## **AvMed**

## PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST\*

<u>Directions</u>: <u>The prescribing physician must sign and clearly print name (preprinted stamps not valid)</u> on this request. All other information may be filled in by office staff; <u>fax to 1-305-671-0200</u>. No additional phone calls will be necessary if all information (<u>including phone and fax #s</u>) on this form is correct. <u>If the information provided is not complete, correct, or legible, the authorization process can be delayed.</u>

MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete.

<u>Drug Requested</u>: <u>Doptelet</u><sup>®</sup> (avatrombopag)

Member Name:

Quantity Limit: 15 tablets

Dosage: 3 tablets (60mg) by mouth daily for 5 days

Member AvMed #:	Date of Birth:	
Prescriber Name:		
Prescriber Signature:		
Office Contact Name:		
Phone Number:		
DEA OR NPI #:		
<b>DRUG INFORMATION:</b> Authorization may be o		
Drug Form/Strength:		
Dosing Schedule:	Length of Therapy:	
Diagnosis:	ICD Code, if applicable:	
Weight:	Date:	
Recommended Dosage and Quantity Limits:		
Chronic liver disease and scheduled to undergo a procedure	Chronic immune thrombocytopenia	
Platelet count 40,000 to <50,000/mm <sup>3</sup>	20 mg Once Daily (Initial Dose Regimen);	
Quantity Limit: 10 tablets	MAXIMUM, 2 tablets (40 mg) once daily	
Dosage: 2 tablets (40mg) by mouth daily for 5 days  Platelet count <40,000/ mm <sup>3</sup>		

**CLINICAL CRITERIA:** Check below all that apply. All criteria must be met for approval. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied.

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	☐ Diagnosis: Chronic Liver Disease-Associated Thrombocytopenia					
<u>O</u>	ONE (1) TIME Service/Procedure-Date Approval					
	Member has a diagnosis of chronic liver disease					
	AND					
	Member is $\geq 18$ years of age					
	<u>AND</u>					
	The requesting provider is a gastroenterologist or hematologist, or has been in consultation with one					
	AND  The member is scheduled for an invasive procedure associated with moderate to high risk for bleeding [Moderate Risk: Liver biospsy, bronchoscopy, Ethanol ablation therapy or chemoembolization for hepatocellular carcinoma]  [High Risk: Vascular catheterization (including right-side procedures in patients with pulmonary hypertension), Transjugular intrahepatic portosystemic shunt, Dental procedures, Renal biopsy, Biliary interventions, Nephrostomy tube placement, Radiofrequency ablation, Laparoscopic interventions]					
	Name of procedure: Procedure date:					
	NOTE: Begin Doptelet 10-13 days prior to procedure (undergo procedure 5-8 days after the last dose)					
	AND					
	The member has a baseline platelet count of $\leq 55 \text{ x} 10^9 / \text{L}$					
	Document platelet count prior to therapy initiation: x10 <sup>9</sup> /L					
	<u>AND</u>					
	Select the corresponding dosing regimen for the member:  Platelet count 40 x10 <sup>9</sup> /L to <50 x10 <sup>9</sup> /L  Quantity Limit: 10 tablets  Dosage: 2 tablets (40mg) by mouth daily for 5 days  Platelet count <40 x10 <sup>9</sup> /L  Quantity Limit: 15 tablets  Dosage: 3 tablets (60mg) by mouth daily for 5 days					
	Diagnosis: Chronic Immune Thrombocytopenia (ITP)					
Int	tial Authorization Approval: 6 months					
	Member has a diagnosis of chronic ITP for at least 6 months (OR meets the corticosteroid requirement below)					
	<u>AND</u>					
□ 1						
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		AND	
_	Documentation of	f platelet levels within the last 30 days has	been submitted confirming < 30 x 10 <sup>9</sup> /L
_		AND	00011 0001111111111
_	The requesting pro	ovider is a hematologist, or has been in con	nsultation with one
		AND	
		e per day (failure defined as not having a re	corticosteroid at a recommended dose of 0.5-2.0 esponse to at least a 3-month trial or is
	DRUG/DOSE:	Date	es of therapy:
		AND	
		ve failed one (1) of the following therapies ill require different prior authorization to	` I U
		ablished dosing level recommendations based	unless otherwise indicated, AND the provider sed on platelet count [see dose table(s) in
R	<b>Reauthorization</b>	<b>Approval of Chronic Immune Th</b>	rombocytopenia Diagnosis ONLY:
cł		below all that apply. All criteria must be rentation, including lab results, diagnostics,	met for approval. To support each line and/or chart notes, must be provided or request
		t count has not reached target level to reco	mmend discontinuation of therapy
		AND	
	Document platele	t count 2 weeks after therapy initiation:	x10 <sup>9</sup> /L
		AND	
	Document current	t platelet count [lab work measured within x109/L	the date of this reauthorization request]:
		AND	
_	Based on current 1	platelet count, enter dose level (see tables	below) at which therapy will continue:
ΓΑ	ABLE 1: Dose Adju	ustment and Corresponding Platelet Count	

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Platelet Count (x10 <sup>9</sup> /L)	Dose Adjustment or Action	
Less than 50 after at least 2 weeks	Increase One Dose Level (according to dose table below)	
of DOPTELET	[Wait 2 weeks to assess the effects of this regimen and any subsequent	
	dose adjustments]	
Between 200 and 400	Decrease One Dose (according to dose table below)	
	[Wait 2 weeks to assess the effects of this regimen and any subsequent	
	dose adjustments]	
Greater than 400	Stop DOPTELET.	
	[Increase platelet monitoring to twice weekly. When platelet count is	
	less than 150 x10 <sup>9</sup> /L, decrease One Dose Level (according to dose table	
	below) and reinitiate therapy.]	
Less than 50 after 4 weeks of	Discontinue DOPTELET.	
DOPTELET 40 mg once daily		
Greater than 400 after 2 weeks of	Discontinue DOPTELET.	
DOPTELET 20 mg weekly		

TABLE 2: Dosage Adjustment Recommendations

Dose	Dose Level
40 mg Once Daily	6
40 mg Three Times a Week AND 20 mg on the Four Remaining Days of Each Week	5
20 mg Once Daily (Initial Dose Regimen)	4
20 mg Three Times a Week	3
20 mg Twice a Week OR 40 mg Once Weekly	2
20 mg Once Weekly	1

## Medication being provided by Specialty Pharmacy - PropriumRx

## Not all drugs may be covered under every Plan

If a drug is non-formulary on a Plan, documentation of medical necessity will be required.

\*\*Use of samples to initiate therapy does not meet step edit/preauthorization criteria. \*\*

\*Previous therapies will be verified through pha rmacy paid claims or submitted chart notes.\*