LOUISIANA UNIFORM PRESCRIPTION DRUG PRIOR AUTHORIZATION FORM

Submitted to: Optum Rx			Phone: 1-800-711-4555	Fax: 1-	Date:		
0	plannik				011 100 1021		
SECTION II — PI	RESCRIBER INFORMATION	J					
Last Name, First Name MI:		NPI# or	Plan Provider #:	Specialty:			
Address:		City:	City:		State:	ZIP Code:	
Phone: Fax:		Office C	Office Contact Name:		Contact Phone:		
	PATIENT INFORMATION						

Last Name, First Name MI:	DOE	3:	Phone:				lale ther	Female
Address:	Ci	ity:					State:	ZIP Code:
Plan Name (if different from Section I):	Member o	r Medicaid ID #:	Plan Prov	ider ID:				
Patient is currently a hospital inpatient getti	ng ready fo	or discharge?	Yes	No	Date	of Disch	narge:	
Patient is being discharged from a psychiatr	ic facility?		Yes	No	Date	of Discl	harge:	
Patient is being discharged from a residentia	al substanc	e use facility?	Yes	No	Date	of Discl	harge:	
Patient is a long-term care resident? EPSDT Support Coordinator contact informa		-	ne and pho	ne numb	er:			

SECTION IV — PRESCRIPTION DRUG INFORMATION

Requeste	d Drug Name:									
Strength:	Dosage Form:	Route of Admin:	Quantity:	Days' Supply:	Dosage Interval/Directions for Us	e: Expected Therapy Durat	ion/Start Date:			
		-	ation is:		herapy/Initial request uation of therapy/Reauthor	zation request				
For Provi	der Administere	ed Drugs only:								
HCPCS/	HCPCS/CPT-4 Code:NDC#:Dose Per Administration:									
Other C	Other Codes:									
		drug in the phy								
	-	If no, list name a	ind NPI of	servicing pr	rovider/facility:					
SECTION	V – PATIENT	CLINICAL INFO	RMATIO	N						
Primary o	Primary diagnosis relevant to this request: ICD-10 Diagnosis Code: Date Diagnosed:									
Secondar	Secondary diagnosis relevant to this request: ICD-10 Diagnosis Code: Date Diagnosed:									
	-	es, pain is: elated diagnose			_Chronic					
Pertinent laboratory values and dates (attach or list below):										
Date Name of Test						Va	Value			
		•				•				

SEC	CTION V	I - This S	Section For Opioid Medications Only						
Cum	nulative dai	ily MME	sted exceed the max quantity limit allowed?YesNo (If yes, provide justification below.)						
DS	YES (True)	NO (False)	THE PRESCRIBER ATTESTS TO THE FOLLOWING:						
Yes (True) (False) A. A complete assessment for pain and function was performed for this patient. B. The patient has been screened for substance abuse / opioid dependence. (Not required for relations)									
B. The patient has been screened for substance abuse / opioid dependence. (Not required for recipients in long-term care facility.)									
F T			C. The PMP will be accessed each time a controlled prescription is written for this patient.						
-9NO-			D. A treatment plan which includes current and previous goals of therapy for both pain and function has been developed for this patient.						
SHORT AND LONG-ACTING			E. Criteria for failure of the opioid trial and for stopping or continuing the opioid has been established and explained to the patient.						
F. Benefits and potential harms of opioid use have been discussed with this patient.									
SH			G. An Opioid Treatment Agreement signed by both the patient and prescriber is on file. (<i>Not required for recipients in long-term care facility.</i>)						
SOIO			H. The patient requires continuous around the clock analgesic therapy for which alternative treatment options have been inadequate or have not been tolerated.						
S OPIOIDS			 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 pharmacologic/non-pharmacologic treatment section below. 						
LONG-ACTING			J. Medication has not been prescribed to treat acute pain, mild pain, or pain that is not expected to persist for an extended period of time.						
7-5			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 NC	FOR ANY (OF THE ABC	OVE (A-L), PLEASE EXPLAIN:						

SECTION VII - Pharmacologic & non-pharmacologic treatment(s) used for this diagnosis (both previous & current):

Drug name	Strength	Frequency	Dates Started and Stoppe or Approximate Duratior	Describe Response, Reason	
Drug Allergies:		Height (if applicable):	Weight (if applicable):		

Is there clinical evidence or patient history that suggests the use of the plan's pre-requisite medication(s), e.g. step medications, will be ineffective or cause an adverse reaction to the patient? _____Yes _____No (If yes, please explain in Section VIII below.)

SECTION VIII — JUSTIFICATION (SEE INSTRUCTIONS)

By signing this request, the prescriber attests that the information provided herein is true and accurate to the best of his/her knowledge. Also, by signing and submitting this request form, the prescriber attests to statements in the 'Attestation' section of the criteria specific to this request, if applicable.

Signature of Prescriber:____

Date:_____