FAX this page



PRESCRIPTION AND ENROLLMENT FORM

Complete and Fax: 1-888-525-2431 | Phone: 1-833-30NAPGO (1-833-366-2746)

Patient Name:	DOB:
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Please read the following carefully, then sign and date below.

I. Patient Authorization for Release of Health Information Form

By signing this Authorization, I authorize my healthcare provider, my health insurance company, and my pharmacy providers ("Healthcare Entities") and each of their respective representatives, employees, and agents (collectively "Providers") to disclose information relating to my insurance benefits, medical condition, treatment, and prescription details ("Protected Health Information" or "PHI") to Supernus Pharmaceuticals and its agents, contractors, and other partners, including companies working with Supernus, which may be branded as Circle of Care™ (collectively, "Supernus") for Supernus to (i) provide me with support and related information and materials on any of Supernus' products, including, but not limited to, educational support provided in-person, online or by telephone, financial assistance, and medication adherence support ("Programs"), (ii) conduct data analytics, market research and other internal business activities including, but not limited to, evaluating the Programs provided, and (iii) provide me with information about Supernus' products, services, and programs and other topics of interest for marketing, educational or other purposes.

I understand I have the option and agree to having Supernus perform, on my behalf, an electronic benefit verification and soft credit check under the Fair Credit Reporting Act "FCRA" for the purpose of determining financial eligibility for Patient Assistance Program "PAP" for the provision of free medication, if applicable and I qualify. Additionally, if free medication is provided via PAP or a Bridge Program, I will not seek reimbursement from my insurance plan.

For purposes of clarification, Supernus includes but is not limited to brand specific support through a hub service provider, specialty pharmacy service providers, Circle of Care Clinical Nurse Navigators, as well as other entities under contract with Supernus to support these or similar aspects of the Programs. For purposes of providing support through the Programs, I thereby authorize Supernus to contact me via text messaging, phone, fax, and/or mail – including texts and calls made using an automatic telephone dialing system or prerecorded or artificial voice messages – and to leave a detailed message that includes reference to ONAPGO treatment, as needed.

Once my PHI has been disclosed to Supernus, I understand that state and federal privacy laws no longer protect the information. However, Supernus agrees to protect my PHI by using and disclosing it only for purposes authorized in this Authorization or as required by law or regulations. I understand that my pharmacy provider may receive remuneration from Supernus in exchange for the PHI and/or for any support services provided to me.

I understand that I am not required to sign this Authorization and that my Providers will not condition my treatment, payment, enrollment, or eligibility for benefits on whether I sign this Authorization. This Authorization will expire in 10 years or a shorter period if required by state law, unless I revoke it sooner by writing Circle of Care/Supernus, c/o PharmaCord, PO Box 5490, Louisville, KY 40255. I understand that revoking my Authorization will not affect any use of my information that occurred before my request was processed. I am entitled to a copy of this signed authorization. I certify the information provided on this form is complete and accurate, to the best of my knowledge and I understand that Supernus can revise, change or terminate the Programs at any time.

I have read and understand the Patient Authorization for Release of Health Information Form and agree to the terms. A signature is required in order to receive Supernus services.

Signature of Patient	Date
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addition, I authorize the disclosure of my Protected He	alth Information to the following designated individual(s) (option
addition, I authorize the disclosure of my Protected He	aith information to the following designated individual(s) (optio

II. Marketing/Other Communications Opt-In (optional)

I further authorize Supernus, and companies working with Supernus, any of which may be branded as Circle of Care (collectively "Supernus"), to contact me by mail, email, fax, telephone call, and text message for marketing purposes or otherwise provide me with information about Supernus' products, services, and programs or other topics of interest, conduct market research or otherwise ask me about my experience with or thoughts about such topics. I understand and agree that any information that I provide may be used by Supernus to help develop new products, services, and programs. Note that Supernus will not sell or transfer my personal data to any unrelated third party for marketing purposes without my express permission.

I have read and understand the Marketing/Other Communications Opt-In and agree to the terms.		
Signature of Patient:	Date:	
Designated Individual (print name):	Relationship:	



PRESCRIPTION AND ENROLLMENT FORM



nd Fav: 1-888-525-2/(31 | Phone: 1-833-30NAPGO (1-833-366-27/(6)

Complete and Fax. 1-666-323-2431 Filo	injection, for subcutaneous use • 4.9 mg/mL
1. PATIENT INFORMATION	
Patient Name:	Alternate Contact Name:
Date of Birth: Gender: Male Female	Relationship to Patient:
Address:	Cell Phone: Other Phone: Alternate Contact Email:
	rred Patient Language: English Spanish Other:
Patient Email: Prefe	
2. INSURANCE INFORMATION Please fax a copy of all insurance ca	
Primary Medical Insurance:	
ID #:	·
Phone #:	Phone #:
3. PRESCRIBER INFORMATION	
Prescriber Name (First, MI, Last):	
NPI #:	
Office Name:	
Office Phone:Address:	
	State Zir
4. ONAPGO PRESCRIPTION INFORMATION	_
Confirm Diagnosis: ICD-10: G20.A2 Parkinson's disease without dyskinesia, with fluctuations ICD-10	O: G20.B2 Parkinson's disease with dyskinesia, with fluctuations Other:
FILL OUT ALL SECTIONS BELOW - A, B, C, & D	
Required for all new patients	If replacement needed:
ONAPGO Infusion Device & Kit (1)	Replacement Infusion Device & Kit (1)
Provided to patient at start, programmed by HCP and/or Circle of Care Clinical Nurse Navigator	As determined by manufacturer and insurance eligibility
B Continuous Dosage Prescription	Under medical supervision, infuse subcutaneously Lowest continuous dosage is 1.0 mg/hr
Select one: 16 hours or less orhours	Max continuous dosage 6.0 mg/hr Max recommended daily dose, including extra doses, is 98 mg
Continuous dosage: 1.0 mg/hr or mg/hr	(1 cartridge per day)
	INSTRUCTIONS: Titrate every 1-7 days to therapeutic effect & tolerability
Titrate continuous dosage by: 0.5 mg/hr or 1 mg/h	That every 17 days to the appealed greet a tolerability
Extra Dose(s) Per Day – Select one:	INSTRUCTIONS: Max recommended extra dose is 2 mg per extra dose
☐ None ☐ 1 extra dose ☐ 2 extra doses ☐ 3 extra doses	•
Hold initiating/programming extra dose until continuous dosage is	S No more than 3 extra doses per day over 16 hours with at least 3 hours between extra doses. If 3 extra doses are routinely required consider
optimized Yes No If extra dose selected, complete below	further adjustment of the continuous dosage.
Extra dose: 0.5 mg or 1 mg or mg	
Titrate extra dose by: □ 0.5 mg or □ 1 mg	Titrate every 1-7 days to therapeutic effect & tolerability
D Dispense Quantity: 30 Cartridges (1 cartridge per da	ay) Refills: 11 refills or refills
	e holders One (1) gallon sharps container Four (4) boxes of Infusion Sets [proximal female luer
lock connector, line 42" or less, 90-degree cannula insertion angle, and subcutaneous need Titration Orders: Continuous and extra dose titration procedures, per full Prescribing Info by prescriber, at start, every few days and as needed per patient response until patient r	ormation, under medical supervision. Titrate by dose increments prescribed above, as directed
Prescriber Signatur	re and Date — No Stamps
SIGN and Datte	or Sign and Date
Dispense As Written / Brand Medically Necessary / Do Not Date: Substitute / No Substitution / DAW / May Not Substitute	May Substitute / Product Selection Permitted / Date: Substitution Permissible
CA, MA, NC & PR: Interchange is mandated unless Prescriber writes the	words "No Substitution"
ATTN: New York and Iowa providers, please submit electronic prescrip	tion
	my knowledge and that I prescribed ONAPGO based on medical necessity. I understand that

Supernus Pharmaceuticals and the Circle of CareTM program for benefits eligibility, coverage, and support. I agree to provide additional information if requested and authorize the program to transmit the prescription to the pharmacy. I will not seek reimbursement for any free product from federal or third-party insurers and will only provide the medication to the patient at no cost. I understand that Supernus products and support are complimentary, with no obligation to prescribe, purchase, or recommend them.

FAX this page



PRESCRIPTION AND ENROLLMENT FORM

(apomorphine HCI)
injection, for subcutaneous use · 4.9 mg/mL

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Patient Name:

5. Circle of Care™ Clinical Nurse Navigator Support - Patient will be enrolled in the program as described below:

- Initial Patient Support and Education: Clinical Nurse Navigator schedules and provides education to patient on ONAPGO.
 Clinical Nurse Navigator will review expectations, work to establish proficiency with the ONAPGO device, review Patient Instructions for Use including daily administration and setup, and train and educate on skin health. The patient will be educated to use ONAPGO per the prescribers' orders. Education may be conducted in person, by phone and/or by video call.
- Start (Initiation) Support for Patients: Clinical Nurse Navigator schedules and coordinates ONAPGO start appointment with patient. During start appointment, the patient will be instructed and trained on administering ONAPGO consistent with the prescribers' orders. The Clinical Nurse Navigator will record the patient's response to ONAPGO. In addition, the Clinical Nurse Navigator will educate on skin health, future dose titration and optimization, and daily administration. Start (Initiation) services may be conducted in-home or in-office as indicated by the prescriber.
- Dose Titration Orders: Clinical Nurse Navigator schedules and coordinates ONAPGO titration visits consistent with the prescribers' orders. Titration visits will occur in the patient's home and the ONAPGO continuous dosage and extra dose will be titrated per sections B and C on the prescription form. Titration will continue until an optimal dose has been reached. The device settings will be set by the prescriber and/or a Supernus/MDD US Operations Clinical Nurse Navigator based on sections B and C on the prescription form.
- Ongoing Support and Education: Clinical Nurse Navigator will continue to educate on device proficiency, skin health,
 and daily administration and setup for the patient's entire time on ONAPGO. Clinical Nurse Navigator will also continue to
 educate on dose optimization, as needed, for the patient's entire time on ONAPGO. Clinical Nurse Navigators do not
 give medical advice about a patient's personal treatment plan and will refer patients to their healthcare provider for
 that specific medical advice.

Dose Titration Visits will occur: 🔲 Home 🗌 Office 🗌	Hybrid (In-home and office)	
► Prescriber Preferred Phone Contact Method for Start and Ti	itration:	☐ Cell ☐ Office
Optional – Clinical Nurse Navigator Support Opt-Out: By checking any of the boxes below, you are opting your patient out of full Clinical	ical Nurse Navigator Support	
No, my staff will conduct the Start (Initiation), but a Clini Initial, and Ongoing Support and Education.	ical Nurse Navigator may be used for	Dose Titration Orders,
No, my staff will conduct the Start (Initiation) and Dose T and Ongoing Support and Education.	Fitration , but a Clinical Nurse Naviga	tor may be used for Initial
No, my staff will conduct the Start (Initiation) and Dose T	Fitration, as well as Initial and Ongo	ing Support and Education
SIGN		

NP.SPN.830.2025-0004

or understanding that I would recommend, prescribe, or use ONAPGO or any other Supernus product or service for anyone, (b) my decision to prescribe ONAPGO was based on my determination of medical necessity as set forth herein, (c) all Clinical Nurse Navigator services are intended to go to the full benefit of the patient, and (d) I will provide appropriate medical

care consistent with my professional duties and independent medical judgment. I acknowledge that I cannot bill for activities provided by the Circle of Care program



Patient Copy



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USE

ONAPGO is a prescription medicine used to treat motor fluctuations (OFF episodes) in adults with advanced Parkinson's disease (PD). It is not known if ONAPGO is safe and effective in children.

IMPORTANT SAFETY INFORMATION

Do not take ONAPGO if you are:

- taking certain medicines to treat nausea (ondansetron, granisetron, dolasetron, palonosetron) and alosetron. People taking ondansetron with apomorphine had very low blood pressure and lost consciousness (blacked out).
- allergic to apomorphine or to any ingredients in ONAPGO including sulfite. Sulfites can cause severe, life-threatening allergic reactions, especially in people with asthma.

Call your healthcare provider or get emergency help right away if you have any of the following symptoms of severe life-threatening allergic reaction:

• hives • itching • rash • swelling (eyes, tongue, lips, or mouth) • chest pain • throat tightness • trouble breathing or swallowing.

Before you start using ONAPGO, tell your healthcare provider about all of your medical conditions, including:

- difficulty staying awake during the daytime dizziness, fainting spells, or low blood pressure asthma allergies to any medicines containing sulfites
- heart problems a history of stroke or other brain problems kidney problems liver problems a mental problem called a major psychotic disorder
- drinking alcohol if you are pregnant or plan to become pregnant, or breastfeeding or plan to breastfeed. It is not known if ONAPGO will harm your unborn baby or pass into your breast milk.

Tell your healthcare provider about all the medicines you take, including prescription and non-prescription (over-the-counter) medicines, vitamins, and herbal supplements. ONAPGO and certain other medicines may affect each other and cause serious side effects.

• If you take nitroglycerin under your tongue (sublingual) while using ONAPGO, your blood pressure may decrease and cause dizziness. If possible, lie down before taking it and then try to continue lying down for at least 45 minutes after.

What should I avoid while using ONAPGO?

- Do not drink alcohol. It can increase your chance of developing serious side effects.
- Do not take medicines that make you sleepy.
- Do not drive, operate machinery, or do other dangerous activities until you know how ONAPGO affects you.
- Do not change your position too fast, get up slowly from sitting or lying. ONAPGO can lower blood pressure and cause dizziness or fainting.

What are the possible side effects of ONAPGO?

ONAPGO may cause serious side effects, including:

- blood clots. Infusing ONAPGO into a vein (intravenous) can cause blood clots. Do not infuse ONAPGO in your vein.
- nausea and vomiting are common. May be serious or severe. Your healthcare provider may prescribe medicine (trimethobenzamide) to help decrease nausea/vomiting. Follow your healthcare provider's instructions on how to take/when to stop this medicine.
- sleepiness or falling asleep during the day is common and may be serious. Some people may get sleepy during the day or fall asleep without warning while doing everyday activities such as talking, eating, or driving.
- dizziness is common and may be serious. ONAPGO can lower your blood pressure and cause dizziness. Dizziness can happen when treatment is started or when the dose is increased. **Do not** get up too fast from sitting or lying down, especially if you have been sitting or lying down for a long time.
- falls. Changes that can happen with PD, and effects of some PD medicines, including ONAPGO, as well as trimethobenzamide, can increase your risk of falling.
- infusion site reaction is common and may be serious. Reactions and infections including infusion site nodules, redness, bruising, swelling, rash, and itching may happen.
- hallucinations or psychotic-like behavior. ONAPGO can cause/worsen psychotic-like behavior including hallucinations (seeing or hearing things that are not real), confusion, excessive suspicion, aggressive behavior, agitation, delusional beliefs (believing things that are not real), and disorganized thinking.
- sudden uncontrolled movements (dyskinesia) are common and may be serious. Some people with PD may get sudden, uncontrolled movements after treatment with some PD medicines. ONAPGO can cause/make dyskinesia worse.
- low red blood cells (hemolytic anemia). Tell your healthcare provider if you have: become pale, fast heartbeat, feel more tired or weaker than usual, skin or eyes look yellow, chest pain, shortness of breath or trouble breathing, dark-colored urine, fever, dizziness, or confusion.
- strong (intense) urges. New or increased gambling urges, sexual urges, and other intense urges have been reported.
- heart problems. If you have shortness of breath, fast heartbeat, or chest pain, call your healthcare provider or get emergency help right away.
- serious heart rhythm changes (QT prolongation). Tell your healthcare provider right away if you have a change in your heartbeat (a fast or irregular heartbeat), or faint.
- allergic reaction. Tell your healthcare provider or get medical help right away if you get hives, itching, rash, swelling of the eyes and tongue, or trouble breathing.
- tissue changes (fibrotic complications). Some people have had changes in the tissues of their pelvis, lungs, and heart valves when taking medicines called non-ergot derived dopamine agonists like ONAPGO.
- prolonged painful erections (priapism). May occur. If you have an erection that lasts more than 4 hours, call your healthcare provider or go to the nearest hospital emergency room right away.

Other common side effects of ONAPGO include headache and trouble falling asleep or staying asleep (insomnia).

You are encouraged to report negative side effects of prescription drugs to the FDA. Visit www.fda.gov/medwatch, or call 1-800-FDA-1088.

Patients and care partners must receive complete instructions on the proper use of ONAPGO. Please see Patient Information and talk to your healthcare provider.



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