Revised date: 6/30/2017

Opioid Analgesic Treatment Worksheet (Consolidated)

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	Aetna Better Health of Louisiana Fax: 1-844-699-2889 For questions only, please call 1-855-242-0802.			Fee f	or Service (FFS)		
					Louisiana Legacy Medicaid		
	www.aetnabetterhealth.com/lo		,		866-797-2329		
	www.acthabetternearth.com/ro	aisiaria, providers, priarmacy	_	•	estions, please call 1-86 amedicaid.com	66-730-4357.	
	Amarina			ΙΛЦ	ealthcare Connect	ions	
	Amerigroup Fax: 1-888-346-0102				866-399-0929	10113	
	For questions, please call 1-800-	454-3730.		For questions, please call 1-888-929-3790.			
	www.myamerigroup.com/la/pages/medicaid.aspx			<u>www.louisianahealthconnect.com/for-members/pharmacy</u> services/			
	AmeriHealth Caritas Lou	isiana		Unite	edHealthcare		
	Fax: 1-855-452-9131	504 5500	Fax: 1-866-940-7328				
	For questions, please call 1-800-684-5502. www.amerihealthcaritasla.com/pharmacy/index.aspx		For questions, please call 1-800-310-6826. www.uhccommunityplan.com/health-				
	www.amerinearticartasia.com/	рпаннасу/шиех.азрх			sionals/la/pharmacy.htr		
Dia	ase fax the completed form to the ap	anronriato plan usina the desig	unated fa				
	ase jax the completed jorm to the ap cipient Name:	FFS / MCO ID #:	упасеа ја	ix number	Recipient DOB:	Medication Allergies:	
	sident of long-term care facility: Yes	/ No			Recipient Weight (kg):	Recipient Height (ft/in):	
	es, name and phone number:	Dunnauih au Canainliu.			Madiasid Duaniday ID # au	NDI#.	
Pre	escriber Name:	Prescriber Specialty:			Medicaid Provider ID # or	NPI#:	
Pre	escriber Address:		Call-Bac	k Phone#:			
Off	fice Fax#:	Office Contact:			EPSDT Support Coordinat	or (Name/Address): (optional)	
		DRUG INFORMATIO	ON (one d	lrug per re	equest)		
Drı	ug Name / Dosage Form:			Strei	ngth:	Quantity:	
Re	quested medication is short-acting /	long-acting. (CIRCLE ONE) Dir	ections:_				
Dia	agnoses for which the opioid is prescri	ibed (include primary and seco	ndary dia	agnoses ap	pplicable to this request, IC	CD code and description):	
	ngnosis:			iagnosis:_			
Da	te of Diagnosis:		Da	ate of Dia	gnosis:		
Thi	s medication is being used for:	acute condition	c	chronic co	ndition (check one only)		
			CI		v	N	
Is t	his medication being used for modera	ate to severe neuropathic pain	or fibrom	nyalgia?	Yes	No	
ls t	his medication being used for postop	erative pain?Yes		No If y	es, date of surgery		
		/ NON DESERVED A		CID CLE	ONE)	1	
ın	is medication is a PREFERRE	D / NON-PREFERRED A	gent. (0	CIRCLE	ONE) If PREFERRE	D, CONTINUE to page 2.	
lf r	equest is for a non-preferred agent, r	ecipient must have had treatm	ent failur	re with at	least two (2) preferred age	ents.	
	Previously Tried Pref				Reason for Discont		
*Re	efer to the appropriate MCO / FFS website	at top of page for a list of preferre	d agents.				
			_				
OR	if preferred agents have not been pr	eviously tried, provide explana	tion:				
	es individual require an abuse deterre						
res	ident has active substance abuse disc	order or a history of substance	abuse dis	sorder? _	YesNo)	
Do	es individual require Butrans (bupren	orphine transdermal patch) or	Belbuca ((buprenor	phine buccal film) due to o	concern for abuse or	
	nendence with nure onioid agents?				•		

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Is this request for medication prescribed for treatment of pain related to cancer, palliative care, hospice, or end-of-life care? YesNo If NO, proceed to next section.						
If YES, STOP HERE, sign below and fax form to the appropriate plan above.						
Prescriber's signature:(For FFS and Amerihealth Caritas, appropriate diagnosis code must be entered at POS.)						
DOES QUANTITY REQUESTED EXCEED THE MAXIMUM QUANTITY LIMIT? YES / NO (CIRCLE ONE)						
DOES DAILY MED EXCEED THE MAXIMUM MED ALLOWED PER DAY? YES / NO (CIRCLE ONE)						
If answer is YES to either of the	se questions, continue	to next section and complet	te the form in its entirety.			
Request is for:Initiation of therapy			being tapered?YesNo			
If no, explain:						
Recipient's current CUMULATIVE MED PER	R DAY:(include	MED of medication being requested)				
Note: The Louisiana Prescription Monitoring Program (PMP) provides the cumulative MED for all of the recipient's controlled medications. Information is current through the previous day (the day before the PMP is accessed).						
For quantity limit override OR MED overr	ide, explain in detail the need	for requested quantity/MED:				
List treatments that have been tried or ar	e currently being given for this	s condition, both pharmacological and	non-pharmacological:			
	Pharmacological Treatme	nts (both opioid and nonopioid)				
Drug / Strength	Directions	Start Date / End Date (or Current)	Reason for Discontinuation (if applicable)			
₩	Non-pharmac	cological Treatments	Date (or Course			
Treatment		Start Date / End	Date (or Current)			

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PRESCRIBER ATTESTATION

Please indicate YES/True or NO/False for each of the following attestations. Explanation is required for each 'No/False' answer in order for the request to be considered for approval. For short-acting opioids, complete A – G. for long-acting opioids, complete A – L.

-	YES (True)	NO (False)	THE PRESCRIBER ATTESTS TO THE FOLLOWING:		
			A. A complete assessment for pain and function was performed for this patient and documentation is attached .		
IOIDS			B. The patient has been screened for substance abuse / opioid dependence and documentation is attached . (Not required for recipients in long-term care facility.)		
О			C. The PMP will be accessed each time a controlled prescription is written for this patient.		
CTING			D. A treatment plan which includes current and previous goals of therapy for both pain and function has been developed for this patient.		
ONG-A			E. Criteria for failure of the opioid trial and for stopping or continuing the opioid has been established and explained to the patient.		
SHORT AND LO			F. Benefits and potential harms of opioid use have been discussed with this patient. In addition, if the patient has concurrent comorbidities or is taking medications that could potentially cause drug-drug interactions, an assessment of increased risk for respiratory depression has been completed and discussed with the patient. The risk of combining opioids with other central nervous system depressants, such as benzodiazepines, alcohol, or illicit drugs such as heroin, has also been specifically addressed. The level of risk for opioid abuse/overdose with the dose/duration prescribed to the patient has also been discussed.		
			G. An Opioid Treatment Agreement signed by both the patient and prescriber is on file. (Not required for recipients in long-term care facility.)		
SOIOS			H. The patient requires continuous around the clock analgesic therapy for which alternative treatment options have been inadequate or have not been tolerated.		
B. The patient has been screened for substance abuse / opioid dependence and documentation required for recipients in long-term care facility.) C. The PMP will be accessed each time a controlled prescription is written for this patient. D. A treatment plan which includes current and previous goals of therapy for both pain and funct developed for this patient. E. Criteria for failure of the opioid trial and for stopping or continuing the opioid has been establic explained to the patient. F. Benefits and potential harms of opioid use have been discussed with this patient. In addition, concurrent comorbidities or is taking medications that could potentially cause drug-drug interest assessment of increased risk for respiratory depression has been completed and discussed with risk of combining opioids with other central nervous system depressants, such as benzodiazepi illicit drugs such as heroin, has also been specifically addressed. The level of risk for opioid abuthe dose/duration prescribed to the patient has also been discussed. G. An Opioid Treatment Agreement signed by both the patient and prescriber is on file. (Not required in long-term care facility.) H. The patient requires continuous around the clock analgesic therapy for which alternative treat have been inadequate or have not been tolerated. I. Patient previously utilized at least two weeks of short-acting opioids for this condition. Please of dose, duration and date of trial in Pharmacological Treatment Section on page 1.	I. Patient previously utilized at least two weeks of short-acting opioids for this condition. Please enter drug(s), dose, duration and date of trial in <i>Pharmacological Treatment Section</i> on page 1.				
ACTIN			· · · · · · · · · · · · · · · · · · ·		
<u>-</u>			K. Medication has not been prescribed for use as an as-needed (PRN) analgesic.		
LON			L. Prescribing information for requested product has been thoroughly reviewed by prescriber.		
IF NO	FOR ANY O	F THE ABO	VE (A-L), PLEASE EXPLAIN:		

Opioid overdose reversal medications are a covered benefit. Prior authorization is not required for some products. CDC guidelines recommend offering naloxone to patients at increased risk of overdose, defined as: history of overdose or substance use disorder, doses \geq 50 MED /day, or concurrent use with benzodiazepines. *Please refer to the appropriate MCO / FFS website (top of page 1) for a list of preferred agents.*

I certify that the benefits of opioid treatment for this patient outweigh the risks of treatment and that the information provided herein is true and accurate to the best of my knowledge and may be subject to a routine audit requesting the medical information necessary to verify the accuracy of the information provided.

Please note: An approval is not a guarantee of payment. All edits will apply when medication is processed at point-of-sale (POS). Payment on a claim will only be made when the claim is billed correctly and all conditions for payment are met.

Prescriber's Signature:	Date:	

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