

FAX: 1.800.420.5150 MAIL: 100 College Street New Haven, CT 06510







OneSource™ is a complimentary, personalized patient support program offered by Alexion. It's designed to support patients' specific needs throughout treatment. For more information, visit www.AlexionOneSource.com. Contact OneSource if you have any questions while completing the forms.



INSTRUCTIONS FOR HEALTHCARE PROFESSIONALS:

To enroll your patient in OneSource, please follow these steps:

- Have your patient complete all required sections and read the Authorization to Share Health Information on the Patient Services Enrollment Form
- Complete all required sections on PAGE 1
- 3 Sign the Prescriber Certification on PAGE 2
- FAX PAGES 1-2 of the completed form and copies of the front and back of the patient's medical insurance and pharmacy coverage cards to OneSource. If applicable, fax the Vaccination Order Form (PAGE 3) to OneSource as well.

Fields in red with asterisks are required.*

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STEP 1: PATIENT INFORMATION					
PATIENT NAME (FIRST, LAST)*	DATE OF BIRTH (MM/DD/YYYY)*	MM/DD/YYYY)* PATIENT PHONE NUMBER*		PATIENT EMAIL	
STEP 2: CLINICAL DIAGNOSIS					
SOLIRIS and ULTOMIRIS are FDA approved for antibody-po	sitive status. If a payer requires	prior authorization and/or has a clinic	cal policy, they m	ay require proof of antibody statu	
INDICATION (check one)*:	avis with (acute) exacerbation	ANTIBODY STATUS (check one)*:	ANTI-AQP4 A	NTIBODY POSITIVE (gMG) NTIBODY POSITIVE (NMOSD) ONTACT ONESOURCE FOR QUESTION:	
STEP 3: INSURANCE INFORMATION					
Complete this section OR attach copies of patient's medical ar	nd pharmacy insurance card(s).*				
☐ PLEA	SE PROVIDE SUMMARY OF BENEFIT	INVESTIGATION FOR ULTOMIRIS AND SO	LIRIS		
☐ COPIES OF PATIENT'S INSURANCE CARD(S) ATTACHED ☐ PATIENT DOES NOT HAVE INSURANCE	PRIMARY MEDICAL INSURANCE	PRIMARY MEDICAL SECONDARY MEDI INSURANCE INSURANCE		PHARMACY COVERAGE	
INSURANCE PROVIDER*					
INSURANCE PHONE #*					
CARDHOLDER NAME*					
CARDHOLDER DATE OF BIRTH*					
MEMBER ID*					
POLICY #*					
GROUP #*					
BIN/PCN #					
STEP 4: HEALTHCARE PRESCRIBER INFORM	IATION				
FIRST NAME*	LAST NAME*		PROVIDER EMA	IL*	
ADDRESS*			PHONE NUMBE	HONE NUMBER*	
CITY*	STATE*		ZIP*	ZIP*	
PRACTICE NAME	TAX ID #*		NPI#*		
OFFICE CONTACT NAME	EMAIL		FAX NUMBER*	NUMBER*	
STEP 5: SITE OF CARE					
SELECT OPTION A OR B BELOW*:					
A) PLEASE PROVIDE ASSISTANCE LOCATING AN INFUS	ON SITE. PLEASE COO	ORDINATE DIRECTLY WITH: HEALTH	CARE PROVIDER	☐ PATIENT	
B) ASSISTANCE IS NOT NEEDED. PATIENT WILL BE INFUS	ED AT: PRESCRIBER'S OFFICE	☐ PATIENT'S HOME ☐ PREFER	RRED INFUSION SIT	TE (PLEASE SPECIFY BELOW)	
SITE OF CARE NAME	NPI#	NPI#		TAX ID #	
ADDRESS	,	<u>'</u>			
CITY	STATE		ZIP		
OFFICE CONTACT FOR FOLLOW-UP		PHONE NUMBER			

Please see Indications & Important Safety Information on page 4 and full Prescribing Information and Medication Guide for ULTOMIRIS, including Boxed WARNING regarding serious and life-threatening or fatal meningococcal infections, also available on www.ULTOMIRIS.com.

Please see Indications & Important Safety Information on page 5 and full Prescribing Information and Medication Guide for SOLIRIS, including Boxed WARNING regarding serious and life-threatening or fatal meningococcal infections, also available on www.SOLIRIS.net.

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PHONE: 1.888.765.4747 8:30 AM to 8 PM ET Monday-Friday





Fields in red with asterisks are required.*

PATIE	NT INFORM	ATION								
PATIENT	NAME (FIRST, LA	AST)*				DATE OF BIRTH (MM/DD	/YYYY)*			
STEP	6: CLINICAL	INFORMATION								
☐ AZ	ATHIOPRINE GARTIGIMOD	ENERALIZED MYASTH MYCOPHENOLA PLASMAPHERE PREDNISONE	TE MOFETIL	PYRIDOSTIGMINE RITUXIMAB		ECK ALL PREVIOUS NEUR AZATHIOPRINE CYCLOPHOSPHAMIDE INEBILIZUMAB		E E	PUM DISORDER (NMOSD) THERAPIES: RITUXIMAB	
MGFA CLASSIFICATION:						MBER OF RELAPSES IN LA	ST 12 MONTHS:		24 MONTHS:	
CURREN	IT MG-ADL SCOR	E:			EDS	S SCORE:				
Abbreviat	ions: AChR, acetylch	noline receptor; EDSS, Exp	anded Disabili	ا ty Status Scale; IVIg, intravenous imi	munog	lobulin; MG-ADL, Myasthenia	Gravis Activities of Daily	Living; N	IGFA, Myasthenia Gravis Foundation of Ameri	
STEP	7: PRESCRIF	PTION								
YOU MA	Y USE THIS SEC	TION TO PROVIDE A P	RESCRIPTIO	ON FOR ULTOMIRIS OR SOLIRIS	6, OR	YOU MAY PROVIDE A SEI	PARATE PRESCRIPTI	ON.		
	Rx ULT	OMIRIS 100 mg/mL PATIENT WEIGHT: .				☐ Rx 5	SOLIRIS 10 mg/mL l	HCPCS	CODE: J1300 PER UNIT	
LOADIN	IG DOSE:		MAINTEN	ANCE DOSE:		LOADING DOSE:		MA	INTENANCE DOSE:	
SIG: INFUSE INTRAVENOUSLY mg ON DAY 0. COVERS THE PATIENT FOR THE FIRST 2 WEEKS.		EVERY 8 V	FUSE INTRAVENOUSLY mg 8 WEEKS. START 2 WEEKS AFTER ETION OF LOADING DOSE.		SIG: INFUSE INTRAVENOUSLYmg WEEKLY FOR THE FIRST 4 WEEKS, FOLLOWED BYmg FOR THE 5TH WEEK. OTHER:		D EVI	SIG: INFUSE INTRAVENOUSLYm		
	300 mg/3 mL	QTY 0F 300 mg/3 mL QTY 0F 300 mg/30 mL			QTY 0F 300 mg/30 mL VIALS: REFILLS:					
QTY 0F 1100 mg/11 mL										
		Vaccination d	ates provi	re available only through a ded as part of this form a d?	re us	ed to confirm vaccina	ation prior to star	n and I ting tr	Mitigation Strategy (REMS). eatment.	
		has received or is	schedule	ed to receive the required	lvac	cinations per ACIP g			Patient needs VACCINATION SUPPORT fro	
		•		MenB	MenABCV		всшу		OneSource	
YES		e:// Menactra M	enQuadfi	1st Dose Date:/ Bexsero Trumen	•	1st Dose Date:		NO	✓ Sign prescriber certification below	
		e:// Menactra M		2nd Dose Date:/ ☐ Bexsero ☐ Trumen		2nd Dose Date:	//		✓ Continue to PAGE 3 to fill out a vaccination	
	✓ Sign prescriber certification below 3rd Dose Date:/_ (3rd dose - Trumenba ON		3rd Dose Date:/ (3rd dose - Trumenba ONLY	<u></u>			prescription [†]			
		idelines recomme e a separate pres		nen of MenACWY AND M	enB (doses prior to starti	ng a complement	t inhib	itor treatment.	
		BER CERTIFICAT								
By signin diagnosis and comp iv) I am u v) the in collected	ng below, I attes s identified on t plied with all app under no obligat formation provi d about me (as t	t that: (i) I am prescr his form and I will be olicable prescription ion to prescribe ULT ded on this form is c he prescriber) in acc	ibing ULTOI supervisin requiremei OMIRIS or S omplete, ci ordance wi	MIRIS or SOLIRIS for the pating the patient's treatment; (ii thst; (iii) I am authorizing Alex SOLIRIS and I have not receivurrent, and accurate to the bith the Privacy Notice on the	ent id) I am ion to ed, no est o Alexi	lentified above based on authorized under applion forward the patient's or will I receive, any ben of my knowledge. I also it on website at https://e	n my clinical judgm icable law to prescr prescription to a pl efit from Alexion fo acknowledge that <i>l</i> llexion.com/Legal#	ent the ibe UL narmac or pres Alexion privac	at it is medically necessary for the TOMIRIS or SOLIRIS and I have verif by by any means under applicable la cribing ULTOMIRIS or SOLIRIS; and will use and share the personal dat //.	
ONE*	/ <u>}</u>			PS) - dispense as Writte n			DATE (MM/DD/YYY			
	PRESC	RIBER'S SIGNATURE	(NO STAMI	PS) - MAY SUBSTITUTE			DATE (MM/DD/YYY	Y)		
	Please	verify your local pre	scribina red	quirements (eg, New York pre	escrib	pers must provide a sep	arate prescription)			

Please see Indications & Important Safety Information on page 4 and full Prescribing Information and Medication Guide for ULTOMIRIS, including Boxed WARNING regarding serious and life-threatening or fatal meningococcal infections, also available on www.ULTOMIRIS.com.

Please see Indications & Important Safety Information on page 5 and full Prescribing Information and Medication Guide for SOLIRIS, including Boxed WARNING regarding serious and life-threatening or fatal meningococcal infections, also available on www.SOLIRIS.net.

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VACCINATION ORDER FORM











PATIENT NAME (PRST, MIDDLE NITAL, LAST) ADDRESS CITY STATE ZP PHONE NUMBER HEALTHCARE PRESCRIBER INFORMATION FIRST NAME LAST NAME LAST NAME CITY STATE ZP PHONE NUMBER FAX NUL ADDRESS CITY STATE ZP PHONE NUMBER FAX NUL ADDRESS CITY STATE ZP FIND NUMBER FAX NUL ADDRESS CITY STATE ZP CITY STATE API CITY STATE	ENT INFORM	IATION					
PHONE NUMBER HEALTHCARE PRESCRIBER INFORMATION FIRST NAME LAST NAME PHONE NUMBER FAX NUL ADDRESS CITY STATE TYP OFFICE CONTACT NAME CLINICAL INFORMATION Primary Diagnosis Description: Encounter for Immunization ICD-10 CODE: 223 MENNIGOCOCAL VACCINATIONS ARE INDICATED FOR PATIENTS. INCLIDING PEOPLE OVER 25 VEARS OF AGE, WHEN ON A COMPLEMENT INHIBITION The Advisory Committee on Immunization Practices (ADP) recommends a regimen of ManACWY AND Man B does pit to starting a complement inhibitor. There are two (2) types of meningococcal vaccines available in the should be initiated at least 2 weeks prior to first dose of Alexion Complement Inhibitor. There are two (2) types of meningococcal vaccines available in the Inhibitor of	NT NAME (FIRST	, MIDDLE INITIAL, LAST)			PATIENT DATE OF E	BIRTH (MM/DD/YY	YY)
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CLINICAL INFORMATION Primary Diagnosis Description: Encounter for Immunization ICD-10 CODE: Z23 MENINGOCOCCAL VACCINATIONS ARE INDICATED FOR PATIENTS, INCLUDING PEOPLE OVER 25 YEARS OF AGE, WHEN ON A COMPLEMENT INHIBIT The Advisory Committee on Immunization Practices (ACIP) recommends a regimen of MenACWY AND MenB doses prior to starting a complement inhibitor the should be initiated at least 2 weeks prior to first dose of Alexion Complement Inhibitor. There are two (2) types of meningococcal vaccines available in the MenACWY MenB VACCINES ARE NOT INTERCHANGEABLE, PATIENT MUST RECEIVE THE SAME PRODUCT FOR ALL DOSES DURING A VACCINATION SCHEDULE FOR CHILDREN 130 YEARS OLD, PLEASE REFER TO THE ACIP VACCINE RECOMMENDATIONS OR TO ALEXION MEDICAL Two quadrivalent meningococcal conjugate (MenACWY) vaccines are currently licensed and available in the United States. INDICATE VACCINE THE PATIENT NEEDS TO RECEIVE: MenQuadfi (meningococcal groups A. C. W. and Y polysaccharide tetanus toxiol donjugate vaccine [MenACWY-TI]) 90612 Menvee (meningococcal groups A. C. W. and Y polysaccharide diphtheria CRM conjugate vaccine [MenACWY-CRMI]) 907340 Dose 1: Day 0 Dose 2: At least 8 weeks after Day 0 MenACWY MenACW	ESS		CITY			STATE	ZIP
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MenB VACCINES ARE NOT INTERCHANGEABLE. PATIENT MUST RECEIVE THE SAME PRODUCT FOR ALL DOSES DURING A VACCINATION SCRIES. FOR THE FULL VACCING THE PATIENT NEEDS TO YEARS OLD, PLEASE REFER TO THE ACIP VACCINE RECOMMENDATIONS OR TO ALEXION MEDICAL Two serogroup B meningococcal (MenB) vaccines are currently licinesed and available in the United States. INDICATE VACCINE THE PATIENT NEEDS TO RECEIVE: MenQuadfi (meningococcal groups A, C, W, and Y polysaccharide tetanus toxoid conjugate vaccine (MenACWY-CRM)) 90619 MenACWY MENORITY MenB Dose 1: Day 0 Dose 1: Day 0 Dose 2: At least 8 weeks after Day 0 Dose 2: At least 8 weeks after Day 0 Dose 2: At least 8 weeks after Day 0 Dose 3: (Turnenba and poly); 6 months after Day 0 Dose 3: (Turnenba and p	,	, ,	•				
NCLIDING THE VACCINATION SCHEDULE FOR CHILIDREN'S 10 YEARS OLD, PLEASE REFER TO THE ACIP VACCINE RECOMMENDATIONS OR TO ALEXION MEDICAL Two quadrivalent meningococcal conjugate (MenACWY) vaccines are currently licensed and available in the United States. INDICATE VACCINE THE PATIENT NEEDS TO RECEIVE: MenQuadfi (meningococcal groups A, C, W, and Y polysaccharide tetanus toxoid conjugate vaccine (MenACWY-TI) 906.19 Menveo (meningococcal groups A, C, W, and Y oligosaccharide diphtheria CRM conjugate vaccine (MenACWY-CRM]) 907340 MenACWY		MenACWY	ONE (1) REQUIRED	FROM EACH GRO	IUP	MenB	
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□ Dose 2: At least 8 weeks after Day 0 □ Booster dose □ Dose 3: For Bexsero: At least (or greater than or equal to) 1 mm For Trumenbas: 1-2 months after Day 0 □ Dose 3 (Trumenbas: 1-2 months after Day 0 □ Booster dose Per CDC recommendations, those who remain at increased risk need regular booster doses. MenACWY: For children under the age of 7 years, administer a 3 years after completion of the primary series and every 5 years thereafter. For children 7 years old or older and adults, administer a booster dose 5 years of the primary series and every 5 years thereafter. MenB: Administer a booster dose of vaccine 1 year after series completion and then every 2 to 3 years NOTE: ALL VACCINES LISTED ABOVE ARE ADMINISTERED INTRAMUSCULARLY AT A DOSE OF 0.5 mL ANCILLARY ORDERS (HOME ADMINISTRATION ONLY - USE AS NEEDED) Anaphylaxis Kit - The following items will be dispensed: □ Diphenhydramine 50 mg/mL 1 mL vial x 1. Inject 25 mg IM PRN for allergic reaction. May repeat x 1 dose in 15 min PRN if no improvement □ NS 500 mL bag x 1. Infuse 500 mL IV at KVO rate PRN anaphylaxis □ Epinephrine ampule/vial 1 mg/mL (1:1000) 1 mL x 2 ampules/vials. Inject 0.3 mg SQ PRN for adverse reaction. May repeat x 1 dose in 5 to 15 min PRN General Anaphylaxis Instructions Administer emergency medications as ordered. Administer epinephrine as above and repeat dose if necessary. Administer injectable diphenhydramine as dose if necessary. Place peripheral IV and administer NS. Initiate CPR if needed. Call EMS (activate the emergency medical system). Monitor vital signs—thypotensive. Notify prescriber and Nursing Director or pharmacist. CRIBER CERTIFICATION ning below, I attest that: (i) based on my clinical judgment, the vaccines identified are medically necessary for the patient and diagnosis identified on this applicable law to prescribe the vaccines identified and I have verified and complied with all applicable prescription requirements; (iii) I am authorizing Alexi iption to a pharmacy by any means under applicable law; (iv) I a		MenACWY	DOSINGS	CHEDULE		MenB	
NOTE: ALL VACCINES LISTED ABOVE ARE ADMINISTERED INTRAMUSCULARLY AT A DOSE OF 0.5 mL ANCILLARY ORDERS (HOME ADMINISTRATION ONLY – USE AS NEEDED) Anaphylaxis Kit - The following items will be dispensed: Diphenhydramine 50 mg/mL 1 mL vial x 1. Inject 25 mg IM PRN for allergic reaction. May repeat x 1 dose in 15 min PRN if no improvement NS 500 mL bag x 1. Infuse 500 mL IV at KVO rate PRN anaphylaxis Epinephrine ampule/vial 1 mg/mL (1:1000) 1 mL x 2 ampules/vials. Inject 0.3 mg SQ PRN for adverse reaction. May repeat x 1 dose in 5 to 15 min PRN General Anaphylaxis Instructions Administer emergency medications as ordered. Administer epinephrine as above and repeat dose if necessary. Administer injectable diphenhydramine as dose if necessary. Place peripheral IV and administer NS. Initiate CPR if needed. Call EMS (activate the emergency medical system). Monitor vital signs—thypotensive. Notify prescriber and Nursing Director or pharmacist. CRIBER CERTIFICATION ning below, I attest that: (i) based on my clinical judgment, the vaccines identified are medically necessary for the patient and diagnosis identified on this applicable law to prescribe the vaccines identified and I have verified and complied with all applicable prescription requirements; (iii) I am authorizing Alexington to a pharmacy by any means under applicable law; (iv) I am under no obligation to prescribe the vaccines identified and I have not received, nor will I man authorizing Alexington to the information provided on this form is complete, current, and accurate to the best of my knowledge.	ose 2: At least 8 poster dose DC recommendars after complet	ntions, those who remain at increased risk ion of the primary series and every 5 year	s thereafter. For childre	Dose 2: For For Dose 3 (Tru Booster dos Doses. MenACWY: To years old or ol	Bexsero: At least (or ; Trumenba: 1-2 month menba only): 6 month se For children under the der and adults, admin	ns after Day 0 ns after Day 0 e age of 7 years, a nister a booster do	ndminister a booster dose ose 5 years after completion
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PRESCRIBER SIGNATURE (NO STAMPS) - DISPENSE AS WRITTEN DATE (MM/DD/YYYY)	/% -	PRESCRIBER SIGNATURE (NO STAMPS) - DISF	PENSE AS WRITTEN			DATE (MM/DD/YY	

Please verify your local prescribing requirements (eg, New York prescribers must provide a separate prescription)

PRESCRIBER SIGNATURE (NO STAMPS) - MAY SUBSTITUTE

DATE (MM/DD/YYYY)





FAX: 1.800.420.5150 MAIL: 100 College Street New Haven, CT 06510



PHONE: 1.888.765.4747 8:30 AM to 8 PM ET Monday-Friday





ULTOMIRIS

INDICATIONS & IMPORTANT SAFETY INFORMATION FOR ULTOMIRIS

INDICATIONS

Generalized Myasthenia Gravis (gMG)

ULTOMIRIS is indicated for the treatment of adult patients with generalized myasthenia gravis (gMG) who are anti-acetylcholine receptor (AChR) antibody-positive.

Neuromyelitis Optica Spectrum Disorder

ULTOMIRIS is indicated for the treatment of adult patients with neuromyelitis optica spectrum disorder (NMOSD) who are antiaquaporin 4 (AQP4) antibody-positive.

IMPORTANT SAFETY INFORMATION

WARNING: SERIOUS MENINGOCOCCAL INFECTIONS ULTOMIRIS, a complement inhibitor, increases the risk of serious infections caused by Neisseria meningitidis [see Warnings and Precautions (5.1)] Life-threatening and fatal meningococcal infections have occurred in patients treated with complement inhibitors. These infections may become rapidly life-threatening or fatal if not recognized and treated early.

- Complete or update vaccination for meningococcal bacteria (for serogroups A, C, W, Y, and B) at least 2 weeks prior to the first dose of ULTOMIRIS, unless the risks of delaying ULTOMIRIS therapy outweigh the risk of developing a serious infection. Comply with the most current Advisory Committee on Immunization Practices (ACIP) recommendations for vaccinations against meningococcal bacteria in patients receiving a complement inhibitor. See Warnings and Precautions (5.1) for additional guidance on the management of the risk of serious infections caused by meningococcal bacteria.
- Patients receiving ULTOMIRIS are at increased risk for invasive disease caused by Neisseria meningitidis, even if they develop antibodies following vaccination. Monitor patients for early signs and symptoms of serious meningococcal infections and evaluate immediately if infection is suspected.

Because of the risk of serious meningococcal infections, ULTOMIRIS is available only through a restricted program under a Risk Evaluation and Mitigation Strategy (REMS) called ULTOMIRIS and SOLIRIS REMS [see Warnings and Precautions (5.2)].

CONTRAINDICATIONS

 Initiation in patients with unresolved serious Neisseria meningitidis infection.

WARNINGS AND PRECAUTIONS

Serious Meningococcal Infections

Revaccinate patients in accordance with ACIP recommendations considering the duration of ULTOMIRIS therapy. Note that ACIP recommends an administration schedule in patients receiving complement inhibitors that differs from the administration schedule in the vaccine prescribing information. If urgent ULTOMIRIS therapy is indicated in a patient who is not up to date with meningococcal vaccines according to ACIP recommendations, provide antibacterial drug prophylaxis and administer meningococcal vaccines as soon as possible. Various durations and regimens of antibacterial drug prophylaxis have been considered, but the optimal durations and drug regimens for prophylaxis and their efficacy have not been studied in unvaccinated or vaccinated patients receiving complement inhibitors, including ULTOMIRIS. The benefits and risks of treatment with ULTOMIRIS, as well as those associated with antibacterial drug prophylaxis in unvaccinated or vaccinated patients, must be considered against the known risks for serious infections caused by Neisseria meningitidis.

Vaccination does not eliminate the risk of serious meningococcal infections, despite development of antibodies following vaccination.

Closely monitor patients for early signs and symptoms of meningococcal infection and evaluate patients immediately if infection is suspected. Inform patients of these signs and symptoms and instruct patients to seek immediate medical care if they occur. Promptly treat known infections. Meningococcal infection may become rapidly life-threatening or fatal if not recognized and treated early. Consider interruption of ULTOMIRIS in patients who are undergoing treatment for serious meningococcal infection depending on the risks of interrupting treatment in the disease being treated.

ULTOMIRIS and SOLIRIS REMS

Due to the risk of serious meningococcal infections, ULTOMIRIS is available only through a restricted program called ULTOMIRIS and SOLIRIS REMS.

Prescribers must enroll in the REMS, counsel patients about the risk of serious meningococcal infection, provide patients with the REMS educational materials, assess patient vaccination status for meningococcal vaccines (against serogroups A, C, W, Y, and B) and vaccinate if needed according to current ACIP recommendations two weeks prior to the first dose of ULTOMIRIS. Antibacterial drug prophylaxis must be prescribed if treatment must be started urgently, and the patient is not up to date with both meningococcal vaccines according to current ACIP recommendations at least two weeks prior to the first dose of ULTOMIRIS. Patients must receive counseling about the need to receive meningococcal vaccines and to take antibiotics as directed, signs and symptoms of meningococcal infection, and be instructed to carry the Patient Safety Card at all times during and for 8 months following ULTOMIRIS treatment.

Further information is available at www.UltSolREMS.com or 1-888-765-4747.

Other Infections

Serious infections with Neisseria species (other than Neisseria meningitidis), including disseminated gonococcal infections, have

ULTOMIRIS blocks terminal complement activation; therefore, patients may have increased susceptibility to infections, especially with encapsulated bacteria, such as infections caused by Neisseria meningitidis but also Streptococcus pneumoniae, Haemophilus influenzae, and to a lesser extent, Neisseria gonorrhoeae. Patients receiving ULTOMIRIS are at increased risk for infections due to these organisms, even if they develop antibodies following vaccination.

Thromboembolic Event Management

The effect of withdrawal of anticoagulant therapy during treatment with ULTOMIRIS has not been established. Treatment should not alter anticoagulant management.

Infusion-Related Reactions

Intravenous administration may result in systemic infusion-related reactions, including anaphylaxis and hypersensitivity reactions. In clinical trials, infusion-related reactions occurred in approximately 1 to 7% of patients treated with ULTOMIRIS. These events included lower back pain, drop in blood pressure, limb discomfort, drug hypersensitivity (allergic reaction), dysgeusia (bad taste), and drowsiness. These reactions did not require discontinuation of ULTOMIRIS. If signs of cardiovascular instability or respiratory compromise occur, interrupt ULTOMIRIS infusion and institute appropriate supportive measures.

ADVERSE REACTIONS

Adverse Reactions for gMG

Most common adverse reactions in adult patients with gMG (incidence ≥10%) were diarrhea and upper respiratory tract infection. Serious adverse reactions were reported in 20 (23%) of patients treated with ULTOMIRIS and in 14 (16%) patients receiving placebo. The most frequent serious adverse reactions were infections reported in at least 8 (9%) patients treated with ULTOMIRIS and in 4 (4%) patients treated with placebo. Of these infections, one fatal case of COVID-19 pneumonia was identified in a patient treated with ULTOMIRIS and one case of infection led to discontinuation of ULTOMIRIS.

Adverse Reactions for NMOSD

Most common adverse reactions in adult patients with NMOSD (incidence >10%) were COVID-19, headache, back pain, arthralgia, and urinary tract infection. Serious adverse reactions were reported in 8 (13.8%) patients with NMOSD receiving ULTOMIRIS.

DRUG INTERACTIONS

Plasma Exchange, Plasmapheresis, and Intravenous Immunoglobulins Concomitant use of ULTOMIRIS with plasma exchange (PE), plasmapheresis (PP), or intravenous immunoglobulin (IVIg) treatment can reduce serum ravulizumab concentrations and requires a supplemental dose of ULTOMIRIS.

Neonatal Fc Receptor Blockers

Concomitant use of ULTOMIRIS with neonatal Fc receptor (FcRn) blockers (e.g., efgartigimod) may lower systemic exposures and reduce effectiveness of ULTOMIRIS. Closely monitor for reduced effectiveness of ULTOMIRIS.

To report SUSPECTED ADVERSE REACTIONS, contact Alexion Pharmaceuticals, Inc. at 1-844-259-6783 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

Please accompanying full Prescribing Information for ULTOMIRIS, including Boxed WARNING regarding serious and life-threatening or fatal meningococcal infections.

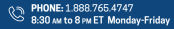
> **KLEXION** AstraZeneca Rare Disease



FAX: 1.800.420.5150



MAIL: 100 College Street New Haven, CT 06510







SOLIRIS

INDICATIONS & IMPORTANT SAFETY INFORMATION FOR SOLIRIS® (eculizumab)

INDICATIONS

Generalized Myasthenia Gravis (gMG)

SOLIRIS is indicated for the treatment of generalized myasthenia gravis (gMG) in adult patients who are anti-acetylcholine receptor (AChR) antibody positive.

Neuromyelitis Optica Spectrum Disorder (NMOSD)

SOLIRIS is indicated for the treatment of neuromyelitis optica spectrum disorder (NMOSD) in adult patients who are anti-aquaporin-4 (AQP4) antibody positive.

IMPORTANT SAFETY INFORMATION

WARNING: SERIOUS MENINGOCOCCAL INFECTIONS

SOLIRIS, a complement inhibitor, increases the risk of serious infections caused by Neisseria meningitidis [see Warnings and Precautions (5.1)]. Life-threatening and fatal meningococcal infections have occurred in patients treated with complement inhibitors. These infections may become rapidly life-threatening or fatal if not recognized and treated early.

- Complete or update vaccination for meningococcal bacteria (for serogroups A, C, W, Y, and B) at least 2 weeks prior to the first dose of SOLIRIS, unless the risks of delaying SOLIRIS therapy outweigh the risk of developing a serious infection. Comply with the most current Advisory Committee on Immunization Practices (ACIP) recommendations for vaccinations against meningococcal bacteria in patients receiving a complement inhibitor. See Warnings and Precautions (5.1) for additional guidance on the management of the risk of serious infections caused by meningococcal bacteria.
- Patients receiving SOLIRIS are at increased risk for invasive disease caused by Neisseria meningitidis, even if they develop antibodies following vaccination. Monitor patients for early signs and symptoms of serious meningococcal infections and evaluate immediately if infection is suspected.

Because of the risk of serious meningococcal infections, SOLIRIS is available only through a restricted program under a Risk Evaluation and Mitigation Strategy (REMS) called ULTOMIRIS and SOLIRIS REMS [see Warnings and Precautions (5.2)].

CONTRAINDICATIONS

•SOLIRIS is contraindicated for initiation in patients with unresolved serious Neisseria meningitidis infection.

WARNINGS AND PRECAUTIONS

Serious Meningococcal Infections

Revaccinate patients in accordance with ACIP recommendations considering the duration of therapy with SOLIRIS. Note that ACIP recommends an administration schedule in patients receiving complement inhibitors that differs from the administration schedule in the vaccine prescribing information.

If urgent SOLIRIS therapy is indicated in a patient who is not up to date with meningococcal vaccines according to ACIP recommendations, provide the patient with antibacterial drug prophylaxis and administer meningococcal vaccines as soon as possible. Various durations and regimens of antibacterial drug prophylaxis have been considered, but the optimal durations and drug regimens for prophylaxis and their efficacy have not been studied in unvaccinated or vaccinated patients receiving complement inhibitors, including SOLIRIS. The benefits and risks of treatment with SOLIRIS, as well as those associated with antibacterial drug prophylaxis in unvaccinated or vaccinated patients, must be considered against the known risks for serious infections caused by Neisseria meningitidis.

Vaccination does not eliminate the risk of serious meningococcal infections, despite development of antibodies following vaccination.

Closely monitor patients for early signs and symptoms of meningococcal infection and evaluate patients immediately if infection is suspected. Inform patients of these signs and symptoms and instruct patients to seek immediate medical care if these signs and symptoms occur. Promptly treat known infections. Meningococcal infection may become rapidly life-threatening or fatal if not recognized and treated early. Consider interruption of SOLIRIS in patients who are undergoing treatment for serious meningococcal infection, depending on the risks of interrupting treatment in the disease being treated.

ULTOMIRIS and SOLIRIS REMS

Due to the risk of serious meningococcal infections, SOLIRIS is available only through a restricted program called ULTOMIRIS and

Prescribers must enroll in the REMS, counsel patients about the risk of meningococcal infection, provide patients with REMS educational materials, assess patient vaccination status for meningococcal vaccines (against serogroups A, C, W,Y, and B) and vaccinate if needed according to current ACIP recommendations two weeks prior to the first dose of SOLIRIS. Antibacterial drug prophylaxis must be prescribed if treatment must be started urgently and the patient is not up to date with both meningococcal vaccines according to current ACIP recommendations at least two weeks prior to the first dose of SOLIRIS. Patients must receive counseling about the need to receive meningococcal vaccines and to take antibiotics as directed, the signs and symptoms of meningococcal infection, and be instructed to carry the Patient Safety Card with them at all times during and for 3 months following SOLIRIS treatment.

Further information is available at www.UltSolREMS.com or 1-888-765-4747.

Other Infections

Serious infections with Neisseria species (other than Neisseria meningitidis), including disseminated gonococcal infections, have been reported.

SOLIRIS blocks terminal complement activation; therefore, patients may have increased susceptibility to infections, especially with encapsulated bacteria, such as infections with Neisseria meningitidis but also Streptococcus pneumoniae, Haemophilus influenzae, and to a lesser extent, Neisseria gonorrhoeae. Additionally, Aspergillus infections have occurred in immunocompromised and neutropenic patients. Patients receiving SOLIRIS are at increased risk for infections due to these organisms, even if they develop antibodies following vaccination.

Infusion-Related Reactions

Administration of SOLIRIS may result in infusion-related reactions, including anaphylaxis or other hypersensitivity reactions. In clinical trials, no patients experienced an infusion-related reaction which required discontinuation of SOLIRIS. Interrupt SOLIRIS infusion and institute appropriate supportive measures if signs of cardiovascular instability or respiratory compromise occur.

ADVERSE REACTIONS

Adverse Reactions for gMG

The most frequently reported adverse reaction in the gMG placebo-controlled clinical trial (≥10%) was: musculoskeletal pain.

Adverse Reactions for NMOSD

The most frequently reported adverse reactions in the NMOSD placebo-controlled trial (≥10%) were: upper respiratory infection, nasopharyngitis, diarrhea, back pain, dizziness, influenza, arthralgia, pharyngitis, and contusion.

To report SUSPECTED ADVERSE REACTIONS contact Alexion Pharmaceuticals, Inc. at 1-844-259-6783 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

Please accompanying full Prescribing Information for SOLIRIS, including Boxed WARNING regarding serious and life-threatening or fatal meningococcal infections.

This material is intended only for residents of the United States.

PATIENT SERVICES ENROLLMENT FORM



EMAIL: OneSource@Alexion.com

PHONE: 1.888.765.4747 8:30 AM to 8 PM ET Monday-Friday



FAX: 1.800.420.5150

MAIL: 100 College St., New Haven, CT 06510

OneSource is a complimentary, personalized patient support program offered by Alexion. It's designed to support patients' specific needs throughout treatment. For more information, visit www.AlexionOneSource.com.



INSTRUCTIONS FOR PATIENTS:

To enroll in OneSource, please follow these steps:



- Sign the Authorization to Share Health Information section on this page
- Email or fax this page and copies of the front and back of your medical insurance and pharmacy coverage cards to OneSource 3 (see the email address and fax number above)

Be sure to complete all required fields and sign and date the form. If information is incomplete, it could delay our ability to enroll you in OneSource. OneSource can start offering you personalized support once you submit this form fully and correctly completed.

Contact OneSource if you have any questions while completing the form. Fields in red with asterisks are required.*

PATIENT IN	IFORMATION					
PATIENT NAM	E (FIRST, MIDDLE INITIAL, LAST)*	DATE OF BIRTH (MM/DD/YYYY)*		GENDER: MALE FEMALE NON-BINARY PREFER TO SELF-DESCRIBE:		
ADDRESS*						
CITY*			STATE*	ZIP*		
PRIMARY PHO		/es □ no □ yes □ no				
PATIENT DIAGNOSIS						
PREFERRED LA	ANGUAGE SPANISH OTHER		PATIENT EMAIL NONE			
LEGAL PATIEN	IT REPRESENTATIVE* (REQUIRED IF A PATIENT IS A	MINOR)	RELATIONSHIP TO PATIENT EMAIL			
NAME:	PHONE:					
DESIGNATED (CARE PARTNER		RELATIONSHIP	TO PATIENT EN	MAIL	
NAME:	PHONE:					
PRESCRIBI	ING PHYSICIAN'S INFORMATION					
PROVIDER NAME PROVIDER PHONE NUM			₹	PROVIDER EMAIL	PROVIDER EMAIL	
AUTHORIZATION TO SHARE HEALTH INFORMATION By signing below, I acknowledge that I have read and agree to the Authorization to Share Health Information terms on the next page.						
HERE*						
In I	SIGNATURE OF PATIENT OR LEGALLY AUTHORIZED REPRESENTATIVE			DATE (MM/DD/YYYY	<u>'</u>)	

CONSENT FOR COPAY PROGRAM (OPTIONAL)

By signing below, I acknowledge that I have read and agree to the Alexion OneSource CoPay Program terms and conditions available at https://alexiononesource.com/CoPay or on request by contacting OneSource at 1.888.765.4747.

SIGNATURE OF PATIENT OR LEGALLY AUTHORIZED REPRESENTATIVE

DATE (MM/DD/YYYY)

CONSENT FOR AUTOMATED TEXT COMMUNICATIONS (OPTIONAL)

By signing below, I give Alexion and companies working at Alexion's direction permission to use automated text (SMS) messages to provide patient support services and to provide by signing below, give already and companies working at Alexton's direction perhassor to use actionable text (owld) messages to produce perhassor to receiving text messages as a condition of any purchase of Alexion products or enrollment in these programs; (ii) my telecommunication services provider may charge me for any text messages that I receive from Alexion; and (iii) I may opt out of receiving automated text messages from Alexion at any time without affecting my enrollment in these programs.

PATIENT SERVICES ENROLLMENT FORM



EMAIL: OneSource@Alexion.com

PHONE: 1.888.765.4747 8:30 AM to 8 PM ET Monday-Friday



FAX: 1.800.420.5150

MAIL: 100 College St., New Haven, CT 06510



AUTHORIZATION TO SHARE HEALTH INFORMATION

Alexion Pharmaceuticals, Inc. ("Alexion") offers patient services including educational resources, case management support, and financial assistance for eligible patients.

By signing the prior page, I give permission for my healthcare providers, health plans, other insurance programs, pharmacies, and other healthcare service providers ("My Healthcare Entities") to share information, including protected health information relating to my medical condition, treatment, and health insurance coverage (collectively "My Information") with Alexion and companies working at its direction so that Alexion may use and disclose My Information to:

- review my insurance coverage and eligibility for benefits for treatment with an Alexion product:
- coordinate treatment with an Alexion product, as well as related services, such as arranging home infusion services or vaccination services;
- provide me with educational and promotional materials, contact me about market research or clinical studies, or otherwise contact me about Alexion products, services, programs, or other topics that Alexion thinks may
- remove identifiers from My Information and combine such resulting information with other information for research, regulatory submissions, business improvement projects, and publication purposes; and
- (as applicable to my Alexion product) review my vaccination and prophylaxis history and provide corresponding patient support, such as sending reminders about potential upcoming vaccinations.

I understand that My Healthcare Entities may receive payment from Alexion in exchange for sharing My Information.

I understand that My Information is also subject to the Alexion Privacy Notice available at https://alexion.com/ Legal#privacy, and that the Alexion Privacy Notice provides additional information about Alexion's privacy practices and the rights that may be available to me. Although Alexion has implemented privacy and security controls designed to help protect My Information, I understand that once My Information has been disclosed to Alexion, the Health Insurance Portability and Affordability Act ("HIPAA") may not apply and My Information may be subject to redisclosure.

I understand that I may refuse to sign this Authorization and that My Healthcare Entities may not condition treatment, payment, enrollment, or eligibility for benefits on whether I sign this Authorization. I also understand that if I do not sign this Authorization, I will not be able to receive support through the Alexion OneSource™ Patient Support Program.

This Authorization expires ten (10) years from the date next to my signature, unless I cancel/revoke it sooner, or unless a shorter time frame is required by applicable law.

I understand that I may revoke my authorization, or unsubscribe or modify the services I receive, at any time by mailing a letter to Alexion OneSource Patient Support Program, 100 College Street, New Haven, CT 06510 or by emailing OneSource@Alexion.com. I also understand that modifying my authorization will not affect any use or disclosure of My Information that occurred before Alexion received notice of my cancellation. I also understand I have a right to receive a copy of this Authorization after it is signed and can request a copy at any time by contacting OneSource at 1.888.765.4747.

OneSource Services

Alexion services and support are subject to change. Participation is voluntary, and person(s) may be removed from Alexion services for code of conduct violations.

