

## **Enrolment Form**

## CAR T-cell Therapy for Relapsed/Refractory Lymphoma (Third or Subsequent Line)

Note: This form should be completed and **funding approved** <u>before</u> apheresis is performed.

Completed form and supporting documentation should be submitted through the online portal: https://mft.cancercare.on.ca.

Username: CARTSubmission

Password: Contact our program at <a href="mailto:OH-CCO\_CARTSubmissions@ontariohealth.ca">OH-CCO\_CARTSubmissions@ontariohealth.ca</a>

Ontario Health collects and uses information on this form in order to determine if the patient meets the eligibility and funding criteria for the CAR T-cell Therapy Program, resulting in reimbursement to the treating facility. They also collect and use information on this form for purposes of analysis or compiling statistical information with respect to the management of, evaluation or monitoring of, the allocation of resources to or planning for all or part of the health system, including the delivery of services, pursuant to Section 45 of the Personal Health Information Protection Act, 2004.

As part of the evaluation of the request, it may be necessary for Ontario Health to disclose the patient's personal health information (PHI) to other administrative programs for health services and insured benefits at the Ministry of Health.

\*Required Fields

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1. Patient Profile									
*Surname:									
*Given Name:									
	Л-YYYY or c	lick arrow	down hutto	n to use ca	lendar to ei	nter the da	te)		
Sate of Shall	N-TITI OF C	nck arrow	aown batto	iii to use cu	ieriaar to er	iter the da			
*Gender:	her	Hei	ight (cm):			Weight			
*Province/Territory of Patient Residence:	○ AB	○ BC	○ MB	○ NB	$\bigcirc$ NL	○ NT	○ NS	$\bigcirc$ NU	ON
	○ PE	○ QC	○ SK	$\bigcirc$ YT					
*Postal Code of Patient Residence:									
*Provincial/Territorial Health Card Number:									
Note: If your patient is not a resident of Ontario, a fund	ling approva	al letter fro	om the patie	ent's provin	cial/territo	rial Ministi	ry of Health	is required.	
2. Enroling Site									
*Enroling Site:									
*Patient Chart Number (MRN) at Enroling Site:									_
*Enroling Physician:									
Enroling Physician CPSO Number (Ontario Only):									
*Enroling Physician Specialty:									
*Enroling Physician Email:									
*Enroling Physician Cell Phone Number:									
*Enroling Physician Fax Number:									
Alternate Contact Email:									
	Note: If an about this			ail is provid	ded, the alt	ernate con	tact will be	copied on a	ll email correspondence

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3. Treatment Centre and Product Information						
	kage. CAR T-cell Therapy Cer		has agreed to treat your patient. Email or fax confirmation is t details are available at <a href="https://www.cancercareontario.ca/en/">https://www.cancercareontario.ca/en/</a>			
*Will this patient receive CAR T-cell therapy in	Ontario?		○ No			
If patient will be treated <b>in Ontario</b> , select CAR T-cell therapy site:		Uravinski Cancer Centre - Hamilton Health Sciences				
		○ King	ston General Hospital - Kingston Health Sciences Centre			
		Princ	cess Margaret Cancer Centre - University Health Network			
		The	Ottawa Hospital			
If patient will be treated in another province of CAR T-cell therapy site name and city/province						
If patient will be treated <b>out of country</b> , please indicate the treating		Rosv	vell Park Comprehensive Cancer Center (Buffalo, New York)			
facility and also complete section 8:		Cleveland Clinic (Cleveland, Ohio)				
		○ Karn	nanos Cancer Institute (Detroit, Michigan)			
*Treating Physician at CAR T-cell therapy site:						
*Requested CAR T-cell therapy product:	<ul><li>Kymriah (tisagenlecleucel)</li><li>Yescarta (axicabtagene ciloleucel)</li></ul>					
	Breyanzi (lisocabtagene maraleucel)					
		require replacement of the original funding letter that was issued. se there is a need to use another product.				
Anticipated date of apheresis :		(DD-N	MMM-YYYY or click arrow down button to use calendar to enter the date)			
4. Funding Criteria						
*A. The patient must meet the following crite	eria:	ny patient	meets the funding criteria outlined below:			
<ul><li>aggressive lymphoma including an a anthracycline or etoposide containing</li><li>Patient is sufficiently stable to facilit</li></ul>	nti-CD20 monoclonal antibong chemotherapy regimen (eate planned CAR T-cell thera noneed for intubation or of performance status.	ody (unless e.g., R-CHO apy (e.g., n	lymphoma after two or more lines of systemic therapy for the tumor is determined to be CD20 negative) and an P) <sup>3</sup> ot rapidly progressing on temporizing therapy, no significant es not require ICU/pressors and does not have active or			
*B. Patient has the following diagnosis <sup>5,6</sup> :						
Notes: As evidence and clinical practice evolve, eligib	ility criteria is subject to change.	Additional r	notes are provided on page 4.			
further defined as per revised Lugano Response Crite	eria for Malignant Lymphoma (Cl	neson et al.,	sy-proven relapse or recurrence. Treatment responses are as follows and are 2014): ased response requirements or meets the complete radiological response			

- a) Complete response (CR) meets the complete metabolic response criteria as per PET scan-based response requirements or meets the complete radiological response criteria as per CT-scan based response requirements.
  - b) Stable disease (SD) or progressive disease (PD) as best response to first-line therapy after at least 3 or more cycles of first-line therapy (e.g., 3 cycles of R-CHOP)
- c) Partial response (PR) as best response after at least 6 cycles and biopsy-proven residual disease or disease progression ≤12 months following first line therapy
- 2. Primary refractory disease indicates progressive or stable disease as the best response to the first line standard therapy for aggressive lymphoma (e.g., R-CHOP). Refractory disease to second or greater line indicates progressive disease or partial response as best response to the most recent therapy regimen.
- 3. Patients must have received standard therapies (e.g., R-CHOP first line and platinum-containing salvage chemotherapy) to be considered for CAR T-cell therapy.
- 4. Patients with primary CNS lymphoma are currently not eligible for funding. For patients who experienced early or isolated CNS relapse or asynchronous systemic and CNS disease and have received or completed systemic and CNS disease treatments separately, standard therapy, or regimen for the treatment of active secondary CNS lymphoma (e.g., HD-methotrexate and cytarabine or MATRIX regimen) may be considered as a separate line of treatment.
- 5. Only diagnoses listed on section 4.B above may be eligible for funding. To be considered for funding, patients with transformations of indolent lymphomas or rare subtypes of large B-cell lymphomas (LBCL) must have received two or more lines of systemic therapy for aggressive lymphoma (i.e., DLBCL) as described on section 4.A above.
- 6. Patients with Richter's transformation from chronic lymphocytic leukemia (CLL) or small lymphocytic lymphoma (SLL) are currently not eligible for funding.

5. Treatment H	listory					
*A. How many lines of systemic therapy against aggressive lymphoma (e.g., DLBCL) has the patient previously received?						3 or More
*B. Did the patient have a previous autologous stem cell transplant (ASCT)?						○No
	de further details in th					
ii.If no, please	e indicate the reason f	or ineligibility or for not undergoing ASCT	:			
	If other, explain:					
Date Initiated	Date Completed	Name of Therapy/Regimen	No	o. of Cycles (if applicable)	Best Res	sponse to Therapy
				<u>, , , , , , , , , , , , , , , , , , , </u>		
*C. Did the patient	t have a previous allog	geneic stem cell transplant?	○ Yes	○ No	1	
i. If yes, provide	the date of the patier	nt's allogeneic stem cell transplant?		(Click arrow down bu	tton to use ca	llendar to enter the date)
		rsus host disease (GvHD)?	○ Yes			
If yes,	a. Does the patient ha	ave active GvHD?		○ No		
b. Is the patient still undergoing treatment for GvHD?			○ Yes	○ No		
*D. Did the patien	t receive any prior noi	n-cellular anti-CD19 therapy?	○ Yes	○ No		
If yes, i. Provi	de the date when the	patient received the therapy:		(Click arrow down bu	tton to use ca	llendar to enter the date)
ii. Speci	fy the non-cellular an	ti-CD19 therapy: O Blinatumomab	) Tafasitama	ab Other:		
6. Confirmation	n of Patient Suita	bility for Therapy				
*A. CNS disease st			O No C	CNS lymphoma		
			Prim	nary CNS lymphoma (not eligi	ble for CAR	T-cell therapy)
				ited secondary CNS lymphom	na - persiste	nt disease (active)
				ated secondary CNS lymphom	na - in remis	sion (inactive)
		cterial, viral (HIV, active hepatitis B or C)	○ No I	nfection		
or fungal infection	:		Cont	trolled Infection		
			OUnce	ontrolled Infection		
*C. Karnofsky Perf	ormance Status (KPS)	≤70%:		○ No		
	Da	te of KPS assessment:	(DD-N	MMM-YYYY or click arrow down b	utton to use	calendar to enter the date)
Renal Function:						
*D. Creatinine ≥14	1.44 μmol/L (1.6 mg/	dL):		○ No		
*E. Estimated glon	nerular filtration rate	(eGFR) ≤45 ml/min/1.73m²:		○ No		
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Liver Function:				
*F. ALT or AST ≥3x upper limit of normal value:		○ No		
*G. Bilirubin ≥2x upper limit of normal value:		○ No		
Pulmonary Function:				
*H. Pulse oxygenation ≤91% on room air:		○ No		
Cardiac Function:				
*I. Left ventricular ejection fraction (LVEF) ≤40% confirmed by echocardiogram or multiple-gated acquisition (MUGA) scan or radionuclide angiography:	Yes	○ No		
Bone Marrow Function:				
*J. Absolute neutrophil count (ANC) ≤1.0x10 <sup>9</sup> /L:		○ No		
*K. Absolute lymphocyte count (ALC) <0.1x10 <sup>9</sup> /L:	Yes	○No		
Note: If ALC is below 0.1x10 <sup>9</sup> /L, application can be considered; but for apheresis to proceed	d, ALC must be at least 0.1x10 <sup>9</sup> /L.			
*L. Hemoglobin ≤80 g/L (8.0 g/dL) and/or transfusion dependent:		○ No		
*M. Platelets ≤50x10 <sup>9</sup> /L:		○ No		
7. Additional Notes				
a. Treatment with tisagenlecleucel or axicabtagene ciloleucel or lisocabtagene maraleucel is a one-time therapy. b. Tisagenlecleucel or axicabtagene ciloleucel or lisocabtagene maraleucel should not be used in combination with other treatments for relapsed/refractory lymphoma. c. At least six weeks must have elapsed from any prior systemic inhibitory/stimulatory immune checkpoint molecule therapy (e.g., nivolumab, pembrolizumab, etc.) to the time of CAR T-cell product infusion. d. A patient with another malignancy may be considered for CAR T-cell therapy if they meet the funding criteria, are suitable for therapy, and are either in complete remission or not undergoing any active drug therapy that could cause serious toxicity and preclude them from receiving CAR T-cell therapy. e. Patients who have had an autologous stem cell transplant in the last 100 days must meet funding criteria at the time of enrolment. f. Patients who have had an allogeneic stem cell transplant and have no active graft versus host disease (GvHD) and are not on immunosuppressive therapy may be eligible for CAR T-cell therapy. g. For CNS lymphomas, active or persistent CNS disease is defined as recent neurologic sign/symptoms, and/or positive imaging studies (MRI, PET scan) and/or positive cerebrospinal fluid (CSF) study. h. Patients with an active, uncontrolled infection should not start treatment with CAR T-cell therapy until the infection has resolved or has been appropriately treated. This includes both the lymphodepleting chemotherapy and the CAR T-cell infusion. i. Patients must meet the funding criteria at the time of enrolment and must continue to be eligible and suitable for therapy at the time of product infusion.				
8. Out-of-Country Applications - Additional Requirements				
<ol> <li>Only complete this section if you are an Ontario physician applying for an Ontario patient to be treated out-of-country:</li> <li>Submit all the documents listed under "Supporting Documents" in section 10.</li> <li>Download, complete and submit the Ministry form "Request for Prior Approval for Full Payment of Insured Out-of-Country (OOC) Health Services."         The form can be found in the Central Forms Repository at: https://forms.mgcs.gov.on.ca/en/dataset/on00314</li> <li>Complete as indicated below:         <ul> <li>Part 1: Patient name, OHIN number, date of birth, mailing address and telephone number</li> <li>Part 2: Physician name, office address, telephone number, email address, and OHIP billing number</li> <li>Part 3: All required fields, check box confirming completion of CCO Questionnaire; in lieu of the questionnaire form, a completed copy of this enrolment form will be submitted</li> <li>Part 4: Auto-completed</li> <li>Part 5: All required fields</li> <li>Part 6: Submit a completed copy of this enrolment form</li> </ul> </li> </ol>				
9. Acknowledgement				
*Yes, I confirm that the patient named above, or relevant substitute decision-maker where applicable, consents that Ontario Health collects and uses information on this form to make funding decisions pursuant to section 38(1)(b) of the Personal Health Information Protection Act, 2004; and for the purpose of analysis or compiling statistical information with respect to the management of, evaluation or monitoring of, the allocation of resources to or planning for all or part of the health system, including the delivery of services, pursuant to section 45 of the Personal Health Information Protection Act, 2004. As part of the evaluation and reimbursement process for the CAR T-cell Therapy Program, it may be necessary for Ontario Health to disclose or share the patient's personal health information to other administrative programs for health services and insured benefits at the Ministry of Health or at Ontario Health.				

10. Supporting Documents						
the	ne enrolment is for an Out-of-Country treatment for an Ontario patient, the following documentation (from <b>Lists A and B</b> ) <b>must be</b> submitted with enrolment form. The Ministry form "Request for Prior Approval for Full Payment of Insured Out-of-Country (OOC) Health Services" must also be luded in the enrolment package.					
	ne enrolment is for in-Ontario treatment, the documents under <b>List A must be</b> submitted and documents under <b>List B</b> should be available upon usest (including for the purpose of audit) to confirm eligibility.					
*Lis	st A: Required upon enrolment					
	If any of the answers to section 6 are "Yes", submit relevant and recent laboratory results showing adequate organ function (e.g., kidney and liver function tests, viral serology, cardiac ECHO/MUGA)					
	Pathology report					
	Recent clinic notes that describe the patient's current clinical status and rationale for CAR T-cell therapy over other treatment options. Include any specialist notes (e.g., BMT, neurology, nephrology, cardiology) that informed the treatment plan					
	Bone Marrow (BM) studies including most recent studies					
	Pre- and post-treatment imaging reports e.g., CT scan (post-treatment imaging reports must be within the last 30 days)					
	If the request is from a treating physician outside an Ontario CAR T-cell treating facility, email or fax from the treating facility/physician confirming that they have capacity and are willing to accept this patient					
	If the request is for treatment out of country, email or fax from the Ontario CAR T-cell treating facilities confirming no capacity and email or fax from the out of country treating facility confirming their capacity and willingness to accept this patient					
	If the request is for a non-Ontario resident, a funding approval letter from the patient's provincial/territorial Ministry of Health is required, specifying CAR T-cell product(s) that is/are funded by the patient's provincial/territorial Ministry of Health					
List	: B: Available upon request					
	Cerebrospinal Fluid (CSF) studies documenting CNS disease status (within the last 30 days)					
	Documentation of CD19 tumour expression in BM or peripheral blood by flow cytometry (if done)					
Ш	Multidisciplinary cancer conference (MCC)/tumour board notes (if available)					
	<u> </u>					
*By	r checking this box, I certify that the information set out in this questionnaire is true and accurate, to the best of my knowledge:					
*Eı	nroling Physician: *Date: (DD-MMM-YYYY or click arrow down button to use calendar to enter the date)					
Nee	ed this information in an accessible format? 1-877-280-8538, TTY 1-800-855-0511, info@ontariohealth.ca					
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