

Scarelon REPETITIVE TRANSCRANIAL MAGNETIC **STIMULATION REQUEST FORM**

Please Fax to Carelon at 1.800.370.1116

□ In Network □ Out of Network									
Member Name: DOB: Gender:									
Health Plan:					Member ID:				
Provider Name:					Provider ID:				
Address:				Email:	:				
Direct Phone:				Fax:					
NPI:				Tax ID:	D:				
Primary Contact:									
1. Has a confirm	med diagnosis of severe	major depressive	disorder (MDD) singl	e or recu	rrent 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 rece	eiving or is a candidate fo	r and has declined e	lectroconvulsive therap	y (ECT) a	nd TMS is considered a less invasive treatment				
	for the TMS procedure ease submit this docum			O), who h	as examined the Member face-to-face and reviewed				
☐ Yes ☐ No, please ex	plain:								
4. Does the Me	mber have a history o	f TMS attempts in	the past?						
☐ Yes ☐ No Dates:									
	a positive outcome? ☐ PhQ-9 outcome sco	ore and date:							
5. Has the Men	nber had an adequate t	rial of evidence-ba	sed psychotherapy,	without s	significant improvement within the past 5 years?				
☐ Yes ☐ No	herapy:								
	e-based psychotherapy t								
If the Member ha	as not had an adequate tr	ial of evidence-base	d psychotherapy, what	is the reas	son?				
6. Please fill in	the Member's psychot	opic medications	taken within the past	five vear	rs: (attach additional information, if needed)				
Medication		Dose	Start and End Dat		Response				
					☐ Improved ☐ Inadequate Response				
					☐ Adverse Response ☐ Intolerability				
					□ Non-Adherence □ Other:				
					☐ Improved ☐ Inadequate Response				
					☐ Adverse Response ☐ Intolerability				
			1		☐ Non-Adherence ☐ Other:				

Medication Name	Dose	Start and End [Date	Res	sponse					
				☐ Improved ☐ Ina	dequate Response					
				☐ Adverse Respons	e □ Intolerability					
				☐ Non-Adherence						
				☐ Improved ☐ Ina						
				☐ Adverse Respons	-					
				☐ Non-Adherence	☐ Other:					
				☐ Improved ☐ Ina						
				☐ Adverse Respons						
				□ Non-Adherence						
				☐ Improved☐ Ina☐ Adverse Response						
				☐ Non-Adherence	•					
	I									
Please list any Augmenting Agents used:										
If none were used, are they contraindicat										
☐ Yes Please Explain:	☐ Yes Please Explain: ☐ No									
7. Were any of these meds used during this depressive episode?										
☐ Yes, list medications:										
□ No										
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:										
□ No										
8. Please check all that apply:										
 □ 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 □ Acute or chronic psychotic symptoms or disorder □ 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 □ Seizure disorder or history of seizures 										
□ Substance abuse at time of treatment □ Pregnant or nursing □ Current suicide plan or suicide attempt □ Non-adherence with previous depression treatments History of: □ Bipolar Disorder □ PTSD □ OCD □ 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.)										
o. That is the member o most recent score on a validated sen-report depression scale (1 mg-s, mapro, ppi, man-b, gps, etc.)										
Rating scale used:	ating scale used: Score: Date completed:									
10. Treatment Request										
Code		Units	S	tart Date	End Date					
90867: initial, including cortical mapping,										
threshold determination, and delivery ma	nagement									
90868: subsequent delivery and manager	nent									
per session										
90869: subsequent motor threshold										
redetermination ended to the control of the control										
with delivery and management										
-					-					