## **Kentucky Medicaid Pharmacy Prior Authorization Form**

- For Drug Requests (unless noted below) Complete ONLY page 1 of this form.
- For ALL Opioid Requests Complete page 1, 2,3 AND page 4 of this form.
- For Hepatitis C Direct Acting Antiviral (DAA) Therapy Complete page 1 AND page 5 of this form.
- For Synagis® Requests Complete page 1 AND page 6 of this form
- For Buprenorphine Products:

Medimpact	
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<ul> <li>For Pain Management Diagnosis — Co</li> </ul>	mplete pag	ge 1 AND page 2	of this form	ո.	
Complete each section legibly and completely. Include any supporting documents as needed (lab results, chart notes, etc.).					
Plan:		Phone numb	none number:		Fax number:
All Kentucky MCO Plans (MedImpact)		1 (844) 336-2676	i		1 (858) 357-2612
Patient Information:					
Member Name:			Date of B	irth:	
Address: City, State, Zip:					
Sex: □Male □Female	Height:			Wei	ght:
Member ID:	Medicatio	n Allergies:			
Prescriber Information:					
Prescriber Name:			NPI:		
Prescriber Address: City, State, Zip:					
Prescriber Specialty:			DEA:		
Phone:			Fax:		
Diagnosis and Medical Information for Requested Med	dication:	☐ INITIAL REQUE	ST 🗆 F	REAUTHORIZATIO	ON (REFILL) Request with current plan
Diagnosis:		ICD-10 Code:			Date of Diagnosis:
Medication Requested (name, strength and dosage for If request is for an opioid, please continue to page 2.	m):				
Quantity: Days' Supply:		Ex	pected Dur	ration of Therapy	:
Directions for Use:		1			
Rationale for Prior Authorization:					
	-	swer the followi	ng questio	ons:	
1) Has the member tried 2 generic manufactures					
2) Please provide medical justification why the p	atient can	not be appropri	ately treate	ed with the gen	eric form of the drug (allergy,
intolerance to inactive ingredient) Injectable Products: Medical vs Pharmacy Coverage?		□No <i>If yes ple</i>		or the following	quartians
Is the medication being self-administered AND appropriate the self-administer of the s					
<ul><li>2) Is the medication being administered by a home in</li></ul>			□No	эссион от инс р	
Please indicate previous treatment outcomes below:					
Previous Medication Strength Quantity	Directions (	Sig) D	ates (from a	and to)	Reason for Discontinuation
Refer to link for List of Preferred Agents:					
https://kyportal.magellanmedicaid.com/public/clier	nt/static/ke	ntucky/document	s/Preferred	dDrugGuide full.	<u>odf</u>
☐ Patient recently hospitalized  If requesting antibiotics, anti-infective, antidepressants, or	nticonvulsant	s, antinsychotic for	discharae to i	complete the cours	se of prescription, provide
duration: (original + refills)		o, amapo, energe.	anoonan ge to	complete the court	(a) (a) (b) (a) (b) (a) (b) (a) (b) (b) (b) (b) (b) (b) (b) (b) (b) (b
Additional Clinical Information or Medical Rationale for	Request:				
Described Dresident Dressiber Dress		Data	of Dogwood		
Requesting Provider: ☐ Prescriber ☐ Phar *Requestor Name (print):	Пасу		of Request: uestor Signa		
Requestor Name (print).		Req	destor Signi	ature.	
*On behalf of the Prescriber or Pharmacy Provider, I certify that the the medication requested above. I understand the desig					
CONTINUE TO PAGE 2 ONLY IF REQUESTING ANY OPIOID					
CONTINUE TO PAGE 5 ONLY IF REQUESTING HEPATITIS C DAA THERAPY OR CONTINUE TO PAGE 6 IF					
REQUESTING SYNAGIS®					

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When requesting ANY OPIOID, provide the following additional information and most recent chart/progress/clinic note: \*\*For members receiving hospice/palliative/end-of-life care or having a diagnosis of active cancer, this page does not need to be completed.\*\* PLEASE NOTE: ALL OPIOID PA REQUESTS MUST BE COMPLETED BY THE PRESCRIBER ONLY INITIAL TREATMENT REQUESTS ONLY (if request is for continuation therapy skip to question 35) Additional Diagnosis (if not stated above): ICD-10 Code: 1. Prescriber has obtained and reviewed the KASPER report for the past 12 months? □Yes □No 2. Urine drug screen (UDS) has been completed within the past 30 days? Documentation (e.g., lab result or progress note) required ☐ Yes ☐ No ☐ Not Applicable (member is in a long-term care (LTC) facility or will not exceed 45 days of opioid therapy) Please indicate if the patient has tried or is using any of the following non-opioid therapies: ☐ Exercise therapy ☐ Cognitive behavioral therapy ☐ Nonsteroidal anti-inflammatory drugs (NSAIDs) or Acetaminophen (APAP) Please indicate if the patient has any of the following baseline risk factors: Respiratory depression (clinically significant) ☐ Acute or severe bronchial asthma ☐ Hypercarbia (clinically significant) ☐ Known or suspected GI obstruction If any of the above are true, does the prescriber attest that benefits of opioid use outweigh the risks?  $\Box$  Yes  $\Box$  No **EXAMPLE: ASSESSING PAIN & FUNCTION USING PEG SCALE PEG score** = average 3 individual question scores **Q1:** What number from 0-10 best describes your **pain** in the past week? 0 = "no pain", 10 = "worst you can imagine" 10 0 1 2 3 Q2: What number from 0 – 10 describes how, during the past week, pain has interfered with your enjoyment of life? 0 = "not at all", 10 = "complete interference" 2 3 5 10 0 1 6 Q3: What number from 0-10 describes how, during the past week, pain has interfered with your general activity? 0 = "not at all", 10 = "complete interference" 2 10 Does the patient meet ONE of the following criteria? 6. The patient is receiving hospice, palliative, or end-of-life care □Yes □No The patient has a diagnosis of active cancer □Yes ΠNo The patient has a diagnosis of sickle cell anemia □Yes □No Does the patient have a diagnosis of severe pain requiring daily, around-the-clock, long-term pain management? ☐Yes ☐No If 'Yes', proceed to 7a, if 'No' proceed to 8 The patient's pain lasts: > 3 consecutive months  $\Box$ Yes  $\Box$ No, or > 6 consecutive months  $\Box$ Yes  $\Box$ No The patient had a trial and failure within the past 90 days of 1 non-opioid analgesic (i.e., NSAIDs, APAP) at maximum tolerated doses without pain relief and/or functional improvement ☐Yes ☐No The patient had a trial and failure within the past 90 days of at least 1 short acting opioid analgesic at maximum tolerated doses without adequate relief of pain □Yes □No 8. Does the patient have a diagnosis of diabetic peripheral neuropathy? 

Yes 

No If 'Yes' proceed to 8a, if 'No' proceed to 9 The patient had a trial and failure of ONE serotonin-norepinephrine reuptake inhibitor (SNRI; such as duloxetine) □No The patient had a trial and failure of ONE tricyclic antidepressant (TCA; such as amitriptyline) ☐Yes ☐No Does the patient have a diagnosis of neonatal abstinence syndrome (NAS) and meet the following criteria? The patient is being discharged from the hospital on a methadone taper □Yes □No 10. The prescriber has proof of consultation with a pain management specialist  $\square$ Yes  $\square$ No OR specialist in an appropriate discipline (e.g., orthopedist, neurologist, spine specialist, etc.) for evaluation of the source of pain and/or treatment of any underlying conditions  $\Box$ Yes 11. The patient does NOT have a history of drug or alcohol abuse/dependence or addiction (drug and alcohol toxicology screen results dated within the past month must be submitted with the PA request) 12. The patient is NOT using more than 1 long-acting opioid and 1 short-acting opioid at a time ☐ Yes ☐ No 13. The patient has ONE of the following headache disorders: Muscular headache, Tension-type headache, or Migraine  $\Box$ Yes 14. For a high strength (e.g., hydromorphone 8mg) or concentrated dosage form (e.g., morphine sulfate 20 mg/mL, oxycodone 20 mg/mL), please submit a rationale as to why lower strength or less-concentrated products cannot be used \_ 15. Is the patient opioid naive (defined as  $\leq$  14 days of opioid use in the past 90 days)? □Yes □No If 'Yes', proceed to 15a, if 'No' proceed to 16 a. The patient is using only 1 short-acting opioid at a time □Yes  $\square$ No Prescribed by a treating physician within 14 days of ONE of the following: major surgery, any operative or invasive procedure or a delivery, significant trauma, being any acute blunt, blast, or penetrating bodily injury that has a risk of death, physical disability, or impairment If treatment with opioids should extend beyond 14 days please provide clinical justification 16. Is Long-term (> 3 months) pain management expected or indicated □Yes □No 17. For non-preferred long acting opioids: Does the patient have a > 1 month trial and therapeutic failure, allergy, contraindication (including potential drug-drug interactions with other medications) or intolerance to TWO preferred agents \quad Yes \quad \quad No, if 'Yes' please see guestion 22.

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18. For non-preferred <b>short acting</b> opioids: The patient had at least a 1-week trial and therapeutic failure, allergy, contraindication (including potential drug-drug interactions with other medications) or intolerance to TWO preferred agents	כ
Female Patients of Child-bearing Age Only:	_
23. Has the patient been counseled on the risk of becoming pregnant while on this medication and the risk of neonatal abstinence syndrome?  ☐ Yes ☐ No	
Naloxone Attestation:	_
24. Are any of the following true?	
a. Patient UDS is positive for illicit or unexpected substances	
b. Morphine milligram equivalent (MME) is over 90 MME per day	
c. Opioid(s) is/are prescribed concurrently with benzodiazepines $\Box$ Yes (clinical justification required) $\Box$ No	
If yes, prescriber attests that a naloxone prescription and associated counseling on its use, was or will be <b>given</b> to the patient: $\Box$ Yes $\Box$ No	
25. Are any of the following true?	
a. Opioid(s) is/are concurrently prescribed with a skeletal muscle relaxant   Yes   No	
b. Opioid(s) is/are concurrently prescribed with a sedative hypnotic   Yes   No	
c. Opioid(s) is/are concurrently prescribed with gabapentin or pregabalin	
d. Patient has a history of opioid or other controlled substance overdose   Yes  No	
e. Patient has a history of substance use disorder (SUD)	
If yes, prescriber attests that a naloxone prescription and associated counseling on its use was, or will be, <b>offered</b> to the patient:	
zo. For non-preferred agents. Please provide clinical rationale to constitute the use of the requested formulation	
Requests over 90 or 200 MME per day:	
27. For requests over 90 MME: Prescriber is, or has proof of consultation with, a Pain Management Specialist OR a specialist in an appropriate discipline	
(e.g., orthopedist, neurologist, spine specialist, etc.) for evaluation of the source of pain and/or treatment of any underlying conditions.  ☐ Yes ☐ No	
28. For requests over 200 MME: The prescriber is, or has proof of consultation with, a Pain Management Specialist □Yes □No 29. Clinical justification for exceeding 90 or 200 MME per day	
30. Prescriber attests that a naloxone prescription and associated counseling on its use was or will be offered to the member	
31. For requests over 200 MME: prescriber submitted documentation (e.g., progress notes) showing attempts and/or plans to taper below 200 MME/day	
as well as other non-opioid components (e.g., NSAIDs, physical therapy, etc.) of the treatment plan	
Concomitant use of Opioids and Benzodiazepines:	
32. Has the member and/or caregiver(s) been counseled about the increased risks of slowed or difficult breathing and/or excessive sedation, and the	
associated signs and symptoms?   Yes  No	
33. The prescriber attests that a naloxone prescription and associated counseling on its use was or will be given to the member	
34. Clinical justification for the concurrent use of benzodiazepine(s) and opioid(s)	

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REAUTHORIZATION (REFILL) REQUESTS ONLY (with current plan) PLEASE NOTE: ALL OPIOID PA REQUESTS MUST BE COMPLETED BY THE PRESCRIBER ONLY			
35.	Does the patient meet ONE of the following criteria?		
	a. The patient is receiving hospice, palliative, or end-of-life care $\square$ Yes $\square$ No		
	b. The patient has a diagnosis of active cancer ☐Yes ☐No		
	c. The patient has a diagnosis of sickle cell anemia ☐Yes ☐No		
36.	Prescriber has obtained and reviewed the KASPER report within the past 3 months?		
	Urine drug screen (UDS) results: ☐ Positive ☐ Negative Date:		
	Prescriber has assessed risk (check box) and documents (e.g., lab result, progress note) a urine drug screen (UDS) within the listed timeframe:		
	□Low Risk (12 months) □ Moderate Risk (6 Months) □ High Risk (3 Months) □ Not Applicable (member is in a long-term care facility)		
39.	If patient UDS is positive for illicit or unexpected substances:		
	a. Please provide explanation		
	b. Will naloxone prescription and counseling be provided  Yes  No		
40.	Prescriber has reassessed pain and function. Provide PEG scoreor clinical documentation (e.g., progress note):		
	The patient has demonstrated a 30% improvement from baseline to continue current dose  \[ \subseteq \text{Yes} \] \[ \subseteq \text{No} \]		
	OR includes the rationale for continued opioid therapy at the current dose		
42.	Has the patient required use of opioid rescue medication (e.g., naloxone), been hospitalized, or otherwise treated for opioid or other controlled		
	substance overdose in the past 6 months?   Yes 'please provide plan for preventing future overdose  No		
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F	tala Datianta of Child haaring too Out.		
	nale Patients of Child-bearing Age Only:		
43.	Does the PRESCRIBER attest that the patient has been counseled on the risk of becoming pregnant while on this medication and the risk of neonatal		
	abstinence syndrome?		
Ren	uests over 90 or 200 MME per day:		
_	For requests over 90 MME: Prescriber is, or has proof of consultation with, a Pain Management Specialist OR a specialist in an appropriate discipline		
	(e.g., orthopedist, neurologist, spine specialist, etc.) for evaluation of the source of pain and/or treatment of any underlying conditions.		
	Yes No		
45			
	For requests over 200 MME: The prescriber is, or has proof of consultation with, a Pain Management Specialist   Yes  No		
	Clinical justification for exceeding 90 or 200 MME per day		
	Prescriber attests that a naloxone prescription and associated counseling on its use was or will be offered to the member		
48.	For requests over 200 MME: prescriber submitted documentation (e.g., progress notes) showing attempts and/or plans to taper below 200 MME/day		
	as well as other non-opioid components (e.g., NSAIDs, physical therapy, etc.) of the treatment plan		
	ncomitant use of Opioids and Benzodiazepines:		
49.	Does the PRESCRIBER attest that the member and/or caregiver(s) been counseled about the increased risks of slowed or difficult breathing and/or		
	excessive sedation, and the associated signs and symptoms? $\Box$ Yes $\Box$ No		
50.	The prescriber attests that a naloxone prescription and associated counseling on its use was or will be given to the member		
51.	Clinical justification for the concurrent use of benzodiazepine(s) and opioid(s)		
۸۸۰	ditional Clinical Information or Medical Rationale for Request (please attach additional pages/documentation as needed):		
Aut	anional Chinean mormation of Medical Nationale for Nequest (please attach additional pages) documentation as needed).		
CC	ONTINUE TO PAGE 5 <b>ONLY</b> IF REQUESTING <b>HEPATITIS C DAA THERAPY</b> OR CONTINUE TO PAGE 6 IF REQUESTING		
SY	NAGIS®		

When requesting Hepatitis C Direct-Acting Antiviral (DAA) Therapy, provide the following additional information:					
	Date of Hepatitis C diagnosis (or earliest record):	Female Patients of Child-bearing Age Only: Is the patient pregnant or nursing? □Yes □No  If yes, Prescriber attests that the benefits of HCV treatment outweigh potential risks □Yes □No			
Diagnosis Criteria and Simplified Treatment Eligibility	<ul> <li>If 'Yes', is it</li></ul>	details below.			
Repeat DAA Therapy Questions (complete only if requesting repeated DAA therapy)	services, or seeing an addiction specialist as part b. Patient has been evaluated for alcohol and subst	□Yes □No (if no justification must be provided: □Yes □No □No (proceed to 5) □Yes □No □Yes □No □Yes □No □Yes □No □Yes □No □Xerce abuse counseling □Yes □No □Xerce abuse using a validated screening tool □Yes □No □Xerce abuse using a validated screening tool □Yes □No □Xerce abuse using a validated screening tool □Yes □No □Xerce abuse using a validated screening tool □Yes □No □Xerce abuse using a validated screening tool □Yes □No □Xerce abuse □Xerce			

Not Syn	nen requesting Synagis®, provide the following additional information:  te: Therapy may begin November 1 with last date of therapy no later than March 31 (end of RSV season).  tagis is available in 50mg and 100mg vials. Always coordinate dosing appropriately to reduce waste.  requests may be accepted beginning October 1 (for a November 1 effective date).
1. 2.	Patient's gestational age at birth:weeksdays  Does the patient have Chronic Lung Disease of Prematurity (formerly called bronchopulmonary dysplasia)? Yes (proceed to 2a)No (proceed to 3)  a. Did the patient receive oxygen immediately following birth?Yes (proceed to 2b)No (proceed to 3)  b. Please indicate the % oxygen received:
3.	Does the patient have a diagnosis of Cystic Fibrosis?
4.	Please indicate if the patient has any of the following:  Anatomic Pulmonary Abnormality  Neuromuscular Disorder  Congenital anomaly that impairs the ability to clear secretions  Specify:  Specify:  Specify:
5.	Please indicate if the patient has any of the following:  HIV  Cancer, receiving chemotherapy  Organ transplant receiving immunosuppressant therapy or hematopoietic stem cell transplant  Other medical condition that is severely immunocompromising. Specify:
6.	Has this patient received a heart transplant? □Yes (Date:) □No
7.	Does patient have hemodynamically significant congenital heart disease?
8.	□Other:
9.	Please list any pharmaceutical therapies for cardiovascular disease and the most recent date administered:  Cardiovascular medication(s):
10.	If this is a request for a sixth dose of Synagis® during the RSV season, has the patient had an ECMO or cardiac bypass during the RSV season?  □Yes (Date:) □No