

# **Clinical Policy: Epoprostenol (Flolan, Veletri)**

Reference Number: CP.PHAR.192 Effective Date: 03.16 Last Review Date: 02.21 Line of Business: Commercial, HIM, Medicaid

Coding Implications Revision Log

See <u>Important Reminder</u> at the end of this policy for important regulatory and legal information.

# Description

Epoprostenol (Flolan<sup>®</sup>, Veletri<sup>®</sup>) is a prostacyclin.

#### FDA Approved Indication(s)

Flolan and Veletri are indicated for the treatment of pulmonary arterial hypertension (PAH) (World Health Organization [WHO] Group 1) to improve exercise capacity.

Studies establishing effectiveness included predominantly patients with New York Heart Association (NYHA) Functional Class III-IV symptoms and etiologies of idiopathic or heritable PAH or PAH associated with connective tissue diseases.

#### **Policy/Criteria**

*Provider must submit documentation (such as office chart notes, lab results or other clinical information) supporting that member has met all approval criteria.* 

It is the policy of health plans affiliated with Centene Corporation<sup>®</sup> that Flolan and Veletri are **necessary** when the following criteria are met:

#### I. Initial Approval Criteria

#### A. Pulmonary Arterial Hypertension (must meet all):

- 1. Diagnosis of PAH;
- 2. Prescribed by or in consultation with a cardiologist or pulmonologist;
- 3. Failure of a calcium channel blocker (*see Appendix B*), unless member meets one of the following (a or b):
  - a. Inadequate response or contraindication to acute vasodilator testing;
  - b. Contraindication or clinically significant adverse effects to calcium channel blockers are experienced;
- 4. If request is for brand Flolan or brand Veletri, medical justification supports inability to use generic epoprostenol sodium (e.g., contraindication to excipients);
- 5. Provider must submit treatment plan detailing pump rate, dose and quantity (in mL). Approval duration:

**Medicaid/HIM** – 6 months **Commercial** – Length of Benefit

#### **B.** Other diagnoses/indications

1. Refer to the off-label use policy for the relevant line of business if diagnosis is NOT specifically listed under section III (Diagnoses/Indications for which coverage is

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NOT authorized): CP.CPA.09 for commercial, HIM.PA.154 for health insurance marketplace and CP.PMN.53 for Medicaid.

### **II.** Continued Therapy

- A. Pulmonary Arterial Hypertension (must meet all):
  - 1. Currently receiving medication via Centene benefit or member has previously met all initial approval criteria;
  - 2. Member is responding positively to therapy;
  - 3. Provider must submit treatment plan detailing pump rate, dose and quantity (in mL). Approval duration:

**Medicaid/HIM** – 12 months **Commercial** – Length of Benefit

### **B.** Other diagnoses/indications (must meet 1 or 2):

1. Currently receiving medication via Centene benefit and documentation supports positive response to therapy.

Approval duration: Duration of request or 6 months (whichever is less); or

2. Refer to the off-label use policy for the relevant line of business if diagnosis is NOT specifically listed under section III (Diagnoses/Indications for which coverage is NOT authorized): CP.CPA.09 for commercial, HIM.PA.154 for health insurance marketplace and CP.PMN.53 for Medicaid.

#### III. Diagnoses/Indications for which coverage is NOT authorized:

A. Non-FDA approved indications, which are not addressed in this policy, unless there is sufficient documentation of efficacy and safety according to the off label use policies – CP.CPA.09 for commercial, HIM.PA.154 for health insurance marketplace and CP.PMN.53 for Medicaid or evidence of coverage documents.

# **IV. Appendices/General Information**

Appendix A: Abbreviation/Acronym Key	
FC: functional class	PAH: pulmonary arterial hypertension
FDA: Food and Drug Administration	PH: pulmonary hypertension
NYHA: New York Heart Association	WHO: World Health Organization

# Appendix B: Therapeutic Alternatives

This table provides a listing of preferred alternative therapy recommended in the approval criteria. The drugs listed here may not be a formulary agent and may require prior authorization.

Drug Name	Dosing Regimen	Dose Limit/ Maximum Dose
nifedipine (Adalat <sup>®</sup> CC, Afeditab <sup>®</sup> CR, Procardia <sup>®</sup> , Procardia XL <sup>®</sup> )	60 mg PO QD; may increase to 120 to 240 mg/day	240 mg/day
diltiazem (Dilacor XR <sup>®</sup> , Dilt-XR <sup>®</sup> , Cardizem <sup>®</sup> CD, Cartia XT <sup>®</sup> , Tiazac <sup>®</sup> , Taztia XT <sup>®</sup> , Cardizem <sup>®</sup> LA, Matzim <sup>®</sup> LA)	720 to 960 mg PO QD	960 mg/day



Drug Name	Dosing Regimen	Dose Limit/ Maximum Dose	
amlodipine (Norvasc <sup>®</sup> )	20 to 30 mg PO QD	30 mg/day	

*Therapeutic alternatives are listed as Brand name*<sup>®</sup> (generic) when the drug is available by brand name only and generic (Brand name<sup>®</sup>) when the drug is available by both brand and generic.

#### Appendix C: Contraindications/Boxed Warnings

- Contraindication(s):
  - Congestive heart failure due to severe left ventricular systolic dysfunction
  - Pulmonary edema
  - Hypersensitivity to the drug or to structurally related coumpounds
- Boxed warning(s): none reported

#### Appendix D: Pulmonary Hypertension: WHO Classification

- Group 1: PAH (pulmonary arterial hypertension)
- Group 2: PH due to left heart disease
- Group 3: PH due to lung disease and/or hypoxemia
- Group 4: CTEPH (chronic thromboembolic pulmonary hypertension)
- Group 5: PH due to unclear multifactorial mechanisms

Treatment Approach*	FC	Status at Rest	Tolerance of Physical Activity (PA)	PA Limitations	Heart Failure
Monitoring for progression of	Ι	Comfortable at rest	No limitation	Ordinary PA does not cause undue dyspnea	
PH and				or fatigue, chest pain,	
treatment of co- existing conditions				or near syncope.	
conditions	II	Comfortable	Slight	Ordinary PA causes	
		at rest	limitation	undue dyspnea or	
				fatigue, chest pain, or	
Advanced treatment of PH	III	Comfortable	Marked	near syncope. Less than ordinary PA	
with PH-	111	at rest	limitation	causes undue dyspnea	
targeted therapy				or fatigue, chest pain,	
- see Appendix				or near syncope.	
F**	IV	Dyspnea or	Inability to	Discomfort is	Signs
		fatigue may	carry out any	increased by any PA.	of right
		be present at	PA without		heart
*DI1		rest	symptoms		failure

#### Appendix E: Pulmonary Hypertension: WHO/NYHA Functional Classes (FC)

\*PH supportive measures may include diuretics, oxygen therapy, anticoagulation, digoxin, exercise, pneumococcal vaccination. \*\*Advanced treatment options also include calcium channel blockers.



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Mechanism of Action	Drug Class	Drug Subclass	Drug	Brand/Generic Formulations
	Prostacyclin* pathway agonist	Prostacyclin	Epoprostenol	Veletri (IV) Flolan (IV) Flolan generic (IV)
	*Member of the prostanoid class of fatty acid derivatives.	Synthetic prostacyclin analog	Treprostinil	Orenitram (oral tablet) Remodulin (IV) Tyvaso (inhalation)
			Iloprost	Ventavis (inhalation)
Reduction of pulmonary arterial		Non-prostanoid prostacyclin receptor (IP receptor) agonist	Selexipag	Uptravi (oral tablet)
pressure through	Endothelin receptor antagonist (ETRA)	Selective receptor antagonist	Ambrisentan	Letairis (oral tablet)
vasodilation		Nonselective dual action receptor	Bosentan	Tracleer (oral tablet)
		antagonist	Macitentan	Opsumit (oral tablet)
	Nitric oxide- cyclic guanosine monophosphate enhancer	Phosphodiesterase type 5 (PDE5) inhibitor	Sildenafil	Revatio (IV, oral tablet, oral suspension)
			Tadalafil	Adcirca (oral tablet)
		Guanylate cyclase stimulant (sGC)	Riociguat	Adempas (oral tablet)

Appendix F: Pulmonary Hypertension: Targeted Therapies

# V. Dosage and Administration

Drug Name	Dosing Regimen	Maximum Dose		
Epoprostenol (Flolan)	l (Flolan) 2 ng/kg/min IV, increased by 1-2 ng/kg/min			
	at intervals of at least 15 minutes	response		
Epoprostenol (Veletri)	2 ng/kg/min IV, increased by 2 ng/kg/min	Based on clinical		
	every 15 minutes or longer	response		

# VI. Product Availability

Drug Name	Availability
Epoprostenol (Flolan)	Vial with powder for reconstitution: 0.5 mg, 1.5 mg
Epoprostenol (Veletri)	Vial: 0.5 mg/10 mL, 1.5 mg/10 mL



# VII. References

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# **Coding Implications**

Codes referenced in this clinical policy are for informational purposes only. Inclusion or exclusion of any codes does not guarantee coverage. Providers should reference the most up-to-date sources of professional coding guidance prior to the submission of claims for reimbursement of covered services.

HCPCS Codes	Description	
J1325	Injection, epoprostenol, 0.5 mg	



Reviews, Revisions, and Approvals	Date	P&T Approval Date
FC II is added to the prostanoid class of PH drugs. Safety criteria were removed unless they 1) represent contraindications or black box warnings not covered by a REMS program, and 2) provide specific lab/imaging parameters that must be met prior to initiation of therapy. An efficacy statement is added to the continuation criteria. Initial and continuation durations increased to 6 and 12 months respectively. Appendices covering PH group, functional class and therapy reorganized.	02.17	03.17
1Q18 annual review: Policies combined for commercial and Medicaid; No significant changes from previous corporate approved policy; Medicaid: removed WHO/NYHA classifications from initial criteria since specialist is involved in care; References reviewed and updated.	11.21.17	02.18
1Q 2019 annual review: no significant changes; references reviewed and updated.	11.20.18	02.19
Added HIM line of business due to addition of agent(s) to the HIM formulary with PA	03.15.19	
1Q 2020 annual review: no significant changes; added statement that treatment plan detailing dose, quantity, and frequency; references reviewed and updated.	11.26.19	02.20
1Q 2021 annual review: no significant changes; references to HIM.PHAR.21 revised to HIM.PA.154; references reviewed and updated.	10.12.20	02.21

# **Important Reminder**

This clinical policy has been developed by appropriately experienced and licensed health care professionals based on a review and consideration of currently available generally accepted standards of medical practice; peer-reviewed medical literature; government agency/program approval status; evidence-based guidelines and positions of leading national health professional organizations; views of physicians practicing in relevant clinical areas affected by this clinical policy; and other available clinical information. The Health Plan makes no representations and accepts no liability with respect to the content of any external information used or relied upon in developing this clinical policy. This clinical policy is consistent with standards of medical practice current at the time that this clinical policy was approved. "Health Plan" means a health plan that has adopted this clinical policy and that is operated or administered, in whole or in part, by Centene Management Company, LLC, or any of such health plan's affiliates, as applicable.

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contract of insurance, etc.), as well as to state and federal requirements and applicable Health Plan-level administrative policies and procedures.

This clinical policy is effective as of the date determined by the Health Plan. The date of posting may not be the effective date of this clinical policy. This clinical policy may be subject to applicable legal and regulatory requirements relating to provider notification. If there is a discrepancy between the effective date of this clinical policy and any applicable legal or regulatory requirement, the requirements of law and regulation shall govern. The Health Plan retains the right to change, amend or withdraw this clinical policy, and additional clinical policies may be developed and adopted as needed, at any time.

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#### Note:

**For Medicaid members**, when state Medicaid coverage provisions conflict with the coverage provisions in this clinical policy, state Medicaid coverage provisions take precedence. Please refer to the state Medicaid manual for any coverage provisions pertaining to this clinical policy.

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