

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.)

non-preferred. The preferred 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** Does patient have other coverage? Aetna Member ID #: ☐ 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: Phone: St Lic #: NPI#: DEA #: UPIN: Fax: Provider E-mail: Office Contact Name: Phone: D. DISPENSING PROVIDER/ADMINISTRATION INFORMATION **Dispensing Provider/Pharmacy:** Place of Administration: ☐ 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): TIN: ____ Address: _____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 and failure, intolerance, or contraindication to docetaxel or conventional paclitaxel? Please explain if there are any medical reason(s) that the patient cannot use docetaxel or conventional 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 Ohio MMP:

1-855-734-9389 PHONE: 1-855-364-0974

For other lines of business:

Note: Abraxane and generic paclitaxel (protein bound) are

Please use other form.

FΔX·

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 Ohio MMP:

FAX: 1-855-734-9389 **PHONE:** 1-855-364-0974

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.

atient First Name		Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL	INFORMATION (continued) – Re	equired clinical information must be comple	ted in its entirety for all precertific	cation requests.			
	Cervical cancer as a single agent 2nd line therapy Local/regional recurrence OR distant metastases						
	☐ Intrahepatic/Extrahepatic cholangiocarcinoma in combination with gemcitabine as primary treatment						
	☐ Unresectable disease Ol						
	Cutaneous melanoma as a single agent second line/subsequent therapy with performance status of 0-2 for						
	☐ Unresectable disease OR ☐ metastatic disease						
_		ression OR 🗌 after maximum clinical be	enefit from BRAF targeted thera	ару			
Ш	☐ 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						
	☐ receiv	ved prior external beam radiation therapy	y (EBRT) to the site of recurren	ce			
		a, clear cell carcinoma, serous carcinom		ntiated carcinoma			
	As primary treatment for disease not suitable for primary surgery						
	☐ As ac	Iditional treatment for disease suitable fo					
		With vaginal brachytherapy fro Stag		IV disease			
		ngle agent with histologic grade 3 tumors		thereny (EDDT)			
		se with vaginal brachytherapy and/or sec e with sequential external beam radiatio		т тегару (ЕВКТ)			
	☐ Adjuvant treatment as si		ir trierapy (LBIXT)				
	Stage IIIA-IVA						
	Epithelial Ovarian Cancer for p						
		ith carboplatin for persons with confirme	d taxane hypersensitivity				
	Fallopian tube cancer for persi						
		ith carboplatin for persons with confirme					
		SCLC) for recurrent or metastatic dise	ase as a single agent for perf	ormance status 2 OR in			
	combination with carboplatin f	or performance status 0-2					
	☐ 1st Line therapy	OS1, BRAF, and PD-L1 negative or unk	nown DRRAE V600E-mutati	on positive tumors			
	Subsequent therapy for	OOT, DIVAL, and I D-LI negative of unki	TIOWIT DIVAL VOOCE-ITIGLALIS	on positive turnors			
	_ , ,,	nutation positive tumors					
		n positive and prior erlotinib/afatinib/gefit	inib/osimertinib therapy				
		mors and prior crizotinib/ceritinib/alectin					
		ement positive tumors and prior crizotin					
	-	(≥50%) tumor, EGFR, ALK, ROS1, and		or pembrolizumab therapy.			
		CLC) 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.)

For Ohio MMP:

FAX: 1-855-734-9389 **PHONE:** 1-855-364-0974

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 - Required clinical	information must be completed for ALL precertification	on 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						
greate <u>r</u> 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 and 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 ☐ M		ilciupy ioi					
☐ 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 ☐ Unresecta	able disease						
For Continuation of Therapy: (clinical docume	ntation required):						
Is this a continuation request a result of the patient receiving samples of Abraxane® (paclitaxel protein-bound particles)?							
Is there clinical documentation supporting disease stability? Yes No							
Is there clinical documentation supporting disease improvement? \Begin{array}{c} Yes \Boxed{D} No							
H. ACKNOWLEDGEMENT							
Request Completed By (Signature Required	d):	Date	:				
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.