Eligibility Form

Raltitrexed - Metastatic Esophageal, Gastroesophageal Junction, or Gastric Cancer

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

1. Patient Profile				
* Surname: * Given Name:				
* OHIN:	* Chart Number:			
* Postal Code:				
* Height (cm):	* Weight (kg):			
* BSA (m ²):	* Gender: O Male O Female O Other			
* Date of Birth:	Day Month Year			
* Site:				
* Attending Physician	(MRP- Most Responsible Physician):			
Requested Prior App	oroval ☐ Yes * Patient on Clinical Trial ○ Yes ○ No			
Other (specify):	<u></u>			
Specify Arm: Standard of care Blinded / Unknown	·			
Prior Approval R	equest			
Select the appropriate prior	1-Unknown primary (submit pathology report			
approval scenario:	O 3-Regimen modification - schedule (complete			
	○ 5-Withholding a drug in combination therapy ○ 6-Maintenance therapy delay (submit clinic note from start of treatment (complete questions d, e and f)			
	 7-Prior systemic therapy clinical trials (comple 8-Modification due to supply interruption/drug question g) Other (specify) 			

pathology report,	ciiiic iiote, aiiu/oi	OT Scalis.		
a. Co-morbidities / toxid	city / justification:			
b. Intended regimen schedule:	<u></u>			
c. Intended regimen:				
d. Drug(s) to be held:				
e. Rationale for holding drug(s):				
f. Intention to introduce drug at a later date?	☐ Yes			
g. Prior clinical trial identifier (e.g., NCT ID, trial name) and treatment description (e.g., arm, drug/regimen):				
h. Anticipated date of first treatment:				
	Day Month	Year		
i. Additional comments	3:			

All relevant supporting documentation must be submitted at the time of prior approval. Documentation may include a

2. Eligibility Criteria

 Raltitrexed will be used to treat pati cancer 	ents with metastatic esophageal, gastroeso	phageal junction, or gastric
b. The patient has complete dihydropy	○ Yes ○ No	
☐ lives more than 60 km from the	toxicity with fluorouracil chemotherapy, and/ treatment centre/hospital, and/or s (e.g., ambulance or special vehicle)	or
3. Baseline Information		
ECOG Performance Status at the time of enrolment	O 0 O 1 O 2	
b. The patient has metastatic cancer	○ Esophageal○ Gastroesopha○ Gastric	geal junction
c. Raltitrexed will be used in combination with	N/A (using monotherapy) Irinotecan	Oxaliplatin
d. If the patient has a complete DPD deficiency, please indicate both DPYD genetic variants	○ c.1905+1G>A (*2A)○ c.1129-5923C>G, c.1236G>A (HapB3)○ Other	○ c.1679T>G (*13) ○ c.2846A>T
If other, please specify		
	c.1905+1G>A (*2A)c.1129-5923C>G, c.1236G>A (HapB3)Other	○ c.1679T>G (*13)○ c.2846A>T
If other, please specify		
e. Is the patient transitioning from a non-publicly funded program?	○ Yes ○ No	
f. If yes to 3e, please indicate the date of the last administered dose	Day Month Year	
4. Funded Dose		
Raltitrexed 3 mg/m ² intravenously (IV) every 21 days.	
Treatment should continue until disc	ease progression or unacceptable toxicity, w	hichever comes first.
[ST-QBP regimen codes: RALT, OX	ALRALT, IRINRALT]	
5. Notes		

1. As per the Clinical Pharmacogenetics Implementation Consortium (CPIC) Guideline (2017), <i>DPYD</i> poor metabolizers are defined as a patient carrying two no function alleles OR a patient carrying one no function allele plus one decreased function allele. Patients with a <i>DPYD</i> poor metabolizer phenotype have complete DPD deficiency.
6. FAQs
1. My patient is currently receiving raltitrexed through non-publicly funded means (e.g., patient support program, private insurance). Can my patient be transitioned to receive funding for raltitrexed through the New Drug Funding Program (NDFP)?
Provided the eligibility criteria were met at the time of treatment initiation and the patient's disease has not progressed, your patient may be eligible for continued coverage of raltitrexed through the NDFP.
Patients who meet the eligibility criteria may be transitioned to NDFP funding through a regular eClaims enrolment. If there is clinical uncertainty regarding eligibility, these requests may be submitted as a prior approval including a clinic note from the time of initiation as well as the most recent clinic note outlining the response to treatment (if able to assess).
5. Supporting Documents
None required at the time of enrolment.
 In the event of an audit or upon request, the following should be available to document eligibility: a clinic note detailing patient and treatment history/response. pharmacogenetic report showing DPYD genotyping result(s) including the specific variants (if applicable).
Signature of Attending Physician (MRP-Most Responsible Physician):

Day

Month Year

Form 1016