

Fax completed form to: (855) 840-1678
If this is an URGENT request, please call (800) 882-4462 (800.88.CIGNA)

Jemperli (dostarlimab)

PHYSICIAN INFORMATION		PATIENT INFORMATION					
* Physician Name: Specialty:	* DEA, NPI or TIN:		*Due to privacy regulations we will not be able to respond via fax with the outcome of our review unless all asterisked (*) items on				
	BEA, IN TO		this form are completed.*				
Office Contact Person:			* Patient Name:				
Office Phone:			* Cigna ID:	* Date of Birth:	* Date of Birth:		
Office Fax:			* Patient Street Address:				
Office Street Address:		City:	State:	Zip:			
City:	State:	Zip:	Patient Phone:				
Urgency: ☐ Standard ☐ Urgent (In checking this box, I attest to the fact that applying the standard review time frame may seriously jeopardize the customer's life, health, or ability to regain maximum function)							
Medication Requested:	☐ Jemperli 500	mg/10mL solution fo	r injection				
Dose: Frequency of therapy: ICD10: Duration of therapy:							
Where will this medication be obtained? Accredo Specialty Pharmacy** Prescriber's office stock (billing on a medical claim form) Retail pharmacy Other (please specify): **Medication orders can be placed with Accredo via E-prescribe - Accredo (1620 Century Center Pkwy, Memphis, TN 38134-8822 NCPDP 4436920), Fax 888.302.1028, or Verbal 866.759.1557							
Facility and/or doctor dispensing and administering medication: Facility Name: State: Tax ID#: Address (City, State, Zip Code):							
NOTE: Per some Cigna plans, infusion of medication MUST occur in the lowest cost, medically appropriate setting.							
Is this infusion occurring i	n a facility affi	iliated with hospital	outpatient setting?		Yes 🗌 No 🗌		
If yes- Is this patient a candidate for re-direction to an alternate setting (such as AIS, MDO, home) with assistance of a Specialty Care Option Case Manager? Yes ☐ No ☐ (provide medical necessity rationale):							
Is the requested medication for a chronic or long-term condition for which the prescription medication may be necessary for the life of the patient?							
Diagnosis related to use:							
ampullary adenocarcinon appendiceal adenocarcin breast cancer colon cancer endometrial cancer esophageal cancers esophagogastric junction other (please specify:	oma	☐ hepato☐ occult☐ ovarial☐ rectal o	c cancer obiliary cancers primary n cancer cancer cowel adenocarcinoma umors				

Clinical Information (if esophageal/esophagogastric junction/gastric) How will the requested medication be used in this patient? ☐ as palliative therapy ☐ as second-line therapy ☐ as subsequent therapy ☐ other					
(if palliative therapy) Is your patient a surgical candidate?	Yes 🗌 No 🗌				
(if yes) Does the patient have unresectable locally advanced, recurrent, or metastatic disease?	Yes ☐ No ☐				
(if palliative therapy) Does the patient have a Karnofsky performance score of at least 60% or an E performance score of 2 or less?	ECOG Yes □ No □				
(if second-line therapy or subsequent therapy) Did your patient's disease progress while on/followitreatment?	ing the previous Yes				
(if second-line therapy or subsequent therapy) Has the patient been previously treated with immur therapy (for example, Bavencio, Imfinzi, Keytruda, Libtayo, Opdivo, Tecentriq, Yervoy)?	no-oncology Yes				
(if endometrial, solid tumors) Does your patient have recurrent or advanced disease?	Yes 🗌 No 🗌				
(if endometrial) Does your patient have mismatch repair deficient (dMMR) disease?	Yes 🗌 No 🗌				
(if yes) Was the patient's dMMR disease determined by an FDA-approved test?	Yes 🗌 No 🗌				
(if endometrial) Has your patient been treated with a platinum-containing regimen before this medication?	Yes 🗌 No 🗌				
(if yes) Did your patient's disease progress while on or following the platinum-containing regimen?	Yes 🗌 No 🗌				
(if endometrial) Does your patient have microsatellite instability-high (MSI-H) disease?	Yes 🗌 No 🗌				
(if endometrial) Did the patient previously complete a course of this medication in combination therapy with carbopla					
(if yes) Will this medication be used as a monotherapy in the frontline setting?	Yes No Yes No No				
(if no) Will this medication be used in combination therapy with carboplatin and paclitaxel?	Yes 🗌 No 🗌				
(if yes) After completion of combination therapy, will this medication be used as a monotherapy in the frontl	ine setting? Yes				
(if solid tumors) Does your patient have mismatch repair deficient (dMMR) disease as determined by an FDA-approv					
(if solid tumors) Has your patient been treated with any other treatments before this medication for this diagnosis?	Yes ☐ No ☐ Yes ☐ No ☐				
(if yes) Did your patient's disease progress while on or following the previous treatment?	Yes ☐ No ☐				
(if solid tumors) Are there any alternative treatment options available that would be an option for your patient?	Yes ☐ No ☐				
(if endometrial, solid tumors) Will this medication be used as monotherapy?	Yes ☐ No ☐				
(if yes) Is the patient currently already receiving this medication?	Yes ☐ No ☐				
(if no) The covered alternative is Keytruda (pembrolizumab). If your patient has tried this medication, please provide strength, date(s) taken and for how long, and what the documented results were of taking this medication, including any intolerances or adverse reactions your patient experienced. If your patient has NOT tried this medication, please provide details why your patient can't try this alternative.					
Per the information provided above, which of the following is true for your patient in regards to the covered The patient has had a trial with this medication. The patient is able to try the alternative, but has not done so yet. The patient tried the alternative, but they did not tolerate it. The patient cannot try the alternative because of a contraindication to this medication. Other	alternative?				
(if ampullary adenocarcinoma, appendiceal adenocarcinoma, breast cancer, colon cancer, esophageal/esophagogastric junction/gastric cancer, hepatobiliary cancer) Is the requested medication being used as single-agent therapy? Yes ☐ No ☐					
(if ampullary adenocarcinoma, appendiceal adenocarcinoma, breast cancer) Has your patient been treated with any other treatments					

before this medication for this diagnosis?	Yes 🗌	No 🗌			
(if yes) Did your patient's disease progress while on/following the previous treatment?	Yes 🗌	No 🗌			
(if colon cancer) Does your patient have progression of advanced or metastatic disease?	Yes 🗌	No 🗌			
(if hepatobiliary cancers) Did your patient's disease progress while on/after systemic treatment?	Yes 🗌	No 🗌			
(if hepatobiliary cancers) Does your patient have unresectable or metastatic disease?	Yes 🗌	No 🗌			
(if ampullary adenocarcinoma, appendiceal adenocarcinoma, breast cancer, esophageal/esophagogastric junction/gastric cancer, hepatobiliary cancers) Are there any alternative treatment options available that would be an option for your patient? Yes \(\subseteq \) No \(\subseteq \)					
(if appendiceal adenocarcinoma, breast cancer) Does your patient have recurrent unresectable or stage IV disease?	Yes 🗌	No 🗌			
(if ovarian cancer) Does the patient have persistent disease or recurrence?	Yes 🗌	No 🗌			
(if ovarian cancer) Does the patient have recurrent or advanced tumors?	Yes 🗌	No 🗌			
(if rectal cancer) Does your patient have progression of advanced or metastatic disease?	Yes 🗌	No 🗌			
(if colon cancer, rectal cancer) Has your patient previously received oxaliplatin- irinotecan- and/or fluoropyrimidine-base					
(if small bowel adenocarcinoma) Does your patient have advanced or metastatic disease?	=	No □ No □			
(if colon cancer, hepatobiliary cancers, rectal cancer, small bowel adenocarcinoma) Has the patient been previously checkpoint inhibitor (for example, Yervoy, Keytruda, Opdivo)?					
(if small bowel adenocarcinoma) Has your patient previously received oxaliplatin in the adjuvant setting or has a contraindication to oxaliplatin?					
(if occult primary, ovarian, rectal, small bowel adenocarcinoma) Is the requested medication being used as single-agent therapy? Yes □ No □					
(if appendiceal adenocarcinoma, breast cancer, colon cancer, esophageal/esophagogastric junction/gastric cancer, cancers, occult primary, ovarian, rectal, small bowel adenocarcinoma) Does your patient have a microsatellite instable or mismatch repair deficient (dMMR) disease?					
Additional pertinent information (please include disease stage, prior therapy, performance status, and names/dose schedule of any agents to be used concurrently):	es/admin				
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:		ı			
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