

MEDICARE FORM

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

Page 1 of 3

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

Continuation of therapy: Date of last treatment (All fields must be completed and legible for presented indicate: Start of treatment: Start date//					,			paclitaxel. Docetaxel and paclitaxel do not require precertification.			
Precertification R	Requested	l By:				Phone: _			Fax:		
A. PATIENT INFOR	RMATION										
First Name:				Last Name	e:				DOB:		
Address:				City:					State:	ZIF	P:
Home Phone:		Wor	k Phone:	•	Cell I	Phone:			E-mail:	<u>'</u>	
Current Weight:	lbs or	kgs	Height:	inches or	cms	Allergies	»:		•		
B. INSURANCE IN	FORMATIC	ON									
Aetna Member ID	#:			Does patient ha	ave other	coverage?	☐ Yes	i □ No			
Group #:				_ If yes, provide			Carrie	r Name: _			
Insured:				Insured:							
C. PRESCRIBER I	NFORMAT	ION									
First Name:				Last Name:	1		(Check On	<u> </u>	1	N.P. □ P.A.
Address:		1		1	C	City:		1	State:	ZIF	P:
Phone:		Fax:		St Lic #:	١	IPI#:		DEA #:		UPIN:	
Provider E-mail:			0	ffice Contact Nam	e:			Phone:			
D. DISPENSING P	ROVIDER/	ADMINISTRAT	TION INFOR	MATION		1					
Place of Administ		_				_	ng Provide		_		
☐ Self-administer		☐ Physiciar				-	cian's Office		Retail I	=	
Outpatient Infu	ision Cente ame:	er Pnon	e:			∐ Specia	alty Pharma	асу	☐ Other _		
☐ Home Infusion			e:								
Agency N						Address:					
☐ Administration	code(s) (C	PT):									
Address:									PIN: _		
NPI:						NPI:					
E. PRODUCT INFO											
Request is for: Al									HCI	PCS Code:	
F. DIAGNOSIS INF											
Primary ICD Code:				-			<u> </u>		<u> </u>		
G. CLINICAL INFO											
Note: Abraxane a				ınd) are non-pref	erred. The	preferred	products	are docet	axel or pa	clitaxel. Do	cetaxel and
paclitaxel do not				Abravana (naalita	wal protair	a bound\ wi	ithin tha lac	+ 265 days	.2		
☐ Yes ☐ No H										>	
Please explain if the	•								paomazor	•	
		,									
For Initiation Req	uests (clir	nical docume	ntation rec	uired for all requ	ests):						
Will Abraxane be	used to trea	at any of the f	ollowing? (p	lease mark all tha	t apply)						
				equent therapy g							
_	•	-] cutaneous, 🔲 d	oral, ∐ v	isceral, OR	. ∐ nodal	disease			
_		metastatic br		e r al growth factor re	centor 2 (F	HFR2\-negg	ative diseas	se OR			
		-	-	(Herceptin) for HE	-				zumab-exr	osed disea	se
				ral disease OR vis	-						
		hormone re				•					
				tive & endocrine th	erapy refr	actory					
				aclitaxel or doceta despite premedica							

For Illinois MMP:

FAX: 1-855-320-8445 **PHONE:** 1-866-600-2139

Please use other form.

Note: Abraxane and generic

For other lines of business:

paclitaxel (protein bound) are non-preferred. The preferred

are contraindicated



MEDICARE FORM

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

Page 2 of 3

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

For Illinois MMP:

FAX: 1-855-320-8445 **PHONE:** 1-866-600-2139

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	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	ent 2nd line therapy							
	☐ Local/regional recurrence OR ☐ distant metastases							
Intrahepatic/Extrahepatic chola	Intrahepatic/Extrahepatic cholangiocarcinoma in combination with gemcitabine as primary treatment							
	☐ Unresectable disease OR ☐ metastatic disease							
	Cutaneous melanoma as a single agent second line/subsequent therapy with performance status of 0-2 for							
	☐ Unresectable disease OR ☐ metastatic disease							
	☐ status post disease progression OR ☐ after maximum clinical benefit from BRAF targeted therapy							
_	Endometrial Carcinoma							
	☐ Primary treatment as a single agent for endometrioid adenocarcinoma							
	☐ Disease not suitable for primary surgery							
	that is limited to the uterus, with cervical involvement, OR extra-uterine disease							
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 recurrence in persons with gross upper abdominal residual disease							
☐ 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								
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 								
7.0 4.0	With vaginal brachytherapy fro Stag		IV disease					
☐ Adiuvant treatment as si	ngle agent with histologic grade 3 tumors							
	☐ 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 si	☐ Adjuvant treatment as single agent for							
☐ Stage IIIA-IVA								
Epithelial Ovarian Cancer for p								
	ith carboplatin for persons with confirme	d taxane hypersensitivity						
☐ Fallopian tube cancer for persi		14						
	ith carboplatin for persons with confirme		iormonos etetus 2 OR in					
combination with carboplating	SCLC) for recurrent or metastatic dise	ase as a single agent for peri	ormance status 2 OR in					
Ist Line therapy	ior performance status 0-2							
	OS1, BRAF, and PD-L1 negative or unki	nown BRAF V600F-mutatio	on positive tumors					
☐ Subsequent therapy for	COT, Brown, and I B ET Hogawood and	Iowii Brott voooz matatio	n postavo tamoro					
_ · · · · · · · · · · · · · · · · · · ·	nutation positive tumors							
	n positive and prior erlotinib/afatinib/gefiti	inib/osimertinib therapy						
	☐ ALK positive tumors and prior crizotinib/ceritinib/alectinib/brigatinib therapy							
•	☐ ROS1 rearrangement positive tumors and prior crizotinib therapy							
-	(≥50%) tumor, EGFR, ALK, ROS1, and	* *	or pembrolizumab therapy.					
	SCLC) when substituted for either pac							
	r receiving paclitaxel or docetaxel des		•					
hypersensitivity premedication	= -							
, , , , , , , , , , , , , , , , , , ,								

Continued on next page



MEDICARE FORM

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

Page 3 of 3

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

FAX: 1-855-320-8445 PHONE: 1-866-600-2139

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	Patient Last Name	Patient Phone	Patient DOB		
G. CLINICA	L INFORMATION - Required clinica	I information must be completed for ALL precerti	ification requests.			
G. CLINICAL INFORMATION - Required clinical information must be completed for ALL precertification requests. Pancreatic cancer in combination with gemcitabine As neoadjuvant therapy Biopsy positive borderline resectable disease OR resectable disease with high-risk features (ie, very highly elevated CA 19-9, large primary tumors, large regional lymph nodes, excessive weight loss, extreme pain) As first line chemotherapy or as induction therapy followed by chemoradiation in persons with good performance status (KPS greater than or equal to 70) Without systemic metastases in locally advanced unresectable disease First-line therapy in metastatic disease As second-line therapy for persons with good performance status (KPS greater than or equal to 70) For locally advanced unresectable /metastatic disease progression following fluoropyrimidine-based therapy Local recurrence in the pancreatic bed after resection OR For metastatic disease Primary carcinoma of the urethra used as a single agent as subsequent systemic therapy for Recurrent disease OR Metastatic disease Primary peritoneal cancer for persistent disease or recurrence in combination with carboplatin for persons with confirmed taxane hypersensitivity OR as a single agent Upper genitourinary tract tumors used as a single agent as subsequent systemic therapy for metastatic disease						
 ☐ Urothelial carcinoma of the prostate used as a single agent as subsequent systemic therapy for metastatic disease ☐ Uveal melanoma as a single agent therapy for ☐ Metastatic OR ☐ Unresectable disease 						
Is this a con	nation of Therapy: (clinical docume	entation required): ent receiving samples of Abraxane® (paclitaxel pages stability?	rotein-bound particles)?	□ Yes □ No		
H. ACKNO	WLEDGEMENT					
Request C	ompleted By (Signature Require	d):	Da	ate: / /		
Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive any insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent insurance act, which is a crime and subjects such person to criminal and civil penalties.						

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