SENTARA HEALTH PLANS

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 may be delayed.</u>

Drug Requested: Gattex® (teduglutide [rDNA Origin]) Injection

request may be denied.

Member Name:				
	Date of Birth:			
Prescriber Name:				
	Date:			
Office Contact Name:				
Phone Number:	Fax Number:			
DEA OR NPI #:				
	authorization may be delayed if incomplete.			
Drug Form/Strength:				
Dosing Schedule:	Length of Therapy:			
Diagnosis:	ICD Code, if applicable:			
Member's current weight:	kg			
Final dose per day:	mg (Max 3.8mg dose per vial)			
SrCr:	r renal impairment [CrCl <50ml/min] dose must be reduced by 50%)			
Recommended Dosage:				
Recommended Dosage: Maximum approval for adults	.05mg/kg once daily			

☐ Member has been dependent on parenteral nutrition/intravenous fluids (PN/IV) therapy \geq 3 times per week for \geq 12 continuous months and failed previous trials of weaning (attach supportive documentation demonstrating the requirement of parenteral support)

Initial Approval Length - 6 months. (All information must be noted or submitted with request form.)

(Continued on next page)

	• Frequency of current PN/IV use:	/week					
	Baseline of volume:	L/week or per infusion					
	• Member's Body Mass Index (BMI): _	kg/m ²					
	AND						
	Member must have diagnosis of short bow	rel syndrome					
	<u>OR</u>						
	•	ease with documentation of clinical remission of Crohn's on demonstrating the clinical remission of Crohn's disease)					
	AND						
	Member received a colonoscopy or alternation of therapy	ate imaging with removal of polyps (if necessary) within six (6					
	Date of colonoscopy (must be within 6 months):						
EXC	LUSIONS:						
•	Age <1 year old OR ≤ 10 kg Diagnosis of active cancer within the last 5 Body Mass Index (BMI) is <15 kg/m ² Member received human growth hormone Member has had four or more SBS-related Member has an active intestinal obstruction	(e.g. Zorbtive) within the last 6 months hospital admissions within the last 12 months					
Firs	t Continuation of Therapy - 6 mont	ths. (All lines below must be completed)					
•	Has member had at least 20% reduction from	om baseline in parenteral nutrition/intravenous fluid (PN/IV)? □ YES □ NO					
•	Frequency of current PN/IV use:	/week					
•		L/week or per infusion (supportive documentation must					
•	Member's Body Mass Index (BMI):	kg/m^2					
•	Member does not have any FDA labeled c	ontraindications to therapy:					
•	Labs must be submitted every six (6) months (supportive documentation must be atta	ths and colonoscopy one (1) year after initiation of therapy ached)					
	ond Continuation of Therapy - 1 year to be completed)	ar after initial approval: 6 months. (All lines below					

• Has i	member had at least 20% redu	action from last parenteral nutrition/intrave		d (PN/ YES		NO				
• Freq	uency of current PN/IV use:		/week							
• Volu										
Member's Body Mass Index (BMI):			$_{\rm max}$ kg/m ²							
• N	Member does not have any FD	A-labeled contraindications to therapy:		YES		NO				
	 Labs must be submitted every 6 months and colonoscopy one 1 year after initiation of therapy and then every 5 years after (supportive documentation must be attached) 									
Continua need to be	1 4 1)	years after initial approval: <u>6 mor</u>		All lines	belov	W				
	*	utrition/intravenous fluid (PN/IV) stabilized NO is checked, it will be denied)		increa YES	sed fr					
• Freque	Frequency of current PN/IV use:/week									
	Volume: L/week or per infusion (supportive documentation must be attached)									
• Mem	iber's Body Mass Index (BM	I):	kg/m	12						
• Mem	iber does not have any FDA-l	abeled contraindications to therapy:		□ YI	ES (□ NO				
	•	nonths and colonoscopy 1 year after initial locumentation must be attached)	ation of (therap	y and	then				
Medication	on being provided by Sr	oecialty Pharmacy - PropriumRx								
If a dru		gs may be covered under every Plan Plan, documentation of medical nec		vill be	requ	uired.				

**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. *

^{*}Approved by Pharmacy and Therapeutics Committee: 11//20/2014
REVISED/UPDATED/REFORMATTED: 4/26/2015; 5/21/2015; 12/27/2015; 12/26/2016; 8/43/2017; 3/31/2018; 10/14/2019