



## **FAX FORM AND CLINICAL DOCUMENTATION**

## **HEPATITIS C AGENTS PRIOR AUTHORIZATION FORM**

Prior authorization guidelines for **Hepatitis C Agents** and **Quantity Limits/Daily Dose Limits** are available on the DHS Pharmacy Services website

at	https://www.dns.pa.gov/prov	viders/Pharmacy-Serv	rices/Pages/default.a	<u>SDX</u> .	
Office contact name/phone:		Prescriber na	Prescriber name:		
LTC facility		State license	#:	NPI:	
contact/phone:					
Total # pages:		Street addres	Street address:		
Beneficiary name:		City/state/zip:	City/state/zip:		
Beneficiary ID#:	DOB:	Phone:		Fax:	
Requested drug #1:	Directions:		Qty:	☐ 8 weeks ☐ 16 weeks ☐ 12 weeks ☐ Other:	
Requested drug #2:	Directions:		Qty:	8 weeks 16 weeks 12 weeks Other:	
Is the beneficiary currently being treated with the requested drug?			☐ No ☐ Yes – The	☐ No ☐ Yes – Therapy start date:	
SUBMIT DOCUMENTATION from the medical record for all items below.					
For requests for NON-PREFERRED Hepatitis C Agents:					
1. Documentation that the beneficiary tried and failed or has a contraindication or intolerance to the preferred Hepatitis C Agents. (See the Preferred Drug List for the list of preferred Hepatitis C Agents at: <a href="https://papdl.com/preferred-drug-list">https://papdl.com/preferred-drug-list</a> .)					
Cirrhosis assessment documented by a recent noninvasive test and date of testing.					
<ul> <li>3. Genotype if one of the following (check the appropriate box for the beneficiary):  The beneficiary is prescribed a non-pangenotypic regimen.  The beneficiary is hepatitis C sofosbuvir-based, sofosbuvir/velpatasvir/voxilaprevir, or sofosbuvir plus glecaprevir/pibrentasvir treatment experienced.  The beneficiary has decompensated cirrhosis and is prescribed ledipasvir/sofosbuvir.  The beneficiary is treatment-naïve (with cirrhosis) and prescribed sofosbuvir/velpatasvir.</li> </ul>					
4. RAS (resistance-associated substitutions) testing and date of testing if one of the following (check the appropriate box for the beneficiary):  The beneficiary is genotype 1a and prescribed elbasvir/grazoprevir.  The beneficiary is genotype 1a, treatment-experienced, and prescribed ledipasvir/sofosbuvir.  The beneficiary is genotype 3, treatment-naïve (with cirrhosis) or treatment-experienced (without cirrhosis) and prescribed 12 weeks of sofosbuvir/velpatasvir.					
For requests for THERAPEUTIC DUPLICATION of Hepatitis C Agents direct-acting antivirals (DAAs):					
For a beneficiary taking more than 1 Hepatitis C Agents DAA product concomitantly:  The beneficiary has a medical reason for concomitant use of the requested products that is supported by peer-reviewed medical literature or national treatment guidelines.					
ATTESTATION from the prescriber for one of the items below.					
Check the appropriate box for the beneficiary.					
☐ The beneficiary is hepatitis C treatment naïve. ☐ The beneficiary has been treated for hepatitis C with the following treatment regimen:					
PLEASE FAX COMPLETED FORM WITH REQUIRED CLINICAL DOCUMENTATION TO 866-240-3712					
Prescriber Signature:	CENTIONE DOCUM	HENTATION 10	Date:		

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