# **FAX or EMAIL**

# **PATIENT STICKER**

**SUBJECT:** COVID-19 Monoclonal Antibody Infusion Order **TO:** Comanche County Memorial Hospital Infusion Services

**FAX:** (580) 585-5472

ADDRESS: 3126 NW Arlington Blvd Lawton, OK 73505

EMAIL: infusion@ccmhhealth.com

PHONE: (580) 355-8699 Option 1, Ext 4756 or Ext. 6194 for Scheduling

# FORMS MUST BE COMPLETE (NO BLANKS) AND SIGNED BY THE PROVIDER FOR THE PATIENT TO BE CONSIDERED FOR Monoclonal Antibody Infusion

	<u>P</u>	LEASE PRINT		
DATE:	VACC	CINATION STA	TUS:	
PATIENT NAME:				DOB:
PHONE:	HEIGHT (inches):	<u>INCHES</u>	WEIGHT:	KG (at least 3kg)
ALLERGIES:				
DIAGNOSIS CODE:		NOSIS NAME (	REQUIRED)	
PROVIDER NAME (PRINT L	,			
PROVIDER OFFICE PHONE			OFFICE FA	X #
CONTACT PERSON AT PRO	OVIDER OFFICE:			
FDA approval has been expanded	positive results of direct SARS-	lesivir) to include its CoV-2 viral testing	s use in adults and pe , and who are not hos	idelines ediatric patients (28 days of age and older epitalized and have mild-to-moderate
•	et available at: <u>HIGHLIGH</u>		RIBING INFORM	MATION These highlights do not
include all the information ne (01212022) 	eded to use VEKLURY Sa	alely and ellec o	x <u>EUA 046 Vekiu</u>	ry (remdesivir) FS for HCPs
			•	drug interactions) and prescrib Medication safety consideration
	d is contraindicated then Covid-19 otpt therap Covid-19 Pediatric otpt the	peutics decision	on guide Dec 22	.PNG
<u>s</u>	SARS-CoV-2 Active infec	ction Remdesi	vir CRITERIA FO	DR USE
Symptom Onset Date: _		Positive	COVID-19 test d	ate
	Patients must m	neet <b>ALL</b> criteri	<mark>a to be eligible</mark>	
at least 28 Days of age a	and weighing at least 3 kg	J		
COVID-19 positive by Po	CR or Antigen testing			

#### Within 7 days of symptom onset

Meets all of the following requirements:

- at high risk for progressing to severe COVID-19 and/or hospitalization
- is NOT hospitalized,

### High risk - defined as meeting one or more of the following criteria (select all that apply):

Body Mass Index (BMI > 25) Cardiovascular disease Chronic Kidney Disease

Hypertension Diabetes COPD/other chronic respiratory disease Immunosuppressive Disease

Pregnancy Medical related technology dependence e.g., gastrostomy)

Receiving immunosuppressive treatment Sickle cell disease (e.g., gastrostomy)

Age ≥ 65 years

Neurodevelopmental disorders or other conditions that confer medical complexity(e.g., genetic, or metabolic syndrome)

#### Patient or caregiver received a copy of the patient antiviral fact sheet

- Remdesivir fact sheet: 1 PATIENT INFORMATION VEKLURY® (VEK-lur-ee) (remdesivir) for injection VEKLURY® (VEK-lur-ee) (remdesivir) injection What is VE
- Patient was informed of risks and benefits of therapy, availability of alternatives and that the drug is FDA approved.
- Patients understand they have the option to accept or refuse treatments and, understanding the risks, benefits and alternatives, have agreed to accept treatment.

#### **SARS-CoV-2 Remdesivir DOSING**

# Remdesivir (Veklury)

# Age 12 yo or great and at least 40 kg:

remdesivir 200 mg IV in NS 40mL on 
Day 1 followed by remdesivir 100mg IV in NS 60mL on 
Day 2 & Day 3

Infuse over 30-120 minutes

## Age 28 days to less than 12 years, weighing at least 3 kg to less than 40 kg:

remdesivir 5 mg/kg IV on Day 1 followed by 2.5 mg/kg IV on Day 2 & Day 3

Infuse over 30-120 minutes

If CKD is present or an eGFR of <30 is suspected, please obtain renal panel prior to infusion

#### **Post-Infusion:**

- Flush administration set with 0.9% sodium chloride to deliver residual volume.
- Leave IV in place for observation period; remove prior to discharge.
- Monitor patients for hypersensitivity reactions for a period of 60 minutes following infusion.
- Record vital signs immediately following infusion and prior to discharge.
- Provide patients with discharge instructions. Send record of treatment to prescriber at fax number as appropriate.
- If signs and symptoms of a clinically significant hypersensitivity reaction or anaphylaxis occur, immediately
  discontinue administration and initiate appropriate medications and/or supportive therapy (see ADVERSE
  REACTIONS below)

## **ADVERSE REACTIONS**

MINOR REACTIONS	SEVERE REACTIONS		
(e.g. nausea, itching, joint pain, rash)	(e.g. bronchospasm, loss of airway, fainting, severe flushing)		
STOP infusion	CALL A CODE OR RAPID RESPONSE		
diphenhydrAMINE 50 mg IV Push Once	STOP infusion		
famotidine 20 mg IV Push Once	EPINEPHrine 0.3 mg/0.3 mL SubCutaneous Once		
dexaMETHasone 10 mg IV Push Once	Oxygen PRN		
Notify Physician	Notify Physician		