

	orm to: 877.974.4411 toll fr		
This form applies to:	☑ Commercial (Tradition☐ Medicald	onal) 🛛 Commercial (In	dividual/Optimized)
This request is:) Non-Urgent (standard re	view)
	Urgent means the standard review to regain maximum function.	time may seriously jeopardize the life or h	nealth of the patient or the patient's ability
Stivarga [®]	· ·		
	(regeratorne)		
Member			
Last Name: ID #:		First Name:	Candari
	:		Gender:
Requesting Provider:		Prov. Phone:	Prov. Fax:
Provider Address:			
Provider NPI:		Contact Name:	
Provider Signature:		Date:	
Product Information	on		
Drug product:	☐ Stivarga 40 mg tablet	Start date (or date of next	dose):
		Date of last dose (if applicable):	
		Dosing frequency:	
Drug cost informa	tion		
The wholesale acquisthan \$172,200.	sition cost for Stivarga is \$157.70	for each tablet. The annual cost o	of treatment with this drug is more
Precertification Re	equirements		
For this drug to be co	vered, the patient must have one of	f the following listed conditions:	
Must have one	of the following conditions and satisf	y any specific criteria pertaining to th	e condition:
 a. Colorecta 	cancer (CRC)		
	have previously been treated with flu- therapy, and, if RAS wild-type, an a		ecan-based chemotherapy, an anti-
	vanced or metastatic gastrointestinal		
	have had previously been treated wit llular carcinoma (HCC)	n imaunid and suniunid.	
	have had previously been treated wit	h sorafenib.	
Priority Health Pre	ecertification Documentation		
	is this drug being requested for	?	
☐ Colorectal	cancer vanced or metastatic GIST		
☐ Hepatocell	ular carcinoma		
	ionale for use:		



Please submit all relevant documentation to support medical necessity including previous therapies tried (with dates of use and associated outcomes), genetic or other laboratory testing, and any other pertinent information.

Additional information

Stivarga is covered for a 21-day supply (based on a 28-day treatment cycle).

Note: Authorization for indications, dosing, or a route of administration not approved by the Food and Drug Administration (FDA) or recognized in CMS-accepted compendia (e.g. DrugDex, AHFS, U.S. Pharmacopeia, and also Clinical Pharmacology for oncology indications only) require supporting evidence for coverage. Please provide two published peer-reviewed literature articles supporting the appropriateness of the drug, the dosing of the drug, or the route of administration to be used for the identified indication.