This sample document shows how the ONWARD ZILBRYSQ Start Form (US-ZL-2400131) could be completed. For illustrative purposes only. Not an actual patient.



This document (US-ZL-2400248) is a sample

Start Form for ZILBRYSQ® (zilucoplan) Injection for Subcutaneous Use

Instructions for Prescribers

To get a patient who has been prescribed ZILBRYSQ started in ONWARD™, please follow these steps:

- **1.** Enroll in ZILBRYSQ REMS at www.ZILBRYSQREMS.com. (Once enrolled, you do not need to re-enroll for subsequent patients.)
- 2. Review the REMS safety information with your patient.
- 3. Complete the ZILBRYSQ Start Form, providing all required information.
- 4. Have your patient read and sign the Patient Authorization sections of the Start Form.
- **5.** Fax the completed Start Form to 1-833-FAX-UCB1 (**1-833-329-8221**) or email it to **ucbonward@rxallcare.com**.

Instructions for Patients

- Read and sign the Patient Authorization sections of the Start Form to enroll in ONWARD.
- 2. If you would like us to communicate with you via email and/or text, make sure to check the appropriate consent boxes and provide your email address and/or mobile phone number.
- **3.** You will receive a call from an ONWARD Care Coordinator to discuss the services being requested by you and/or your physician. Please note that when you receive this call, you may see "ONWARD" on your caller ID.

If you have any questions, please call us at 1-844-ONWARD1 (**1-844-669-2731**). A program associate is available to help you **Monday through Friday**, **8 AM to 8 PM**, **Eastern Time**.

Please see Important Safety Information including Boxed Warning at the end of the document.

ONWARD is provided as a service of UCB and is intended to support the appropriate use of UCB medicines. ONWARD may be amended or canceled at any time without notice. Some program and eligibility restrictions may apply.

ZILBRYSQ® is a registered trademark, and ONWARD™ is a trademark, of the UCB Group of Companies.

ZILBRYSQ® (zilucoplan) Injection for Subcutaneous Use Start Form

FAX: 3-FAX-UCB1 (1-833-329-8221	ENROLL ONLINE:) ONWARDhcp-enroll-ZILBRYSQ	-	ONS? CALL: 1 (1-844-669-2	EMAIL: 731) ucbonward@rxallcare.
ices Requested: 🗹 Financial A	ssistance 🗹 Care Coordinator Su	oport 🖊 Patient Assista	nce Program (PA	AP) - Consent Required: see pa
Step 1: Patient Inform	mation * Required field		☑ New t	to Therapy 🔲 On Therapy
First Name* Jane	Middle In	itial ^M Last Name	* Smith	
Date of Birth* 08/13/198				nder: M 🔲 F 🗹 Other 🔲
	arter Street	L Holli	e <u>ma</u> cella dell	Apt#
City* Kansas City	State* KS ZIP* 66101	Preferred Language:] Fnalish ∏Spa	
Communication Preference: Please check here to author you/your caregiver's voicen	rize ONWARD Care Coordinators to	mail janemarysf3810	@gmail.com	
Caregiver Information By providing this information, you as	uthorize ONWARD™ to communicate with	this person regarding your h	ealth condition and	d services provided by the program.
First Name	Last Name	Reli	ationship to Pati	ent
Phone	Home Cell	Primary Point of Cont	act: 🔲 Patient	☐ Caregiver
Email		Caregiver is the a	uthorized legal r	representative of the patient
NOTE: You may attach copies of Check here if patient does not have insurance Check here to investigate meningococal vaccine coverage	the front and back of the patient's insu PRIMARY PHARMACY INSURANCE	PRIMARY MEDI INSURANCE (only required for vaccing	CAL	OTHER (e.g., Secondary)
INSURANCE PROVIDER	BCBS			
INSURANCE PHONE#	913-550-1234			
CARDHOLDER NAME	Jane M Smith			
RELATIONSHIP TO PATIENT	Self			
MEMBER ID	1234567-8904			
GROUP#	3456			
BIN#	1234			
PCN#	7891			
Step 3: Prescriber In	formation * Required fi	e <i>ld</i>		
Prescriber First Name* Jessica		Prescriber Last Na	ame* Barry	
Specialty Neurology NPI#* 5678912345		Tax ID# 1234567893		
Supervising Physician		Supervising Physician NPI		
Practice/Clinic Name Barry Me	edical			
Address* 9087 Field Stre	et			
City Kansas City	State KS		ZIP 66401	
Office Phone#* 913-902-2038		Office Fax# 913-902-2037		
Office Contact Name		Office Contact Email john.doe@barrymedical.com		
Office Contact Phone#		Office Contact Communication Preference: Phone Email		

STEP 4: Clinical Information	* Required field					
ICD-10 Diagnosis*	1 Other	MGFA Classification	n (I, II, III, IV, V):			
MG-ADL Score 19	Date of A	ssessment 03/20)/2024			
AChR Antibody Test: Positive Neg	ative Not Known					
Current Therapies:	Non-steroidal ISTs	Previous The	erapies:	Non-steroidal ISTs		
Eculizumab	Azathioprine	Eculizumab	Rituximab	Azathioprine		
☐ Efgartigimod ☐ IVIG	Cyclophosphamide	Efgartigimod	✓ IVIG	Cyclophosphamide		
Ravulizumab SCIG	Cyclosporine	Ravulizumab	_	Cyclosporine		
	= ' '	_ =		_ = ' ' '		
✓ Oral Corticosteroids	Methotrexate	Oral Corticos		Methotrexate		
Acetylcholinesterase Inhibitors	Tacrolimus	Acetylcholine	sterase Inhibitors	Tacrolimus		
Other	Mycophenolate	Other		Mycophenolate		
Medical Allergies: No allergies						
STEP 5: ZILBRYSQ® Prescription * Required field						
ZILBRYSQ prescription will be triaged to PANTHERx RARE. Your patient will be contacted by PANTHERx to arrange for delivery of ZILBRYSQ.						
Patient First & Last Name* Jane M Si	mith	Г	Date of Birth* (MM/DD/YY	_{YY)} 08/13/1981		
Patient Weight (kg)* 72	Date Weight Measured	d* 7/23/2024				

MEDICATION	WEIGHT CLASS	STRENGTH/FORM	DIRECTIONS FOR ADMINISTRATION	QUANTITY	REFILLS
	Less than 56 kg	16.6 mg/0.416 mL prefilled syringe NDC 50474-990-80	Administer 1 prefilled syringe subcutaneously once daily as directed	28 prefilled syringes	11
ZILBRYSQ (zilucoplan)	56 kg to less than 77 kg	23 mg/0.574 mL prefilled syringe NDC 50474-991-80			
	77 kg and above	32.4 mg/0.81 mL prefilled syringe NDC 50474-992-80			

Appropriate ancillary supplies will be dispensed with the medication unless otherwise indicated.

Physician Attestation and Signature

By signing below, I certify: 1) The therapy is medically necessary and that this information is accurate to the best of my knowledge; 2) I am disclosing this information to UCB, their affiliates, agents, representatives, business partners, and service providers (together "UCB") to help enable treatment for this Patient; 3) The Patient is aware of, has consented to, and has directed my disclosure of their information to UCB so that UCB may contact the Patient to further enable services for those purposes and that such consent and direction applies to disclosures made through the duration of the Patient's therapy; 4) I will not seek reimbursement from any third party for the support UCB provides; and 5) I am licensed to prescribe the prescription medication identified in this form, the prescription complies with my state-specific prescribing requirements and I appoint UCB as my agent for the limited purposes of conveying this prescription by any means under applicable law only to the dispensing pharmacy. I understand that by signing this form, I am requesting support from UCB for Patients receiving ZILBRYSQ®. PRESCRIBER SIGNATURE: PRESCRIBER MUST MANUALLY SIGN AND DATE. RUBBER STAMPS AND SIGNATURE BY OTHER OFFICE PERSONNEL FOR THE PRESCRIBER WILL NOT BE ACCEPTED.

and Last Name* Jessica Barry
Date* 7/23/2024

ATTN: If your local prescribing requirements call for submission of an electronic prescription, please submit to PANTHERx, NPI# 1659762524.

Patient Vaccination History (including Antibiotic Prophylaxis History) & **Prescription(s)** (to be completed by healthcare provider)

The **ZILBRYSQ REMS program requires** the certified dispensing pharmacy (PANTHERX Rare) to contact the prescriber to assess the patient's vaccination status. ZILBRYSQ REMS program requirements are available at: www.ZILBRYSQREMS.com. Full ZILBRYSQ Prescribing Information is available at: www.ZILBRYSQ.com.

Please ensure your patient has had the following in accordance with ZILBRYSQ REMS program requirements:

- Complete and up-to-date vaccinations against MenACWY & MenB, and/or provide prophylactic antibiotics when urgent start needed
- A copy of the ZILBRYSQ REMS Patient Guide & Safety Card

Please provide all vaccination records and prophylactic antihiptic records associated with ZII RRYSO to the dispensing pharmacy

Patient Name			DOB	_		
meningococcal vacci Information. DOES PATIENT NEED	ne(s) as soon as possible TO START ON ZILBRYS	e and p	patient who is not up to dat provide the patient with antib SENTLY? O YES O NO ED. Start date of antibiotic	oacterial dru	g prophylaxis. R	denB vaccines, administer efer to full ZILBRYSQ Prescribin <u>c</u>
			de the dose date details. As nB-4C (Bexsero) and MenB-F			
Me	nACWY	MenB		MenABCWY		
Menveo Menactra ✓ PnQuadfi 1st Dose Date: 4/15/24	Menveo Menactra MenQuadfi 2 nd Dose Date: 6/10/2	24	Bexsero Trumenba 1st Dose Date: 6/14/24	2 nd Dose [Date: <u>7/19/24</u>	Penbraya 1 st Dose Date:
Menveo Menactra MenQuadfi Most Recent	If MenACWY history is available, please sel below: History Unknown No Vaccine Doses	ect	3 rd Dose Date: (Only applicable to Trumenba)	available below: Histor No Va	story is not , please select y Unknown ccine Doses	Penbraya 2 nd Dose Date:
Booster Date:	Received		Most Recent Booster Date:	Receiv	ea	
the retail pharmacy (According to the full	(as applicable) to help co ZILBRYSQ Prescribing Int veeks prior to administeri ection	ordinate formatio	the needed vaccination(s) on, complete or update meni	ngococcal vac s the risk of c	cination (for serog	ith your office, the patient, and roups A, C, W, and Y, and sero- utweighs the risk of developing
_				PICILD KX. (_	
Menveo® (MenACWY-CRM) MenQuadfi® (MenACWY-TT)		Bexsero® (MenB-4C) • 2-dose primary series MenB-4C at least 1 month apart Trumenba® (MenB-FHbp) • 3-dose primary series MenB-FHbp at 0, 1-2, 6 months				
Directions: Administer Intra	amuscularly as directed	Direct	ions: Administer Intramuscu	larly as direct	ed	
Quantity: Refills#:		Quant	city: Refill	s#:		
Revaccinate every 5 years if risk remains		 Note: MenB-4C (Bexsero) and MenB-FHbp (Trumenba) are not interchangeable 1 dose booster 1 year after primary series Revaccinate every 2-3 years if risk remains 				
Prescriber Signature	(Stamps not accepte	ed)*	Gessiea Barry	Di	ate* 7/23/2024	
I certify that the com			is accurate based on n			al judgment. Please
REMS vaccination shipment be heard allow for after at least and MenB; or (1) possible and page of the state o	on requirements andeld with addition administration of weeks post com 2) if urgent therapy then is provided withing and a second control of the control of t	d my al folion miss pletic is ind th ant	patient's vaccination hilow-up to my office to ing vaccinations as pon of the vaccination solicated, after administribiotic prophylaxis.	istory, and to confirm prescribed eries (or beation of me	I request that appropriate I. Patient is to coster if applications of the property of the prop	e timing for dispense start ZILBRYSQ (1) cable) for both MenACWY vaccine(s) as soon as
DISPENSE: I h	nave reviewed the Z	ILBRY	'SQ REMS vaccination i	requiremer	its and my pai	tient's vaccination history

Prescriber Signature (Stamps not accepted)* Jessica Barry

start date must be provided as requested in the urgent ZILBRYSQ therapy request above).

and certify that my patient has been or will be vaccinated as per the full ZILBRYSQ prescribing information prior to beginning treatment with ZILBRYSQ. Patient is to start ZILBRYSQ after at least 2 weeks post completion of the vaccination series (or booster if applicable) for both MenACWY and MenB. PANTHERX Rare is authorized to dispense as soon as possible if urgent therapy is indicated above (antibiotic prophylaxis

Patient Authorization

Please see next page for required HIPAA Authorization

PAP Consent* (Required for the Patient Assistance Program):

By checking here, applicants authorize ONWARD™ PAP and its Administrators to obtain a consumer report. The consumer report, and the information derived from public and other sources, will be used to estimate income as part of the process to decide eligibility to receive free medication from the ONWARD PAP. Upon request, the ONWARD PAP will provide applicants with the name and address of the consumer reporting agency that provides the consumer report. For additional questions about eligibility, please call ONWARD at 1-844-669-2731 (1-844-ONWARD1).

Text Message Consent Language:

By checking here, you agree to receive text messages from UCB ONWARD for patient support. Message and data rates may apply. Message frequency will vary based on need. Text "**HELP**" to **844669** for help. Text "**STOP**" to **844669** to cancel. If you have questions, call 1-844-669-2731 (1-844-ONWARD-1). For more information on how UCB will use your information, please view our privacy policy at www.ucb-usa.com/policy and our text messaging terms and conditions at www.ucbONWARD.com/Text-Terms-Conditions.

If giving consent, please provide your mobile number in Step 1: Patient Information to receive SMS communication.

Terms of Use

UCB ONWARD Patient Support Text Messaging Terms of Use:

The UCB ONWARD Patient Support Text Messaging Program will send adherence and benefit information, as well as allow your Care Coordinator to check-in periodically. This text service is not medical information or medical advice and is not a replacement for seeking medical advice or care from your healthcare provider. Please consult your healthcare provider for any medical advice or information on your condition or treatments.

ONWARD text service is available on the following carriers: Participating carriers: AT&T, Verizon, Sprint, Boost, Virgin, T-Mobile. T-Mobile is not liable for delayed or undelivered messages. Minor carriers: Aio Wireless, Alaska Communications Systems (ACS), Appalachian Wireless (EKN), Bluegrass Cellular, Boost Mobile, Carolina West Wireless, CellCom, Cellular One of East Central IL (ECIT), Cellular One of Northeast Arizona, Cellular One of Northeast Pennsylvania, Chariton Valley Cellular, Coral Wireless (Mobi PCS), Cricket, Cross, C-Spire (CellSouth), Duet IP (Maximum Communications New Core Wireless), Element Mobile (Flat Wireless), Epic Touch (Elkhart Telephone), GCI, Golden State, Google Voice, Hawkeye (Chat Mobility), Hawkeye (NW Missouri), Illinois Valley Cellular, Inland Cellular, iWireless (Iowa Wireless), Keystone Wireless (Immix Wireless/PC Man), Metro PCS, Mosaic (Consolidated or CTC Telecom), MTA Communications, MTPCS (Cellular One Nation), Nex-Tech Wireless, Panhandle Communications, Peoples Wireless, Pine Cellular, Pioneer, RINA, Sagebrush Cellular (Nemont), SI Wireless/Mobile Nation, Simmetry (TMP Corporation), SouthernLinc, SRT Wireless, Thumb Cellular, Union Wireless, United Wireless, U.S. Cellular, Viaero Wireless, Virgin Mobile, and West Central (WCC or 5 Star Wireless). Message and data rates may apply. Alert frequency may vary.

Subscribers will opt-in via the patient enrollment form from "UCB, Inc". UCB will only share this information with its representatives and agents. Message frequency will vary based on need. Text "HELP" to 844669 for help. Text "STOP" to 844669 to cancel. Messages and data rates may apply for any messages sent to you from us and to us from you. If you have any questions about your text plan or data plan, it is best to contact your wireless provider. For all questions about the services provided by this short code, you can send an email to ONWARD@ucb.com or call 1-844-669-2731.

Alerts sent via SMS may not be delivered to you if your phone is not in range of a transmission site, or if sufficient network capacity is not available at a particular time. Even within a coverage area, factors beyond the control of your wireless carrier may interfere with message delivery, including the customer's equipment, terrain, proximity to buildings, foliage, and weather. You acknowledge that urgent alerts may not be timely received and that your wireless carrier does not guarantee that alerts will be delivered. Carriers are not liable for delayed or undelivered messages.

UCB reserves the right to terminate this service in whole or in part, at any time, without notice.

Marketing Consent (Optional):

By checking here & providing your information, you acknowledge you are a U.S. resident and give UCB and its business partners permission to send you information or contact you and/or your healthcare provider regarding your disease as well as information on other related treatments, products and services, and for marketing and informational purposes by phone, email, or mail. You understand that UCB or its business partners will not sell your name, address, email address, or any other information to another party for their own marketing use.

Please ensure an email address and phone number are provided in Step 1: Patient Information.

Patient Authorization to Use/Disclose Health Information (Required)

HIPAA Authorization:*

By signing this form, I hereby authorize each of my physicians, pharmacists (including any specialty pharmacy) that receives my prescription for a UCB medication, and other of my healthcare providers (together, "Providers") and each of my health insurers (together, "Insurers") to disclose information related to my medical condition and treatment (including prescription information), my health insurance coverage and policy number, my name, mailing and email addresses, telephone number, date of birth and Social Security Number (together, "Protected Health Information"), to UCB, Inc. and its agents, service providers, contractors and representatives (together, "UCB"), so that UCB may:

- (i) enroll me in, and contact me about, UCB medication support programs.
- (ii) provide me with educational materials, information, and services related to UCB medications.
- (iii) verify, investigate, assist with, and coordinate my coverage for a UCB medication with my Insurers and Providers.
- (iv) conduct market analyses/research or other commercial activity, including aggregating my Protected Health Information with other data for such analyses.
- (v) assist with analysis related to quality, efficacy, and safety for UCB medication, and, in some cases, contact you to follow up on adverse events in order to obtain additional information.
- (vi) de-identify my Protected Health Information for use for any purpose under applicable law.

I understand that once my Protected Health Information has been disclosed to UCB, federal privacy laws may no longer protect the information and it may be subject to re-disclosure. I understand that one or more Provider and/or Insurer may receive payment from UCB for disclosing my Protected Health Information for some or all of the purposes listed above.

I understand that I am not required to sign this Patient Authorization to Use/Disclose Health Information Authorization, and that if I decline to sign, that will not affect my treatment (including the receipt of UCB medication), payment for treatment, insurance enrollment, or eligibility for insurance benefits, but it may mean that I will not receive the other services described above.

I understand that I may cancel (revoke) this Authorization at any time by calling ONWARD at 1-844-669-2731 (1-844-ONWARD-1) or mailing a letter with my notice of withdrawal to ONWARD, 50 Bearfoot Road, Northborough, MA 01532.

UCB shall provide timely notification of my cancellation (revocation) to my Providers and Insurers. Once my Providers and Insurers receive and process the notice of cancellation (revocation) of this Authorization, my Providers and Insurers may no longer make disclosures of my Protected Health Information to UCB as permitted by this Authorization.

However, canceling this Authorization will not affect any action(s) taken by my Providers or Insurers based on this Authorization before receipt of my notice of cancellation. This authorization expires 10 years from the date it was signed, or such earlier date as required by applicable law unless I cancel it beforehand. I understand that I have the right to receive a copy of this Authorization when it is signed.

I agree to this Patient Authorization Form.
Print Patient First and Last Name: Jane Smith
Date of Birth: 8/13/1981
(If applicable) Print Legal Representative First and Last Name:

Patient or Legal Representative Signature:

If patient is not signing, indicate legal representative's authority to act on patient's behalf (e.g., legal quardian):

Signature*	Jane Smith	
Date:* 7/23/	2024	

INDICATION

ZILBRYSQ® (zilucoplan) is indicated for the treatment of generalized myasthenia gravis (gMG) in adult patients who are antiacetylcholine receptor (AChR) antibody positive.

IMPORTANT SAFETY INFORMATION INCLUDING BOXED WARNING

WARNING: SERIOUS MENINGOCOCCAL INFECTIONS

Life-threatening and fatal meningococcal infections have occurred in patients treated with complement inhibitors; ZILBRYSQ is a complement inhibitor. Meningococcal infection may become rapidly life-threatening or fatal if not recognized and treated early.

- Complete or update meningococcal vaccination (for serogroups A, C, W, and Y, and serogroup B) at least 2 weeks prior to administering the first dose of ZILBRYSQ, unless the risk of delaying therapy outweighs the risk of developing a meningococcal infection. Comply with the most current Advisory Committee on Immunization Practices (ACIP) recommendations for meningococcal vaccinations in patients receiving a complement inhibitor.
- Persons receiving ZILBRYSQ are at increased risk for invasive disease caused by N. meningitidis, even if they develop antibodies following vaccination. Monitor patients for signs of meningococcal infections and evaluate immediately if infection is suspected.

Because of the risk of serious meningococcal infections, ZILBRYSQ is available only through a restricted program under a Risk Evaluation and Mitigation Strategy (REMS) called ZILBRYSQ REMS.

CONTRAINDICATIONS

ZILBRYSQ is contraindicated in patients with unresolved Neisseria meningitidis infection.

WARNINGS AND PRECAUTIONS

Serious Meningococcal Infections

Life-threatening and fatal meningococcal infections have occurred in both vaccinated and unvaccinated patients treated with complement inhibitors; ZILBRYSQ is a complement inhibitor. The use of ZILBRYSQ increases a patient's susceptibility to serious and life-threatening meningococcal infections (septicemia and/or meningitis) caused by any serogroup, including non-groupable strains.

Complete or update meningococcal vaccination (for both serogroups A, C, W, and Y [MenACWY] and serogroup B [MenB]) at least 2 weeks prior to administering the first dose of ZILBRYSQ, according to current ACIP recommendations for meningococcal vaccinations in patients receiving a complement inhibitor.

If urgent ZILBRYSQ therapy is indicated in a patient who is not up to date with both MenACWY and MenB vaccines according to ACIP recommendations, administer meningococcal vaccine(s) as soon as possible and provide the patient with antibacterial drug prophylaxis.

Closely monitor patients for early signs and symptoms of meningococcal infection and evaluate patients immediately if infection is suspected. Withhold administration of ZILBRYSQ in patients who are undergoing treatment for meningococcal infection until the infection is resolved.

ZILBRYSQ REMS

Due to the risk of meningococcal infections, ZILBRYSQ is available only through a restricted program under a REMS called ZILBRYSQ REMS

Under the ZILBRYSQ REMS, prescribers must enroll in the program. Prescribers must counsel patients about the risk of meningococcal infection, provide the patients with the REMS educational materials, and ensure patients are vaccinated with meningococcal vaccines. Additional information on the REMS requirements is available at www.ZILBRYSQREMS.com or 1-877-414-8353.

Other Infections

ZILBRYSQ 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*. Administer vaccinations for the prevention of *Streptococcus pneumoniae* and *Haemophilus influenzae type b* (Hib) infections according to ACIP guidelines. Persons receiving ZILBRYSQ are at increased risk for infections due to these bacteria, even after vaccination.

Pancreatitis And Other Pancreatic Conditions

Pancreatitis and pancreatic cysts have been reported in patients treated with ZILBRYSQ. Patients should be informed of this risk before starting ZILBRYSQ. Obtain lipase and amylase levels at baseline before starting treatment with ZILBRYSQ. Discontinue ZILBRYSQ in patients with suspected pancreatitis and initiate appropriate management until pancreatitis is ruled out or has resolved.

ADVERSE REACTIONS

In a placebo-controlled study, the most common adverse reactions (reported in at least 10% of gMG patients treated with ZILBRYSQ) were injection site reactions, upper respiratory tract infections, and diarrhea.

Please refer to the full Prescribing Information provided by the UCB representative and visit www.ZILBRYSQHCP.com

