<b>◆</b> aet	Hercep (trastu (trastu Kanjin dkst), (pertuz Precer Page 1 o	MEDICARE FORM Herceptin <sup>®</sup> (trastuzumab), Herceptin Hylecta <sup>™</sup> (trastuzumab and hyaluronidase-oysk), Herzuma (trastuzumab-pkrb), Kadcyla <sup>®</sup> (ado-trastuzumab), Kanjinti (trastuzumab-anns), Ogivri (trastuzumab- dkst), Ontruzant (trastuzumab-dttb), Perjeta <sup>®</sup> (pertuzumab) and Trazimera (trastuzumab-qyyp) Precertification Request Page 1 of 3 (All fields must be completed and legible for precertification review.)					For Medicare Advantage Part B: PHONE: 1-866-503-0857 FAX: 1-844-268-7263 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:		eatment: St	art date _									
Precertification R				of last treatment _	/				Fax:			
A. PATIENT INFOR									1 ux.			
First Name:					Last	Name:						
Address:					City:				State:		ZIP:	
Home Phone:			Work	Phone:			Ce	ell Phone:	1		-	
DOB:	Alle	ergies:					E-	mail:				
Current Weight:	lbs	or	kgs	Heigh	t:	inches or		cm	5			
B. INSURANCE INF	ORMATION											
Aetna Member ID				Does patient have		•		s 🗌 No				
Group #: Insured:				If yes, provide ID# Insured:		(	Carrie	r Name:				
C. PRESCRIBER IN												
First Name:	FORMATION			Last Name:				(Check O	ne): 🗆 M.D.		D.O. 🗌 N.P. 🔲	ΡA
Address:					С	Sity:		(00	State:	· 🖵 -	ZIP:	
Phone:	Fax	k.		St Lic #:		IPI #:		DEA #:		UP	'IN:	
Provider Email:		<u></u>	Offi	ce Contact Name:				Phone:				
D. DISPENSING P	ROVIDER/AD	MINISTRATI										
Place of Administr Self-administer Outpatient Infus	ration: ed 🛛 🗌 I	Physician's ( Phone: _	Office			Dispensing Pro	Office arma	cy [	☐ Retail Pha ☐ Other		-	
Home Infusion						Name:						
Agency Na						Address:						
Address:						Phone:						
Administration code(s) (CPT):					TIN:			PIN:				
	Herceptin (tra Ontruzant (tra	istuzumab-c uzumab-ani	lttb) 🔲 H ns) 🗌 Tra	a (pertuzumab)  [ lerzuma (trastuzu azimera (trastuzu ency:	mab-p mab-q	krb) 🗌 Hercept yyp)		lecta (tras		nd hy		
F. DIAGNOSIS INF	ORMATION - P	Please indicate					able.					
Primary ICD Code:			Secon	dary ICD Code:			(	Other ICD	Code:			
	linical docume es the patient ha Check all that a ☐ Immunohistoo	ntation requ ave HER2 pro pply: chemistry (IH	<b>ired):</b> tein overe: C) Assay le	xpression document	ed by c	one of the following	l?	·	ests.			
[ r	] Positive Fluor	rescent in situ	ı hybridizat	tion (FISH) HER2 ge tion (FISH) HER2 ge	ene cop	by of greater than 6	signa	als/nucleus				
ا Note: Herzuma, Og Preferred products	Results Results	zant are non	-preferred			Date of Test:		/ /		nd Tra	azimera.	
	the patient had	a trial and fa astuzumab)	ilure, intole		ication	to any of the follow	/ing?			o-ann:	s)	



## MEDICARE FORM

Herceptin<sup>®</sup> (trastuzumab), Herceptin Hylecta<sup>™</sup> (trastuzumab and hyaluronidase-oysk), Herzuma (trastuzumab-pkrb), Kadcyla<sup>®</sup> (ado-trastuzumab), Kanjinti (trastuzumab-anns), Ogivri (trastuzumabdkst), Ontruzant (trastuzumab-dttb), Perjeta<sup>®</sup> (pertuzumab) and Trazimera (trastuzumab-qyyp) Precertification Request Page 2 of 3 
 For Medicare Advantage Part B:

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(All fields must be completed and legible for precertification review.)							
Patient First Name	Patient Last Name	Patient Phone	Patient DOB				
Please explain if there are any or diagnosis (select all that apply)	( <i>continued</i> ) – Required clinical information mus other medical reason(s) that the patient cannot u (trastuzumab) a (trastuzumab-qyyp)	se any of the following preferred pro	oducts when indicated for the patient's				
☐ Yes ☐ No Will Hercer ☐ Yes ☐ No Will Hercer	ma Gastric adenocarcinoma Esopha, tin (trastuzumab) be used as palliative therapy? tin (trastuzumab) be used in combination with sy rovide the name of the systemic chemotherapy:	vstemic chemotherapy?	ioma				
☐ Yes       ☐ No       Does the patient         ☐ Yes       ☐ No       Does the patient	nt have advanced (stage III/IV) disease? It have a documented diagnosis of uterine serou It have recurrent disease? trastuzumab) be used in combination with carbo						
	nt have recurrent disease with distant metastase trastuzumab) will be used:		f systemic chemotherapy:				
(as intracerebr → □ Yes □ 1 □ Yes □ 1	<ul> <li>at have recurrent, metastatic, stage IV disease of ospinal fluid treatment)? ☐ recurrent disease ☐ leptomeningeal metabolic Will Herceptin (trastuzumab) be used as pre</li> <li>Please select in which of the following se</li> <li>☐ Node-positive disease likely to becom</li> <li>☐ Locally advanced disease ☐ Individ</li> <li>☐ None of the above</li> <li>No Will Herceptin (trastuzumab) be used as adjio</li> <li>Will Herceptin (trastuzumab) be used as par</li> </ul>	☐ metastatic disease ☐ stage IV tastases from breast cancer (as intu- operative (neoadjuvant) systemic t titings Herceptin (trastuzumab) will the ne node-negative with pre-operative duals who fulfill criteria for breast-co uvant therapy?	/ disease racerebrospinal fluid treatment) herapy? be used: a systemic therapy nserving surgery except for tumor size				
HER2 positive breast cancer Please select which of the follow Early stage HER2-overexpre	Vill Herceptin Hylecta (trastuzumab and hyaluror	iidase-oysk) be used as adjuvant th	erapy?				
Please select which type of trea Adjuvant therapy Yes No Is Preoperative (neoadjuvant) Please select in v Node Indiv Loca	structions for both drugs are documented in Itment Perjeta (pertuzumab) and Herceptin (tras Is the patient's disease node-positive or at high-ris Please select: Node-positive At high-risk	tuzumab) is being used for: sk for recurrence? s for recurrence	l be used: py				
Yes 🗌 No 🛛	hich applies to the patient's disease: ☐ Recurre loes the patient have symptomatic visceral diseas Please specify: ☐ Symptomatic visceral diseas	ase or visceral crisis?					



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Herceptin<sup>®</sup> (trastuzumab), Herceptin Hylecta<sup>™</sup> (trastuzumab and hyaluronidase-oysk), Herzuma (trastuzumab-pkrb), Kadcyla<sup>®</sup> (ado-trastuzumab), Kanjinti (trastuzumab-anns), Ogivri (trastuzumabdkst), Ontruzant (trastuzumab-dttb), Perjeta<sup>®</sup> (pertuzumab) and Trazimera (trastuzumab-qyyp) Precertification Request Page 3 of 3

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

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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 <u>entirety</u> for all	precertification requests.
KADCYLA (ado-trastuzumab emtansine):         Yes       No         Yes       No         Is the patient         Yes       No         Is the patient         Yes         No         Is the patient         Is the patient      <	Imented diagnosis of HER2-positive normation in the sendence of the terms of the sendence of	on-small cell lung cancer? surrent or metastatic breast cance emtansine) be used as adjuvan ijuvant therapy containing a taxi of use:/to al disease after receiving neoad ast cancer [] metastatic brease hatic visceral disease or visceral east cancer: [] Hormone recepti [] Unknown [] C cancer refractory to endocrine the ect which of the following endoce roidal aromatase inhibitors (ana al aromastase inhibitors (exemption en receptor (ER) antagonists (tai vn-regulators (fulvestrant) [] H ens (fluoxymesterone) [] Other tic visceral disease [] visceral emtansine) be used as a single	cer? It systemic therapy? ane (with or without anthracycline) / / ljuvant therapy? st cancer I crisis? tor- negative Hormone receptor-positive Other herapy? crine therapy the patient is refractory to: I strozole and letrozole) estane) moxifen or toremifene) High-dose estrogen (ethinyl estradiol) er: Please explain: I crisis a agent?
For Continuation Requests (clinical docum □ Yes □ No Has the patient experienced			
HERCEPTIN (trastuzumab): For HER2-positive breast cancer only: Yes No Is there clinical evidence of di Please provide initial start d			
HERCEPTIN HYLECTA (trastuzumab and h Yes No Will Herceptin Hylecta (trastu Please provide the initial sta	zumab and hyaluronidase-oysk) be us	sed in adjuvant settings?	
PERJETA (pertuzumab) with HERCEPTIN ( Yes No Is there clinical evidence of d Please provide initial start of	istant metastatic disease?		
KADCYLA (ado-trastuzumab emtansine):         Yes       No       Is Kadcyla (ado-trastuzumab         Yes       No       Is there clinical evidence of m         Please provide initial start d	netastatic disease?	with Herceptin (trastuzumab), 7	⊺ykerb (lapatinib), or Perjeta (pertuzumab)?
H. ACKNOWLEDGEMENT			
Request Completed By (Signature Requ	ired):		Date: / /
Any person who knowingly files a request for	or authorization of coverage of a me	dical procedure or service with	n the intent to injure, defraud or deceive any

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.