

Nevada Medicaid – Molina Healthcare

Cystic Fibrosis Agents Prior Authorization Request Form

Please provide the information below, please print your answer, attach supporting documentation, sign, date, and return to our office as soon as possible to expedite this request. Please FAX responses to: (844) 259-1689. Phone: (833) 685-2103.

	Member Info	rmation _{(r}	equired)	Provider Information (required)				
Me	ember Name:	Provider Name:						
Molina ID#:				NPI#:		Specialty:		
Date of Birth:				Office Phone:				
Street Address:				Office Fax:				
Cit	y:	State:	Zip:	Office Street Addres	SS:			
Ph	one:			City:		State:	Zip:	
			Medication In	formation (require	ed)			
Medication Name:				Strength:	, a ,	Dosage Form:		
	Check if requesting brand	Directions for Use:						
	□ Check if request is for continuation of therapy							
	Clinical Information (required)							
	elect the diagnosis belo	w:						
	Cystic Fibrosis Other diagnosis:			ICD-10 Code(s):				
	Other diagnosis.						_	
	Drug-Specific Information (required)							
	Kalydeco® (ivacaftor)							
	The recipient is six months of age or older							
	☐ There is documentation that the recipient has had an FDA-approved cystic fibrosis mutation test confirming the presence of one of the gene mutations listed in the FDA-approved package insert (please attach documentation to request)							
							- /	
	☐ If the request is for continuation of therapy, the recipient has documentation of positive clinical response to Kalydeco®							
	therapy							
Oı	rkambi® (lumacaftor/i	ivacaftor)						
	The recipient is two year		older					
	, ,	gous for the	e F508del mutation	n in the cystic fibrosi	s transmembran	e conducta	nceregulator	
	(CTFR) gene	over 10 he	al iro					
		•		of severe henatic i	mnairment			
		-	dis in the presence	or severe nepation	трантенс			
Sy	/mdeko® (tezacaftor/i							
	The recipient is six year	•						
 □ The medication is prescribed by or in consultation with a pulmonolo □ The recipient is homozygous for the F508del mutation as detected 					•			
	approved facility	, 9000 101 111	c . ooodoi matatioi	. as actooled by all	. Di Colouica Oi	matation to	OST OF OED T	
	-	of the FDA a	ipproved package i	nsert listed mutation	ns on at least on	e allele in tl	he CF	

transmembrane conductance regulator (CFTR) gene as detected by FDA cleared CF mutation test or CLIA approved

☐ If the request is for continuation of therapy, the recipient has documentation of positive clinical response to Symdeko®

therapy (e.g., improvement in lung function or decreased number of pulmonary exacerbations)

facility

Trikafta®	(elexacaftor/tezacaftor/ivacaftor and ivacaftor)			
☐ The rec	ipient is 12 years of age or older ipient has at least one F508del mutation in the CFTR gene as detected by an FDA cleared CF mutation test or a formed at a CLIA approved facility			
☐ If the re therapy	The medication is prescribed by or in consultation with a pulmonologist or a specialist associated with a CF care center If the request is for continuation of therapy, the recipient has documentation of positive clinical response to Trikafta® therapy (e.g., improvement in lung function [percent predicted forced expiratory volume in one second {PPFEV1}] or decreased number of pulmonary exacerbations)			
Are there any o	other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to			

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