# AvMed

## PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST\*

**Directions:** The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request. All other information may be filled in by office staff; <u>fax to 1-305-671-0200</u>. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. <u>If the information provided is not</u> complete, correct, or legible, the authorization process can be delayed.

## Drug Requested: Spravato<sup>®</sup> (esketamine) (S0013)

## Mark the benefit you would like the PA entered under:

- □ Pharmacy Benefit
- □ Medical Buy and Bill submit prior authorization request via fax to 1-877-535-1391

#### MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete.

| Member Name:              |                                      |
|---------------------------|--------------------------------------|
| Member AvMed #:           | Date of Birth:                       |
| Prescriber Name:          |                                      |
|                           | Date:                                |
| Office Contact Name:      |                                      |
| Phone Number:             |                                      |
| NPI #:                    |                                      |
| DRUG INFORMATION: Authori | zation may be delayed if incomplete. |
| Drug Name/Form/Strength:  |                                      |
| Dosing Schedule:          | Length of Therapy:                   |
| Diagnosis:                | ICD Code, if applicable:             |
| Weight (if applicable):   | Date weight obtained:                |

## **Quantity Limit:**

- Major Depressive Disorder with Acute Suicidal Ideation or Behavior: 8 kits/month; 1 month of treatment
- Treatment-Resistant Depression: 4 kits/month (\*induction dose requires 8 kits/month)

**CLINICAL CRITERIA:** Check below all that apply. All criteria must be met for approval. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied. Check box below for the Diagnosis that applies.

Choose <u>ONE</u> of the following applicable diagnoses below. <u>Provider Please Note</u>: Any indication that is <u>NOT</u> FDA approved will be considered experimental/investigational and <u>NOT</u> medically necessary

## **Treatment Resistant Depression.** <u>ALL</u> the following criteria must be met:

### **Reauthorization is <u>NOT</u> required**

- □ Member must be 18 years of age or older
- **\Box** Spravato<sup>®</sup> must be prescribed by <u>**ONE**</u> of the following:
  - Psychiatrist
  - Provider who has consulted with a psychiatrist (include name/date):
- □ Member must have a diagnosis of treatment resistant depression (TRD) without psychotic features defined by current DSM criteria made or verified by a psychiatrist

#### □ ICD Code/Diagnosis: \_\_\_\_\_

Member must be experiencing moderate to severe symptomology documented by a standardized rating scale that reliably measures depressive symptoms. A current baseline (within previous 30 days, prior to starting Spravato<sup>®</sup>) scale with scoring <u>must be attached</u>.

□ Scale: \_\_\_\_\_

- Date Administered:
- Member must have experienced clinical failure or intolerance with at least two (2) antidepressant therapies from at least two (2) different drug classes (verified by pharmacy paid claims and/or chart notes)
  - Failures must be of adequate dose (maximally tolerated)
  - Failures must be of adequate duration (at least 6 weeks)
  - Adherent fills required (verified by pharmacy claims)
  - Failures must occur during current depressive episode
  - Antidepressant therapy would include any of the following classes:
    - Selective serotonin reuptake inhibitors (e.g., citalopram, fluoxetine, paroxetine, sertraline)
    - Serotonin norepinephrine reuptake inhibitors (e.g., duloxetine, venlafaxine)
    - Bupropion
    - Tricyclic antidepressants (e.g., amitriptyline, clomipramine, nortriptyline)
    - Mirtazapine
    - Monoamine oxidase inhibitors (e.g., selegiline, tranylcypromine)
    - Serotonin modulators (e.g., nefazodone, trazodone)
  - 1. Drug:
     Dose:
     Duration:

     Reason for Discontinuation:
     Dose:
     Duration:

     2. Drug:
     Dose:
     Duration:

     Reason for Discontinuation:
     Duration:
     Duration:

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- □ Member must have experienced clinical failure or intolerance with at least one (1) augmentation therapy (e.g., lithium, liothyronine, antipsychotics or anticonvulsants) (verified by pharmacy paid claims and/or chart notes)
  - Failures must be of adequate dose (maximally tolerated)
  - Failures must be of adequate duration (at least 6 weeks)
  - Adherent fills required (verified by pharmacy claims)
  - Failures must occur during current depressive episode
- Member does <u>NOT</u> have aneurysmal vascular disease (including thoracic and abdominal aorta, intracranial and peripheral arterial vessels), arteriovenous malformation, or a history of intracerebral hemorrhage
- Prescriber must have assessed the member's risk for abuse of controlled substances (i.e., review of medical history, review of state prescription monitoring program (PMP))
- □ Member must be enrolled in the Spravato<sup>®</sup> REMS program
- □ Administering site/provider must be certified in the Spravato<sup>®</sup> REMS program:

**Diagnosis: Major Depressive Disorder with Suicidal Ideation or Behavior** 

**Continuation of inpatient Spravato**<sup>®</sup> therapy, <u>ALL</u> the following criteria must be met:

**One-time authorization per episode for remaining doses required for continuation. Maximum allowable duration = 1 month** 

- Provider <u>MUST</u> submit date of therapy initiation and number of doses administered up to point of request
  - Date Saravati<sup>®</sup> therapy initiated: \_\_\_\_\_\_
  - Number of doses administered since initiation:
- □ Member must be 18 years of age or older

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**Diagnosis: Major Depressive Disorder with Suicidal Ideation or Behavior** 

□ Initiation of outpatient Spravato<sup>®</sup> therapy, <u>ALL</u> the following criteria must be met:

One-time authorization per episode for a duration of 1 month, total of 8 kits/month

- □ Member must be 18 years of age or older
- □ Spravato<sup>®</sup> must be prescribed by or in consultation with a psychiatrist
  - Psychiatrist
  - □ Provider who has consulted with a psychiatrist (include name/date): \_\_\_\_\_
- □ Member must have a diagnosis of major depressive disorder <u>with</u> acute suicidal ideation or behavior verified by a psychiatrist
- □ Spravato<sup>®</sup> must be used in combination with a daily oral antidepressant. **Documentation (pharmacy claims or chart notes) required.**

Drug:\_

- Prescriber must have assessed the member's risk for abuse of controlled substances (i.e., review of medical history, review of state prescription monitoring program (PMP))
- $\hfill\square$  Member must be enrolled in the Spravato<sup>®</sup> REMS program
- □ Administering site/provider must be certified in the Spravato<sup>®</sup> REMS program:

Medication being provided by (check applicable box(es) below):

Physician's office OR Specialty Pharmacy – Proprium Rx

Not all drugs may be covered under every Plan.

If a drug is non-formulary on a Plan, documentation of medical necessity will be required. \*\*Use of samples to initiate therapy does not meet step edit/ preauthorization criteria.\*\* \*<u>Previous therapies will be verified through pharmacy paid claims or submitted chart notes.</u>\*