Myocarditis associated with immune checkpoint inhibitors: Practical considerations in diagnosis and management

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Abstract

Immune checkpoint inhibitors (ICI) have caused radical changes in the treatment scheme of many types of cancer in the past 10 years. ICIs are specific monoclonal antibodies that increase T-cell mediated immune response against cancer cells. Despite important advances in cancer treatment, uncontrolled activation of cytotoxic T cells has brought along many autoimmune clinical side effects, especially acute myocarditis. Although the incidence of ICI-related myocarditis is about 1%, it is remarkable in terms of mortality rate reaching 46% and demonstrating the necessity of rapid diagnosis and multidisciplinary approach. The present review aimed to summarize the heterogeneous symptomatology of ICI-associated myocarditis, clinical presentation ranging from elevated asymptomatic cardiac enzyme levels to cardiogenic shock, prominent diagnostic value of cardiac magnetic resonance imaging, and current information on the effectiveness of immunosuppressants in therapy.

Keywords: cardiotoxicity, immune checkpoint inhibitor, myocarditis

Introduction

Immune checkpoint inhibitors (ICIs) have been introduced into clinical practice as specific monoclonal antibodies that demonstrate significant antitumor activity through enhancement of T-cell-mediated immune response against tumor cells (1). These agents have been considered as the most important breakthrough of cancer management over the past 10 years with a significant potential to induce radical changes in the management algorithm of many cancer types including malignant melanoma, non-small cell lung carcinoma, and renal cell carcinoma (1-3). Mechanistically, ICIs, instead of direct anticancer action, primarily inhibit specific pathways that downregulate T lymphocyte response to the tumoral cells potentially rendering them as the target of cytotoxic T cells. ICIs exert their actions through three basic pathways: cytotoxic T lymphocyte-associated antigen-4 (CTLA-4), programmed cell death protein 1 (PD-1), and programmed cell death ligand 1 (PD-L1) (Fig. 1) (4). However, with the gradually expanding use of ICIs in clinical practice, various autoimmune-mediated adverse events including acute myocarditis have been increasingly encountered possibly due to the uncontrolled activation of cytotoxic T cells (5). Table 1 lists major cardiac toxic events reported to be associated with the use of ICIs in the literature (1-10). This study aimed to discuss diagnostic and therapeutic implications of ICI-associated myocarditis in clinical practice.

Figure 1. Major types of immune checkpoint inhibitor-related cardiotoxicity

CTLA-4 - cytotoxic T lymphocyte-associated antigen-4; PD-1 - programmed cell death protein receptor; PD-L-1 - programmed cell death protein ligand
ICI and immune myocarditis

Mechanisms

The mechanism of ICI-related myocarditis has not been clearly elucidated yet. However, immunological homeostasis failure has been suggested to be caused by the inhibition of CTLA-4, PD-1, and PD-L1 receptors, known as the checkpoint immune system, and reduced T cell tolerance to normal tissues initiate all immune-related adverse events, including myocarditis (8, 10-12). CTLA-4 and PD-1 proteins are receptors on cytotoxic T lymphocytes that enable the control of the immune response by suppressing the excessive activation of these cells (1, 2). PD-L1 receptors, on the other hand, are ligands found in normal cells such as cardiomyocytes and bind to receptors on cytotoxic T lymphocytes and play a cardio protective role against excessive activation of these cells (11, 12). In the literature, it has been reported that cytotoxic T cell infiltration and PD-L1 inhibition are observed in myocardial tissue in postmortem analysis of ICI-related lethal myocarditis cases (8, 12). Additionally, lethal myocarditis and dilated cardiomyopathy associated with cytotoxic T-lymphocyte infiltration have been reported in studies with CTLA-4 and PD-1 knockout mice investigating the effects of immunotherapeutics (13). Another study reported that macrophage and B cell infiltrations, mainly CD4+ and CD8+ T cells, were less common in the myocardium of cynomolgus monkeys given with the combination of ipilimumab and nivolumab (14).

As a summary, in the light of data from postmortem analyses and animal studies, the primary mechanisms responsible for the development of ICI-related myocarditis are as follows: 1. Cross-reaction infiltration of normal myocardial tissue, which was detected as a tumor cell by immune system cells (especially cytotoxic T lymphocytes) and 2. Myocardial damage due to inhibition of PD-1/PD-L1 receptors, which play a cardio protective role against uncontrolled immune response (10, 11, 15).

Clinical features

Even though the overall incidence of ICI-associated immune myocarditis appears to be around 1%, its mortality rate might potentially reach 46% absolutely mandating a rapid diagnostic process with a multidisciplinary approach (5-7). Clinical presentation generally ranges from an asymptomatic elevation of cardiac enzymes to sudden death due to heart failure. Accordingly, symptoms generally present with a heterogeneous clinical spectrum that might include dyspnea (usually accompanied by a reduced left ventricular ejection fraction (LVEF) value), palpitations, nausea, fatigue, weight loss, and chest pain (6). In full-blown cases, all aspects of an acute heart failure scenario are usually fully established. However, clinical findings in this setting might be easily overlooked since patients receiving ICI treatment already appear to suffer significant frailty generally manifesting as malaise, low exercise capacity, and dyspnea associated with the primary disease itself (lung cancer, etc.).

Temporal emergence

In particular, the first three months of ICI treatment is considered as a high-risk period in the development of immune myocarditis (7, 16, 17). According to a large case series study by Mahmood et al. (7), the median time to onset of myocarditis development from the first ICI dose was reported to be 34 days. Importantly, another study by Atallah-Yunes et al. (17) reported that 93% of subjects with fatal myocarditis received only one or two doses of ICI. Of note, previous cases of immune myocarditis were reported to develop even within the first 15 days of therapy, particularly in those who had received combination therapy (ICI-tyrosine kinase inhibitors (TKI) and/or previous PD-1 inhibition) (6).

Accompanying clinical conditions

It is well known that a second immune-related condition can potentially develop with ICI-associated immune myocarditis (7, 16, 17). Accompanying conditions in this setting have been reported such as myositis, colitis, nephritis, and hepatitis. Interestingly, these concomitant conditions might just precede the evolution of myocarditis and, hence, might serve as a predictor of an impending myocarditis (7, 16). However, varied symptoms in these conditions can lead to significant delays in their diagnosis. Particularly in myositis (the most frequently accompanying condition), attribution of symptoms (including fatigue and widespread myalgia) to a metastatic malignancy might delay the diagnosis of an existing myositis (that strongly warrants interruption of ICI therapy) and, hence, also substantially increase the risk for myocarditis with uninterrupted ICI use. Therefore, treating clinicians should also be fully aware of these immune-related events and their clinical implications during ICI therapy (7, 16, 17).

Diagnosis

The diagnosis of an immune-related myocarditis generally requires a high index of suspicions and mandates close monitoring of ICI recipients. It should be carefully considered that a timely diagnosis in this setting generally translates into a timely

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**Table 1. Major types of ICI-related cardiotoxicity**

| Cardiotoxicity       | Incidence*       |
|----------------------|------------------|
| Myocarditis          | 0.09%-2.4%       |
| Pericarditis         | <1%-2%           |
| Pericardial effusion | 2%               |
| Cardiac arrhythmia   | 4%               |
| Myocardial infarction| <1%-2%           |
| Heart failure        | 0.4%             |
| Takotsubo cardiomyopathy | Rarely reported |
| Cardiac arrest       | Rarely reported  |

*The frequency of ICI-related cardiotoxicity varies according to the molecule used and the method of treatment (monotherapy/combination)*
initiation of life-saving therapeutic strategies for a given patient. In fact, there exists only limited information in the literature regarding the diagnosis and management of ICI-related myocarditis, most of which are primarily based on expert opinion. The fundamental reason for diagnostic delay in ICI-related myocarditis is due to the existing nonspecific symptomatology (frequently encountered in cancer patients) along with the frequent coexistence of noncardiac factors associated with cardiac biomarker elevation in this group of patients. Figure 2 shows an algorithm for the differential diagnosis of ICI-related myocarditis (10, 11, 15, 18-23).

Identification of high-risk groups
In ICI therapy, the possibility of an ICI-related myocarditis should be considered by the treating clinician even in the presence of asymptomatic troponin elevation and/or myositis that presents with fatigue, widespread muscle pain, and elevated creatine kinase (CK). Particularly in combination therapy (CI+TKI) and/or previous PD-1 inhibitor, asymptomatic troponin elevation should raise a high index of suspicion (8, 17). It has been reported that the risk of myocarditis is fivefold in combination therapy (nivolumab and ipilimumab) compared with nivolumab only (8, 17). It is also noteworthy that myositis occurs in approximately one-third of those with ICI-associated myocarditis. Accordingly, previous studies suggested that an existing myositis should be considered as a “red flag” for the diagnosis of ICI-related myocarditis (8, 16, 17).

Electrocardiogram (ECG)
Currently, ECG finding for patients with ICI-related myocarditis remains nonspecific. However, nearly 90% of these patients have been reported to have various new-onset ECG changes (7). An ECG finding has been suggested to include sinus tachycardia, bundle branch block, atrial fibrillation, AV block, bigeminy ventricular premature beats, ventricular tachycardia, and ST-segment elevation (8, 17, 24, 25). Even though ECG generally has a limited diagnostic value in ICI-associated myocarditis, it is an easily accessible and low-cost test and, hence is recommended at the beginning and before every consecutive dose of ICI treatment (26) so potential changes can be monitored in time.

Cardiac biomarkers
Troponin: Studies showed that almost all patients with ICI-associated myocarditis have elevated high-sensitivity troponin levels (1). High troponin levels strongly suggest the presence of an ongoing damage due to myocardial inflammation (4). As expected, a strong correlation was observed between troponin levels and the risk of major adverse cardiac events including cardiovascular death, shock, and AV complete block in patients with ICI-related myocarditis (8, 17, 24). However, troponins are quite nonspecific in this setting as it is well known that various noncardiac causes of troponin elevation also exist in the clinical setting.

Natriuretic peptides: Similar to troponins, these peptides are also nonspecific in this setting (7, 27). Importantly, owing to

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**Figure 2.** How to make the differential diagnosis in patients using ICI and presenting nonspecific symptoms: An algorithm for the clinician

- **BNP/NT-proBNP** - brain natriuretic peptide; **LV** - left ventricle; **EF** - ejection fraction; **MRI** - magnetic resonance imaging; **CT** - computed tomography; **PET-CT** - positron emission tomography-computed tomography; **EMB** - endomyocardial biopsy
the significant inflammatory process induced by cancer itself, N-terminal pro-brain natriuretic peptide (NT-proBNP) might be chronically elevated among cancer patients. Moreover, all other acute cardiac stress conditions that the cancer patient potentially suffers might have elevated NT-proBNP levels (9, 27). However, it is well known that increased myocardial wall stress appears to be the primary trigger for natriuretic peptide release. Therefore, these peptides usually increase to substantial levels in myocarditis accompanied by acute heart failure.

**Echocardiography**

Results of previous studies on the clinical relevance of echocardiography in diagnosis and long-term risk stratification of ICI-related myocarditis are conflicting. A study by Escudier et al. (9) reported that three-fourth of patients diagnosed with ICI-associated myocarditis developed left ventricular dysfunction on echocardiography, while another study by Mahmoud et al. (7) reported that ejection fraction was within normal limits in more than half of the patients who suffered ICI-associated myocarditis. Echocardiography also has a limited value in predicting cardiovascular outcomes among patients with ICI-associated myocarditis (17). An existing normal ejection fraction value does not exclude ICI-related myocarditis (7). Moreover, a previous study reported that the left ventricle might not dilate in fulminant myocarditis (7, 9). Even though an existing normal systolic function potentially denotes a relatively benign and self-limiting course, this does not warrant an absolute sign of perfect prognosis. Therefore, in patients with suspected myocarditis, further tests including cardiac magnetic resonance imaging (MRI) are necessary to detect myocardial inflammation particularly in the presence of normal systolic functions (4).

**MRI**

Cardiac MRI, a noninvasive method, plays a pivotal role in the early detection and serial monitoring of myocardial inflammation. In T1-/T2-weighted MRI of patients with ICI-related myocarditis, mid-myocardial edema and late gadolinium enhancement that does not correspond to coronary anatomy are of particular importance (25, 26, 28). The specificity and sensitivity of MRI in the diagnosis of myocarditis are 91% and 67%, respectively. It should be carefully considered that MRI may be completely normal in 20–30% of ICI-related myocarditis with diffuse or low-grade inflammation (9, 25, 26, 28), complementing the results for MRI in the diagnosis of ICI-related myocarditis. Furthermore, FDG-PET can also be utilized to monitor the degree of inflammation in myocarditis and manage immunosuppressive therapy accordingly (29).

**Endomyocardial biopsy**

Endomyocardial biopsy has been suggested as the gold standard diagnostic method in ICI-related myocarditis (17, 30, 31). Detection of lymphocytic infiltration along with myocyte necrosis in the biopsy material is of diagnostic value. Inflammatory and fibrotic changes characterized by infiltration of CD 4+ and CD 8+ T cells (CD 8+ involvement is reported to be more dominant in the conduction system), macrophages, and occasionally B and plasma cells (with the particular absence of eosinophilic granuloma or giant cells) are quite typical in this setting (4, 17, 32). Since endomyocardial biopsy is an invasive procedure with significant false-negative results (particularly when the biopsy material is taken from disease-free territories), its diagnostic value is limited in this setting (28). Table 2 summarizes diagnostic and follow-up methods for the prevention or early diagnosis of myocarditis in cancer patients receiving ICI therapy.

**Management Strategies**

**Immunosuppressive therapy**

Postmortem analysis in patients who succumbed to ICI-related myocarditis has reported significant infiltration of various T cell populations (7). This substantiates the idea that the fundamental mechanism accounting for myocyte damage results from the over activation of cytotoxic T cells. Although no strong evidence-based recommendations has been reported, suppression of high-grade inflammatory response within the myocardium largely through immunosuppressive agents has been suggested as a milestone in the treatment of ICI-associated myocarditis (24, 30, 33, 34). Within this context, even though corticosteroids (7) appear to be the basis of immunosuppressive therapy, cases have been reported where treatment regimens including antithymocyte globulin (6), mycophenolate mofetil (6), abatacept (33), alemtuzumab (34), tacrolimus (30), and rituximab (30) have been initiated depending on LVEF, hs-troponin level along with hemodynamic status. The common mechanism of action of these agents is to induce the inactivation or destruction of T cells, that are considered to play a pivotal role in the evolution of myocarditis, as confirmed by endomyocardial biopsy findings of patients with ICI-related myocarditis. Therefore, since initiation of immunosuppressive therapy in ICI-related myocarditis implies the reversal of anti-tumor immunity, careful consideration of risks and benefits is strongly warranted beforehand.

In ICI-related myocarditis with left ventricular systolic dysfunction, heart failure treatment as per current guidelines should be immediately started (6). However, clear consensus on the optimal dose of corticosteroids (the first-line treatment to start as an immunosuppressive) and on how and when to add other immunosuppressive agents has not been reported yet. In this regard, the initial dose of corticosteroid ([methyl prednisolone]) is usually recommended as 1–2 mg/kg in the present study (6, 7, 9). Some studies showed strong association of high-dose steroid therapy with a significantly lower rate of major adverse cardiac events (6, 7). In fact, mostly based on hs-Troponin levels of the patient, continuing with a higher steroid dose during the maintenance period is highly recom-
mended (28). Previous studies reported on the use of myco-
phenate mofetil (1 g oral BD), intravenous immunoglobulin (IVIG) (3 days, 2 g/day), antithymocyte globulin, and infliximab in ste-
roid-refractory myocarditis cases. However, given the relation
between infliximab and heart failure, this agent should not to be
given in high doses (6, 7, 35, 36). Figure 3 shows an algorithm for
the treatment of patients according to their response to corti-
costeroid (7, 11, 17, 22, 31).

Prevention of ICI-related myocarditis: General strategies be-
fore, during, and after treatment
Currently, evidence for the role of cardiac monitoring, opti-
mal follow-up strategy, or cardio protective strategies in patients
with ICI-related myocarditis is nonspecific. Diagnosis and treat-
ment methods suggested for the prevention and early diagnosis
of ICI-related myocarditis (Table 2) should apply to all cancer pa-
tients receiving ICI therapy (1, 4, 6, 7, 30, 31, 37).

Table 2. Diagnosis and follow-up strategies for the prevention or early diagnosis of ICI-related myocarditis

| Potential risk factors for the evolution of ICI-related myocarditis | - The combination of dual ICIs (such as ipilimumab+nivolumab) or ICI+cardio toxic VEGF tyrosine kinase inhibitors
| | - Accompanying immune-related adverse event, especially myositis
| | - History of myocardial infarction, heart failure, myocarditis, previous treatment with an anthracycline, history of cancer therapy-associated left ventricular dysfunction
| | - Presence of autoimmune diseases such as systemic lupus erythematosus, rheumatoid arthritis, and sarcoidosis.
| Perform basic cardiovascular assessment | - Clinical history
| | - ECG
| | - Cardiac biomarkers: troponin, BNP, or NT-proBNP
| | - Creatinine kinase
| | - Echocardiogram
| Close follow-up (particularly during the first 3 months when the risk is high) | - In high-risk patients, ECG, cardiac troponin, and NT-proBNP should be checked before 2<sup>nd</sup> or 4<sup>th</sup> doses.
| | - In high-risk patients, echocardiography should be performed before 2<sup>nd</sup> or 3<sup>rd</sup> doses, and echocardiography should be repeated every 3–6 months if there is initial left ventricular or right ventricular dysfunction.
| | - Question the emerging symptoms: fatigue, muscle pain, fever, chest pain, shortness of breath, palpitations, presyncope, or syncope, etc.
| | Follow multidisciplinary approach. Refer to the cardio-oncology outpatient clinic in case of recently elevated troponin, NT-proBNP increase, and ECG or echocardiographic abnormality.
| Suspected myocarditis | - The patient should be referred to the cardio-oncology outpatient clinic
| | - ECG (new changes?)
| | - Repeat echocardiography
| | - In the presence of palpitations, presyncope, or syncope, a 24-hour rhythm Holter test is indicated
| | - Troponin (normal troponin levels during follow-up does not exclude myocarditis)
| | - BNP/NT-proBNP (It should be noted that these peptides may increase in every condition associated with acute cardiac stress and might even remain chronically high due to cancer-related inflammatory process)
| | - CKMB/CK (particularly in the setting of suspected false-positive troponin values)
| | - Cardiac catheterization (±endomyocardial biopsy), cardiac MRI or cardiac CT based on clinical situation

ICI - immune checkpoint inhibitor; ECG - electrocardiography; BNP - B-type natriuretic peptide; CK - creatine kinase; MRI - magnetic resonance imaging; CT - computed tomography
**Figure 3. Follow-up of the patient according to the response to corticosteroid treatment in ICI-related myocarditis**

ICI - immune checkpoint inhibitors; ECG - electrocardiogram; LVEF - left ventricular ejection fraction; IVIg - intravenous immunoglobulin; ATG - anti-thymocyte globulin

*There are no prospective or randomized controlled studies evaluating treatment options for ICI-related myocarditis. The available information is based on case series experience. There is no clear information about the optimal duration and dose for corticosteroid therapy. However, based on the experience, it is recommended to continue the treatment for 4-6 weeks and to reduce it according to clinical and troponin levels.

**The American Society of Clinical Oncology (ASCO) guidelines recommend the initial dose for corticosteroid therapy as 1 mg/kg/day oral or intravenous (IV). In case series in the literature, it is reported that faster recovery in clinical and troponin levels and lower rates of major adverse cardiovascular events (MACE) are observed during follow-up period with a high initial dose (1000 mg/day/IV 3-5 days) of corticosteroid treatment.

**Figure 4. Management of immune checkpoint inhibitor-related myocarditis-ASCO recommendations**

- **Grade 1**
  - High cardiac biomarkers or abnormal ECG findings
  - *Interrupt ICI*
  - *Perform cardiac monitoring and follow-up*
  - *Treatment of symptoms, if any*

- **Grade 2**
  - Elevated cardiac biomarkers or abnormal ECG or abnormal ECHO findings on top of mild symptoms
  - *Stop ICI permanently*
  - *Perform cardiac monitoring and follow-up*
  - *Transfer the patient to the coronary intensive care unit if there is an increase in troponin level or a new-onset changes on ECG*
  - *Start high dose (methyl) prednisolone orally or intravenously at a dose of 1-2 mg/kg depending on the clinical symptoms and troponin elevation. Continue and taper the dose within 4-6 weeks until the patient reaches Grade 1.***
  - *Initiate heart failure management strategies simultaneously.*

- **Grade 3**
  - Symptoms on mild activity or moderately impaired screening tests
  - *Recommendations in Grade 2 hold true.*

- **Grade 4**
  - Moderate signs of decompensation, life-threatening conditions where intravenous medication and intervention are required
  - *Recommendations in Grade 2 hold true.*
  - *However, if there is no response to high-dose steroid therapy, (methylprednisolone should be given as 1 g/day (3 days), followed by 1 mg/kg/day for 4-6 weeks) within 24 hours, additional immunosuppressive strategies (mycophenolate or anti-thymocyte globulin or infliximab) should be considered.*
Society of Clinical Oncology (ASCO) recommendations (based on expert opinions in the presence of possible or definitive myocarditis) are presented in Figure 4 (31).

Clinical course and prognosis
Future evolution of dilated cardiomyopathy has been reported in one-third of subjects with ICI-related myocarditis (38, 39). Mortality rate with the used of monotherapy (nivolumab) is around 10%, while it exceeds 60% in those receiving combination therapy (nivolumab + ipilimumab) (25). On the other hand, whether the occurrence of immune-mediated adverse events during ICI therapy is associated with the degree of antitumor activity still remains to be established (40, 41). Clinical decision-making for resumption of ICI following successful immunosuppression was suggested to be based on the severity of initial ICI-related myocarditis. It seems noteworthy that half of the patients were reported to suffer myocarditis recurrence or another immune-related event when ICI treatment was re-initiated following recovery from myocarditis. In general, restarting ICIs in survivors of life-threatening cardiotoxicity associated with these agents is not recommended (39, 41).

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