

## REPETITIVE TRANSCRANIAL MAGNETIC STIMULATION REQUEST FORM

Please fax to: 866-878-4250

□ In Network			□ Out of	Network						
Member Name:			OB:	Gender:						
Health Plan:				Member ID:						
					Brovider ID:					
Provider Name:					Provider ID:					
Address:				Email:	:					
Direct Phone:				Fax:						
NPI:				Tax ID:						
Primary Contact:										
1. Has a confirmed diagnosis of severe major depressive disorder (MDD) single or recurrent episode:										
□ F32.2	Major Depressive Disorder, Single Episode, Severe (Without Psychotic Features)									
□ F33.2	Major Depressive Disorder, Recurrent Episode, Severe (Without Psychotic Features)									
2. Does the Member have one or more of the following?										
Resistance to treatment with psychopharmacologic agents as evidenced by a lack of clinically significant response to a trial of psychopharmacological medications in the current depressive episode										
Inability to tolerate psychopharmacologic agents as evidenced by intolerable side effect(s) that are not expected to diminish or resolve with continued administration of the medication										
Currently receiving or is a candidate for and has declined electroconvulsive therapy (ECT) and TMS is considered a less invasive treatment option										
3. Has the order for the TMS procedure been written by a Psychiatrist (MD or DO), who has examined the Member face-to-face and reviewed the record? (Please submit this documentation with the request form)										
□ Yes										
□ No, please explain:										
	mber have a history o	f TMS attempts in th	ne past?							
□ Yes □ No Dates:										
If yes, was there a positive outcome?										
5. Has the Member had an adequate trial of evidence-based psychotherapy, without significant improvement within the past 5 years?										
□ Yes □ No Type of Psychotherapy:										
Dates of evidence-based psychotherapy trial:										
If the Member has not had an adequate trial of evidence-based psychotherapy, what is the reason?										
6. Please fill in the Member's psychotropic medications taken within the past five years: (attach additional information, if needed)										
Medicati	on Name	Dose	Start and End Dat	e	Response					
					□ Improved □ Inadequate Response □ Adverse Response □ Intolerability □ Non-Adherence □ Other:					
					Improved Inadequate Response Adverse Response Intolerability Non-Adherence Other:					

Medication Name	Dose	Start and End D	ate	Res	ponse					
				☐ Improved ☐ Inac ☐ Adverse Response						
				□ Non-Adherence						
				□ Adverse Response □ Non-Adherence						
				□ Improved □ Inac						
				<ul> <li>Adverse Respons</li> <li>Non-Adherence</li> </ul>						
				□ Improved □ Inac						
				□ Adverse Respons						
				□ Non-Adherence	□ Other:					
Please list any Augmenting Agents used:	Please list any Augmenting Agents used:									
If none were used, are they contraindicat	ed?									
Yes Please Explain:					🗆 No					
7. Were any of these meds used during this depressive episode?										
□ Yes, list medications:										
If yes, was improvement inadequate at adequate dose and duration?										
□ Yes □ No										
If yes, was the medication discontinued due to side effects?										
Yes, list medications and side effects:										
8. Please check all that apply:										
<ul> <li>The presence of a medically implanted magnetic-sensitive device or other implanted metal items including, but not limited to, a cochlear implant, implanted cardiac defibrillator (ICD), pacemaker, vagus nerve stimulator (VNS), metal aneurysm clips/coils, staples, or stents, that are located less than or equal to 30 cm from the TMS magnetic coil</li> <li>Acute or chronic psychotic symptoms or disorder</li> <li>Neurological conditions that include epilepsy, cerebrovascular disease, dementia, increased intracranial pressure, having a history of repetitive or severe head trauma, or with primary or secondary tumors in the central nervous system</li> </ul>										
□ Seizure disorder or history of seizures										
□ Substance abuse at time of treatment										
Pregnant or nursing     Current suiside plan or suiside attempt										
Current suicide plan or suicide attempt Non-adherence with previous depression treatments										
History of:  Bipolar Disorder  PTSD  CD Eating Disorder										
□ None of the above										
9. What is the Member's most recent score on a validated self-report depression scale (PHQ-9, MADRS, BDI, HAM-D, GDS, etc.)										
Rating scale used: Score: Date completed:										
10. Treatment Request										
Code		Units	S	tart Date	End Date					
<b>90867</b> : initial, including cortical mapping,										
threshold determination, and delivery ma	nagement									
90868: subsequent delivery and management										
per session										
90869: subsequent motor threshold					 					
redetermination										
with delivery and management										