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 can be delayed.</u>

NON-PREFERRED

Acthar® Gel (repository corticotropin) 80 USP

□ **Acthar**[®] **Gel** (repository corticotropin) 40 USP Units/0.5 mL single-dose prefilled SelfJect

Units/mL 5 mL multi-dose vial

Drug Requested: Repository Corticotropin Medications - Nephrotic Syndrome (NS)

PREFERRED

□ Purified Cortrophin[™] Gel

(repository corticotropin)

adults only.

	injector	
	Acthar® Gel (repository corticotropin) 80 USP	
	Units/mL single-dose prefilled SelfJect injector	
	*Member must have tried and failed preferred Purified Cortrophin [™] Gel and meet all applicable PA criteria below	
MEMBER & PRESCRIBER INFORM	IATION: 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 #:		
DRUG INFORMATION: Authorization r	may be delayed if incomplete.	
Drug Name/Form/Strength:		
osing Schedule: Length of Therapy:		
Diagnosis:	sis: ICD Code, if applicable:	
eight (if applicable): Date weight obtained:		

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Acthar Gel single-dose pre-filled SelfJect injector is for subcutaneous administration by

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. Check box below for the Diagnosis that applies.

Member MUST have a documented diagnosis of Nephrotic Syndrome with ONE of the following			
□ Focal Segmental Glomerulosclerosis (FSGS) OR □ Membranous Nephropathy (MPGN)			
☐ Minimal Change Disease			
The following MUST be noted:			
1. Baseline current kg:			
2. Baseline (prior to corticosteroid and calcineurin inhibitor) urine protein/creatinine ratio with collection date:; mg/mg (> 3-3.5 mg/mg nephrotic range proteinuria)			
Member <u>MUST</u> have tried and failed both a corticosteroid <u>AND</u> a calcineurin inhibitor (CNI) taken concurrently within the year of request. Failure is defined as no change or an increase fro baseline proteinuria levels after 90 consecutive days of concomitant corticosteroid and calcineur therapy trial. Approval will be based on proteinuria increase from baseline after 90 consecutive days of concomitant corticosteroids and calcineurin inhibitor therapy.			
3. 90 days post concurrent corticosteroid and calcineurin inhibitor trial, urine protein/creatinine ratio;			
Date:;(mg/mg nephrotic range proteinuria)			
□ 1 mg/kg (max 80 mg)			
AND			
☐ Member MUST have had concurrent trial and failure of calcineurin inhibitor for a minimum of 90 days consecutive days within last 12 months (must note therapy tried and trial MUST be no in pharmacy paid claims):			
□ Cyclosporine □ Tacrolimus □ Cyclophosphamide			
OR			
If member has a relative <u>contraindication or intolerance to high dose corticosteroids</u> (e.g., uncontrolled diabetes BS > 200, or GI BLEED within the last 30 days):			
Member has had trial and failure of calcineurin inhibitor only (therapy tried MUST be noted in			
pharmacy paid claims):			
Cyclosporine: mg (4 to 5 mg/kg/day in 2 divided doses for at least 12 months OR mg/m²/day in 2 divided doses; adjust doses based on trough levels {(pediatrics): 80 to 100 ng			
☐ Tacrolimus: mg			
□ Cyclophosphamide: mg			

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PA Repository Corticotropin-NS (CORE)
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☐ Progress notes MUST be su	omitted with documentation of ALL	THREE (3) of the following labs:			
□ Proteinuria	□ Serum Albumin	□ Cyclosporine levels			
Dose Regimen: Anticipated Length of therapy:					
NOTE: Approval will be for a period of 6 weeks with a follow up Proteinuria lab required to be submitted. IF additional therapy is needed; the prescribing physician will need to submit a second request for continuation of therapy.					
Medication being provided b	v Snecialty Pharmacy – Pronriu	ım Rv			
Medication being provided by Specialty Pharmacy – Proprium Rx					

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 pharmacy paid claims or submitted chart notes.