

FAX: 1.800.420.5150 MAIL: 1.00 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:

- 1 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)*	PATIENT PHONE NUMBER* PATIEN		EMAIL	
STEP 2: CLINICAL DIAGNOSIS					
SOLIRIS and ULTOMIRIS are FDA approved for antibody-po	sitive status. If a payer requires p	orior authorization and/or has a clinic	al policy, they	may require proof of antibody st	
INDICATION (check one)*: GENERALIZED MYASTHENIA GRAV			ANTI-AChR AN	ITIBODY POSITIVE (gMG)	
= '	gravis without (acute) exacerbation] UNKNOWN (C	ONTACT ONESOURCE FOR QUESTIO	
☐ ICD-10: G70.01 Myasthenia (gravis with (acute) exacerbation				
Complete this section OR attach copies of patient's medical ar	nd pharmacy insurance card(s).*				
☐ PLEA:	SE PROVIDE SUMMARY OF BENEFIT I	NVESTIGATION FOR ULTOMIRIS AND SOI	LIRIS		
☐ COPIES OF PATIENT'S INSURANCE CARD(S) ATTACHED ☐ PATIENT DOES NOT HAVE INSURANCE	PRIMARY MEDICAL INSURANCE	SECONDARY MEDICAL 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 EM	IAIL*	
ADDRESS*	l l		PHONE NUME	HONE NUMBER*	
CITY*	STATE*		ZIP*		
PRACTICE NAME	TAX ID #*		NPI #*		
OFFICE CONTACT NAME	EMAIL		FAX NUMBER	X NUMBER*	
STEP 5: SITE OF CARE					
SELECT OPTION A OR B BELOW*:					
A) PLEASE PROVIDE ASSISTANCE LOCATING AN INFUSI	ON SITE. PLEASE COOF	RDINATE DIRECTLY WITH: HEALTH	CARE PROVIDE	R PATIENT	
B) ASSISTANCE IS NOT NEEDED. PATIENT WILL BE INFUSI	ED AT: PRESCRIBER'S OFFICE	PATIENT'S HOME PREFER	RED INFUSION S	CITE (PLEASE SPECIFY BELOW)	
SITE OF CARE NAME	NPI#		TAX ID#		
ADDRESS	'				
CITY	STATE	;	ZIP		
OFFICE CONTACT FOR FOLLOW-UP			PHONE NUMBE	?	

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 meningococcal infections/sepsis, 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 meningococcal infections, also available on www.SOLIRIS.net.

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



PHONE: 1.888.765.4747





Fields in red with asterisks are required.*

	NAME (FIRST, LAST)*			DATE OF BIRTH (MM/DD/YYYY)*			
	ATTENT PARIE VINOT, EACT						
STEP 6	6: CLINICAL INFORMATION						
CHECK A	ALL PREVIOUS GENERALIZED MYASTI	HENIA GRAVIS (gMG) THERA	PIES:				
		PHENOLATE MOFETIL	☐ PREDN	ISONE RITU	KIMAB	☐ OTHER	
☐ IVIg	☐ PLASM	IAPHERESIS	PYRIDO	OSTIGMINE EFGA	RTIGIMO	 D	
MGFA CL	LASSIFICATION:			CURRENT MG-ADL SCORE:			
	Abbreviations: IVIg, intrave	nous immunoglobulin; MG-AD	DL, Myasthenia Grav	is Activities of Daily Living; MGFA, Myasth	enia Grav	ris Foundation of America.	
STEP 7	7: PRESCRIPTION						
YOU MAY	Y USE THIS SECTION TO PROVIDE A F	PRESCRIPTION FOR ULTOMII	RIS OR SOLIRIS, OR	YOU MAY PROVIDE A SEPARATE PRESC	RIPTION		
	Rx ULTOMIRIS 100 mg/mL PATIENT WEIGHT:	HCPCS CODE: J1303 PER	UNIT	Rx SOLIRIS 10 mg	J/mL HCI	PCS CODE: J1300 PER UNIT	
LOADIN	IG DOSE:	MAINTENANCE DOSE:		LOADING DOSE:		MAINTENANCE DOSE:	
SIG: INFl	USE INTRAVENOUSLY mg	SIG: INFUSE INTRAVENOR	USLYmg	SIG: INFUSE INTRAVENOUSLY	mg	EVERY 2 WEEKS. START 2 WEEKS AFT	
	O. COVERS THE PATIENT FOR THE	EVERY 8 WEEKS. START		WEEKLY FOR THE FIRST 4 WEEKS, FOLL			
FIRST 2		COMPLETION OF LOADING		BYmg FOR THE 5TH WEEK.		THE 5TH WEEK'S DOSE IS COMPLETE.	
	ER:	OTHER:		QTY 0F 300 mg/30 mL		☐ OTHER: QTY OF 300 mg/30 mL VIALS: REFILLS:	
QTY OF 3	300 mg/3 mL REFILLS: 0	QTY OF 300 mg/3 mL VIALS:					
VIALO: _	KEFILLS: U	QTY 0F 1100 mg/11 mL					
		VIALS:					
	Alexion complement-inhibitor t Vaccination	herapies are available or dates provided as part o		lelines below.* trictive program under a Risk Evalu sed to confirm vaccination prior to	ation ai	nd Mitigation Strategy (REMS). g treatment.	
	Antibiotic prophylaxis adı			s, start date://			
	Patient has received or is		he required vacc	inations per ACIP guidelines.		—	
	Patient has received or is	scheduled to receive th	he required vacc llowing informat	inations per ACIP guidelines. ion: MenB			
YES	Patient has received or is	scheduled to receive the following the follo	he required vacc llowing informat 1st Dose Dat	inations per ACIP guidelines. ion:	NO	☐ VACCINATION SUPPORT fro	
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Please verify your local prescribing requirements (eg, New York prescribers must provide a separate prescription).

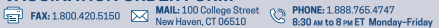
Please see Indications & Important Safety Information on page 4 and full <u>Prescribing Information</u> and <u>Medication Guide</u> for ULTOMIRIS, including Boxed WARNING regarding serious and life-threatening meningococcal infections/sepsis, also available on www.ULTOMIRIS.com.

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 meningococcal infections, also available on www.SOLIRIS.net.

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VACCINATION ORDER FORM









PATIENT NAME (CIT	RMATION					
PATIENT NAME (FIRST, MIDDLE INITIAL, LAST)			PATIENT DATE OF BIRTH (MM/DD/YYYY)			
ADDRESS		CITY			STATE	ZIP
PHONE NUMBER				HEIGHT	WEIGHT	
HEALTHCARE	PRESCRIBER INFORMATION					
FIRST NAME		LAST NAME		PHONE NUMBER		FAX NUMBER
ADDRESS		CITY			STATE	ZIP
OFFICE CONTACT N	NAME			NPI		
CLINICAL INFO	DRMATION					
Primary Diagno	sis Description: Encounter for Immun	ization		ICD-10 CODE: Z2	23	
MENINGOCOC	CAL VACCINATIONS ARE INDICATED FOR P	PATIENTS, INCLUDING PE	OPLE OVER 25 YI	EARS OF AGE, WHEN	ON A COMPLEME	NT INHIBITOR TREATMENT.
	nmittee on Immunization Practices (ACIP) rel ated at least 2 weeks prior to first dose of Al					
	MenACWY	ONE (1) REQUIRED	FROM EACH GRO	IUP	MenB	
	ES ARE NOT INTERCHANGEABLE. PATIENT MUS					
	THE VACCINATION SCHEDULE FOR CHILDREN ≤1					
, , , , , , , , , , , , , , , , , , , ,			Two serogroup B meningococcal (MenB) 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-TT]) 90619 Menveo (meningococcal groups A, C, W, and Y oligosaccharide diphtheria CRM conjugate vaccine [MenACWY-CRM]) 907340 Bexsero (MenB-4C) 90620 Trumenba (MenB-FHbp) 90621 						
	MenACWY	DOSING	SCHEDULE		MenB	
□ Dose 1: Day 0 □ Dose 2: At leas □ Booster dose	st 8 weeks after Day 0		For	Bexsero: At least (or Trumenba: 1-2 montl menba only): 6 mont	ns after Day 0	ual to) 1 month after Day 0
3 years after comp	ndations, those who remain at increased ris oletion of the primary series and every 5 yea ies and every 5 years thereafter. MenB: Adm	ars thereafter. For childre	n 7 years old or ol	der and adults, admi	nister a booster do	
pa. , 0011		illilistel a boostel dose of	vaccine T year at	ter series completio	n and then every 2	se 5 years after completion to 3 years thereafter.
par y 0011	NOTE: ALL VACCINES LISTED		-	·	-	to 3 years thereafter.
	NOTE: ALL VACCINES LISTED RDERS (HOME ADMINISTRATION O	O ABOVE ARE ADMINI	STERED INTRA	·	-	to 3 years thereafter.
ANCILLARY OF		D ABOVE ARE ADMINI DNLY – USE AS NEEL	STERED INTRA	MUSCULARLY AT	A DOSE OF 0.5	to 3 years thereafter.
ANCILLARY OF Anaphylaxis Kit - Diphenhydrami NS 500 mL bag	RDERS (HOME ADMINISTRATION C The following items will be dispensed: ine 50 mg/mL 1 mL vial x 1. Inject 25 mg IM I g x 1. Infuse 500 mL IV at KVO rate PRN anap	DABOVE ARE ADMINI ONLY - USE AS NEED PRN for allergic reaction. ohylaxis	STERED INTRA DED) May repeat x 1 do	MUSCULARLY AT	A DOSE OF 0.5	to 3 years thereafter.
ANCILLARY OF Anaphylaxis Kit - Diphenhydrami NS 500 mL bag Epinephrine am General Anaphyla Administer emerge dose if necessary.	RDERS (HOME ADMINISTRATION C The following items will be dispensed: ine 50 mg/mL 1 mL vial x 1. Inject 25 mg IM I g x 1. Infuse 500 mL IV at KVO rate PRN anap npule/vial 1 mg/mL (1:1000) 1 mL x 2 ampul	DABOVE ARE ADMINI DNLY - USE AS NEED PRN for allergic reaction. chylaxis les/vials. Inject 0.3 mg So chinephrine as above and reate CPR if needed. Call EN	STERED INTRA DED) May repeat x 1 do PRN for adverse epeat dose if necessity.	MUSCULARLY AT ose in 15 min PRN if reaction. May repeatessary. Administer in	A DOSE OF 0.5 of the control of the	to 3 years thereafter. mL 5 min PRN dramine as above and repeat
ANCILLARY OF Anaphylaxis Kit - 1 Diphenhydrami NS 500 mL bag Epinephrine am General Anaphyla Administer emerge dose if necessary, hypotensive. Notif	RDERS (HOME ADMINISTRATION C The following items will be dispensed: ine 50 mg/mL 1 mL vial x 1. Inject 25 mg IM I g x 1. Infuse 500 mL IV at KVO rate PRN anapapule/vial 1 mg/mL (1:1000) 1 mL x 2 ampul xis Instructions ency medications as ordered. Administer ep Place peripheral IV and administer NS. Initia fy prescriber and Nursing Director or pharma	DABOVE ARE ADMINI DNLY - USE AS NEED PRN for allergic reaction. phylaxis les/vials. Inject 0.3 mg So pinephrine as above and reate CPR if needed. Call EN acist.	STERED INTRA DED) May repeat x 1 do PRN for adverse epeat dose if neodis (activate the emedically necessarial) applicable pro	muscularly at ose in 15 min PRN if reaction. May repeatessary. Administer in the immergency medical starts for the patient and escription requirements.	a DOSE OF 0.5 in the improvement in the x 1 dose in 5 to 18 injectable diphenhydystem). Monitor vit in the thing i diagnosis identifints: (iii) I am author	to 3 years thereafter. mL 5 min PRN dramine as above and repeat al signs—elevate legs if ed on this form; (ii) I am authorizing Alexion to forward the pat

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

DATE (MM/DD/YYYY)

PRESCRIBER SIGNATURE (NO STAMPS) - MAY SUBSTITUTE



FAX: 1.800.420.5150 🔀



MAIL: 100 College Street New Haven, CT 06510







INDICATION & IMPORTANT SAFETY INFORMATION FOR **ULTOMIRIS®** (ravulizumab-cwvz)

INDICATION

What is ULTOMIRIS?

ULTOMIRIS is a prescription medicine used to treat adults with a disease called generalized Myasthenia Gravis (gMG) who are anti-acetylcholine receptor (AChR) antibody positive.

It is not known if ULTOMIRIS is safe and effective for the treatment of gMG in children.

IMPORTANT SAFETY INFORMATION

What is the most important information I should know about ULTOMIRIS?

ULTOMIRIS is a medicine that affects your immune system and can lower the ability of your immune system to fight infections.

- ULTOMIRIS increases your chance of getting serious and life-threatening meningococcal infections that may quickly become life-threatening and cause death if not recognized and treated early.
- 1. You must receive meningococcal vaccines at least 2 weeks before your first dose of ULTOMIRIS if you are not vaccinated.
- 2. If your healthcare provider decided that urgent treatment with ULTOMIRIS is needed, you should receive meningococcal vaccination as soon as possible.
- 3. If you have not been vaccinated and ULTOMIRIS therapy must be initiated immediately, you should also receive 2 weeks of antibiotics with your vaccinations.
- 4. If you had a meningococcal vaccine in the past, you might need additional vaccination. Your healthcare provider will decide if you need additional vaccination.
- 5. Meningococcal vaccines reduce but do not prevent all meningococcal infections. Call your healthcare provider or get emergency medical care right away if you get any of these signs and symptoms of a meningococcal infection: headache with nausea or vomiting, headache and fever, headache with a stiff neck or stiff back, fever, fever and a rash, confusion, muscle aches with flu-like symptoms and eyes sensitive to light.

Your healthcare provider will give you a Patient Safety Card about the risk of meningococcal infection. Carry it with you at all times during treatment and for 8 months after your last ULTOMIRIS dose. It is important to show this card to any healthcare provider or nurse to help them diagnose and treat you quickly.

ULTOMIRIS is only available through a program called the **ULTOMIRIS REMS.** Before you can receive ULTOMIRIS, your healthcare provider must: enroll in the ULTOMIRIS REMS program; counsel you about the risk of meningococcal infection; give you information and a Patient Safety Card about the symptoms and your risk of meningococcal infection (as discussed above); and make sure that you are vaccinated with a meningococcal vaccine, and if needed, get revaccinated with the meningococcal vaccine. Ask your healthcare provider if you are not sure if you need to be revaccinated.

ULTOMIRIS may also increase the risk of other types of serious infections. Call your healthcare provider right away if you have any new signs or symptoms of infection.

Who should not receive ULTOMIRIS?

Do not receive ULTOMIRIS if you have a meningococcal infection or have not been vaccinated against meningococcal infection unless your healthcare provider decides that urgent treatment with ULTOMIRIS is needed.

Before you receive ULTOMIRIS, tell your healthcare provider about all of your medical conditions, including if you: have an infection or fever, are pregnant or plan to become pregnant, and are breastfeeding or plan to breastfeed. It is not known if ULTOMIRIS will harm your unborn baby or if it passes into your breast milk. You should not breastfeed during treatment and for 8 months after your final dose of ULTOMIRIS.

Tell your healthcare provider about all the vaccines you receive and medicines you take, including prescription and over-the-counter medicines, vitamins, and herbal supplements which could affect your treatment.

What are the possible side effects of ULTOMIRIS? ULTOMIRIS can cause serious side effects including infusion-related reactions. Symptoms of an infusionrelated reaction with ULTOMIRIS may include lower back pain, tiredness, feeling faint, discomfort in your arms or legs, bad taste, or drowsiness. Stop treatment of ULTOMIRIS and tell your healthcare provider or nurse right away if you develop these symptoms, or any other symptoms during your ULTOMIRIS infusion that may mean you are having a serious infusion reaction, including: chest pain, trouble breathing or shortness of breath, swelling of your face, tongue, or throat, and feel faint or pass out.

The most common side effects of ULTOMIRIS in people with gMG are diarrhea and upper respiratory tract infections.

Tell your healthcare provider about any side effect that bothers you or that does not go away. These are not all the possible side effects of ULTOMIRIS. For more information, ask your healthcare provider or pharmacist. Call your healthcare provider right away if you miss an ULTOMIRIS infusion or for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.

Please see full Prescribing Information and Medication Guide for ULTOMIRIS, including Boxed WARNING regarding serious and life-threatening meningococcal infections/sepsis, also available on www.ULTOMIRIS.com.





FAX: 1.800.420.5150









SOLIRIS

INDICATION & IMPORTANT SAFETY INFORMATION FOR SOLIRIS® (eculizumab)

INDICATION

What is SOLIRIS?

SOLIRIS is a prescription medicine used to treat adults with a disease called generalized myasthenia gravis (gMG) who are anti-acetylcholine receptor (AChR) antibody positive.

It is not known if SOLIRIS is safe and effective in children with gMG.

IMPORTANT SAFETY INFORMATION

What is the most important information I should know about SOLIRIS?

SOLIRIS is a medicine that affects your immune system and can lower the ability of your immune system to fight infections.

- SOLIRIS increases your chance of getting serious and life-threatening meningococcal infections that may quickly become life-threatening and cause death if not recognized and treated early.
- 1. You must receive meningococcal vaccines at least 2 weeks before your first dose of SOLIRIS if you are not vaccinated.
- 2. If your doctor decided that urgent treatment with SOLIRIS is needed, you should receive meningococcal vaccination as soon as possible.
- 3. If you have not been vaccinated and SOLIRIS therapy must be initiated immediately, you should also receive two weeks of antibiotics with your vaccinations.
- 4. If you had a meningococcal vaccine in the past, you might need additional vaccination. Your doctor will decide if you need additional vaccination.
- 5. Meningococcal vaccines reduce but do not prevent all meningococcal infections. Call your doctor or get emergency medical care right away if you get any of these signs and symptoms of a meningococcal infection: headache with nausea or vomiting, headache and fever, headache with a stiff neck or stiff back, fever, fever and a rash, confusion, muscle aches with flu-like symptoms, and eyes sensitive to light.

Your doctor will give you a Patient Safety Card about the risk of meningococcal infection. Carry it with you at all times during treatment and for 3 months after your last SOLIRIS dose. It is important to show this card to any doctor or nurse to help them diagnose and treat you quickly.

SOLIRIS is only available through a program called the **SOLIRIS REMS.** Before you can receive SOLIRIS, your doctor must enroll in the SOLIRIS REMS program; counsel you about the risk of meningococcal infection; give you information and a Patient Safety Card about the symptoms and your risk of meningococcal infection (as discussed above); and make sure that you are vaccinated with the meningococcal vaccine and, if needed, get revaccinated with the meningococcal vaccine. Ask your doctor if you are not sure if you need to be revaccinated.

SOLIRIS may also increase the risk of other types of serious infections. Certain people may be at risk of serious infections with gonorrhea. Certain fungal infections (Aspergillus) may occur if you take SOLIRIS and have a weak immune system or a low white blood cell count.

Who should not receive SOLIRIS?

Do not receive SOLIRIS if you have a meningococcal infection or have not been vaccinated against meningitis infection unless your doctor decides that urgent treatment with SOLIRIS is needed.

Before you receive SOLIRIS, tell your doctor about all of your medical conditions, including if you: have an infection or fever, are pregnant or plan to become pregnant, and are breastfeeding or plan to breastfeed. It is not known if SOLIRIS will harm your unborn baby or if it passes into your breast milk.

Tell your doctor about all the vaccines you receive and medicines you take, including prescription and over-thecounter medicines, vitamins, and herbal supplements which could affect your treatment. It is important that you have all recommended vaccinations before you start SOLIRIS, receive 2 weeks of antibiotics if you immediately start SOLIRIS, and stay up-to-date with all recommended vaccinations during treatment with SOLIRIS.

What are the possible side effects of SOLIRIS? **SOLIRIS** can cause serious side effects including serious infusion-related reactions. Tell your doctor or nurse right away if you get any of these symptoms during your SOLIRIS infusion: chest pain, trouble breathing or shortness of breath, swelling of your face, tongue, or throat, and feel faint or pass out. If you have an infusion-related reaction to SOLIRIS, your doctor may need to infuse SOLIRIS more slowly, or stop SOLIRIS.

The most common side effects in people with gMG treated with SOLIRIS include: muscle and joint (musculoskeletal) pain.

Tell your doctor about any side effect that bothers you or that does not go away. These are not all the possible side effects of SOLIRIS. For more information, ask your doctor or pharmacist. Call your doctor for medical advice about side effects. You are encouraged to report negative side effects of prescription drugs to the FDA. Visit MedWatch, or call 1-800-FDA-1088.

Please see full Prescribing Information and Medication Guide for SOLIRIS, including Boxed WARNING regarding serious and life-threatening meningococcal infections, also available on www.SOLIRIS.net.

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:







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.

PATIENT IN	NFORMATION						
PATIENT NAM	ME (FIRST, MIDDLE INITIAL, LAST)*	DATE OF BIRTH (MM/D	D/YYYY)*	GENDER:	MALE FEMALE NON-BINARY		
		PREF		PREFER TO SE	ELF-DESCRIBE:		
ADDRESS*							
CITY*			STATE*	ZIP*			
	OK TO SEND A TEXT MESSAGE? YES NO OBILE HOME OK TO LEAVE A PHONE MESSAGE? YES NO						
PATIENT DIAG	BNOSIS						
PREFERRED LANGUAGE □ ENGLISH □ SPANISH □ OTHER			PATIENT EMAIL NONE				
LEGAL PATIENT REPRESENTATIVE* (REQUIRED IF A PATIENT IS A MINOR)			RELATIONSHIP TO PATIENT EMAIL				
NAME:	PHONE:						
DESIGNATED	CARE PARTNER		RELATIONSHIP	TO PATIENT	EMAIL		
NAME:	PHONE:						
PRESCRIB	ING PHYSICIAN'S INFORMATION						
PROVIDER NA	AME P	ROVIDER PHONE NUMBER	ł	PI	ROVIDER 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*							
my /	SIGNATURE OF PATIENT OR LEGALLY AUTHORIZED REPRESENTATIVE			DATE (MM/DD/YYYY)			

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 information to me about Alexion products, services, programs, or other topics that Alexion thinks may interest me. I understand that (i) I am not required to consent 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.

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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 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 vaccine services;
- access my credit information and information from other sources to estimate my income, if needed, to assess eligibility for financial assistance programs;
- remove identifiers from Mv Information and combine such resulting information with other information for research, regulatory submissions, business improvement projects, and publication purposes; and
- contact me about market research or clinical studies, provide me with educational and promotional materials, or otherwise contact me about Alexion products, services, programs, or other topics that Alexion thinks may interest me.

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 may no longer protect the information.

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.

