

SPRAVATO® REMS

For Healthcare Setting Use Place Patient Label or Barcode Here

Fax: 1-877-778-0091

Patient Monitoring Form - Outpatient Use Only

INSTRUCTIONS:

This form is intended only for use by outpatient medical offices or clinics, **excluding emergency departments**. You must also submit the **Patient Enrollment Form** if this is the patient's first treatment session.

- 1. Monitor the patient for any signs of sedation, dissociation, or respiratory depression during the 2-hour monitoring period as a requirement of the REMS.
- 2. Complete all required fields on this form after every treatment session for all outpatients enrolled in the SPRAVATO® REMS.
- 3. Submit completed patient monitoring forms within 7 days online at www.SPRAVATOrems.com.

*Indicates Required Field

Patient											
First Name*:	MI:	Last Name*:			Birthdate* (MM/DD/YYYY):	Sex*: Male					
That Name.	IVII.	Last Name .			Diffidate (MIM/DD/1111).	Sex*: Male Other	☐ Female				
Is this the patient's first treatment*? ☐ Yes ☐ No											
If YES, is the patient enrolled*? ☐ Yes ☐ No If NO is selected, please submit the Patient Enrollment Form at www.SpravatoREMS.com or by fax.											
Concomitant Medication											
Is the patient currently taking any of the (including but not limited to benzodiazep	•	,	, .			r blood pressure	changes				
If yes, list medications here:											
Monitoring Healthcare Provider											
First Name*:			Last Name*:								
Telephone*:			Email*:								
·											
Healthcare Setting Information (PRINT)											
Healthcare Setting Name*:											
Healthcare Setting Address 1*:			Healthcare Setting Address 2:								
City*:	State*:	4	ZIP*:	Healthcare S	etting DEA Number*:						
Patient Treatment Session I	nformat	tion (Adm	ninistration and Mor	nitoring)							
Treatment Date* (MM/DD/YYYY):											
Dose Administered*	☐ 56 mg ☐ 84 mg ☐ Other: Lot Number*:										
Treatment Duration* (Patient must be monitored for at least 2 hours)	Total treatment duration minutes (from 1st device administration to completion of monitoring)										
	If not monitored for at least 2 hours, provide reason why:										
Monitoring of Vital Signs*:			Monitoring of Pulse Oxir	metry*:							
Were vital signs in acceptable range prior to:			Was pulse oximetry at an acceptable level prior to administration? ☐ Yes ☐ No								
Administration? ☐ Yes ☐ No			During treatment? ☐ Yes ☐ No								
Treatment session completion? ☐ Yes ☐ No			At treatment session completion? ☐ Yes ☐ No;								
			If a Serious Adverse Event (SAE) occurred during the session, describe in the following section								
Serious Adverse Events of I	nterest										
For this SPRAVATO® REMS, a Serior respiratory depression, or hyperte or is life-threatening.							ent+,				
Did the patient experience a SAE of	interest as	defined ab	ove?								
☐ Yes (describe in following section) ☐ No											



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Patient											
First Name*:	MI:	Last Name*:			Birthdate* (MM/DD/YYYY):	Sex*: ☐ Ma	le □ Female ner				
Monitoring Healthcare Provider											
First Name*:			Last Na	me*:							
Phone*:			Email:								
Treatment Date (MM/DD/YYYY):											
Serious Adverse Events of Inter	est – <i>I</i>	Additional Details (PRINT)								
Janssen Pharmaceuticals, Inc., Safety Depart If needed, add additional pages to document S		ill follow up to obtain more	informa	<u>tion about e</u>	vents reported in this table.						
Event Outcome (Check all that apply)		Event Timing		Description of Serious Adverse Event of Intere (include relevant details such as clinical course therapeutic interventions, comorbidities, prescription/nonprescription medications)		ical course, bidities,	Event Resolution				
The SAE resulted in one or more of the following outcomes: Death Life-threatening Hospitalization Disability/permanent damage Important Medical Event		Date of Event (MM/DD During treatment sess ☐ Yes ☐ No	,		tion □ Dissociation iratory depression □ Hyper	tension	☐ Yes ☐ No ☐ Unknown If yes, time to resolution (min):				
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*Defined as any event that may jeopard	ize the	e patient or may requi	e inter	vention to	prevent one of the abo	ve outcome	S				
Report other product quality complaints or adverse events that are not defined above to: Janssen at 1-800-JANSSEN (1-800-526-7736) or the FDA at 1-800-FDA-1088 or www.fda.gov/medwatch .											