

INSTRUCTIONS

To be completed in full, signed, and dated, then faxed to 844-394-7155.
For additional assistance, call 84-INGREZZA (844-647-3992), 8 AM – 8 PM EST, M – F.

1 PATIENT INFORMATION

First Name*:	Last 4 digits of the SSN:	
Last Name*:	DOB*: / /	
Address:		
City:	State:	ZIP:
Patient Residence: <input type="checkbox"/> At Home <input type="checkbox"/> LTC <input type="checkbox"/> Group Home <input type="checkbox"/> Other		
US Resident: <input type="checkbox"/> Yes <input type="checkbox"/> No		Gender: <input type="checkbox"/> Male <input type="checkbox"/> Female

Email: _____

Preferred Phone: _____

Is Preferred Phone a Mobile Number? Yes No

Ship Prescription to (optional): Care Partner HCP office LTC facility

I consent to have my Rx shipped to the preference noted and for the INBRACE Program Pharmacy to contact the Care Partner or healthcare provider.

Patient/Authorized Representative Signature:

Date: _____

Description of Authorized Representative's Authority: _____

Alternate Contact/Care Partner Name: _____

Alternate Contact/Care Partner Phone: _____

3 LTC/SNF/ASSISTED LIVING RESIDENTS[†] ONLY:

Resident Room Number:	Ship Prescription to: <input type="checkbox"/> Facility Contact <input type="checkbox"/> Facility Pharmacy		
Facility Pharmacy Name:	Facility Pharmacy Phone:		
Facility Pharmacy Address:	City:	State:	ZIP:

[†]Residents currently covered under Medicare Part A stay are not eligible.

4 CLINICAL INFORMATION

Tardive Dyskinesia (G24.01) Other diagnosis: _____ Allergies: _____

5 INGREZZA START PROGRAM*

Free Trial Program Rx (New Patients)

This program is only available to adults diagnosed with tardive dyskinesia and is not contingent on a purchase of any kind. Product dispensed under this free trial program may not be submitted for reimbursement to any third party payer. Neurocrine reserves the right to modify or cancel the program at any time. I authorize the INBRACE Program Pharmacy to dispense a free one-time, 1-month supply of INGREZZA.

Select one of the following (**NO REFILLS**):

40 mg once a day x 7 days and 80 mg once a day x 21 days **OR**

40 mg once a day x 30 days

Other Rx: _____

Sig: _____ Quantity: _____

6 PRESCRIBER CERTIFICATION

I certify that the information provided in this INGREZZA[®] (valbenazine) capsules Start Program Form is complete and accurate to the best of my knowledge, I have prescribed INGREZZA based on my judgment of medical necessity, and I will supervise the patient's medical treatment. I certify that, where required by federal and/or state law, I have obtained my patient's written legal permission to share identifiable information with Neurocrine Biosciences, Inc., its agents and pharmacies, including but not limited to the INBRACE Support Program Pharmacy. I authorize the forwarding of this prescription and information to a dispensing pharmacy for the INGREZZA Start Program. I understand that neither I nor the patient should seek reimbursement for any free or discounted product received under the program. If the patient has requested shipment to my office, LTC facility, or pharmacy, I agree not to receive any compensation for dispensing the product and I will clearly label and dispense only for use by the patient.

Prescriber or Authorized Agent Signature: *

Date*: _____

(Original signature required—If required by applicable law, please attach copies of all prescriptions on official state prescription forms)

*Indicates required fields.

Please see Indication and Important Safety Information on next page and accompanying INGREZZA full Prescribing Information.

Important Information

INDICATION & USAGE

INGREZZA[®] (valbenazine) capsules is indicated for the treatment of adults with tardive dyskinesia.

IMPORTANT SAFETY INFORMATION

CONTRAINDICATIONS

INGREZZA is contraindicated in patients with a history of hypersensitivity to valbenazine or any components of INGREZZA. Rash, urticaria, and reactions consistent with angioedema (e.g., swelling of the face, lips, and mouth) have been reported.

WARNINGS & PRECAUTIONS

Somnolence

INGREZZA can cause somnolence. Patients should not perform activities requiring mental alertness such as operating a motor vehicle or operating hazardous machinery until they know how they will be affected by INGREZZA.

QT Prolongation

INGREZZA may prolong the QT interval, although the degree of QT prolongation is not clinically significant at concentrations expected with recommended dosing. INGREZZA should be avoided in patients with congenital long QT syndrome or with arrhythmias associated with a prolonged QT interval. For patients at increased risk of a prolonged QT interval, assess the QT interval before increasing the dosage.

Parkinsonism

INGREZZA may cause parkinsonism in patients with tardive dyskinesia. Parkinsonism has also been observed with other VMAT2 inhibitors. Reduce the dose or discontinue INGREZZA treatment in patients who develop clinically significant parkinson-like signs or symptoms.

ADVERSE REACTIONS

The most common adverse reaction ($\geq 5\%$ and twice the rate of placebo) is somnolence. Other adverse reactions ($\geq 2\%$ and $>$ Placebo) include: anticholinergic effects, balance disorders/falls, headache, akathisia, vomiting, nausea, and arthralgia.

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

Please see accompanying INGREZZA full Prescribing Information.