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## PROCEDURES FOR PRIOR AUTHORIZATION

Completed forms can be faxed in confidence to 1-514-286-8480 for residents of Quebec and 1-844-661-2640 for residents of all other provinces

Upon receipt, this request will be confidentially reviewed according to payment criteria developed by Medavie Blue Cross in consultation with independent health care consultants. In some cases, additional clinical or diagnostic information may be required to process your claim.

For Quebec residents, the criteria for prior authorization are adjusted to meet the requirements of the Régie de l'assurance maladie du Québec (RAMQ).

- Prior Authorization is a pre-approval process to determine if certain products will be reimbursed under a member's benefit plan.
- Please complete entire form. Incomplete forms cannot be processed.
- For certain medications, approval for reimbursement may be conditional on confirmation of enrollment in the patient support program.
- Prior Authorization may be limited to a specified period or quantity of medication.
   Some Medavie Blue Cross plans may require you to purchase a drug requiring prior authorization from a preferred pharmacy\*. If your prior authorization request is approved, a case manager may contact you, your physician, or Patient Assistance Program to provide information about the program and to arrange to have your prescription transferred to the preferred pharmacy.
  - \*Not applicable in Quebec.
- In cases where a request for Prior Authorization is declined, Medavie Blue Cross is denying payment for a product and is not challenging the medical opinion of the physician nor rendering a medical opinion.
- Any costs associated with the completion of this form or obtainment of additional medical information are the responsibility of the member.
- Renewal of the Prior Authorization will be considered by Medavie Blue Cross upon request from the patient/member. The renewal request should include information from the physician supporting continued use of the medication.
- Prior Authorization coverage is contingent on your continued status as a Medavie Blue Cross cardholder or beneficiary.
- If this is a request under the *Mesure du patient d'exception* for a Quebec resident, please include a completed *Patient d'exception* form that can be found here: <a href="www.medaviebc.ca/en/resources">www.medaviebc.ca/en/resources</a>, in addition to this document.
- If you would like more information about our Patient First Network, including how your Patient Support Program can become integrated with our new enhanced Prior Authorization processes, please send an e-mail to: patientfirstnetwork@medavie.bluecross.ca.

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1 PHARMACY INFORMATION	
This section is to be completed by the Professional coordinating the re Pharmacy)	quest on behalf of the member (PSP, Cancer Care Navigator or
Decision communication preference:	Telephone, Number:
Name of Program/Pharmacy:	
Contact Name:	Contact E-mail:
2 PATIENT INFORMATION	
Part A	
Patient Name:	Date of Birth:
E-mail address of patient (or of legal guardian if patient is underage):	(mm/dd/yyyy)
Address:	Suite: City:
Province: Postal Code:	Telephone Number:
Policy Number: ID Number:	
Do you have valid Medicare coverage in your current province of residence?	☐ Yes ☐ No
Have you already purchased this prescription?	☐ Yes ☐ No
Please attach your paid-in-full receipt with this request form. If you have alre your receipt to Medavie Blue Cross, please indicate the date of the oldest receipt to Medavie Blue Cross, please indicate the date of the oldest receipt to Medavie Blue Cross, please indicate the date of the oldest receipt to Medavie Blue Cross, please indicate the date of the oldest receipt to Medavie Blue Cross, please indicate the date of the oldest receipt to Medavie Blue Cross, please indicate the date of the oldest receipt to Medavie Blue Cross, please indicate the date of the oldest receipt to Medavie Blue Cross, please indicate the date of the oldest receipt to Medavie Blue Cross, please indicate the date of the oldest receipt to Medavie Blue Cross, please indicate the date of the oldest receipt to Medavie Blue Cross, please indicate the date of the oldest receipt to Medavie Blue Cross, please indicate the date of the oldest receipt to Medavie Blue Cross, please indicate the date of the oldest receipt to Medavie Blue Cross, please indicate the date of the oldest receipt to Medavie Blue Cross, please indicate the date of the oldest receipt to the oldest receipt to the oldest receipt to the oldest receipt the date of the oldest receipt to the oldest	
Part B – Coordination of Benefits	
Do you or any dependant have coverage for this drug under any other plan	or program?
Policy Number:	Carrier:
(If applicable, please attach Explanation of Benefits from prior carrier with complete for	m)
If the patient is a dependent, provide the birth day and month of the cardhold	ler for the other carrier:(mm/dd)
Public-Funded Program – Have you applied for coverage through a public-fu	
If No, please indicate why:	
Part C – Authorization	
	d and used by Medavie Blue Cross to administer the terms of my policy or the ducts and services that I am eligible for as a member of a policy, and other nent at www.medaviebc.ca.
released to following third parties as required for the purposes of administe	ch as claim, health and/or financial related data may be collected from and/or tring and managing the benefits outlined in the policy of which I am an eligible surance companies, regulatory authorities and investigative bodies, services nt.
Where allowed by law, my information may be shared with Medavie Blue collected. If I am a resident of Quebec, this includes transferring or disciproviders outside of that province.	Cross employees or service providers in jurisdictions other than where it was osing my personal information to Medavie Blue Cross employees or service
I understand that my consent is only valid for the time it is needed to achieve my consent at any time. However, in some instances doing so may preven may be useful to me and/or my dependents. This consent complies with federal consent	the purposes outlined herein, unless I withdraw it. I understand I may withdraw t Medavie Blue Cross from providing me with certain products or services that eral and provincial privacy laws.
For more details about our information practices, including how your persor you have concerns or questions, please see our Medavie Blue Cross Privacy	nal information is protected, how to access or correct personal information, or if y Statement available at <a href="https://www.medaviebc.ca">www.medaviebc.ca</a> or call 1-800-667-4511.
Signature of Patient:	<b>Date:</b>

Residents of All Other Provinces
PO BOX 220, MONCTON (NB) E1C 8L3
TEL.: 1-800-667-4511 FAX: 1-844-661-2640

Residents of Quebec PO BOX 3300, STATION B, MONTREAL (QC) H3B 4Y5 TEL.: 1-888-588-1212 FAX: 1-514-286-8480



3 SPECIALTY DRUG INFORM	ATION			
Name of patient:				
Policy Number:				
E-mail address of patient or of leg	al guardian if patient	is underage:		
3A Patient Support Program (PS	SP) Enrollment			
•			, , ,	
Indicate the name of the Patient Su	· · · · · · · · · · · · · · · · · · ·			
PSP phone #:			PSP Fax #:	
Product Name	Strength	Dosage		Diagnosis
OPDIVO (NIVOLUMAB)				
Patient weight:	☐ lbs ☐ kg	Numbe	r of vials / syringes per do	ose:
Date treatment was initiated:	(mm/dd/yyyy)		Expected duration of tre	eatment:
Date of diagnosis:			Was treatment initiated	in hospital? ☐ Yes ☐ No
(mm	/dd/yyyy)			. – –
Where is medication being admining admining a special type of the physical type of the physic			he treatment:	
Indicate if the disease or injury is			ne treatment.	
For Initial Request, please comp	ete sections 3B and	d 3D. For Rene	wals, please complete s	sections 3C and 3D.
3B Initial Request				
				indications. The information for the on the contraindication if applicable.
<ul> <li>Renal-cell carcinoma: pazopani</li> <li>Non-small cell lung cancer: plat afatinib, erlotinib, gefitinib, certi</li> <li>Squamous cell carcinoma of the Hepatocellular carcinoma: sora</li> </ul>	inum doublet combin nib, alectinib, pembro head and neck: plati	ation chemother olizumab, doceta	apy (cisplatin or carboplati xel, ramucirumab, pemetre	in based), cytotoxic chemotherapy, osimertinib, exed, gemcitabine, vinorelbine
Category Prod	uct Name	Dosage	Duration of Treatment	Response to Treatment or Contraindication
Category	uct Name	Dosage	Duration of Treatment	response to Treatment of Contramucation
Is the drug being prescribed acc *NOTE: Do not provide genetic test res Approved indications from Health C	ults.	n Canada prod	uct monograph? 🗌 Ye	s 🗌 No
Clinical information				
		,		
ECOG score (date):		(mm/d	<u>d/yyyy)</u>	
Pharmacological treatment:				
☐ First-line treatment	П	Second-line tre	atment	☐ Third-line treatment or more
i not-mio dodunont	Ш	2000 IU-IIIIO IIG	aunont	

3	SPECIALTY	DRUG INFOR	MATION		
	•				
	•				
E-I	mail address o	of patient or of le	gal guardian if patient is underage	:	_
3B	Initial Requ	est (cont'd)			
	Clinical info	rmation (cont'd	)		
	Administration As mono	n of treatment:			
			metyx (cabozantinib)		
			by (ipilimumab) and 2 cycles of pla	tinum-based chemoth	erapy
	☐ In combir	nation with other	medications. Specify:		
	The cancer is	2.			
	☐ Metastati		☐ Nonmetastatic	☐ Resectable	e Unresectable
	_	dvanced	☐ Recurrent	<del></del>	cify):
1.	Renal-cell	carcinoma			
	Processo of	activo motactaco	es to the central nervous system:	□ Vos. □ No.	
	Tresence or	active metastase	es to the central hervous system.		
2.	Non-small	cell lung cance	er		
	Stage of can	cer: 🔲 I 🔠 I	I ☐ III ☐ IV ☐ Other (spe	ecify):	<u> </u>
	•	-	ific genomic tumour aberration? │ nich gene: ☐ EGFR gene* ☐ A		o not provide genetic test results.
				-	, •
			oimmune disease?		
	Does the pat	ient nave sympto	omatic interstitial lung disease?	」Yes □ No	
	Does the pat	ient have brain r	metastases?		
	$\hookrightarrow$	• •	answer the following questions:		
		•	t returned to baseline neurological related toxicity, at least 2 weeks p	•	☐ Yes ☐ No
		Has the patien	t been titrated down to a lower dai	ly steroid dose?	☐ Yes ☐ No
		→ If yes, sp	ecify current dose of prednisone p	er day:	mg
3.	Squamous	s cell carcinoma	a of the head and neck		
•				□ Voc. □ No.	
			es to the central nervous system:		
	•	·	ım-based therapy? ☐ Yes ☐ N		
	<b>→</b>	ıт yes, has the di	sease progressed within 6 months	or naving chemothera	apy? ∐ Yes ∐ No

3	SPECIALTY DRUG INFORMATION
	ame of patient: Date of Birth:
	olicy Number: mail address of patient or of legal guardian if patient is underage:
3B	Initial Request (cont'd)
4.	Hepatocellular carcinoma
	Is the cancer amenable to curative surgery or local treatment?   Yes   No
5.	Melanoma
	Adjuvant treatment of melanoma:
	Stage of cancer:
	Did the patient have a complete resection? ☐ Yes ☐ No
6.	Urothelial carcinoma
	Adjuvant treatment:    Yes    No
	Has the patient undergone radical resection of urothelial carcinoma? ☐ Yes ☐ No
	→ If yes, indicate the date of surgery:  (mm/dd/yyyy)
	Is the patient at high risk of recurrence?
3C	Renewal Request
Plea	ase provide information on the evolution of the disease to evaluate the response to treatment.
Da	ate of initial evaluation (pretreatment): Date of most recent evaluation: (mm/dd/yyyy)
	Information necessary to the assessment of this request
	Initial ECOG score (pretreatment): Most recent ECOG score (current):
	Response to treatment:
	☐ Partial ☐ Complete ☐ Stabilization of the disease ☐ Progression of the disease ☐ Confirmed by radiographic evidence: ☐ Yes ☐ No
	Response as per RECIST criteria:
	Does the patient continue to receive clinical benefit from the medication?   Yes   No
	Is the patient still tolerating the treatment with minimal / manageable side effects?

3 SPECIALTY DRUG INFORMATION	
Name of patient: Date of Birth:	
Policy Number: ID Number: E-mail address of patient or of legal guardian if patient is underage:	
3D Additional Information	
Please indicate any additional information pertaining to this request.	





First Name:	Last Name:	Permit Number:
Specialty:		
Clinic Name:		
Address:		Suite:
City:	Province:	Postal Code:
E-mail:	Telephone:	Fax:
Signature:		Date:
		(mm/dd/yyyy)

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No.