# SENTARA COMMUNITY PLAN (MEDICAID)

### 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 1-800-750-9692</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>

## Weight Loss Management

**Drug Requested:** (check box below that applies)

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PREFERRED MEDICATIONS					
*Requires prior					
□ Adipex-P®/Suprenza <sup>™</sup> (phentermine HCl)	u	Xenical® (orlistat)			
□ Bontril®/Bontril PDM® (phendimetrazine)		Radtue® (diethylpropion)			
□ <b>Didrex</b> ®/ <b>Regimex</b> ® (benzphetamine)					
NON-PREFERRED	M	EDICATIONS			
*Saxenda, Wegovy, Zepbound requires trial and failure of one (1) oral preferred agent For Wegovy and Zepbound: the member has tried and failed the selected product, Saxenda®, as indicated on the PDL.					
□ Saxenda <sup>®</sup> (liraglutide) *ages 12 and older		Wegovy® (semaglutide) *ages 12 and older			
☐ Imcivree <sup>™</sup> (setmelanotide) *ages 2 and older		<b>Zepbound</b> <sup>™</sup> (tirzepatide) *ages 18 and older			
MEMBER & PRESCRIBER INFORMATIO	<b>N:</b>	Authorization may be delayed if incomplete.			
Member Name:					
Member Sentara #:		Date of Birth:			
Prescriber Name:					
Prescriber Signature: Date:					
Office Contact Name:					
Phone Number:		Fax Number:			
NPI #:					
<b>DRUG INFORMATION:</b> Authorization may be d	alar	rod if in commutate			
DRUG INFORMATION. Authorization may be d	eray	red if incomplete.			
Drug Name/Form/Strength:					
Dosing Schedule:		Length of Therapy:			
Diagnosis:		ICD Code, if applicable:			
Weight (if applicable):	Date weight obtained:				

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**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.

<u>Initia</u>	l Request Req	uirements:				
	Provider must submit current height and weight measurements (verified by chart notes)					
	Height:	Current Weight:	BMI:	Date:		
Cove	rage for all me	edications will be limited to th	e following:			
Absen	ce of medical co	ntraindications:				
	No contraindicate products); <b>AND</b>	tions to use; (i.e. uncontrolled hypert	ension, hyperthyroidis	m etc. for stimulant based		
	No malabsorption syndromes, cholestasis, pregnancy and/or lactation (for orlistat); <b>AND</b>					
	No history of an eating disorder (e.g., anorexia, bulimia); <b>AND</b>					
	No acute pancreatitis, acute suicidal behavior/ideation, personal or family history of medullary thyroid cancer or multiple endocrine neoplasia 2 syndrome (if requesting a GLP-1 Receptor Agonists); <b>AND</b>					
Addit	ional qualifying	criteria to include the following: (e	excludes Imcivree <sup>TM</sup> )			
	Participation in nutritional counseling; <b>AND</b>					
	Participation in physical activity program, unless medically contraindicated; <b>AND</b>					
	Commitment to	continue the above weight-loss treat	ment plan; AND			
-	rovider attests th or high-morbidit	nat the patient's obesity is disabling conditions):	g and life threatening	(i.e., puts the patient at		
	Yes					
	No					
Writt		n must include the following:				
	specific reduced	status and weight loss plan. An indi calorie meal plan, recommended rolle modification as needed to improve	utine physical activity,	and behavioral intervention		
	Current dated ac	curate height and weight measurement	ents			
	nent details of <u>pr</u> tting a copy of th	revious weight loss treatment plans ne plan:	s to include diet and e	xercise plans, in addition to		
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## Drug Specific Requirements (Minimum ages are per FDA approvals):

1.	. For phentermine (min age 17), phendimetrazine tablet (min age 18), phendimetrazine ER capsule (min age 17) and orlistat (min age 12):						
	Body mass index (BMI) $\geq 30 \text{kg/m}^2$ , if no applicable risk factors; <b>OR</b>						
		coronary heart disease	□ dyslipidemia	□ hypertension	□ sleep apnea	☐ Type II Diabetes	
2.		For benzphetamine (min age 17), diethylpropion (min age 16):					
		Body mass index (BM	,				
3.	Foi	r Imcivree® (min age 2)	)				
		BMI $\geq$ 30 kg/m <sup>2</sup> ; <b>AN</b>	D				
	□ Prescribed by or in consultation with an endocrinologist or geneticist; <b>AND</b>						
	□ Member has Bardet-Biedl syndrome (BBS); <b>OR</b>						
	☐ Member has proopiomelanocortin (POMC), proprotein convertase subtilisin/kexin type 1 (PCSK1), or leptin receptor (LEPR) deficiency, as confirmed by a genetic test; <b>AND</b>						
		☐ Member's genetic variants are interpreted as pathogenic, likely pathogenic, or of uncertain significance (VUS)					
4.	For GLP-1 receptor agonists indicated for weight loss (Wegovy/Saxenda min age 12, Zepbound Min age 18):					12, Zepbound	
	$\square$ BMI > 40 kg/m <sup>2</sup> , if no applicable risk factors; <b>OR</b>						
	$\square$ BMI > 37 kg/m <sup>2</sup> with one or more of the following risk factors:						
		□ dyslipidemia	□ h	ypertension	☐ Type II Diabete	S	
	<ul> <li>□ Member has tried and failed one of the non-GLP1 weight-loss medications*; <b>OR</b></li> <li>□ Member is intolerant to all non-GLP1 weight-loss medications*; <b>AND</b></li> </ul>						
		☐ The member has tried and failed* the selected product, Saxenda®, as indicated on the					
	PDL: https://www.virginiamedicaidpharmacyservices.com/provider/preferred-drug-list/						
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*Definitions of Accepted Drug Trial			
Drug	Trial		
Benzphetamine, diethylpropion, phendimetrazine, phentermine	3-month trial without a weight loss of 10lbs		
Orlistat	6-month trial without a weight loss of 10lbs		
GLP-1 Receptor Agonist	6-month trial without a body weight reduction of 5%		

#### **Length of Authorization:**

- ☐ <u>Initial request</u>: Varies (drug specific)
  - Benzphetamine, diethylpropion, phendimetrazine, phentermine -- 3 months
  - GLP-1 receptor agonists 6 month
  - Xenical®(orlistat) 6 months
  - Imcivree<sup>™</sup> -- 4 months
- ☐ Renewal requests: Renewals will no longer be granted once a member reaches a BMI < 25. See additional requirements below (drug specific):
  - **Benzphetamine**, **diethylpropion**, **phendimetrazine**, **phentermine** If patient achieves at least a 10-lb weight loss during initial 3 months of therapy, an additional 3-month prior authorization may be granted. Maximum length of continuous drug therapy = 6 months (waiting period of 6 months before next request)
  - Alli®/Xenical® (orlistat)- If patient achieves at least a 10-lb weight loss, an additional 6-month prior authorization may be granted. Maximum length of continuous drug therapy = 24 months (waiting period of 6 months before next request)
  - **Imcivree**<sup>™</sup> If the member has experienced ≥ 5% reduction in body weight (or ≥ 5% of baseline BMI in those with continued growth potential), an additional 1-year prior authorization may be granted
  - GLP-1 Receptor Agonists If the member achieves a weight loss of  $\geq 5\%$  reduction in body weight compared to the most recent authorization, an additional 6-month SA may be granted

Current measu	rements: BMI (Adult	t) or % of 95" perce	entile weight (12-17)	<b>7/0):</b>
Height:	Weight:	BMI:	DATE:	(attach chart notes)
Previous author	rization measuremen	ts: BMI (Adult) or	% of 95 <sup>th</sup> percentile	weight (12-17 y/o):
Height:	Weight:	BMI:	DATE:	(attach chart notes)

All approvals are subject to the criteria on this form. Existing authorizations will be honored until renewal.

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

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