# **PRESCRIPTION REFERRAL FORM**

Confidential—Protected Health Information All fields with an \* are required.

AllianceRx Walgreens Fax: (877) 231-8302 P	hone: <u>(800) 480-9052</u>	Optum Fax: (844) 249-0014	Phone: <u>(877) 719-6330</u>	PANTHERx Rare Fax: (855) 246-3986	Phone: <u>(833) 599-2245</u>
1) Patient Demographic					
First Name*		Last Name*		MI DOB*	
Address					
City			State:	ZIP	
Gender: M F Preferre	d Language	I	Email		
Phone*(    )	-	·	Mobile ( )	-	
Please attach a copy of your patient's c	current insurance card as well as a	n updated medication list.	tandard mobile carrier rates for	voice and text messaging apply.	
2) Prescription Information					
	tolvaptan) tablets, b.i.d., take of a sweekly blister packs tolvaptan) tablets, b.i.d., take of a sweekly blister packs tolvaptan) tablets, b.i.d., take of a sweekly blister packs a sweekly blister packs a sweekly blister packs tolvaptan) tablets, b.i.d., take of a sweekly blister packs tolvaptan) tablets, b.i.d., take of a sweekly blister packs tolvaptan) tablets, b.i.d., take of a sweekly blister packs tolvaptan) tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tolvaptan tablets, b.i.d., take of a sweekly blister packs tablets, b.i.d., take of a	nore than 4 weekly blister packs one 45-mg tablet p.o. upon wa s 2 weekly blister one 60-mg tablet p.o. upon wa s 2 weekly blister one 90-mg tablet p.o. upon wa s 2 weekly blister moderate CYP3A inhibitors (see D one 15-mg tablet p.o. upon wa s 2 weekly blister one 30-mg tablet p.o. upon wa	aking, one 15-mg tablet p.o. 8     packs   1 weekly b     aking, one 30-mg tablet p.o. 8     packs   1 weekly b     aking, one 30-mg tablet p.o. 8     packs   1 weekly b     aking, one 30-mg tablet p.o. 8     packs   1 weekly b     osage and Administration [2.4]). Pa     aking, one 15-mg tablet p.o. 8     packs   1 weekly b     aking, one 15-mg tablet p.o. 8	lister pack Refills	
1 weekly blister pack, 7-day supply, 2 Titration Directions (if needed Special Instructions Known Food/Drug Allergies	1)	s, 14-day supply, 28 tablets, 3 w		ply, 42 tablets, 4 weekly blister pac	ks, 28-day supply, 56 tablets 
Rx Date*	NPI #*		Prescriber Name*		
Yes No prescribed prod authority. I attee   Yes No If the patient instore cover.	ense as Written/Do Not Substitut erapy with JYNARQUE <sup>®</sup> (tolvaptan) is uct. I certify that the information pro st that I am not on the HHS/OIG list surance plan allows the pharmacy to age and initiate the insurance prior a	medically necessary for this patient vided in this form is complete and ac of Excluded Individuals and that I am submit an authorization request on t	based on my best professional judg curate to the best of my knowledge presently authorized under State la sehalf of prescriber, I authorize the S	nent, and I have reviewed the current Pro and medical expertise. I understand that w to prescribe and dispense the request pecialty Pharmacy and its representative ired forms on my behalf as my authorized	t I may not delegate signature ed medication. Is to act as my authorized agent
PRESCRIBER SIGNA			Date*		
Prescriber's signature required	(NO STAMPS).				
First Name*		Last Nan	ne*		МІ
State License #*		DEA #			
Site Name and Address					
			CL_L_*	710*	
City*			State*	<u>ZIP*</u>	
Phone <sup>*</sup> ( )	-		Fax ( )   Required for SPs	-	

Office Contact Name\* (and contact information if different from above)

## INDICATION and IMPORTANT SAFETY INFORMATION for JYNARQUE® (tolvaptan)

### **INDICATION:**

JYNARQUE is indicated to slow kidney function decline in adults at risk of rapidly progressing autosomal dominant polycystic kidney disease (ADPKD).

#### **IMPORTANT SAFETY INFORMATION:**

#### WARNING: RISK OF SERIOUS LIVER INJURY

- JYNARQUE (tolvaptan) can cause serious and potentially fatal liver injury. Acute liver failure requiring liver transplantation has been reported
- Measure transaminases (ALT, AST) and bilirubin before initiating treatment, at 2 weeks and 4 weeks after initiation, then monthly for the first 18 months and every 3 months thereafter. Prompt action in response to laboratory abnormalities, signs, or symptoms indicative of hepatic injury can mitigate, but not eliminate, the risk of serious hepatotoxicity
- Because of the risks of serious liver injury, JYNARQUE is available only through a Risk Evaluation and Mitigation Strategy program called the JYNARQUE REMS Program

### **CONTRAINDICATIONS:**

- History, signs or symptoms of significant liver impairment or injury. This contraindication does not apply to uncomplicated polycystic liver disease
- Taking strong CYP3A inhibitors
- With uncorrected abnormal blood sodium concentrations
- Unable to sense or respond to thirst
- Hypovolemia
- Hypersensitivity (e.g., anaphylaxis, rash) to JYNARQUE or any component of the product
- Uncorrected urinary outflow obstruction
- Anuria

**Serious Liver Injury:** JYNARQUE can cause serious and potentially fatal liver injury. Acute liver failure requiring liver transplantation has been reported in the post-marketing ADPKD experience. Discontinuation in response to laboratory abnormalities or signs or symptoms of liver injury (such as fatigue, anorexia, nausea, right upper abdominal discomfort, vomiting, fever, rash, pruritus, icterus, dark urine or jaundice) can reduce the risk of severe hepatotoxicity. To reduce the risk of significant or irreversible liver injury, assess ALT, AST and bilirubin prior to initiating JYNARQUE, at 2 weeks and 4 weeks after initiation, then monthly for 18 months and every 3 months thereafter.

Hypernatremia, Dehydration and Hypovolemia: JYNARQUE therapy increases free water clearance which can lead to dehydration, hypovolemia and hypernatremia. Instruct patients to drink water when thirsty, and throughout the day and night if awake. Monitor for weight loss, tachycardia and hypotension because they may signal dehydration. Ensure abnormalities in sodium concentrations are corrected before initiating therapy. If serum sodium increases above normal or the patient becomes hypovolemic or dehydrated and fluid intake cannot be increased, suspend JYNARQUE until serum sodium, hydration status and volume status parameters are within the normal range.

Inhibitors of CYP3A: Concomitant use of JYNARQUE with drugs that are moderate or strong CYP3A inhibitors (e.g., ketoconazole, itraconazole, lopinavir/ritonavir, indinavir/ritonavir, ritonavir, and conivaptan) increases tolvaptan exposure. Use with strong CYP3A inhibitors is contraindicated; dose reduction of JYNARQUE is recommended for patients taking moderate CYP3A inhibitors. Patients should avoid grapefruit juice beverages while taking JYNARQUE.

Adverse Reactions: Most common observed adverse reactions with JYNARQUE (incidence >10% and at least twice that for placebo) were thirst, polyuria, nocturia, pollakiuria and polydipsia.

#### Other Drug Interactions:

- Strong CYP3A Inducers: Co-administration with strong CYP3A inducers reduces exposure to JYNARQUE. Avoid concomitant use of JYNARQUE with strong CYP3A inducers
- V<sub>2</sub>-Receptor Agonist: Tolvaptan interferes with the V<sub>2</sub>-agonist activity of desmopressin (dDAVP). Avoid concomitant use of JYNARQUE with a V<sub>2</sub>-agonist.

**Pregnancy and Lactation:** Based on animal data, JYNARQUE may cause fetal harm. In general, JYNARQUE should be discontinued during pregnancy. Advise women not to breastfeed during treatment with JYNARQUE.

To report SUSPECTED ADVERSE REACTIONS, contact Otsuka America Pharmaceutical, Inc. at <u>1-800-438-9927</u> or FDA at <u>1-800-FDA-1088</u> (www.fda.gov/medwatch).

Please see FULL PRESCRIBING INFORMATION, including BOXED WARNING.

