

Fax completed form to: (855) 840-1678
If this is an URGENT request, please call (800) 882-4462 (800.88.CIGNA)

Hepatitis Prior Authorization

PHYSICIAN I	NFORMATION		PATIEN	T INFOR	MATION	
* Physician's Name:			*Due to privacy regulations we will not be able to respond via fax with the outcome of our review unless all asterisked (*) items on this			
Specialty: * DEA, NPI or TIN:			form are completed.*			
Office Contact Person:			* Patient Name:			
Office Phone:			* Cigna ID: * Date of Birth:			Birth:
Office Fax:			* Patient Street Address:			
Office Street Address:			City State Zip			Zip
City	State	Zip	Patient Phone:			1
Urgency: ☐ Standard		Urgent (In chec	king this box, I attest to the fact tha jeopardize the customer's life, healt	t applying the h, or ability to	standard r regain ma	review time frame may eximum function)
Medication requested: □ Daklinza 30mg tablets □ Daklinza 60mg tablets □ Daklinza 90mg tablets □ Daklinza 90mg tablets □ Epclusa 200mg-50mg tablet □ Epclusa 400mg-100mg tablet □ Harvoni 45mg-200mg tablet □ Harvoni oral pellets 33.75m □ Harvoni oral pellets 45mg-2 □ ledipasir/sofosbuvir 90mg-4 □ Mavyret 100mg-40mg tablet □ Sovaldi 200mg tablets □ Sovaldi 400mg tablets □ Sovaldi oral pellets 150mg □ Sovaldi oral pellets 200mg □ sofosbuvir/velpatasvir 400m □ Viekira Pak □ Vosevi □ Zepatier 50mg-100mg tablet □ Ribavirin 200mg □ Rebetrol 200mg capsules □ Copegus 200mg tablets	lets ets ng-150mg 200mg 400mg tablets (au ets		Pegasys Pegasys Pegasys Pegasys Peg-Intro Intron A		5mg prefill 5ml Proclic 1 vial 5ml vial 5.5ml vial 0.5ml vial 5ml Redip 5.5ml Red 0.5ml Red nits multid	ed syringe ck pen pen ipen ipen
TREATMENT LENGTH: 8 weeks 12 weeks 16 weeks 24 weeks Other (please specify):						
Is the requested medication for the patient?	a chronic or long	g-term conditior	for which the prescription med	ication may	be neces	sary for the life of ☐ Yes ☐ No
Diagnosis related to use: ☐ acute Hepatitis C ☐ recurrent hepatitis c virus (H☐ Other (please specify): **Please fill out questions for		ansplantation —	□acute Hepatitis B**		□chro	nic Hepatitis B**
Clinical Information: What is the patient's current we	eight?		☐ lbs ☐ kg			

(if not already indicated above) Will y (if no) Please explain:	our patient also b	e taking ribavirin?			Yes	□No
For Harvoni 45/200 or Solvaldi 200 (your patient (examples could include					dosing/qu	antity for
FOR HEPATITIS C:						
		☐ 1b ☐ 5	☐ 1 (unknown subtype) ☐ 6 ☐ Other:			
Does the patient have HIV/AIDS?					☐ Yes	☐ No
Does the patient have Hepatitis B?					☐ Yes	□No
Does the patient have recurrent Hepa If yes, is your patient treatm					☐ Yes ☐ Yes	_ □ No □ No
Does the patient have hepatocellular If yes, has your patient prev If yes, is your patient waiting If yes, does your patient	iously had a liver g to undergo a live	transplant? er transplant?			☐ Yes ☐ Yes ☐ Yes ☐ Yes	No No No No
Does your patient have severe renal	impairment (a cre	eatinine clearance	less than 30 mL/min)?		☐ Yes	□No
☐ liver biopsy ☐ ultrasound-base ☐ specific blood te ☐ other (please sp	ed transient elaste est (for example, becify): sis) Which best detected the following later the following later the state of the following later	ography FibroTest, FibroSi escribes the patie	gree of cirrhosis or hepatic fib ure, etc.) nt's grade/Child-Pugh score c	, ,		ing:
total bilirubin		Date ta	ken:	_		
INR		_ Date taken:				
(if patient does NOT have FibroSure, IASL, Is			ne score and scoring system ເ 	used (for examp	le: Batts-	Ludwig,
OR - the following: AST	☐ Yes ☐No	_ Date tak _ Date tak _ Date tak _ weight _	en: en: en: en:			

Is your patient committed to pa	rticipate	in a hepatitis C disease s	state management progra	m?	☐ Yes ☐ No	
The following HCV RNA le	vels ar					
Week of Therapy		Level in	iu/ml	Date T	aken	
Pretreatment Baseline**						
4						
8						
12						
24						
other						
**Pretreatment H	CV RNA	levels should be basel	ine (e.g. within 3 month	s of planned therap	y initiation).	
patient has used in previ	Please indicate all drugs the patient has used in previous treatments (tx) or if use is Stopped tx early (Response: none (N), partial (P), reatments (tx) or if use is Relapsed / Recurrence? to any of these?** Y/N Y/N					
☐ Alferon N (interferon alfa-n	3)					
☐ Daklinza (daclatasvir)	10)					
☐ Epclusa (sofosbuvir/velpat	asvir)					
☐ Harvoni (ledipasvir-sofosb	uvir)					
Harvoni oral pellets						
Incivek (telaprevir)	<u> </u>					
☐ Intron A (interferon alfa 2b☐ ledipasvir/sofosbuvir (auth						
generic for Harvoni)	onzeu					
Mavyret (glecaprevir/pibre						
☐ Pegasys (peginterferon alf						
PegIntron (peginterferon a	lfa 2b)					
Ribavirin						
sofosbuvir/valpatasvir (aut generic for Epclusa)	norized					
Sovaldi (sofosbuvir)						
Sovaldi oral pellets						
☐ Victrelis (boceprevir)						
☐ Viekira-Pak	.:\					
(ombitasvir/paritaprevir/ritonav	vir)					
(sofosbuvir/velpatas/voxilapre	x)					
Zepatier (elbasvir/grazopre						
Other:						
*If stopped early, please explain why: **If contraindicated, please explain why:						
For Harvoni or ledipasvir/sofosbuvir (authorized generic for Harvoni) requests: (if requesting Harvoni or its authorized generic and the patient has the following: genotype 1, non-cirrhotic, AND treatment-naïve) Which of the following applies to this patient? HIV positive HCV RNA level 6 million IU/mL or higher none of the above						

For Vosevi resquests: (if requesting Vosevi and the patient has genotype 3) Has you yes and detected yes and NOT detected no, this testing was not done	ur patient been screened for Y93H substitution?	
For Zepatier requests: (if requesting Zepatier and the patient has genotype 1a) Has y mutations at codons 28, 30, 31, or 93? not requesting Zepatier or not genotype 1a yes, and NS5A polymorphism was detected at codons 28, yes, and NS5A polymorphism was NOT detected at codonn no, this testing was not done	30, 31, or 93	d found to have
FOR HEPATITIS B:		
What is the patient's genotype?		
(if age 3-17 AND new start for Pegasys) Does your patient have	ve Hepatitis B e-antigen (HBeAg)-positive disease?	☐ Yes ☐ No
(if age 3-17 AND new start for Pegasys) Does your patient havaminotransferase (ALT)?	ve evidence of viral replication and elevations in serum	
Does the patient have cirrhosis? (if yes) Which best describes the patient's grade/Chil Grade A/ score 5-6 Grade B/ score 7-9 Grade C/ score 10-15 unknown – Please provide the following labs: albumin Da	d-Pugh score of hepatic impairment?	☐ Yes ☐ No
total bilirubin	Date taken:	
INR Date to	aken:	
Is the patient currently on the requested therapy? If yes, how many weeks has the patient completed:_ Date started therapy://	weeks	☐ Yes ☐ No
For Other Diagnoses:		
adult T-cell leukemia/lymphoma (ATLL) AIDS-related Kaposi sarcoma chronic myelocytic leukemia (CML) condylomata acuminata desmoid tumor (aggressive fibromatosis) essential thrombocythemia (ET) follicular lymphoma giant cell tumor of the bone (GCTB) hairy cell leukemia neuroendocrine tumors of the gastrointestinal tract, lung ar primary cutaneous CD30+ T-cell lymphoproliferative disord		
(if ATLL) Which of the following applies to your patient? ☐ Intron A is being given as first-line therapy. ☐ Intron A is being given second-line therapy. ☐ other/unknown (if first-line) Is Intron-A being used in combination with (if second-line) Is Intron-A being used in combination		☐ Yes ☐ No ☐ Yes ☐ No
(if CML) Which of the following applies to your patient? ☐ Your patient was unable to tolerate one of the following: Bo (nilotinib). ☐ Patient is post-transplant and relapsed. ☐ other/unknown	osulif (bosutinib), Gleevec (imatinib), Sprycel (dasatinib	o), Tasigna

(if condylomata) Does your patient have failure, contraindication or intolerance to podofilox (Condylox)? (if desmoid tumor) Does your patient have primary, recurrent or progressive disease? (if desmoid tumor or hairy cell leukemia) Will Intron-A be used as single-agent treatment?	☐ Yes ☐ No ☐ Yes ☐ No ☐ Yes ☐ No
(if GCTB) Will Pegasys be used as single-agent therapy? (if melanoma) Is Intron-A being used as an adjuvant to surgical treatment? (if no) Is Intron-A being used for intralesional treatment? (if intralesional) Does your patient have unresectable disease? (if intralesional) Does your patient have recurrent or metastatic disease? (if NOT intralesional) Does your patient have metastatic or unresectable disease? (if not intralesional) Is your patient using Intron-A in combination with platinum-based chemotherapy (such or cisplatin)?	Yes No Yes No Yes No Yes No Yes No Yes No as carboplatin Yes Yes No
(if yes) Has your patient tried only one other treatment option for this diagnosis before using Intron-with platinum-based chemotherapy? (if no) Did your patient have disease progression while on BRAF therapy (such as dabrafenib [Tafini [Zelboraf])? (if disease progression) Does your patient have performance status (PS) 0-2? (if meningioma) Does your patient have surgically unresectable recurrent or progressive disease? (if meningioma) Was your patient previously treated with radiation for this diagnosis? (if myelofibrosis) Is your patient symptomatic? (if myelofibrosis) Does your patient have low-risk myelofibrosis? (if RCC) Does your patient have relapsed or stage IV (4) disease? (if RCC) What is the histology of the disease? non-clear cell	☐ Yes ☐ No
Additional Pertinent Information: Please provide clinical support for the use of this drug in your patient (includin prior therapy, performance status, and names/doses/admin schedule of any agents to be used concurrently).	g disease stage,
Attestation: I attest the information provided is true and accurate to the best of my knowledge. I unders Health Plan or insurer its designees may perform a routine audit and request the medical information necesthe accuracy of the information reported on this form. Prescriber Signature: Date:	
Save Time! Submit Online at: www.covermymeds.com/main/prior-authorization-forms/cigna/ or via SureScrip	ots in your EHR.
Our standard response time for prescription drug coverage requests is 5 business days. If your request is urgent, it	is important that

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