

State of Oklahoma SoonerCare

Yervoy[®] (Ipilimumab) Prior Authorization Form

Member Name:	Date of Birth:	Member ID#:		
Drug Information				
Physician billing (HCPCS code:) Star	rt Date (or date of next dose):		
Dose:		gimen:		
	Billing Provider I			
Provider NPI:	Provider	r Name:		
Provider Phone:	Provi	ider Fax:		
Prescriber Information				
Prescriber NPI:	_ Prescriber Nam	ne:		
Prescriber Phone:	Prescriber Fax:	Specialty:		
	Criteri			
Page 1 of 2—Please complete and return <u>all</u> pages. Failure to complete all pages will result in processing delays.* Please note: If Yervoy® (ipilimumab) is to be used in combination with Opdivo® (nivolumab), please completely fill out and submit the Opdivo® (nivolumab) prior authorization form (PHARM-64) that is available at: https://oklahoma.gov/ohca/rxforms.html				
For Initial Authorization:				
 Please indicate the diagnosis and informati Unresectable or Metastatic Melanom 				
A. Will ipilimumab be used in combina B. Will ipilimumab be used in combina progression if nivolumab was not p i. If answer to previous question a. Has the member previous a. Has the member previous a. Has the member previous question a. Will ipilimumab be used as a single be ii. If answer to previous question a. Did member experience a. Did member experience a. Did disease progress and course of ipilimumab, and the provide member's weight (kg. Please indicate member's ECOG providemember's ECOG providemember	ation with nivolumabation	as second-line or subsequent therapy for disease S No ovide the following: -L1 inhibitors? Yes No therapy? Yes No ne or subsequent lines of therapy? Yes No ent? Yes No ovide the following: c toxicity during prior ipilimumab therapy? greater than six months following completion of a prior revening therapy has been administered? O-5): n lymphadenectomy? Yes No les of >1 mm and no in-transit metastasis? Yes No		
C. Will ipilimumab be used as a single-agent? Yes No D. Please provide member's weight (kg):				
 Mesothelioma A. Is diagnosis malignant pleural mesothelioma that cannot be surgically removed? Yes No B. Will ipilimumab be used as first-line therapy? Yes No C. Will ipilimumab be used in combination with nivolumab? Yes No ii. Will ipilimumab be used in combination with nivolumab and 2 cycles of platinum-doublet chemotherapy? Yes No iii. Does tumor express PD-L1 ≥1%? Yes No 				
 Esophageal Squamous Cell Carcinom A. Is diagnosis unresectable advance B. Will ipilimumab be used as first-line C. Will ipilimumab be used in combination 	d or metastatic ESC therapy? Yes	No		
PLEASE PROVIDE THE INFORMATION REQUESTED AND RETURN TO: CONFIDENTIALITY NOTICE				
University of Oklahoma College of Physics	_	This document, including any attachments, contains information which is		

University of Oklahoma College of Pharmacy Pharmacy Management Consultants Product Based Prior Authorization Unit Fax: 1-800-224-4014

Phone: 1-800-522-0114 Option 4

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Member Name:	Date of Birth:	Member ID#:
	Criteria	
	urn <u>all</u> pages. Failure to complete ส	all pages will result in processing delays.*
For Initial Authorization (continued)		
 Please indicate the diagnosis and info 	ormation (continued):	
Small Cell Lung Cancer		
 A. Did disease relapse within 6 m 	onths of initial chemotherapy? Yes	No
B. Did disease progress on initial	chemotherapy? Yes No bination with nivolumab? Yes N	
C. Will ipilimumab be used in com	ibination with nivolumab? YesN	0
	OG performance status (0-5)	
■ Non-Small Cell Lung Cancer (NS		_
A. Is diagnosis recurrent, advance	ed, or metastatic disease? YesN	No
B. Will ipilimumab be used as firs	t-line therapy? Yes No	
	eptor (EGFR) or anaplastic lymphom	a kinase (ALK) genomic tumor aberrations?
YesNo		de la falla Carrara de la la falla de la constitución de la Carrara de la constitución de
	combination with nivolumab and 2 cyc	cles of platinum-doublet chemotherapy?
YesNo	> 40/ 2 V/aa Na	
iii. Does tumor express PD-L1	21%? Yes NO	
☐ Hepatocellular Carcinoma	i i di	A to a source of O. Maria
	ble disease and is not a candidate for	
	disease or extensive liver tumor burg	
	cond-line or greater therapy? Yes	
D. Will ipilimumab be used in com	led other checkpoint inhibitors? Yes	lo No
Renal Cell Cancer	led other checkpoint inhibitors: res_	NO
	ally upresentable stage IV disease in	the initial treatment of a member with
	renal cell cancer? YesNo	the initial treatment of a member with
	estion is 'yes', please provide the follo	owing:
i. Il aliswel to previous qu ☐ Intermediate		owing.
☐ Poor risk	lisk	
☐ Other:		
B Will initimumah he used in com	nbination with nivolumab? Yes N	0
C. Has the member previously fai	led PD-L1 or PD-1 inhibitors? Yes	No.
D. Please provide member's weig	ht (ka):	140
☐ Colorectal Cancer	··· (Ng)	
	etastatic microsatellite instability-high	n (MSI-H) or mismatch repair deficient (dMMR)
colorectal cancer? Yes No		r (INOI-11) of mismater repair denoient (divinity)
	bination with nivolumab? Yes N	lo.
☐ If diagnosis is not listed above, ple		
Additional Information:	sase maleate diagnosis.	
Additional information.		
For Continued Authorization:		
Date of last dose:		
2. Does member have any evidence of	progressive disease while on ipilimum	nab? Yes No
3. Has the member experienced advers	e drug reactions related to ipilimumal	b therapy? YesNo
	eactions:	
Drocoribor Signature:	D.	240.
Prescriber Signature:	Di	ate:
knowledge. Failure to complete this form	i medically necessary and all infort n in full will result in processing delay	mation is true and correct to the best of my
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