

#### MEDICARE FORM

### Abraxane® (paclitaxel protein-bound particles) Injectable Medication **Precertification Request**

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non-preferred. The preferred (All fields must be completed and legible for precertification review.) products are docetaxel or paclitaxel. Docetaxel and Please indicate: Start of treatment: Start date \_\_\_\_/ paclitaxel do not require Continuation of therapy: Date of last treatment / / precertification. Precertification Requested By: Phone: Fax: A. PATIENT INFORMATION First Name: Last Name: DOB: ZIP: Address: City: State: Home Phone: Work Phone: Cell Phone: E-mail: Current Weight: kgs Height: inches or cms Allergies: lbs or **B. INSURANCE INFORMATION** Aetna Member ID #: Does patient have other coverage? ☐ Yes ☐ No Group #: If yes, provide ID#: Carrier Name: Insured: Insured: C. PRESCRIBER INFORMATION First Name: Last Name: (Check One): M.D. D.O. N.P. P.A. ZIP: Address: Citv: State: Fax: St Lic #: NPI#: DEA #: UPIN: Phone: Phone: Office Contact Name: D. DISPENSING PROVIDER/ADMINISTRATION INFORMATION Place of Administration: **Dispensing Provider/Pharmacy:** ☐ Physician's Office ☐ Self-administered ☐ Physician's Office ☐ Retail Pharmacv Phone: Outpatient Infusion Center ☐ Specialty Pharmacy ☐ Other \_\_\_\_\_ Center Name: \_\_\_\_ Name: \_\_\_ ☐ Home Infusion Center Phone: Address: Agency Name: \_ Phone: \_\_\_\_\_ Fax: \_\_\_\_\_ Administration code(s) (CPT): Address: **TIN:** \_\_\_\_\_\_ PIN: \_\_\_\_\_ NPI: NPI: E. PRODUCT INFORMATION HCPCS Code: Request is for: Abraxane (paclitaxel protein-bound): Dose: Frequency: F. DIAGNOSIS INFORMATION – Please indicate primary ICD Code and specify any other where applicable. Primary ICD Code: Secondary ICD Code: Other ICD Code: G. CLINICAL INFORMATION - Required clinical information must be completed in its entirety for all precertification requests. Note: Abraxane and generic paclitaxel (protein bound) are non-preferred. The preferred products are docetaxel or paclitaxel. Docetaxel and paclitaxel do not require precertification. ☐ Yes ☐ No Has the patient had prior therapy with Abraxane (paclitaxel protein-bound) within the last 365 days? Yes No Has the patient had a trial, intolerance, or contraindication to docetaxel or generic paclitaxel? Please explain if there are any medical reason(s) that the patient cannot use docetaxel or generic paclitaxel: For Initiation Requests (clinical documentation required for all requests): Will Abraxane be used to treat any of the following? (please mark all that apply) ☐ AIDS-related Kaposi sarcoma as subsequent therapy given with anti-retroviral therapy (ART) ☐ relapsed/refractory advanced, ☐ cutaneous, ☐ oral, ☐ visceral, OR ☐ nodal disease ☐ Recurrent OR metastatic breast cancer ☐ Single agent for human epidermal growth factor receptor 2 (HER2)-negative disease OR ☐ In combination with trastuzumab (Herceptin) for HER-2 positive recurrent or metastatic trastuzumab-exposed disease with symptomatic visceral disease OR visceral crisis, ☐ hormone receptor negative, OR hormone receptor positive & endocrine therapy refractory

> ☐ Substituted for either paclitaxel or docetaxel in persons who have experienced hypersensitivity reactions after receiving paclitaxel or docetaxel despite premedication, or for persons in whom standard hypersensitivity pre-medications

For Michigan MMP:

PHONE: 1-855-676-5772

Please use other form.

For other lines of business:

Note: Abraxane and generic paclitaxel (protein bound) are

FAX.

1-844-241-2495

are contraindicated



### **MEDICARE FORM**

# Abraxane® (paclitaxel protein-bound particles) Injectable Medication Precertification Request

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(All fields must be completed and legible for precertification review.)

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB				
G. CLINICAL INFORMATION (continued) – Re	equired clinical information must be comple	ted in its <u>entirety</u> for all precertific	ation requests.				
☐ Cervical cancer as a single age			•				
	☐ Local/regional recurrence OR ☐ distant metastases						
	Intrahepatic/Extrahepatic cholangiocarcinoma in combination with gemcitabine as primary treatment						
Unresectable disease Of		, ,					
Cutaneous melanoma as a sing	<b>-</b>						
☐ Unresectable disease Of	☐ Unresectable disease OR ☐ metastatic disease						
☐ status post disease prog	☐ status post disease progression OR ☐ after maximum clinical benefit from BRAF targeted therapy						
Endometrial Carcinoma							
☐ Primary treatment as a s	☐ Primary treatment as a single agent for endometrioid adenocarcinoma						
☐ Disease not su	☐ Disease not suitable for primary surgery						
	☐ that is limited to the uterus, ☐ with cervical involvement, OR ☐ extra-uterine disease						
☐ Pre-operatively	☐ Pre-operatively for disease that is suitable for primary surgery with abdominal/pelvic confined disease						
☐ For distant metastases							
☐ Single agent therapy for endometrioid adenocarcinoma							
☐ Distant/isolated metastases ☐ disseminated metastases that have progressed on hormonal therapy OR							
☐ are grade 2, 3, or large volume disseminated metastases OR							
☐ local/regional re	☐ local/regional recurrence in persons with gross upper abdominal residual disease						
☐ With sequentia	☐ With sequential external beam radiation therapy (EBRT) for local/regional recurrence with disease						
☐ Confi	☐ Confined to the vagina or pelvic lymph nodes ☐ in para-aortic or common iliac lymph nodes						
	☐ Local/regional recurrent disease for						
	☐ microscopic residual upper abdominal OR ☐ peritoneal disease						
	☐ received prior external beam radiation therapy (EBRT) to the site of recurrence						
	Carcinosarcoma, clear cell carcinoma, serous carcinoma, or undifferentiated/dedifferentiated carcinoma						
· · · · · · · · · · · · · · · · · · ·	imary treatment for disease not suitable						
∐ As ad	Iditional treatment for disease suitable fo						
	☐ With vaginal brachytherapy fro Stag		IV disease				
	Adjuvant treatment as single agent with histologic grade 3 tumors for						
	☐ Stage IB disease with vaginal brachytherapy and/or sequential external beam radiation therapy (EBRT)						
	☐ Stage II disease with sequential external beam radiation therapy (EBRT)						
	☐ Adjuvant treatment as single agent for ☐ Stage IIIA-IVA ☐ Stage IVB						
<ul> <li>□ Epithelial Ovarian Cancer for persistent or recurrent disease</li> <li>□ As a single agent □ With carboplatin for persons with confirmed taxane hypersensitivity</li> </ul>							
☐ Fallopian tube cancer for persi							
	ith carboplatin for persons with confirme	d taxane hypersensitivity					
	SCLC) for recurrent or metastatic dise		ormance status 2 OR in				
combination with carboplatin f	for performance status 0-2						
☐ 1st Line therapy							
	OS1, BRAF, and PD-L1 negative or unk	nown 🔲 BRAF V600E-mutati	on positive tumors				
☐ Subsequent therapy for							
	nutation positive tumors						
	☐ EGFR mutation positive and prior erlotinib/afatinib/gefitinib/osimertinib therapy						
	ALK positive tumors and prior crizotinib/ceritinib/alectinib/brigatinib therapy						
-	gement positive tumors and prior crizotin						
	( $\geq$ 50%) tumor, EGFR, ALK, ROS1, and						
	SCLC) when substituted for either pac						
	r receiving paclitaxel or docetaxel des	spite premedication, or for pe	rsons in whom standard				
hypersensitivity premedication	ns are contraindicated						

Continued on next page



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## Abraxane® (paclitaxel protein-bound particles) Injectable Medication Precertification Request

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For Michigan MMP: FAX: 1-844-241-2495 PHONE: 1-855-676-5772

For other lines of business: Please use other form.

Note: Abraxane and generic paclitaxel (protein bound) are non-preferred. The preferred products are docetaxel or paclitaxel. Docetaxel and paclitaxel do not require

precertification.

Patient First Name	Pa	atient Last Name	Patient Phone	Patient DOB		
O OLINIOAL INFORMAT						
G. CLINICAL INFORMAT	ION - Required clinical into	ormation must be completed for ALL precertification	on requests.			
	ancer in combination wit	th gemcitabine				
☐ As ne	eoadjuvant therapy	_				
		rline resectable disease OR 🗌 resectable diseas	•			
<u>_</u>		y tumors, large regional lymph nodes, excessive v	•	•		
_	1.7	s induction therapy followed by chemoradiation in	persons with good perfo	rmance status (KPS		
great	ter than or equal to 70)					
		astases in locally advanced unresectable disease		n metastatic disease		
∐ As se		sons with good performance status (KPS greater the				
		unresectable /metastatic disease and disease pro		pyrimidine-based therapy		
_ <b></b>		ne pancreatic bed after resection OR  For metas				
<ul> <li>□ Primary carcinoma of the urethra used as a single agent as subsequent systemic therapy for</li> <li>□ Recurrent disease OR □ Metastatic disease</li> </ul>						
<b>—</b>	_					
<ul> <li>□ Primary peritoneal cancer for persistent disease or recurrence</li> <li>□ in combination with carboplatin for persons with confirmed taxane hypersensitivity OR □ as a single agent</li> </ul>						
	-	•				
<ul> <li>☐ Upper genitourinary tract tumors used as a single agent as subsequent systemic therapy for metastatic disease</li> <li>☐ Urothelial carcinoma of the prostate used as a single agent as subsequent systemic therapy for metastatic disease</li> </ul>						
	-		c therapy for metastat	ic disease		
	oma as a single agent th					
☐ Meta	static OR  Unresectable	e disease				
For Continuation of Thera	apy: (clinical documenta	tion required):				
Is this a continuation reque	est a result of the patient re	eceiving samples of Abraxane® (paclitaxel proteir	n-bound particles)?	Yes 🗌 No		
Is there clinical documenta	ation supporting disease st	tability? 🗌 Yes 🔲 No				
Is there clinical documenta	ation supporting disease im	nprovement?				
H. ACKNOWLEDGEMEN	Т					
Request Completed By	(Signature Required):		Date	. , ,		
		zation of coverage of a medical procedure or servi				
		nformation or conceals material information for the erson to criminal and civil penalties.	ne purpose or misteadil	ig, commis a maudulent		
insurance act, which is a c	anne and subjects such pe	erson to criminal and civil penalties.				

The plan may request additional information or clarification, if needed, to evaluate requests.