

MEDICARE FORM

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

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Please indicate:	Continuation of therapy: Date of last treatment / /						non-preferred. The preferred products are docetaxel or paclitaxel. Docetaxel and paclitaxel do not require precertification.				
Precertification R						Phone: _			Fax: _		
A. PATIENT INFOR	RMATION										
First Name:				Last Name:					DOB:		
Address:		_		City:					State:	ZIP:	
Home Phone:		Work Phone:			Cell	Phone:			E-mail:		
Current Weight:	Ibs or	_kgs Height:	in	ches or	_ cms	Allergies	s:				
B. INSURANCE IN	FORMATION										
Aetna Member ID	#:		_ Doe	es patient have	e other	coverage?	☐ Ye	s 🗌 No			
Group #:				es, provide ID#			Carrie	er Name:			
Insured:			_ Ins	ured:							
C. PRESCRIBER II	NFORMATION										
First Name:			Las	t Name:				(Check One	e):	. 🗌 D.O. 🗌 N.P.	
Address:					(City:			State:	ZIP:	
Phone:	Fax:		St L	ic#:	1	NPI #:		DEA #:		UPIN:	
Provider E-mail:		0	ffice C	ontact Name:	•			Phone:			
D. DISPENSING PI	ROVIDER/ADMINI	STRATION INFOR	MATIC	N							
Place of Administ Self-administer Outpatient Infus Center Na Home Infusion Agency N Administration Address: NPI: E. PRODUCT INFO Request is for: At	red Phesion Center ame:Center ame:code(s) (CPT):	Phone:				☐ Physic ☐ Special Name: Address: Phone: TIN: NPI:	cian's Officialty Pharm	acy	Retail F Other _ Fax: _ PIN: _		
F. DIAGNOSIS INF									-		
Primary ICD Code:									Code:		
G. CLINICAL INFO	RMATION - Requ	ired clinical informa	tion mu	ust be complete	ed in its	entirety for a	all precertifi	cation reque	ests.		
Note: Abraxane and paclitaxel do not in the paclitaxel do not in the packet of the pac	nd generic paclit require precertifi as the patient had as the patient had	caxel (protein bou cation. prior therapy with a trial and failure,	nd) ar Abrax intole	e non-preferi ane (paclitaxe rance, or conti	red. The lprotein raindica	e preferred n-bound) wi	I products ithin the la	are doceta st 365 days onventional	axel or pac		el and
☐ Recu	used to treat any c -related Kaposi s -relapsed/refrac irrent OR metast - Single agent fo - In combination - with s - horm - horm	of the following? (parcoma as subsectory advanced, atic breast cance or human epiderma with trastuzumab symptomatic visce one receptor positions.	lease equen cutan cutan Herce (Herce ral dise tive, C	mark all that a t therapy give eous,	ipply) en with il, \[\]\ ptor 2 (i) -2 positeral criseraly reference.	risceral, OR HER2)-nega ive recurrer is, ractory	t ☐ noda ative disea nt or metas	l disease se OR static trastuz			
		tituted for either pa									

Virginia (HMO D-SNP) FAX: 1-833-280-52

PHONE: 1-855-463-0933 For other lines of business: Please use other form.

Note: Abraxane and generic paclitaxel (protein bound) are

1-833-280-5224

are contraindicated



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For other lines of business:

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
G. CLINICAL INFORMATION (continued) -	Required clinical information must be comple	ted in its <u>entirety</u> for all precertific	cation requests.					
☐ Cervical cancer as a single a	gent 2nd line therapy							
☐ Local/regional recurrer	☐ Local/regional recurrence OR ☐ distant metastases							
☐ Intrahepatic/Extrahepatic cho	Intrahepatic/Extrahepatic cholangiocarcinoma in combination with gemcitabine as primary treatment							
Unresectable disease	☐ Unresectable disease OR ☐ metastatic disease							
	Cutaneous melanoma as a single agent second line/subsequent therapy with performance status of 0-2 for							
☐ Unresectable disease	☐ Unresectable disease OR ☐ metastatic disease							
status post disease pro	☐ status post disease progression OR ☐ after maximum clinical benefit from BRAF targeted therapy							
Endometrial Carcinoma								
☐ Primary treatment as a	Primary treatment as a single agent for endometrioid adenocarcinoma							
	suitable for primary surgery							
	is limited to the uterus, \square with cervical in							
	☐ 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/regiona	☐ local/regional recurrence in persons with gross upper abdominal residual disease							
☐ With sequent	☐ With sequential external beam radiation therapy (EBRT) for local/regional recurrence with disease							
☐ Confined to the vagina or pelvic lymph nodes ☐ in para-aortic or common iliac lymph nodes								
☐ Local/regional recurrent disease for								
☐ mic	☐ 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							
	☐ As primary treatment for disease not suitable for primary surgery							
☐ As additional treatment for disease suitable for primary surgery								
_	☐ 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							
	persistent or recurrent disease With carboplatin for persons with confirme	ed tayana hyporeansitivity						
☐ Fallopian tube cancer for per		d taxarie riyperserisitivity						
	With carboplatin for persons with confirme	ed taxane hypersensitivity						
	ISCLC) for recurrent or metastatic dise		ormance status 2 OR in					
combination with carboplation		ace ac a congre agent for per-						
☐ 1st Line therapy	·							
	ROS1, BRAF, and PD-L1 negative or unk	nown 🔲 BRAF V600E-mutation	on positive tumors					
☐ Subsequent therapy fo								
☐ BRAF V600E	mutation positive tumors							
☐ EGFR mutati	on positive and prior erlotinib/afatinib/gefit	inib/osimertinib therapy						
☐ ALK positive tumors and prior crizotinib/ceritinib/alectinib/brigatinib therapy								
ROS1 rearrangement positive tumors and prior crizotinib therapy								
	re (≥50%) tumor, EGFR, ALK, ROS1, and		or pembrolizumab therapy.					
	ISCLC) when substituted for either pac		· · · · · · · · · · · · · · · · · · ·					
	er receiving paclitaxel or docetaxel des							
hypersensitivity premedication	<u> </u>							
- · · · · · · · · · · · · · · · · · · ·								

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G. CLINICAL INFORMATION - Required clinical	information must be completed for ALL precertification	on requests.	
Pancreatic cancer in combination As neoadjuvant therapy Biopsy positive bo CA 19-9, large prin As first line chemotherapy o greater than or equal to 70) Without systemic r As second-line therapy for p For locally advanc Local recurrence in Primary carcinoma of the urethra Recurrent disease OR M Primary peritoneal cancer for pers in combination with carbopla Upper genitourinary tract tumors	with gemcitabine rderline resectable disease OR resectable disease nary tumors, large regional lymph nodes, excessive or as induction therapy followed by chemoradiation in metastases in locally advanced unresectable disease ersons with good performance status (KPS greater the dunresectable /metastatic disease and disease product the pancreatic bed after resection OR For metastatic disease as single agent as subsequent systemic the etastatic disease or recurrence status for persons with confirmed taxane hypersensitivities used as a single agent as subsequent systemic that eused as a single agent as subsequent systemic the used as a single agent as subsequent systemic the therapy for	e with high-risk features weight loss, extreme pai persons with good perform First-line therapy in han or equal to 70) orgression following fluore static disease herapy for by OR as a single age herapy for metastatic of	n) primance status (KPS n metastatic disease ppyrimidine-based therapy ent disease
For Continuation of Therapy: (clinical docume) Is this a continuation request a result of the patier Is there clinical documentation supporting disease Is there clinical documentation supporting disease H. ACKNOWLEDGEMENT Request Completed By (Signature Required)	nt receiving samples of Abraxane® (paclitaxel proteir e stability? ☐ Yes ☐ No e improvement? ☐ Yes ☐ No	n-bound particles)? ☐	
Any person who knowingly files a request for auth	orization of coverage of a medical procedure or servi		

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