Approximately 5% or fewer metastatic colorectal cancers (CRCs) have defective DNA mismatch repair systems (dMMR) caused by either sporadic or genetic DNA mutations. Characterizing dMMR status is critical in understanding the genetic makeup of cancer cells and possible immunotherapy treatment opportunities and should be included in the diagnostic workup of all patients with CRC. Defective DNA mismatch repair systems cause cellular hypermutations and high microsatellite instability (MSI-H), which makes the cancer cells susceptible to immune checkpoint inhibitors. Patients with metastatic CRC and dMMR status have recently been found in clinical trials to have significantly improved objective response rates and progression-free survival rates when given pembrolizumab (Keytruda®), an anti-programmed death receptor-1 (PD-1) monoclonal antibody checkpoint inhibitor.

PD-1 is a negative regulator of T-cell activation and proliferation, meaning it “turns the immune response off,” essentially acting as a break. This type of inhibitory role is necessary to prevent excessive immune reaction and autoimmunity. For this reason, PD-1 and other regulators acting in this manner are known as immune checkpoints. We now understand that some tumors can exploit the PD-1 pathway, enabling them to evade the immune response. Pembrolizumab selectively binds to PD-1, thus blocking the inhibitory pathway and releasing the immune system. This allows the immune response to occur.

Pembrolizumab is approved as monotherapy in adult and pediatric patients with MSI-H or dMMR unresectable or metastatic CRC that has progressed following exposure to a fluopyrimidine, oxaliplatin, or irinotecan.

This document is part of an overall HCP toolkit intended to assist providers in optimizing management of d-MMR and MSI-H metastatic colorectal cancer patients receiving pembrolizumab.
DRUG DOSAGE/ADMINISTRATION

- For dMMR and MSI-H metastatic CRC in adults, the recommended dosage of pembrolizumab (Keytruda®) is 200 mg administered as an intravenous infusion over 30 minutes every 3 weeks until disease progression or unacceptable toxicity, with a maximum treatment length of 24 months without disease progression.

- For pediatric patients, the recommended dosage of pembrolizumab is 2 mg/kg, up to a maximum of 200 mg/kg, administered as an intravenous infusion over 30 minutes until disease progression or unacceptable toxicity, or up to 24 months in patients without disease progression.

- Pembrolizumab solution is clear to slightly opalescent, colorless to slightly yellow. Discard the vial if visible particles are observed.

- Pembrolizumab is provided as a 50 mg lyophilized powder in a single-dose vial for reconstitution or as a 100 mg/4 mL (25 mg/mL) solution in a single-dose vial. When reconstituting pembrolizumab for injection, slowly swirl the vial. Do NOT shake the vial.

- Pembrolizumab is classified as an irritant and may be safely administered via a central or peripheral line. It is important to assure IV access before administration. Pembrolizumab should be administered through an intravenous line containing a sterile, non-pyrogenic, low-protein-binding in-line or add-on filter (pore size of 0.2 – 5 micrometers). Do not coadminister pembrolizumab with other drugs through the same intravenous line.
SIDE EFFECTS AND THEIR MANAGEMENT

Because pembrolizumab is an immunotherapy that works by enhancing the patient’s immune system, most adverse reactions associated with pembrolizumab are related to overactivity of the patient’s immune system (ie, immune-related adverse events [irAEs]). Various organ systems (often more than one) or tissues may be affected.

- Keys to toxicity management:
  - Proactive assessment for early signs/symptoms of toxicity
  - Prompt intervention
  - irAEs are typically managed with treatment interruption and selective use of corticosteroids
  - In rare instances, toxicity may be steroid refractory, and additional immunosuppressive agents may be necessary (infliximab, mycophenolate mofetil, cyclophosphamide, etc)
  - Pembrolizumab will likely be held or discontinued depending on severity and/or persistence
  - Referral to organ specialist should be considered, given that unique testing and management strategies may be required

*irAEs associated with pembrolizumab treatment can be categorized into those that are most common, less common but serious, and others that are easily overlooked. (Table 1; Appendix 1). Other adverse events associated with pembrolizumab therapy are listed in Appendix 2

### Table 1. Care Step Pathways for the management of immune-related AEs associated with pembrolizumab monotherapy.

<table>
<thead>
<tr>
<th>irAE category</th>
<th>Examples</th>
<th>Location</th>
</tr>
</thead>
<tbody>
<tr>
<td>Most common</td>
<td>Skin toxicities (pruritis, rash, etc)</td>
<td>Appendix 1</td>
</tr>
<tr>
<td></td>
<td>Gastrointestinal toxicity: Diarrhea and colitis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Thyroiditis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hepatic toxicities</td>
<td></td>
</tr>
<tr>
<td>Less common but serious</td>
<td>Additional endocrinopathies</td>
<td>Appendix 1</td>
</tr>
<tr>
<td></td>
<td>- Hypophysitis (pituitary)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Adrenal insufficiency (adrenalitis)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Diabetes</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Pneumonitis</td>
<td></td>
</tr>
<tr>
<td>Easily overlooked</td>
<td>Arthralgia/arthritis</td>
<td>Appendix 1</td>
</tr>
<tr>
<td></td>
<td>Mucositis/xerostomia</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Neuropathy</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Nephritis</td>
<td></td>
</tr>
</tbody>
</table>
**CLINICAL PEARLS**

- Programmed cell death ligand 1 (PD-L1) status or elevated expression is not a prerequisite for pembrolizumab treatment of dMMR and MSI-H metastatic colorectal cancer as it is in gastric cancer, but genetic testing on the cancer cells to determine dMMR or MSI-H status is a prerequisite prior to initiating treatment, as it is for some lung-cancer indications.

- It is important to monitor laboratory values at the start of treatment, periodically during treatment, and as indicated clinically. Laboratory values commonly monitored include: CBC w/ differential, creatinine, alkaline phosphatase, AST/ALT, bilirubin (direct/total), sodium, potassium, calcium, magnesium, thyroid function, and glucose. See individual irAE CSPs for more detail on laboratory monitoring.

- Pembrolizumab-related irAEs may occur at any time, including after treatment completion or discontinuation.

- Patients sometimes experience signs/symptoms that they think are due to “flu” or a cold, but that actually represent an irAE or an infusion reaction.

- Endocrinopathies often present with vague symptoms (fatigue, headache, and/or depression) that can easily be overlooked or initially misdiagnosed. Hypervigilance and follow-up is important on the part of both HCPs and patients.

- Unlike other irAEs, endocrinopathies usually do not resolve and may require lifelong hormone replacement therapy.

- IrAEs have different time courses. New irAEs may become apparent upon tapering of corticosteroids used to treat an earlier onset irAE, since the new irAE can be suppressed or masked by immunosuppressive therapy. Therefore, during the taper period, patients should be advised to be on the lookout for early signs of new irAEs as well as recurrence of the original irAE that was being treated.

- HCPs should encourage patients to carry information about their pembrolizumab regimen with them at all times. This might be the pembrolizumab-specific wallet card, or at least emergency phone numbers and the side effects associated with the regimen. You may suggest that they paperclip the wallet and insurance cards together so information about their regimen will be shared whenever they show the insurance card.

- Advise patients to take pictures of any skin changes for documentation.
Q. How long will patients stay on pembrolizumab?

A. The prescribing information indicates until disease progression or unacceptable toxicity with a maximum treatment duration of 24 months without disease progression. The interpretation of these criteria varies from institution to institution and from provider to provider.

Q. Are there standard dosage reductions for irAEs associated with pembrolizumab?

A. There are no dosage reductions for irAEs associated with pembrolizumab. The dose is either held until the irAE resolves sufficiently (typically to Grade 0 or Grade 1) or, if the irAE is severe enough, pembrolizumab is discontinued permanently.

Q. Does the safety profile of pembrolizumab differ when it is used in various tumor types?

A. Generally, the safety profile of pembrolizumab is similar across tumor types. However, the context may be different—patients with other tumor types may have differing comorbidities or underlying organ dysfunction. For example, lung cancer patients may have underlying lung disease that will exacerbate shortness of breath associated with pneumonitis, but metastatic colorectal cancer patients may have more problems with diarrhea and/or colitis.

Q. How do I counsel my patients about immunizations?

A. That’s a logical question, given that the checkpoint inhibitors alter the immune response. Advise your patients not to receive live vaccines (eg, measles, mumps, and rubella and the varicella vaccine [Zostavax®]) because they have not been evaluated in this setting. The use of attenuated vaccines has been and continues to be evaluated. Counsel patients to discuss all immunizations with the oncology team prior to administration so the benefits and risks can be weighed on an individual basis. For example, SHINGRIX®, approved in 2017, is an attenuated (non-live) varicella vaccine; its use be discussed with the oncology team if a recommendation is being made for the patient to receive the injection series. Annual influenza vaccination with the inactivated influenza vaccine is recommended. The nasal spray flu vaccine is a live attenuated influenza vaccine and should not be administered to patients treated with immune checkpoint inhibitors.
Q. How is a pembrolizumab-induced rash different from an EGFR-induced rash seen with cetuximab or panitumumab?

A. Generally, an EGFR-induced rash manifests earlier, is more macular-papular, less pruritic and responsive to the STEPP skin protocol. A rash induced from pembrolizumab tends to be more pruritic with tightness/burning sensation, responds to oral corticosteroids, and can occur at any point during treatment.

Q. What are the differences between oxaliplatin-induced peripheral neuropathy (PN) and nerve problems seen with pembrolizumab?

A. Oxaliplatin PN tends to happen on the hands and feet with tingling and numbness. Pembrolizumab-induced nerve problems manifest as unilateral weakness and difficulty walking, but can also manifest with worsening numbness/tinglings/functional impairment. It is imperative to ascertain baseline neurological deficits prior to initiating pembrolizumab for appropriate evaluation of worsening symptoms.

Q. How do we test for MSI and MMR?

A. Polymerase chain reaction for MSI and immunohistochemistry for MMR can be requested to identify patients appropriate for treatment with pembrolizumab. Your pathologist may be able to perform the testing in your hospital/laboratory or may need to request testing from a reference lab.
PATIENT RESOURCES

Financial Assistance
The Merck Access Program
1-855-257-3932
www.keytruda.com/keytruda-cost/

Additional Information Resources
Colorectal Cancer Alliance
http://www.ccalliance.org/

American Cancer Society Resource Section
ADDITIONAL RESOURCES


ADDITIONAL RESOURCES


APPENDIX 1
The 12 Care Step Pathways (CSPs) referenced in this Appendix are housed in the CSP section of the AIMWithImmunotherapy.com IO Essentials website. These CSP are currently universally applicable (i.e., they don’t differ across tumor types).

Please click the link below to access the CSPs, which can also be printed from that section of the site.

http://aimwithimmunotherapy.org/care-step-pathways/
APPENDIX 2
### Management of other AEs associated with pembrolizumab monotherapy

<table>
<thead>
<tr>
<th>Adverse event</th>
<th>Common symptoms</th>
<th>Common management/anticipatory guidance</th>
</tr>
</thead>
</table>
| Anorexia                 | Decreased appetite                                                                                        | • Monitor weight; query patient about appetite/eating habits; advise dietary modification if necessary (should improve with time)  
  • Anticipate standard dose holds/discontinuations*  
  • Consider referral to nutrition services for counseling on best food choices to avoid excessive weight loss |
| Constipation/abdominal pain | Infrequent stools/difficulty stooling, abdominal pain                                                   | • Consider other causes, such as opioid-induced constipation  
  • Increase fluid, fiber; use laxatives with caution; suggest stool softeners and physical activity  
  • Consider appropriate testing to evaluate bowel obstruction  
  • Anticipate standard dose holds/discontinuations* for Grade 3 and Grade 4 (constipation with manual evacuation indicated, severe abdominal pain, or life-threatening consequences) |
| Embryo-fetal toxicity    | —                                                                                                        | • Advise of risk to fetus and recommend use of effective contraception during treatment and for 4 months after pembrolizumab is discontinued  
  • Advise patients to tell their HCPs immediately if they or their partners suspect they are pregnant while taking therapy |
| Encephalitis             | Headache, fever, tiredness, confusion, memory problems, sleepiness, hallucinations, seizures, stiff neck | • New-onset, moderate-to-severe symptoms: rule out infectious or other causes  
  • Counsel neurologist, obtain brain MRI, and lumbar puncture  
  • Anticipate standard dose-holds and discontinuations* |
| Fatigue                  | Feeling tired; lack of energy                                                                             | • Query patients regarding energy level; evaluate possible contributory factors, including infection, disease progression, and hematological and metabolic abnormalities; standard supportive care  
  • Anticipate standard dose holds/discontinuations*  
  • Fatigue that interferes with ADLs is concerning and should be evaluated for underlying causes, such as possible endocrinopathy |
| Headache                 | Head pain                                                                                                | • Need to rule out brain metastases, encephalitis, or hypophysitis; otherwise, standard supportive care (should improve with time)  
  • Headache occurring in conjunction with fatigue could be indicative of hypophysitis; obtain MRI of brain with pituitary slices/pituitary protocol to confirm/rule out  
  • Anticipate standard dose holds/discontinuations* |
### Management of other AEs associated with pembrolizumab monotherapy

(Continued)

<table>
<thead>
<tr>
<th>Adverse event</th>
<th>Common symptoms</th>
<th>Common management/anticipatory guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Infusion reaction</strong></td>
<td>Chills/shaking, back pain, itching, flushing, difficulty breathing, hypotension, fever</td>
<td>* Monitor patients for signs and symptoms. For grade 3 or 4 reactions: stop infusion and permanently discontinue pembrolizumab</td>
</tr>
</tbody>
</table>
| **Insomnia** | Difficulty falling or staying asleep | * Counsel patients on good sleep habits; prescription medications can be used if needed (should improve over time)  
* Anticipate standard dose holds/discontinuations* |
| **Myocarditis** | Shortness of breath; arrhythmia; light-headedness; chest pain; fatigue; nausea; edema | * Obtain baseline ECG  
* Assess cardiac biomarkers (BNP; troponin)  
* Control cardiac diseases (and risk factors) optimally  
* Consult cardiologist and consider corticosteroids if myocarditis is suspected  
* Add additional immunosuppressive agents in severe, refractory cases  
* Institute standard dose holds/discontinuations (in consultation with cardiologist) |
| **Nausea/vomiting** | Vomiting, queasiness, RUQ or LUQ pain | * Rule out brain metastases and gastroenteritis  
* Provide standard supportive care, since it is adequate in most cases  
* Check LFTs/lipase/amylase if hepatotoxicity or pancreatitis are suspected  
* Anticipate standard dose holds/discontinuations* |
| **Upper respiratory tract infection** | Cough, runny nose, sore throat, nasal congestion | * Evaluate potential causes—a dry cough and shortness of breath would increase concern for pneumonitis  
* Standard supportive care  
* Anticipate standard treatment holds* |
| **Vision changes** | Eye redness, pain, blurred vision, photophobia | * Test and evaluate for uveitis and episcleritis (by ophthalmologist, preferably)  
* Urgency of ophthalmology referral increases with grade  
* G1: continue immunotherapy, use artificial tears  
* G2: hold immunotherapy; ophthalmic and systemic corticosteroids (under ophthalmologist guidance)  
* G3 or G4: permanently discontinue immunotherapy; treatment by ophthalmologist to include ophthalmic and systemic corticosteroids |

*Withhold pembrolizumab for any Grade 3 (severe) AE. Permanently discontinue for any Grade 4 (life-threatening) AE, persistent Grade 2–3 AE, any severe (Grade 3) AE that recurs, or when ≥10 mg/d prednisone or equivalent is required for 12 weeks. Resume treatment when AE returns to Grade 0 or 1.