## **Cancer Care** Ontario **Action Cancer** Ontario

# Trastuzumab (Biosimilar) with First Line Docetaxel - Metastatic Breast Cancer

This policy version is for trastuzumab biosimilars Herzuma, Kanjinti, Ogivri and Trazimera.

(This form must be completed <u>before</u> the first dose is dispensed.)

1. Patient Profile			,
* Surname:			
* Given Name:			
* OHIN:	* Chart Nu	mber:	
* Postal Code:			
* Height (cm):	* Weight (kg):	······	
* BSA (m <sup>2</sup> ):	* Gender:	O Male O Female O Other	
* Date of Birth:			
	Day Month Year		
* Site:			
* Attending Physician	n (MRP- Most Responsible Physician)		
Requested Prior Ap	pproval  Yes * Patient on Clini	cal Trial O Yes O No	
Other (specify):	<u></u>		
Specify Arm:			
<ul><li>Standard of care</li><li>Blinded / Unkno</li></ul>	· ·	erimental arm	
O Billided / Official	) VVIII		
Prior Approval F	Request		
* Select the appropria	ate		
prior approval			
scenario:			

	<ul> <li>and clinic note)</li> <li>2-Clinical document review (identify the patient history that needs to be reviewed against eligibility criteria in Additional Comments below)</li> </ul>	
	3-Regimen modification - schedule (complete questions a and b)	
	<ul> <li>4-Regimen modification - drug substitutions         (complete questions a and c)</li> <li>5-Withholding a drug in combination therapy</li> </ul>	
	from start of treatment (complete questions d, e and f)	
	<ul> <li>6-Maintenance therapy delay (submit clinic note)</li> <li>7-Prior systemic therapy clinical trials (complete question g)</li> <li>8-Modification due to supply interruption/drug</li> </ul>	
	shortage  Other (specify)	
	rting documentation must be submitted at the time of prior approval. Documentation may include clinic note, and/or CT scans.	а
a. Co-morbidities / toxic	ity / justification:	
a. Co-morbidities / toxic	eity / justification:	
a. Co-morbidities / toxic	eity / justification:	
<ul><li>a. Co-morbidities / toxic</li><li>b. Intended regimen schedule:</li></ul>	city / justification:	
b. Intended regimen	bity / justification:	
b. Intended regimen schedule:	bity / justification:	
<ul><li>b. Intended regimen schedule:</li><li>c. Intended regimen:</li></ul>		
<ul><li>b. Intended regimen schedule:</li><li>c. Intended regimen:</li><li>d. Drug(s) to be held:</li><li>e. Rationale for holding</li></ul>		

h. Anticipated date of first treatment:	Day Month	Year		
i. Additional comments:				
2. Eligibility Criteria	ì			
The patient must meet	the following crite	ria:		
a. Her2/neu Status				
Patient must test positi	ve for Her2/neu a	per Cancer Care Ontario criteria. A p	photocopy of the pathology report of the	
Her2/neu test must be	submitted to Can	er Care Ontario. The report must state	e clearly the hospital, date of biopsy and	
the hospital pathology	specimen number	of the original material used for the H	er2/neu test.	
Please specify t	ests used for dete	ction of Her2/neu:		
IHC 3+		☐ Yes		
	FISH/ SISH ≥ 2		☐ Yes	
b. Patient has metastatic breast cancer.			☐ Yes	
c. Patient will <u>not</u> be receiving an anthracycline.			Yes	
	·		e cost of docetaxel as part of this regimen and is included in the band level pricing.	
3. Precautions				
a. The patient has a norm	nal cardiac ejectio	ı fraction	☐ Yes	
b. There is no evidence of extensive lung involvement				
4. Funded Dose				
Therapy should be initi	ated with a loadin	g dose of 4 mg/kg, followed by 2 mg/k	g IV weekly.	

• Patients may be switched to 6 mg/kg IV q3 weeks after they are adequately loaded over a reasonable period of time with weekly dosing.

#### 5. Notes

a. Reimbursement will be discontinued for patients whose disease progresses while being treated with trastuzumab in the metastatic setting.

#### 6. FAQs

i. My patient is currently receiving trastuzumab (Herceptin). Can my patient stay on the reference biologic (i.e., trastuzumab (Herceptin))?

Yes, patients currently on trastuzumab (Herceptin) or initiated on trastuzumab (Herceptin) before December 15, 2019 may continue with the reference biologic until their treatment course has ended.

Patients who are continuing treatment with trastuzumab (Herceptin) after December 15, 2019 must have an enrolment form and treatment claim(s) submitted in eClaims prior to December 15, 2019 to be eligible for continued reimbursement of trastuzumab (Herceptin).

ii. My patient is currently receiving trastuzumab (Herceptin). Can my patient be switched to a trastuzumab biosimilar for the remainder of their treatment cycles?

At the discretion of the treating physician or based on individual hospital policy, patients currently on trastuzumab (Herceptin) may be switched over to a trastuzumab biosimilar for the remainder of the funded doses if a trastuzumab biosimilar is funded for the specific indication.

If the patient is already enrolled in an NDFP policy for trastuzumab, please re-enroll the patient in the updated trastuzumab enrolment form in order to submit treatments for trastuzumab biosimilars.

iii. Will the reimbursement rate be the same for all trastuzumab biosimilars?

Reimbursement in eClaims will be adjudicated based on the Drug Identification Number (DIN) submitted, and may be unique to each trastuzumab biosimilar.

Any incentives or promotions accepted by hospitals related to purchase and/or selection decisions of trastuzumab will be interpreted as lowering the local acquisition cost of trastuzumab, and, consistent with other NDFP policies, are subject to recovery.

### 7. Supporting Documents

To ensure reimbursement of your claim, both the completed enrolment form and a copy of the required documentation (where applicable) must be submitted through CCO e-Claims.

In the event of an audit, the following should be available to document eligibility:

A clinic note detailing treatment history and, if requested, MAR confirming treatment was given in combination.

Signature of Attending Physician (MRP-Most Responsible Physician):				
	Day	Month		

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