Clinical Criteria Worksheet: Spravato® (esketamine) Nasal Spray

Claim Submission

For Pharmacy and Medical billing:

- Claim processing may be delayed if the information submitted in this worksheet is illegible.
- If the worksheet is left blank or information is missing the claim will be rejected for not enough documentation and reimbursement will be delayed.

For Medical billing only:

• A claim should not be submitted until the drug has been administered to the patient.

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Clinical Criteria – Drug Information Drug Administration: Provide the date of drug administration (MM/DD/YYYY): Provide the expiration date of the drug if the invoice date is greater than 6 months from the date of drug administration (MM/DD/YYYY): **Drug Name and Strength:** Spravato 56 mg Dose Kit: Two 28 mg nasal spray devices Spravato 84 mg Dose Kit: Three 28 mg nasal spray devices Directions: Quantity: **Initiation of Therapy:** Yes No Date therapy initiated: _____ **Continuation of Therapy:** Yes No **Clinical Criteria – Diagnosis** 1. Treatment-resistant depression (TRD) OR Depressive symptoms associated with acute suicidal ideation or behavior **Enrollee Last Name: Enrollee First Name:**

Clinical Criteria – Initiation of Therapy

1.	Before initiating esketamine nasal therapy, was a baseline score on a depression assessment tool (e.g., 17 item Hamilton Rating Scale for Depression [HAMD17], 16-item Quick Inventory of Depressive Symptomatology [QIDS-C16], 10-item Montgomery-Asberg Depression Rating Scale [MADRS]) obtained?
	☐ Yes ☐ No
2.	Has the healthcare outpatient site and the patient been enrolled in the Spravato Risk Evaluation and Mitigation Strategy (REMS)? Yes No
3.	Before prescribing esketamine nasal spray was the New York State Prescription Monitoring Program reviewed? Yes No
4.	For the initial request for patients with a diagnosis of TRD , has the patient had a trial of at least two oral antidepressants prior to initiating esketamine intranasal therapy? Yes No
	Please provide the names of the most recent antidepressant therapies and dates of the trials:
	Antidepressant and strength:
	Date of use:
	Antidepressant and strength:
	Date of use:
5.	Confirm patient observation by a healthcare practitioner for 2 hours during and after esketamine administration.
	☐ Yes ☐ No
6.	Is the patient on an oral antidepressant in conjunction with esketamine nasal spray? Yes No Antidepressant and Strength:
	Directions for Use:

Clinical Criteria – Continuation of Therapy

1. Utilizing the same baseline depression assessment tool, was there an improvement in the patient's score while receiving esketamine treatment?

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	Directions for use:
	Antidepressant and Strength:
	Please provide the patient's current antidepressant therapy and directions for use:
4.	Is the patient on an antidepressant in conjunction with esketamine intranasal therapy? Yes No
	☐ Yes ☐ No
3.	Confirm patient observation by a healthcare practitioner for 2 hours during and after esketamine administratio
	☐ Yes ☐ No
۷.	Before prescribing esketamine nasal spray was the New York State Prescription Monitoring Program reviewed?
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