## SENTARA HEALTH PLANS

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

<u>Prections:</u> The prescribing physician must sign and clearly print name (preprinted stamps not valid) 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 information provided is not complete, correct, or legible, authorization may be delayed.</u>

**NON-PREFERRED** 

□ HP Acthar® Gel (repository corticotropin)

\*Member must have tried and failed preferred

**Drug Requested: Repository Corticotropin Medications – Ocular Diseases** 

**PREFERRED** 

□ Purified Cortrophin<sup>™</sup> Gel (repository corticotropin)

	Purified Cortrophin <sup>™</sup> Gel and meet all applicable PA criteria below			
MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete.				
Member Name:				
Member Sentara #:				
Prescriber Name:				
Prescriber Signature:				
Office Contact Name:				
Phone Number:				
DEA OR NPI #:				
DRUG INFORMATION: Authoriz				
Drug Form/Strength:				
Dosing Schedule:				
Diagnosis:	ICD Code, if applicable:			
Weight:	Date:			

Adverse effects that may occur with repository corticotropin are related primarily to its steroidogenic effects and are similar to corticosteroids. There may be increased susceptibility to new infection and increased risk of reactivation of latent infections.
 Adrenal insufficiency may occur after abrupt withdrawal of the drug following prolonged therapy.

**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.

## PA Repository Corticotropin\_Ocular Diseases (Continued from previous page)

SECTION A:							
Slit lamp examination used to r	□ Yes	□ No					
Intraocular pressure (IOP) mea	□ Yes	□ No					
Baseline IOP results							
Visual Acuity Test results _							
Labs and documentation to rule	Labs and documentation to rule out infectious etiology						
Anterior Chamber cells present	t?		□ Yes	□ No			
SECTION B:  PREDNISONE MUST HAVE B IMMUNOSUPPRESIVE DRUC THE LAST 12 MONTHS.							
☐ Please note therapy tried (pa failure of prednisone plus co							
□ methotrexate	□ cyclosporine	□ mycophenolate	<ul><li>azathioprir</li></ul>	ne			
□ cyclophosphamide	□ tacrolimus	□ sirolimus	Other:				
<ul> <li>NON-INFECTIOUS</li> <li>approval. To support each chart notes, must be provided</li> <li>Initial Authorization: 3 n</li> </ul>	line checked, all docume led or request may be der	entation, including lab res					
Use of repository corticotropin injection is considered <b>NOT medically necessary</b> as treatment of corticosteroid responsive conditions. <b>Please note member's diagnosis</b> . **NOTE if member is only diagnosed with Anterior Uveitis additional comorbidities including etiology will be required for approval**							
□ Anterior Uveitis □	Intermediate Uveitis	□ Posterior Uveitis	□ Pan Uveiti	S			
☐ Is this member positive for I	HLA-B27 antigen?		□ Yes	□ No			
Please include other diagnosis that contributes to Anterior Uveitis <b>ONLY</b> diagnosis:							
□ Completed SECTION A  AND							

	PAID CLAIMS MUST MATCH	STATEMENT BELOW:			
	Member must have tried and failed the therapies below for at least 3 months consecutively within the last 12 months. Failure will be defined as no improvement in symptoms while on high dose corticosteroid <b>and</b> immunosuppressant agent concomitantly. Please note therapies tried:				
	Member tried and maximized topic	Member tried and maximized topical steroid treatment for at least 4 weeks resulting in ineffective therapy:			
	□ prednisolone acetate (Pred Forte <sup>®</sup> )	□ difluprednate (Durezo	ol®) 🗖 lotepred	dnol (Lotemax®)	
	☐ Fluoromethalone (FML®)	<ul><li>Dexamethasone</li></ul>	Other:		
	AND				
	Prednisone 1 mg/kg/day oral (or an	n equivalent high dose steroid	1)		
	Name, dose and dates of the equiva-	alent high does steroid trials:			
	AND				
	Completed SECTION B				
	AND				
	☐ Member tried and failed at least 2 different BIOLOGICS for a minimum of 3 months due to toxicity OR failure to stabilize disease. Submit supporting document on toxicities and progression. (Include labs - CBC, BUN, SCr, AST, ALT and albumin). Check ALL that apply:				
	□ adalimumab (Humira <sup>®</sup> ) □	etanercept (Enbrel®)	□ infliximab	□ rituximab	
	□ golimumab (Simponi <sup>®</sup> ) □	tocilizumab (Actemra®)	□ IVIG	□ Other:	
	AND				
	CBC, CMP, HbA1C, TB, Hepatitis therapy have been submitted	s B and C labs collected prior	to initiation of re	pository corticotropin	
	AND				
	Medication is prescribed by an oph	thalmologist or rheumatolog	ist		
	NON-INFECTIOUS KERA approval. To support each line cl chart notes, must be provided or a	hecked, all documentation, in			
*:	*Note approval will not excee	ed 16 weeks for this indi	ication**		
	Complete SECTION A				
	AND				
	Provider attests all infectious etiologies have been ruled out (e.g., bacterial, fungal or viral eye infection)  (Attach labs and culture and sensitivity reports to support)				
	AND				

	Positive fluorescein stain has been obtain	ned			
	AND Corneal Scraping used to stain and cultur	ro er	agaiman has baan gamplatad t	0 111	le out infactious atiologies
	AND	ic s <sub>k</sub>	becimen has been completed t	J I U.	le out infectious etiologies
	☐ prednisolone acetate (Pred Forte <sup>®</sup> )		difluprednate (Durezol®)		loteprednol (Lotemax®)
	☐ Fluoromethalone (FML®)		Artificial tears		Cyclosporine (Restasis®)
	□ Dexamethasone		Other:		
	AND				
	Medication is prescribed by an ophthalm	olog	gist		
	<b>OPTIC NEURITIS.</b> Check below each line checked, all documentation, i provided or request may be denied.				
*	*Note approval will not exceed 14	da	ys for this indication**		
	☐ MRI of brain and orbital region has been completed for lesions consistent with MS and visualizing the optic chiasm for negative pituitary tumors (submit imaging results)				
	AND				
	Provider attests all other primary etiolog	ies l	nave been ruled out (e.g., infec	ctiou	us, neuromyelitis optica)
	AND				
	Member is contraindicated or has failed	metl	nylprednisolone IV use for 3-:	5 da	ys
	AND				
	Member is contraindicated or has failed omethylprednisolone	oral	prednisone (1 mg/kg) use for	2 w	eeks after IV
	AND				
	Member tried and failed IVIG for a mini	mur	n of 3 months		
	Proof of inability to improve vision with				·
	Visual Acuity Baseline:				
	Contrast Sensitivity:				
	other ophthalmic distance approval. To support each line checked chart notes, must be provided or requestions below including literations.	d, al st m	l documentation, including la ay be denied. <b>(Please subm</b>	b res	sults, diagnostics, and/or upporting document to

	Diagnosis:
	AND
	Completed SECTION A
	AND
	Completed SECTION B
	Other previously failed therapies along with dates tried have been documented below:
	<b>REAUTHORIZATION FOR NON-INFECTIOUS UVEITIS.</b> 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.
If	f member is in remission for 2 years reduced dose is indicated until discontinuation
	Completed SECTION A
	AND
	Signs and symptoms have improved within 3 months of use (Submit supporting labs and documentation)  Current IOP results:
	Current acuity:
	Anterior Chamber cells present?
	AND
	No toxicities or severe adverse reactions have developed
	AND
	Medication is prescribed by a specialist in treatment of the disease/condition (rheumatologist or ophthalmologist)
N	Tedication 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. \*\*