

MEDICARE FORM

Herceptin[®] (trastuzumab), Herceptin Hylecta[™] (trastuzumab and hyaluronidase-oysk), Herzuma (trastuzumab-pkrb), Kadcyla® (ado-trastuzumab), Kanjinti (trastuzumab-anns), Ogivri (trastuzumabdkst), Ontruzant (trastuzumab-dttb), Perjeta® (pertuzumab) and Trazimera (trastuzumab-qyyp)

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

Precertification Request

Please use other form. Note: Herzuma, Ogivri, and Ontruzant are non-preferred. The preferred products are Herceptin, Herceptin Hylecta, Kanjinti, and Trazimera.

For other lines of business:

1-855-320-8445

PHONE: 1-866-600-2139

For Illinois MMP:

FAX:

Please indicate:	☐ Start of treatment:	·							
D		rapy: Date of last treat	tment		-		_		
	equested By:			Pho	ne:		Fax:		
A. PATIENT INFOR	MATION		Los	st Name:					
Address:			City				State:	ZIP:	
Home Phone:		Work Phone:	City	y.	-	Cell Phone:	State.	ZIF.	
DOB:	Allergies	Work Priorie.							
	Allergies:	Long	11.1.14		1	-mail:			
	lbs or	kgs	Height:	inches	s or	cms			
B. INSURANCE INF		Doos natio	ant have other	or coverage?	□ v/	os 🗆 No			
	#:		Does patient have other coverage? If yes, provide ID#			Carrier Name:			
Insured:			, ido 15 //		ouiii				
C. PRESCRIBER IN									
First Name:		Last Name	e:			(Check On	e): 🔲 M.D.	☐ D.O. ☐ N.P. ☐	P.A.
Address:		.		City:			State:	ZIP:	
Phone:	Fax:	St Lic #:		NPI #:		DEA #:		UPIN:	
Provider Email:	'	Office Contact	Name:			Phone:		Ш	
D. DISPENSING PI	ROVIDER/ADMINISTRA	TION INFORMATION							
Place of Administr ☐ Self-administere ☐ Outpatient Infus	s Office					/: ☐ Retail Pharmacy ☐ Other			
	me:			Name:					
Home Infusion C		:							
Address:	ime:								
	ode(s) (CPT):								
E. PRODUCT INFOR									
Request is for: Herceptin (trastuzumab) Perjeta (pertuzumab) Kadcyla (ado-trastuzumab emtansine) Ogivri (trastuzumab-dkst) Ontruzant (trastuzumab-dttb) Herzuma (trastuzumab-pkrb) Herceptin Hylecta (trastuzumab and hyaluronidase-oysk) Kanjinti (trastuzumab-anns) Trazimera (trastuzumab-qyyp)									
Dose:		Frequency:					PCS Code:		_
	DRMATION – Please indic						_		
G. CLINICAL INFORMATION – Required clinical information must be completed in its entirety for all precertification requests.									
For All Requests (clinical documentation required): Yes No Does the patient have HER2 protein overexpression documented by one of the following? Check all that apply: Immunohistochemistry (IHC) Assay level of 3+ Results Positive Fluorescent in situ hybridization (FISH) HER2 gene copy of greater than 6 signals/nucleus Results Positive Fluorescent in situ hybridization (FISH) HER2 gene/ chromosome 17 ratio greater than or equal to 2.0									
	Results			Date of ⁻	Test:	1 1	<u> </u>		
Preferred products	ivri, and Ontruzant are n may vary based on indi	cation.		•		ptin Hylecta,	Kanjinti, and	d Trazimera.	
Yes No Has	the patient had prior thera the patient had a trial and Herceptin (trastuzumab Trazimera (trastuzumab	failure, intolerance, or co	ontraindicatio	n to any of the fo	ollowing			·	
								Continued on next	nanc



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION (continued) –	Required clinical information must be	completed in its entirety for all p	precertification requests.
Please explain if there are any other medical rediagnosis (select all that apply)	eason(s) that the patient cannot use a	any of the following preferred pro	oducts when indicated for the patient's
HERCEPTIN (trastuzumab): Esophageal adenocarcinoma Gastri Yes No Will Herceptin (trastuzuma Yes No Will Herceptin (trastuzuma Please provide the name	ab) be used as palliative therapy?		oma
Endometrial carcinoma ☐ Yes ☐ No Does the patient have advance ☐ Yes ☐ No Does the patient have a docur ☐ Yes ☐ No Does the patient have recurre ☐ Yes ☐ No Will Herceptin (trastuzumab) b	mented diagnosis of uterine serous cant disease?		
Salivary gland tumors Yes No Does the patient have recurre Please indicate how Herceptin (trastuzumab) v	vill be used: ☐ single agent ☐ Oth		systemic chemotherapy:
Yes No Will Hercel Please: No Company Note Note Note Note Note Note Yes No Will Hercel	nt, metastatic, stage IV disease or let atment)?	otomeningeal metastases from be metastatic disease stage IV tases from breast cancer (as intrerative (neoadjuvant) systemic the systemic that the system of	reast cancer disease acerebrospinal fluid treatment) nerapy? e used:
HERCEPTIN HYLECTA (trastuzumab and h HER2 positive breast cancer Please select which of the following applies to Early stage HER2-overexpressing breast cancer Yes No Will Herceptin H Metastatic HER2-overexpressing breast cancer Other	the patient's disease stage: ancer lylecta (trastuzumab and hyaluronida ncer	se-oysk) be used as adjuvant th	erapy?
Please select: Preoperative (neoadjuvant) therapy Please select in which of the foll Node-positive disea Individuals who des Locally advanced d	both drugs are documented in sec (pertuzumab) and Herceptin (trastuzudisease node-positive or at high-risk for Node-positive At high-risk for the Nowing settings Perjeta (pertuzumab) ase likely to become node-negative wire breast preservation and fulfill criterisease None of the above	umab) is being used for: or recurrence? recurrence Other: with Herceptin (trastuzumab) wil ith pre-operative systemic theraperia for breast-conserving surger	l be used:
☐ Yes ☐ No Does the patien	the patient's disease: ☐ Recurrent of t have symptomatic visceral disease : ☐ Symptomatic visceral disease	or visceral crisis?	



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB						
$\textbf{G. CLINICAL INFORMATION} \ \textit{(continued)} - \\$	Required clinical information must be	completed in its <u>entirety</u> for all pr	ecertification requests.						
KADCYLA (ado-trastuzumab emtansine):									
Yes Yes	at being treated for HER2-positive recul No Will Kadcyla (ado-trastuzumab et No Will Kadcyla (ado-trastuzumab et Has the patient received neoadjust and trastuzumab? Please provide the date range of Does the patient have a residual icate which applies: Tecurrent breast No Does the patient have symptomaticate which applies: Please indicate the type of breast complete in Nonsteron Steroidatic St	rrent or metastatic breast cancer mansine) be used as adjuvant uvant therapy containing a taxar of use:	systemic therapy? ne (with or without anthracycline) / / Ivant therapy? cancer crisis? r- negative						
For Continuation Requests (clinical docume									
HERCEPTIN (trastuzumab): For HER2-positive breast cancer only: Yes No Is there clinical evidence of dis Please provide initial start da									
HERCEPTIN HYLECTA (trastuzumab and hy Yes No Will Herceptin Hylecta (trastuz	rumab and hyaluronidase-oysk) be use rt date:/	ed in adjuvant settings?							
PERJETA (pertuzumab) with HERCEPTIN (t	stant metastatic disease?								
KADCYLA (ado-trastuzumab emtansine): Yes □ No Is Kadcyla (ado-trastuzumab e	etastatic disease?	vith Herceptin (trastuzumab), Ty	kerb (lapatinib), or Perjeta (pertuzumab)?						
Request Completed By (Signature Requi	red):		Date: / /						
Any person who knowingly files a request fo insurance company by providing materially	r authorization of coverage of a med	ical procedure or service with t	the intent to injure, defraud or deceive any						

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.