Management of Immune-Related Adverse Events in Patients Treated With Immune Checkpoint Inhibitor Therapy: American Society of Clinical Oncology Clinical Practice Guideline


ABSTRACT

Purpose
To increase awareness, outline strategies, and offer guidance on the recommended management of immune-related adverse events in patients treated with immune checkpoint inhibitor (ICPi) therapy.

Methods
A multidisciplinary, multi-organizational panel of experts in medical oncology, dermatology, gastroenterology, rheumatology, pulmonology, endocrinology, urology, neurology, hematology, emergency medicine, nursing, trialist, and advocacy was convened to develop the clinical practice guideline. Guideline development involved a systematic review of the literature and an informal consensus process. The systematic review focused on guidelines, systematic reviews and meta-analyses, randomized controlled trials, and case series published from 2000 through 2017.

Results
The systematic review identified 204 eligible publications. Much of the evidence consisted of systematic reviews of observational data, consensus guidelines, case series, and case reports. Due to the paucity of high-quality evidence on management of immune-related adverse events, recommendations are based on expert consensus.

Recommendations
Recommendations for specific organ system–based toxicity diagnosis and management are presented. While management varies according to organ system affected, in general, ICPi therapy should be continued with close monitoring for grade 1 toxicities, with the exception of some neurologic, hematologic, and cardiac toxicities. ICPI therapy may be suspended for most grade 2 toxicities, with consideration of resuming when symptoms revert to grade 1 or less. Corticosteroids may be administered. Grade 3 toxicities generally warrant suspension of ICPIs and the initiation of high-dose corticosteroids (prednisone 1 to 2 mg/kg/d or methylprednisolone 1 to 2 mg/kg/d). Corticosteroids should be tapered over the course of at least 4 to 6 weeks. Some refractory cases may require infliximab or other immunosuppressive therapy. In general, permanent discontinuation of ICPIs is recommended with grade 4 toxicities, with the exception of endocrinopathies that have been controlled by hormone replacement. Additional information is available at www.asco.org/supportive-care-guidelines and www.asco.org/guidelineswiki.

ASSOCIATED CONTENT

Appendix
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Data Supplement
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INTRODUCTION

Immune checkpoint inhibitors (ICPIs) have revolutionized the treatment of many different types of cancers. These inhibitors work by blocking pathways called checkpoints. These checkpoint pathways are mechanisms for the human immune system to control the immune response. The immune checkpoint proteins cytotoxic T-lymphocyte–associated-4 (CTLA-4) and programmed cell death protein 1 (PD-1) are
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Guideline Question
How should clinicians manage immune-related adverse events (irAEs) in adult patients with cancer treated with immune checkpoint blockade antibodies?

Target Population
Adult patients with cancer receiving treatment with immune checkpoint blockade inhibitors alone.

Target Audience
Health care practitioners, including oncologists, medical specialists, emergency medicine, family practitioners, nurses, and pharmacists, who provide care to patients with cancer as well as patients receiving immune checkpoint inhibitors (ICPis) and their caregivers.

Methods
An Expert Panel was convened to develop clinical practice guideline recommendations based on a systematic review of the medical literature.

Recommendations
The following are general recommendations that should be followed irrespective of affected organs. For organ-specific management, see Tables 1, 2, 3, 4, 5, 6, 7, 8, 9 and 10. (Note: Definition of grades are found in each table and, for the most part, follow the Common Terminology Criteria for Adverse Events [version 5.0]).

It is recommended that clinicians manage toxicities as follows:

- Patient and family caregivers should receive timely and up-to-date education about immunotherapies, their mechanism of action, and the clinical profile of possible irAEs prior to initiating therapy and throughout treatment and survivorship.
- There should be a high level of suspicion that new symptoms are treatment related.
- In general, ICPi therapy should be continued with close monitoring for grade 1 toxicities, with the exception of some neurologic, hematologic, and cardiac toxicities.
- Hold ICPis for most grade 2 toxicities and consider resuming when symptoms and/or laboratory values revert to grade 1 or less. Corticosteroids (initial dose of 0.5 to 1 mg/kg/d of prednisone or equivalent) may be administered.
- Hold ICPis for grade 3 toxicities and initiate high-dose corticosteroids (prednisone 1 to 2 mg/kg/d or methylprednisolone IV 1 to 2 mg/kg/d). Corticosteroids should be tapered over the course of at least 4 to 6 weeks. If symptoms do not improve with 48 to 72 hours of high-dose corticosteroid, infliximab may be offered for some toxicities.
- When symptoms and/or laboratory values revert to grade 1 or less, rechallenging with ICPis may be offered; however, caution is advised, especially in those patients with early-onset irAEs. Dose adjustments are not recommended.
- In general, grade 4 toxicities warrant permanent discontinuation of ICPis, with the exception of endocrinopathies that have been controlled by hormone replacement.

All recommendations in this guideline are based on expert consensus, benefits outweigh harms, moderate strength of recommendation.

ASCO believes that cancer clinical trials are vital to inform medical decisions and improve cancer care and that all patients should have the opportunity to participate.
receptors expressed on the surface of cytotoxic T cells that interact with their ligands CD80/CD86 in the case of CTLA-4 and programmed death-ligand 1 (PD-L1) in the case of PD-1. These pathways can be co-opted to help cancer cells to evade cytotoxic T-cell–mediated death. ICPis work by preventing the receptors and ligands from binding to each other, thereby disrupting signaling.1

Currently, there are several ICPis approved by the US Food and Drug Administration. Ipilimumab, an anti–CTLA-4 antibody, was the first agent approved for use in patients with advanced melanoma.2 Pembrolizumab and nivolumab target PD-1 and have been approved for melanoma, metastatic non–small-cell lung cancer (NSCLC), head and neck squamous cancers, urothelial carcinoma, gastric adenocarcinoma, and mismatch-repair–deficient solid tumors as well as for classic Hodgkin lymphoma.2 Nivolumab is approved for use for hepatocellular carcinoma and patients with renal cell carcinoma. The combination of ipilimumab and nivolumab for patients with advanced melanoma has also received US Food and Drug Administration approval.2 Most recently, PD-L1 antibodies atezolizumab (approved for use in urothelial cancers and NSCLC), durvalumab (approved for use in urothelial cancers), and avelumab (approved for use in Merkel cell carcinoma and urothelial carcinoma) have also been developed to block the PD-1 pathway. The indications for use continue to expand at a rapid pace. Development of novel ICPi agents and combinations continue to be evaluated for multiple indications. Thus, this field is rapidly changing.

Despite the often durable clinical benefits of the immune checkpoint blockade therapy, ICPi use is associated with a spectrum of adverse effects related to the mechanism of action that is quite different from other systemic therapies such as cytotoxic chemotherapy. The adverse effects can affect multiple organs of the body and are most commonly seen in the skin; GI tract; lungs; and endocrine, thyroid, adrenal, pituitary, musculoskeletal, renal, nervous, hematologic, cardiovascular, and ocular systems, and there should be a high level of suspicion that any changes are treatment-related (Appendix Fig A1, online only). ICPi therapy can usually continue in the presence of mild immune-related adverse events (irAEs) with close monitoring. However, moderate to severe irAEs may be associated with severe declines in organ function and quality of life, and fatal outcomes have been reported; hence, these toxicities require early detection and proper management. Use of ICPis in patients with preexisting autoimmune disease or history of prior organ transplant requires an especially thoughtful discussion of potential risks and benefits.

In recognition of an increasing need for guidance, ASCO and the National Comprehensive Cancer Network partnered to develop guidelines on the management of irAEs. Organizational representation from the Society for Immunotherapy of Cancer, the American Society of Hematology, and the Oncology Nursing Society and informal collaboration with the Friends of Cancer Research and the Parker Institute helped to ensure coordination of efforts and a harmonization of recommended care options for this patient population. With the increasing use of immunotherapy in cancer treatment regimens, it is imperative that clinicians are knowledgeable about the symptoms associated with these agents, their recommended management, and how best to monitor for them.

This clinical practice guideline addresses one overarching clinical question: How should clinicians manage irAEs in adult patients with cancer treated with immune checkpoint blockade antibodies?

**GUIDELINE QUESTION**

This clinical practice guideline addresses one overarching clinical question: How should clinicians manage irAEs in adult patients with cancer treated with immune checkpoint blockade antibodies?

**METHODS**

**Guideline Development Process**

A multidisciplinary, multi-organizational panel of experts in medical oncology, dermatology, gastroenterology, rheumatology, pulmonology, endocrinology, urology, neurology, hematology, emergency medicine, nursing, trialist, and advocacy was convened to develop the clinical practice guideline (Appendix Table A1, online only). The Expert Panel met in person, via teleconference, and webinar and corresponded through e-mail. Based on the consideration of the evidence, the authors were asked to contribute to the development of the guideline, provide critical review, and finalize the guideline recommendations. Members of the Expert Panel were responsible for reviewing and approving the penultimate version of guideline, which was then circulated for external review and submitted to *Journal of Clinical Oncology* for editorial review and consideration for publication. All ASCO guidelines are ultimately reviewed and approved by the Expert Panel and the ASCO Clinical Practice Guideline Committee prior to publication. All funding for the administration of this project was provided by ASCO.

ASCO guidelines are based on systematic reviews of the literature. A protocol for each systematic review defines parameters for a targeted literature search. Additional parameters include relevant study designs, literature sources, types of reports, and prespecified inclusion and exclusion criteria for literature identified. The protocol for this guideline was reviewed and approved by the ASCO Clinical Practice Guidelines Committee's Supportive Care Guideline Advisory Group.

Study eligibility was guided by the population, intervention, comparator, and outcome (PICO) framework as described in the Cochrane Handbook for Systematic Reviews of Interventions. In addition, the review took into account specific timing, setting, and study design as appropriate. The PICO criteria for the studies that were included in this review are as follows:

- **Population:** Adult patients with cancer receiving treatment with immune checkpoint blockade inhibitors alone (not in combination with chemotherapy)
- **Intervention:** Corticosteroids; immunosuppressive therapy; dose modification or discontinuation of therapy; organ-specific management, including hormone replacement, disease-modifying antirheumatic drugs (DMARDs), plasmapheresis, hospitalization, consultation to subspecialties, and best supportive care
- **Comparator:** No intervention or best supportive care
- **Outcomes:** Hospitalization, discontinuations of immunotherapy due to AE, AE-related morbidity or mortality, organ dysfunction based on organ system affected, required treatment due to irAEs, retreatment with immunotherapy, recovery from AEs, and health-related quality of life

The searches were designed and conducted by a team of expert medical librarians at Doctor Evidence in established clinical and medical bibliographic databases by using a range of Medical Subject Headings, EMTREE, and free-text terms based on the PICO criteria. All searches were peer reviewed by a senior Doctor Evidence (DOC) librarian. Bibliographic sources included MEDLINE In-Process via PubMed, Embase via OvidSP, and Cochrane Central Register of Control Trial via Wiley.
All study selection and screening were conducted using the DOC Library software platform (Doctor Evidence). DOC Library is a Web-based platform featuring duplicate removal, keyword emphasis (coloring or bolding of keywords), and search and ranking functionalities and can assign and manage reasons for exclusion. Before screening began, duplicate studies and those that did not meet language or date restrictions were excluded. Screening guidelines based on the protocol were then developed by consensus between methodology staff and the lead librarian and checked by a senior methodologist.

The screening procedure was conducted based on a two-step process: (1) title/abstract screening and (2) full-text screening. At both stages, the reasons for exclusion were documented. Full-text screening was conducted by two reviewers. Discrepancies between reviewers were resolved by an independent third reviewer. Articles were excluded from the systematic review if they were editorials, commentaries, letters, news articles, narrative reviews, or published in a non-English language.

The guideline recommendations are crafted, in part, by using the Guidelines Into Decision Support (GLIDES) methodology. In addition, a guideline implementability review was conducted. Based on the implementability review, revisions were made to the draft to clarify recommended actions for clinical practice.

Detailed information about the methods used to develop this guideline is available in the Methodology Supplement at www.asco.org/supportive-care-guidelines, including an overview (eg, panel composition, development process, revision dates), literature search and data extraction, recommendation development process, and quality assessment.

The ASCO Expert Panel and guidelines staff will work with co-chairs to keep abreast of any substantive updates to the guideline. Based on formal review of the emerging literature, ASCO will determine the need to update. The Methodology Supplement (available at www.asco.org/supportive-care-guidelines) provides additional information about the signals approach to guideline updating.

This is the most recent information as of the publication date. Visit the ASCO Guidelines Wiki at www.asco.org/guidelinewiki to submit new evidence.

All abbreviations used in this Guideline can be found in Appendix Table A3, online only.

**Guideline Disclaimer**

The Clinical Practice Guidelines and other guidance published herein are provided by the American Society of Clinical Oncology, Inc. (ASCO) to assist providers in clinical decision making. The information herein should not be relied upon as being complete or accurate, nor should it be considered as inclusive of all proper treatments or methods of care or as a statement of the standard of care. With the rapid development of scientific knowledge, new evidence may emerge between the time information is developed and when it is published or read. The information is not continually updated and may not reflect the most recent evidence. The information addresses only the topics specifically identified therein and is not applicable to other interventions, diseases, or stages of diseases. This information does not mandate any particular course of medical care. Further, the information is not intended to substitute for the independent professional judgment of the treating provider, as the information does not account for individual variation among patients. Recommendations reflect high, moderate, or low confidence that the recommendation reflects the net effect of a given course of action. The use of words like “must,” “must not,” “should,” and “should not” indicates that a course of action is recommended or not recommended for either most or many patients, but there is latitude for the treating physician to select other courses of action in individual cases. In all cases, the selected course of action should be considered by the treating provider in the context of treating the individual patient. Use of the information is voluntary. ASCO provides this information on an “as is” basis and makes no warranty, express or implied, regarding the information. ASCO specifically disclaims any warranties of merchantability or fitness for a particular use or purpose. ASCO assumes no responsibility for any injury or damage to persons or property arising out of or related to any use of this information, or for any errors or omissions.

**Guideline and Conflicts of Interest**

The Expert Panel was assembled in accordance with ASCO’s Conflict of Interest Policy Implementation for Clinical Practice Guidelines (“Policy,” found at http://www.asco.org/rwc). All members of the Expert Panel completed ASCO’s disclosure form, which requires disclosure of financial and other interests, including relationships with commercial entities that are reasonably likely to experience direct regulatory or commercial impact as a result of promulgation of the guideline. Categories for disclosure include employment; leadership; stock or other ownership; honoraria, consulting or advisory role; speaker’s bureau; research funding; patents, royalties, other intellectual property; expert testimony; travel, accommodations, expenses; and other relationships. In accordance with the Policy, the majority of the members of the Expert Panel did not disclose any relationships constituting a conflict under the Policy.

**RESULTS**

A total of 38 systematic reviews and 166 primary studies met the eligibility criteria of the systematic review. Much of the evidence consisted of systematic reviews of observational data, consensus guidelines, case series, and case reports. Due to the limitations of the available evidence, the guideline relied on informal consensus for the recommendations. Use of formal consensus methodology was deemed unnecessary, favoring open discussion that allowed for articulation of views and opinions instead. Dissenting opinions, when raised, are noted.

**RECOMMENDATIONS**

**Clinical Question**

How should clinicians manage irAEs in adult patients with cancer treated with immune checkpoint blockade antibodies? All recommendations in this guideline are expert consensus based, with benefits outweighing harms, and a moderate strength of recommendation.

**1.0 Skin Toxicities**

Please refer to Table 1 for a complete set of recommendations, definition of grades, and additional considerations.

**1.1 RASH/INFLAMMATORY DERMATITIS**

**Recommendation 1.1a – Diagnostic Work-up.** It is recommended that for all grades, the diagnostic work-up should include the following:

- Pertinent history and physical examination.
- Rule out any other etiology of the skin problem, such as an infection, an effect of another drug, or a skin condition linked to another systemic disease or unrelated primary skin disorder.
- A biologic checkup, including a blood cell count, liver, and kidney tests, may be performed if needed.
- Directed serologic studies if an autoimmune condition is suspected, such as lupus or dermatomyositis: a screening antinuclear antibody (ANA) test, SS-A/Anti-Ro, and SS-B/Anti-La if the rash is predominantly photodistributed or demonstrating photosensitivity. Consider expanding serologic studies or diagnostic work-up if other autoimmune conditions are considered.
- Skin biopsy, clinical photography may be performed when indicated.
### Table 1. Management of Skin irAEs in Patients Treated With ICPis

#### 1.0 Skin Toxicities

#### 1.1 Rash/inflammatory dermatitis

**Definition:** Erythema multiforme minor is a targetoid reaction in the skin and mucous membranes usually triggered by infections, such as herpes simplex viruses, but can be associated with an immune-related drug eruption and if progresses to erythema multiforme major, it can be a harbinger of SCAR, such as SJS, lichenoid (resembling the flat-topped, polygonal, and sometimes scaly or hypertrophic lesions of lichen-planus), eczematous (inflammatory dermatitis characterized by pruritic, erythematous, scaly, or crusted papules or plaques on the skin, which is vulnerable to superinfection, psoriasisform (resembling the well-demarcated, erythematous, and scaly papules and plaques of psoriasis), morbilliform (a nonpustular, nonbullous measles-like exanethematous rash of the skin often referred to as "maculopapular" and without systemic symptoms or laboratory abnormalities, excluding occasional isolated peripheral eosinophilia, palmoplantar erythrodysthesia [hand-foot syndrome; redness, numbness, burning, itching, and superficial desquamation of the palms and soles], neutrophilic dermatoses [eg, Sweet syndrome], and others)

#### Diagnostic work-up

**Pertinent history and physical examination**
- Rule out any other etiology of the skin problem, such as an infection, an effect of another drug, or a skin condition linked to another systemic disease or unrelated primary skin disorder
- If needed, a biologic checkup, including a blood cell count and liver and kidney tests

**Directed serologic studies**
- If an autoimmune condition is suspected, such as lupus or dermatomyositis: a screening antinuclear antibody test, SS-A/Anti-Ro, SS-B/Anti-La if predominantly photodistributed/photosensitivity, antihistone, double-stranded DNA, and other relevant serologies. Consider expanding serologic studies or diagnostic work-up if other autoimmune conditions are considered based on signs, symptoms

**Skin biopsy**
- Consider clinical monitoring with use of serial clinical photography
- Review full list of patient medications to rule out other drug-induced cause for photosensitivity

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
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<tbody>
<tr>
<td><strong>G1:</strong> Symptoms do not affect the quality of life or controlled with topical regimen and/or oral antipruritic</td>
<td>Continue ICPi</td>
</tr>
<tr>
<td><strong>G2:</strong> Inflammatory reaction that affects quality of life and requires intervention based on diagnosis</td>
<td>Treat with topical emollients and/or mild-moderate potency topical corticosteroids</td>
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<tr>
<td><strong>G3:</strong> As G2 but with failure to respond to indicated interventions for a G 2 dermatitis</td>
<td>Systemic corticosteroids: IV (methyl)prednisolone (or equivalent) dosed at 1-2 mg/kg, tapering over at least 4 weeks</td>
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<tr>
<td><strong>G4:</strong> All severe rashes unmanageable with prior interventions and intolerable</td>
<td>Immediately hold ICPi and consult dermatology to determine appropriateness of resuming ICPi therapy upon resolution of skin toxicity and once corticosteroids are reduced to prednisone (or equivalent) ≤ 10 mg</td>
</tr>
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#### 1.2 Bullous dermatoses

**Definition:** Including bullous pemphigoid or other autoimmune bullous dermatoses, bullous drug reaction

#### Diagnostic work-up

**Physical examination**
- Rule out any other etiology of the skin problem, such as an infection, an effect of another drug, or a skin condition linked to another systemic disease
- If needed, a biologic checkup, including a blood cell count, liver, and kidney tests; consider serum antibody tests to rule out bullous pemphigoid or other autoimmune bullous dermatoses

**Referral to dermatology for bullae that are not explained by infectious or transient other causes (eg, herpes simplex virus, herpes zoster, bullous impetigo, bullous insect bite, friction or pressure blister)**
- Consider skin biopsy (both hematoxylin and eosin evaluation of lesional skin and direct immunofluorescence evaluation of perilesional skin)

<table>
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<tr>
<th>Grading</th>
<th>Management</th>
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<tr>
<td><strong>G1:</strong> Asymptomatic, bullae covering &lt; 10% BSA and no associated erythema</td>
<td>If bullae are &lt; 10% BSA, asymptomatic, and noninflammatory (such as the case with friction bullae or pressure bullae), cessation of ICPi is not necessary, and only observation and/or local wound care is warranted. When symptomatic bullae or erosions, which are denoted vesicles or bullae, are observed on the skin or mucosal surfaces, the cutaneous irAE is by definition considered at least G2</td>
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(continued on following page)
Table 1. Management of Skin irAEs in Patients Treated With ICPs (continued)

1.0 Skin Toxicities

<table>
<thead>
<tr>
<th>G2: Blistering that affects quality of life and requires intervention based on diagnosis not meeting criteria for grade &gt; 2</th>
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<tr>
<td>Blisters covering 10%-30% BSA</td>
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<tr>
<td>Hold ICP therapy and consult with dermatology for work-up and to determine appropriateness of resuming</td>
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<tr>
<td>Attention given to general local wound care, which includes plain petrolatum ointment and bandages or plain petrolatum ointment gauze and bandage over any open erosions, which are left over on the skin after the blister has popped or if the roof of the blister easily sloughs off</td>
</tr>
<tr>
<td>Counsel patients to avoid skin irritants and overexposure to sun, wear protective clothing, use sunscreens</td>
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<tr>
<td>Work-up for autoimmune bullous disease as above</td>
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<tr>
<td>Initiate class 1 high-potency topical corticosteroid (eg, clobetasol, betamethasone or equivalent) and reassess every 3 days for progression or improvement</td>
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<tr>
<td>Low threshold to initiate treatment with prednisone (or equivalent) at 0.5-1 mg/kg dosing and taper over at least 4 weeks</td>
</tr>
<tr>
<td>Monitor patients with G2 irAEs closely for progression to involvement of greater BSA and/or mucous membrane involvement. Consider following patients closely using serial photography</td>
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<tr>
<td>Primer on monitoring for complicated cutaneous adverse drug reactions:</td>
</tr>
<tr>
<td>• Review of systems: Skin pain (like a sunburn), fevers, malaise, myalgias, arthralgias, abdominal pain, ocular discomfort or photophobia, sores or discomfort in the nares, sores or discomfort in the oropharynx,odynophagia, hoarseness, dysuria, sores or discomfort in the vaginal area for women or involving the meatus of the penis for men, sores in the perianal area, or pain with bowel movements</td>
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<tr>
<td>• Physical examination: Include vital signs and a full skin examination specifically evaluating all skin surfaces and mucous membranes (eyes, nares, oropharynx, genitals, and perianal area). Assess for lymphadenopathy, facial or distal extremity swelling (may be signs of DIHS/DRESS). Assess for pustules or blisters or erosions in addition to areas of &quot;dusky erythema,&quot; which may feel painful to palpation. To assess for a positive Nikolsky sign, place a gloved finger tangentially over erythematous skin and apply friction parallel to the skin surface. Nikolsky sign is positive if this results in detached or sloughing epidermis demonstrating poor attachment of the epidermis to the dermis, which is the case in some autoimmune disorders (eg, pemphigus) and SJS/TEN</td>
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<tr>
<th>G3: Skin sloughing covering &gt; 30% BSA with associated pain and limiting self-care ADL</th>
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<tbody>
<tr>
<td>Hold ICP therapy and consult with dermatology to determine appropriateness of resuming</td>
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<tr>
<td>Administer IV (methyl)prednisolone (or equivalent) 1-2 mg/kg, tapering over at least 4 weeks</td>
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<tr>
<td>If bullous pemphigoid is diagnosed, it may be possible to avoid long-term use of systemic corticosteroids and treat with rituximab, as an alternative approach to treating the irAE</td>
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<tr>
<td>Seek infectious disease consultation if patient might have secondary cellulitis or if patient has other infection risk factors, such as neutropenia, etc</td>
</tr>
<tr>
<td>Work-up for autoimmune bullous disease as above</td>
</tr>
<tr>
<td>Primarily evaluate all skin surfaces and mucous membranes (eyes, nares, oropharynx, genitals, and perianal area). Assess for lymphadenopathy, facial or distal extremity swelling (may be signs of DIHS/DRESS) Assess for pustules or blisters or erosions in addition to areas of &quot;dusky erythema,&quot; which may feel painful to palpation. To assess for a positive Nikolsky sign, place a gloved finger tangentially over erythematous skin and apply friction parallel to the skin surface. Nikolsky sign is positive if this results in detached or sloughing epidermis demonstrating poor attachment of the epidermis to the dermis, which is the case in some autoimmune disorders (eg, pemphigus) and SJS/TEN</td>
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<tr>
<th>G4: Blister covering &gt; 30% BSA and associated fluid or electrolyte abnormalities</th>
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<tr>
<td>Permanently discontinue ICP</td>
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<tr>
<td>Admit patient immediately and place under supervision of a dermatologist</td>
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<tr>
<td>Administer IV (methyl)prednisolone (or equivalent) 1-2 mg/kg with tapering over at least 4 weeks when the toxicity resolves</td>
</tr>
<tr>
<td>If bullous pemphigoid is diagnosed, it may be possible to avoid long-term use of systemic corticosteroids and treat with rituximab as an alternative approach to treating the irAE</td>
</tr>
<tr>
<td>Seek infectious disease consultation if patient might have secondary cellulitis or if patient has other infection risk factors, such as neutropenia, etc</td>
</tr>
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1.3 SCARs, including SJS, TEN, acute generalized exanthematous pustulosis, and DRESS/DIHS

Definition: Severe changes in either structure or functions of the skin, the appendages or the mucous membranes due to a drug

Diagnostic work-up

Total body skin examination with attention to examining all mucous membranes as well as complete review of systems

Rule out any other etiology of the skin problem, such as an infection, an effect of another drug, or a skin condition linked to another systemic disease

A biologic checkup, including a CBC with differential test, and liver and kidney function tests, including urinalysis, in addition to the blood work; if the patient is febrile, blood cultures should be considered as well

Skin biopsies to assess for full-thickness epidermal necrosis, as is seen in SJS/TEN, as well as other possible etiologies like paraneoplastic pemphigus or other autoimmune blistering dermatoses or other drug reactions, such as acute generalized exanthematous pustulosis

Consider following patients closely using serial clinical photography

If mucous membrane involvement or blistering is observed on the skin, consider early admission to a burn center for further monitoring and management

Primarily evaluate all skin surfaces and mucous membranes (eyes, nares, oropharynx, genitals, and perianal area). Assess for lymphadenopathy, facial or distal extremity swelling (may be signs of DIHS/DRESS). Assess for pustules or blisters or erosions in addition to areas of "dusky erythema," which may feel painful to palpation. To assess for a positive Nikolsky sign, place a gloved finger tangentially over erythematous skin and apply friction parallel to the skin surface. Nikolsky sign is positive if this results in detached or sloughing epidermis demonstrating poor attachment of the epidermis to the dermis, which is the case in some autoimmune disorders (eg, pemphigus) and SJS/TEN
Clinicians manage grade 1 toxicities as follows:

1. **G1: NA**
   - For SCAFs, there is no G1 category; if lower BSA is involved with bullae or erosions, there should remain a high concern that this reaction will progress to G3 or G4.

2. **G2: Morbilliform ("maculopapular") exanthem covering 10%-30% BSA with systemic symptoms, lymphadenopathy, or facial swelling**
   - Hold ICPI and monitor patients closely every 3 days with G2 irAEs for progression to involvement of greater BSA and/or mucous membrane involvement.
   - Consider following patients closely by serial photography.
   - Initiate therapy with topical emollients, oral antihistamines, and medium- to high-potency topical corticosteroids.
   - Consider initiation of prednisone (or equivalent) 0.5-1 mg/kg tapered over at least 4 weeks.

3. **G3: Skin sloughing covering < 10% BSA with mucosal involvement associated signs (eg, erythema, purpura, epidermal detachment, mucous membrane detachment)**
   - Hold ICPI therapy and consult with dermatology.
   - Treat skin with topical emollients and other petrolatum emollients, oral antihistamines, and high-potency topical corticosteroids; dimethicone may also be offered as an alternative to petrolatum.
   - Administer IV (methyl)prednisolone (or equivalent) 0.5-1 mg/kg and convert to oral corticosteroids on response, clean over at least 4 weeks.
   - Admit to burn and/or consult wound services with attention to supportive care, including fluid and electrolyte balance, minimizing insensible water losses, and preventing infection.
   - Given the immune mechanism of action of these medicines, use of immune suppression is warranted and should be offered.
   - For mucous membrane involvement of SJS or TEN, appropriate consulting services should be offered to guide management in preventing sequelae from scarring (eg, ophthalmology; ear, nose, and throat; urology; gynecology; etc, as appropriate).

4. **G4: Skin erythema and blistering/sloughing covering > 10% BSA with associated signs (eg, erythema, purpura, epidermal detachment, mucous membrane detachment) and/or systemic symptoms and concerning associated blood work abnormalities (eg, liver function test elevations in the setting of DRESS/DIHS)**
   - Permanently discontinue ICPI.
   - Admit patient immediately to a burn unit or ICU with consulted dermatology and wound care services.
   - Consider further consultations based on management of mucosal surfaces (eg, ophthalmology; urology; gynecology; ear, nose, and throat surgery; etc).
   - Admit patient immediately to a burn unit or ICU with consulted dermatology and wound care services.
   - Consider pain palliative consultation and/or admission in patients presenting with unresponsive cases.
   - IVIG or cyclosporine may also be considered in severe or corticosteroid-unresponsive cases.
   - Consider pain palliative consultation and/or admission in patients presenting with unresponsive cases.
   - Adequate suppression is necessary with corticosteroids or other agents and may be prolonged in cases of DRESS/DIHS.

**Additional considerations:** The usual prohibition of corticosteroids for SJS is not relevant here, as the underlying mechanism is a T-cell immunodirected toxicity. Adequate suppression is necessary with corticosteroids or other agents and may be prolonged in cases of DRESS/DIHS.

**Management:**

**1.0 Skin Toxicities**

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>All grades</td>
<td>In cases of suspected SJS or any mucous membrane involvement, discontinue ICPI treatment and monitor closely for improvement, regardless of grade.</td>
</tr>
<tr>
<td>G1: NA</td>
<td>For SCAFs, there is no G1 category; if lower BSA is involved with bullae or erosions, there should remain a high concern that this reaction will progress to G3 or G4.</td>
</tr>
<tr>
<td>G2: Morbilliform (&quot;maculopapular&quot;) exanthem covering 10%-30% BSA with systemic symptoms, lymphadenopathy, or facial swelling</td>
<td>Hold ICPI and monitor patients closely every 3 days with G2 irAEs for progression to involvement of greater BSA and/or mucous membrane involvement. Consider following patients closely by serial photography. Initiate therapy with topical emollients, oral antihistamines, and medium- to high-potency topical corticosteroids. Consider initiation of prednisone (or equivalent) 0.5-1 mg/kg tapered over at least 4 weeks.</td>
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<tr>
<td>G3: Skin sloughing covering &lt; 10% BSA with mucosal involvement associated signs (eg, erythema, purpura, epidermal detachment, mucous membrane detachment)</td>
<td>Hold ICPI therapy and consult with dermatology. Treat skin with topical emollients and other petrolatum emollients, oral antihistamines, and high-potency topical corticosteroids; dimethicone may also be offered as an alternative to petrolatum. Administer IV (methyl)prednisolone (or equivalent) 0.5-1 mg/kg and convert to oral corticosteroids on response, clean over at least 4 weeks. Admit to burn and/or consult wound services with attention to supportive care, including fluid and electrolyte balance, minimizing insensible water losses, and preventing infection. Given the immune mechanism of action of these medicines, use of immune suppression is warranted and should be offered. For mucous membrane involvement of SJS or TEN, appropriate consulting services should be offered to guide management in preventing sequelae from scarring (eg, ophthalmology; ear, nose, and throat; urology; gynecology; etc, as appropriate).</td>
</tr>
<tr>
<td>G4: Skin erythema and blistering/sloughing covering &gt; 10% BSA with associated signs (eg, erythema, purpura, epidermal detachment, mucous membrane detachment) and/or systemic symptoms and concerning associated blood work abnormalities (eg, liver function test elevations in the setting of DRESS/DIHS)</td>
<td>Permanently discontinue ICPI. Admit patient immediately to a burn unit or ICU with consulted dermatology and wound care services. Consider further consultations based on management of mucosal surfaces (eg, ophthalmology; urology; gynecology; ear, nose, and throat surgery; etc). Admit patient immediately to a burn unit or ICU with consulted dermatology and wound care services. Consider pain palliative consultation and/or admission in patients presenting with unresponsive cases. IVIG or cyclosporine may also be considered in severe or corticosteroid-unresponsive cases. Consider pain palliative consultation and/or admission in patients presenting with unresponsive cases. Adequate suppression is necessary with corticosteroids or other agents and may be prolonged in cases of DRESS/DIHS.</td>
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**Recommendation 1.1b – Management.** It is recommended that clinicians manage grade 1 toxicities as follows:

- Should continue to offer ICPI.
- Should treat skin with topical emollients (if predominately dry skin is observed) and/or mild to moderate potency (hydrocortisone 2.5% or equivalent to triamcinolone 0.1% or equivalent) topical corticosteroids (signs of inflammation/redness with or without itching).
- Should counsel patients to avoid skin irritants and sun exposure.
- It is recommended that clinicians manage grade 2 toxicities, including intermittent pruritus, as follows:
  - May hold ICPI and monitor weekly for improvement. If not resolved, interrupt treatment until skin AE has reverted to grade 1 or less and consider dermatology referral.
  - Should treat skin with topical emollients, oral antihistamines, and medium- to high-potency topical corticosteroids.
  - In addition, consider initiating prednisone (or equivalent) at dosing 1 mg/kg tapering over at least 4 weeks, depending on primary skin lesions observed on examination.
- It is recommended that clinicians manage grade 3 toxicities, including constant pruritus, as follows:
  - Should hold ICPI therapy and consult with dermatology, if available, to determine appropriateness of resuming.
  - Should treat skin with topical emollients, oral antihistamines, and high-potency topical corticosteroids.
  - Initiate intravenously (IV) (methyl)prednisolone (or equivalent) dosed at 1 to 2 mg/kg and taper over at least 4 weeks.
  - If not resolved, refer to dermatology.
- It is recommended that clinicians manage grade 4 toxicities as follows:
  - Should immediately hold ICPI and consult dermatology to determine appropriateness of resuming ICPI therapy upon resolution of skin toxicity and once corticosteroids are reduced to prednisone (or equivalent) 10 mg or less.

**Abbreviations:** ADL, activities of daily living; BSA, body surface area; CTCAE, Common Terminology Criteria for Adverse Events; DIHS, drug-induced hypersensitivity syndrome; DRESS, drug reaction with eosinophilia and systemic symptoms; G, grade; ICPI, immune checkpoint inhibitor; ICU, intensive care unit; irAE, immune-related adverse event; IV, intravenous; IVIG, intravenous immunoglobulin; NA, not applicable; SCAR, severe cutaneous adverse reactions; SJS, Stevens-Johnson syndrome; TENS, toxic epidermal necrolysis.

**Table 1. Management of Skin irAEs in Patients Treated With ICPIs (continued)**

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<td>G2: Morbilliform (&quot;maculopapular&quot;) exanthem covering 10%-30% BSA with systemic symptoms, lymphadenopathy, or facial swelling</td>
<td>Hold ICPI and monitor patients closely every 3 days with G2 irAEs for progression to involvement of greater BSA and/or mucous membrane involvement. Consider following patients closely by serial photography. Initiate therapy with topical emollients, oral antihistamines, and medium- to high-potency topical corticosteroids. Consider initiation of prednisone (or equivalent) 0.5-1 mg/kg tapered over at least 4 weeks.</td>
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<td>G3: Skin sloughing covering &lt; 10% BSA with mucosal involvement associated signs (eg, erythema, purpura, epidermal detachment, mucous membrane detachment)</td>
<td>Hold ICPI therapy and consult with dermatology. Treat skin with topical emollients and other petrolatum emollients, oral antihistamines, and high-potency topical corticosteroids; dimethicone may also be offered as an alternative to petrolatum. Administer IV (methyl)prednisolone (or equivalent) 0.5-1 mg/kg and convert to oral corticosteroids on response, clean over at least 4 weeks. Admit to burn and/or consult wound services with attention to supportive care, including fluid and electrolyte balance, minimizing insensible water losses, and preventing infection. Given the immune mechanism of action of these medicines, use of immune suppression is warranted and should be offered. For mucous membrane involvement of SJS or TEN, appropriate consulting services should be offered to guide management in preventing sequelae from scarring (eg, ophthalmology; ear, nose, and throat; urology; gynecology; etc, as appropriate).</td>
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<td>Permanently discontinue ICPI. Admit patient immediately to a burn unit or ICU with consulted dermatology and wound care services. Consider further consultations based on management of mucosal surfaces (eg, ophthalmology; urology; gynecology; ear, nose, and throat surgery; etc). Admit patient immediately to a burn unit or ICU with consulted dermatology and wound care services. Consider pain palliative consultation and/or admission in patients presenting with unresponsive cases. IVIG or cyclosporine may also be considered in severe or corticosteroid-unresponsive cases. Consider pain palliative consultation and/or admission in patients presenting with unresponsive cases. Adequate suppression is necessary with corticosteroids or other agents and may be prolonged in cases of DRESS/DIHS.</td>
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**All recommendations are expert consensus based, with beneﬁts outweighing harms, and strength of recommendations are moderate**
• Should administer IV (methyl)prednisolone (or equivalent) dosed at 1 to 2 mg/kg, with slow tapering when the toxicity resolves.
• Should monitor closely for progression to severe cutaneous adverse reaction (SCAR).
• Should admit patient immediately with direct oncology involvement and with an urgent consult by dermatology.
• Consider alternative antineoplastic therapy over resuming ICPIs if the skin irAE does not resolve to grade 1 or less. If ICPIs are the patient’s only option, consider restarting once these adverse effects have resolved to a grade 1 level.

1.2 Bullous Dermatoses

Recommendation 1.2a – Diagnostic Work-up. It is recommended that for all grades of irAEs the diagnostic work-up should include the following:
• Comprehensive physical examination, including evaluation of all mucous membranes.
• Rule out any other etiology of the skin problem, such as an infection, an effect of another drug, or a skin condition linked to another systemic disease or unrelated primary skin disorder.
• If needed, a biologic checkup may be performed, including a blood cell count, liver and kidney tests, hepatitis antibody tests, and tuberculosis (TB) testing.
• Referral to dermatology for blisters that are not explained by infectious/transient other causes (eg, herpes simplex, herpes zoster infections, pressure/ friction bullae).
• Skin biopsy (lesional biopsy of inflamed skin or the edge of a bulla or vesicle) for hematoxylin and eosin histology and biopsy of a perilesional or “near-inflamed” area for direct immunofluorescence testing.
• If the biopsy demonstrates a subepidermal blister and/or the direct immunofluorescence testing is suspicious or positive for a diagnosis of bullous pemphigoid (BP), or in cases where skin biopsies are not possible, consider serum testing to further evaluate tense bullae (BP 230 and BP 130 enzyme-linked immunosorbent assay serum testing). If negative, under the guidance of dermatology, sending the patient serum for indirect immunofluorescent testing to rule out other autoimmune blistering diseases could be considered.

Recommendation 1.2b – Management. If blisters are < 10% body surface area (BSA), asymptomatic, and noninflammatory (eg, the case with friction blisters or pressure blisters), cessation of ICPI is not necessary, and only observation and/or local wound care is warranted. Once a blister or erosion, which is essentially a deroofed blister, is observed on examination, with associated erythema or blister, is essentially a deroofed blister, the reaction should be considered due to ICPI therapy and graded 2 or above. It is recommended that clinicians manage grade 2 toxicities as follows:
• Should hold ICPI therapy and consult with dermatology (or skin care team, which may include general surgeon) to determine appropriateness of resuming ICPI and initiate general local skin/wound care, which includes plain petrolatum ointment and bandages or plain petrolatum ointment gauze and bandage over any open erosions that are left over on the skin after the blister has popped or if the roof of the blister easily sloughs off.
• Should counsel patients to avoid skin irritants and over-exposure to sun, wear protective clothing, and use sunscreens.
• Should order work-up for autoimmune bullous disease as above.
• Initiate class 1 high-potency topical corticosteroid (eg, clobetasol, betamethasone or equivalent) and reassess every 3 days for progression or improvement.
• Lower threshold to initiate treatment with prednisone (or equivalent) at 0.5 to 1 mg/kg dosing and taper over at least 4 weeks.
• Monitor patients with grade 2 irAEs closely for progression to involvement of greater BSA and/or mucous membrane involvement. Consider following patients closely using serial photography.
• Primer on monitoring for complicated cutaneous adverse drug reactions:
  o Review of systems: skin pain (“like a sunburn”), fevers, malaise, myalgias, arthralgias, abdominal pain, ocular discomfort or photophobia, sores or discomfort in the nares, sores or discomfort in the oropharynx, dysuria, hoarseness, dysuria, sores or discomfort in the vaginal area for women or involving the meatus of the penis for men, sores in the perianal area, or pain with bowel movements.
  o Physical examination: include vital signs and a full skin examination, specifically evaluating all skin surfaces and mucous membranes (eyes, nares, oropharynx, genitalia, and perianal area). Assess for lymphadenopathy, facial or distal extremity swelling (may be signs of drug-induced hypersensitivity syndrome [DIHS]/drug reaction with eosinophilia and systemic symptoms [DRESS]). Assess for pustules or blisters or erosions in addition to areas of “dusky erythema,” which may feel painful to palpation. To assess for a positive Nikolsky sign, place a gloved finger tangentially over erythematous skin and apply friction parallel to the skin surface. Nikolsky sign is positive if this results in detached or sloughing epidermis, demonstrating poor attachment of the epidermis to the dermis, which is the case in some autoimmune disorders (eg, pemphigus) and Stevens-Johnson syndrome (SJS)/toxic epidermal necrolysis (TEN).

It is recommended that clinicians manage grade 3 toxicities as follows:
• Should hold ICPI therapy and consult with dermatology to determine appropriateness of resuming.
• Should administer IV (methyl)prednisolone (or equivalent) at 1 to 2 mg/kg dosing tapered over at least 4 weeks.
• If BP is diagnosed, it may be possible to avoid long-term use of systemic corticosteroids and treat with rituximab as an alternative approach to treating the irAE.
• Seek infectious disease consultation if patient might have secondary cellulitis or if patient has other infection risk factors, such as neutropenia, etc.

It is recommended that clinicians manage grade 4 toxicities as follows:
• Should permanently discontinue ICPI.
• Should admit patient immediately and place under supervision of a dermatologist.
• Should administer IV (methyl)prednisolone (or equivalent) 1 to 2 mg/kg/d. When toxicity improves to grade 2 or less, start corticosteroid taper. Taper should be at least 4 weeks.
• If BP is diagnosed, it may be possible to avoid long-term use of systemic corticosteroids and treat with rituximab as an alternative to treating the irAE.

• Seek infectious disease consultation if patient might have secondary cellulitis or if patient has other infection risk factors, such as neutropenia, etc.

1.3 Severe Cutaneous Adverse Reactions

Recommendation 1.3a – Diagnostic work-up. Severe cutaneous adverse reactions, or SCARs, include, but are not limited to, SJS/TEN and DRESS (also called DIHS).

It is recommended that for all grades of irAEs, the diagnostic work-up should include the following:

• Total body skin examination with attention to ALL mucous membranes as well as a complete review of systems.

• Rule out any other etiology of the skin problem, such as an infection, an effect of another drug, or a skin condition linked to another systemic disease.

• A biologic checkup, including a CBC with differential test, and liver and kidney function tests, including urinalysis, in addition to the blood work. If the patient is febrile, blood cultures should be considered as well.

• Skin biopsies to assess for full-thickness epidermal necrosis, as is seen in SJS/TEN, as well as other possible etiologies like paraneoplastic pemphigus or other autoimmune blistering dermatoses, or other drug reactions, such as acute generalized exanthematous pustulosis.

• Consider following patients closely using serial clinical photography.

• If mucous membrane involvement or blistering is observed on the skin, consider early admission to a burn center for further monitoring and management.

• Primer on monitoring for complicated cutaneous adverse drug reactions:

  o Review of systems: skin pain (“like a sunburn”), fevers, malaise, myalgias, arthralgias, abdominal pain, ocular discomfort or photophobia, sores or discomfort in the nares, sores or discomfort in the oropharynx, odynophagia, hoarseness, dysuria, sores or discomfort in the vaginal area for women or involving the meatus of the penis for men, sores in the perianal area, or pain with bowel movements.

  o Physical examination: include vital signs and a full skin examination specifically evaluating all skin surfaces and mucous membranes (eyes, nares, oropharynx, genitals, and perianal area). Assess for lymphadenopathy, facial or distal extremity swelling (may be signs of DIHS/DRESS). Assess for pustules or blisters or erosions in addition to areas of dusky erythema, which may feel painful to palpation. To assess for a positive Nikolsky sign, place a gloved finger tangentially over erythematous skin and apply friction parallel to the skin surface. Nikolsky sign is positive if this results in detached or sloughing epidermis, demonstrating poor attachment of the epidermis to the dermis, which is the case in some autoimmune disorders (eg, pemphigus) and SJS/TEN.

Recommendation 1.3b – Management. In cases of suspected SJS or any mucous membrane involvement, it is recommended that clinicians should discontinue ICPI treatment and refer to dermatology. It would not be advisable to restart ICPI unless “cleared” by a dermatologist if SJS/TEN is suspected.

For SCARs, there is no grade 1 category. If lower BSA is involved with bullae or erosions, there should remain a high concern that this reaction will progress to grade 3 or 4.

It is recommended that clinicians manage grade 2 toxicities as follows:

• Hold ICPI and monitor patients closely every 3 days with grade 2 irAEs for progression to involvement of greater BSA and/or mucous membrane involvement.

• Consider following patients closely by using serial photography.

• Initiate therapy with topical emollients, oral antihistamines, and medium- to high-strength topical corticosteroids.

• Consider initiation of prednisone or equivalent at 0.5 to 1 mg/kg tapered over at least 4 weeks.

It is recommended that clinicians manage grade 3 toxicities as follows:

• Should hold ICPI therapy and consult with dermatology.

• Should treat skin with topical emollients and other petrolatum emollients, oral antihistamines, and high-strength topical corticosteroids. Dimethicone may also be offered as an alternative to petrolatum.

• Administer IV (methyl)prednisolone (or equivalent) at doses of 1 to 2 mg/kg and taper over at least 4 weeks.

• Admit to burn and/or consult wound services with attention to supportive care, including fluid and electrolyte balance, minimizing insensible water losses, and preventing infection.

• Given the immune mechanism of action of these medicines, use of immune suppression, such as with systemic corticosteroids, is warranted and should be offered, though the use of systemic corticosteroids has been more controversial for the treatment of SJS/TEN, in general. For DRESS/DIHS, high-dose and usually prolonged courses of systemic corticosteroids is first-line therapy following cessation of the offending drug.

• For mucous membrane involvement of SJS or TEN, appropriate consulting services should be offered to guide management in preventing sequelae from scarring (eg, ophthalmology; ear, nose, and throat; urology; gynecology; etc, as appropriate).

• Seek infectious disease consultation if patient might have secondary cellulitis or if patient has other infection risk factors, such as neutropenia, etc.

It is recommended that clinicians manage grade 4 toxicities as follows:

• Should permanently discontinue ICPI.

• Should admit patient immediately with consideration to a burn unit or ICU in the case of SJS/TEN and consult dermatology.

• Administer IV (methyl)prednisolone or equivalent 1 to 2 mg/kg with tapering when the toxicity resolves to normal.

• May consider IV immunoglobulin (IVIG) or cyclosporine as an alternative or in corticosteroid-refractory cases.

• Seek infectious disease consultation if patient might have secondary cellulitis or if patient has other infection risk factors, such as neutropenia, etc.
Qualifying statement. The usual prohibition of corticosteroids for SJS is not relevant here, as the underlying mechanism is a T-cell immunodirected toxicity. Adequate suppression is necessary with corticosteroids or other agents and may be prolonged in cases of DRESS/DIHS. Additionally, patients with DRESS/DIHS may experience other autoimmune diseases as long-term sequelae, such as thyroid disease and recurrences as systemic corticosteroids are tapered or discontinued are not uncommon. Thus, continued monitoring should be considered for these patients. It is generally advisable to avoid rechallenge with an offending drug when a patient experiences SCARs, such as SJS/TEN or DRESS/DIHS. It is advisable to consider alternate antineoplastic therapies because rechallenge in these cases may result in an even more severe SCAR.

Discussion. The dramatic and durable responses seen with ICPIs are often at the cost of increased toxicities due to unrestrained activity of T cells. Among the diverse irAEs, cutaneous toxicities such as rash, pruritus, and vitiligo are by far the most common and the earliest to occur. Although most cutaneous toxicities are transient, they can cause significant morbidity and impairment of patients’ health-related quality of life.

Cutaneous toxicities are reported in 30% to 50% of patients treated with ICPIs. Our understanding of cutaneous toxicities stems mostly from the ipilimumab experience wherein the overall incidence ranges from 37% to 70% for all-grade and 1% to 3% for grade 3 or higher cutaneous toxicities. Cutaneous toxicities are less frequently reported with anti–PD-1 agents (17% to 37%); however, the incidence of grade 3 or higher toxicities is the same as with ipilimumab.

Cutaneous toxicities pose a myriad of challenges. Rash is the most common cutaneous toxicity reported with ICPIs. They span a variety of inflammatory conditions, including spongiotic, psoriasiform, and lichenoid dermatitides, mimicking eczema, psoriasis, and lichen planus, respectively. The clinical presentations vary with focal to diffuse distributions, including flexural, inverse, and erythrodermic variants. Pruritus can be severe and is the most common associated symptom. Vitiligo presents as well-demarcated depigmented macules and patches, reported exclusively in patients with melanoma. Besides varying clinical presentation, the time to onset varies greatly among these rashes, as vitiligo can appear months after treatment initiation; however, the inflammatory dermatoses usually present within the first one to two cycles of treatment. This mandates constant vigilance for signs and symptoms of cutaneous toxicities. In addition, these irAEs are increasingly recognized as a contributing factor to treatment noncompliance, discontinuation, or dose modification. As targeted systemic therapies are available for eczema and psoriasis, correlating the inflammatory patterns of the cutaneous toxicities with the inflammatory patterns that they mimic may lead to more efficacious treatments, fewer drug interruptions and dose modifications, and increased compliance and efficacy of the immune ICPIs. However, classification of rashes has not been undertaken prospectively, and histologic characterization of the cutaneous toxicities is lacking.

Interestingly, emerging data suggest that development of cutaneous toxicity, especially rash and vitiligo, may correlate with response to ICPI therapy in patients with metastatic melanoma. In a retrospective analysis of 148 patients with melanoma treated with nivolumab plus peptide vaccine or nivolumab, survival benefit was reported in patients who developed rash (n = 64) or vitiligo (n = 19). Overall survival (OS) was significantly longer in patients who developed rash (hazard ratio [HR], 0.423; 95% CI, 0.243 to 0.735; P = .001) and vitiligo (HR, 0.184; 95% CI, 0.036 to 0.94; P = .012). Objective response rate (ORR) was also significantly higher in patients with rash (P = .03) or vitiligo (P = .009). In a prospective study evaluating pembrolizumab in treatment of patients with melanoma, ORR was higher in patients who developed vitiligo than in those who did not (71% vs 28%; P = .002). Similarly, in a phase I study of ipilimumab for patients with melanoma, rash was the most common irAE reported among responders. Furthermore, in a meta-analysis of 27 studies in patients with melanoma treated with various immunotherapeutic agents, vitiligo was significantly associated with both progression-free survival (HR, 0.51; 95% CI, 0.32 to 0.82; P = .005) and OS (HR, 0.25; 95% CI, 0.10 to 0.61; P = .003). These findings from large clinical development programs suggest that cutaneous irAEs may be a surrogate for clinical benefit, and it would be important to correctly identify these skin changes so that the ICPI therapy is not discontinued in these cases with good prognoses. In addition, many cutaneous toxicities may be managed without the discontinuation of therapy. With early diagnosis and prompt management of cutaneous toxicities, patients may be able to stay on ICPI therapy, which could be crucial for improved treatment outcomes. However, little is known about the underlying mechanisms and the relationship between cutaneous toxicity and clinical outcome in patients with advanced cancer other than melanoma. This lack of knowledge presents challenges for prompt diagnosis and hampers the development of strategies to mitigate or minimize the occurrence of cutaneous toxicities in patients treated with ICPIs.

With increasing use of ICPIs in the clinic, characterization and development of sensitive and robust markers of cutaneous toxicity is a priority.

2.0 GI Toxicities

Please refer to Table 2, for a complete set of recommendations, definition of grades, and additional considerations.

2.1 Colitis

Recommendation 2.1a – Diagnostic work-up. No specific diagnostic work-up is recommended for grade 1 adverse events. It is recommended that the diagnostic work-up should include the following for grade 2 toxicity:

- Work-up of blood (CBC, comprehensive metabolic panel, thyroid-stimulating hormone [TSH], erythrocyte sedimentation rate [ESR], C-reactive protein [CRP]), stool (culture, Clostridium difficile, parasite, cytomegalovirus [CMV] or other viral etiology, ova and parasite) should be performed.
- May test for lactoferrin for patient stratification to determine who needs urgent endoscopy, and calprotectin may be offered to follow up on disease activity.
- Screening laboratories (HIV, hepatitis A and B, and blood quantifier for TB) to prepare patients to start infliximab should be routinely done in patients at high risk for those infections and in appropriately selected patients based on infectious disease expert’s evaluation.
### 2.1 Colitis

**Definition:** A disorder characterized by inflammation of the colon. 

**Diagnostic work-up**

| G2 | Work-up of blood (CBC, comprehensive metabolic panel, TSH, ESR, CRP), stool (culture, *Clostridium difficile*, parasite, CMV or other viral etiology, ova and parasite) should be performed. Consider testing for lactoferrin (for patient stratification to determine who needs more urgent endoscopy) and calprotectin (to follow up on disease activity). Screening laboratories (HIV, hepatitis A and B, and blood quantiferon for TB) to prepare patients to start immunosuppressive agents. Imaging (eg, CT scan of abdomen and pelvis and GI endoscopy with biopsy) should be considered as there is evidence showing that the presence of ulceration in the colon can predict a corticosteroid-refractory course, which may require early infliximab. Consider repeating endoscopy for patients who do not respond to immunosuppressive agents. Repeating endoscopy for disease monitoring can be considered when clinically indicated and when planning to resume therapy. 
| G3-4 | All the work-up listed for G2 (blood, stool, imaging, and scope with biopsy) should be completed immediately. Consider repeating endoscopy for patients who do not respond to immunosuppressive agents. Repeating endoscopy for disease monitoring should only be considered when clinically indicated and when planning to resume ICPI. 

<table>
<thead>
<tr>
<th>Grading (based on CTCAE for diarrhea, as most often used clinically)</th>
<th>Management</th>
</tr>
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</table>
| All patients | Counsel all patients to be aware of and inform their health care provider immediately if they experience:
Abdominal pain, nausea, cramping, blood or mucus in stool or changes in bowel habits
Fever, abdominal distention, obstruction, constipation
For G2 or higher, consider permanently discontinuing CTLA-4 agents and may restart PD-1, PD-L1 agents if patient can recover to G1 or less; concurrent immunosuppressant maintenance therapy should be considered only if clinically indicated in individual cases. May also include supportive care with medications such as lomudim if infection has been ruled out.
Should consult with gastroenterology for G2 or higher.
Administer corticosteroids, unless diarrhea is transient, starting with initial dose of 1 mg/kg/day prednisone or equivalent.
When symptoms improve to G1 or less, taper corticosteroids over at least 4-6 weeks before resuming treatment, although resuming treatment while on low-dose corticosteroid may also be an option after an evaluation of the risks and benefits.
EGD/colonoscopy, endoscopy evaluation should be highly recommended for cases grade ≥ 2 to stratify patients for early treatment with infliximab based on the endoscopic findings and to determine the safety of resuming PD-1, PD-L1 therapy.
Stool inflammatory markers can be considered (lactoferrin and calprotectin) in cases of G2 or higher to differentiate functional v inflammatory diarrhea, and use calprotectin to monitor treatment response if provider prefers.
Repeat colonoscopy is optional for cases of G2 or higher for disease activity monitoring to achieve complete remission, especially if there is a plan to resume ICPI.
May obtain gastroenterology consult for prolonged G1 cases.

| G1 | Increase of fewer than four stools per day over baseline; mild increase in ostomy output compared with baseline | Continue ICPI; alternatively, ICPI may be held temporarily and resumed if toxicity does not exceed G1.
Monitor for dehydration and recommend dietary changes.
Facilitate expedited phone contact with patient/caregiver.
May obtain gastroenterology consult for prolonged G1 cases.

| G2 | Increase of four to six stools per day over baseline; moderate increase in ostomy output compared with baseline | Should hold ICPI temporarily until patient’s symptoms recover to G1; can consider permanently discontinuing CTLA-4 agents and may restart PD-1, PD-L1 agents if patient can recover to G1 or less.
Concurrent immunosuppressant maintenance therapy (< 10 mg prednisone equivalent dose) may be offered only if clinically indicated in individual cases.
May also include supportive care with medications such as lomudim if infection has been ruled out.
Should consult with gastroenterology for G2 or higher.
Administer corticosteroids, unless diarrhea is transient, starting with initial dose of 1 mg/kg/day prednisone or equivalent.
When symptoms improve to G1 or less, taper corticosteroids over at least 4-6 weeks before resuming treatment, although resuming treatment while on low-dose corticosteroid may also be an option after an evaluation of the risks and benefits.
EGD/colonoscopy, endoscopy evaluation should be highly recommended for cases grade ≥ 2 to stratify patients for early treatment with infliximab based on the endoscopic findings and to determine the safety of resuming PD-1, PD-L1 therapy.
Stool inflammatory markers can be considered (lactoferrin and calprotectin) in cases of G2 or higher to differentiate functional v inflammatory diarrhea, and use calprotectin to monitor treatment response if provider prefers.
Repeat colonoscopy is optional for cases of G2 or higher for disease activity monitoring to achieve complete remission, especially if there is a plan to resume ICPI.

| G3 | Increase of seven or more stools per day over baseline, incontinence, hospitalization indicated, severe increase in ostomy output compared with baseline, limiting self-care ADL | Should consider permanently discontinuing CTLA-4 agents and may restart PD-1, PD-L1 agents if patient can recover to G1 or less.
Administer corticosteroids (initial dose of 1-2 mg/kg/d prednisone or equivalent).
Consider hospitalization or outpatient facility for patients with dehydration or electrolyte imbalance.
If symptoms persist ≥ 3-5 days or recur after improvement, consider administering IV corticosteroid or noncorticosteroid (eg, infliximab).
Consider colonoscopy in cases where patients have been on immunosuppression and may be at risk for opportunistic infections as an independent cause for diarrhea (ie, CMV colitis) and for those who are anti-TNF or corticosteroid refractory.

| G4 | Life-threatening consequences; urgent intervention indicated | Permanently discontinue treatment.
Should admit patient when clinically indicated; patients managed as outpatients should be very closely monitored.
Administer 1-2 mg/kg methylprednisolone or equivalent until symptoms improve to G1, and then taper over 4-6 weeks.
Consider early infliximab 5-10 mg/kg if symptoms refractory to corticosteroid within 2-3 days.
Consider lower GI endoscopy if symptoms are refractory despite treatment or there is concern of new infections.

(continued on following page)
### Additional considerations

The use of vedolizumab may be considered in patients refractory to infliximab and/or contraindicated to TNF-α blocker. The decision should be made on an individual basis from gastroenterology and oncology evaluation. This is based on case series showing promising results. Patients with hepatitis and irAE colitis are rare, and management should include permanently discontinuing ICPi and offering other immunosuppressant agents that work systemically for both conditions. Currently, enteritis alone as the cause of diarrhea is uncommon and requires small bowel biopsy as the evaluation tool. It may be managed similar as colitis, including corticosteroid and/or infliximab, etc.

### 2.2 Hepatitis

**Definition:** A disorder characterized by a viral pathologic process involving the liver parenchyma.

**Diagnostic work-up**

Monitor patient for abnormal liver blood tests: AST, ALT, and bilirubin prior to each infusion and/or weekly if G1 liver function test elevations. No treatment is recommended for G1 liver function test abnormality.

For G2 or higher:

- Work-up for other causes of elevated liver enzymes should be tested, viral hepatitis, alcohol history, iron study, thromboembolic event, liver ultrasound, cross-sectional imaging for potential liver metastasis from primary malignancy. If suspicion for primary autoimmune hepatitis is high, can consider ANAs, antismooth muscle antibodies, antineutrophil cytoplasmic antibodies. If patients with elevated alkaline phosphatase alone, γ-glutamyl transferase should be tested. For isolated elevation of transaminases, consider checking CK for other etiologies.

#### Grading Management

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
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| All patients | Counsel all patients to be aware of and inform their health care provider immediately if they experience:  
- Yellowing of skin or whites of the eyes  
- Severe nausea or vomiting  
- Pain on the right side of the abdomen  
- Drowsiness  
- Dark urine (tea colored)  
- Bleeding or bruising more easily than normal  
- Feeling less hungry than usual |
| G1: Asymptomatic (AST or ALT > ULN to 3.0 × ULN and/or total bilirubin > ULN to 1.5 × ULN) | Continue ICPi with close monitoring; consider alternate etiologies  
Monitor laboratories one to two times weekly  
Manage with supportive care for symptom control |
| G2: Asymptomatic (AST or ALT > 3.0 to ≤ 5 ULN and/or total bilirubin > 1.5 to ≤ 3 ULN) | Hold ICPi temporarily and resume if recover to G1 or less on prednisone ≤ 10 mg/d  
For grade 2 hepatic toxicity with symptoms, monitor corticosteroid 0.5-1 mg/kg prednisone or equivalent if the abnormal elevation persists with significant clinical symptoms in 3-5 days  
Increase frequency of monitoring to every 3 days  
Infliximab might not be the most appropriate treatment option in the situation of immune-mediated hepatitis given the potential risk of idiosyncratic liver failure (Note: No clear evidence shows the liver toxicity from infliximab from other studies)  
In follow-up, may resume ICPi treatment followed by taper only when symptoms improve to G1 or less and corticosteroid = 10 mg/d; taper over at least 1 month  
 Patients should be advised to stop unnecessary medications and any known hepatotoxic drugs |
| G3: Symptomatic liver dysfunction, fibrosis by biopsy, compensated cirrhosis, reactivation of chronic hepatitis (AST or ALT 5-20 × ULN and/or total bilirubin 3-10 × ULN) | Permanently discontinue ICPi  
Immediately start corticosteroid 1-2 mg/kg methylprednisolone or equivalent  
If corticosteroid refractory or no improvement after 3 days, consider mycophenolate mofetil or azathioprine (if using azathioprine should test for thiopurine methyltransferase deficiency)  
Laboratories at daily or every other day; consider inpatient monitoring for patients with AST/ALT > 8 × ULN and/or elevated TB 3 × ULN  
Increase frequency of monitoring to every 1-2 days  
Infliximab might not be the most appropriate treatment option in the situation of immune-mediated hepatitis given the potential risk of liver failure (Note: No clear evidence shows the liver toxicity from infliximab from other studies); alternatives include non-TNF-α agents as systemic immunosuppressants  
If no improvement is achieved with corticosteroids or for patients on combination therapy with a novel agent, with standard chemotherapy, or with targeted therapy, refer to hepatologist for further pathologic evaluation of hepatitis  
Corticosteroid taper can be attempted around 4-6 weeks; re-escalate if needed; optimal duration unclear |
| G4: Decompensated liver function (eg, ascites, coagulopathy, encephalopathy, coma; AST or ALT > 20 × ULN and/or total bilirubin > 10 × ULN) | Permanently discontinue ICPi  
Administer 2 mg/kg/d methylprednisolone equivalents  
If corticosteroid refractory or no improvement after 3 days, consider mycophenolate mofetil  
Monitor laboratories daily; consider inpatient monitoring  
Avoid the use of infliximab in the situation of immune-mediated hepatitis  
Hepatology consult if no improvement was achieved with corticosteroid  
Corticosteroid taper can be attempted around 4-6 weeks when symptoms improve to G1 or less; re-escalate if G1 or less; optimal duration unclear  
Consider transfer to tertiary care facility if necessary |

**Abbreviations:** ADL, activities of daily living; ANA, antinuclear antibody; CK, creatine kinase; CMV, cytomegalovirus; CRP, C-reactive protein; CT, computed tomography; CTCAE, Common Terminology Criteria for Adverse Events; CTLA-4, cytotoxic T-cell lymphocyte-4; EGD, esophagogastroduodenoscopy; ESR, erythrocyte sedimentation rate; G, grade; ICPi, immune checkpoint inhibitor; irAE, immune-related adverse event; IV, intravenous; PD-1, programmed death 1; PD-L1, programmed death ligand 1; TB, tuberculosis; TNF, tumor necrosis factor; TSH, thyroid-stimulating hormone; ULN, upper limit of normal.
• Imaging with computed tomography (CT) scan of abdomen and pelvis and GI endoscopy with biopsy may be performed as there is evidence showing that the presence of ulceration in the colon can predict a corticosteroid-refractory course, which may require early infliximab. Infliximab or other tumor necrosis factor (TNF)–blocking agent should not be delayed while awaiting the results of these screening tests.
• Repeat endoscopy may be offered to patients who do not respond to immunosuppressive agents. Repeating endoscopy for disease monitoring should only be offered when clinically indicated and when planning to resume therapy.

It is recommended that the diagnostic work-up should include the following for grade 3 to 4 toxicity:
• All the work-up listed for G2 (blood, stool, imaging, and scope with biopsy) should be completed immediately.
• Repeat endoscopy may be offered for patients who do not respond to immunosuppressive agents. Repeating endoscopy for disease monitoring should only be offered when clinically indicated and when planning to resume ICPI.

Recommendation 2.1b – Management. It is recommended that clinicians counsel all patients to be aware of and inform their health care provider immediately if they experience:
• Abdominal pain, nausea, cramping, blood or mucus in stool, or changes in bowel habits
• Fever, abdominal distention, obstipation, constipation

It is recommended that clinicians manage grade 1 toxicities as follows:
• May continue ICPI. Alternatively, ICPI may be held temporarily and resumed if toxicity does not exceed grade 1.
• Should monitor for dehydration and recommend dietary changes.
• Should facilitate expedited phone contact with patient/caregiver.
• May obtain gastroenterology consult for prolonged grade 1 cases.

It is recommended that clinicians manage grade 2 toxicities as follows:
• Should hold the ICPI until patient’s symptoms recover to grade 1 or less. Can consider permanently discontinuing CTLA-4 agents and may restart PD-1, PD-L1 agents if patient can recover to grade 1 or less.
• Concurrent immunosuppressant maintenance therapy (< 10 mg prednisone equivalent dose) may be offered only if clinically indicated in individual cases.
• May also include supportive care with medications such as Imodium if infection has been ruled out.
• Should consult with gastroenterology for grade 2 or higher.
• Should administer corticosteroids, unless diarrhea is transient, starting with an initial dose of 1 mg/kg/d prednisone or equivalent.
• When symptoms improve to grade 1 or less, should taper corticosteroids over at least 4 to 6 weeks before resuming treatment, although resuming treatment while on low-dose corticosteroid may also be an option after an evaluation of the risks and benefits.
• Should offer esophagogastroduodenoscopy/colonoscopy, endoscopy evaluation for cases of grade 2 or higher to stratify patients for early treatment with infliximab based on the endoscopic findings and to determine the safety of resuming PD-1, PD-L1 therapy.
• Testing for stool inflammatory markers, lactoferrin, or calprotectin may be offered in cases of grade 2 or higher to differentiate functional versus inflammatory diarrhea. Calprotectin testing may also be offered to monitor treatment response.
• Repeat colonoscopy is optional and may be offered for cases of grade 2 or higher for disease activity monitoring to achieve complete remission, especially if there is a plan to resume ICPI.

Qualifying statement. Starting infliximab before colonoscopy is reasonable if negative infectious stool work-up is confirmed. However, prompt access to colonoscopy is advised to justify the dose and duration of infliximab. Once infliximab is indicated, patients most often have grade 2 and higher diarrhea/colitis, and most are hospitalized.

It is recommended that clinicians manage grade 3 toxicities as follows:
• Should consider permanently discontinuing CTLA-4 agents and may restart PD-1, PD-L1 agents if patient can recover to grade 1 or less.
• Should administer corticosteroids (initial dose of 1 to 2 mg/kg/d prednisone or equivalent).
• Should refer to hospitalization or outpatient facility for patients with dehydration or electrolyte imbalance.
• If symptoms persist ≥ 3 to 5 days or recur after improvement, may administer IV corticosteroid or noncorticosteroid (eg, infliximab).
• May offer colonoscopy in cases where patients have been on immunosuppression and may be at risk for opportunistic infections as an independent cause for diarrhea (ie, CMV colitis) and for those who are anti-TNF or corticosteroid refractory.

It is recommended that clinicians manage grade 4 toxicities as follows:
• Should permanently discontinue all ICPI treatment.
• Should admit patient when clinically indicated. Patients managed as outpatients should be very closely monitored.
• Should administer IV corticosteroid until symptoms improve to grade 1 and then start taper over 4 to 6 weeks.
• May offer early infliximab 5 to 10 mg/kg if symptoms are refractory to corticosteroid within 2 to 3 days.
• May offer lower GI endoscopy if symptoms are refractory despite treatment or there is concern of new infections.

Qualifying statement. The use of vedolizumab may be offered to patients refractory to infliximab and/or contraindicated to TNF-α blocker. The decision should be made on an individual basis from gastroenterology and oncology evaluation. This is based on case series showing promising results.

2.2 Hepatitis

Recommendation 2.2a – Diagnostic work-up. It is recommended that work-up should include the following:
Monitor patient for abnormal liver blood tests: AST, ALT, and bilirubin prior to each infusion and/or weekly if there are grade 1 liver function test elevations. No treatment is recommended for grade 1 liver function test abnormality.

For grade 2 or higher toxicity:

- Work-up for other causes of elevated liver enzymes, viral hepatitis, alcohol history, iron study, thromboembolic event, liver ultrasound, cross-sectional imaging for potential liver metastasis. If suspicion for primary autoimmune hepatitis is high, can consider ANA, anti-smooth muscle antibodies, and antineutrophil cytoplasmic antibodies. If patients with elevated alkaline phosphatase alone, γ-glutamyl transferase should be tested. For isolated elevation of transaminases, consider checking creatine kinase (CK) for other etiologies.

**Recommendation 2.2b – Management.** It is recommended that clinicians counsel all patients to be aware of and inform their health care provider immediately if they experience:

- Yellowing of skin or whites of the eyes
- Severe nausea or vomiting
- Pain on the right side of the abdomen
- Drowsiness
- Dark urine (tea colored)
- Bleeding or bruising more easily than normal
- Feeling less hungry than usual

It is recommended that clinicians manage grade 1 toxicities as follows:

- Should continue to offer ICPI, but with close monitoring.
- Should rule out alternate etiologies.
- Should monitor laboratories one to two times weekly.
- Should offer supportive care for symptom control.

It is recommended that clinicians manage grade 2 toxicities as follows:

- Should hold ICPI treatment temporarily and resume if recover to grade 1 or less on prednisone ≤ 10 mg/d.
- For grade 2 hepatic toxicity with symptoms, may administer corticosteroid 0.5 to 1 mg/kg/d prednisone or equivalent if the abnormal elevation persists with significant clinical symptoms in 3 to 5 days.
- Should increase frequency of monitoring to every 3 days.
- Inflimab might not be the most appropriate treatment option in the situation of immune-mediated hepatitis given the potential risk of idiosyncratic liver failure (Note: No clear evidence shows liver toxicity from infliximab from other studies).
- In follow-up, may resume ICPI treatment followed by taper only when symptoms improve to grade 1 or less on corticosteroid ≤ 10 mg/d. Taper over at least 1 month.
- Patients should be advised to stop unnecessary medications and any known hepatotoxic drugs.

It is recommended that clinicians manage grade 3 toxicities as follows:

- Should permanently discontinue treatment with ICPI.
- Should immediately administer corticosteroid 1 to 2 mg/kg methylprednisolone or equivalent.
- If corticosteroid refractory or no improvement after 3 days, may offer mycophenolate mofetil or azathioprine (if using azathioprine, should test for thiopurine methyltransferase deficiency).
- Should order laboratories daily or every other day; may offer inpatient monitoring for patients with AST/ALT more than eight times the upper limit of normal (ULN) and/or elevated TB three times ULN.
- Should increase frequency of monitoring to every 1 to 2 days.
- Infliximab might not be the most appropriate treatment option in the situation of immune-mediated hepatitis given the potential risk of liver failure (Note: No clear evidence shows liver toxicity from infliximab from other studies). Alternatives include non–TNF-α agents as systemic immunosuppressants.
- If no improvement is achieved with corticosteroids or for patients on combination therapy with a novel agent, with standard chemotherapy, or with targeted therapy, refer to hepatologist for further pathologic evaluation of hepatitis.
- Corticosteroid taper should be attempted over a period of 4 to 6 weeks, re-escalate if needed, optimal duration unclear.

It is recommended that clinicians manage grade 4 toxicities as follows:

- Should permanently discontinue treatment with ICPI.
- Should administer 2 mg/kg/d methylprednisolone equivalents.
- If corticosteroid refractory or no improvement after 3 days, may offer mycophenolate mofetil.
- Should monitor laboratories daily; inpatient monitoring may be offered.
- Should not offer infliximab in the situation of immune-mediated hepatitis.
- Should refer to hepatology if no improvement is achieved with corticosteroid.
- Corticosteroid taper should be attempted over a period of 4 to 6 weeks when symptoms improve to grade 1 or less, re-escalate if needed, optimal duration unclear.
- Consider transfer to tertiary care facility if necessary.

**Discussion.** GI toxicities are some of the most common complications reported with ICPI use. While the frequency of colitis reported in the literature ranges from 8% to 27%, the incidence of diarrhea is as high as 54% in patients treated with anti–CTLA-4 antibodies, especially in patients who receive anti–CTLA-4 and anti–PD-1 combination therapy. GI toxicity is less common with anti–PD-1 monotherapy, with the incidence of diarrhea reported to be ≤ 19%. In a recent meta-analysis of patients with cancer treated with ICPIs, the relative risk (RR) of all-grade diarrhea and colitis was 1.64 (95% CI, 1.19 to 2.26; P = .002) and 10.35 (95% CI, 5.78 to 18.53; P < .001), the RR of high-grade diarrhea and colitis is reported to be 4.46 (95% CI, 1.46 to 13.57; P = .008) and 15.81 (95% CI, 6.34 to 39.42; P < .001), respectively. On the contrary, RR of upper-GI symptoms (eg, vomiting) was not significant. Frequency of intestinal perforation has been described at approximately 1%. The most common clinical presentations of immune-related GI toxicities vary from very frequent and/or loose stools to colitis symptoms (eg, mucus in the stools, abdominal pain, fever, rectal bleeding). The onset of these GI symptoms is most often in the range of 5 to 10 weeks after initiation of ICPI but can occur or recur months after discontinuation of immunotherapy. While clinical factors associated with ICPI-induced colitis have not been well...
established, nonsteroidal anti-inflammatory drug (NSAID) use is reported to be associated with an increase in ICPI-induced enterocolitis, and care should be taken with NSAID use in this setting. There is a lot of similarity between ICPI-induced colitis and inflammatory bowel disease (eg, clinical presentations, radiologic findings). CT findings of ICPI-induced colitis include mesenteric vessel engorgement; bowel wall thickening; and fluid-filled colonic distention; and on positron emission tomography/CT scan, diffuse colonic wall thickening is observed. The distribution of colitis has been reported to involve descending colon more often than other parts of the colon. On the other hand, pathology from patients with colitis demonstrated changes that were more than what classic inflammatory bowel disease shows. The histologic picture is often characterized by marked mixed inflammatory cell infiltrates in the lamina propria, consisting of neutrophils, lymphocytes, plasma cells, and eosinophils. Inflammatory changes also tend to be more diffuse (75%).

For patients with mild diarrhea symptoms (grade 1), usually conservative observation and maintenance of hydration are recommended rather than more-aggressive evaluation. Once diarrhea symptoms are grade 2 or higher or with apparent colitis symptoms, corticosteroid at 1 to 2 mg/kg is still the first-line treatment option if the stool infectious work-up is negative. Endoscopic evaluation can be considered if clinically deemed critical. If the symptoms are not improving after 3 to 5 days of corticosteroid treatment, stronger immunosuppressive agents (eg, TNF-α blocker infliximab, anti-integrin α4β7 antibody vedolizumab) have been shown in multiple case reports and case series to be very effective at managing grade 2 toxicities. No significant adverse effects or negative effect on the overall survival on the patient’s outcomes were identified.

Compared with lower-GI toxicities, upper-GI toxicity, characterized by dysphagia, nausea/vomiting, and epigastric pain, is much less common. Pathology can present as patchy chronic duodenitis or chronic gastritis with rare granulomas. It can coexist with lower-GI toxicity or as an isolated condition. The treatment strategy is similar to colitis: corticosteroid followed by TNF-α blockers for refractory cases based on case studies.

Hepatotoxicity has been reported to occur in 2% to 10% of patients treated with ipilimumab, nivolumab, and pembrolizumab monotherapy. Combination treatment with ipilimumab and nivolumab has resulted in a reported 25% to 30% all-grade hepatitis and approximately 15% incidence of grade 3 toxicity. Onset develops predominately within the first 6 to 12 weeks after treatment initiation. The mainstay of treatment is prednisone or equivalent at 1 to 2 mg/kg with frequent monitoring of liver tests. However, other etiologies that could contribute to liver dysfunction have to be thoroughly evaluated and ruled out. For corticosteroid refractory cases, mycophenolate mofetil has been reported in a case study with some success. The TNF-α blocker infliximab is not recommended given the concern of liver toxicity, despite lack of evidence. Other alternative immunosuppressive agents still need further data proof for efficacy and safety. The patient with pre-existing hepatitis who experiences ICPI-induced colitis is rare but represents a management challenge. Available options are more limited and should include permanent cessation of anti–CTLA-4 and possibly other ICPI treatment.

Acute pancreatitis has also been reported in the literature, but it is rare. Routine monitoring of amylase/lipase in asymptomatic patients is not recommended unless pancreatitis is clinically suspected. In the absence of symptoms, corticosteroid treatment is not indicated for modest elevations in serum amylase and lipase.

In terms of resumption of ICPI after toxicities have occurred, the recommendation is different based on the grade level of toxicities. For grade 4 toxicities, ICPI treatment should be permanently discontinued. In patients with grade 3 or less toxicities who improve from their irAE after adequate treatment, the risk of recurrent toxicities with rechallenge appears to vary with organ toxicity and type of ICPI therapy. Only a small proportion of patients with ICPI-related colitis are reported to experience recurrences with anti–PD-1 resumption alone. Toxicities such as hepatitis and pancreatitis also have some risk of recurrence. These most often occur early and are generally low grade and manageable with standard treatments. Nonetheless, care should be taken to ensure that proper monitoring and management strategies are implemented.

### 3.0 Lung Toxicity

Please refer to Table 3 a complete set of recommendations, definition of grades, and additional considerations.

#### 3.1 Pneumonitis

**Recommendation 3.1a – Diagnostic work-up.** It is recommended that the diagnostic work-up should include the following:

- Chest x-ray (CXR), CT, pulse oximetry
- For grade 2 or higher, may include the following infectious work-up: nasal swab, sputum culture and sensitivity, blood culture and sensitivity, and urine culture and sensitivity.

**Recommendation 3.1b – Management.** It is recommended that clinicians manage grade 1 toxicities as follows:

- Should hold ICPI with radiographic evidence of pneumonitis progression.
- May offer one repeat CT in 3 to 4 weeks. In patients who have had baseline testing (Appendix Table A2, online only), may offer a repeat spirometry/diffusing capacity of lung for carbon monoxide in 3 to 4 weeks.
- May resume ICPI with radiographic evidence of improvement or resolution. If no improvement, should treat as grade 2.
- Should monitor patients weekly with history and physical examination and pulse oximetry; may also offer CXR.

It is recommended that clinicians manage grade 2 toxicities as follows:

- Should hold immunotherapy until resolution to grade 1 or less.
- Should administer prednisone 1 to 2 mg/kg/d and taper by 5 to 10 mg/wk over 4 to 6 weeks per institutional guidelines.
- May offer bronchoscopy with bronchoalveolar lavage.
- May prescribe empirical antibiotics.
- Should monitor every 3 days with history and physical examination and pulse oximetry; may also offer CXR. No clinical improvement after 48 to 72 hours of prednisone, should treat as grade 3.
management of immune-related adverse events

3.0 Lung Toxicities

3.1 Pneumonitis

Definition: Focal or diffuse inflammation of the lung parenchyma (typically identified on CT imaging)

- No symptomatic, pathologic, or radiographic features are pathognomonic for pneumonitis

Diagnostic work-up
- Should include the following: CXR, CT, pulse oximetry
- For G2 or higher, may include the following infectious work-up: nasal swab, sputum culture and sensitivity, blood culture and sensitivity, urine culture and sensitivity

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<th>Grading</th>
<th>Management</th>
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<tr>
<td>G1: Asymptomatic, confined to one lobe of the lung or &lt; 25% of lung parenchyma, medical intervention indicated, limiting instrumental ADL</td>
<td>Hold ICPI until radiographic evidence of pneumonitis progression. May offer one repeat CT in 3-4 weeks; in patients who have had baseline testing, may offer a repeat spirometry/DLCO in 3-4 weeks. May resume ICPI with radiographic evidence of improvement or resolution.</td>
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<tr>
<td>G2: Symptomatic, involves more than one lobe of the lung or 25%-50% of lung parenchyma, medical intervention indicated, limiting instrumental ADL</td>
<td>Hold ICPI until resolution to G1 or less. Prednisone 1-2 mg/kg/d and taper by 5-10 mg/week over 4-6 weeks. Consider bronchoscopy with BAL. Consider empirical antibiotics. Monitor patients weekly with history and physical examination and pulse oximetry; consider CXR. No clinical improvement after 48-72 hours of prednisone, treat as G3</td>
</tr>
<tr>
<td>G3: Severe symptoms, hospitalization required, involves all lung lobes or &gt; 50% of lung parenchyma, limiting self-care ADL, oxygen indicated</td>
<td>Permanently discontinue ICPI. Empirical antibiotics; (methyl)prednisolone IV 1-2 mg/kg/d; no improvement after 48 hours, may add infliximab 5 mg/kg or mycophenolate mofetil IV 1 g twice a day or IVIG for 5 days or cyclophosphamide; taper corticosteroids over 4-6 weeks. Pulmonary and infectious disease consults if necessary. Bronchoscopy with BAL ± transbronchial biopsy. Patients should be hospitalized for further management.</td>
</tr>
<tr>
<td>G4: Life-threatening respiratory compromise, urgent intervention indicated (intubation)</td>
<td>May resume ICPI with radiographic evidence of improvement or resolution. Consider bronchoscopy with BAL; if consistent picture is consistent with pneumonitis, no need for biopsy.</td>
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Additional considerations

- GI and Pneumocystis prophylaxis with PPI and Bactrim may be offered to patients on prolonged corticosteroid use (> 12 weeks), according to institutional guidelines.
- Consider calcium and vitamin D supplementation with prolonged corticosteroid use.
- The role of prophylactic fluconazole with prolonged corticosteroid use (> 12 weeks) remains unclear, and physicians should proceed according to institutional guidelines.
- Bronchoscopy + biopsy; if clinical picture is consistent with pneumonitis, no need for biopsy.
- All recommendations are expert consensus based, with benefits outweighing harms, and strength of recommendations are moderate.

Abbreviations: ADL, activities of daily living; BAL, bronchoalveolar lavage; CT, computed tomography; CXR, chest x-ray; DLCO, diffusing capacity of lung for carbon monoxide; G, grade; ICPI, immune checkpoint inhibitor; irAE, immune-related adverse event; IV, intravenous; IVIG, intravenous immunoglobulin; PPI, proton pump inhibitor.

It is recommended that clinicians manage grade 3 to 4 toxicities as follows:

- Should permanently discontinue ICPI.
- Should prescribe empirical antibiotics and administer (methyl)prednisolone IV 1 to 2 mg/kg/d. No improvement after 48 hours, may add infliximab 5 mg/kg or mycophenolate mofetil IV 1 g twice a day or IVIG for 5 days or cyclophosphamide. Taper corticosteroids over 4 to 6 weeks.
- Should consult pulmonary and infectious disease if necessary.
- Should offer bronchoscopy with bronchoalveolar lavage with or without transbronchial biopsy.
- Patients should be hospitalized for further management.

Qualifying statement. The role of prophylactic fluconazole with prolonged corticosteroid use (> 12 weeks) remains unclear, and physicians should proceed according to institutional guidelines.

Discussion. ICPI-related pneumonitis is an uncommon but potentially serious toxicity. The reported incidence of pneumonitis in studies investigating anti–PD-1/PD-L1 is variable and ranges from 0% to 10%,48,163 with an overall incidence of 2.7% reported in a recent meta-analysis of 20 studies with PD-1 inhibition.39

The toxicity seems to be less common with anti–CTLA-4 treatment, with pneumonitis reported in < 1% of trial participants receiving ipilimumab.40-44 A higher incidence was seen in patients who received combination therapy than those who received monotherapy (10% vs 3%, respectively; P < .001),38 and patients treated with combination immunotherapy may be less likely to experience resolution of the irAE compared with patients treated with monotherapy.40,45 In patients who improve from their irAE, rechallenging with anti–PD-(L)1 therapy was associated with recurrent or new irAEs in half of patients and was more common in early-onset irAEs.46 The majority of such patients were managed successfully, but two deaths have been reported.46

The evidence on whether the risk of ICPI-related pneumonitis and pneumonitis-related deaths varies by tumor type remains equivocal. The odds of all-grade pneumonitis was higher in patients with NSCLC than in those with melanoma (odds ratio [OR], 1.43; 95% CI, 1.08 to 1.89; P = .005) according to the Nishino et al.39 meta-analysis. Similarly, patients with renal cell carcinoma...
were also significantly more likely to experience all-grade pneumonitis than patients with melanoma (OR, 1.59; 95% CI, 1.32 to 1.92; P < .001). In contrast, other studies have reported similar rates of grades 3 to 4 pneumonitis across tumor types but with more treatment-related deaths due to pneumonitis seen in patients with NSCLC. In a multicenter, large retrospective analysis, pneumonitis was reported to develop in both former/current smokers (56%) and never smokers (44%). Recent evidence also demonstrated no significant difference in the rates of irAEs, including pneumonitis, between patients who received thoracic radiotherapy in addition to checkpoint inhibitors.

The median onset of ICPI-related pneumonitis can vary, with a range of 2 to 24 months and a median time to onset of approximately 3 months reported in the literature. However, onset does occur earlier with combination therapy versus monotherapy. Clinical symptoms can include dyspnea (53%), cough (35%), fever (12%), and chest pain (7%). Hypoxia may occur and progress rapidly, leading to respiratory failure. Ground-glass opacities or patchy nodular infiltrates, predominantly in the lower lobes, are common manifestations on chest imaging. Radiologic abnormalities vary but are often reported to be focal and very different from the diffuse pneumonitis associated with targeted agents. Naidoo et al recently reported on five distinct radiologic subtypes: chronic obstructive pneumonia like, ground-glass opacities, hypersensitivity type, interstitial type, and pneumonitis not otherwise specified.

When the clinical picture is consistent with pneumonitis, biopsy is generally unnecessary. However, transbronchial biopsy may have a role in assisting to rule out other etiologies like lymphangitic spread of tumor or infection. The decision to perform lung biopsy in the evaluation of immune-related pulmonary reactions is based on the probability that this examination will yield a specific diagnosis, leading to a change in management. Yet, there is no specific pathology to confirm immune-related pneumonitis. Ultimately, the decision to proceed with biopsy should be taken after a careful risk-benefit analysis, with the optimal technique, number, size, and location of biopsies depending upon the suspected diagnosis, the anatomic distribution of the disease process, and the availability of pulmonologists.

In addition to typical findings of pneumonitis, sarcoid-like granulomatous reactions, including subpleural micronodular opacities and hilar lymphadenopathy, as well as pleural effusions have been associated with both CTLA-4- and PD-1/PD-L1–targeted therapies. Clinical manifestations are diverse and often patient-specific and can include cough, wheezing, fatigue, chest pain, or no symptoms at all. With varying clinical presentation, it is prudent for clinicians to be aware of the possibility of such immune-related pulmonary reactions, as they may mimic disease progression on imaging and examination. Biopsy may assist in confirming the diagnosis.

### 4.0 Endocrine Toxicities

Please refer to Table 4 for a complete set of recommendations, definition of grades, and additional considerations.

**Recommendation 4.0 – Endocrine general.** It is recommended that clinicians counsel patients to inform their health care provider and team immediately if they experience any changes in their health since their last visit, especially any of the following:

- Headaches that will not go away or unusual headache patterns
- Vision changes
- Rapid heartbeat
- Increased sweating
- Extreme tiredness or weakness
- Muscle aches
- Weight gain or weight loss
- Dizziness or fainting
- Feeling more hungry or thirsty than usual
- Hair loss
- Changes in mood or behavior, such as decreased sex drive, irritability, or forgetfulness
- Feeling cold
- Constipation
- Voice gets deeper
- Urinating more often than usual
- Nausea or vomiting
- Abdominal pain

### 4.1 Thyroid

#### 4.1.1 Primary Hypothyroidism

**Recommendation 4.1.1a – Diagnostic work-up.** It is recommended that the diagnostic work-up should include the following:

- Testing for TSH and free thyroxine (FT4) every 4 to 6 weeks as part of routine clinical monitoring on therapy or for case detection in symptomatic patients.

**Recommendation 4.1.1b – Management.** It is recommended that clinicians manage grade 1 toxicities as follows:

- Should continue to offer ICPI with close follow-up and monitoring of TSH, FT4.

It is recommended that clinicians manage grade 2 toxicities as follows:

- May hold ICPI until symptoms resolve to baseline.
- May consult endocrinology.
- Should prescribe thyroid hormone supplementation in symptomatic patients with any degree of TSH elevation or in asymptomatic patients with TSH levels that persist > 10 mIU/L (measured 4 weeks apart).
- Should monitor TSH every 6 to 8 weeks while titrating hormone replacement to normal TSH.
- FT4 can be used in the short term (2 weeks) to ensure adequacy of therapy in those with frank hypothyroidism where the FT4 was initially low.
- Once adequately treated, should monitor thyroid function (at least TSH) every 6 weeks while on active therapy or as needed for symptoms to ensure appropriate replacement. Repeat testing annually or as indicated by symptoms once stable.

It is recommended that clinicians manage grade 3 to 4 toxicities as follows:

- May hold ICPI until symptoms resolve to baseline with appropriate supplementation.
- Should consult endocrinology.
- May admit for IV therapy if signs of myxedema (bradycardia, hypothermia).
### Table 4. Management of Endocrine irAEs in Patients Treated With ICPIs

#### 4.0 Endocrine Toxicity

Counsel patients to inform their health care provider immediately if they experience any changes in their health since their last visit, especially any of the following:

- Headaches that will not go away or unusual headache patterns
- Vision changes
- Rapid heartbeat
- Increased sweating
- Extreme tiredness or weakness
- Muscle aches
- Weight gain or weight loss
- Dizziness or fainting
- Feeling more hungry or thirsty than usual
- Hair loss
- Changes in mood or behavior, such as decreased sex drive, irritability, or forgetfulness
- Feeling cold
- Constipation
- Voice gets deeper
- Urinating more often than usual
- Nausea or vomiting
- Abdominal pain

#### 4.1 Thyroid

##### 4.1.1 Primary hypothyroidism

**Definition:** Elevated TSH, normal or low FT4

**Diagnostic work-up**

- TSH and FT4 every 4-6 weeks as part of routine clinical monitoring on therapy or for case detection in symptomatic patients

**Grading Management**

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1: TSH &lt; 10 mIU/L and asymptomatic</td>
<td>Should continue ICPI with close follow-up and monitoring of TSH, FT4</td>
</tr>
<tr>
<td>G2: Moderate symptoms; able to perform ADL; TSH persistently &gt; 10 mIU/L</td>
<td>May hold ICPI until symptoms resolve to baseline</td>
</tr>
<tr>
<td></td>
<td>Consider endocrine consultation</td>
</tr>
<tr>
<td></td>
<td>Prescribe thyroid hormone supplementation in symptomatic patients with any degree of TSH elevation or in asymptomatic patients with TSH levels that persist &gt; 10 mIU/L (measured 4 weeks apart)</td>
</tr>
<tr>
<td></td>
<td>Monitor TSH every 6-8 weeks while titrating hormone replacement to normal TSH levels</td>
</tr>
<tr>
<td></td>
<td>Once adequately treated, should monitor thyroid function at least TSH level every 6 weeks while on active ICPI therapy or as needed for symptoms to ensure appropriate replacement; repeat testing annually or as indicated by symptoms</td>
</tr>
<tr>
<td>G3-4: Severe symptoms, medically significant or life-threatening consequences, unable to perform ADL</td>
<td>Hold ICPI until symptoms resolve to baseline with appropriate supplementation</td>
</tr>
<tr>
<td></td>
<td>Endocrine consultation</td>
</tr>
<tr>
<td></td>
<td>May admit for IV therapy if signs of myxedema (bradycardia, hypothermia)</td>
</tr>
<tr>
<td></td>
<td>Thyroid supplementation and reassessment as in G2</td>
</tr>
</tbody>
</table>

**Additional considerations**

- For patients without risk factors, full replacement can be estimated with an ideal body weight–based dose of approximately 1.6 μg/kg/d
- For elderly or fragile patients with multiple comorbidities, consider titrating up from low dose, starting at 25-50 μg
- Extreme elevations of TSH can be seen in the recovery phase of thyroiditis and can be watched in asymptomatic patients to determine whether there is recovery to normal within 3-4 weeks
- Under guidance of endocrinology, consider tapering hormone replacement and retesting in patients with a history of thyroiditis (initial thyrotoxic phase) Adrenal dysfunction, if present, must always be replaced before thyroid hormone therapy is initiated

##### 4.1.2 Hyperthyroidism

**Definition:** Suppressed TSH and high normal or elevated FT4 and/or triiodothyronine

**Diagnostic work-up**

- Monitor TSH, FT4 every 4-6 weeks from the start of therapy or as needed for case detection in symptomatic patients
- Consider TSH receptor antibodies if there are clinical features and suspicion of Grave disease (eg, ophthalmopathy)
- Close monitoring of thyroid function every 2-3 weeks after diagnosis to catch transition to hypothyroidism in patients with thyroiditis and hyperthyroidism

**Grading Management**

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1: Asymptomatic or mild symptoms</td>
<td>Can continue ICPI with close follow-up and monitoring of TSH, FT4 every 2-3 weeks until it is clear whether there will be persistent hyperthyroidism (see below) or hypothyroidism (see 4.1.1)</td>
</tr>
<tr>
<td>G2: Moderate symptoms, able to perform ADL</td>
<td>Consider holding ICPI until symptoms return to baseline</td>
</tr>
<tr>
<td></td>
<td>β-Blocker (eg, atenolol, propranolol) for symptomatic relief</td>
</tr>
<tr>
<td></td>
<td>Hydration and supportive care</td>
</tr>
<tr>
<td></td>
<td>Corticosteroids are not usually required to shorten duration</td>
</tr>
<tr>
<td></td>
<td>For persistent hyperthyroidism (&gt; 6 weeks) or clinical suspicion, work-up for Graves disease (TSI or TRAb) and consider thionamide (methimazole or PTU)</td>
</tr>
<tr>
<td></td>
<td>Refer to endocrinology for Graves disease</td>
</tr>
</tbody>
</table>

(continued on following page)
### Table 4. Management of Endocrine irAEs in Patients Treated With ICPis (continued)

#### 4.0 Endocrine Toxicity

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>G3-4: Severe symptoms, medically significant or life-threatening consequences, unable to perform ADL</td>
<td>Hold ICPi until symptoms resolve to baseline with appropriate therapy&lt;br&gt;Endocrine consultation&lt;br&gt;β-Blocker (eg, atenolol, propranolol) for symptomatic relief&lt;br&gt;For severe symptoms or concern for thyroid storm, hospitalize patient and initiate prednisone 1-2 mg/kg/d or equivalent tapered over 1-2 weeks; consider also use of SSKI or thionamide (methimazole or PTU).</td>
</tr>
</tbody>
</table>

**Additional considerations**
- Thyroiditis is transient and resolves in a couple of weeks to primary hypothyroidism or normal. Hypothyroidism can be treated as above.
- Graves disease is generally persistent and is due to increased thyroid hormone production that can be treated with antithyroid medical therapy.
- Physical examination findings of ophthalmopathy or thyroid bruit are diagnostic of Graves and should prompt early endocrine referral.

#### 4.2 Adrenal – primary adrenal insufficiency

**Definition:** Adrenal gland failure leading to low morning cortisol, high morning ACTH, as well as hyponatremia and hyperkalemia with orthostasis and volume depletion due to loss of aldosterone.

**Diagnostic work-up for patients in whom adrenal insufficiency is suspected:**
- Evaluate ACTH (am), cortisol level (am), and midnight cortisol
- Basic metabolic panel (Na, K, CO2, glucose)
- Consider ACTH stimulation test for indeterminate results

If primary adrenal insufficiency (high ACTH, low cortisol) is found biochemically:
- Evaluate for precipitating cause of crisis such as infection
- Perform an adrenal CT for metastasis/hemorrhage
- Consider evaluating LH, FSH, and testosterone levels in males or estrogen in premenopausal females with fatigue, loss of libido, and mood changes
- Consider MRI of the brain with or without contrast with pituitary/sellar cuts in patients with multiple endocrine abnormalities
- Evaluate ACTH, cortisol (am), TSH, FT4, electrolytes

**Emergent therapy for someone with suspected adrenal insufficiency:**
- Initiate outpatient treatment at two to three times maintenance dose of prednisone (10-20 mg orally every morning, 5-10 mg orally every evening) in the morning, and 10-20 mg in the afternoon) to manage acute symptoms.
- Taper stress-dose corticosteroids down to maintenance doses over 5-10 days
- Maintenance therapy as in G1.

**Additional considerations**
- Primary and secondary adrenal insufficiency can be distinguished by the relationship between ACTH and cortisol. If the ACTH is low with low cortisol, then management is as per 4.3.
- Patients on corticosteroids for management of other conditions will have low morning cortisol as a result of iatrogenic, secondary adrenal insufficiency. ACTH will also be low in these patients. A diagnosis of adrenal insufficiency is challenging to make in these situations (see next section on hypophysitis).
- Emergent therapy for someone with suspected adrenal insufficiency is best done with dexamethasone as a stimulation test can still be performed. If the diagnosis is not clear and stimulation testing will be needed)
- May require hydrocortisone (0.1 mg/d) for mineralocorticoid replacement in primary adrenal insufficiency
- Titrating dose up or down as symptoms dictate

**Table 4 (continued on following page)**

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**4.3 Pituitary - hypophysitis**

**Definition:** Inflammation of the pituitary with varying effects on hormone function. Most commonly presenting with central adrenal insufficiency. May also have central hypothyroidism, diabetes insipidus, and hypogonadism.

**Diagnostic work-up**
- Diagnosis: Low ACTH with a low cortisol. Low or normal TSH with a low FT4. Hyponatremia and volume depletion with diabetes insipidus. Low testosterone or estradiol with low LH and FSH.
- Testing:
  - Evaluate ACTH, cortisol (am), TSH, FT4, electrolytes
  - Consider evaluating LH, FSH, and testosterone levels in males or estrogen in premenopausal females with fatigue, loss of libido, and mood changes
  - Consider MRI of the brain with or without contrast with pituitary/ellar cuts in patients with multiple endocrine abnormalities ± new severe headaches or complaints of vision changes

**(continued from previous page)**
Management of Immune-Related Adverse Events

Table 4. Management of Endocrine irAEs in Patients Treated With ICPIs (continued)

4.0 Endocrine Toxicity

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1: Asymptomatic or mild symptoms</td>
<td>Considering holding ICPI until patient is stabilized on replacement hormones</td>
</tr>
<tr>
<td></td>
<td>Hormonal supplementation as needed, using dosing as above for primary hypothyroidism and adrenal insufficiency (eg, hydrocortisone 10-20 mg orally in the morning, 5-10 mg orally in early afternoon; levothyroxine by weight)</td>
</tr>
<tr>
<td></td>
<td>Testosterone or estrogen therapy as needed in those without contraindications</td>
</tr>
<tr>
<td></td>
<td>Endocrine consultation</td>
</tr>
<tr>
<td></td>
<td>Always start corticosteroids several days before thyroid hormone to prevent precipitating adrenal crisis</td>
</tr>
<tr>
<td></td>
<td>Follow FT4 for thyroid hormone replacement titration (TSH is not accurate)</td>
</tr>
<tr>
<td>G2: Moderate symptoms, able to perform ADL</td>
<td>Consider holding ICPI until patient is stabilized on replacement hormones</td>
</tr>
<tr>
<td></td>
<td>Endocrine consultation</td>
</tr>
<tr>
<td></td>
<td>Hormonal supplementation as in G1</td>
</tr>
<tr>
<td>G3-4: Severe symptoms, medically significant or life-threatening consequences, unable to perform ADL</td>
<td>Hold ICPI until patient is stabilized on replacement hormones</td>
</tr>
<tr>
<td></td>
<td>Endocrine consultation</td>
</tr>
<tr>
<td></td>
<td>Hormonal supplementation as in G1</td>
</tr>
<tr>
<td></td>
<td>Consider initial pulse dose therapy with prednisone 1-2 mg/kg oral daily (or equivalent) tapered over at least 1-2 weeks</td>
</tr>
</tbody>
</table>

Additional considerations

- Be aware of the need to start corticosteroids first when planning hormone replacement therapy for multiple deficiencies
- All patients need instruction on doubling doses for illness (stress dosing) and a medical alert bracelet for adrenal insufficiency to trigger stress-dose corticosteroids by EMS
- Corticosteroid use can cause isolated central adrenal insufficiency
- Work-up cannot be done with a simple AM cortisol in a patient on corticosteroids for other conditions
- Laboratory confirmation of adrenal insufficiency should not be attempted until treatment with corticosteroids for other disease is ready to be discontinued

4.4 Diabetes

Definition: T2DM is a combination of insulin resistance and insulin deficiency that may require oral or insulin therapy. It may be new onset or exacerbated during therapy for nonimmunologic reasons, such as corticosteroid exposure.

Autoimmune T1DM results from islet cell destruction and is often acute onset, with ketosis and an insulin requirement

Diagnostic work-up

- Monitor patients for hyperglycemia or other signs and symptoms of new or worsening DM, including measuring glucose at baseline and with each treatment cycle during induction for 12 weeks, then every 3-6 weeks thereafter. To guide the work-up in new-onset hyperglycemia, clinicians should consider a patient’s medical background, exposure history, and risk factors for each subtype of DM.
- Laboratory evaluation in suspected T1DM should include testing for ketosis in urine and an assessment of the anion gap on a metabolic panel. Anti–glutamic acid decarboxylase, anti–islet cell, or anti–insulin antibodies are highly specific for autoimmune diabetes. Insulin and C-peptide levels can also assist in the diagnosis.

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1: Asymptomatic or mild symptoms; fasting glucose value &gt; ULN (180 mg/dL); fasting glucose value &gt; ULN (8.9 mmol/L); no evidence of ketosis or laboratory evidence of T1DM</td>
<td>Can continue ICPI with close clinical follow-up and laboratory evaluation</td>
</tr>
<tr>
<td></td>
<td>May initiate oral therapy for those with new-onset T2DM</td>
</tr>
<tr>
<td></td>
<td>Screen for T1DM if appropriate, for example, acute onset with prior normal values or clinical concern for ketosis</td>
</tr>
<tr>
<td>G2: Moderate symptoms, able to perform ADL; fasting glucose value &gt; 160-250 mg/dL; fasting glucose value &gt; 8.9-13.9 mmol/L; ketosis or evidence of T1DM at any glucose level</td>
<td>May hold ICPI until glucose control is obtained</td>
</tr>
<tr>
<td></td>
<td>Titrate oral therapy or add insulin for worsening control in T2DM</td>
</tr>
<tr>
<td></td>
<td>Should administer insulin for T1DM (or as default therapy if there is confusion about type)</td>
</tr>
<tr>
<td></td>
<td>Urgent endocrine consultation for any patient with T1DM; in the absence of endocrinology, internal medicine may suffice</td>
</tr>
<tr>
<td>G3-4: Severe symptoms, medically significant or life-threatening consequences, unable to perform ADL</td>
<td>Hold ICPI until glucose control is obtained on therapy with reduction of toxicity to G1 or less</td>
</tr>
<tr>
<td>G3: &gt; 250-500 mg/dL (&gt; 13.9-27.8 mmol/L)</td>
<td>Urgent endocrine consultation for all patients</td>
</tr>
<tr>
<td>G4: &gt; 500 mg/dL (&gt; 27.8 mmol/L)</td>
<td>Initiate insulin therapy for all patients</td>
</tr>
<tr>
<td></td>
<td>Admit for inpatient management: Concerns for developing DKA, Symptomatic patients regardless of diabetes type, New-onset T1DM unable to see endocrinology</td>
</tr>
</tbody>
</table>

Additional considerations

- Insulin therapy can be used as the default in any case with hyperglycemia.
- Long-acting therapy alone is not usually sufficient for T1DM, where half of daily requirements are usually given in divided doses as prandial coverage and half as long acting.
- Insulin doses will be lower in T1DM because of preserved sensitivity (total daily requirement can be estimated at 0.3-0.4 units/kg/d).
- In T2DM, sliding-scale coverage with meals over a few days provides data to estimate a patient’s daily requirements and can be used to more rapidly titrate basal needs.

All recommendations are expert consensus based, with benefits outweighing harms, and strength of recommendations are moderate.

Abbreviations: ACTH, adrenocorticotropic hormone; ADL, activities of daily living; CT, computed tomography; DKA, diabetic ketoacidosis; DM, diabetes mellitus; EMS, emergency medical services; FSH, follicle-stimulating hormone; FT4, free thyroxine; G, grade; ICPI, immune checkpoint inhibitor; irAE, immune-related adverse event; LH, luteinizing hormone; MRI, magnetic resonance imaging; PTU, propylthiouracil; SSKI, potassium iodide; T1DM, type 1 diabetes mellitus; T2DM, type 2 diabetes mellitus; TRAb, thyroid-stimulating hormone receptor antibody; TSH, thyroid-stimulating hormone; TSI, thyroid-stimulating immunoglobulin; ULN, upper limit of normal.
4.1.2 Hyperthyroidism

**Recommendation 4.1.2a – Diagnostic work-up.** It is recommended that the diagnostic work-up should include the following:

- Monitor TSH and FT4 every 4 to 6 weeks from the start of therapy or as needed for case detection in symptomatic patients.
- Test for TSH receptor antibodies if there are clinical features and suspicion of Grave disease (eg, ophthalmopathy).
- Should closely monitor thyroid function every 2 to 3 weeks after diagnosis to catch transition to hypothyroidism in patients with thyroiditis and hyperthyroidism.

**Recommendation 4.1.2b – Management.** It is recommended that clinicians manage grade 1 toxicities as follows:

- Should continue to offer ICPi with close follow-up and monitoring of TSH and FT4 every 2 to 3 weeks until it is clear whether there will be persistent hyperthyroidism (see below) or hypothyroidism (see 4.1.1).

It is recommended that clinicians manage grade 2 toxicities as follows:

- May hold ICPi until symptoms resolve to baseline.
- May consult endocrinology.
- Should offer a β-blocker (eg, atenolol, propranolol) for symptomatic relief.
- Should offer hydration and supportive care.
- Should note that corticosteroids are not usually required to shorten duration.
- For persistent hyperthyroidism (> 6 weeks) or clinical suspicion, clinicians should work up for Graves disease (thyroid-stimulating immunoglobulin or TSH receptor antibody) and consider thionamide (methimazole or propylthiouracil).
- Should refer to endocrinology for Graves disease.

It is recommended that clinicians manage grade 3 to 4 toxicities as follows:

- Should hold ICPi until symptoms resolve to baseline with appropriate therapy.
- Should consult endocrinology.
- Should offer a β-blocker (eg, atenolol, propranolol) for symptomatic relief.
- For severe symptoms or concern for thyroid storm, should hospitalize patient and initiate prednisone 1 to 2 mg/kg/d or equivalent tapered over 1 to 2 weeks. May also use saturated solution of potassium iodide or thionamide (methimazole or propylthiouracil).

4.2 Adrenal – Primary Adrenal Insufficiency

**Recommendation 4.2a – Diagnostic work-up.** It is recommended that the diagnostic work-up should include the following for patients in whom adrenal insufficiency is suspected:

- Evaluate adrenocorticotropic hormone (ACTH; AM), cortisol level (AM).
- Basic metabolic panel (Na, K, CO₂, glucose).
- Consider ACTH stimulation test for indeterminate results.

- For evidence of primary adrenal insufficiency (high ACTH, low cortisol), evaluate for a precipitating cause of crisis, such as infection, and perform an adrenal CT scan for metastasis/hemorrhage.

**Recommendation 4.2b – Management.** It is recommended that clinicians manage grade 1 toxicities as follows:

- May hold ICPi until patient is stabilized on replacement hormone.
- Should consult endocrinology.
- Should offer replacement therapy with prednisone (5 to 10 mg daily) or hydrocortisone (10 to 20 mg orally in the morning, 5 to 10 mg orally in early afternoon)
- May prescribe fluordrocortisone (0.1 mg/d) for mineralocorticoid replacement in primary adrenal insufficiency.
- Should titrate dose up or down as symptoms dictate.

It is recommended that clinicians manage grade 2 toxicities as follows:

- May hold ICPi until patient is stabilized on replacement hormone.
- Should consult endocrinology.
- Should initiate outpatient treatment at two to three times maintenance (eg, if prednisone, 20 mg daily; if hydrocortisone, 20 to 30 mg in the morning and 10 to 20 mg in the afternoon) to manage acute symptoms.
- Should taper stress-dose corticosteroids down to maintenance doses over 5 to 10 days.
- Should offer maintenance therapy as in grade 1.

It is recommended that clinicians manage grade 3 to 4 toxicities as follows:

- Should hold ICPi until patient is stabilized on replacement hormone.
- Should consult endocrinology.
- Should see in clinic or, for after hours, make an emergency department referral for normal saline (at least 2 L) and IV stress-dose corticosteroids on presentation (hydrocortisone 100 mg or dexamethasone 4 mg if the diagnosis is not clear and stimulation testing will be needed).
- Should taper stress-dose corticosteroids down to maintenance doses over 7 to 14 days after discharge.
- Should offer maintenance therapy as in grade 1.

**Qualifying statement.** Primary and secondary adrenal insufficiency can be distinguished by the relationship between ACTH and cortisol. If the ACTH is low with low cortisol, then management is as per 4.3. Emergent therapy for someone with suspected adrenal insufficiency is best done with dexamethasone, as a stimulation test can still be performed. If the diagnosis is already confirmed, can use hydrocortisone 100 mg.

4.3 Pituitary – Hypophysitis

**Recommendation 4.3a – Diagnostic work-up.** It is recommended that the diagnostic work-up should include the following:

- Evaluate ACTH, cortisol (AM), TSH, FT4, and electrolytes.
- Consider evaluating luteinizing hormone, follicle-stimulating hormone, and testosterone levels in males or estrogen in premenopausal females with fatigue, loss of libido, and mood changes.
• Consider magnetic resonance imaging (MRI) of brain with or without contrast with pituitary/sellar cuts in patients with multiple endocrine abnormalities with or without new severe headaches or complaint of vision changes.

Recommendation 4.3b—Management. It is recommended that clinicians manage grade 1 toxicities as follows:

- May hold ICPI until patient is stabilized on replacement hormones.
- Should offer hormonal supplementation as needed, using dosing as specified for primary hypothyroidism and adrenal insufficiency (eg, hydrocortisone 10 to 20 mg orally in the morning, 5 to 10 mg orally in early afternoon; levothyroxine by weight).
- Testosterone or estrogen therapy as needed in those without contraindications.
- Should consult endocrinology.
- Should always start corticosteroids several days before thyroid hormone to prevent precipitating adrenal crisis.
- Should follow FT4 for thyroid hormone replacement titration (TSH is not accurate). It is recommended that clinicians manage grade 2 toxicities as follows:
  - May hold ICPI until patient is stabilized on replacement hormones.
  - Should consult endocrinology.
  - Should offer hormonal supplementation as in grade 1.

It is recommended that clinicians manage grade 3 to 4 toxicities as follows:

- Should hold ICPI until patient is stabilized on replacement hormones.
- Should consult endocrinology.
- Should offer hormonal supplementation as in grade 1.
- May administer initial pulse dose therapy with prednisone 1 to 2 mg/kg oral daily (or equivalent) tapered over at least 1 to 2 weeks.

Qualifying statement. Be aware of the need to start corticosteroids first when planning hormone replacement therapy for multiple deficiencies. All patients need instruction on doubling doses for illness (stress dosing) and a medical alert bracelet for adrenal insufficiency to trigger stress-dose corticosteroids by emergency medical services.

4.4 Diabetes

Recommendation 4.4a—Diagnostic work-up. It is recommended that the diagnostic work-up should include the following:

- Monitor patients for hyperglycemia or other signs and symptoms of new or worsening diabetes mellitus (DM), including measuring glucose at baseline and with each treatment cycle during induction for 12 weeks, then every 3 to 6 weeks thereafter.
- To guide the work-up in new-onset hyperglycemia, clinicians should consider a patient’s medical background, exposure history, and risk factors for each subtype of DM.
- Laboratory evaluation in suspected type 1 DM (T1DM) should include testing for ketosis in urine and an assessment of the anion gap on a metabolic panel. Anti-glutamic acid decarboxylase, anti–islet cell, or anti–insulin antibodies are highly specific for autoimmune diabetes. Insulin and C-peptide levels can also assist in the diagnosis.

Recommendation 4.4b—Management. It is recommended that clinicians manage grade 1 toxicities as follows:

- May continue to offer ICPI with close clinical follow-up and laboratory evaluation.
- May initiate oral therapy for those with new-onset type 2 DM (T2DM).
- Should screen for T1DM if appropriate, for example, acute onset with prior normal values or clinical concern for ketosis.

It is recommended that clinicians manage grade 2 toxicities as follows:

- May hold ICPI until glucose control is obtained.
- Should titrate oral therapy or add insulin for worsening control in T2DM.
- Should administer insulin for T1DM (or as default therapy if there is confusion about type).
- Should seek urgent endocrine consultation for any patient with T1DM. In the absence of endocrinology, internal medicine may suffice.
- May admit for T1DM if early outpatient evaluation is not available or signs of ketoacidosis are present.

It is recommended that clinicians manage grade 3 to 4 toxicities as follows:

- Should hold ICPI until glucose control is obtained on therapy with reduction of toxicity to grade 1 or less.
- Should seek urgent endocrine consultation for all patients.
- Should initiate insulin therapy for all patients.
- Should admit for inpatient management for any of the following: concern for developing diabetic ketoacidosis (DKA), symptomatic patients regardless of diabetes type, new onset T1DM unable to see endocrinology.

Discussion. Endocrine adverse events with immune checkpoint therapy present a unique clinical challenge for the non-endocrinologist who faces the possibility of central as well as primary endocrine dysfunction in a patient with symptoms or abnormal laboratories. Diverse therapies and combinations have varied rates of targeting individual organs, for example, hypophysitis is most commonly seen when ipilimumab is used, and primary ovarian failure has not yet been reported. However, with sporadic autoimmune disease known for all endocrine organs, we anticipate the possibility of any condition as the use becomes more widespread. In a recent systematic review and meta-analysis that included 7,551 patients in 38 randomized trials, the overall incidence of clinically significant endocrinopathies was approximately 10% of patients treated with checkpoint inhibitors. Clinical measure of both the primary hormone and the pituitary hormone are needed to localize disease. For example, a low morning cortisol suggests adrenal insufficiency but not whether the problem is pituitary or adrenal. We would look for hypophysitis if a simultaneously measured ACTH is low, whereas in primary adrenal insufficiency (eg, Addison), the ACTH will be elevated. The same applies for systems where we typically screen with the pituitary hormone—low TSH suggests hyperthyroidism if the thyroid hormone level (FT4 is typically sufficient) is elevated and central hypothyroidism if FT4 is low. Drawing both hormones is
especially important when hypophysitis is suspected because TSH can be at low-normal levels but lack function with pituitary disease.

Distinguishing primary from secondary hormonal problems is necessary because there are treatment implications. Perhaps most importantly for preventing harm is recognizing that hypophysitis often causes both central hypothryoidism and secondary adrenal insufficiency. If thyroid hormone is replaced first when cortisol is low, the increase in cortisol metabolism can trigger an adrenal crisis. Fludrocortisone is needed in addition to hydrocortisone in most cases of primary adrenal insufficiency, which involves the loss of mineralocorticoid as well as glucocorticoid, leading to more-profound blood pressure and electrolyte abnormalities. Monitoring is also affected by localization, as pituitary hormones are not reliable indicators of status with central disease. TSH, therefore, is not helpful in monitoring therapy with levothyroxine in central hypothryoidism, and FT4 should be used instead.

Diagnosis of endocrine dysfunction is complicated by any acute illness and the administration of medications that have an effect on pituitary function, including many therapies that patients with cancer are on, such as narcotics and megestrol acetate. Most relevant for the patients administered ICPIs is the effect of corticosteroids, given for many irAEs, which will directly suppress ACTH and may cause persistent central adrenal insufficiency when stopped. Cortisol levels should not be routinely measured while patients are on corticosteroid therapy because of variable assay effects from synthetic corticosteroids, low endogenous levels from the exposure, and the fact that the patient is on supraphysiologic doses and therefore treated for any underlying adrenal insufficiency that might have developed. If a diagnosis is needed, for example, after acute treatment of presumed adrenal crisis is initiated, ACTH stimulation testing may be performed on dexamethasone, which is not measured by most assays. Endogenous levels can be directly measured 24 hours after the last dose of physiologic hydrocortisone replacement to assess for functional recovery. High-dose corticosteroids can also cause a low TSH and a pattern similar to nontyroidal illness, neither of which are thought to benefit from therapy. Especially in difficult cases, endocrinology consult is recommended.

The response of the oncologist to the development of endocrine dysfunction may be different from other irAEs because organ failure can be managed with hormone replacement. It is not a given that the patient benefits from stopping cancer therapy to get immunosuppressive therapy to reverse the autoimmune disease. For example, there is no good evidence at this time that high-dose corticosteroids improve the rate of pituitary hormone recovery. Therefore, a clinical judgment is needed to balance benefits, such as the possibility of improved headache, with risks, such as corticosteroid adverse effects, on glycemic control and delay of therapy.

Rare cases of T1DM present an analogous challenge to the clinician in the need to distinguish these from the much more common cases of worsening glycemic control attributable to insulin resistance and T2DM. The acute risks of DKA in T1DM require vigilance on the part of treating oncologists, despite the very low occurrence rate. New-onset hyperglycemia in a patient without risk factors for T2DM (eg, preexisting disease, corticosteroid exposure) should raise the level of concern for T1DM. Acute onset of polyuria, polydipsia, weight loss, and lethargy are characteristic presenting features of T1DM. Urine ketones and acid base status can be evaluated as screening for DKA and the need for inpatient evaluation. Antibodies, insulin, and C-peptide levels can be sent to support diagnosis, although the initiation of therapy should not be delayed pending results. Insulin should be used to treat hyperglycemia in anyone where the diagnosis is in question. Endocrinology consultation is appropriate where the diagnosis of T1DM is suspected even without evidence of DKA.

5.0 Musculoskeletal Toxicities

Please refer to Table 5 for a complete set of recommendations, definition of grades, and additional considerations.

5.1 Inflammatory Arthritis

Recommendation 5.1a – Diagnostic work-up. It is recommended that the diagnostic work-up should include the following for grade 1:

- Complete rheumatologic history and examination of all peripheral joints for tenderness, swelling, and range of motion.
- Examination of the spine.
- Consider plain x-ray/ imaging to exclude metastases and evaluate joint damage (erosions), if appropriate.
- Consider autoimmune blood panel, including ANA, rheumatoid factor (RF), and anti–citrullinated protein antibody (anti-CCP), and inflammatory markers (ESR and CRP) if symptoms persist. If symptoms are suggestive of reactive arthritis or affect the spine, consider HLA B27 testing.

It is recommended that the diagnostic work-up should include the following for grade 2:

- Complete history and examination as above; laboratory tests as above.
- Consider ultrasound with or without MRI of affected joints if clinically indicated (eg, persistent arthritis unresponsive to treatment, suspicion for differential diagnoses such as metastatic lesions or septic arthritis).
- Consider early referral to a rheumatologist if there is joint swelling (synovitis) or if symptoms of arthralgia persist > 4 weeks.

It is recommended that the diagnostic work-up should include the following for grades 3 to 4:

- As for grade 2.
- Seek rheumatologist advice and review.

It is recommended that all patients with inflammatory arthritis be monitored with serial rheumatologic examinations, including inflammatory markers, every 4 to 6 weeks after treatment is instituted.

Recommendation 5.1b – Management. It is recommended that clinicians should follow reports of new joint pain to determine whether inflammatory arthritis is present. Clinicians should question whether symptoms are new since receiving ICPI.

It is recommended that clinicians manage grade 1 toxicities as follows:

- Should continue to offer ICPI.
- Should initiate analgesia with acetaminophen and/or NSAIDs.
5.1 Inflammatory arthritis

Definition: A disorder characterized by inflammation of the joints.

Clinical symptoms: Joint pain accompanied by joint swelling; inflammatory symptoms, such as stiffness after inactivity or in the morning, lasting > 30 minutes to 1 hour; improvement of symptoms with NSAIDs or corticosteroids but not with opioids or other pain medications may also be suggestive of inflammatory arthritis.

Diagnostic work-up

G1: Complete rheumatologic history and examination of all peripheral joints for tenderness, swelling, and range of motion; examination of the spine.

Consider plain x-ray/imaging to exclude metastases and evaluate joint damage (erosions), if appropriate.

Consider autoimmune blood panel including ANA, RF, and anti-CCP, and anti-inflammatory markers (ESR and CRP) if symptoms persist; if symptoms are suggestive of reactive arthritis or affect the spine, consider HLA B27 testing.

G2: Complete history and examination as above; laboratory tests as above.

Consider US ± MRI of affected joints if clinically indicated (eg, persistent arthritis unresponsive to treatment, suspicion for differential diagnoses such as metastatic lesions or septic arthritis).

Consider early referral to a rheumatologist, if there is joint swelling (synovitis) or if symptoms of arthralgia persist > 4 weeks.

G3-4: As for G2.

Consider paraneoplastic autoantibody testing for myositis and neurologic conditions, such as myasthenia gravis.

Referral to rheumatology, if suspicion for reactive arthritis or erosive joint damage.

Monitoring: Patients with inflammatory arthritis should be monitored with serial rheumatologic examinations, including inflammatory markers, every 4-6 weeks after treatment is instituted.

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1: Mild pain with inflammation, erythema, or joint swelling</td>
<td>Continue ICPi</td>
</tr>
<tr>
<td>G2: Moderate pain associated with signs of inflammation, erythema, or joint swelling, limiting instrumental ADL</td>
<td>Hold ICPi and resume upon symptom control and on prednisone ≤ 10 mg/d. Escalate analgesia and consider higher doses of NSAIDs as needed. If inadequately controlled, initiate prednisone or prednisolone 10-20 mg/d or equivalent for 4-6 weeks. If improvement, slow taper according to response during the next 4-6 weeks; if no improvement after initial 4-6 weeks, treat as G3. If unable to lower corticosteroid dose to &lt; 10 mg/d after 3 months, consider DMARD. Consider intra-articular corticosteroid injections for large joints.</td>
</tr>
<tr>
<td>G3-4: Severe pain associated with signs of inflammation, erythema, or joint swelling; irreversible joint damage; disabling; limiting self-care ADL</td>
<td>Hold ICPi temporarily and may resume in consultation with rheumatology, if recover to G1 or less. Initiate oral prednisone 0.5-1 mg/kg/day. If failure of improvement after 4 weeks or worsening in meantime, consider synthetic or biologic DMARD. Synthetic: methotrexate, leflunomide. Biologic: consider anticytokine therapy such as TNF-α or IL-6 receptor inhibitors. (Note: As caution, IL-6 inhibition can cause intestinal perforation; while this is extremely rare, it should not be used in patients with colitis.) Test for viral hepatitis B, C, and latent/active TB test prior to DMARD treatment.</td>
</tr>
</tbody>
</table>

**Additional considerations**

Early recognition is critical to avoid erosive joint damage.

Corticosteroids can be used as part of initial therapy in inflammatory arthritis, but due to likely prolonged treatment requirements, physicians should consider starting corticosteroid-sparing agents earlier than one would with other irAEs.

Oligoarthritis can be treated early on with intra-articular corticosteroids; consider early referral.

Consider PCP prophylaxis for patients treated with high dose of corticosteroids for > 12 weeks, as per local guidelines.

5.2 Myositis

Definition: A disorder characterized by muscle inflammation with weakness and elevated muscle enzymes (CK). Muscle pain can be present in severe cases. Can be life threatening if respiratory muscles or myocardium are involved.

**Diagnostic work-up**

Complete rheumatologic and neurologic history regarding differential diagnosis; rheumatologic and neurologic examination, including muscle strength; and examination of the skin for findings suggestive of dermatomyositis. Muscle weakness is more typical of myositis than pain. Consider preexisting conditions that can cause similar symptoms.

Blood testing to evaluate muscle inflammation.

CK, transaminases (AST, ALT), LDH, and aldolase can also be elevated.

Troponin to evaluate myocardial involvement and other cardiac testing, such as echocardiogram, as needed.

Inflammatory markers (ESR and CRP).

Consider EMG, imaging (MRI), and/or biopsy on an individual basis when diagnosis is uncertain and overlap with neurologic syndromes, such as myasthenia gravis, is suspected.

Consider paraneoplastic autoantibody testing for myositis and neurologic conditions, such as myasthenia gravis.

Monitoring: CK, ESR, CRP.

(continued on following page)
### 5.0 Musculoskeletal Toxicities

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1: Mild weakness with or without pain</td>
<td>Continue ICPi if CK is elevated and patient has muscle weakness, may offer oral corticosteroids, and treat as G2. Offer analgesia with acetaminophen or NSAIDs if there are no contraindications.</td>
</tr>
<tr>
<td>G2: Moderate weakness with or without pain, limiting age-appropriate instrumental ADL</td>
<td>Hold ICPi temporarily and may resume upon symptom control, if CK is normal and prednisone dose &lt; 10 mg; if worsens, treat as per G3 NSAIDs as needed.</td>
</tr>
<tr>
<td>G3-4: Severe weakness with or without pain, limiting self-care ADL</td>
<td>Hold ICPi until G1 or less while off immune suppression and permanently discontinue if any evidence of myocardial involvement.</td>
</tr>
</tbody>
</table>

**Table 5. Management of Musculoskeletal irAEs in Patients Treated With ICPis (continued)**

**5.3 Polymyalgia-like syndrome**

**Definition:** Characterized by marked pain and stiffness in proximal upper and/or lower extremities and no signs of true muscle inflammation such as CK elevation or EMG findings of myositis. No true muscle weakness, difficulty in active motion related to pain.

**Diagnostic work-up**

**G1**
- Complete rheumatologic history regarding differential diagnosis and examination of all joints and skin.
- Check for symptoms of temporal arteritis, such as headache or visual disturbances; refer to ophthalmologist if present, and consider temporal artery biopsy.
- CK to evaluate differential diagnosis of myositis.
- Inflammatory markers (ESR, CRP).
- Monitoring: ESR, CRP.

**G2**
- Complete history and examination as above; autoimmune myositis blood panel; EMG, MRI of affected joints.

**G3-4**
- As for G2; consider holding ICPi and resuming upon symptom control, prednisolone 10 mg; if worsens, treat as per G3.

**Additional considerations:** Caution is advised with rechallenging.

**Hold ICPi until G1 or less while off immune suppression and permanently discontinue if any evidence of myocardial involvement.**

**Consideration:** Referral to rheumatologist or neurologist.

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**Abbreviations:** ADL, activities of daily living; ANA, antinuclear antibodies; CCP, citrullinated protein antibody; CK, creatine kinase; CRP, C-reactive protein; DMARD, disease-modifying antirheumatic drug; EMG, electromyography; ESR, erythrocyte sedimentation rate; ICPi, immune checkpoint inhibitor; IL, interleukin; irAE, immune-related adverse event; IV, intravenous; IVIG, intravenous immunoglobulin; LDH, lactate dehydrogenase; NSAID, nonsteroidal anti-inflammatory drug; PCP, Pneumocystis pneumonia; RF, rheumatoid factor; TNF, tumor necrosis factor.
It is recommended that clinicians manage grade 2 toxicities as follows:
- Should hold ICPI and resume upon symptom control and on prednisone ≤ 10 mg/d.
- Should escalate analgesia and consider higher doses of NSAIDs as needed.
- If inadequately controlled, should initiate oral prednisone 10 to 20 mg/d or equivalent for 4 to 6 weeks.
- If improvement, slow taper according to response during the next 4 to 6 weeks. If no improvement after initial 4 to 6 weeks, treat as grade 3.
- If unable to lower prednisone dose to < 10 mg/d after 3 months, may offer DMARD.
- May offer intra-articular corticosteroid injections for large joints.
- Should refer to rheumatology.

It is recommended that clinicians manage grade 3 to 4 toxicities as follows:
- Should hold ICPI and may resume in consultation with rheumatology, if recover to grade 1 or less.
- Should initiate oral prednisone 0.5 to 1 mg/kg.
- If failure of improvement after 4 weeks or worsening in meantime, may offer synthetic or biologic DMARD:
  - Synthetic: methotrexate, leflunomide.
  - Biologic: consider anticytokine therapy, such as TNF-α or interleukin-6 (IL-6) receptor inhibitors. (Note: As caution, IL-6 inhibition can cause intestinal perforation. While this is extremely rare, it should not be used in patients with colitis).
- Should refer to rheumatology.

5.2 Myositis

Recommendation 5.2a – Diagnostic work-up. It is recommended that the diagnostic work-up should include the following:
- Complete rheumatologic and neurologic history regarding differential diagnosis; rheumatologic and neurologic examination, including muscle strength; and examination of the skin for findings suggestive of dermatomyositis. Muscle weakness is more typical of myositis than pain. Consider preexisting conditions that can cause similar symptoms.
- Blood testing to evaluate muscle inflammation.
- CK, transaminases (AST, ALT), lactate dehydrogenase (LDH), and aldolase can also be elevated.
- Troponin to evaluate myocardial involvement and other cardiac testing, such as echocardiogram, as needed.
- Inflammatory markers (ESR and CRP).
- Consider electromyography (EMG), imaging (MRI), and/or biopsy on an individual basis when diagnosis is uncertain and overlap with neurologic syndromes, such as myasthenia gravis, is suspected.
- Consider paraneoplastic autoantibody testing for myositis and neurologic conditions, such as myasthenia gravis.

It is recommended that the following should be included for monitoring:
- CK, ESR, CRP

It is recommended that the diagnostic work-up should include the following for grade 1:
- Complete examination and laboratory work-up as specified in Section 5.2a

It is recommended that the diagnostic work-up should include the following for grade 2:
- Complete history and examination as above; autoimmune myositis and neurologic panel; EMG, MRI of affected proximal limbs as needed. Consider muscle biopsy if diagnosis is uncertain.
- Early referral to a rheumatologist or neurologist.

It is recommended that the diagnostic work-up should include the following for grade 3:
- As for grade 2
- Urgent referral to a rheumatologist or neurologist

Recommendation 5.2b – Management. It is recommended that clinicians manage grade 1 toxicities as follows:
- Should continue to offer ICPI.
- If CK is elevated and patient has muscle weakness, may offer oral corticosteroids and treat as grade 2.
- Should offer analgesia as needed for pain with acetaminophen or NSAIDs if there are no contraindications.

It is recommended that clinicians manage grade 2 toxicities as follows:
- Should hold ICPI and may resume upon symptom control, if CK is normal, and prednisone dose < 10 mg; if worsens, treat as per grade 3. Permanently discontinue if there is evidence of myocardial involvement.
- Should offer NSAIDs as needed.
- Referral to rheumatologist or neurologist.
- If CK is elevated (three times or more), should initiate prednisone or equivalent at 0.5 to 1 mg/kg.
- May require permanent discontinuation of ICPI therapy in most patients with grade 2 symptoms and objective findings (elevated enzymes, abnormal EMG, abnormal muscle MRI or biopsy).

It is recommended that clinicians manage grade 3 to 4 toxicities as follows:
- Should hold ICPI until grade 1 or less while off immune suppression and permanently discontinue if any evidence of myocardial involvement.
- Consider hospitalization for severe weakness.
- Referral to rheumatologist or neurologist.
- Should initiate prednisone 1 mg/kg or equivalent. Consider 1 to 2 mg/kg of methylprednisolone IV or higher-dose bolus if severe compromise (weakness severely limiting mobility, cardiac, respiratory, dysphagia).
- May offer plasmapheresis.
- May offer IVIG therapy.
- May offer other immunosuppressant therapy, such as methotrexate, azathioprine, or mycophenolate mofetil, if symptoms and laboratory findings do not improve or worsen after 4 to 6 weeks. Rituximab is used in primary myositis, but caution is advised given its long biologic duration.
### 5.3 Polymyalgia-Like Syndrome

**Recommendation 5.3a – Diagnostic work-up.** It is recommended that the diagnostic work-up should include the following for grade 1:

- Complete rheumatologic history regarding differential diagnosis and examination of all joints and skin.
- Check for symptoms of temporal arteritis, such as headache or visual disturbances; refer to ophthalmologist. If temporal arteritis is suspected, consider temporal artery biopsy.
- ANA, RF, anti-CCP.
- CK to evaluate differential diagnosis of myositis.
- Inflammatory markers (ESR, CRP).

It is recommended that the following should be included for monitoring:

- ESR, CRP

It is recommended that the diagnostic work-up should include the following for grade 2:

- Complete history and examination as above
- Autoimmune tests as above and others as required for differential diagnosis
- Early referral to a rheumatologist

It is recommended that the diagnostic work-up should include the following for grades 3 to 4:

- **As for grade 2**
- Seek rheumatologist advice and review

**Recommendation 5.3b – Management.** It is recommended that clinicians manage grade 1 toxicities as follows:

- Should continue to offer ICPi.
- Should initiate analgesia with acetaminophen and/or NSAIDs if there are no contraindications.

It is recommended that clinicians manage grade 2 toxicities as follows:

- May hold ICPi and resume upon symptom control, prednisone $< 10$ mg; if worsens, treat as per grade 3.
- Should initiate prednisone $20$ mg/d or equivalent. If symptoms improve, start to taper dose after 3 to 4 weeks.
- If no improvement or need for higher dosages after 4 weeks, escalate to grade 3.
- Consider referral to rheumatologist.

It is recommended that clinicians manage grade 3 to 4 toxicities as follows:

- Should hold ICPi and may resume, in consultation with rheumatology, if recover to grade 1 or less. However, note that cases of toxicity returning upon rechallenge have been reported.
- Referral to rheumatologist.
- Should initiate prednisone $20$ mg/d or equivalent. If no improvement or need for higher dosages for prolonged time, consider a corticosteroid-sparing agent such as methotrexate or IL-6 inhibition with tocilizumab. (Note: As caution, IL-6 inhibition can cause intestinal perforation. While this is extremely rare, it should not be used in patients with colitis or GI metastases).
- Consider admission for pain control.

### Discussion.

Musculoskeletal symptoms such as arthralgia and myalgia are common in patients receiving ICPi therapy, as reported in up to 40% of those treated in clinical trials. More-severe inflammatory AEs are not as frequent but can have an important effect on patients’ quality of life because of their effect on function and daily activities. The most common musculoskeletal and rheumatic irAEs are arthritis, polymyalgia-like syndromes, and myositis. These events can occur with either CTLA-4 or anti–PD-1/PD-L1 antagonists, but seem to be more frequent with the latter class of drugs and when these agents are used in combination.

The clinical presentation of patients with immune-related arthritis secondary to ICPi can vary and affect large and/or small joints. Some patients present with oligoarthritides of large joints, such as knees, ankles, or wrists. These patients can also have other features commonly seen with reactive arthritis, such as conjunctivitis or urethritis, and occasionally complain of back pain or cervical pain suggestive of saccroilitis. Other patients present with symmetrical polyarthritis resembling rheumatoid arthritis and can have autoantibodies such as RF present in their sera. Many patients also develop sicca symptoms, with dry eyes and dry mouth; autoantibodies, such as anti-SSA and anti-SSB, have occasionally been found, but most patients tend to be seronegative. Of interest, arthritis can occur at any time during treatment. Some patients have developed arthritis for the first time many months after initiation of ICPi therapy. Most common differential diagnoses include other causes of joint pain, including degenerative joint disease or osteoarthritis and soft tissue rheumatic disorders, such as rotator cuff tendinitis, crystal arthropathies (gout and pseudogout), and septic arthritis. Diagnostic evaluation should include serum inflammatory markers (ESR, CRP), evaluation of autoantibodies (ANA, RF, and anti-CCP), and imaging as needed (x-rays, ultrasound, and/or MRI). Inflammatory markers are usually very elevated in patients with ICPi-induced arthritis and are useful to differentiate these events from other rheumatic syndromes. NSAIDs alone are usually not sufficient to control symptoms, and corticosteroids and synthetic or biologic DMARDs might be required.

Intra-articular corticosteroid injections are an option if only one or two joints are affected.

Patients receiving ICPis can develop severe myalgia in their proximal upper and lower extremities, with severe fatigue resembling polymyalgia rheumatica. These patients can also have arthralgia but typically do not have definite synovitis, although ultrasound or MRI might show a mild effusion in the shoulder joints. Patients experiencing a polymyalgia-like syndrome have pain but not true weakness. Differential diagnoses include inflammatory myositis, fibromyalgia, statin-induced myopathy, and other types of arthritis or soft tissue rheumatic syndromes. RF and anti-CCP are negative, and inflammatory markers are highly elevated. CK levels should generally be within normal limits, differentiating this condition from myositis. Imaging with MRI and EMG should not show any evidence of myopathy or muscle inflammation.

Myositis is a rare complication of ICPis but can be severe and fatal. It is more common with PD-1/PD-L1 inhibitors than with ipilimumab. It can present as reactivation of preexisting paraneoplastic polymyositis or dermatomyositis or as a de novo myositis. The main symptom of inflammatory myositis is
weakness, primarily in the proximal extremities, with difficulties in standing up, lifting arms, and moving around. In severe cases, patients can complain of myalgia as well. Patients with de novo myositis do not develop the typical rash seen with paraneoplastic dermatomyositis. Myositis can have a fulminant necrotizing course with rhabdomyolysis and can involve vital skeletal muscle, such as the myocardium, in which case it requires urgent treatment to avoid fatal complications.\textsuperscript{75,76} Laboratory tests include measurement of muscle enzymes, especially CK, which often is markedly elevated, and inflammatory markers; autoantibody panels for myositis can be considered, although there is no evidence that any specific autoantibodies have a role in ICPI-associated myositis. Other diagnostic tests that may be useful include EMG, which can show muscle fibrillations indicative of myopathy, and/or MRI, which shows increased intensity and edema in affected muscles. Finally, biopsy can be performed to confirm the diagnosis. Differential diagnoses include generalized fatigue, polymyalgia rheumatica, fibromyalgia, adverse events from concomitant therapies (eg, statins, corticosteroids), and muscle dystrophies. These other disorders (other than some muscle dystrophies or drug-induced myopathy) have normal CK. The cornerstone of initial treatment is high-dose corticosteroids that should be administered as a bolus in severe cases. Plasmapheresis should be considered in cases with poor response to corticosteroids or in life-threatening situations. The use of other immunosuppressants and IVIG may also be indicated, as they are used for treatment of polymyositis/dermatomyositis. However, their efficacy in ICPI-induced myositis is not clearly documented.

A number of other rheumatic disorders have been occasionally documented as case reports of patients receiving ICPIs.\textsuperscript{77} These include vasculitis and lupus-like syndromes, among others. Management and treatment principles are similar to those reported for other ICPI-induced rheumatic syndromes.

Patients with preexisting autoimmune rheumatic conditions may be at higher risk of toxicity as either irAEs or flares of their preexisting disease.\textsuperscript{78} Many of these patients, nevertheless, can continue ICPI therapy or be rechallenged after their AE is properly managed, so having preexisting autoimmune disease does not represent an absolute contraindication for treatment. Close monitoring and multidisciplinary management is required for these patients, as they frequently need concomitant treatment of their preexisting autoimmune disease once they develop an AE. Management principles are similar to those described for irAE.

### 6.0 Renal Toxicities

Please refer to Table 6 for a complete set of recommendations, definition of grades, and additional considerations.

#### 6.1 Nephritis

**Recommendation 6.1a – Diagnostic work-up.** It is recommended that the diagnostic work-up should include the following:

- For any suspected immune-mediated adverse reactions, should exclude other causes.
- Monitor patients for elevated serum creatinine prior to every dose.

**Qualifying Statement.** Routine urinalysis is not necessary other than to rule out urinary tract infections, etc. Nephrology may be considered. If no potential alternative cause of acute kidney injury (AKI) is identified, then one should forego biopsy and proceed directly with immunosuppressive therapy. The swift treatment of the autoimmune component is important.

**Recommendation 6.1b – Management.** It is recommended that clinicians manage grade 1 toxicities as follows:

- May hold treatment temporarily, pending consideration of potential alternative etiologies (recent IV contrast, medications, fluid status) and baseline renal function. (Note: A change that is still < 1.5 ULN could be meaningful).

It is recommended that clinicians manage grade 2 toxicities as follows:

- Should hold treatment temporarily.
- Should consult nephrology.
- Should evaluate for other causes (recent IV contrast, medications, fluid status, etc). If other etiologies are ruled out, should administer 0.5 to 1 mg/kg/d prednisone equivalents.
- If worsening or no improvement, should administer 1 to 2 mg/kg/d prednisone or equivalent and permanently discontinue ICPI.
- If improved to grade 1 or less, taper corticosteroids over 4 to 6 weeks.
- If no recurrence of chronic renal insufficiency, discuss resumption of ICPI with patient after taking into account the risks and benefits.

It is recommended that clinicians manage grade 3 to 4 toxicities as follows:

- Should permanently discontinue ICPI.
- Should consult nephrology.
- Should evaluate for other causes (recent IV contrast, medications, fluid status, etc).
- Should administer corticosteroids (initial dose of 1 to 2 mg/kg/d prednisone or equivalent).

#### 6.2 Symptomatic Nephritis

**Recommendation 6.2a – Management.** It is recommended that clinicians manage grade 1 toxicities as follows:

- If improved to baseline, should resume routine creatinine monitoring.

It is recommended that clinicians manage grade 2 toxicities as follows:

- If improved to grade 1, should taper corticosteroids over at least 3 weeks before resuming treatment with routine creatinine monitoring.
- If elevations persist > 7 days or worsen and no other cause found, should treat as grade 3.

It is recommended that clinicians manage grade 3 toxicities as follows:

- If improved to grade 1, should taper corticosteroids over at least 4 weeks.
- If elevations persist > 3 to 5 days or worsen, may offer additional immunosuppression (eg, mycophenolate).
It is recommended that clinicians manage grade 4 toxicities as follows:

- If improved to grade 1, should taper corticosteroids over at least 4 weeks.
- If elevations persist > 2 to 3 days or worsen, may offer additional immunosuppression (eg, mycophenolate).

**Discussion.** AKI is an uncommon complication of checkpoint inhibitor immunotherapy. The estimated incidence of any-grade AKI was 1% to 2% in patients treated with a single agent (ipilimumab, nivolumab, pembrolizumab) and 4.5% in those treated with the combination of nivolumab plus ipilimumab. The incidence of grade 3 or 4 AKI was < 1% with single agents and 1.6% with the combination of nivolumab plus ipilimumab. While initial studies had quoted a small incidence of AKI with ICPi use, emerging data suggest a higher incidence rate of AKI (range, 9.9% to 29%) with ICPi. The vast majority of this extra toxicity is stage I based on AKI network criteria and typically involves electrolyte disturbances rather than declines in renal function.

In a retrospective series of 13 patients who underwent kidney biopsy at seven centers, renal toxicity was diagnosed a median of 91 days after initiation of checkpoint inhibitor immunotherapy (range, 21 to 245 days). The median peak serum creatinine was 4.5 mg/dL. Pathology from the kidney biopsies revealed acute

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### Table 6. Management of Renal irAEs in Patients Treated With ICPis

<table>
<thead>
<tr>
<th>6.0 Renal Toxicities</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Nephritis and renal dysfunction: diagnosis and monitoring</strong></td>
</tr>
<tr>
<td>For any suspected immune-mediated adverse reactions, exclude other causes</td>
</tr>
<tr>
<td>Monitor patients for elevated serum creatinine prior to every dose</td>
</tr>
<tr>
<td>Routine urinalysis is not necessary, other than to rule out UTIs, etc; nephrology may consider further</td>
</tr>
<tr>
<td>If no potential alternative cause of AKI identified, then one should forego biopsy and proceed directly with immunosuppressive therapy</td>
</tr>
<tr>
<td>Swift treatment of autoimmune component important</td>
</tr>
</tbody>
</table>

**6.1 Nephritis**

**Definition:** Inflammation of the kidney affecting the structure

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1: Creatinine level increase of &gt; 0.3 mg/dL; creatinine 1.5-2.0 × over baseline</td>
<td>Consider temporarily holding ICPi, pending consideration of potential alternative etiologies (recent IV contrast, medications, fluid status) and baseline renal function. A change that is still &lt; 1.5 ULN could be meaningful</td>
</tr>
<tr>
<td>G2: Creatinine 2-3 × above baseline</td>
<td>Hold ICPi temporarily; consult nephrology; evaluate for other causes (recent IV contrast, medications, fluid status, etc); if other etiologies ruled out, administer 0.5-1 mg/kg/d prednisone equivalents; if worsening or no improvement: 1 to 2 mg/kg/d prednisone equivalents and permanently discontinue treatment. If improved to G1 or less, taper corticosteroids over 4-6 weeks. If no recurrence of chronic renal insufficiency, discuss resumption of ICPi with patient after taking into account the risks and benefits.</td>
</tr>
<tr>
<td>G3: Creatinine &gt; 3 × baseline or &gt; 4.0 mg/dL; hospitalization indicated</td>
<td>Permanently discontinue ICPi</td>
</tr>
<tr>
<td>G4: Life-threatening consequences; dialysis indicated</td>
<td>Consult nephrology; evaluate for other causes (recent IV contrast, medications, fluid status, etc); administer corticosteroids (initial dose of 1-2 mg/kg/d prednisone or equivalent)</td>
</tr>
</tbody>
</table>

**Additional considerations**
- Monitor creatinine weekly
- Reflex kidney biopsy should be discouraged until corticosteroid treatment has been attempted

**6.2 Symptomatic nephritis: follow-up**

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1</td>
<td>If improved to baseline, resume routine creatinine monitoring</td>
</tr>
<tr>
<td>G2</td>
<td>If improved to G1, taper corticosteroids over at least 3 weeks before resuming treatment with routine creatinine monitoring; if elevations persist &gt; 7 days or worsen and no other cause found, treat as G3</td>
</tr>
<tr>
<td>G3</td>
<td>If improved to G1, taper corticosteroids over at least 4 weeks; if elevations persist &gt; 3-6 days or worsen, consider additional immunosuppression (eg, mycophenolate)</td>
</tr>
<tr>
<td>G4</td>
<td>If improved to G1, taper corticosteroids over at least 4 weeks; if elevations persist &gt; 2-3 days or worsen, consider additional immunosuppression (eg, mycophenolate)</td>
</tr>
</tbody>
</table>

All recommendations are expert consensus based, with benefits outweighing harms, and strength of recommendations are moderate.

Abbreviations: AKI, acute kidney injury; G, grade; ICPi, immune checkpoint inhibitor; irAE, immune-related adverse event; IV, intravenous; ULN, upper limit of normal; UTI, urinary tract infection.
tubulointerstitial nephritis in 12 patients and thrombotic microangiopathy in one patient. Two of 13 patients required transient hemodialysis, and two remained on hemodialysis at the time of publication. Checkpoint inhibitor immunotherapy was discontinued in all 13 patients. Eleven patients were treated with corticosteroids, and among these 11, nine improved. One patient with thrombotic microangiopathy did not improve, despite glucocorticoids, and another patient transiently improved but then worsened. Two additional patients did not receive immunosuppression and did not recover renal function.

Checkpoint inhibitor therapy appears to be safe in patients with baseline renal impairment from a nonimmune basis (eg, prior nephrectomy, old age, hypertension); however, patients with a renal allograft are at high risk of rejecting the transplanted kidney and requiring dialysis. Limited data suggest that the risk of renal allograft rejection with anti-CTLA-4 antibodies may be less than requiring dialysis. Limited data suggest that the risk of renal allograft rejection with anti-CTLA-4 antibodies may be less than requiring dialysis. Although some patients may be able to be treated with PD-1 pathway blockers with preservation of their allografts by having adjustments in their immunosuppressive agents, this approach should only be considered with multidisciplinary input from the renal transplant nephrology team.

Patients should have their renal function (serum creatinine) checked prior to every dose of checkpoint inhibitor therapy. For those with new elevations in creatinine, one should consider holding therapy while other potential causes are evaluated (eg, recent IV radiographic contrast administration, dehydration, other medicines, urinary tract infection) and if identified, treat appropriately. Patients without other obvious causes or who do not respond to alternative treatment measures should be presumed to have immune-related renal toxicity and treated empirically according to the algorithm.

7.0 Nervous System Toxicities

Please refer to Table 7 for a complete set of recommendations, definition of grades, and additional considerations.

7.1 Myasthenia Gravis

Recommendation 7.1a – Diagnostic work-up. It is recommended that the diagnostic work-up for all grades should include the following:

- Acetylcholine receptor (AChR) and antistriated muscle antibodies in blood. If AChR antibodies are negative, consider muscle-specific kinase and lipoprotein-related 4 antibodies in blood.
- Pulmonary function assessment with negative inspiratory force and vital capacity.
- Creatine phosphokinase (CPK), aldolase, ESR, and CRP for possible concurrent myositis.
- Consider MRI of brain and/or spine, depending on symptoms, to rule out CNS involvement by disease or alternate diagnosis.
- If respiratory insufficiency or elevated CPK, troponin T, perform cardiac examination with ECG and transthoracic echocardiogram (TTE) for possible concomitant myocarditis.
- Neurology consultation.

- Electrodiagnostic studies, including neuromuscular junction testing with repetitive stimulation and/or jitter studies, nerve conduction study (NCS), to exclude neuropathy, and needle EMG to evaluate for myositis.

Recommendation 7.1b – Management. All grades warrant work-up and intervention given potential for progressive myasthenia gravis to lead to respiratory compromise. (Note: There is no grade 1 toxicity).

It is recommended that clinicians manage grade 2 toxicities as follows:

- Should hold ICPi and may resume in grade 2 patients (Myasthenia Gravis Foundation of America 1 and 2) only if symptoms resolve.
- Should consult neurology.
- Should offer pyridostigmine starting at 30 mg PO three times a day and gradually increase to a maximum of 120 mg orally four times day as tolerated and based on symptoms.
- May go directly to corticosteroids (prednisone 1 to 1.5 mg/kg PO daily) if symptoms grade 2. Wean based on symptom improvement.

It is recommended that clinicians manage grade 3 to 4 toxicities as follows:

- Should permanently discontinue ICPi.
- Should admit patient, may need intensive care unit monitoring.
- Consult neurology.
- Continue corticosteroids and initiate IVIG 2 g/kg over 5 days or plasmapheresis for 5 days.
- Should offer frequent pulmonary function assessment.
- Should offer daily neurologic evaluation.

Qualifying statement. Avoid medications that can worsen myasthenia, such as β-blockers, IV magnesium, fluoroquinolones, aminoglycosides, and macrolides. ICPi-associated myasthenia gravis may be monophasic; therefore, additional corticosteroid-sparing agents may not be required.

7.2 Guillain-Barré Syndrome

Recommendation 7.2a – Diagnostic work-up. It is recommended that the diagnostic workup should include the following:

- Neurology consultation.
- MRI of spine with and without contrast (rule out compressive lesion and evaluate for nerve root enhancement/thickening).
- Lumbar puncture: CSF typically has elevated protein and often elevated WBCs; even though this is not typically seen in classic Guillain-Barré syndrome (GBS), cytology should be sent with any CSF sample from a patient with cancer.
- Serum antiganglioside antibody tests for GBS and its subtypes (eg, anti-GQ1b for Miller Fisher variant associated with ataxia and ophthalmoplegia).
- Electrodiagnostic studies to evaluate polynuropathy.
- Pulmonary function testing (negative inspiratory force/vital capacity).
- Frequent neurochecks.

Recommendation 7.2b – Management. All grades warrant work-up and intervention given the potential for progressive GBS to lead to respiratory compromise. (Note: There is no grade 1 toxicity).
### 7.0 Nervous System Toxicities

#### 7.1 Myasthenia gravis

**Definition:** Fatigable or fluctuating muscle weakness, generally more proximal than distal. Frequently has ocular and/or bulbar involvement (ptosis, extracocular movement abnormalities resulting in double vision, dysphagia, dysarthria, facial muscle weakness). May have neck and/or respiratory muscle weakness. (Note: May occur with myositis and/or myocarditis. Respiratory symptoms may require evaluation to rule out pneumonitis, myocarditis. Miller Fisher variant of Guillain-Barré syndrome (ophthalmoparesis) and the oculobulbar myositis (ptosis, ophthalmoparesis, dysphagia, neck and respiratory weakness) with ICPi may have overlapping symptoms.)

**Diagnostic work-up**

- AChR and antistriated muscle antibodies in blood; if AChR antibodies are negative, consider muscle specific kinase and lipoprotein-related 4 antibodies in blood
- CPK, aldolase, ESR, CRP for possible concurrent myositis
- Consider MRI of brain and/or spine, depending on symptoms to rule out CNS involvement by disease or alternate diagnosis
- If respiratory insufficiency or elevated CPK, troponin T, perform cardiac examination with ECG and TTE for possible concomitant myocarditis
- Neurologic consultation
- Electrodiagnostic studies, including neuromuscular junction testing with repetitive stimulation and/or jitter studies, NCS to exclude neuropathy, and needle EMG to evaluate for myositis

**Grading**

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>All grades</td>
<td>All grades warrant work-up and intervention given potential for progressive myasthenia gravis to lead to respiratory compromise</td>
</tr>
<tr>
<td>No G1</td>
<td>Hold ICPi and may resume in G2 patients (MGFA 1 and 2) only if symptoms resolve(^{23}) Should consult neurology Pyridostigmine starting at 30 mg orally three times a day and gradually increase to maximum of 120 mg orally four times a day as tolerated and based on symptoms Administer corticosteroids (prednisolone, 1-1.5 mg/kg orally daily) if symptoms G2; wean based on symptom improvement</td>
</tr>
<tr>
<td>G2: Some symptoms interfering with ADL MGFA severity class 1 (ocular symptoms and findings only) and MGFA severity class 2 (mild generalized weakness)</td>
<td>Permanent discontinuation of ICPi Admit patient, may need ICU-level monitoring Neurology consult Continue corticosteroids and initiate IVIG 2 g/kg IV over 5 days (0.4 g/kg/d) or plasmapheresis for 5 days Frequent pulmonary function assessment Daily neurologic review</td>
</tr>
<tr>
<td>G3-4: Limiting self-care and aids warranted, weakness limiting walking, ANY dysphagia, facial weakness, respiratory muscle weakness, or rapidly progressive symptoms, or MGFA severity class 3-4 moderate to severe generalized weakness to myasthenic crisis</td>
<td>Admit patient, may need ICU-level monitoring Neurology consult Continue corticosteroids and initiate IVIG 2 g/kg IV over 5 days (0.4 g/kg/d) or plasmapheresis for 5 days Frequent pulmonary function assessment Daily neurologic review</td>
</tr>
</tbody>
</table>

**Additional considerations**

- Avoid medications that can worsen myasthenia: β-blockers, IV magnesium, fluoroquinolones, aminoglycosides, and macrolides
- Initially a 5-day course of plasmapheresis or a 2 g/kg course of IVIG over 5 days
- 1-2 mg/kg methylprednisolone daily, wean based on symptom improvement
- Pyridostigmine, wean based on improvement
- ICPi-associated myasthenia gravis may be monophasic, and additional corticosteroid-sparing agents may not be required

**Diagnosis work-up**

- MRI of spine with or without contrast (rule out compressive lesion and evaluate for nerve root enhancement/thickening)
- Lumbar puncture: CSF typically has elevated protein and often elevated WBCs; even though this is not typically seen in classic Guillain-Barré syndrome, cytology should be sent with any CSF sample from a patient with cancer.
- Serum antiganglioside antibody tests for Guillain-Barré syndrome and its subtypes (eg, anti-GQ1b for Miller Fisher variant associated with ataxia and ophthalmoplegia)
- Electrodiagnostic studies to evaluate polyneuropathy
- Pulmonary function testing (NIF/VC)
- Frequent neurochecks

**Grading**

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>All grades</td>
<td>Warrant work-up and intervention given potential for progressive Guillain-Barré syndrome to lead to respiratory compromise Note: There is no G1 toxicity</td>
</tr>
<tr>
<td>G1: Mild, none</td>
<td>NA</td>
</tr>
<tr>
<td>G2: Severe, some interference with ADL, symptoms concerning to patient</td>
<td>Discontinue ICPi Admission to inpatient unit with capability of rapid transfer to ICU-level monitoring Start IVIG (0.4 g/kg/d for 5 days for a total dose of 2 g/kg) or plasmapheresis. Corticosteroids are usually not recommended for idiopathic Guillain-Barré syndrome; however, in ICPi-related forms, a trial is reasonable (methylprednisolone 2-4 mg/kg/d), followed by slow corticosteroid taper Pulse corticosteroid dosing (methylprednisolone 1 g/d for 5 days) may also be considered for G3-4 along with IVIG or plasmapheresis Frequent neurochecks and pulmonary function monitoring Monitor for concurrent autonomic dysfunction Nonopioid management of neuropathic pain Treatment of constipation/ileus</td>
</tr>
<tr>
<td>G3-4: Severe, limiting self-care and aids warranted, weakness limiting walking, ANY dysphagia, facial weakness, respiratory muscle weakness, or rapidly progressive symptoms</td>
<td>(continued on following page)</td>
</tr>
</tbody>
</table>
Additional considerations
Slow prednisone taper after corticosteroid pulse plus IVIG or plasmapheresis
May require repeat IVIG courses
Caution with rechallenging for severe cases

7.3 Peripheral neuropathy
Definition: Can present as asymmetric or symmetric sensory, motor, or sensory motor deficit. Focal mononeuropathies, including cranial neuropathies (eg, facial neuropathies/Bell palsy) may be present. Numbness and paresthesias may be painful or painless. Hypo- or areflexia or sensory ataxia may be present.

Diagnostic work-up
G1
Screen for reversible neuropathy causes: diabetic screen, B12, folate, TSH, HIV, consider serum protein electrophoresis, and other vasculitic and autoimmune screen
Neurologic consultation
Consider MRI of spine with or without contrast
G2: in addition to above
MRI spine advised/MRI of brain if cranial nerve
Consider EMG/NCS
Consider neurology consultation
G3-4: go to Guillain-Barré syndrome algorithm

Grading Management
G1: Mild, no interference with function and symptoms not concerning to patient. Note: Any cranial nerve problem should be managed as moderate
Low threshold to hold ICPi and monitor symptoms for a week
If to continue, monitor very closely for any symptom progression
G2: Moderate, some interference with ADL, symptoms concerning to patient (ie, pain but no weakness or gait limitation)
Hold ICPi and resume once return to G1
Initial observation OR initiate prednisone 0.5-1 mg/kg (if progressing from mild)
Neurontin, pregabalin, or duloxetine for pain
G3-4: Severe, limiting self-care and aids warranted, weakness limiting walking or respiratory problems (ie, leg weakness, foot drop, rapidly ascending sensory changes).
Severe may be Guillain-Barré syndrome and should be managed as such
Permanently discontinue ICPi
Admit patient
Neurologic consultation
Initiate IV methylprednisolone 2-4 mg/kg and proceed as per Guillain-Barré syndrome management

7.4 Autonomic neuropathy
Definition: Nerves that control involuntary bodily functions are damaged. This may affect blood pressure, temperature control, digestion, bladder function, and sexual function. A case of severe enteric neuropathy with ICPi has been reported. Can present with GI difficulties such as new severe constipation, nausea, urinary problems, sexual difficulties, sweating abnormalities, sluggish pupil reaction, and orthostatic hypertension.

Diagnostic work-up
An evaluation by neurologist or relevant specialist, depending on organ system, with testing that may include
Screen for other causes of autonomic dysfunction: diabetic screen, adrenal insufficiency, HIV, paraproteinemia, amyloidosis, botulism, consider chronic diseases such as Parkinson’s and other autoimmune screen
• Orthostatic vital signs
• Consider electrodagnostic studies to evaluate for concurrent polyneuropathy
• Consider paraneoplastic autonomic dysautonomia antibody testing (eg, anti-ganglionic acetylcholine receptor, antineuronal nuclear antibody type 1 [ANNA-1], and N-type voltage gated calcium channel antibodies)

Grading Management
G1: Mild, no interference with function and symptoms not concerning to patient
Low threshold to hold ICPi and monitor symptoms for a week; if to continue, monitor very closely for any symptom progression
G2: Moderate, some interference with ADL, symptoms concerning to patient
Hold ICPi and resume once return to G1
Initial observation OR initiate prednisone 0.5-1 mg/kg (if progressing from mild)
Neurologic consultation
G3-4: Severe, limiting self-care and aids warranted
Permanently discontinue ICPi
Admit patient
Initiate methylprednisolone 1 g daily for 3 days followed by oral corticosteroid taper
Neurologic consultation

7.5 Aseptic meningitis
Definition: may present with headache, photophobia, and neck stiffness; often afebrile but may be febrile. There may be nausea/vomiting. Mental status should be normal (distinguishes from encephalitis).

Diagnostic work-up
MRI of brain with or without contrast + pituitary protocol
Consider lumbar puncture: measure opening pressure; check cell count and protein glucose; and perform Gram stain, culture, PCR for HSV, and other viral PCRs depending on suspicion, cytology
May see elevated WBC count with normal glucose, normal culture, and Gram stain; may see reactive lymphocytes or histiocytes on cytology

(continued on following page)
It is recommended that clinicians manage all-grade toxicities as follows:

- Should discontinue ICPi.
- Admission to inpatient unit with capability for rapid transfer to intensive care unit—level monitoring.
- Corticosteroids are not usually recommended for idiopathic GBS; however, in ICPi-related forms, a trial is reasonable. Should start IVIG 0.4 g/kg/d for 5 days for a total dose of 2 g/kg or plasmapheresis plus concurrent corticosteroids (methylprednisolone 2 to 4 mg/kg/d).
- Should offer frequent neurochecks and pulmonary function monitoring.
- Should monitor for concurrent autoimmune dysfunction.
- Nonopioid management of neuropathic pain.
- Treatment of constipation/ileus.

### 7.3 Peripheral Neuropathy

**Recommendation 7.3a – Diagnostic work-up.** It is recommended that the diagnostic work-up for grade 1 should include the following:

1. Admission to inpatient unit with capability for rapid transfer to intensive care unit—level monitoring.
2. Corticosteroids are not usually recommended for idiopathic GBS; however, in ICP-related forms, a trial is reasonable. Should start IVIG 0.4 g/kg/d for 5 days for a total dose of 2 g/kg or plasmapheresis plus concurrent corticosteroids (methylprednisolone 2 to 4 mg/kg/d).
3. Should offer frequent neurochecks and pulmonary function monitoring.
4. Should monitor for concurrent autoimmune dysfunction.
5. Nonopioid management of neuropathic pain.
6. Treatment of constipation/ileus.

### Table 7. Management of Nervous System irAEs in Patients Treated With ICPis (continued)

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1: Mild, no interference with function and symptoms not concerning to patient. Note: Any cranial nerve problem should be managed as moderate.</td>
<td>Hold ICPi and discuss resumption with patient only after taking into account the risks and benefits.</td>
</tr>
<tr>
<td>G2: Moderate, some interference with ADL, symptoms concerning to patient (ie, pain but no weakness or gait limitation)</td>
<td>Consider empirical antiviral (IV acyclovir) and antibacterial therapy until CSF results obtained and negative.</td>
</tr>
<tr>
<td>G3-4: Severe, limiting self-care and aids warranted</td>
<td>Trial of methylprednisolone 1-2 mg/kg.</td>
</tr>
</tbody>
</table>

**7.6 Encephalitis**

**Definition:** As for aseptic meningitis, need to exclude infectious causes, especially viral (ie, HSV).

<table>
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<tr>
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<td>Hold ICPi and discuss resumption with patient only after taking into account the risks and benefits.</td>
</tr>
<tr>
<td>G2: Moderate, some interference with ADL, symptoms concerning to patient (ie, pain but no weakness or gait limitation)</td>
<td>As above for aseptic meningitis, suggest concurrent IV acyclovir until PCR results obtained and negative.</td>
</tr>
<tr>
<td>G3-4: Severe, limiting self-care and aids warranted</td>
<td>If severe or progressing symptoms or oligoclonal bands present, consider pulse corticosteroids methylprednisolone 1 g IV daily for 3-5 days plus IVIG 2 g/kg over 5 days.</td>
</tr>
</tbody>
</table>

**7.7 Transverse myelitis**

**Definition:** Acute or subacute weakness or sensory changes bilateral, often with increased deep tendon reflexes.

<table>
<thead>
<tr>
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</tr>
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<tbody>
<tr>
<td>G1: Mild, no interference with function and symptoms not concerning to patient. Note: Any cranial nerve problem should be managed as moderate.</td>
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</tr>
<tr>
<td>G2: Moderate, some interference with ADL, symptoms concerning to patient (ie, pain but no weakness or gait limitation)</td>
<td>As above for aseptic meningitis, suggest concurrent IV acyclovir until PCR results obtained and negative.</td>
</tr>
<tr>
<td>G3-4: Severe, limiting self-care and aids warranted</td>
<td>If severe or progressing symptoms or oligoclonal bands present, consider pulse corticosteroids methylprednisolone 1 g IV daily for 3-5 days plus IVIG 2 g/kg over 5 days.</td>
</tr>
</tbody>
</table>

**Abbreviations:** AChR, acetylcholine receptor; ACTH, adrenocorticotropic hormone; ADL, activities of daily living; ANCA, antineutrophil cytoplasmic antibodies; CPK, creatine phosphokinase; CRP, C-reactive protein; EMG, electromyography; HSV, herpes simplex virus; ICPi, immune checkpoint inhibitor; ICU, intensive care unit; IgG, immunoglobulin G; IV, intravenous; IVIG, intravenous immunoglobulin; irAE, immune-related adverse event; MGFA, Myasthenia Gravis Foundation of America; MRI, magnetic resonance imaging; NA, not applicable; NCS, nerve conduction study; NIF, negative inspiratory force; PCR, polymerase chain reaction; RPR, rapid plasma reagin; TPO, thyroid peroxidase; TSH, thyroid-stimulating hormone; TTE, transthoracic echocardiogram; VC, vital capacity.
• Screen for reversible neuropathy causes: diabetic screen, B12, folate, TSH, HIV. Consider serum protein electrophoresis and other vasculitic and autoimmune screen.
• Consider MRI of spine with or without contrast.
• Consider neurology consultation.

It is recommended that the diagnostic work-up for grade 2 should include the following, in addition to what is recommended for grade 1:
• MRI of spine advised; MRI of brain if cranial nerve.
• Consider EMG/NCS.
• Consider neurology consultation

It is recommended that the diagnostic work-up for grades 3 to 4 should follow that of GBS.

**Recommendation 7.3b – Management.** It is recommended that clinicians manage grade 1 toxicities as follows:
• Should have a low threshold to hold ICPI and monitor symptoms for a week. If to continue, should monitor very closely for any symptom progression.
• Should hold ICPI and resume once return to grade 1.
• Should offer initial observation OR may initiate prednisone 0.5 to 1 mg/kg (if progressing from mild).
• Should monitor off corticosteroids or consider oral prednisone 0.5 to 1 mg/kg or IV methylprednisolone 1 g daily for 3 days followed by oral corticosteroid taper.

**7.4 Autonomic Neuropathy**

**Recommendation 7.4a – Diagnostic work-up.** It is recommended that the diagnostic work-up should include the following:
An evaluation by neurologist or relevant specialist, depending on organ system, with testing that may include:
• Screening for other causes of autonomic dysfunction: diabetic screen, adrenal insufficiency, HIV, paraproteinemia, amyloidosis, botulism. Consider chronic diseases such as Parkinson and other autoimmune screening.
• Orthostatic vital signs.
• Consider electrodiagnostic studies to evaluate for concurrent polyneuropathy.
• Consider paraneoplastic autoimmune dysautonomia antibody testing (eg, anti-ganglionic acetylcholine receptor, antineuronal nuclear antibody type 1 [ANNA-1], and N-type voltage gated calcium channel antibodies).

**Recommendation 7.4b – Management.** It is recommended that clinicians manage grade 1 toxicities as follows:
• Should have a low threshold to hold ICPI and monitor symptoms for a week. If to continue, should monitor very closely for any symptom progression.
• Should offer initial observation OR may initiate prednisone 0.5 to 1 mg/kg (if progressing from mild).
• Should consult neurology.
• Should offer neurontin, pregabalin, or duloxetine for pain.
• Should proceed as per GBS management.

**Recommendation 7.5a – Diagnostic work-up.** It is recommended that the diagnostic work-up for grade 2 toxicities should include the following:
• MRI of brain with or without contrast may reveal T2/fluid-attenuated inversion changes typical of what is seen in autoimmune encephalopathies or limbic encephalitis or may be normal.
• Lumbar puncture: check cell count and protein glucose and perform Gram stain, culture, and polymerase chain reaction (PCR) for herpes simplex virus and other viral PCRs depending on suspicion, cytology.
• May see elevated WBC count with normal glucose, normal culture, and Gram stain. May see reactive lymphocytes or histiocytes on cytology.

**Recommendation 7.5b – Management.** It is recommended that clinicians manage grade 2 toxicities as follows:
• Should permanently discontinue ICPI.
• Should admit patient.
• Should initiate methylprednisolone 1 g daily for 3 days followed by oral corticosteroid taper.
• Should consult neurology.

**7.5 Aseptic Meningitis**

**Recommendation 7.5a – Diagnostic work-up.** It is recommended that the diagnostic work-up should include the following:
• MRI of brain with or without contrast and pituitary protocol.
• Cortisol (AM), adrenocorticotropic hormone (ACTH) to rule out adrenal insufficiency.
• Consider lumbar puncture: measure opening pressure; check cell count and protein glucose; and perform Gram stain, culture, and polymerase chain reaction (PCR) for herpes simplex virus and other viral PCRs depending on suspicion, cytology.
• Consider MRI of brain with or without contrast and pituitary protocol.
• Should permanently discontinue ICPI.
• Should hold corticosteroids or consider oral prednisone 0.5 to 1 mg/kg or IV methylprednisolone 1 mg/kg if moderate/severe symptoms.

**7.6 Encephalitis**

**Recommendation 7.6a – Diagnostic work-up.** It is recommended that the diagnostic work-up should include the following:
• Neurologic consultation.
• MRI of brain with or without contrast may reveal T2/fluid-attenuated inversion recovery changes typical of what is seen in autoimmune encephalopathies or limbic encephalitis or may be normal.
• Lumbar puncture: check cell count and protein glucose and perform Gram stain, culture, PCR for herpes simplex virus, and other viral PCRs depending on suspicion, cytology, oligoclonal bands, autoimmune encephalopathy, and paraneoplastic panels.
• May see elevated WBC count with lymphocytic predominance and/or elevated protein.
• EEG to evaluate for subclinical seizures.
• Blood: metabolic; CBC; ESR; CRP; antineutrophil cytoplasmic antibodies (if suspect vasculitic process); and thyroid panel, including thyroid peroxidase and thyroglobulin.
• Rule out concurrent anemia/thrombotic thrombocytopenic purpura (TTP) as cause of encephalopathy; check peripheral smear.

**Recommendation 7.6b – Management.** It is recommended that clinicians manage all-grade toxicities as follows:
- Should hold ICPi.
- As above for aseptic meningitis, suggest concurrent IV acyclovir until PCR results obtained and negative.
- Should offer a trial of methylprednisolone 1 to 2 mg/kg.
- If severe or progressing symptoms or oligoclonal bands present, may offer pulse corticosteroid methylprednisolone 1 g IV daily for 3 to 5 days plus IVIG 2 g/kg over 5 days.
- If positive for autoimmune encephalopathy or paraneoplastic antibody and limited or no improvement, may offer rituximab or plasmapheresis in consultation with neurology.

### 7.7 Transverse Myelitis

**Recommendation 7.7a – Diagnostic work-up.** It is recommended that the diagnostic work-up should include the following:
- Neurologic consultation
- MRI of spine (with thin axial cuts through the region of suspected abnormality) and MRI of brain done with and without contrast
- Lumbar puncture: cell count, protein, glucose, oligoclonal bands, viral PCRs, cytology, onconeural antibodies
- Blood: B12, HIV, rapid plasma reagin, ANA, Ro/La, TSH, aquaporin-4 immunoglobulin G
- Evaluation for urinary retention and constipation

**Recommendation 7.7b – Management.** It is recommended that clinicians manage all-grade toxicities as follows:
- Should permanently discontinue ICPi.
- Should administer methylprednisolone 2 mg/kg.
- Should strongly consider higher doses of 1 g/d for 3 to 5 days.
- Should strongly consider IVIG.

**Discussion.** ICPi-related neurologic toxicities were originally reported with 1% incidence; however, more-recent analyses suggest that they are more common.\(^8^8\)-\(^9^0\) An analysis of 59 trials totaling 9,208 patients reported the overall incidence of neurologic irAEs to be 3.8% in patients receiving anti–CTLA-4 antibodies, 6.1% in patients receiving anti–PD-1 antibodies, and 12.0% in patients receiving a combination of both.\(^9^0\) However, the incidence of grade 3 and 4 irAEs was <1% across all ICPis. A review of patients who received ICPi therapy for melanoma at the Royal Marsden Hospital found the rate of neurologic irAEs to be 2.4%.\(^9^0\) The EORTC 18071 trial reported neurologic irAEs at a rate of 4% in the adjuvant ipilimumab arm.\(^9^1\) Most neurologic events are mild; headache and peripheral sensory neuropathy are the most commonly encountered symptoms.\(^8^8\) Severe neurologic irAEs grade 3 or higher occur in <1% of patients and may involve the peripheral nervous system or CNS. They encompass a broad spectrum of neurologic syndromes, including myasthenia gravis/myasthenic syndrome, aseptic meningitis, encephalitis, sensory motor neuropathy or Guillain-Barré-like syndromes, painful sensory neuropathy, enteric neuropathy, transverse myelitis, and posterior reversible encephalopathy syndrome.

The first step in management is to rule out CNS progression of cancer, seizure activity, infection, and metabolic derangement as causes of neurologic symptoms. Consultation with a neurologist is advised for all neurologic irAEs grade 2 or higher to help to determine the type and severity of neurologic impairment and guide selection and interpretation of further neurologic tests and management. In patients presenting with headache (which, in isolation, could suggest aseptic meningitis), it is important to evaluate for new confusion, altered behavior, aphasia, seizure-like activity, or short-term memory loss, any of which might suggest encephalitis. The distinction is important because suspected encephalitis triggers a distinct work-up and management from aseptic meningitis, including autoimmune encephalitis and paraneoplastic antibody evaluation and consideration of pulse-dose corticosteroids.\(^9^2\)-\(^9^5\)

For most neurologic irAEs, diagnostic work-up should include MRI of the brain and/or spine with and without contrast and CSF analysis, including cytology, to rule out leptomeningeal metastasis. CSF analysis is helpful in cases of clinical suspicion of encephalitis, aseptic meningitis, and sensory motor neuropathy or GBS, revealing lymphocytic pleocytosis and elevated protein in many cases. Abnormal leptomeningeal enhancement on neuroimaging may occur in aseptic meningitis, encephalitis, and sensory motor neuropathy, underscoring the importance of checking CSF cytology, which should be negative. NCSs and EMG may assist in diagnosis of sensory symptoms or weakness. Autonomic neuropathy may occur along with other neuropathy symptoms and should be screened for. EEG is helpful for ruling out seizure activity in cases of encephalopathy. Paraneoplastic neurologic syndromes and autoimmune encephalopathies should also be considered.\(^9^2\)

For mild (grade 1) neurologic symptoms, checkpoint inhibitor therapy may be continued under close observation. For grade 2 or higher neurologic symptoms, checkpoint inhibitor therapy should be held until the nature of the irAE and symptom progression is defined. In the event of significant neurologic toxicity of grade 2 or higher, a corticosteroid equivalent of methylprednisolone 1 to 4 mg/kg, depending on the symptoms, should be started. For more-severe grade 3 or higher toxicity, immunotherapy should be discontinued. Symptom control may require escalation of corticosteroid therapy to pulse-dose methylprednisolone (1 g daily for 5 days) in addition to IVIG, or plasma exchange (PEX). Pyridostigmine may be helpful for myasthenia gravis in addition to corticosteroids.

### 8.0 Hematologic Toxicities

Please refer to **Table 8** for a complete set of recommendations, definition of grades, and additional considerations.

#### 8.1 Autoimmune Hemolytic Anemia

**Recommendation 8.1a – Diagnostic work-up.** It is recommended that the diagnostic work-up should include the following:
- History and physical examination (with special consideration of history of new drugs and insect, spider, or snake bites)
Management of Immune-Related Adverse Events

8.0 Hematologic Toxicities

8.1 Autoimmune hemolytic anemia

Definition: A condition in which RBCs are destroyed and removed from the blood stream before their normal lifespan is over. Symptoms include weakness, paleness, jaundice, dark-colored urine, fever, inability to do physical activity, and heart murmur.

Diagnostic work-up
- History and physical examination (with special consideration of history of new drugs and insect, spider, or snake bites)
- Blood chemistry, CBC with evidence of anemia, macrocytosis, evidence of hemolysis on peripheral smear; LDH, haptoglobin, bilirubin, reticulocyte count; free Hgb
- DIC panel, which could include PTINR infectious causes
- Autoimmune serology
- Paroxysmal nocturnal hemoglobinuria screening
- Direct and indirect bilirubin; LDH; direct agglutinin test; and if no obvious cause, bone marrow analysis, cytogenetic analysis to evaluate for myelodysplastic syndromes
- Evaluation for viral/bacterial (mycoplasma, etc) causes of hemolysis studies
- Protein electrophoresis, cryoglobulin analysis
- Work-up for bone marrow failure syndrome if refractory, including B12, folate, copper, parvovirus, FE, thyroid, infection
- Glucose-6-phosphate dehydrogenase
- Evaluation of common drug causes (ribavirin, rifampin, dapsone, interferon, cephalosporins, penicillins, NSAIDs, quinine/quinidine, lorazepam, diclofenac, etc)
- Assessment of methemoglobinemia

Table 8. Management of Hematologic irAEs in Patients Treated With ICPis

<table>
<thead>
<tr>
<th>Grade</th>
<th>Management</th>
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<tbody>
<tr>
<td>G1: Hgb &lt; LLN to 10.0 g/dL; &lt; LLN to 6.2 mmol/L; &lt; LLN to 100 g/L</td>
<td>Continue ICPi with close clinical follow-up and laboratory evaluation</td>
</tr>
<tr>
<td>G2: Hgb &lt; 10.0 to 8.0 g/dL; &lt; 6.2 to 4.9 mmol/L; &lt; 100 to 80 g/L</td>
<td>Hold ICPi and strongly consider permanent discontinuation Administer 0.5-1 mg/kg/d prednisone equivalents</td>
</tr>
<tr>
<td>G3: Hgb &lt; 8.0 g/dL; &lt; 4.9 mmol/L; &lt; 80 g/L; transfusion indicated</td>
<td>Permanently discontinue ICPi Should use clinical judgment and consider admitting the patient Hematology consult Prednisone 1-2 mg/kg/d (oral or IV depending on symptoms/speed of development) If worsening or no improvement, 1-2 mg/kg/d prednisone equivalents and permanently discontinue ICPi treatment Consider RBC transfusion per existing guidelines; do not transfuse more than the minimum number of RBC units necessary to relieve symptoms of anemia or to return a patient to a safe Hgb range (7-8 g/dL in stable, noncardiac inpatients) Should offer patients supplementation with folic acid 1 mg once daily</td>
</tr>
<tr>
<td>G4: Life-threatening consequences, urgent intervention indicated</td>
<td>Permanently discontinue ICPi Admit patient Hematology consult IV prednisone corticosteroids 1-2 mg/kg/d If no improvement or if worsening while on corticosteroids or severe symptoms on presentation, initiate other immunosuppressive drugs, such as rituximab, IVIG, cyclosporin A, and mycophenolate mofetil RBC transfusion per existing guidelines; discuss with blood bank team prior to transfusions that a patient with possible ICPi serious AE is in house.</td>
</tr>
</tbody>
</table>

Additional considerations: Monitor Hgb levels on a weekly basis until the corticosteroid tapering process is complete; thereafter, less-frequent testing is needed164

8.2 Acquired TTP

Definition: A disorder characterized by the presence of microangiopathic hemolytic anemia, thrombocytopenic purpura, fever, renal abnormalities, and neurologic abnormalities, such as seizures, hemiplegia, and visual disturbances. It is an acute or subacute condition.

Diagnostic work-up
- History with specific questions related to drug exposure (eg, chemotherapy, sirolimus, tacrolimus, opana ER antibiotics, quinine)
- Physical examination, peripheral smear
- ADAMTS13 activity level and inhibitor titer
- LDH, haptoglobin, reticulocyte count, bilirubin, urinalysis to rule out other causes
- PT, activated PTT, fibrinogen
- Blood group and antibody screen, direct antiglobulin test, CMV serology
- Consider CT/MRI brain, echocardiogram, ECG
- Viral studies
- Note: This disorder is usually associated with a severe drop in platelets and hemolysis/anemia precipitously

Table 8. Management of Hematologic irAEs in Patients Treated With ICPis (continued on following page)

<table>
<thead>
<tr>
<th>Grade</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>All grades</td>
<td>The first step in the management of TTP is a high index of suspicion for the diagnosis and timely recognition; hematology consult should immediately be called, as delay in identification is associated with increased mortality/morbidity. Initially, the patient should be stabilized and any critical organ dysfunction stabilized</td>
</tr>
<tr>
<td>G1: Evidence of RBC destruction (schistocytosis) without anemia, renal insufficiency, or thrombocytopenia clinically</td>
<td>Hold ICPi and discuss resumption with patient only after taking into account the risks and benefits, noting that there are currently no data to recommend restarting ICPi therapy Hematology consult Administer 0.5-1 mg/kg/d prednisone</td>
</tr>
<tr>
<td>G2: Evidence of RBC destruction (schistocytosis) without clinical consequence with G2 anemia and thrombocytopenia</td>
<td>(continued on following page)</td>
</tr>
</tbody>
</table>
### 8.0 Hematologic Toxicities

<table>
<thead>
<tr>
<th>Grade</th>
<th>Clinical Findings</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1</td>
<td>Laboratory findings with clinical consequences (G3 thrombocytopenia, anemia, renal insufficiency &gt; 2)</td>
<td>Hold ICPi and discuss resumption with patient only after taking into account the risks and benefits, noting that there are currently no data to recommend restarting ICPi therapy</td>
</tr>
<tr>
<td>G2</td>
<td>Laboratory findings with clinical consequences (eg, renal insufficiency, petechiae)</td>
<td>Permanently discontinue ICPi</td>
</tr>
<tr>
<td>G3</td>
<td>Life-threatening consequences (eg, CNS thrombosis/embolism or renal failure)</td>
<td>Begin therapy with eculizumab therapy 900 mg weekly for four doses, 1,200 mg week 5, then 1,200 mg every 2 weeks</td>
</tr>
<tr>
<td>G4</td>
<td>Life-threatening consequences (eg, CNS hemorrhage or thrombosis/embolism or renal failure)</td>
<td>Red blood transfusion according to existing guidelines</td>
</tr>
</tbody>
</table>

### 8.3 Hemolytic Uremic Syndrome

**Definition:** A disorder characterized by a form of thrombotic microangiopathy with renal failure, hemolytic anemia, and severe thrombocytopenia. Signs and symptoms of hemolytic uremic syndrome can include:
- Bloody diarrhea
- Decreased urination or blood in the urine
- Abdominal pain, vomiting, and occasionally fever
- Pallor
- Small, unexplained bruises or bleeding from the nose and mouth
- Fatigue and irritability
- Confusion or seizures
- High blood pressure
- Swelling of the face, hands, feet, or entire body

**Diagnostic work-up**
- History and physical examination (special consideration for new history of high-risk drugs, hypertension, or cardiac causes)
- CBC with indices
- Blood smear morphology. Note that the presence of schistocytes on smear is critical for diagnosis.
- Serum creatinine
- ADAMTS13 (to rule out TTP)
- Homocysteine/methylmalonic acid
- Complement testing C3, C4, CH50 (complement inhibitory antibodies for suspected familial)
- Evaluate reticulocyte count and mean corpuscular volume
- Evaluation of infectious cause, including screening for EBV, CMV, HHV6
- Evaluation for nutritional causes of macrocytosis (B12 and folate)
- Pancreatic enzymes
- Evaluation for diarrheal causes, shiga toxin, *Escherichia coli* 0157, etc
- Direct antibody test (Coombs test), haptoglobin, LDH, and other etiologies of anemia
- Evaluation for common drugs causing hemolysis (tacrolimus, cyclosporine, sirolimus, etc)
- Evaluation for concurrent confusion
- Complement testing C3, C4, CH50 (complement inhibitory antibodies for suspected familial)
- Evaluate reticulocyte count and mean corpuscular volume
- Evaluation of infectious cause, including screening for EBV, CMV, HHV6
- Evaluation for nutritional causes of macrocytosis (B12 and folate)
- Pancreatic enzymes
- Evaluation for diarrheal causes, shiga toxin, *Escherichia coli* 0157, etc
- Direct antibody test (Coombs test), haptoglobin, LDH, and other etiologies of anemia
- Evaluation for common drugs causing hemolysis (tacrolimus, cyclosporine, sirolimus, etc)
- Evaluation for concurrent confusion

### 8.4 Aplastic Anemia

**Definition:** Condition in which the body stops producing enough new blood cells.

**Diagnostic work-up**
- History and physical examination (close attention to medications, exposure to radiation, toxins, recent viral infections)
- CBC, smear, reticulocyte count
- Viral studies, including CMV, HHV6, EBV, parvovirus
- Nutritional assessments including B12, folate, iron, copper, ceruloplasmin, vitamin D
- Serum LDH, renal function
- Work-up for infectious causes
- Identify marrow hypoplasia
- Bone marrow biopsy and aspirate analysis
- Peripheral blood analysis, including neutrophil count, proportion of GPI-negative cells by flow for PNH
- Flow cytometry to evaluate loss of GPI-anchored proteins
- Type and screen patient for transfusions and notify blood bank that all transfusions need to be irradiated and filtered

### Grading

<table>
<thead>
<tr>
<th>Grade</th>
<th>Clinical Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1</td>
<td>Nonsevere, &gt; 0.5 polymorphonuclear cells × 10^9/L, hypopcellular marrow, with marrow cellularity &lt; 25%, peripheral platelet count &gt; 20,000, reticulocyte count &gt; 20,000</td>
</tr>
<tr>
<td>G2</td>
<td>Severe, hypopcellular marrow &lt; 25% and two of the following: ANC &lt; 900, peripheral platelet &lt; 20,000, and reticulocyte &lt; 20,000</td>
</tr>
</tbody>
</table>

**Management**
- Hold ICPi and discuss resumption with patient only after taking into account the risks and benefits, noting that there are currently no data to recommend restarting ICPi therapy
- Supportive transfusions as per local guidelines
- Supportive care with granulocyte colony-stimulating factor may be added in addition
- Supportive care with granulocyte colony-stimulating factor may be added in addition
- Supportive care with granulocyte colony-stimulating factor may be added in addition
- Supportive care with granulocyte colony-stimulating factor may be added in addition
- Supportive care with granulocyte colony-stimulating factor may be added in addition

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Table 8: Management of Hematologic irAEs in Patients Treated With ICPis (continued)

<table>
<thead>
<tr>
<th>Grade</th>
<th>Clinical Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1</td>
<td>Nonsevere, &gt; 0.5 polymorphonuclear cells × 10^9/L, hypopcellular marrow, with marrow cellularity &lt; 25%, peripheral platelet count &gt; 20,000, reticulocyte count &gt; 20,000</td>
</tr>
<tr>
<td>G2</td>
<td>Severe, hypopcellular marrow &lt; 25% and two of the following: ANC &lt; 900, peripheral platelet &lt; 20,000, and reticulocyte &lt; 20,000</td>
</tr>
</tbody>
</table>

**Management**
- Hold ICPi and discuss resumption with patient only after taking into account the risks and benefits, noting that there are currently no data to recommend restarting ICPi therapy
- Supportive transfusions as per local guidelines
- Supportive care with granulocyte colony-stimulating factor may be added in addition
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- Supportive care with granulocyte colony-stimulating factor may be added in addition
- Supportive care with granulocyte colony-stimulating factor may be added in addition
- Supportive care with granulocyte colony-stimulating factor may be added in addition

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**Hematology consult**
- In conjunction with hematology, initiate PEX according to existing guidelines with further PEX dependent on clinical progress
- Administer methylprednisolone 1 g IV daily for 3 days, with the first dose typically administered immediately after the first PEX
- May offer rituximab

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**Table 8. Management of Hematologic irAEs in Patients Treated With ICPis (continued)**
### Table 8. Management of Hematologic irAEs in Patients Treated With ICPis (continued)

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>G3-4: Very severe. ANC &lt; 200; platelet count &lt; 20,000; reticulocyte count &lt; 20,000, plus hypocellular marrow &lt; 25%</td>
<td>Hold ICPi and monitor weekly for improvement; if not resolved, discontinue treatment until AE has reverted to G1. Hematology consult, growth factor support. Horse ATG plus cyclosporine. If no response, repeat immunosuppression with rabbit ATG plus cyclosporine, cyclophosphamide. For refractory patients, consider eltrombopag plus supportive care.</td>
</tr>
</tbody>
</table>

#### 8.5 Lymphopenia

**Definition:** An abnormally low level of lymphocytes in PB; for adults, counts of < 1,500/mm³

**Diagnostic work-up**
- History and physical examination (special attention for lymphocyte-depleting therapy such as fludarabine, ATG, corticosteroids, cytotoxic chemotherapy, radiation exposure, etc., as well as history of autoimmune disease, family history of autoimmune disease)
- Evaluation of nutritional state as cause
- Spleen size
- CBC with differential, peripheral smear and reticulocyte counts
- CXR for evaluation of presence of thymoma
- Bacterial cultures and evaluation for infection (fungal, viral, bacterial specifically CMV/HIV)

**Grading**
- G1-2: 500-1,000 PB lymphocyte count
- G3: 250-499 PB lymphocyte count
- G4: < 250 PB lymphocyte count

**Management**
- G1-2: Continue ICPi
- G3: Continue ICPi; checking CBC weekly for monitoring, initiation of CMV screening
- G4: Consider holding ICPi
- Initiate Mycobacterium avium complex prophylaxis and Pneumocystis jirovecii prophylaxis, CMV screening. HIV/hepatitis screening if not already done
- May consider EBV testing if evidence of lymphadenopathy/hepatitis, fevers, hemolysis consistent with lymphoproliferative disease

#### 8.6 Immune thrombocytopenia

**Definition:** An autoimmune disorder characterized by immunologic destruction of otherwise normal platelets

**Diagnostic work-up**
- History and physical examination (special attention for lymphocyte-depleting therapy, such as fludarabine, ATG, corticosteroids, cytotoxic therapy)
- Family history of autoimmunity or personal history of autoimmune disease
- History of viral illness
- CBC
- Peripheral blood smear, reticulocyte count
- Bone marrow evaluation only if abnormalities in the above test results and further investigation is necessary for a diagnosis
- Patients with newly diagnosed immune thrombocytopenia should undergo testing for HIV, hepatitis C virus, hepatitis B virus, and Helicobacter pylori
- Direct antigen test should be checked to rule out concurrent Evan syndrome
- Nutritional evaluation
- Bone marrow evaluation if other cell lines affected and concern for aplastic anemia

**Grading**
- G1: Platelet count < 100/µL
- G2: Platelet count < 75/µL
- G3: Platelet count < 50/µL
- G4: Platelet count < 25/µL

**Management**
- G1: Continue ICPi with close clinical follow up and laboratory evaluation
- G2: Hold ICPi but monitor for improvement; if not resolved, interrupt treatment until AE has reverted to G1. Administer prednisone 1 mg/kg/d (dosage range, 0.5-2 mg/kg/d) orally for 2-4 weeks after which time this medication should be tapered over 4-6 weeks to the lowest effective dose. IVIG may be used in conjunction with corticosteroids if a more-rapid increase in platelet count is required.
- G3: Hold ICPi but monitor for improvement; if not resolved, interrupt treatment until AE has reverted to G1. Hematology consult. Prednisone 1-2 mg/kg/d (oral or IV depending on symptoms). If worsening or no improvement, 1-2 mg/kg/d prednisone equivalents and permanently discontinue treatment. IVIG used with corticosteroids when a more-rapid increase in platelet count is required. If IVIG is used, the dose should initially be 1 g/kg as a one-time dose. This dosage may be repeated if necessary. If previous treatment with corticosteroids and/or IVIG unsuccessful, subsequent treatment may include rituximab, thalidomide, cyclosporine, or more-potent immunosuppression (From American Society of Hematology guideline on immune thrombocytopenia97; consult for further details).
- G4: Hold ICPi

#### 8.7 Acquired hemophilia

**Definition:** Disorder caused by the development of autoantibodies (inhibitors) directed against plasma coagulation factors

**Diagnostic work-up**
- Full blood count to assess platelet number, fibrinogen, PT, PTT, INR; the typical finding in patients with acquired hemophilia A is a prolonged activated PTT with a normal PT
- MRI, CT, and ultrasonography may be indicated to localize, quantify, and serially monitor the location and response of bleeding
- Medication review to assess for alternative causes
- Determination of Bethesda unit level of inhibitor

(continued on following page)
Blood chemistry, CBC with evidence of anemia, macrocytosis, evidence of hemolysis on peripheral smear. LDH, haptoglobin, bilirubin, reticulocyte count, free hemoglobin.

Disseminated intravascular coagulation panel, which could include Prothrombin Time and International Normalized Ratio (PT/INR), infectious causes.

Autoimmune serology.

Paroxysmal nocturnal hemoglobinuria screening.

Direct and indirect bilirubin; LDH; direct agglutinin test; and if no obvious cause, bone marrow analysis, cytogenetic analysis to evaluate for myelodysplastic syndromes.

Evaluation for viral/bacterial (mycoplasma, etc) causes of hemolysis studies.

Protein electrophoresis, cryoglobulin analysis.

Work-up for bone marrow failure syndrome if refractory, including B12, folate, copper, parvovirus, iron, thyroid, infection.

Glucose-6-phosphate dehydrogenase.

Evaluation of common drug causes (ribavirin, rifampin, dapsone, interferon, cephalosporins, penicillins, NSAIDs, quinine/quinidine, fludarabine, ciprofloxacin, lorazepam, diclofenac, etc).

Assessment of methemoglobinemia.

**Recommendation 8.1b — Management.** It is recommended that clinicians manage all grade 1 toxicities as follows:

- Should continue to offer ICPI with close clinical follow-up and laboratory evaluation.

It is recommended that clinicians manage all grade 2 toxicities as follows:

- Should hold ICPI and strongly consider permanent discontinuation.
- Should administer 0.5 to 1 mg/kg/d prednisone equivalents. It is recommended that clinicians manage all grade 3 toxicities as follows:

- Should permanently discontinue ICPI.
- Should use clinical judgment and consider admitting the patient.
- Should consult hematology.
- Should administer prednisone 1 to 2 mg/kg/d (oral or IV depending on symptoms/speed of development).
- If worsening or no improvement, should administer 1 to 2 mg/kg/d prednisone equivalents and permanently discontinue ICPI treatment.
- May offer RBC transfusion per existing guidelines. Do not transfuse more than the minimum number of RBC units necessary to relieve symptoms of anemia or to return a patient to a safe hemoglobin range (7 to 8 g/dL in stable, noncardiac inpatients).
- Should offer patients supplementation with folic acid 1 mg once daily.

It is recommended that clinicians manage all grade 4 toxicities as follows:

- Should permanently discontinue ICPI.
- Should admit patient.
- Should consult hematology.
- Should administer IV prednisone 1 to 2 mg/kg/d.
- If no improvement or if worsening while on corticosteroids or severe symptoms on presentation, should initiate other...
• Should offer RBC transfusion per existing guidelines. Discuss with blood bank team prior to transfusions that a patient with a possible ICPI severe AE is in house.

8.2 Acquired Thrombotic Thrombocytopenic Purpura

Recommendation 8.2a – Diagnostic work-up. It is recommended that the diagnostic work-up should include the following:

• History with specific questions related to drug exposure (eg, chemotherapy, sirolimus, tacrolimus, opuna ER antibiotics, quinine).
• Physical examination, peripheral smear.
• ADAMTS13 activity level and inhibitor titer.
• LDH, haptoglobin, reticulocyte count, bilirubin.
• Prothrombin time (PT), activated partial thromboplastin time (APTT), fibrinogen.
• Blood group and antibody screen, direct antiglobulin test, CMV serology.
• Consider CT scan/MRI of brain, echocardiogram, ECG.
• Viral studies.
• Note: This disorder is usually associated with a severe drop in platelets and hemolysis/anemia precipitously.

Recommendation 8.2b – Management. It is recommended that clinicians manage all-grade toxicities as follows:

• The first step in the management of TTP is a high index of suspicion for the diagnosis and timely recognition. Hematology consult should immediately be called, as delay in identification is associated with increased mortality/morbidity.
• Initially, the patient should be stabilized and any critical organ dysfunction stabilized.

It is recommended that clinicians manage grade 1 to 2 toxicities as follows:

• Should hold ICPI and discuss resumption with patient only after taking into account the risks and benefits, noting that currently, there are no data to recommend restarting ICPI therapy.
• Should consult hematology.
• Should administer 0.5 to 1 mg/kg/d prednisone.

It is recommended that clinicians manage grade 3 to 4 toxicities as follows:

• Should hold ICPI and discuss resumption with patient only after taking into account the risks and benefits, noting that currently there are no data to recommend restarting ICPI therapy.
• Should consult hematology.
• In conjunction with hematology, should initiate PEX according to existing guidelines, with further PEX dependent on clinical progress.
• Should administer methylprednisolone 1 g IV daily for 3 days, with the first dose typically administered immediately after the first PEX.
• May offer rituximab.

8.3 Hemolytic Uremic Syndrome

Recommendation 8.3a – Diagnostic work-up. It is recommended that the diagnostic work-up should include the following:

• History and physical examination (special consideration for new history of high-risk drugs, hypertension, or cardiac causes).
• CBC with indices.
• Blood smear morphology. Note that the presence of schistocytes on smear is critical for diagnosis.
• Serum creatinine.
• ADAMTS13 (to rule out TTP).
• Homocysteine/methyalomalonic acid.
• Complement testing C3, C4, CH50 (complement inhibitory antibodies for suspected familial).
• Evaluate reticulocyte count and mean corpuscular volume.
• Evaluation of infectious cause, including screening for Epstein-Barr virus (EBV), CMV, human herpesvirus 6.
• Evaluation for nutritional causes of macrocytosis (B12 and folate).
• Pancreatic enzymes.
• Evaluation for diarrheal causes, shiga toxin, Escherichia coli 0157, etc
• Direct antibody test (Coombs test), haptoglobin, LDH, and other etiologies of anemia.
• Evaluation for common drugs causing hemolysis (tacrolimus, cyclosporine, sirolimus, etc).
• Evaluation for concurrent confusion.

Recommendation 8.3b – Management. It is recommended that clinicians manage grade 1 to 2 toxicities as follows:

• Should continue to offer ICPI with close clinical follow-up and laboratory evaluation.
• Should offer supportive care.

It is recommended that clinicians manage grade 3 to 4 toxicities as follows:

• Permanently discontinue ICPI.
• Begin eculizumab therapy 900 mg weekly for four doses, 1,200 mg on week 5, then 1,200 mg every 2 weeks.
• Red blood transfusion according to existing guidelines.

8.4 Aplastic Anemia

Recommendation 8.4a – Diagnostic work-up. It is recommended that the diagnostic work-up should include the following:

• History and physical examination (close attention to medications, exposure to radiation, toxins, recent viral infections).
• CBC, smear, reticulocyte count.
• Viral studies, including CMV, human herpesvirus 6, EBV, parvovirus.
• Nutritional assessments including B12, folate, iron, copper, ceruloplasmin, vitamin D.
• Serum LDH, renal function.
• Work-up for infectious causes.
• Identify marrow hypo/aplasia.
• Bone marrow biopsy and aspirate analysis.
• Peripheral blood analysis, including neutrophil count, proportion of glycosylphosphatidylinositol-negative cells.
• Flow cytometry to evaluate loss of glycosylphosphatidylinositol-anchored proteins.
• Type and screen patient for transfusions and notify blood bank that all transfusions need to be irradiated and filtered.
Recommendation 8.4b – Management. It is recommended that clinicians manage grade 1 toxicities as follows:

- Should hold ICPI and provide growth factor support, close clinical follow-up, and laboratory evaluation.
- Supportive transfusions as per local guidelines.

It is recommended that clinicians manage grade 2 toxicities as follows:

- Should hold ICPI and provide growth factor support and close clinical laboratory evaluations daily.
- Should administer ATG + cyclosporine. HLA typing and evaluation for bone marrow transplantation if patient is candidate. All blood products should be irradiated and filtered.
- May also offer supportive care with granulocyte colony-stimulating factor.

It is recommended that clinicians manage grade 3 to 4 toxicities as follows:

- Should hold ICPI and monitor weekly for improvement. If not resolved, should discontinue treatment until AE has reverted to grade 1.
- Should consult hematology.
- Should offer horse ATG plus cyclosporine.
- If no response, should repeat immunosuppression with rabbit ATG plus cyclosporine, alemtuzumab.
- For refractory patients, may offer eltrombopag plus supportive care.

8.5 Lymphopenia

Recommendation 8.5a – Diagnostic work-up. It is recommended that the diagnostic work-up should include the following:

- History and physical examination (special attention for lymphocyte-depleting therapy such as fludarabine, ATG, corticosteroids, cytotoxic chemotherapy, radiation exposure, etc, as well as history of autoimmunity disease, family history of autoimmune disease).
- Evaluation of nutritional state as cause.
- Spleen size.
- CBC with differential and reticulocyte counts
- CXR for evaluation of presence of thymoma.
- Bacterial cultures and evaluation for infection (fungal, viral, bacterial, specifically CMV/HIV).

Recommendation 8.5b – Management. It is recommended that clinicians manage grade 1 to 2 toxicities as follows:

- Should continue to offer ICPI.
- It is recommended that clinicians manage grade 3 toxicities as follows:
  - Continue ICPI, checking CBC weekly for monitoring, initiation of CMV screening.
- It is recommended that clinicians manage grade 4 toxicities as follows:
  - Should hold ICPI and discuss resumption with patient only after taking into account the risks and benefits.
  - Initiate Mycobacterium avium complex prophylaxis and Pneumocystis jirovecii pneumonia prophylaxis, CMV screening. HIV/hepatitis screening if not already done.

- May consider EBV testing if evidence of lymphadenopathy/hepatitis, fevers, hemolysis consistent with lymphoproliferative disease.

8.6 Immune Thrombocytopenia

Recommendation 8.6a – Diagnostic work-up. It is recommended that the diagnostic work-up should include the following:

- History and physical examination (special attention for lymphocyte-depleting therapy, such as fludarabine, ATG, corticosteroids, cytotoxic therapy).
- Family history of autoimmunity or personal history of autoimmune disease.
- History of viral illness.
- CBC.
- Peripheral blood smear, reticulocyte count.
- Bone marrow evaluation only if abnormalities in the above test results and further investigation is necessary for a diagnosis.
- Patients with newly diagnosed immune thrombocytopenia should undergo testing for HIV, hepatitis C virus, hepatitis B virus, and Helicobacter pylori.
- Direct antigen test should be checked to rule out concurrent Evan syndrome.
- Nutritional evaluation.
- Bone marrow evaluation if other cell lines affected and concern for aplastic anemia.

Recommendation 8.6b – Management. It is recommended that clinicians manage grade 1 toxicities as follows:

- Should continue ICPI with close clinical follow-up and laboratory evaluation.

It is recommended that clinicians manage grade 2 toxicities as follows:

- Should hold ICPI but monitor for improvement. If not resolved, should interrupt treatment until AE has reverted to grade 1.
- Should administer prednisone 1 mg/kg/d (dosage range, 0.5 to 2 mg/kg/d) orally for 2 to 4 weeks after which time this medication should be tapered over 4 to 6 weeks to the lowest effective dose.
- IVIG may be used in conjunction with corticosteroids if a more-rapid increase in platelet count is required.

It is recommended that clinicians manage grade 3 to 4 toxicities as follows:

- Should hold ICPI but monitor for improvement. If not resolved, should interrupt treatment until AE has reverted to grade 1.
- Should consult hematology.
- Should administer prednisone 1 to 2 mg/kg/d (oral or IV depending on symptoms).
- If worsening or no improvement, should administer 1 to 2 mg/kg/d prednisone equivalents and permanently discontinue treatment.
- IVIG may be used with corticosteroids when a more-rapid increase in platelet count is required.
- If IVIG is used, the dose should initially be 1 g/kg as a one-time dose. This dosage may be repeated if necessary.
If previous treatment with corticosteroids and/or IVIG has been unsuccessful, subsequent treatment may include splenectomy, rituximab, thrombopoietin receptor agonists, or more-potent immunosuppression.

Adapted from the American Society of Hematology guideline on immune thrombocytopenia; consult for further details.

8.7 Acquired Hemophilia

Recommendation 8.7a – Diagnostic work-up. It is recommended that the diagnostic work-up should include the following:

- Full blood count to assess platelet number, fibrinogen, PT, PTT, international normalized ratio. The typical finding in patients with acquired hemophilia A is a prolonged activated PTT with a normal PT.
- MRI, CT, and ultrasonography may be indicated to localize, quantify, and serially monitor the location and response of bleeding.
- Medication review to assess for alternative causes.
- Determination of Bethesda unit level of inhibitor.

Recommendation 8.7b – Management. It is recommended that clinicians manage grade 1 toxicities as follows:

- Should hold ICPi and discuss resumption with patient only after taking into account the risks and benefits.
- Should administer 0.5 to 1 mg/kg/d prednisone.
- Transfusion support as required.
- May treat bleeding episodes in consultation with a hematologist and/or hemophilia center experienced in the treatment of inhibitors.

It is recommended that clinicians manage grade 2 toxicities as follows:

- Should hold ICPi and discuss resumption with patient only after taking into account the risks and benefits.
- Should consult hematology.
- Should administer 1 mg/kg/d prednisone ± rituximab (dose, 375 mg/m² weekly for 4 weeks) and/or cyclophosphamide (dose, 1 to 2 mg/kg/d). Choice of rituximab versus cyclophosphamide is patient specific and should be done with assistance of hematology consult. Prednisone, rituximab, and cyclophosphamide should be given for at least 5 weeks.

It is recommended that clinicians manage grade 3 to 4 toxicities as follows:

- Should permanently discontinue ICPi.
- Should admit patient.
- Should consult hematology.
- Administration of factor replacement, choice based on Bethesda unit level of inhibitor.
- Bypassing agents may be used (factor VII, factor VIII inhibitor bypass activity). Caution should be taken in the elderly and those with coronary artery disease.
- Prednisone 1 to 2 mg/kg/d (oral or IV depending on symptoms) ± rituximab (dose, 375 mg/m² weekly for 4 weeks) and/or cyclophosphamide (dose, 1 to 2 mg/kg/d).
- Transfusion support as required for bleeding.
- If worsening or no improvement, should add cyclosporine or immunosuppression/immunosorption.

Qualifying statement. Acquired hemophilia A requires specialist clinical and laboratory expertise. Consult and/or transfer to a specialist center is often appropriate. If consultation with or transfer to a hemophilia center is not immediately possible, then investigation and treatment should be initiated while a liaison is being established.

Discussion. Review of literature for hematologic toxicities of checkpoint inhibitors revealed evidence of toxicity but relatively little in the form of comprehensive evaluation. A recent review has incorporated a systematic review of all phase I to III prospective clinical trials for Food and Drug Administration–approved ICPis and collated the incidence of common toxicities (unpublished data, J. Holter Chakrabarty, 2017).

Anemia (grades 1 to 4) occurs in approximately 11% of patients, with grades 3/4 at approximately 5.4% (1.1% to 17%).

If anemia progresses to pancytopenia or multiple cell lines are affected, evaluation for pure red cell aplasia, autoantibodies, aplastic anemia, and myelodysplasia must be considered. Toxicities between checkpoint inhibitors appear relatively similar. The majority of patients respond to withdrawal and are managed successfully with corticosteroids, IVIG, and growth factor support. Hemolytic anemia has been described as having development of autoantibodies and can commonly be treated by withholding ICPi, corticosteroids, and IVIG.

Thrombocytopenia is also relatively uncommon, occurring in approximately 8% (1% to 28%) of patients for all grades and 4.3% (3% to 6%) for grades 3/4.

Evaluation for causes of thrombocytopenia must be undertaken, including evaluation of TTP, disseminated intravascular coagulation, myelodysplastic syndrome, as well as immune-mediated thrombocytopenia related to ICPi. Corticosteroids have been shown to be effective with transfusion support as required.

Factor-related acquired bleeding disorders have been described with factor VIII. Involvement of hematologic expertise should be considered, including evaluation for antibody titer formation and choice of factor replacement. At low titer levels, simple factor replacement and corticosteroids may be effective; however, at high Bethesda unit levels > 5, bypassing agents such as factor VIII inhibitor bypass activity or factor VII may be required. Care in elderly patients when using these agents should be considered.

In most cases of mild hematologic toxicities, ICPi can be safely continued. However, cases of more-severe hemolytic anemia, pure red cell anemia, aplastic anemia, severe thrombocytopenia, or coagulation factor deficiencies have been described. In these cases, corticosteroids should be started and supportive care measures instituted. Of note, lymphopenia is not an uncommon event, and the degree of lymphopenia should be assessed with CD4 count and appropriate prophylaxis/assessment started for Pneumocystis and CMV undertaken.

Checkpoint inhibitors have been used in both organ and hematopoietic stem-cell transplantation. In both, caution is advised, and immediate involvement with subspecialty care is advised secondary to increased toxicities that have been seen in these populations.

9.0 Cardiovascular Toxicities

Please refer to Table 9 for a complete set of recommendations, definition of grades, and additional considerations.
9.1 Myocarditis, Pericarditis, Arrhythmias, Impaired Ventricular Function With Heart Failure and Vasculitis

**Recommendation 9.1a – Diagnostic work-up.** It is recommended that the diagnostic work-up should include the following:

- **At baseline:**
  - ECG
  - Consider troponin, especially in patient treated with combination immune therapies
  - Upon signs/symptoms (consider cardiology consult):
    - ECG
    - Troponin
    - Brain natriuretic peptide (BNP)
    - Echocardiogram
    - CXR

  Additional testing to be guided by cardiology and may include:
  - Stress test
  - Cardiac catheterization
  - Cardiac MRI

**Recommendation 9.1b – Management.** It is recommended that clinicians manage all-grade toxicities as follows, as all grades warrant work-up and intervention given potential for cardiac compromise:

- Should hold ICPi and permanently discontinue after grade 1.
- Should administer high-dose corticosteroids (1 to 2 mg/kg of prednisone) initiated rapidly (oral or IV depending on symptoms).
- Should admit patient and consult cardiology.
- Should manage cardiac symptoms according to American College of Cardiology (ACC)/AHA guidelines and with guidance from cardiology.
- May offer immediate transfer to a coronary care unit for patients with elevated troponin or conduction abnormalities.
- In patients without an immediate response to high-dose corticosteroids, may offer early institution of cardiac transplant rejection doses of corticosteroids (methylprednisolone 1 g every day) and the addition of either mycophenolate, infliximab, or ATG.

**Qualifying statement.** Treatment recommendations are based on anecdotal evidence and the life-threatening nature of cardiovascular complications. Holding checkpoint inhibitor therapy is recommended for all grades of complications. The appropriateness of rechallenging remains unknown. Note that infliximab has been associated with heart failure and is contraindicated at high doses in patients with moderate-severe heart failure.

9.2 Venous Thromboembolism

**Recommendation 9.2a – Diagnostic work-up.** It is recommended that the diagnostic work-up should include the following:

- An evaluation of signs and symptoms of pulmonary embolism (PE) or deep vein thrombosis (DVT), which may include a clinical prediction rule to stratify patients with suspected venous thromboembolism, venous ultrasound for suspected DVT, and CT pulmonary angiography for suspected PE.
- May also offer d-dimer for low-risk patients based on risk stratification by clinical prediction rule for DVT/PE when CT or Doppler are not available or appropriate.
- Ventilation/perfusion scan is also an option when CT pulmonary angiography is not appropriate.
- May make use of other testing, including ECG, CXR, BNP and troponin levels, and arterial blood gas.

**Recommendation 9.2b – Management.** It is recommended that clinicians manage grade 1 toxicities as follows:

- Should continue to offer ICPi.
- Should offer warm compress.
- Should offer clinical surveillance.

It is recommended that clinicians manage grade 2 to 3 toxicities as follows:

- Should continue to offer ICPi.
- Should manage according to CHEST, ACC, and/or AHA guidelines and consider consult from cardiology or other relevant specialties.
- Low-molecular-weight heparin is suggested over vitamin K agonist, dabigatran, rivaroxaban, apixaban, or edoxaban for initial and long-term treatment.
- IV heparin is an acceptable alternative for initial use, and oral anticoagulants are acceptable for the long term.

It is recommended that clinicians manage grade 4 toxicities as follows:

- Should permanently discontinue ICPi.
- Should admit patient and manage according to CHEST, ACC, and/or AHA guidelines and with guidance from cardiology.
- Should seek respiratory and hemodynamic support.
- Low-molecular-weight heparin is suggested over vitamin K agonist, dabigatran, rivaroxaban, apixaban, or edoxaban for initial and long-term treatment.
- IV heparin is an acceptable alternative for initial use, and oral anticoagulants are acceptable for the long term.
- Should offer further clinical management as indicated based on symptoms.

**Qualifying statement.** While it may be impossible to determine the etiology of thromboembolic disease in patients with advanced cancer and the role, if any, that ICPi treatment plays, it is reasonable to remove the potential inciting agents given the severity and life-threatening potential of grade 4 complications. Clinicians are to use clinical judgment and take into account the risks and benefits when deciding whether to discontinue ICPi treatment. Anticoagulant therapy duration should continue for a minimum of 9 to 12 months to indefinitely in the setting of active cancer unless the patient is asymptomatic, doing well, or in remission.

**Discussion.** Cardiovascular complications of ICPi therapy are rare but potentially life-threatening and/or of devastating clinical consequences. They have been reported with all currently approved agents. However, due to their rarity and involvement of major organs leading to rapidly fatal consequences, data are sparse and generally have included case reports or small case series. Cardiovascular irAEs occur in < 0.1% of patients receiving these therapies based on a review of pharmaceutical safety databases. The risk may be increased when combination therapy is
### 9.0 Cardiovascular Toxicities

#### 9.1 Myocarditis, pericarditis, arrhythmias, impaired ventricular function with heart failure and vasculitis

**Definition:** Signs and symptoms may include chest pain, arrhythmia, palpitations, peripheral edema, progressive or acute dyspnea, pleural effusion, fatigue

**Diagnostic work-up**
- **At baseline**
  - ECG
  - Consider troponin, especially in patients treated with combination immune therapies
- **Upon signs/symptoms (consider cardiology consult)**
  - ECG
  - Troponin
  - BNP
  - Echocardiogram
  - CXR
- Additional testing to be guided by cardiology and may include
  - Stress test
  - Cardiac catheterization
  - Cardiac MRI

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1: Abnormal cardiac biomarker testing, including abnormal ECG</td>
<td>All grades warrant work-up and intervention given potential for cardiac compromise</td>
</tr>
<tr>
<td>G2: Abnormal screening tests with mild symptoms</td>
<td>Consider the following:</td>
</tr>
<tr>
<td>G3: Moderately abnormal testing or symptoms with mild activity</td>
<td>Hold ICPI and permanently discontinue after G1</td>
</tr>
<tr>
<td>G4: Moderate to severe decompensation, IV medication or intervention required, life-threatening conditions</td>
<td>High-dose corticosteroids (1-2 mg/kg of prednisone) initiated rapidly (oral or IV depending on symptoms)</td>
</tr>
<tr>
<td></td>
<td>Admit patient, cardiology consultation</td>
</tr>
<tr>
<td></td>
<td>Management of cardiac symptoms according to ACC/AHA guidelines and with guidance from cardiology</td>
</tr>
<tr>
<td></td>
<td>Immediate transfer to a coronary care unit for patients with elevated troponin or conduction abnormalities</td>
</tr>
<tr>
<td></td>
<td>In patients without an immediate response to high-dose corticosteroids, consider early institution of cardiac transplant rejection doses of corticosteroids (methylprednisolone 1 g every day) and the addition of either mycophenolate, infliximab, or antithymocyte globulin</td>
</tr>
</tbody>
</table>

**Qualifying statement:** Treatment recommendations are based on anecdotal evidence and the life-threatening nature of cardiovascular complications. Holding checkpoint inhibitor therapy is recommended for all grades of complications. The appropriateness of rechallenging remains unknown. Note that infliximab has been associated with heart failure and is contraindicated at high doses in patients with moderate-severe heart failure.108

#### 9.2 Venous thromboembolism

**Definition:** A disorder characterized by occlusion of a vessel by a thrombus that has migrated from a distal site via the blood stream. Clinical signs and symptoms are variable and may include pain, swelling, increased skin vein visibility, erythema, and cyanosis accompanied by unexplained fever for DVT and dyspnea, pleuritic pain, cough, wheezing, or hemopty of PE

**Diagnostic work-up**
- Evaluation of signs and symptoms of PE or DVT may include
  - Clinical prediction rule to stratify patients with suspected venous thromboembolism
  - Venous ultrasound for suspected DVT
  - CTPA for suspected PE
- Can also consider d-dimer for low-risk patients based on risk stratification by clinical prediction rule for DVT/PE when CT or Doppler are not available or appropriate
- Ventilation/perfusion scan is also an option when CTPA is not appropriate
- Consider other testing, including ECG, CXR, BNP and troponin levels, and arterial blood gas

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1: Venous thrombosis (eg, superficial thrombosis)</td>
<td>Continue ICPI</td>
</tr>
<tr>
<td></td>
<td>Warm compress</td>
</tr>
<tr>
<td></td>
<td>Clinical surveillance</td>
</tr>
<tr>
<td>G2: Venous thrombosis (eg, uncomplicated DVT), medical intervention indicated</td>
<td>Continue ICPI</td>
</tr>
<tr>
<td></td>
<td>Management according to CHEST, ACC, and/or AHA guidelines and consider consult from cardiology or other relevant specialists</td>
</tr>
<tr>
<td></td>
<td>LMWH is suggested over VKA, dabigatran, rivaroxaban, apixaban, or edoxaban for initial and long-term treatment</td>
</tr>
<tr>
<td></td>
<td>IV heparin is an acceptable alternative for initial use, and oral anticoagulants are acceptable for the long term</td>
</tr>
<tr>
<td>G3: Thrombosis (eg, uncomplicated PE [venous], nonembolic cardiac mural [arterial] thrombus), medical intervention indicated</td>
<td>Permanently discontinue ICPI</td>
</tr>
<tr>
<td></td>
<td>Admit patient and management according to CHEST, ACC, and/or AHA guidelines and with guidance from cardiology</td>
</tr>
<tr>
<td></td>
<td>Respiratory and hemodynamic support</td>
</tr>
<tr>
<td></td>
<td>LMWH is suggested over VKA, dabigatran, rivaroxaban, apixaban, or edoxaban for initial and long-term treatment</td>
</tr>
<tr>
<td></td>
<td>IV heparin is an acceptable alternative for initial use, and oral anticoagulants are acceptable for the long term</td>
</tr>
<tr>
<td></td>
<td>Further clinical management as indicated based on symptoms</td>
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</tbody>
</table>

(continued on following page)
used. In these safety data, combination therapy of ipilimumab and nivolumab had greater rates of cardiovascular complications than nivolumab alone (0.28% vs 0.06%).

Mortality is high, with death frequently secondary to refractory arrhythmia or cardiogenic shock.75,113,114

One review of compiled case reports and case series by Jain et al.111 found that the onset of cardiovascular irAEs can be as soon as 2 weeks and as long as 32 weeks after initiation of therapy, with a median onset of 10 weeks after initiation. Based on results of myocardial biopsies, these complications are thought to be caused by lymphocytic infiltration of the myocardium and myocardial conduction system.75 Pathology has shown lymphocytic infiltration in the tumor specimens.

A wide range of cardiovascular complications have been reported. Pathologic review shows occurrences of myocarditis; myocardial fibrosis; cardiomyopathy; heart failure; conduction abnormalities, including heart block; and cardiac arrest.111 Pericarditis and pericardial effusions have been described as well.85,115 There has also been a case report of irAE-associated acute coronary syndrome.116

Immune-mediated myocarditis may result in heart failure or arrhythmia. The myocarditis may be fulminant, progressive, and life-threatening.75,117 Acute heart failure may occur secondary to decreased cardiac function and diminished ejection fraction.75,114

Conduction abnormalities can include complete heart block75,114 and arrhythmias. A variety of dysrhythmias may occur from the more benign (supraventricular tachycardias) to more fatal and can lead to sudden death (ventricular tachycardias).75,76,112-114,118-120

Presentation of cardiovascular complications of checkpoint inhibitors could include arrhythmia, palpitations, chest pain, or signs and symptoms of heart failure (shortness of breath, peripheral edema, pleural effusion, fatigue). Severe cases can present with cardiogenic shock or sudden death. Patients can also present with fatigue, malaise, myalgia, and/or weakness alone or in combination with more-specific cardiovascular symptoms. Symptoms can often be masked by other irAEs (eg, pneumonitis, hypothyroidism) or symptoms related to disease (eg, pulmonary symptoms).

Initial evaluation of patients with potential cardiovascular toxicity should include ECG, troponin, BNP, and CXR. Reported cases have invariably had elevations of troponin, CK, and CK-MB.112 BNP will also be elevated in cases with decrease ejection fraction. Diagnostic evaluation should consider the possibility of other etiologies of the patient’s symptoms and could include, for example, cardiac stress testing, heart catheterization, or cardiac MRI. Due to the possibility of arrhythmia and progression to life-threatening arrhythmias or heart block, continuous telemetry monitoring should be instituted. Typically, many of these patients will often be admitted to an inpatient unit and worked up there given the severity of the symptoms. Patients with mild shortness of breath of unclear etiology should get typical outpatient testing (ECG, BNP, troponin).

Echocardiogram to evaluate for cardiac function should be performed in symptomatic patients. Echocardiogram may reveal decreased left or right ventricular ejection fraction (with global or regional abnormalities). Cardiac MRI can demonstrate evidence of myocarditis but is less sensitive than endomyocardial biopsy.112,117 Endomyocardial biopsy should be considered for patients who are unstable or failed to respond to initial therapy or in whom the diagnosis is in doubt. Typically, initial diagnostic testing reveals issues, and treatment is often administered empirically before confirmatory pathologic testing is obtained.

There is no clear evidence regarding the efficacy or value of routine baseline or serial ECGs or troponin measurements in patients receiving checkpoint inhibitor therapy. Some centers obtain baseline testing, and others continue this through the initial period of therapy. Some centers stratify management based on magnitude of troponin changes.112 Baseline information can potentially be useful when patients present acutely with nonspecific symptoms and have equivocal diagnostic testing.

Treatment recommendations are based on anecdotal evidence and the life-threatening nature of cardiovascular complications of irAEs due to either malignant arrhythmia or the possibility of fulminant myocarditis with heart failure. Holding checkpoint inhibitor therapy is recommended for all grades of complications, including grade 1 (asymptomatic biomarker elevations), with reinitiation of treatment almost never happening.111,112

For patients with mild to moderate symptoms (grades 2 to 3), systemic prednisone or methylprednisolone is indicated at 1 to 2 mg/kg/day.6,75,113 Those with more severe disease (grades 3 to 4), including clinical decompensation, highly abnormal testing, fulminant disease, cardiogenic shock, and acute heart failure, or with life-threatening arrhythmia should be considered for more-aggressive therapy, as should those who fail to respond to initial corticosteroid dosing within 3 to 5 days. This could include therapy with higher doses of corticosteroids (methylprednisolone at 1 g daily) and the possible addition of mycophenolate, infliximab, or ATG.75,111-113,117 Management of symptoms of arrhythmia and

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**Table 9. Management of Cardiovascular irAEs in Patients Treated With ICPIs (continued)**

<table>
<thead>
<tr>
<th>Cardiovascular Toxicities</th>
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</thead>
<tbody>
<tr>
<td>9.0 Cardiovascular Toxicities</td>
</tr>
</tbody>
</table>

**Additional considerations**

While it may be impossible to determine the etiology of thromboembolic disease in patients with advanced cancer and the role, if any, that ICPI treatment plays, it is reasonable to remove the potential inciting agents given the severity and life-threatening potential of G4 complications. Clinicians are to use clinical judgment and take into account the risks and benefits when deciding whether to discontinue ICPI treatment.

Anticoagulant therapy duration should continue for a minimum of 9-12 months to indefinitely in the setting of active cancer unless patient is asymptomatic, doing well, or in remission.109,112

All recommendations are expert consensus based, with benefits outweighing harms, and strength of recommendations are moderate.

**Abbreviations:** ACC, American College of Cardiology; AHA, American Heart Association; BNP, brain natriuretic peptide; CT, computed tomography; CTPA, computed tomography pulmonary angiography; CXR, chest x-ray; DVT, deep vein thrombosis; ICPI, immune checkpoint inhibitor; irAE, immune-related adverse event; IV, intravenous; LRWH, low-molecular-weight heparin; MRI, magnetic resonance imaging; PE, pulmonary embolism; VKA, vitamin K antagonist.
heart failure should be as per national cardiology guidelines and clinical judgment.\textsuperscript{112}

Although some diseases are fulminant and progress to death, with appropriate therapy and holding of checkpoint inhibitors, cardiac contractility and conduction abnormalities can improve.\textsuperscript{119} There have not been sufficient cases in the literature to determine the proportion expected to progress or improve. Given the potential severity of the symptoms, the patient’s disease status must be taken into account before excessive support measures are performed (eg, defibrillator, resuscitation, balloon pump).

The evidence on how to distinguish among risk factors in patients with cancer treated with ICPi therapy is limited. Furthermore, determining the true cause of thromboembolic disease in such patients is difficult, if not impossible, given the thrombogenicity of both the disease and the treatment. Treating physicians are urged to use clinical judgment in the management of these patients.

10.0 Ocular Toxicities

Please refer to Table 10 for a complete set of recommendations, definition of grades, and additional considerations.

Recommendation 10.0 – Diagnostic work-up for all ocular toxicities. It is recommended that clinicians counsel all patients to inform their health care provider immediately if they experience any of the following ocular symptoms:

- Blurred vision
- Change in color vision
- Photophobia
- Distortion
- Scotomas
- Visual field changes
- Double vision
- Tenderness
- Pain with eye movement
- Eyelid swelling
- Proptosis

It is recommended that the diagnostic work-up should include the following, under the guidance of ophthalmology:

- Check vision in each eye separately
- Color vision
- Red reflex
- Pupil size, shape, and reactivity
- Fundoscopic examination
- Inspection of anterior part of eye with penlight

Qualifying statement. Clinicians should be aware that ocular irAEs are many times seen in the context of other organ irAEs, and there should be a high level of clinical suspicion as symptoms may not always be associated with severity. It is best to treat ocular irAEs after ophthalmologist eye examination.

10.1 Uveitis/Iritis

Recommendation 10.1 – Management. It is recommended that clinicians manage grade 1 toxicities as follows:

- Should continue to offer ICPi.
- Should refer to ophthalmology within 1 week.
- Should offer artificial tears.

It is recommended that clinicians manage grade 2 toxicities as follows:

- Should hold ICPi temporarily until after ophthalmology consult.
- Should make an urgent ophthalmology referral.
- Should administer topical corticosteroids, cycloplegic agents, systemic corticosteroids.
- May resume ICPi treatment once off systemic corticosteroids, which are purely indicated for ocular adverse effects or once corticosteroids for other concurrent systemic irAEs are reduced to $\leq 10$ mg. Continued topical/ocular corticosteroids are permitted when resuming therapy to manage and minimize local toxicity. Should re-treat after return to grade 1 or less.

It is recommended that clinicians manage grade 3 toxicities as follows:

- Should permanently discontinue ICPi.
- Should make an urgent ophthalmology referral.
- Should administer systemic corticosteroids and intravitreal/periocular/topical corticosteroids.

It is recommended that clinicians manage grade 4 toxicities as follows:

- Should permanently discontinue ICPi.
- Should make an emergent ophthalmology referral.
- Should administer systemic corticosteroids and intravitreal/perioocular/topical corticosteroids.

10.2 Episcleritis

Recommendation 10.2 – Management. It is recommended that clinicians manage grade 1 toxicities as follows:

- Should continue to offer ICPi.
- Should refer to ophthalmology within 1 week.
- Should offer artificial tears.

It is recommended that clinicians manage grade 2 toxicities as follows:

- Should hold ICPi until after ophthalmology consult.
- Should make an urgent ophthalmology referral.
- Should administer topical corticosteroids, cycloplegic agents, systemic corticosteroids.

It is recommended that clinicians manage grade 3 toxicities as follows:

- Should permanently discontinue ICPi.
- Should make an urgent ophthalmology referral.
- Should administer systemic corticosteroids and topical corticosteroids with cycloplegic agents.

It is recommended that clinicians manage grade 4 toxicities as follows:

- Should permanently discontinue ICPi.
- Should make an emergent ophthalmology referral.
- Should administer systemic corticosteroids and topical corticosteroids with cycloplegic agents.
Table 10. Management of Ocular irAEs in Patients Treated With ICPIs

10.0 Ocular Toxicities

Counsel all patients to inform their health care provider immediately if they experience any of the following ocular symptoms:
- Blurred vision
- Change in color vision
- Photophobia
- Distortion
- Scotomas
- Visual field changes
- Double vision
- Tenderness
- Pain with eye movement
- Eyelid swelling
- Proptosis

Evaluation, under the guidance of ophthalmology:
- Check vision in each eye separately
- Color vision
- Red reflex
- Pupil size, shape, and reactivity
- Fundoscopic examination
- Inspection of anterior part of eye with penlight

Prior conditions:
- Exclude patients with history of active uveitis
- History of recurrent uveitis requiring systemic immunosuppression or continuous local therapy

Additional considerations:
- Ocular irAEs are many times seen in the context of other organ irAEs
- High level of clinical suspicion as symptoms may not always be associated with severity
- Best to treat after ophthalmologist eye examination

10.1 Uveitis/iritis

Definition: Inflammation of the middle layer of the eye

Diagnostic work-up: as per above

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1: Asymptomatic</td>
<td>Continue ICPI. Refer to ophthalmology within 1 week. Artificial tears.</td>
</tr>
<tr>
<td>G2: Medical intervention required, anterior uveitis</td>
<td>Hold ICPI temporarily until after ophthalmology consult. Topical corticosteroids, cycloplegic agents, systemic corticosteroids. May resume ICPI treatment once off systemic corticosteroids, which are purely indicated for ocular adverse effects or once corticosteroids for other concurrent systemic irAEs are reduced to ≤ 10 mg; continued topical/ocular corticosteroids are permitted when resuming therapy to manage and minimize local toxicity. Re-treat after return to G1 or less.</td>
</tr>
<tr>
<td>G3: Posterior or panuveitis</td>
<td>Permanently discontinue ICPI. Urgent ophthalmology referral. Systemic corticosteroids and intravitreal/pericocular/topical corticosteroids.</td>
</tr>
<tr>
<td>G4: 20/200 or worse</td>
<td>Permanently discontinue ICPI. Emergent ophthalmology referral. Systemic corticosteroids (IV prednisone 1-2 mg/kg or methylprednisolone 0.8-1.6 mg/kg) and intravitreal/pericocular/topical corticosteroids per ophthalmologist opinion.</td>
</tr>
</tbody>
</table>

Additional considerations: Consider use of infliximab or other TNF-α blockers in cases that are severe and refractory to standard treatment.

10.2 Episcleritis

Definition: Inflammatory condition affecting the episcleral tissue between the conjunctiva and the sclera that occurs in the absence of an infection

Diagnostic work-up: As per 10.0

<table>
<thead>
<tr>
<th>Grading</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1: Asymptomatic</td>
<td>Continue ICPI. Refer to ophthalmology within 1 week. Artificial tears.</td>
</tr>
<tr>
<td>G2: Vision 20/40 or better</td>
<td>Hold ICPI therapy temporarily until after ophthalmology consult. Urgent ophthalmology referral. Topical corticosteroids, cycloplegic agents, systemic corticosteroids.</td>
</tr>
<tr>
<td>G3: Symptomatic and vision worse than 2/40</td>
<td>Permanently discontinue ICPI. Urgent ophthalmology referral. Systemic corticosteroids and topical corticosteroids with cycloplegic agents.</td>
</tr>
<tr>
<td>G4: 20/200 or worse</td>
<td>Permanently discontinue ICPI. Emergent ophthalmology referral. Systemic corticosteroids and topical corticosteroids with cycloplegic agents.</td>
</tr>
</tbody>
</table>

Additional considerations: Consider use of infliximab or other TNF-α blockers in cases that are severe and refractory to standard treatment.
10.0 Ocular Toxicities

10.3 Blepharitis

**Recommendation 10.3 – Management.** It is recommended that clinicians manage all-grade toxicities as follows:

- Should offer warm compresses and lubricant drops.
- Should continue to offer ICPi unless irAE is persistent and serious.

**Discussion.** In the context of irAEs as a result of ICPi therapy in cancer, ocular toxicities are considered uncommon and less complex in their management compared with other immune-related toxicities.

A variety of ocular events have been reported with CTLA-4, anti–PD-1, and anti–PD-L1–inhibiting agents, including uveitis, iritis, episcleritis, and blepharitis. Like in other irAEs, the principal mechanism of the encountered toxicity is inflammatory and the principal management is immunosuppression with corticosteroids.

The overall incidence of uveitis with ICPis includes ipilimumab, anti–PD-1 antibodies, and anti–PD-L1 agents is up to 1%, although the incidence may be higher in patients receiving combination ICPis. Presenting symptoms include blurred vision and photophobia, change in color vision, and distortion as well as physical signs like tenderness, swelling, and pain with eye movement, among others. The practitioner should be aware that symptoms of uveitis may not indicate severity of the syndrome and thus should seek consultation with ophthalmology and slit lamp examination. Rarely, a panuveitis is induced that can lead to exudative retinal detachment and can be vision threatening. Milder forms of uveitis respond to temporary holding of ICPis and topical corticosteroids, and any symptomatic presentation should prompt urgent ophthalmology evaluation. Typical management includes topical corticosteroids and often the addition of cycloplegic agents, and in rare cases, systemic corticosteroid administration is necessary.

Episcleritis is a rare, but clinically important event, occurring in < 1% of treated patients. The management is similar to the one recommended for uveitis, and any visual compromise (vision < 20/40) should prompt urgent ophthalmology referral to assess the need for more-specific interventions. We recommend ophthalmology referral for all cases of episcleritis even if asymptomatic, and holding immune checkpoint therapy until such evaluation is completed. Artificial tears, topical corticosteroids, and cycloplegic agents are typically used and highly effective in managing this toxicity, but in rare cases, systemic corticosteroids may be required. In case of recurrent events or a grade 4 presentation (vision 20/200 or worse), permanent discontinuation of ICPi is advised. Infliximab may be considered for severe and treatment-refractory cases, although the data on this intervention rely on case reports only.

Blepharitis is equally rare as other ocular toxicities and is encountered < 1% of patients treated with ICPis. This toxicity is managed with warm compresses and artificial tears for lubrication. Disruption of therapy is not typically necessary but may be advised by the consulting ophthalmologist if symptoms are severe and treatment refractory.

In most cases of ocular toxicities, ICPi can be safely continued as most presenting grades are mild and manageable with topical corticosteroids. However, ocular toxicity is commonly associated with other systemic immune-related events, and systemic corticosteroids are often used in these patients to manage the more-prominent toxicities outside the eye. Modification and possible cessation of ICPi may need to be considered in these cases as well as in cases of higher grade, treatment-refractory, or recurrent ocular toxicity.

**DISCUSSION**

While identifying patients at an increased risk for irAEs would help to determine the need for surveillance and prompt, aggressive treatment, the evidence of who is at an elevated risk remains unclear. Patients with preexisting autoimmune diseases, such as ulcerative colitis, Crohn disease, lupus, and active rheumatoid arthritis, are usually not offered therapy with checkpoint inhibitors and typically have been excluded from clinical trials involving these agents. However, data suggest that they may be safely treated. Indeed, a systematic review of case reports of patients with pre-existing autoimmune diseases treated with ICPis found that 40% of patients did not experience an irAE or exacerbation of their autoimmune disease, despite many having active disease. Ultimately, cautious use of ICPi therapy may be acceptable with close monitoring for recurrence of the underlying autoimmune condition.

The pattern of toxicity based on tumor type and location has not been well established. Some reports have claimed higher incidences of pneumonitis in patients with NSCLC compared with melanoma, but other analyses found no statistically significantly differential effects according to cancer type. Treatment-naive
patients are reported to have a higher incidence of pneumonitis compared with those previously treated. \(^{132}\) Other evidence is also emerging on patient-related modifiers of risk. Personal ecologic factors, such as the patient’s microbiome, may also play a role in the susceptibility to specific irAEs, such as enterocolitis. \(^{27,135,136}\) Further studies are needed to investigate whether a patient’s biologic profile predisposes to the occurrence of irAEs. \(^{137}\)

Possible treatment-specific risks for increased irAEs include dose of therapy, individual checkpoint inhibitor (CTLA-4 \(^{n}\) PD-1), and combination checkpoint blockade. Model-based pooled estimates from 498 trial patients who received ipilimumab monotherapy at 0.3, 3, or 10 mg/kg doses indicated that higher doses produce higher rates of irAEs. \(^{138}\) Grade 3 or higher irAEs are reported to occur more frequently in patients receiving anti–CTLA-4 monotherapy (ipilimumab, 15% to 42%) than in those receiving anti–PD-1 (nivolumab, 8%; pembrolizumab, 5% to 10%) or anti–PD-L1 (atezolizumab, 5% to 7%; durvalumab, 2%; avelumab, 1% to 2%) monotherapy. \(^{139}\) Evidence also exists for the elevated risk with combination therapy. A recent meta-analysis revealed the OR of all-grade pneumonitis was 3.7 (95% CI, 1.6 to 8.5; \(P = .002\)), with an anti–CTLA-4 and anti–PD-1 therapy combination (ipilimumab and nivolumab) versus anti–CTLA-4 monotherapy. \(^{140}\) Combination anti–CTLA-4 and anti–PD-1 therapy also significantly increased the risk of grade 3 and 4 rash and fatigue. \(^{140-142}\) As the use of ICPi therapy increases and incidences of irAEs are further collected, the understanding of which patient is at an elevated risk is sure to become clearer. In the meantime, clinicians should maintain a high level of suspicion for immune-related toxicities with checkpoint inhibitors, with early recognition and treatment of upmost importance in mitigating the severity of irAEs. \(^{143}\)

While treatment with ICPIs is sometimes well tolerated, the potential for life-disabling irAEs that are severe and/or irreversible exists. \(^{137}\) A recent meta-analysis of approximately 6,000 patients with solid tumors reported a statistically significant increased risk of fatal AE for patients treated with ipilimumab (pooled Peto OR, 2.3; 95% CI, 1.4 to 3.6; \(P < .001\)). \(^{144}\) Among the specific causes of fatal AEs, ipilimumab was associated with an increased risk of fatal GI toxicity (OR, 4.5; 95% CI, 1.5 to 13.6). \(^{144}\)

The decision to resume ICPI therapy after resolution of toxicity is complicated because the optimal duration of ICPI therapy is not defined. Early trials of ICPI used 1 year of therapy; later trials used 2 years of therapy or continued ICPI treatment until disease progression or patient intolerance. Recent evidence suggests that patients who discontinued induction immunotherapy due to AEs did just as well as those who continued treatment uninterrupted. \(^{145}\) In a pooled analysis of randomized trials of patients with advanced melanoma who received nivolumab plus ipilimumab combination therapy, Schadendorf et al \(^{145}\) found an ORR of approximately 60% in patients who discontinued compared with approximately 50% in those who completed induction therapy. Progression-free survival was also similar between the two groups. While these data are intriguing, prospective evidence is still required to gain a better understanding of the merits, liability, and optimal duration of ongoing anti–PD-1 therapy after discontinuing induction therapy due to irAEs. \(^{146}\) A patient’s tumor response status is an important factor in deciding whether to resume ICPI. If a patient has achieved objective response to initial ICPI, there is a reasonable likelihood that the response will be durable and that resumption of therapy (with attendant risk of recurrence of toxicity) may not be advisable. Conversely, for patients who have not yet responded or whose response is deemed inadequate, consideration of resumption of ICPI therapy after resolution of toxicity is reasonable.

Whether the appearance of irAEs is associated with efficacy parameters still remains unclear. \(^{147}\) After adjusting for differences in number of nivolumab doses received, baseline LDH, and tumor PD-L1 expression, one analysis found that the ORR was significantly better in patients who experienced irAEs of any grade compared with those who did not, with the greatest benefit seen in patients who reported three or more irAEs. \(^{148}\) No significant difference in ORR on the basis of the occurrence of grade 3 to 4 irAEs was observed. \(^{148}\)

There are important studies under way that are evaluating the efficacy of various strategies in mitigating toxicities while maintaining efficacy, such as alternative dosing strategies or increasing the interval between treatment infusions. \(^{146}\) Until such evidence becomes available, dose reductions of ICPI therapy should be avoided. Rather, therapeutic adjustments by way of temporary interruption or permanent discontinuation of treatment are recommended.

Guidance on the management of toxicities related to ICPI therapy is in demand. This guideline and its recommendations is intended to arm the clinician with strategies and best practices to rapidly recognize, diagnose, coordinate with other medical subspecialties, and manage these sets of unique toxicities.

### PATIENT AND CLINICIAN COMMUNICATION

As immunotherapeutic treatment of cancer continues to evolve with single agents and in new combinations, it is imperative that patients and family caregivers receive timely and up-to-date education about immunotherapies, their mechanism of action, and the clinical profile of possible irAEs. Patient and caregiver education should occur prior to initiating therapy and continue throughout treatment and survivorship. It should be emphasized that immunotherapy works differently than traditional chemotherapy and that these treatments elicit unique therapeutic responses and corresponding irAEs. \(^{149}\) This response can be unique to each patient, and irAEs may occur across the treatment trajectory from the start of treatment and into survivorship. Most notably, the ability to influence immune response even after discontinuation of the immunotherapeutic agent is a unique feature, and important education point for patients and their caregivers. As such, patients should be encouraged to alert all health care providers that they are receiving or have received an immunotherapeutic agent and to report any changes in health status to each provider. This is important as patients are often seen by multiple providers, and each provider should be aware of the potential for irAEs.

In most cases, irAEs can be managed with treatment interruption and/or supportive care and for some patients, will involve a multidisciplinary team (eg, endocrinologist, pulmonologist, gastroenterologist) to address specific symptoms. \(^{150}\) Patients and caregivers need to know that AEs can often be managed effectively, especially when they are identified early. In addition, education addressing the safe handling
Management of Immune-Related Adverse Events

of medications, infection control, and safe sexual practices is important to supporting optimal management of irAEs.149

Using a questionnaire or standard assessment may assist the provider and patient to recognize possible irAEs. In addition, health care professionals should ask patients about any new symptoms or changes in their health, no matter how small they may seem. Minor changes in how a patient is feeling may indicate early signs of an AE, and patients may not attribute the change to their cancer treatment.152 Consistent assessment and documentation at each encounter will also enable the clinical team to note changes that may occur over time. Close monitoring throughout treatment is important as minimal changes in a patient’s baseline status may indicate an early irAE. Wallet cards detailing symptoms to watch for and how to notify their health care provider may be an effective tool in empowering patients and their caregivers to recognize and manage irAEs and may be useful to other health care providers (eg, emergency department staff) caring for patients with a history of immunotherapy.150 The Oncology Nursing Society has an immunotherapy wallet card available for patients and providers (Fig 1). Copies of the card or additional information can be obtained by e-mail at clinical@ons.org.

For recommendations and strategies to optimize patient-clinician communication, see Patient-Clinician Communication: American Society of Clinical Oncology Consensus Guideline.152

HEALTH DISPARITIES

Although ASCO clinical practice guidelines represent expert recommendations on the best practices in disease management to provide the highest level of cancer care, it is important to note that many patients have limited access to medical care. Racial and ethnic disparities in health care contribute significantly to this problem in the United States. Patients with cancer who are members of racial/ethnic minorities suffer disproportionately from comorbidities, experience more substantial obstacles to receiving care, are more likely to be uninsured, and are at greater risk of receiving care of poor quality than other Americans.153-156 Many other patients lack access to care because of their geographic location and distance from appropriate treatment facilities. Moreover, with evidence to suggest that patients with a higher mutation burden are at an increased likelihood of responding to IPCs,157-158 African American patients with lung cancer may be affected as the burden of somatic mutations appears to be different in such patients.157 Awareness of these disparities in access to care should be considered in the context of this clinical practice guideline, and health care providers should strive to deliver the highest level of cancer care to these vulnerable populations.

While concerns about racial disparities in access to trials of new cancer drugs have been raised, including trials of anti–PD-1, whether these disparities extend to patients in real-world practice has only recently been investigated.159 In a retrospective analysis of electronic health records of 4,643 patients treated with anti–PD-1, investigators found that racial distributions differed for anti–PD-1–treated patients compared with non–anti–PD-1–treated patients in a cohort of patients with advanced NSCLC (P < .01) but not in a cohort of patients with metastatic renal cell carcinoma (P = .84) or advanced melanoma (P = .96). In bivariate analyses of patients with advanced NSCLC, the use of anti–PD-1 treatment was associated with race, male sex, stage II at diagnosis, squamous histology, smoking history, and line of therapy (all P < .05).159 Adjusted models showed that there were no significant differences in likelihood of receiving anti–PD-1s when comparing black and white patients undergoing systemic therapy for NSCLC.159

IMMUNOTHERAPY WALLET CARD

NAME: __________________________
CANCER DX: __________________________
I-O AGENTS RCVD: ☐ CHECKPOINT INHIBITOR(S)
☐ CAR-T ☐ VACCINES ☐ ONCOLYTIC VIRAL THERAPY
☐ MONOCLONAL ANTIBODIES
DRUG NAME(S): __________________________
immunotherapy TX START DATE: __________________________
OTHER CANCER MEDICATIONS: __________________________

NOTE: IMMUNOTHERAPY AGENTS ARE NOT CHEMOTHERAPY AND SIDE EFFECTS MUST BE MANAGED DIFFERENTLY (SEE BACK)

IMMUNE-MEDIATED SIDE EFFECTS*, COMMON WITH CHECKPOINT INHIBITORs VARY IN SEVERITY AND MAY REQUIRE REFERRAL AND STEROIDS. PATIENTS HAVE A LIFETIME RISK OF IMMUNE-RELATED SIDE EFFECTS.

*May present at all sites: diarrhea, abnormal liver function test, dermatitis, fatigue, headache, kidney problems, liver problems, low blood pressure, low platelets, low red blood cells, low white blood cells, muscle pain, nausea, infection, pancreatitis, kidney problems, rash, cough, skin rash, fever, fatigue, hair loss, shortness of breath, taste changes, trouble swallowing, blisters, mouth sores, ulcers, itching, joint pain, eye problems, and may involve other body systems. Patients have a lifetime risk of immune-related side effects.

ONCOLOGY PROVIDER NAME __________________________
ONCOLOGY PROVIDER NO. __________________________
EMERGENCY CONTACT __________________________
CONTACT PHONE NO. __________________________

MULTIPLE CHRONIC CONDITIONS

Creating evidence-based recommendations to inform treatment of patients with additional chronic conditions, a situation in which the patient may have two or more such conditions—referred to as multiple chronic conditions (MCC)—is challenging. Patients with MCC are a complex and heterogeneous population, making it difficult to account for all the possible permutations to develop specific recommendations for care. In addition, the best available evidence for treating index conditions, such as cancer, is often from clinical trials whose study selection criteria may exclude patients with MCC, such as preexisting autoimmune diseases, to avoid potential interaction effects or confounding of results associated with MCC. As a result, the reliability of outcome data from these
studies may be limited, thereby creating constraints for expert groups to make recommendations for care in this heterogeneous patient population.

As many patients for whom guideline recommendations apply present with MCCs, including preexisting autoimmune diseases, any treatment plan needs to take into account the complexity and uncertainty created by the presence of MCCs and highlights the importance of shared decision making regarding guideline use and implementation. Therefore, in consideration of recommended care for the target index condition, clinicians should review all other chronic conditions present in the patient and take those conditions into account when formulating the management and follow-up plan.

In light of the above considerations, practice guidelines should provide information on how to apply the recommendations for patients with MCCs, perhaps as a qualifying statement for recommended care. This may mean that some or all of the recommended care options are modified or not applied, as determined by best practice in consideration of any MCCs.

**REFERENCES**


**AUTHORS’ DISCLOSURES OF POTENTIAL CONFLICTS OF INTEREST**

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**AUTHOR CONTRIBUTIONS**

Manuscript writing: All authors
Final approval of manuscript: All authors

**ADDITIONAL RESOURCES**

More information, including a Data Supplement with additional evidence tables, a Methodology Supplement with information about evidence quality and strength of recommendations, slide sets, and clinical tools and resources, is available at www.asco.org/supportive-care-guidelines and www.asco.org/guidelineswiki. Patient information is available at www.cancer.net. Visit www.asco.org/guidelineswiki to provide comments on the guideline or to submit new evidence.
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Management of Immune-Related Adverse Events in Patients Treated with Immune Checkpoint Inhibitor Therapy: American Society of Clinical Oncology Clinical Practice Guideline

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All Appendices and Acknowledgment material (including the table of Expert Panel members) are online only. It will appear on the JCO Web site but not in the print version.

Appendix

Fig A1. Distribution of (A) grade 1 to 2 and (B) grade 3 to 5 immune-related adverse events (irAEs) for all tumor types in the main clinical trials with anti–cytotoxic T-cell lymphocyte-4 (anti–CTLA-4), anti–programmed death 1 (PD-1), or anti–PD ligand 1 (PD-L1) antibodies as single therapies. The values quoted are the median (range) irAE rates for the set of clinical trials as a whole. Adapted from European Journal of Cancer, Vol 54, J.M. Michot et al, Immune-Related Adverse Events With Immune Checkpoint Blockade: A Comprehensive Review, 139-149, Copyright 2016, with permission from Elsevier. Endoc, endocrinology; Neurol, neurology; ocul, ocular; Pulm, pulmonary.
### Table A1. Management of Immune-Related Adverse Events Guideline Expert Panel Membership

<table>
<thead>
<tr>
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<th>Role/Area of Expertise</th>
</tr>
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### Table A2. Commonly Conducted Testing at Baseline Prior to ICPI Therapy*

<table>
<thead>
<tr>
<th>Testing</th>
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<tbody>
<tr>
<td>Clinical</td>
</tr>
<tr>
<td>Physical examination, including physical stature, weight, body mass index, heart rate, and blood pressure</td>
</tr>
<tr>
<td>Comprehensive history, including autoimmune, organ-specific disease, endocrinopathy, neuropathy, and infectious disease</td>
</tr>
<tr>
<td>Questioning of general health, including appetite, bowel habits, and asthenia. Preexisting symptoms involving bowel movements, dyspnea, cough, rash, headaches, and arthralgia should be noted.</td>
</tr>
<tr>
<td>Laboratory</td>
</tr>
<tr>
<td>CBC + differential test</td>
</tr>
<tr>
<td>Complete metabolic panel that may include serum electrolytes (Na, K, Ca, CO₂), liver function (AST, ALT, alkaline phosphatase, γ-glutamyl transferase), creatinine, creatine kinase, total bilirubin</td>
</tr>
<tr>
<td>Glucose</td>
</tr>
<tr>
<td>Lactate dehydrogenase and aldolase</td>
</tr>
<tr>
<td>Thyroid-stimulating hormone, free thyroxine</td>
</tr>
<tr>
<td>Luteinizing hormone, follicle-stimulating hormone, and testosterone levels in males or estrogen in premenopausal females with fatigue, loss of libido, and mood changes</td>
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<td>Urinalysis</td>
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<tr>
<td>Surveillance for latent tuberculosis</td>
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<td>Virology including HIV, hepatitis C virus and hepatitis B virus, Epstein-Barr virus, cytomegalovirus</td>
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<tr>
<td>Troponin</td>
</tr>
<tr>
<td>Spirometry/diffusing capacity of lung for carbon monoxide</td>
</tr>
<tr>
<td>Imaging</td>
</tr>
<tr>
<td>Chest x-ray</td>
</tr>
<tr>
<td>Computed tomography</td>
</tr>
<tr>
<td>ECG</td>
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</tbody>
</table>

*Other testing may also be necessary based on patient’s history and preexisting comorbidities and/or risk factors.
<table>
<thead>
<tr>
<th>Acronym</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>ABG</td>
<td>arterial blood gas</td>
</tr>
<tr>
<td>ACC</td>
<td>American College of Cardiology</td>
</tr>
<tr>
<td>ACTH</td>
<td>adrenocorticotropic hormone</td>
</tr>
<tr>
<td>ADL</td>
<td>activities of daily living</td>
</tr>
<tr>
<td>AE</td>
<td>adverse event</td>
</tr>
<tr>
<td>AHA</td>
<td>acquired hemophilia A</td>
</tr>
<tr>
<td>AHA</td>
<td>American Heart Association</td>
</tr>
<tr>
<td>AI</td>
<td>adrenal insufficiency</td>
</tr>
<tr>
<td>ALT</td>
<td>alanine aminotransferase</td>
</tr>
<tr>
<td>ANA</td>
<td>antinuclear antibodies</td>
</tr>
<tr>
<td>ANCA</td>
<td>antineutrophil cytoplasmic antibodies</td>
</tr>
<tr>
<td>ANNA-1</td>
<td>anti-neuronal nuclear antibody 1</td>
</tr>
<tr>
<td>APTT</td>
<td>activated partial thromboplastin time</td>
</tr>
<tr>
<td>ASCO</td>
<td>American Society of Clinical Oncology</td>
</tr>
<tr>
<td>ASH</td>
<td>American Society of Hematology</td>
</tr>
<tr>
<td>ASMA</td>
<td>anti–smooth muscle antibodies</td>
</tr>
<tr>
<td>AST</td>
<td>aspartate aminotransferase</td>
</tr>
<tr>
<td>ATG</td>
<td>antithymocyte globulin</td>
</tr>
<tr>
<td>BAL</td>
<td>bronchoalveolar lavage</td>
</tr>
<tr>
<td>BNP</td>
<td>brain natriuretic peptide</td>
</tr>
<tr>
<td>BSA</td>
<td>body surface area</td>
</tr>
<tr>
<td>CAP-T</td>
<td>chimeric antigen receptor T-cell</td>
</tr>
<tr>
<td>CBC</td>
<td>complete blood count</td>
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<tr>
<td>CMP</td>
<td>comprehensive metabolic panel</td>
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<tr>
<td>CMV</td>
<td>cytomegaloovirus</td>
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<tr>
<td>CNS</td>
<td>central nervous system</td>
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<tr>
<td>CPK</td>
<td>creatine phosphokinase</td>
</tr>
<tr>
<td>CRP</td>
<td>C-reactive protein</td>
</tr>
<tr>
<td>CSF</td>
<td>cerebrospinal fluid</td>
</tr>
<tr>
<td>CT</td>
<td>computed tomography</td>
</tr>
<tr>
<td>CTCaE</td>
<td>Common Terminology Criteria for Adverse Events</td>
</tr>
<tr>
<td>CTLA-4</td>
<td>cytotoxic T-cell lymphocyte-4</td>
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<tr>
<td>CTPI</td>
<td>immune checkpoint inhibitor</td>
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<tr>
<td>ICU</td>
<td>intensive care unit</td>
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<tr>
<td>IgG</td>
<td>immunoglobulin G</td>
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<tr>
<td>iAE</td>
<td>immune-related adverse event</td>
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<td>ITP</td>
<td>idiopathic thrombocytopenic purpura</td>
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<tr>
<td>IV</td>
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<td>IVIG</td>
<td>intravenous immunoglobulin</td>
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<td>LEMS</td>
<td>Lambert-Eaton Myasthenic Syndrome</td>
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<td>LFTs</td>
<td>liver function tests</td>
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<td>LH</td>
<td>lutenizing hormone</td>
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<tr>
<td>LLN</td>
<td>lower limit of normal</td>
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<tr>
<td>LMWH</td>
<td>low molecular weight heparin</td>
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<tr>
<td>MCV</td>
<td>mean corpuscular volume</td>
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<tr>
<td>MDS</td>
<td>myelodysplastic syndromes</td>
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<tr>
<td>MGFA</td>
<td>myasthenia gravis foundation of America</td>
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<tr>
<td>MMF</td>
<td>mycophenolate mofetil</td>
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<tr>
<td>MRI</td>
<td>magnetic resonance imaging</td>
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<tr>
<td>NCS</td>
<td>nerve conduction study</td>
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<tr>
<td>NIF</td>
<td>negative inspiratory force</td>
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<tr>
<td>O&amp;P</td>
<td>ova and parasite</td>
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<tr>
<td>PCP</td>
<td>Pneumocystis pneumonia</td>
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<tr>
<td>PCR</td>
<td>polymerase chain reaction</td>
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<td>PD-1</td>
<td>programmed death 1</td>
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<tr>
<td>PD-L1</td>
<td>programmed death ligand 1</td>
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<tr>
<td>PE</td>
<td>pulmonary embolism</td>
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<tr>
<td>PEX</td>
<td>plasma exchange</td>
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<tr>
<td>PMNs</td>
<td>polymorphonuclear cells</td>
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<tr>
<td>PNH</td>
<td>paroxysmal nocturnal hemoglobinuria</td>
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<tr>
<td>PPI</td>
<td>proton pump inhibitor</td>
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<tr>
<td>PT</td>
<td>prothrombin time</td>
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<tr>
<td>PTU</td>
<td>propylthiouracil</td>
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<tr>
<td>RBC</td>
<td>red blood cell</td>
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<tr>
<td>RPR</td>
<td>rapid plasma reagin</td>
</tr>
<tr>
<td>T1DM</td>
<td>type 1 diabetes mellitus</td>
</tr>
<tr>
<td>T2DM</td>
<td>type 2 diabetes mellitus</td>
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<tr>
<td>TB</td>
<td>tuberculosis</td>
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<tr>
<td>TID</td>
<td>three times a day</td>
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<tr>
<td>TPO</td>
<td>thyroid peroxidase</td>
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<tr>
<td>TSH</td>
<td>thyroid stimulating hormone</td>
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<tr>
<td>TSI</td>
<td>thyroid stimulation immunoglobulin</td>
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<tr>
<td>TTE</td>
<td>transthoracic echocardiogram</td>
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<tr>
<td>TTP</td>
<td>thrombotic thrombocytopenic purpura</td>
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<tr>
<td>ULN</td>
<td>upper limit of normal</td>
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<td>US</td>
<td>ultrasound</td>
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<td>UTIs</td>
<td>urinary tract infections</td>
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<td>V/Q</td>
<td>ventilation-perfusion lung scan</td>
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<td>VC</td>
<td>vital capacity</td>
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<td>VKA</td>
<td>vitamin K antagonists</td>
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<tr>
<td>VTE</td>
<td>venous thromboembolism</td>
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<tr>
<td>WBC</td>
<td>white blood cell</td>
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</table>

(continued in next column)