

# Esketamine Nasal Spray (Spravato<sup>™</sup>) and Intravenous Ketamine for Mental Health Conditions Prior Authorization Request Form

# Medical Policy #087 Esketamine Nasal Spray (SpravatoTM) and Intravenous Ketamine for Mental Health Conditions

## **CLINICAL DOCUMENTATION**

- Clinical documentation that supports the medical necessity criteria for for Esketamine Nasal Spray (Spravato<sup>™</sup>) and Intravenous Ketamine for Treatment-Resistant Depression must be submitted.
- If the patient does not meet all the criteria listed below, please submit a letter of medical necessity with a request for <u>Clinical Exception (Individual Consideration)</u> explaining why an exception is justified.

## **Requesting Prior Authorization Using Authorization Manager**

Providers will need to use <u>Authorization Manager</u> to submit initial authorization requests for services. Authorization Manager, available 24/7, is the quickest way to review authorization requirements, request authorizations, submit clinical documentation, check existing case status, and view/print the decision letter. For commercial members, the requests must meet medical policy guidelines.

To ensure the request is processed accurately and quickly:

- Enter the facility's NPI or provider ID for where services are being performed.
- Enter the appropriate surgeon's NPI or provider ID as the servicing provider, *not* the billing group.

#### **Authorization Manager Resources**

• Refer to our <u>Authorization Manager</u> page for tips, guides, and video demonstrations.

Complete Prior Authorization Request Form for Esketamine Nasal Spray (Spravato) and Intravenous Ketamine (094) using <u>Authorization Manager.</u>

For out of network providers: Requests should still be faxed to 1-888-641-5199.

Patient Information	
Patient Name:	Today's Date:
BCBSMA ID#:	Treatment Start Date:
Date of Birth:	Place of Service: Outpatient  Inpatient

Servicing Physician	Servicing Facility
Name:	Name:
Address:	Address:
Phone #:	Phone #:
Fax#:	Fax#:
This is a secure fax line	□ This is a secure fax line
NPI / TIN#:	NPI / TIN#:

□ Check if the servicing physician is billing entity	Check if the servicing facility is billing entity				
Referring provider if different from servicing provider:	Referring provider if different from servicing provider:				
Name	Phone				
For Esketamine and IV Ketamine requests for TREATME	NT RESISTANT DEPRESSION:				
Initial requests are authorized for up to 28 days - Number	of treatment session requested:				
Esketamine (Spravato) Nasal Spray: Initial, acute therapy	3				
Intravenous Ketamine: Initial, acute treatment 🗖 Subsequ	ient trial				
Individual is 18 or over,					
	sive episode (See Table 1 in medical policy # <u>087</u> ) by a structured				
clinical interview for DSM-5 disorders,					
Current depressive episode is severe depression based	0				
a. Montgomery-Asberg Depression Rating Scale (MA					
b. Hamilton Rating Scale for Depression (HAM-D) sco					
Individual has had an <u>inadequate</u> response to four antidepressant agents from at least:					
<ul> <li>Four antidepressant agents from at least:</li> <li>2 or more different entidepressant alegand (i.e. colective corretorin rountel/e inhibitory, corretorin and</li> </ul>					
<ul> <li>2 or more different antidepressant classes (i.e. selective serotonin reuptake inhibitors, serotonin and</li> <li>acception provide reuptake inhibitora, triggelia antidepressant, humanian, ar mittaganica) AND</li> </ul>					
norepinephrine reuptake inhibitors, tricyclic antidepressants, bupropion, or mirtazapine) AND					
<ul> <li>at least one trial of augmenting agent (i.e. atypical antipsychotic, lithium, or thyroid hormone T3</li> <li>An adequate trial of an antidepressant is defined by <u>BOTH</u> of the following:</li> </ul>					
a. The trial length was at least 6 weeks at generally accepted doses or of sufficient duration as determined by the					
treating physician at the generally accepted doses; AND the Individual was ≥80% adherent to the agent during the					
trial; <u>AND</u>					
<ul> <li>Individual is to receive <u>Esketamine Nasal Spray</u> or <u>Intr</u></li> <li>Individual <u>does not have any</u> of the following:</li> </ul>	avenous Ketamine in conjunction with an oral antidepressant,				
a. Current substance use disorder unless in remission	n (for example, complete abstinence for one month)				
b. Hypersensitivity to esketamine, ketamine, or any or					
c. Previous treatment that was determined not to redu	•				
d. Current episode of delirium					
e. Not currently pregnant or breastfeeding					
f. Aneurysmal vascular disease (including thoracic and abdominal aorta, intracranial and peripheral arterial vessels) or					
arteriovenous malformation					
g. Intracerebral hemorrhage, AND					
Administration of Esketamine (Spravato) or Intravenous	Ketamine is to occur in a provider's office or hospital setting and				
must be monitored by a specialist in the area of a patient's	diagnosis (e.g., psychiatrist), <b>OR</b>				
Request for reauthorization after initial therapy. Request	s will be authorized for up to 1 year when the following				
Request for reauthorization after initial therapy. Requests will be authorized for up to 1 year when the following conditions are met:					
Esketamine (Spravato) Nasal Spray: Initial, acute therapy  Subsequent trial					
Intravenous Ketamine: Initial, acute treatment D Subsequ	uent trial				
Number of treatment session requested:					
Individual has had improvement in depression symptoms as evaluated with an appropriate depression rating scale (e.g. Patient					
	come Scale, Quick Inventory of Depressive Symptomatology-Self				
Report 16 Item, MADRS, HAM-D) AND					

Individual is to receive <u>Esketamine Nasal Spray</u> or <u>Intravenous Ketamine</u> in conjunction with an oral antidepressant <u>AND</u>

□ Individuals with substance use disorder have remained in remission (complete abstinence) <u>AND</u>

Individual does <u>NOT</u> develop any FDA labeled contraindications to esketamine nasal spray including aneurysmal vascular disease, intracerebral hemorrhage, or hypersensitivity to Esketamine, ketamine or any of the excipients. Use of Esketamine is intended to be used consistently with the FDA approved label including meeting Spravato REMS program requirement.

Administration of <u>Esketamine (Spravato) or Intravenous Ketamine</u> is to occur in a provider's office or hospital setting and must be monitored by a specialist in the area of a patient's diagnosis (e.g., psychiatrist)

# For Esketamine or IV Ketamine requests for MAJOR DEPRESSIVE DISORDER WITH ACUTE SUICIDAL IDEATION:

Initial requests are authorized for up to 28 days - Number of treatment session requested: \_

Esketamine (Spravato) Nasal Spray: Initial, acute treatment 
Subsequent trial
Intravenous Ketamine: Initial, acute treatment
Subsequent trial

# □ Individual is 18 or over,

Individual is currently hospitalized and is at an imminent risk for suicide as documented by:

- a. Individual response to a structured assessment for suicidal ideation indicative of imminent risk of suicide (see policy guidelines) AND,
- b. Confirmation of imminent risk of suicide by clinical assessment by a mental health professional/psychiatrist (see policy guidelines)
- □ Individual current depressive episode is moderate or severe based on either of the following scales:
  - c. Montgomery-Asberg Depression Rating Scale (MADRS) ≥ 28\* OR,
    - d. Hamilton Rating Scale for Depression (HAM-D) score ≥ 17\*\*

□ Individual is to receive Esketamine (Spravato<sup>™</sup>) nasal spray or IV Ketamine in conjunction with standard-of-care treatment based on clinical judgment and practice guidelines that may be comprised of oral antidepressant(s), an atypical antipsychotic, or a mood stabilizer.

□ Individual does NOT have any U.S. Food and Drug Administration (FDA) labeled contraindications to the requested agent and esketamine nasal spray is intended to be used consistently with the FDA approved label (see policy guidelines) including meeting Spravato Risk Evaluation and Mitigation Strategy (REMS) program requirements (see policy guidelines).

The prescriber is a specialist in the area of the patient's diagnosis (e.g. psychiatrist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis.

# CPT Codes/ HCPCS Codes/ ICD Codes for Esketamine only (for IV Ketamine see BH and SUD Payment Policy)

	HCPCS codes	Code Description
	G2082	Office or other outpatient visit for the evaluation and management of an established patient that requires the supervision of a physician or other qualified health care professional and provision of up to 56 mg of esketamine nasal self-administration, includes 2 hours post-administration observation
	G2083	Office or other outpatient visit for the evaluation and management of an established patient that requires the supervision of a physician or other qualified health care professional and provision of greater than 56 mg esketamine nasal self-administration, includes 2 hours post-administration observation

# Providers should enter the relevant diagnosis code(s) below:

Code	Description
F33.2	Major Depressive Disorder, Severe