

## **MEDICARE FORM**

Herceptin® (trastuzumab), Herceptin Hylecta<sup>TM</sup> (trastuzumab and hyaluronidase-oysk), Herzuma (trastuzumab-pkrb), Kadcyla® (ado-trastuzumab), Kanjinti (trastuzumab-anns), Ogivri (trastuzumab-dkst), Ontruzant (trastuzumab-dttb), Perjeta® (pertuzumab) and Trazimera (trastuzumab-qyyp) Precertification Request

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

FAX: 1-833-322-0034 PHONE: 1-844-362-0934 For other lines of business: Please use other form.

For New Jersey HMO D-SNP:

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

	☐ Start of treatment: S☐ Continuation of ther			1	1				
Precertification Re	quested By:				Phone:		Fax:	:	
A. PATIENT INFORM	MATION								
First Name:			L	.ast	Name:		1	<b>.</b>	
Address:			C	City:		T	State:	ZIP:	
Home Phone:		Work	Phone:			Cell Phone:			
DOB:	Allergies:					E-mail:			
Current Weight:	lbs or	kgs	Height: _		inches or	cms	i		
B. INSURANCE INFO									
Aetna Member ID #:			Does patient have other coverage?						
				If yes, provide ID#: Carrier Name:					
Insured:	CORMATION		Insured:						
C. PRESCRIBER INI First Name:	FORMATION		Last Name:			(Check Or	ы). П M D	. 🔲 D.O. 🔲 N.P	
Address:			Last Name.		ity:	(Check Of	State:	ZIP:	. <u></u>
	Fax:		St Lic #:	_	PI #:	DEA #.	State.	UPIN:	
Phone:	rax.	Offi		IN	PI #:	DEA #:		UPIN:	
Provider Email:	ROVIDER/ADMINISTRA		ce Contact Name:			Phone:			
☐ Home Infusion C Agency Nat Address:	d Physician's on Center Phone: ne: center Phone: me:			_	Dispensing Provi Physician's Of Specialty Phar Name: Address: Phone: TIN:	fice [ macy [	Retail Ph Other Fax:		
E. PRODUCT INFOR	RMATION								
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)									
	, , , , , , , , , , , , , , , , , , , ,	•	ency:			нс	PCS Code:	:	
F. DIAGNOSIS INFO	RMATION - Please indica								
Primary ICD Code:		Second	dary ICD Code:			Other ICD (	Code:		
G. CLINICAL INFOR	MATION – Required clinic	al informatio	n must be completed i	n its	entirety for all prece	rtification reque	sts.		
For All Requests (clinical documentation required):    Yes									
	<u> </u>		•		-			Continued on a	



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION (conti	<b>nued)</b> – Required clinical information mu	st be completed in its <u>entirety</u> for all	precertification requests.
Please explain if there are any other n	nedical reason(s) that the patient cannot		
diagnosis (select all that apply)  ☐ Herceptin (trast	uzumab) 🔲 Herceptin Hylecta (trastuzu	ımab and hyaluronidase-oysk) 🔲 ŀ	Kanjinti (trastuzumab-anns)
☐ Trazimera (tras	tuzumab-qyyp)	, , , –	,
HERCEPTIN (trastuzumab):			
	☐ Gastric adenocarcinoma ☐ Esopha astuzumab) be used as palliative therapy′		noma
☐ Yes ☐ No Will Herceptin (tra	stuzumab) be used in combination with s	systemic chemotherapy?	
	the name of the systemic chemotherapy	·	
Endometrial carcinoma  ☐ Yes ☐ No Does the patient have	e advanced (stage III/IV) disease?		
☐ Yes ☐ No Does the patient have	e a documented diagnosis of uterine sero	us carcinoma?	
☐ Yes ☐ No Does the patient have	e recurrent disease? zumab) be used in combination with carb	oplatin and paclitaxel?	
Salivary gland tumors	,		
	e recurrent disease with distant metastas		
Please indicate now Herceptin (trastuz	zumab) will be used:   single agent   in combination w	ith systemic chemotherapy: Name o	f systemic chemotherapy:
HER2 positive breast cancer			
	e recurrent, metastatic, stage IV disease of Ifluid treatment)?     recurrent disease		
	☐ leptomeningeal m	etastases from breast cancer (as int	racerebrospinal fluid treatment)
	ill Herceptin (trastuzumab) be used as pr Please select in which of the following se		
	<ul><li>☐ Node-positive disease likely to beco</li><li>☐ Locally advanced disease</li><li>☐ Indiv</li></ul>		
	☐ None of the above	iduais wilo idilili criteria foi breast-co	onserving surgery except for turnor size
	ill Herceptin (trastuzumab) be used as ad ill Herceptin (trastuzumab) be used as pa	, , ,	
HERCEPTIN HYLECTA (trastuzuma		int of a complete treatment regiment:	
HER2 positive breast cancer	<u> </u>		
Please select which of the following a Early stage HER2-overexpressing	pplies to the patient's disease stage: breast cancer		
	rceptin Hylecta (trastuzumab and hyalurc	nidase-oysk) be used as adjuvant th	nerapy?
Other	neast cancel		
PERJETA (pertuzumab) with HERC			
"	ions for both drugs are documented ir Perjeta (pertuzumab) and Herceptin (tras	, .	cancer
Adjuvant therapy	ationt's disease node positive or at high	rial for requirement	
	atient's disease node-positive or at high- e select: ☐ Node-positive ☐ At high-ris		
Preoperative (neoadjuvant) therap	y of the following settings Perjeta (pertuzum	iah) with Hercentin (trastuzumah) wi	Il he used:
☐ Node-posit	ive disease likely to become node-negati	ve with pre-operative systemic thera	ру
_ =	who desire breast preservation and fulfill ranced disease \quid \text{None of the above}	criteria for breast-conserving surger	ry except for tumor size
Other	<del>_</del>		
	pplies to the patient's disease:		
	e specify: Symptomatic visceral disea		



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Dational First Name	D-#414 N	D-tit Db	D-4:4 DOD
Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION (continued) – I	Required clinical information must be	completed in its entirety for all pr	recertification requests
KADCYLA (ado-trastuzumab emtansine):	required clinical information must be	completed in its <u>entirety</u> for all pr	ecertification requests.
Yes   Yes	t being treated for HER2-positive rect No Will Kadcyla (ado-trastuzumab e No Has the patient received neoadj and trastuzumab? Please provide the date range o No Does the patient have a residua cate which applies:  Tecurrent brea No Does the patient have symptom Please indicate the type of bre  Yes No Is the breast o Please sele Nonster Steroida Estroge ER dow Androge Please specify:  Symptomati	urrent or metastatic breast cance emtansine) be used as adjuvant uvant therapy containing a taxar of use: / to I disease after receiving neoadjust cancer metastatic breast atic visceral disease or visceral disease in Unknown Other contains of the following endocripoidal aromatase inhibitors (anastal aromastase inhibitors (exemes in receptor (ER) antagonists (tamin-regulators (fulvestrant) Highers (fluoxymesterone) Other ic visceral disease visceral disease	systemic therapy?  the (with or without anthracycline)  / / Ivant therapy? cancer crisis? r- negative  Hormone receptor-positive ner erapy? ne therapy the patient is refractory to: crozole and letrozole) tane) looxifen or toremifene) gh-dose estrogen (ethinyl estradiol) The Please explain: crisis
☐ Yes ☐ No Will Kadcyla	No Will Kadcyla (ado-trastuzumab e (ado-trastuzumab emtansine) be use ertuzumab)?	emtansine) be used as a single a	igent?
For Continuation Requests (clinical docume  Yes No Has the patient experienced di  Please indicate: Disease			
HERCEPTIN (trastuzumab):  For HER2-positive breast cancer only:  Yes No Is there clinical evidence of dist  Please provide initial start da			
HERCEPTIN HYLECTA (trastuzumab and hy Yes No Will Herceptin Hylecta (trastuz	umab and hyaluronidase-oysk) be us t date:/	ed in adjuvant settings?	
PERJETA (pertuzumab) with HERCEPTIN (to	stant metastatic disease?		
KADCYLA (ado-trastuzumab emtansine):	etastatic disease?	with Herceptin (trastuzumab), Ty	kerb (lapatinib), or Perjeta (pertuzumab)?
Request Completed By (Signature Require	red):		Date: / /
Any person who knowingly files a request for insurance company by providing materially	authorization of coverage of a med		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.