



Medicaid | Marketplace | Medicare

Sunflower Product Line:

- ☐ KanCare (Medicaid)
☐ Ambetter (Health Insurance Marketplace)
☐ Wellcare By Allwell (Medicare Advantage)

SUBMIT TO

Utilization Management Department

Phone: 1-877-644-4623 Fax: 1-844-824-7705

TRANSCRANIAL MAGNETIC STIMULATION (TMS) REQUEST FORM

Please print clearly - incomplete or illegible forms will delay processing. Please consider Sunflower Clinical and Payment Policies (www.sunflowerhealthplan.com/providers/resources/clinical-payment-policies.html), as appropriate.

PATIENT INFORMATION

Name
Date of Birth
Member ID#
Social Security #
Health Plan #

PROVIDER INFORMATION

Provider Name
Group Name
Provider Tax ID# NPI#
Fax# Phone#
Referral Source

PROVISIONAL DSM-V DIAGNOSIS

The provider must report all diagnoses being considered for this patient.

Primary R/O R/O

CLINICAL INFORMATION

- 1. Will the TMS be administered using a Food and Drug Administration (FDA) cleared device...
2. Is the member experiencing a current major depressive episode?
3. Is the member experiencing any current psychotic symptoms?
4. Has the member received psychotherapy?
5. Did member have lack of significant improvement in depressive symptoms despite adequate trial of evidenced-based psychotherapy?
6. Please describe the reason member did not receive psychotherapy
7. Has member received TMS treatment in the past?
8. Has member had trials of at least four different antidepressants from at least two different pharmacological classes?
9. Has member had trials of at least three different antidepressants from at least two different pharmacological classes?

10. Is antidepressant medications contraindicated for one of the following reasons? (must answer all)
- A. Is there a potential for serious medication adverse effects due to an underlying medical condition?  Yes  No
  - B. Is there a potential for serious worsening of underlying medical condition?  Yes  No
  - C. Is there a potential for serious drug-drug interaction?  Yes  No
  - D. Is member pregnant?  Yes  No  N/A
  - E. Is member postpartum and/or breastfeeding?  Yes  No  N/A
11. Are there any of the following contraindications? (must answer all)
- A. Does the member have a vagus nerve stimulator lead in the carotid sheath?  Yes  No
  - B. Does the member have any implanted stimulators controlled by or that use electrical or magnetic signals?  Yes  No
  - C. Are there any conductive or ferromagnetic or other magnetic-sensitive metals implanted or embedded in member's head or neck?  Yes  No
  - D. Does the member have an acute or chronic psychotic disorder?  Yes  No
  - E. Does the member have a seizure disorder or history of a seizure disorder?  Yes  No
  - F. Does the member abuse any substances at the time of referral or at start of TMS treatments?  Yes  No
  - G. Does member have severe dementia?  Yes  No
  - H. Does the member have a non-adherence with previous treatment for depression?  Yes  No
  - I. Does the member have bullet fragments?  Yes  No
  - J. Does the member have metallic dyes in tattoos?  Yes  No
- 11b. Other implanted stimulators controlled by or that use electrical or magnetic signals such as, but not limited to, the following:
- A. Does the member have deep brain stimulation?  Yes  No
  - B. Does the member have a cardiac pacemaker?  Yes  No
  - C. Does the member have a cardioverter defibrillator?  Yes  No
  - D. Does the member have intracardiac lines?  Yes  No
  - E. Does the member have medication pumps?  Yes  No
12. Which self-reporting rating scale will be used for baseline score and periodic outcome measures?
- Beck Depression Inventory  
Date administered \_\_\_\_\_ Score \_\_\_\_\_
  - PQH-9  
Date administered \_\_\_\_\_ Score \_\_\_\_\_
  - Other  
Date administered \_\_\_\_\_ Score \_\_\_\_\_
  - Date administered \_\_\_\_\_ Score \_\_\_\_\_
13. What other treatment modalities have been tried (example: ECT, EMDR, Ketamine) \_\_\_\_\_
- 
14. Please provide a list of antidepressant medications and/or augmenting agents member has tried in the past as well as current medications:
- 
- 
15. Which treatment sessions are planned?  Repetitive transcranial magnetic stimulation  Deep transcranial magnetic stimulation
16. Is the member experiencing current symptoms of Obsessive Compulsive Disorder (OCD)?  Yes  No
17. Has the member failed to respond to a combination of multiple trials of medication combined with Cognitive Behavioral Therapy (CBT) and/or Exposure and Response Prevention (ERP) for at least 12 weeks during the current episode of illness, as demonstrated by both of the following:
- A. Less than 25% improvement in the Yale Brown Obsessive Compulsive Scale (Y[1]BOCS)  Yes  No
  - B. Failure to respond to psychopharmacologic agents is defined as a lack of clinically significant response in the current OCD episode to four trials of agents from at least two different agent classes, and one of the following:
    - 1. At least two of the treatment trials were administered as an adequate course of mono- or poly-drug therapy with Selective Serotonin Reuptake Inhibitors (SSRIs), Serotonin and Norepinephrine Reuptake Inhibitors (SNRIs), clomipramine, or atypical antipsychotic augmentation involving standard therapeutic doses of at least 12 weeks duration  Yes  No
    - 2. The patient is unable to take SSRI, NSRI, clomipramine, or atypical antipsychotics due to one of the following:
      - a. Drug interactions with medically necessary medications  Yes  No
      - b. Inability to tolerate psychopharmacologic agents, as evidenced by trials of four such agents with distinct side effects in the current episode  Yes  No

18. Are there any of the following contraindications:

- A. History of seizures  Yes  No
- B. Conductive or ferromagnetic or other magnetic-sensitive metals implanted or embedded in head or neck within 30 cm of dTMS coil placement other than dental fillings (e.g. cochlear implants, implanted electrodes/stimulators, aneurysm clips or coils, stents, bullet fragments, metallic dyes in tattoos, deep brain stimulators, vagus nerve stimulators, other implanted electrodes or stimulators)  Yes  No
- C. Vagus nerve stimulator leads in the carotid sheath  Yes  No
- D. Other implanted stimulators controlled by or that use electrical or magnetic signals, (e.g. deep brain stimulation, cardiac pacemaker, cardioverter defibrillator, intracardiac lines and medication pumps)  Yes  No
- E. Substance abuse at time of treatment  Yes  No
- F. Severe dementia  Yes  No
- G. Severe cardiovascular disease  Yes  No
- H. Known non-adherence with previous treatment for OCD  Yes  No
- I. Any mental health and substance use disorders (previously categorized as "Axis I" psychiatric disorders) other than OCD (e.g. including active alcohol or substance abuse, mood disorders, psychotic disorders, other anxiety disorders, etc.); neurological diseases or head injury; or pregnancy  Yes  No

19. Please list the outcome measures from the Yale Brown Obsessive Compulsive Scale (Y-BOCS)

Date administered \_\_\_\_\_ Score \_\_\_\_\_

Date administered \_\_\_\_\_ Score \_\_\_\_\_

20. Notes/comments or additional clinical \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

### SERVICES REQUESTED

CODE REQUESTED	UNITS REQUESTED	HOURS	START AND END DATES
90867			
90868			
90869			

*Not all codes are covered for all lines of business, but may be considered for in lieu of services.*

Please feel free to attach additional documentation to support your request (e.g. updated treatment plan, progress notes, etc.)

**STANDARD REVIEW:**

Standard 14-day time frame will be applied.

**EXPEDITED REVIEW:** By signing below, I certify that applying the standard 14-day time frame could seriously jeopardize the member's health, life or ability to regain maximum function.

\_\_\_\_\_  
Clinician Signature

\_\_\_\_\_  
Date

\_\_\_\_\_  
Clinician Signature

\_\_\_\_\_  
Date

SUBMIT TO  
**Utilization Management Department**  
 Phone: 1-877-644-4623 Fax: 1-844-824-7705