

- malignant pleural mesothelioma (MPM)
- melanoma
- Merkel cell carcinoma (MCC)
- nasopharyngeal carcinoma
- Non-pancreatic neuroendocrine tumor (non-pNET)
- non-small cell lung cancer (NSCLC) first-line treatment
- non-small cell lung cancer (NSCLC) neoadjuvant therapy/treatment
- non-small cell lung cancer (NSCLC) subsequent treatment
- pancreatic adenocarcinoma
- primary mediastinal large B-cell lymphoma (PMLBCL)
- renal cell carcinoma (RCC)
- small bowel adenocarcinoma (SBA)
- soft tissue sarcomas (including angiosarcoma, those of the extremities/body wall/head/neck/retroperitoneal/intra-abdominal, and rhabdomyosarcoma)
- squamous cell vulvar carcinoma
- small cell lung cancer (SCLC)
- urothelial carcinoma (UCC, also transitional cell carcinoma [TCC])
- squamous cell carcinoma of the head and neck (SCCHN)
- other (*please specify*):

Clinical Information

(if nasopharyngeal carcinoma) Has the patient been started on Opdivo? Yes No

(if no, and nasopharyngeal carcinoma) Does the patient have recurrent or metastatic non-keratinizing disease? Yes No

(if yes and nasopharyngeal carcinoma) Is this medication being used as subsequent therapy? Yes No

(if no, and nasopharyngeal carcinoma) Does the patient have recurrent, unresectable, oligometastatic, or metastatic disease? Yes No

(if yes and nasopharyngeal carcinoma, if first line treatment) Will this medication be used in combination with cisplatin and gemcitabine? Yes No

(if yes and nasopharyngeal carcinoma) The covered alternative is Loqtorzi (toripalimab intravenous infusion) [may require prior authorization]. If your patient has tried this drug, please provide drug strength, date(s) taken and for how long, and what the documented results were of taking this drug, including any intolerances or adverse reactions your patient experienced. If your patient has NOT tried this drug, please provide details why your patient can't try this alternative.

(if nasopharyngeal carcinoma) Per the information provided above, which of the following is true for your patient in regard to the covered alternative?

- The patient tried the alternative, but it didn't work well enough
- The patient tried the alternative, but they did not tolerate it
- The patient cannot try the alternative because of a contraindication to this drug
- Other

Is this new start or continuation of therapy? new start continuation of therapy
 (if continuation of therapy) Is your patient responding to therapy OR is your patient NOT having disease progression while on the requested drug? Yes No

*****This drug requires supportive documentation (i.e. genetic testing, chart notes, lab/test results, etc). Supportive documentation for all answers must be attached with this request.**

(if anal cell carcinoma, endometrial, non-pNET, squamous vulvar) Was your patient previously treated with only one other chemotherapy regimen for this diagnosis? Yes No

(if non-pNET and previous first line chemo) Did your patient experience progression (or disease worsening) while on first line chemotherapy? Yes No

(if endometrial) Does your patient have recurrent or metastatic disease? Yes No
 (if not recurrent or metastatic) Does your patient have high-risk mismatch repair deficient (dMMR) tumors? Yes No

(if bone cancer) Does your patient have tissue tumor mutation burden-high (TMB-H) tumors with 10 or more mutations per megabase? Yes No

(if bone cancer) Has your patient previously been treated with any therapy for this diagnosis? Yes No
 (if yes) Did your patient have disease progression with the previous treatment? Yes No

(if bone cancer) Are there any satisfactory alternative options available for treatment? Yes No

(if brain mets) Is melanoma the primary tumor/site? Yes No
 (if no) What is the primary tumor/site _____

(if brain mets) Does your patient have recurrent disease? Yes No

(if CRC) Does your patient have unresectable, advanced, or metastatic disease? Yes No

(if CRC or SBA) Has your patient undergone immunohistochemistry (IHC) or microsatellite instability (MSI) testing? Yes No
 (if yes) What were the results?
 deficient mismatch repair (dMMR) or microsatellite instability-high (MSI-H)
 proficient mismatch repair (pMMR) or microsatellite instability-low or stable (MSI-low or MSI-stable)

(if CRC) Has your patient previously used any type of chemotherapy for this diagnosis? Yes No
 (if no previous chemo) Is intensive therapy appropriate for your patient? Yes No
 (if previous chemo) Has your patient been previously treated with an oxaliplatin-based chemotherapy regimen? Yes No

(if CLL/SLL) Does your patient have the del(17p)/TP53 mutation? Yes No
 (if no) Is your patient refractory to chemotherapy and unable to receive chemoimmunotherapy? Yes No

(if ESCC) Is this the first therapy your patient has received for this diagnosis? Yes No
 (if yes) Will your patient be using Opdivo in combination with Yervoy (ipilimumab)? Yes No
 (if no) Was the patient previously treated with fluoropyrimidine (like capecitabine [Xeloda], floxuridine, and fluorouracil [5-FU, Adrucil]) and platinum-based chemotherapy (like carboplatin or cisplatin) for this diagnosis? Yes No
 (if yes) Does the patient have unresectable advanced, or metastatic disease? Yes No

(if gastric, GEJ or esophageal adenocarcinoma) Does your patient have advanced or metastatic disease? Yes No
 (if yes) Is/Will the requested medication (be)ing given in combination with a fluoropyrimidine (like capecitabine [Xeloda], floxuridine, and fluorouracil [5-FU, Adrucil]) and platinum (like carboplatin or cisplatin)-containing chemotherapy? Yes No

(if esophageal OR GEJ cancer except ESCC) Was your patient treated with chemoradiation followed by surgery to remove the cancer, but some cancer cells were found in the removed tumor or lymph nodes? Yes No
 (if yes) Is this medication being given to help prevent the cancer from coming back? Yes No

(if extranodal NK/T-cell lymphoma [nasal type] or PMLBCL) Does your patient have relapsed or refractory disease? Yes No

(if extranodal NK/T-cell lymphoma, nasal type) Was your patient previously treated with more than 1 regimen of chemotherapy? Yes No
 (if yes) Was one of the lines of therapy an alternate combination chemotherapy regimen (asparaginase-based) that was not previously used? Yes No

(if GTN) Does your patient have recurrent or progressive disease? Yes No
 (if GTN) Was your patient previously treated with a platinum/etoposide-containing regimen? Yes No
 (if HCC) Was your patient previously treated with Nexavar? Yes No

(if HL) Which type of Hodgkin lymphoma does your patient have?
 classical type
 nodular lymphocyte predominant type
 unknown

(if HL) Which of the following applies to your patient?
 relapsed or refractory disease
 palliative therapy and patient is older than 60 years
 neither of the above
 (if relapsed/refractory) Has your patient undergone an autologous stem cell transplant? Yes No
 (if yes) After the transplant, did your patient have therapy with Adcetris? Yes No

(if melanoma) How is this medication being used for this diagnosis?
 Adjuvant treatment for metastatic disease that has spread to the lymph nodes
 Adjuvant treatment for stage IIB/C disease
 Single-agent therapy
 In combination with ipilimumab (generic for Yervoy)
 Other

(if bone cancer OR melanoma & and not adjuvant) Does your patient have metastatic or unresectable disease? Yes No

(if melanoma & adjuvant tx) Did your patient have complete resection of the melanoma? Yes No

(if MPM) Which of the following applies?

- Drug requested is being used as single-agent therapy
- Drug requested is being given in combination with Yervoy
- other

(if NSCLC) Which best describes Opdivo's role in therapy?

- Opdivo is being given as first line treatment
- Opdivo is being given as subsequent therapy.
- Opdivo is being given as neoadjuvant therapy.
- unknown

(if cervical carcinoma) Is this medication being used as a second line or subsequent therapy? Yes No

(if cervical carcinoma) Does the patient have PD-L1 positive disease? Yes No

(if bone cancer, non-pNET or NSCLC [1st line]) Is/Will the requested drug be(ing) used in combination with Yervoy (ipilimumab)?
Yes No

(if anal cell carcinoma, non-pNET or NSCLC [1st line or subsequent]) Does your patient have metastatic disease? Yes No

(if NSCLC, 1st line) Is this medication being used as first-line therapy? Yes No

(if NSCLC, 1st line) Does your patient have PD-L1 expressing (greater than 1%) tumors? Yes No

(if NSCLC, 1st line) Does your patient have presence of EGFR (epidermal growth factor receptor) or ALK (anaplastic lymphoma kinase) genomic tumor aberrations? Yes No

(if NSCLC, neoadjuvant) Does the patient have resectable disease? Yes No

(if NSCLC, neoadjuvant) What is the patient's stage of disease?

- Occult (hidden) cancer
- Stage 0
- Stage 1 (includes: IA1, IA2, IA3, IB)
- Stage 2A (IIA)
- Stage 2B (IIB)
- Stage 3A (IIIA)
- Stage 3B (IIIB)
- Stage 3C (IIIC)
- Stage 4A (IVA)
- Stage 4B (IVB)

(if NSCLC, neoadjuvant) Is/Will the medication be(ing) given with platinum therapy (carboplatin, cisplatin)? Yes No

(if NSCLC, neoadjuvant, Stage 3B (IIIB)) Is/Will the medication be(ing) given with platinum-doublet chemotherapy? Yes No

(if NSCLC, neoadjuvant, Stage 3B (IIIB)) Will the patient receive Opdivo monotherapy as adjuvant therapy after surgery?
Yes No

(if NSCLC, neoadjuvant, Stage 3B (IIIB)) Is the patient previously untreated? Yes No

(if NSCLC, neoadjuvant, Stage 3B (IIIB)) Does your patient have known EGFR (epidermal growth factor receptor) mutations or ALK (anaplastic lymphoma kinase) rearrangements? Yes No

(if NSCLC, subsequent therapy) Does your patient have performance status 0-2? Yes No

(if NSCLC, subsequent) Was your patient previously treated with platinum-base chemotherapy, such as carboplatin or cisplatin?
Yes No

(if no) Which of the following applies to your patient?

- ALK-positive disease
- EGFR mutation-positive disease
- testing did not indicate either EGFR mutation- or ALK- positive disease
- molecular testing was not done

(if ALK-pos) Was your patient previously treated with Xalkori or Zykadia? Yes No

(if EGFR mutation-pos) Was your patient previously treated with Gilotrif, Iressa, or Tarceva (erlotinib)?
Yes No

(if pancreatic adenocarcinoma) Will your patient be using Opdivo in combination with Yervoy (ipilimumab)? Yes No

(if pancreatic adenocarcinoma) Is this medication being used as a second line or subsequent therapy? Yes No

(if pancreatic adenocarcinoma) Has the patient received prior immunotherapy? Yes No

(if pancreatic adenocarcinoma) Does your patient have tumor mutational burden-high (TMB-H) disease? Yes No

(if pancreatic adenocarcinoma) Does your patient have locally advanced or metastatic disease? Yes No

(if pancreatic adenocarcinoma) Does the patient have good performance status? Yes No

(if pancreatic adenocarcinoma) Did your patient have disease progression? Yes No

(if PMLBCL) Which of the following best describes how the requested drug will be given to this patient?

- single agent therapy
 given with Adcetris (brentuximab vedotin)
 neither of the above/unknown

(if EGFR mutation-pos) Was your patient previously treated with Gilotrif, Iressa, or Tarceva (erlotinib)? Yes No

(if SBA) Does your patient have advanced or metastatic disease? Yes No

(if SBA) Which of the following best describes how the requested drug will be given to this patient?

- as single agent therapy
 in combination with Yervoy (ipilimumab)
 neither of the above/unknown

(if SCCHN) Was your patient previously treated with platinum-base chemotherapy, such as carboplatin or cisplatin? Yes No

(if yes) Did your patient have progression of disease afterwards? Yes No

Please provide the following details: drug name, date(s) taken and for how long, and what the documented results were of taking each drug.

(if RCC) Does your patient have advanced, stage IV, or relapsed disease? Yes No

(if RCC) Will the drug requested be used in combination with Yervoy? Yes No

(if yes) Has your patient received any other chemotherapy before for this diagnosis? Yes No

(if RCC, not in combo with Yervoy) Will the drug requested be used in combination with Cabometyx? Yes No

(if RCC, with Cabometyx) Is this the first therapy your patient has received for this diagnosis? Yes No

(if RCC, not in combo with Yervoy or Cabometyx) Has your patient previously received anti-angiogenic therapy (for example: Afinitor, Avastin, Inlyta, Nexavar, Sutent, Votrient)? Yes No

(if squamous vulvar) Does your patient have HPV-related advanced, recurrent or metastatic disease? Yes No

(if anal cell carcinoma, cervical carcinoma, CRC, endometrial, GTN, HL, NSCLC [not in combo with Yervoy], SCCHN, squamous cell vulvar or RCC) Is the drug requested being used as single-agent therapy? Yes No

(if SCLC) Was your patient previously treated with platinum-based chemotherapy (such as carboplatin or cisplatin) AND a least one other line of therapy? Yes No

(if yes) Did your patient have progression of disease after these treatments? Yes No

Please provide the following details: drug name, date(s) taken and for how long, and what the documented results were of taking each drug.

(if UCC/TCC) Which of these best describes the use of the requested medication?

- As adjuvant treatment in patient at high risk of recurrence after undergoing radical resection
 As first line treatment
 For locally advanced or metastatic disease
 None of the above

(if UCC/TCC and locally advanced or metastatic) Was your patient previously treated with platinum-base chemotherapy, such as carboplatin or cisplatin? Yes No

(if yes) Did your patient have progression of disease while on the drug or afterwards? Yes No

(if UCC/TCC, and first line treatment) Will this medication be used in combination with cisplatin and gemcitabine? Yes No

(if UCC/TCC, and first line treatment) Does the patient have metastatic or unresectable disease? Yes No

(if biliary tract carcinoma) Has the patient previously been treated with any therapy for this diagnosis? Yes No

(if biliary tract carcinoma) Has the patient previously been treated with any systemic therapy for this diagnosis? Yes No

(if biliary tract carcinoma) Is this medication being used as a single agent? Yes No

(if biliary tract carcinoma) Will your patient be using Opdivo in combination with ipilimumab (generic for Yervoy)? Yes No

(if biliary tract carcinoma) Was your patient previously treated with a checkpoint inhibitor? Yes No

(if biliary tract carcinoma) Does your patient have tumor mutation burden-high (TMB-H) tumors? Yes No

(if biliary tract carcinoma) Did your patient have progression of disease during or after previous systemic treatments? Yes No

(if biliary tract carcinoma) Does the patient have unresectable, resected gross residual disease, or metastatic disease? Yes No

(if Kaposi sarcoma) Does your patient have relapsed OR refractory disease? Yes No

(if Kaposi sarcoma) Does your patient have advanced cutaneous, oral, visceral, or nodal disease? Yes No

(if Kaposi sarcoma) Did your patient have disease progression while on, or did not respond to, first line systemic therapy?

Yes No

(if Kaposi sarcoma) Will your patient be using Opdivo in combination with ipilimumab (generic for Yervoy)?

Yes No

(if anaplastic thyroid) Is this medication being used as a single-agent?

Yes No

(if anaplastic thyroid) Does the patient have stage IVC (metastatic) disease?

Yes No

(if anaplastic thyroid) How will this drug be used?

- As aggressive first-line therapy
 As second-line therapy
 None of the above

****This drug requires supportive documentation (i.e. genetic testing, chart notes, lab/test results, etc). Supportive documentation for all answers must be attached with this request.**

Additional Pertinent Information: *(including disease stage, prior therapy, performance status, and names/doses/admin schedule of any agents to be used concurrently).*

Attestation: I attest the information provided is true and accurate to the best of my knowledge. I understand that the Health Plan or insurer its designees may perform a routine audit and request the medical information necessary to verify the accuracy of the information reported on this form.

Prescriber Signature: _____ **Date:** _____

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