. His initial troponin I of 26 ng/L rose to 674 ng/L within 2 hours. Blood tests revealed diabetic ketoacidosis (DKA) with a blood glucose level of 24.6 mmol/L (normal range 3–7.8 mmol/L), ketones of...
6.6 mmol/L (range <1 mmol/L) and a pH of 7.12 (normal range 7.32–7.43). He was managed with nitrates, antiplatelet agents and enoxaparin once again together with intravenous rehydration, but an angiogram was not performed. An echocardiogram demonstrated ongoing hypokinesis of the apical segments.

The troponin levels returned to normal soon after the symptoms got resolved. Clinical features were not suggestive of myocarditis.

**DIFFERENTIAL DIAGNOSIS**

To exclude the possibility of ICI-induced myocarditis, a cardiac MRI was performed. The MRI showed normal systolic function, an ejection fraction of 66% and normal left ventricular myocardial mass and ventricular wall thickness. There was mild hypokinesis of the apical segments consistent with previous history of TS. There was no myocardial gadolinium enhancement to suggest myocarditis.

In this clinical setting, it is clear that the TS and subsequent ketoacidosis were secondary to the infusion of ipilimumab and nivolumab as there was no other diagnosis or aetiology identified. The development of ketoacidosis is consistent with emplagliflozin therapy and immune-stress response associated with ICI therapy.

**TREATMENT**

Following our patient’s first episode of TS, he was placed on aspirin, a long-acting beta-blocker (bisoprolol) and an ACE inhibitor (ramipril) for management of his TS and their cardio-protective benefits. On his initial presentation, it was not clear that there was a direct correlation between the ICI and his TS. The managing oncology team felt that due to the severity of his disease and the unclear correlation of his TS with his previous ICI infusion, there was no barrier in receiving his second ICI infusion. Following his second infusion and subsequent DKA and TS, he was determined to be for no further ICI infusions due to complications and commenced on corticosteroids as treatment for his IRAEs.

**OUTCOME AND FOLLOW-UP**

Our patient made a successful recovery from his TS and DKA. His cardiac MRI showed a normalised LVEF and systolic function post his second presentation. There were no further DKA or cardiac events. Despite receiving no further ICI treatments, our patient unfortunately had further IRAEs experiencing ICI-induced colitis and hepatitis, requiring months of high dose corticosteroids and mycophenolate.

**DISCUSSION**

Our patient had a classic presentation of TS. A literature review of TS found that an emotional or physical stress is present as the trigger in 39% and 35% of patients, respectively.\(^\text{10}\) Having excluded acute coronary syndrome, myocarditis and physical and emotional stress in conjunction with the observation that two sequential episodes of TS followed the same dual ICI therapy confirms the observation that TS was secondary to ipilimumab and nivolumab treatment.

Historically TS has been considered a benign condition with a favourable prognosis.\(^\text{11}\) Recent data shows that indeed TS secondary to emotional triggers has a favourable outcome compared with CAD, however overall TS has comparable
mortality outcomes to age-controlled patients with CAD. TS secondary to a physical trigger has a worse 5-year mortality than CAD, likely due to the underlying comorbid state of the patient population.12–13

There have been reports in the literature of TS secondary to ICIs and a proportionality study found that there was an increased OR of TS when compared with positive and negative controls.6–8 The precise pathophysiology of TS in the setting of ICIs is unclear and imaging studies have not consistently shown myocardial oedema.7 8 The proposed mechanism of injury in ICI-induced myocarditis is lymphocyte (CD4+ and CD8+ T cells) and macrophage infiltration of the myocardium.14 There has also been a case report of lymphocytic myocarditis mimicking TS, which may imply that some of these cases are atypical myocarditis.15 The lack of gadolinium enhancement and myocardial oedema in our patient is consistent with the previously published case reports, suggesting that ICIs can cause TS without myocarditis.

Our literature review revealed no case reports of ICI-induced euglycaemic or hyperglycaemic DKA in the setting of concurrent SGLT2 inhibitor. However, considering our patient’s hyperglycaemia and his significant ketoacidotic state, it is possible that SGLT2 inhibitor therapy contributed to the metabolic disturbance in the setting of ICI, and it remains unknown whether this is an atypical manifestation of euglycaemic DKA. ICI-induced diabetes (type 1) has been reported with an estimated incidence of 1%18 and approximately 76% of new cases present with DKA.19 Our patient had established type 2 diabetes.

An antibody screen showed antibodies to glutamic acid decarboxylase (anti-GAD) antibodies of 1012 (upper limit of normal <5). Anti-GAD antibodies have been reported as the most common antibody associated with ICI-induced type 1 diabetes.20 It is not possible to determine whether the development of ketoacidosis was related to the presence of TS or a manifestation of ICI immune-mediated toxicity, or alternately unrelated to both as ketoacidosis associated with SGLT2 therapy, although not too common, is well recognised.

In conclusion, ICIs are showing exceptional promise as the new frontier in treating malignancies. It is important to be vigilant for IRAEs and recognise that a patient may have multiple IRAEs affecting multiple organ systems. Over time, new case reports and data are bringing to light the many manifestations of IRAEs. Knowledge and science related to ICI therapy are improving with enhanced efficacy as well as safety. More research is required to identify best means of prevention and management of such adverse effects (IRAEs) as the use of these agents become more prevalent.

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