SUBMIT TO

Utilization Management Department

Phone: 1.866.912.6285 Fax: 1.866.694.3649



INJECTABLE ANTIPSYCHOTICS AUTHORIZATION FORM

Fax completed form to Cepatico at 866.694.3649. Upon receipt of all necessary information Cenpatico will contact you by fax or phone within two business dates after receipt. If you are a participating provider, no authorization is required for Haldol, Haldol-D, Prolixin-D or Geodon.

			Date				
MEMBER INFORMATION			PROVIDER INFORMATIO	PROVIDER INFORMATION			
Name			Provider Name (print)	Provider Name (print)			
DOB				Provider/Agency Tax ID #			
Memb	oer ID #			_			
			Provider/Agency NPI Sub Pr	ovider #			
		norization:	Phone	Fax			
CURR	ENT ICD DIAGNOSIS						
Primar	Additional CHECK ALL APPLICABLE CRITERIA SPECIFIC TO THIS MEDIC These criteria apply to all injectable medications. Additional Member is under a court order for outpatient treatment and members at least 18 years of age. C. The medication is being prescribed by a psychiatrist (MD/D)			t OB Gontinuing Medication Request			
Secon	econdary		□ New Medication Request O □	R Continuing Medication Request			
CHEC	CK ALL APPLICABLE C	RITERIA SPECIFIC TO THIS MED	ICATION REQUEST				
The	ese criteria apply to all in	jectable medications. Additional <i>t</i>	Medication specific to criteria are listed ur	nder each medication:			
□ A.	Member is under a co	urt order for outpatient treatment	and medications. Date of court order (Ple	ease attach the Order):			
□ В.	Member is at least 18	years of age.					
□ C.	The medication is beir	ng prescribed by a psychiatrist (ME	D/DO), Nurse Practioner (ARNP, NP), or Clir	nical Nurse Specialist (CNS).			
□ D.				and other Psychotic Disorders," or is being			
		,	'				
□ E.							
	reduce or discontinue	e it.	, , , , , , , , , , , , , , , , , , , ,	·			
12794	1 RISPERDAL CONSTA						
32//-	T KISI ERBAE CONSTA						
Dose	age	Units Requested	Frequency	Total Units			
25m		50	Q 2 weeks				
37.5		75	Q 2 weeks				
50m		100	Q2 weeks				
	.9	100	42				
CHECK	ALL APPLICABLE CRITERI	A SPECIFIC TO THIS REQUEST FOR RI	SPERDAL CONSTA:				
☐ F.	Member had a docu hospitalization(s).	mented response to Risperdal, but	was non-compliant to the oral form of this	medication, which resulted in inpatient			
☐ G.	Dosage planned is 50 mg or less Q 2 weeks.						
☐ H.	For continuing requests, the member is currently being prescribed requested medication, is stable and has been compliant with treatment; or, the patient was previously prescribed Risperdal Consta by another provider and was stable.						
l.	For new requests, who final drug regimen.	ere the member is titrating from ord	al to injectable medication. Describe the o	cross titration schedule and intended			
	<u> </u>						

J2426 INVEGA SUSTENNA							
Dosage	Units Required	Frequency	Total Units				
39mg	39	Q 1 month					
117mg	117	Q 1 Month					
156mg	156	Q 1 Month					
234mg	234	Q 1 Month					
390mg	390	Initial Dose					
	LIFIC TO THIS REQUEST FOR INVEGA SUS	STENNA:					
A. The member has had prior u	nsuccessful trial of Risperdal Consta. T ty, adverse effects, clinical contraindi	he provder also indicates whether it i					
B. Member had a documented hospitalization.	I response to Invega, but was non-cor	mpliant on the oral form of this medica	ation, which resulted in inpatient				
or, the patient was previously	ember is currently being prescibed requ prescribed Invega Sustenna by anot practice. Please include information o	ther provider, and was stable on the r	medication when he/she began				
D. For new requests, where the r drug regimen.	nember is titrating from oral to injecta	able medication, describe the cross ti	ration schedule and intended				
J2426 INVEGA TRINZA							
J2426 INVEGA TRINZA Dosage	Units Requested	Frequency	Total Units				
	Units Requested 273	Frequency Once every 3 months	Total Units				
			Total Units				

Once every 3 months

819

819mg

CHECK	ALL APPLICABLE CRITERIA						
A.	The member has a documented diagnosis of schizophrenia and has been shown to have been adequately treated with Invega Sustenna for at least 4 months or more than 4 months.						
□ B.	Member has a docume contraindications and re	on the oral form of this medication. The following pted by the provider:					
		Dementia-related psychosis					
	* * * * * * * * * * * * * * * * * * * *	, , , , , , , , , , , , , , , , , , , ,	done, or to any excipients in the for				
		•		ld re-establish treatment with Invega Sustenna x			
_		efore reinitiating Invega Trinza t	. ,				
∐ C.		stable, and has been compliant with treatmen table on the medication when he/she began as available.					
☐ D.	For new requests, where regimen.	the member is titrating from orc	ıl to injectable medication, describe	e the cross titration schedule and intended drug			
J3490	ZYPREXA RELPREVV						
J3490 Dosag		Units Requested	Frequency	Total Units			
	je	Units Requested	Frequency Q2 weeks	Total Units			
Dosag	j e	-		Total Units			
Dosag	j e	150	Q2 weeks	Total Units			
Dosag 150mg	i e	150	Q2 weeks Q2 weeks	Total Units			
150mg 210mg 300mg		150 210 300	Q2 weeks Q2 weeks Q2 weeks	Total Units			
210mg 300mg 405mg		150 210 300 300	Q2 weeks Q2 weeks Q2 weeks Q4 weeks Q4 weeks	Total Units			
210mg 300mg 405mg	ALL APPLICABLE CRITERIA	150 210 300 300 405 SPECIFIC TO THIS REQUEST FOR Z	Q2 weeks Q2 weeks Q2 weeks Q4 weeks Q4 weeks				
210mg 300mg 405mg	ALL APPLICABLE CRITERIA The member has had p	300 300 405 SPECIFIC TO THIS REQUEST FOR Z	Q2 weeks Q2 weeks Q2 weeks Q4 weeks Q4 weeks Q4 weeks	Total Units Provided the second of the seco			
210mg 300mg 405mg	ALL APPLICABLE CRITERIA The member has had part member due to hypers	300 300 405 SPECIFIC TO THIS REQUEST FOR Z prior unsuccessful trial of Risperdorensitivity, adverse effects, clinical	Q2 weeks Q2 weeks Q4 weeks Q4 weeks Q4 weeks Q4 consta. The provder also indicate all contraindications, or ineffective/s	es whether it is clinically contraindicated for this			

☐ C.	For continuing requests, the member is currenlty being prescribed requested medication, is stable, and has been compliant with treatment or, the patient was previously prescribed Invega Sustenna by another provider, and was stable on the medication when he/she began receiving services from your practice. Please include information on the previous provider, as available.					
☐ D.	For new requests, where th regimen.	e member is titrating from oral to injec	ctable medication, describe the cross	titration schedule and intended drug		
☐ E.	Delirium/Sedation Syndrome	a Relprevv are at risk for severe sediate) and must be observed for at least 3 with the se requirements will be met:				
	Services. Fledse describe fle	w mese requirements will be met.				
☐ F.	Provider has identified whi	ch one of 3 possible medication regin	nens will be used for this patient:			
		g/day: 210 IM q2wk or 405 mg IM q4wl	0 .	r 300 mg q4wk.		
		ng/day: 300 mg IM q2wk for 1st 8 weeks, then 210 mg q2w or 405 mg q4wk. ng/day: 300 mg IM q2wk for 1st 8 weeks, continue with 300 mg q2wk thereafter.				
	ABILIFY MAINTENA	L	l -	1		
Dosa		Units Requested	Frequency	Total Units		
300m	ng 	300	Q4 weeks			
400m	ng	400	Q4 weeks			
OUEOK	A.I. A.D.I.G.A.D.I.F. G.D.ITT.D.I.A. G.D.F.					
A.	ALL APPLICABLE CRITERIA SPECIFIC TO THIS REQUEST FOR Abilify Maintena: The member has had a prior unsuccessful trial of Risperdal Consta. Or, the member has had a prior unsuccessful trial of oral Risperdal, making it inappropriate to attempt Risperdal Consta. The provider indicates whether it is clinically contraindicated for this member due to hypersensitivity, adverse effects, clinical contraindications, or ineffective/sub-optimal response to maximized dosing.					
□ В.	,	ed response to Abilify but was non-cor	·	G		
□ c.	For continuing requests, the member was prescribed the medication by this provider, is currently stable, and has been compliant with treatment. Or, the member was prescribed Abilify Maintena by another provider, and was stable on the medication when he/she began receiving services from the most recent provider; the current request includes the information about the previous provider if available.					
□ D.	For new requests, where the	e member is receiving this injectable for		·		

C9470	O ARISTADA							
Dosage		Units Requeste	d	Freque	Frequency		Total Units	
441m	ng	441		Q4 we	eeks			
662m	ng	662		Q4 we	eeks			
882m	ng	882	Q4 wee		weeks		1	
CHECK	ALL APPLICABLE CRITERIA	SPECIFIC TO THIS REQU	UEST FOR Abilify Ari	istada				
□ A.□ B.	For initial (new) reques injectable medication, have been diagnosed whether it is clinically a sub-optimal response to the Member has a documhospitalization (s).	, the provider must des by a psychiatrist with s contraindicated for this to maximized dosing.	scribe the cross titro schizophrenia, and s member due to h	ation sched I has had d Typersensiti	dule and inte a prior succe: ivity, adverse	nded final drug ssful trial of Abilif effects, clinical	regimen. Also, the y Maintena. The pr contraindications,	member must rovider indicates or ineffective/
☐ C.	For continuing request	entia-related psychosits, the member was pre	is or known hyperse escribed the medic	ensitivity re cation by t	eaction to arigonal action to arigonal this provider,	piprazole.	e, and has been co	ompliant
	D. For continuing requests, the member was prescribed the medication by this provider, is currently stable, and has been compliant with treatment. Or, the member was prescribed Abilify Aristada by another provider, and was stable on the medication when he/she began receiving services from the most recent provider; the current request includes information about the previous provider if available. The member must also meet the following criteria for continuing requests: documented adherence to Aristada; demonstrate therapeutic response; and 1. If currently taking 441 mg of Aristada and > 6 weeks have elapsed since the last injection, the plan include concomitant oral aripiprazole; Or 2. If currently taking 662 mg or 882 mg of Aristada and > 8 weeks have elapsed since the last injection the plan includes concomitant oral aripiprazole; 3. No contraindications to or reasons to discontinue Aristada						provider if a; demonstrated he plan includes	
lf you c	are a non-participatir				andication	codo vou gra	a requesting:	
ii you c	are a non-participati	g provider, piedse	indicate which	- Onlei II	realcallon		requesting.	
		486 J0400	J3230	J2680	J0780	J1630	J2060	J3360
Dosag Units F	ge Requested		+					+
Frequ	· ·							+
Total I	· ·							
Physicio	an Signature		Physician Printe	ed Name	ed Name Date			
					SUBMIT TO Utilization Management Department			ment

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