## NC Medicaid Pharmacy Prior Approval Request for



Pharmacy PA Call Center: (866) 246-8505

## Viekira Pak

Beneficiary Information			
1. Beneficiary Last Name:	2. First Name:		
3. Beneficiary ID #:	4. Beneficiary Date of Birth:	5. Beneficiary Gender:	
Prescriber Information			
7. Requester Contact Information - N	ame:Pho	ne #: Ext	
Drug Information			
8. Drug Name:	9. Strength:	10. Quantity Per 30 Days: <u>112</u>	
11. Length of Therapy (in days):	] 365 Days		
Clinical Information			
Total Length of Therapy (Ch	eck ONE):		
☐ 12 weeks = Genotype 1a,	, without cirrhosis, or genotype 1b, wit	th cirrhosis	
☐ 24 weeks = Genotype 1a,	, with compensated cirrhosis		
1. What is the beneficiary's	Genotype?		
2. Is the beneficiary is 18 ye	ars of age or older with a diagnosis of	chronic hepatitis C (CHC) infection with	
confirmed genotype 1b v	vithout cirrhosis or with compensated	cirrhosis or confirmed genotype 1a	
without cirrhosis or with	compensated cirrhosis in combination	າ with ribavirin? 🗆 Yes 🗆 No	
3. For all treatment courses	except genotype 1b, will treatment in	nclude the use of ribavirin? $\square$ Yes $\square$ No	
4. As the provider, are you	easonably certain that treatment will	improve the beneficiary's overall health	
status? ☐ Yes ☐ No			
5. Has the provider assessed	d for laboratory and clinical evidence of	of hepatic decompensation?   Yes   No	
6. Does the beneficiary have cirrhosis? $\square$ Yes $\square$ No If answer is yes, please answer the following:			
· ·		nptoms of hepatic decompensation (such	
	cephalopathy, variceal hemorrhage)?		
•	, , ,	uding direct bilirubin levels at baseline and	
_	weeks of starting treatment and as clir		
	•	hibitors used to treat CHC (i.e. boceprevir,	
		eotide NS5B polymerase inhibitor such as	
Sovaldi® (sofosbuvir)? □		ANGEA SALIES OF THE PARTY OF THE	
, -	iekira Pak in combination with anothe		
		either failed to achieve a SVR (defined as a SVR during a prior successfully completed	
	sting of Sofosbuvir? $\square$ Yes $\square$ No	3VN during a prior successfully completed	
•	S	d either failed to achieve a SVR (defined as	
	of 25 IU/mL) or relapsed after achieving		
	egimen consisting of Ledipasvir? $\square$ Yes		
Completed treatment re	semen consisting of Leulpasvii!	) LI INU	

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•	isease as defined by Child-Pugh classification score of ed in beneficiaries with moderate to severe hepatic
<ul><li>12. Has the beneficiary attempted a previous course</li><li>13. Does the beneficiary have any FDA labeled contra</li></ul>	• •
Signature of Prescriber:	Date:
(Prescriber Signa	ature Mandatory)
any falsification, omission, or concealment of material fac	implete to the best of my knowledge, and I understand that it may subject me to civil or criminal liability.