

Please complete and email or fax toll-free

Phone: 888-435-2284

FAX: 877-937-2284 EMAIL: intake@supportandaccess.com

PATIENT & PRESCRIBER PATIENT INFORMATION	(repository corticotrop	oin inje	ction) 80 U/mL		FAX: 877-937-2284 EMAIL: intake@su	pportandaccess.com	
Last Name:		Ac	Idress:				
First Name and Middle Initial:			ty:				
Date of Birth:			-	·γο.			
		State: Zip Code:					
Mobile Phone: Phone:			Email:				
Preferred Language (other than English):		Ca	aregiver:				
Allergies: NKDA - No known drug allergies					Patient Sex:	Male Fen	
(Additional space provided on pg 2)	Pharmacy Benefit Provider:	acy Benefit Provider:		Primary M	Medical Insurance:		
INSURANCE INFORMATION Include a copy of the front and back of the patient's	Subscriber #:		Subscribe				
rescription benefit and insurance card(s) when	Group #:		Group				
submitting this form OR complete the fields to the right.	Phone #:		Phone #:				
PRESCRIBER INFORMATION		1					
HCP Name:		NF	PI #:		Tax ID #:		
Specialty:		Of	Office Contact Name:				
Address:		Co	Contact Phone:		Extensi	Extension:	
City: State:	Zip Code:	Co	ntact Fax:				
State License Number:		Co	Contact Email:				
PRESCRIPTION: ACTHAR® GEL SUBCUTANEOUS INJECTION ICD-10 Code (Required):	Preferred Spot (SEE PG 2 FOR PRIMARY DIAGNIFOR A MORE COMPLETE LIST, SI	OSIS CO	DES;	E LIST])		PLEASE ENROLL PATIEN ACTHAR PATIENT SUPPO	
B0 Units/mL NDC 63004-8711-4				Units y 24 hours	Syringe: 1 ML Needle for Drawin Needle for Injectic Sharps Container OTHER: Pharmacy to dispense st course of therapy. Pharmalternate supplies as nec	nn: 25 G, 5/8" Ifficient supplies to complet acist may elect to dispense	
COMMERCIAL STARTER PROGRAM ICD-10 Code (Required):	The Acthar Gel	Comm			ligible, commercially-insure		
Acthar Gel Single-Dose Pre-filled SelfJect™ Injector Subcutaneous Injection Acthar Gel 5 mL multi-dose vial (80 USP Units/mL): NDC 63004- Subcutaneous Injection			IDC 63004-8710-1	following unless "C	Pharmacy to supply to THER" is specified		
80 Units/mL NDC 63004-8711-4 40 Units/0.5 mL NDC 6300	94-8712-4 80 Units 4	80 Units 40 Units Other: Ur		Units	Syringe: 1 ML		
Frequency:	Frequency:	Frequency:		Needle for Drawing: 20 G			
Every 72 hours Every 48 hours Every 24	4 hours Every 72 hours	s Every 72 hours Every 48		ry 48 hours Every 24 hours		Needle for Injection: 25 G, 5/8"	
Other:	Other:	Other			Sharps Container		
Number of Refills:	Number of Refills:				Pharmacy to dispense su course of therapy. Pharm alternate supplies as nec	ufficient supplies to complet acist may elect to dispense essary.	
THER INSTRUCTIONS: (Attach taper schedule and pro	vide additional instructions, if appli	cable)					
RESCRIBER SIGNATURE: Please sign only ONI		,	agreeing to the Droseriha	r Consont oo	action on page 4 of this do	cument)	
Brand Medically Necessary / Do Not Substitute / No Substitution	(,, , , ,	Ju ale	May Substitute / Product There is no A/B rated sub	Selection Perr	mitted / Substitution Permissi har. This space is required by	ble	
X Disperse of Weither	D.J.	or	X			Data	
Dispense as Written	Date		Substitutions Allowed			Date	

Prescriber signature required for consent and to validate prescriptions. Prescriber attests that this is her/his signature. NO STAMPS. By signing, Prescriber certifies that the above is medically necessary. ATTN: New York and Iowa providers, please submit electronic prescription. CA, MA, NC & PR: Interchange is mandated unless Prescriber writes the words "No Substitution." For questions, please call: 1-888-435-2284 Monday through Friday (8:00 AM to 9:00 PM ET) Saturday (9:00 AM to 2:00 PM ET).



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DIAGNOSIS AND MEDICAL INFORMATION \ Patient N

	2.11. 112. 11.10.10@50pportariod050505105111
Patient Name:	Date of Birth:

THESE CODES HAVE BEEN PROVIDE	D FOR CONVENIENCE ON	LY. THESE ARE NOT ALL POSS	IBLE DIAGNOSIS CODES, AND N	I BE FOUND IN APPENDIX A, PAGES 6 AND 7. IOT INTENDED TO INFLUENCE A DIAGNOSIS. agnosis in the "OTHER DIAGNOSIS" section.	
☐ NEUROMYELITIS OPTICA [DEVIC] G36.0	l	☐ DIFFUSE INTERSTITIAL KEI	RATITIS, LEFT EYE	UNSPECIFIED CHORIORETINAL INFLAMMATION	
UNSPECIFIED SCLERITIS, UNSPE	ECIFIED EYE	DIFFUSE INTERSTITIAL KE	RATITIS, BILATERAL	H30.93 ☐ RETINAL VASCULITIS, BILATERAL	
SCLERITIS WITH CORNEAL INVOING RIGHT EYE	LVEMENT,	OTHER KERATITIS H16.8		H35.063 ☐ PANUVEITIS, RIGHT EYE	
H15.041 UNSPECIFIED SUPERFICIAL KERATITIS, BILATERAL H16.103 FILAMENTARY KERATITIS, BILATERAL H16.123 PUNCTATE KERATITIS, RIGHT EYE H16.141		☐ PRIMARY IRIDOCYCLITIS, LEFT EYE H20.012 ☐ RECURRENT ACUTE IRIDOCYCLITIS, LEFT EYE H20.022 ☐ SECONDARY NONINFECTIOUS IRIDOCYCLITIS, RIGHT EYE		H44.111 ☐ PANUVEITIS, LEFT EYE	
				H44.112 ☐ PANUVEITIS, BILATERAL	
				H44.113 SYMPATHETIC UVEITIS, UNSPECIFIED EYE	
		H20.041		H44.139 RETROBULBAR NEURITIS, RIGHT EYE	
□ PUNCTATE KERATITIS, LEFT EYE H16.142		CHRONIC IRIDOCYCLITIS, RIGHT EYE H20.11		H46.11 ☐ RETROBULBAR NEURITIS, LEFT EYE	
□ PUNCTATE KERATITIS, BILATERA H16.143	L	CHRONIC IRIDOCYCLITIS, LEFT EYE H20.12 CHRONIC IRIDOCYCLITIS, BILATERAL		H46.12	
☐ OTHER KERATOCONJUNCTIVITIS, BILATERAL H16.293 ☐ DIFFUSE INTERSTITIAL KERATITIS, RIGHT EYE H16.321		H20.13 UNSPECIFIED IRIDOCYCLI		H46.8 UNSPECIFIED OPTIC NEURITIS	
		H20.9		H46.9 OTHER DIAGNOSIS:	
HISTORY OF CORTICOSTEROID USE	E (if applicable). Please a	dd details in the section below	I		
PLEASE CHECK ALL THAT APPLY: A corticosteroid was tried with the following r Corticosteroid use failed, but same respo Patient hypersensitive or allergic to cortic Patient intolerant of corticosteroids Other:	nse not expected with Acthar	OR	A corticosteroid was not tried due to Corticosteroid use is contraindice Intravenous access is not possib Patient has known intolerance to Other:	ated for this patient le for this patient	
CONCURRENT MEDICATIONS					
	RY (Including recent c	orticosteroid history. Attac	h additional case notes as ı	necessary.)	
RELEVANT TREATMENT HISTO				EVEL AND OUTCOME MUTU DETAIL	
RELEVANT TREATMENT HISTO THERAPY NAME	DOSE	START DATE	STOP DATE (if applicable	EXPLAIN OUTCOME WITH DETAIL (eg. type of outcome)	
	DOSE	START DATE	STOP DATE (if applicable		
	DOSE	START DATE	STOP DATE (if applicable		
	DOSE	START DATE	STOP DATE (if applicable		
THERAPY NAME				(eg. type of outcome)	
	RY (Including recent co			(eg. type of outcome)	

Y

Signature



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EMAIL: intake@supportandaccess.com

FOR COMPLETION BY PATIENT OR THEIR REPRESENTATIVE

	Patient Name:	Date of Birth:
PATIENT AUTHORIZATION(S)		

Patient Consent to allow Acthar Patient Support Team to work together with your insurance provider, pharmacy, advocacy organization and others to provide support on your behalf.

By signing this authorization, I authorize my physician(s), my health insurance company and my pharmacy providers (collectively, "Designated Parties") to use, disclose, and redisclose to Mallinckrodt ARD LLC ("Mallinckrodt"), the distributor of Acthar, and its agents, authorized designees and contractors, including Mallinckrodt reimbursement support personnel and United BioSource LLC ("UBC") or any other operator of Acthar Patient Support on behalf of Mallinckrodt (collectively, "Manufacturer Parties"), health information relating to my medical condition, treatment and insurance coverage (my "Health Information") in order for them to (1) provide certain services to me, including reimbursement and coverage support, patient assistance and access programs, medication shipment tracking, and home injection training, (2) provide me with support services and information associated with my Acthar therapy, (3) serve internal business purposes, such as marketing research, internal financial reporting and operational purposes, and (4) carry out the Manufacturer Parties' respective legal responsibilities.

Once my Health Information has been disclosed to Manufacturer Parties, I understand that it may be redisclosed by them and no longer protected by federal and state privacy laws. However, Manufacturer Parties agree to protect my Health Information by using and disclosing it only for the purposes detailed in this authorization or as permitted or required by law.

I understand that I may refuse to sign this authorization and that my physician and pharmacy will not condition my treatment on my agreement to sign this authorization form, and my health plan or health insurance company will not condition payment for my treatment, insurance enrollment or eligibility for insurance benefits on my agreement to sign this authorization form. I understand that my pharmacies and other Designated Parties may receive payment in connection with the disclosure of my Health Information as provided in this authorization. I understand that I am entitled to receive a copy of this authorization after I sign it.

I may revoke (withdraw) this authorization at any time by mailing a letter to Acthar Patient Support, 680 Century Point, Lake Mary, FL 32746. Revoking this authorization will end further disclosure of my Health Information to Manufacturer Parties by my pharmacy, physicians, and health insurance company when they receive a copy of the revocation, but it will not apply to information they have already disclosed to Manufacturer Parties based on this authorization. I also know I may cancel my enrollment in a patient support program at any time in writing by contacting Mallinckrodt via fax at 1-877-937-2284 or by calling Acthar Patient Support at 1-888-435-2284. This authorization is in effect for 5 years unless a shorter period is provided for by state law (MARYLAND HEALTHCARE PROVIDERS, under Maryland Code HG § 4-303(b)(4) this authorization expires ONE YEAR from the date of signature) or until the conclusion of any ongoing coverage support, whichever is longer, once I have signed it unless I cancel it before then.

THIS SECTION MUST BE COMPLETED IN ITS ENTIRETY, INCLUDING DATE



X

PATIENT NAME OR LEGAL REPRESENTATIVE

PATIENT OR LEGAL REPRESENTATIVE SIGNATURE

IF LEGAL REPRESENTATIVE, RELATIONSHIP TO PATIENT

DATE

Patient Consent to receive additional information from Mallinckrodt such as education on your disease and Acthar.

I authorize Mallinckrodt and its partners to use, disclose, and/or transfer the personal information I supply (1) to contact me and provide me with informational and marketing materials and clinical trial opportunities related to my condition or treatment by any means of communication, including but not limited to text, email, mail, or telephone; (2) to help Mallinckrodt improve, develop, and evaluate products, services, materials, and programs related to my condition or treatment; (3) to enroll me in and provide me with Acthar-related programs and services that I may select or refuse at any time; (4) to disclose my enrollment and use of these services to my prescriber and insurers; and (5) to use my information that cannot identify me for scientific and market research. This authorization will remain in effect until I cancel it, which I may do at any time in writing by contacting Mallinckrodt via fax at 1-877-937-2284 or by calling Acthar Patient Support at 1-888-435-2284. I may request a copy of this signed authorization.

THIS SECTION MUST BE COMPLETED IN ITS ENTIRETY, INCLUDING DATE



<u>X</u>

PATIENT NAME OR LEGAL REPRESENTATIVE

PATIENT OR LEGAL REPRESENTATIVE SIGNATURE

IF LEGAL REPRESENTATIVE, RELATIONSHIP TO PATIENT

DATE

Scan the QR Code below to save the Acthar Patient Support phone number to your mobile device's contacts (see steps below).



SIEP1

Open the camera on your mobile device

STEP 2

Hold your camera over the QR code to scan



Save your Acthar Patient Support Team information to your contacts

If patient is not present to sign the form, send them to

Acthar Consent.com

and have them sign electronically.

ACTHAR GEL COMMERCIAL STARTER PROGRAM TERMS & CONDITIONS: Eligible patients for this Program must meet the following criteria: have a valid prescription for the FDA-approved indication of severe acute and chronic allergic and inflammatory processes involving the eye and its adnexa such as. keratitis, iritis, iridocyclitis, diffuse posterior uveitis and choroiditis, optic neuritis, optic neuritis, chorioretinitis, or anterior segment inflammation, have verified commercial or private insurance, and are not participating in Medicare, Medicare, Medicaid, or any government-funded healthcare plan. This Program is valid for one vial of Acthar Gel at at time as needed; however, the patient will no longer receive Acthar Gel under this Program when the patient receives insurance approval or a final denial of coverage. The patient agrees not to seek reimbursement from any third-party payer for all or any part of Acthar Gel dispensed pursuant to this Program. This Program is void where prohibited by law. Mallinckrodt reserves the right to rescind, revoke, or amend this Program any time without notice. By participating in this Program, the patient agrees to these terms and conditions. Other terms and conditions apply. See acthar-pay-now/csp-terms for full details.



Acthar Gel Enrollment/Prescription Form Please complete and email or fax toll-free

Phone: 888-435-2284 FAX: 877-937-2284 EMAIL: intake@supportandaccess.com

RESOURCE PAGE. DO NOT NEED TO FAX BACK.

PRESCRIBER INSTRUCTIONS

- 1. Complete pages 1 and 2 of the Acthar Enrollment/Prescription Form.
- 2. Have your patient read page 3, PATIENT AUTHORIZATION(S). Request that the patient sign both sections to allow Acthar Patient Support to provide a complete level of support both during the approval process and after starting treatment. Alternatively, direct the patient to provide this consent at Acthar Consent.com. Tell your patient to expect a call and save the Acthar Patient Support number, 1-888-435-2284.
- 3. Email or fax pages 1, 2, and 3 of the completed Enrollment/Prescription Form along with clinical notes, any medically relevant documentation, and copies of both the front and back of your patient's medical and prescription benefit card(s) to intake@supportandaccess.com or 1-877-937-2284.

Acthar Patient Support will process the Enrollment/Prescription Form and contact both you and your patient by phone, text, or email. Prior authorization assistance will only be provided for FDA-approved indications. Medicare, Medicaid, and other federal or state healthcare program patients may be ineligible for certain other aspects of Acthar assistance programs.

PRESCRIBER SIGNATURE ON PAGE 1 AUTHORIZES PRESCRIPTION, CONSENT, AND STATEMENT OF MEDICAL NECESSITY

By signing page 1, I certify that Acthar® Gel is medically necessary for this patient and that I have reviewed this therapy with the patient and will be monitoring the patient's treatment. I verify that the patient and Prescriber information on this enrollment form was completed by me or at my direction and that the information contained herein is complete and accurate to the best of my knowledge.

I understand that I must comply with my practicing state's specific prescription requirements, such as e-prescribing, state-specific prescription form, fax language, etc. Noncompliance with state-specific requirements could result in outreach to me by the dispensing pharmacy.

I authorize United BioSource LLC ("UBC"), the current operator of Acthar Patient Support, and other designated operators of the Program, to act on my behalf for the limited purposes of transmitting this prescription to and received by the designated Specialty Pharmacy by any means under applicable law, including via a designated third party or other operator of the Program.

I understand that representatives from the Program or UBC may contact me or my patient for additional information relating to this prescription. I acknowledge and agree that this prescription may be sent to and received by the designated Specialty Pharmacy by any means under applicable law, including via a designated third party or other operator of the Program, and that no additional confirmation of receipt of prescription is required by the designated Specialty Pharmacy.

I request that company-funded Acthar Injection Training Services be arranged for my patient. I understand that Acthar Injection Training Services are available for multiple visits but are NOT a home health nursing service and that I or my patient may opt out of any nursing services by notifying the Acthar Patient Support Team by calling 1-888-435-2284. Patients can contact their Nurse Navigator at any time about injection training.

PATIENT INSTRUCTIONS

Your Prescriber will submit the completed Acthar Enrollment/Prescription Form to Acthar Patient Support. After we receive the form, we will call you so we can help you get your medicine. Please be on the lookout and answer calls from 1-800, 1-888, or blocked numbers. If you have any questions, please call **1-888-435-2284** Monday through Friday from 8 AM to 9 PM ET or Saturday from 9 AM to 2 PM ET.



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IMPORTANT SAFETY INFORMATION

Contraindications

Acthar is contraindicated:

- For intravenous administration
- In infants under 2 years of age who have suspected congenital infections
- With concomitant administration of live or live attenuated vaccines in patients receiving immunosuppressive doses of Acthar
- In patients with scleroderma, osteoporosis, systemic fungal infections, ocular herpes simplex, recent surgery, history of or the presence of a peptic ulcer, congestive heart failure, uncontrolled hypertension, primary adrenocortical insufficiency, adrenocortical hyperfunction, or sensitivity to proteins of porcine origin

Warnings and Precautions

- The adverse effects of Acthar are related primarily to its steroidogenic effects
- Acthar may increase susceptibility to new infection or reactivation of latent infections
- Suppression of the hypothalamic-pituitary-adrenal (HPA) axis may occur following prolonged therapy with the potential for adrenal insufficiency after withdrawal of the medication. Adrenal insufficiency may be minimized by tapering of the dose when discontinuing treatment. During recovery of the adrenal gland patients should be protected from the stress (e.g., trauma or surgery) by the use of corticosteroids. Monitor patients for effects of HPA axis suppression after stopping treatment
- Cushing's syndrome may occur during therapy but generally resolves after therapy is stopped. Monitor patients for signs and symptoms
- Acthar can cause elevation of blood pressure, salt and water retention, and hypokalemia. Monitor blood pressure and sodium and potassium levels
- Acthar often acts by masking symptoms of other diseases/disorders. Monitor patients carefully during and for a period following discontinuation of therapy
- Acthar can cause gastrointestinal (GI) bleeding and gastric ulcer. There is also an increased risk for perforation in patients with certain GI disorders. Monitor for signs of perforation and bleeding
- Acthar may be associated with central nervous system effects ranging from euphoria, insomnia, irritability, mood swings, personality changes, and severe depression to psychosis. Existing conditions may be aggravated
- Patients with comorbid disease may have that disease worsened. Caution should be used when prescribing Acthar in patients
 with diabetes and myasthenia gravis
- Prolonged use of Acthar may produce cataracts, glaucoma, and secondary ocular infections. Monitor for signs and symptoms
- Acthar is immunogenic and prolonged administration of Acthar may increase the risk of hypersensitivity reactions. Cases of anaphylaxis have been reported in the postmarketing setting. Neutralizing antibodies with chronic administration may lead to loss of endogenous ACTH and Acthar activity
- There may be an enhanced effect in patients with hypothyroidism and in those with cirrhosis of the liver
- Long-term use may have negative effects on growth and physical development in children. Monitor pediatric patients
- Decrease in bone density may occur. Bone density should be monitored in patients on long-term therapy

Adverse Reactions

- Commonly reported postmarketing adverse reactions for Acthar include injection site reaction, asthenic conditions (including fatigue, malaise, asthenia, and lethargy), fluid retention (including peripheral swelling), insomnia, headache, and blood glucose increased
- The most common adverse reactions for the treatment of infantile spasms (IS) are increased risk of infections, convulsions, hypertension, irritability, and pyrexia. Some patients with IS progress to other forms of seizures; IS sometimes masks these seizures, which may become visible once the clinical spasms from IS resolve

Pregnancy

Acthar may cause fetal harm when administered to a pregnant woman

Please see accompanying full Prescribing Information for additional Important Safety Information or visit https://www.actharhcp.com/Static/pdf/Acthar-Pl.pdf

INDICATION AND USAGE

 Acthar Gel is indicated for severe acute and chronic allergic and inflammatory processes involving the eye and its adnexa such as: keratitis, iritis, iridocyclitis, diffuse posterior uveitis and choroiditis, optic neuritis, chorioretinitis, anterior segment inflammation.



APPENDIX A

RESOURCE PAGE. DO NOT NEED TO FAX BACK.

OPHTHALMOLOGY

- KERATOCONJUNCTIVITIS DUE TO ACANTHAMOEBA B60.13
- SARCOID IRIDOCYCLITIS
 D86.83
- NEUROMYELITIS OPTICA [DEVIC] G36.0
- DISCOID LUPUS ERYTHEMATOSUS OF RIGHT UPPER EYELID
 H01.121
- DISCOID LUPUS ERYTHEMATOSUS OF RIGHT LOWER EYELID H01.122
- DISCOID LUPUS ERYTHEMATOSUS OF RIGHT EYE, UNSPECIFIED EYELID H01.123
- DISCOID LUPUS ERYTHEMATOSUS OF LEFT UPPER EYELID H01.124
- DISCOID LUPUS ERYTHEMATOSUS OF LEFT LOWER EYELID H01.125
- DISCOID LUPUS ERYTHEMATOSUS OF LEFT EYE, UNSPECIFIED EYELID H01.126
- DISCOID LUPUS ERYTHEMATOSUS OF UNSPECIFIED EYE, UNSPECIFIED EYELID H01.129
- OTHER SPECIFIED INFLAMMATIONS OF EYELID H01.8
- UNSPECIFIED INFLAMMATION OF EYELID H01.9
- CHRONIC DACRYOADENITIS, RIGHT LACRIMAL GLAND H04.021
- CHRONIC DACRYOADENITIS, LEFT LACRIMAL GLAND H04.022
- CHRONIC DACRYOADENITIS, BILATERAL LACRIMAL GLAND
- H04.023
 CHRONIC DACRYOADENITIS, UNSPECIFIED LACRIMAL
- UNSPECIFIED LACRIMAL GLAND H04.029
- CHRONIC DACRYOCYSTITIS OF RIGHT LACRIMAL PASSAGE H04.411
- CHRONIC DACRYOCYSTITIS
 OF LEFT LACRIMAL PASSAGE
 H04 412
- CHRONIC DACRYOCYSTITIS OF BILATERAL LACRIMAL PASSAGES H04.413
- CHRONIC DACRYOCYSTITIS
 OF UNSPECIFIED LACRIMAL
 PASSAGE
- +04.419

 UNSPECIFIED ACUTE INFLAMMATION OF ORBIT +05.00
- TENONITIS OF RIGHT ORBIT H05.041
- TENONITIS OF LEFT ORBIT
 H05.042
- TENONITIS OF BILATERAL ORBITS H05.043
- TENONITIS OF UNSPECIFIED ORBIT H05.049
- UNSPECIFIED CHRONIC INFLAMMATORY DISORDERS OF ORBIT H05.10
- H05.10
 GRANULOMA OF RIGHT ORBIT H05.111
- GRANULOMA OF LEFT ORBIT H05.112
- GRANULOMA OF BILATERAL ORBITS H05.113

- GRANULOMA OF UNSPECIFIED ORBIT H05.119
- ORBITAL MYOSITIS, RIGHT ORBIT H05.121
- ORBITAL MYOSITIS, LEFT ORBIT H05.122
- ORBITAL MYOSITIS, BILATERAL H05.123
- ORBITAL MYOSITIS, UNSPECIFIED ORBIT H05.129
- ACUTE ATOPIC CONJUNCTIVITIS, UNSPECIFIED EYE H10.10
- ACUTE ATOPIC CONJUNCTIVITIS, RIGHT EYE H10.11
- ACUTE ATOPIC CONJUNCTIVITIS, LEFT EYE H10.12
- ACUTE ATOPIC CONJUNCTIVITIS, BILATERAL H10.13
- UNSPECIFIED CHRONIC CONJUNCTIVITIS, RIGHT EYE H10 401
- UNSPECIFIED CHRONIC CONJUNCTIVITIS, LEFT EYE H10.402
- UNSPECIFIED CHRONIC CONJUNCTIVITIS, BILATERAL H10.403
- UNSPECIFIED CHRONIC CONJUNCTIVITIS, UNSPECIFIED EYE H10.409
- CHRONIC GIANT PAPILLARY CONJUNCTIVITIS, RIGHT EYE H10.411
- CHRONIC GIANT PAPILLARY CONJUNCTIVITIS, LEFT EYE H10.412
- CHRONIC GIANT PAPILLARY CONJUNCTIVITIS, BILATERAL H10.413
- CHRONIC GIANT PAPILLARY CONJUNCTIVITIS, UNSPECIFIED EYE H10.419
- SIMPLE CHRONIC CONJUNCTIVITIS, RIGHT EYE H10.421
- SIMPLE CHRONIC CONJUNCTIVITIS, LEFT EYE H10.422
- SIMPLE CHRONIC CONJUNCTIVITIS, BILATERAL H10.423
- SIMPLE CHRONIC CONJUNCTIVITIS, UNSPECIFIED EYE H10.429
- CHRONIC FOLLICULAR CONJUNCTIVITIS, RIGHT EYE H10.431
- CHRONIC FOLLICULAR CONJUNCTIVITIS, LEFT EYE H10.432
- CHRONIC FOLLICULAR CONJUNCTIVITIS, BILATERAL H10.433
- CHRONIC FOLLICULAR CONJUNCTIVITIS, UNSPECIFIED EYE H10.439
- VERNAL CONJUNCTIVITIS H10.44
- OTHER CHRONIC ALLERGIC CONJUNCTIVITIS H10.45
- LIGNEOUS CONJUNCTIVITIS, RIGHT EYE
 H10.511
- LIGNEOUS CONJUNCTIVITIS, LEFT EYE H10.512
- LIGNEOUS CONJUNCTIVITIS, BILATERAL H10.513
- LIGNEOUS CONJUNCTIVITIS, UNSPECIFIED EYE H10.519

- UNSPECIFIED SCLERITIS, RIGHT EYE H15.001
- UNSPECIFIED SCLERITIS, LEFT EYE
 H15.002
- UNSPECIFIED SCLERITIS, BILATERAL H15.003
- UNSPECIFIED SCLERITIS, UNSPECIFIED EYE H15.009
- ANTERIOR SCLERITIS, RIGHT EYE H15.011
- ANTERIOR SCLERITIS, LEFT EYE
- H15.012

 ANTERIOR SCLERITIS, BILATERAL H15.013
- ANTERIOR SCLERITIS, UNSPECIFIED EYE H15.019
- POSTERIOR SCLERITIS, RIGHT EYE
 H15.031
- POSTERIOR SCLERITIS, LEFT EYE
 H15.032
- POSTERIOR SCLERITIS, BILATERAL H15.033
- POSTERIOR SCLERITIS, UNSPECIFIED EYE
 H15 039
- SCLERITIS WITH CORNEAL INVOLVEMENT, RIGHT EYE H15.041
- SCLERITIS WITH CORNEAL INVOLVEMENT, LEFT EYE H15.042
- SCLERITIS WITH CORNEAL INVOLVEMENT, BILATERAL H15.043
- SCLERITIS WITH CORNEAL INVOLVEMENT, UNSPECIFIED EYE H15.049
- OTHER SCLERITIS, RIGHT EYE
 H15.091
- OTHER SCLERITIS, LEFT EYE
 H15.092
- OTHER SCLERITIS, BILATERAL H15.093
- OTHER SCLERITIS, UNSPECIFIED EYE H15.099
- UNSPECIFIED CORNEAL ULCER, RIGHT EYE H16.001
- H16.001UNSPECIFIED CORNEAL ULCER, LEFT EYE
- H16.002

 UNSPECIFIED CORNEAL
- ULCER, BILATERAL
 H16.003

 UNSPECIFIED CORNEAL
- ULCER, UNSPECIFIED EYE
 H16.009

 CENTRAL CORNEAL ULCER,
- RIGHT EYE H16.011
- CENTRAL CORNEAL ULCER, LEFT EYE H16.012
- CENTRAL CORNEAL ULCER, BILATERAL H16.013
- CENTRAL CORNEAL ULCER, UNSPECIFIED EYE H16.019
- RING CORNEAL ULCER, RIGHT EYE H16.021
- RING CORNEAL ULCER, LEFT EYE H16.022
- RING CORNEAL ULCER, BILATERAL H16.023
- RING CORNEAL ULCER, UNSPECIFIED EYE H16.029

- CORNEAL ULCER WITH HYPOPYON, RIGHT EYE **H16.031**
- CORNEAL ULCER WITH HYPOPYON, LEFT EYE H16.032
- CORNEAL ULCER WITH HYPOPYON, BILATERAL **H16.033**
- CORNEAL ULCER WITH HYPOPYON, UNSPECIFIED EYE
- H16.039

 MARGINAL CORNEAL ULCER, RIGHT EYE
- H16.041

 MARGINAL CORNEAL ULCER, LEFT EYE
 H16.042
- MARGINAL CORNEAL ULCER, BILATERAL H16.043
- MARGINAL CORNEAL ULCER, UNSPECIFIED EYE H16.049
- MOOREN'S CORNEAL ULCER, RIGHT EYE
 H16.051
- MOOREN'S CORNEAL ULCER, LEFT EYE H16.052
- MOOREN'S CORNEAL ULCER, BILATERAL H16.053
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