

MEDICARE FORM

Herceptin® (trastuzumab), Herceptin HylectaTM (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

Page 1 of 3

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

Virginia (HMO D-SNP) FAX: 1-833-280-5224 PHONE: 1-855-463-0933

For other lines of business: Please use other form.

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

Please indicate: Start	of treatment: Start	t date _	/ / of last treatment						
Precertification Requested						e:		Fax:	
A. PATIENT INFORMATION									
First Name:				Last	Name:				
Address:				City:				State:	ZIP:
Home Phone:		Work	Phone:			(Cell Phone:	I	
DOB:	Allergies:					E	E-mail:		
Current Weight:	· ·	kgs	Height:		inches o	or	cms		
B. INSURANCE INFORMATION			g			_			
Aetna Member ID #:			Does patient have	other	coverage?	□ Y	es 🗌 No		
Group #:				Carrier Name:					
Insured:			Insured:						
C. PRESCRIBER INFORMATI	ION								
First Name:			Last Name:				(Check One	e): 🔲 M.D.	□ D.O. □ N.P. □ P.A
Address:				С	ity:			State:	ZIP:
Phone:	Fax:		St Lic #:	N	PI #:		DEA #:		UPIN:
Provider Email:		Offi	ce Contact Name:				Phone:		
D. DISPENSING PROVIDER	R/ADMINISTRATIO	N INFO	RMATION						
Place of Administration: Self-administered Physician's Office Outpatient Infusion Center Phone: Center Name: Home Infusion Center Phone:				<u> </u>	Dispensing Provider/Pharmacy: ☐ Physician's Office ☐ Retail Pharmacy ☐ Specialty Pharmacy ☐ Other Name: Address:				
Agency Name:									
Address: Administration code(s) (CPT):					Phone: Fax:				
E. PRODUCT INFORMATION									
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: HCPCS Code:									
F. DIAGNOSIS INFORMATION	N – Please indicate p	rimary I	CD Code and specify	any c	other where appl	icable			
Primary ICD Code:		Secon	dary ICD Code:				Other ICD C	ode:	
G. CLINICAL INFORMATION – Required clinical information must be completed in its entirety for all precertification requests.									
Re	hat apply: histochemistry (IHC) esults Fluorescent in situ hy esults Fluorescent in situ hy esults Ontruzant are non-p based on indicatio t had prior therapy with	Assay loybridizate ybridizate ybridizate referred n. th Herzu re, intole	evel of 3+ tion (FISH) HER2 ger tion (FISH) HER2 ger The preferred produma, Ogivri, or Ontruza	ne cop ne/ chi ducts ant wit ation	Date of Tearly of greater than Date of Tearly Herceptin, I hin the last 365 dto any of the follow	st: n 6 sig st: tio gre st: Herce lays? owing	/ / eater than or eq / / eptin Hylecta, I	Kanjinti, and	
☐ Trazime	ra (trastuzumab-qyyp	o)							Continued on next nee



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB				
G CLINICAL INFORMATION (continued) -	Required clinical information must	he completed in its entirety for all pro	ecertification requests				
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests. Please explain if there are any other medical reason(s) that the patient cannot use any of the following preferred products when indicated for the patient's diagnosis (select all that apply) Herceptin (trastuzumab) Herceptin Hylecta (trastuzumab and hyaluronidase-oysk) Kanjinti (trastuzumab-anns) Trazimera (trastuzumab-qyyp)							
HERCEPTIN (trastuzumab):							
□ Esophageal adenocarcinoma □ Gastric adenocarcinoma □ Esophageal-gastric junction adenocarcinoma □ Yes □ No Will Herceptin (trastuzumab) be used as palliative therapy? □ Yes □ No Will Herceptin (trastuzumab) be used in combination with systemic chemotherapy? → Please provide the name of the systemic chemotherapy: □ Yes □ No Will Herceptin (trastuzumab) be used in combination with systemic chemotherapy?							
Endometrial carcinoma ☐ Yes ☐ No Does the patient have advanc ☐ 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 nt disease?						
Salivary gland tumors							
Yes No Does the patient have recurre Please indicate how Herceptin (trastuzumab) v	vill be used: ☐ single agent ☐ C		vstemic chemotherany				
Yes No Will Hercen Please: No Company Nor Yes No Will Hercen Nor Yes No Will Hercen	nt, metastatic, stage IV disease or atment)?	leptomeningeal metastases from bre ☐ metastatic disease ☐ stage IV d astases from breast cancer (as intrac operative (neoadjuvant) systemic the tings Herceptin (trastuzumab) will be e node-negative with pre-operative s uals who fulfill criteria for breast-cons	east cancer lisease cerebrospinal fluid treatment) erapy? used:				
HERCEPTIN HYLECTA (trastuzumab and hy	yaluronidase-oysk):						
HER2 positive breast cancer Please select which of the following applies to ☐ Early stage HER2-overexpressing breast c ☐ Yes ☐ No Will Herceptin F ☐ Metastatic HER2-overexpressing breast ca	ancer lylecta (trastuzumab and hyaluron	idase-oysk) be used as adjuvant ther	rapy?				
PERJETA (pertuzumab) with HERCEPTIN (t	rastuzumah):						
(please ensure dosing and instructions for Please select which type of treatment Perjeta ☐ Adjuvant therapy ☐ Yes ☐ No Is the patient's of	both drugs are documented in s (pertuzumab) and Herceptin (trastudisease node-positive or at high-ris	uzumab) is being used for:	ncer				
Please select: Node-positive At high-risk for recurrence Other: Other:							
Please select in which of the following settings Perjeta (pertuzumab) with Herceptin (trastuzumab) will be used: Node-positive disease likely to become node-negative with pre-operative systemic therapy Individuals who desire breast preservation and fulfill criteria for breast-conserving surgery except for tumor size Locally advanced disease None of the above							
	the patient's disease: Recurrent have symptomatic visceral disease: Symptomatic visceral disease	se or visceral crisis?					



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION (continu	(ad) - Required clinical information mus	et he completed in its entirety for all	nrecertification requests			
KADCYLA (ado-trastuzumab emtansi		st be completed in its <u>entirety</u> for all	precertification requests.			
Yes No Does the patient have a documented diagnosis of HER2-positive non-small cell lung cancer? Yes No Is the patient being treated for HER2-positive recurrent or metastatic breast cancer? Yes No Will Kadcyla (ado-trastuzumab emtansine) be used as adjuvant systemic therapy? Has the patient received neoadjuvant therapy containing a taxane (with or without anthracycline) and trastuzumab? Please provide the date range of use:						
For Continuation Requests (clinical do ☐ Yes ☐ No Has the patient experien → Please indicate: ☐ Di		ble toxicity while on HER2 therapy?				
HERCEPTIN (trastuzumab): For HER2-positive breast cancer only Yes No Is there clinical evidence Please provide initial s	: of distant metastatic disease?	•				
HERCEPTIN HYLECTA (trastuzumab Yes No Will Herceptin Hylecta (Please provide the init	trastuzumab and hyaluronidase-oysk) b	ne used in adjuvant settings?				
PERJETA (pertuzumab) with HERCEF Yes No Is there clinical evidence Please provide initial s						
Yes No Is there clinical evidence Please provide initial s	umab emtansine) being used concomita e of metastatic disease?	antly with Herceptin (trastuzumab), 1	Гукегb (lapatinib), or Perjeta (pertuzumab)?			
H. ACKNOWLEDGEMENT						
Request Completed By (Signature			Date: /			
			n 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.