



المجلس الصحي السعودي Saudi Health Council

Saudi MoH Protocol for Patients Suspected of/Confirmed with COVID-19

Supportive care and antiviral treatment of suspected or confirmed COVID-19 infection

(Version 2.9) May 19th, 2021



رقم الاعتماد: 1-9-1-1442 التاريخ: 11-10-1442ه



Supportive care and antiviral treatment of suspected or confirmed COVID-19 infection

(Version 2.9) May 19th, 2021

<u>Disclaimer</u>: This is a living guidance that is subject to change as more evidence accumulates. It will be updated regularly and whenever needed. The guidance should be used to assist healthcare practitioners select the best available pharmacotherapy for COVID-19 infection according the best available and current evidence and is not intended to replace clinical judgement but rather to complement it. The evidence is inconclusive regarding the efficacy of most medications for covid-19. It is important to explain this to patient and family and obtain informed consent for use of these medications for unapproved indications. Convalescent plasma transfusion should only be used according to an approved study protocol

COVID-19 Testing*	Category	Supportive Care	Pharmacotherapy	Precautions
Suspicious Cases (follow case definition published in Saudi CDC guidelines)	Mild to Moderate: Symptoms with no shortness of breath	 Treat symptoms If no hospital admission required, need to follow instructions and recommendations published by Saudi CDC <u>https://covid19.cdc.gov.sa/pr</u> ofessionals-health-workers/ 	 Not required Do not stop ACEI/ARBs in patients with hypertension, post-MI, or heart failure 	 Paracetamol (acetaminophen) is the prefered agent for pain/fever see below table <i>"Medication Related Information"</i> Labs and work-up: CBC, Urea/Electrolytes, Creatinine, CRP, LFTs, Chest X-ray, COVID-19 PCR tests
	Mild to Moderate: Symptoms with no shortness of breath in high-risk patients ^S Mild to Moderate: Symptoms with shortness of breath in high-risk patients ^S	 Treat symptoms If hospital admission is not required, follow instructions and recommendations published by Saudi CDC <u>https://covid19.cdc.gov.sa/pr</u> <u>ofessionals-health-workers/</u> Consult Infectious Disease Specialist 	 Case shall be discussed with infectious disease specialist, to initiate empirical antiviral therapy, while awaiting PCR result. Do not stop ACEI/ARBs in patients with hypertension, post-MI, heart failure <u>If decision is to treat empirically, follow the treatment option under confirmed by PCR</u> 	
PCR Confirmed Cases	Asymptomatic	 Follow instructions and recommendations published by Saudi CDC <u>https://covid19.cdc.gov.sa/pr</u> ofessionals-health-workers/ 	- Not required	

وزارة الصحة Ministry of Health Supportive care and antiviral treatment of suspected or confirmed COVID-19 infection

(Version 2.9) May 19th, 2021

COVID-19 Testing*	Category	Supportive Care	Pharmacotherapy	Precautions
PCR Confirmed Cases	Mild to Moderate: Symptoms (no O ₂ requirements/no evidence of pneumonia but with other symptoms of covid- 19 e.g., fever)	 Treat symptoms Follow instructions and recommendations published by Saudi CDC https://covid19.cdc.gov.sa/pr ofessionals-health-workers/ 	 In case of new onset cough and fever or anosmia, or both) within 7 days Consider inhaled budesonide (Pulmicort®) <u>Adult Dosing:</u> 800 µg per actuation (two inhalations) twice a day until symptom resolution Consider starting any of the following according to clinical evaluation and treating consultant's discretion: Consider Favipiravir <u>Adult Dosing</u>: 1800 mg/dose twice a day on the first day; followed by 800 mg/dose twice a day for 7-10 days <u>Pediatric Dosing</u>: 10-15 kg: Loading Dose: One tablet PO BID for One day (maximum 400 mg/day). Maintenance from Day 2: Half tablet (100 mg) PO BID (maximum 200 mg/day) 16-21 kg: Loading Dose: Two tablets PO BID one day (maximum 400 mg/day). Maintenance fromDay2: One Tablet PO BID for One day (maximum 1200 mg/day) 22-35 kg: Loading Dose: Three Tablets PO BID for One day (maximum 1200 mg/day). Maintenance from Day2: One tablet PO BID for One day (maximum 1200 mg/day). 36-45 kg: Loading Dose: Four tablets PO BID for One day (maximum 100 mg/day) 36-45 kg: Loading Dose: Five tablets PO BID for One day (maximum 1600 mg/day). Maintenance from Day2: Two tablets PO BID (maximum 200 mg/day) 46-55 kg: Loading Dose: Five tablets PO BID for One day (maximum 1000 mg/day). 46-55 kg: Loading Dose: Five tablets PO BID for One day (maximum 2000 mg/day). Maintenance from Day2: Two Tablets qAM, Three Tablets qPM (maximum 1000 mg/day) For >55 kg: Can use adult dosing if age ≥16 years, if age <16years use dosing of 46-55 kg range 	 Inhaled budesonide (Pulmicort®) see below table "Medication Related Information" Bronchospasm, oral candidiasis, and vasculitis Favipiravir (non-formulary and non-SFDA registered) see below table "Medication Related Information" Contraindicated in pregnancy Anticoagulation see below "Thromboprophylaxis"
	 Severe: Clinical signs of pneumonia (fever, cough, dyspnea, fast breathing) and one of the following: Respiratory rate >30/min (adults); ≥40/min (children < 5 years) Blood oxygen saturation <90% on room air Severe respiratory distress 	 Treat symptoms Follow instructions and recommendations published by Saudi CDC <u>https://covid19.cdc.gov.sa/pr</u><u>ofessionals-health-workers/</u> ICU admission, decision by ICU treating team Antibiotics and antifungals according to local antibiogram and institutional pneumonia management guidelines/ pathways. 	Consider starting any of the following according to clinical evaluation and treating consultant's discretion: - Consider Favipiravir Adult Dosing: 1800 mg/dose twice a day on the first day; followed by 800 mg/dose twice a day for 7-10 days. Pediatric Dosing: 10-15 kg: Loading Dose: One tablet PO BID for One day (maximum 400 mg/day). Maintenance from Day 2: Half tablet (100 mg) PO BID (maximum 200 mg/day). 16-21 kg: Loading Dose: Two tablets PO BID One day (maximum 800 mg/day). Maintenance fromDay2: One Tablet PO BID (maximum 400 mg/day) 22-35 kg: Loading Dose: Three Tablets PO BID for One day (maximum 1200 mg/day). Maintenance from Day2: One tablet PO TID (maximum 600 mg/day) 36-45 kg: Loading Dose: Four tablets PO BID for One day (maximum 1600mg/day). Maintenance from Day2: Two tablets PO BID (maximum 800 mg/day) 	 Remdesivir (non-formulary and non-SFDA registered) see below table "Medication Related Information" Exclusion criteria evidence of multiorgan failure, need of inotropes, Creatinine clearance < 30 ml/min, dialysis/hemofiltration, transaminases > 5X ULN, or concomitant use of lopinavir/ritonavir Favipiravir (non-formulary and non-SFDA registered) (see precautions above) Systemic Dexamethasone see below table "Medication Related Information"

وزارة الصحة Supportive care and antiviral treatment of suspected or confirmed COVID-19 infection

(Version 2.9) May 19th, 2021

COVID-19 Testing*	Category	Supportive Care	Pharmacotherapy	Precautions
PCR Confirmed Cases			 46-55 kg: Loading Dose: Five tablets PO BID for One day (maximum 2000 mg/day). Maintenance from Day2: Two Tablets qAM, Three Tablets qPM (maximum 1000 mg/day) For >55 kg: Can use adult dosing if age ≥16 years, if age <16years use dosing of 46-55 kg range OR Consider Remdesivir Adult Dosing: 200 mg loading dose (IV, within 30 min), followed by 100 mg once daily for 5 to 10 days Pediatric dosing	 Cardiovascular disease: Use with caution in patients with heart failure and/or hypertension; use has been associated with fluid retention, electrolyte disturbances, and hypertension. Use with caution following acute myocardial infarction; corticosteroids have been associated with myocardial rupture. Diabetes: Use corticosteroids with caution in patients with diabetes mellitus; may alter glucose production/regulation leading to hyperglycemia. Gastrointestinal disease: Use with caution in patients with Gl diseases (diverticulitis, fresh intestinal anastomoses, active or latent peptic ulcer, ulcerative colitis, abscess or other pyogenic infection) due to perforation risk. Myasthenia gravis: Use with caution in patients with myasthenia gravis; exacerbation of symptoms has occurred especially during initial treatment with corticosteroids. Seizure disorders: Use corticosteroids with caution in patients with a history of seizure disorder; seizures have been reported with adrenal crisis. Labs and workup: Hemoglobin, occult blood loss, blood pressure, serum potassium, glucose, weight, and height in children; HPA axis suppression

وزارة الصحة Ministry of Health Supportive care and antiviral treatment of suspected or confirmed COVID-19 infection

(Version 2.9) May 19th, 2021

COVID-19 Testing*	Category	Supportive Care	Pharmacotherapy	Precautions
PCR Confirmed Cases	 Critical: Symptoms of the following: ARDS Respiratory failure requiring ventilation Sepsis Septic Shock Criteria for using tocilizumab: Within 24 hours of ICU admission for MV, NIV, or HFNC oxygen Patients who are exhibiting rapidly increasing oxygen needs while on dexamethasone and have a C-reactive protein level ≥75 mg/L (715 nmol/L). 	 Treat symptoms Follow instructions and recommendations published by Saudi CDC https://covid19.cdc.gov.sa/pr ofessionals-health-workers/ ICU admission and management by ICU treating team Antibiotics and antifungals according to local antibiogram and institutional pneumonia management guidelines/ pathways. 	 Consider starting any of the following according to clinical evaluation and treating consultant's discretion: Consider Remdesivir Adult Dosing: 200 mg loading dose (IV, within 30 min), followed by 100 mg once daily for 5 to 10 days Pediatric dosing <u< th=""><th>Remdesivir (non-formulary and non-SFDA registered) (see precautions above) Systemic Dexamethasone: (see precautions above) Baricitinib see below table "Medication Related Information" Patients treated with baricitinib are at risk for developing serious infections, malignancies, and thrombosis Tocilizumab see below table "Medication Related Information" Should perform IL6 and other inflammatory markers testing prior to start (CRP, Ferritin, D-dimer) Watch for infusion reaction Anticoagulation see below "Thromboprophylaxis"</th></u<>	Remdesivir (non-formulary and non-SFDA registered) (see precautions above) Systemic Dexamethasone: (see precautions above) Baricitinib see below table "Medication Related Information" Patients treated with baricitinib are at risk for developing serious infections, malignancies, and thrombosis Tocilizumab see below table "Medication Related Information" Should perform IL6 and other inflammatory markers testing prior to start (CRP, Ferritin, D-dimer) Watch for infusion reaction Anticoagulation see below "Thromboprophylaxis"

وزارة الصحة Supportive care and antiviral treatment of suspected or confirmed COVID-19 infection

COVID-19 Testing*	Category	Supportive Care	Pharmacotherapy	Precautions
PCR Confirmed Cases			 <u>Preterm infants with a corrected gestation age of <40 weeks:</u> 0.5 mg/kg every 12 hours OR Methylprednisolone sodium succinate (IV): 0.8 mg/kg once daily (max: 32 mg) If rapid respiratory decompensation due to COVID-19, consider tocilizumab with dexamethasone <u>Adult Dosing</u>: Single dose of tocilizumab 8 mg/kg of actual body weight (maximum 800 mg) by IV infusion in combination with dexamethasone 6 mg daily for up to 10 days <u>Pediatric Dosing (<18 years)</u>:	
 Serum IL- Ferritin > Tocilizumab is re off-label. Remde 	6 ≥3x upper normal limit 00 ug/L (or surrogate) with doublin gistered medications in Saudi Ara esivir and favipiravir are not curre	bia and available in MoH formulary in the second seco	 Ferritin >600 ug/L at presentation and LDH >250 Elevated D-dimer (>1 mcg/mL) for other indications but have not shown proven efficacy in many randomized clinical trials as of 	
patient and trea Thrombopropl Recommendatio – All admitt – Laborator – Baseline o – Patients o – Warfarin,	ting team. Refer to the MoH COVI ylaxis: ed patients should be evaluated up y evaluation and monitoring: Base r surveillance imaging are not rec n chronic VTE prophylaxis should DOAC and antiplatelet medicatior	D-19 guidance in pregnancy oon admission, and daily thereafter f line CBC, fibrinogen, PT, aPTT, D-di ommended in the absence of clinica continue as planned before. as are not recommended to be used	al symptoms of VTE	
hospitaliz - Thrombop <u>When to consult</u> - Heparin ir - Platelets - Unexplain - Inherited - Inherited - Previously	ed medically ill patients rophylaxis should continue until th	e time of discharge at least. Contini ombasthenia)	uation of anticoagulation is subject to assessment of VTE risk by the treating medical team.	



وزارة الصحة Ministry of Health Supportive care and antiviral treatment of suspected or confirmed COVID-19 infection

COVID-19				
Testing*	Category	Supportive Care	Pharmacotherapy	Precautions
Adults:				
– Therapeutio	c doses should not be offered b	ecause of the risk of bleeding		
			idered in ALL patients (including non-critically ill) who require hospital admission for COVID-19 ir	nfection, in the absence of any
		telet count less than 25 x 109/L; mon	nitoring is advised in severe renal impairment; abnormal PT or APTT is not a contraindication)	
	prophylaxis doses:			
	ocutaneously once daily			
	MI > 40 kg/m ² : 40 mg subcutar			
	y: 40 mg subcutaneously once	daily		
Renal imp				
	Cl > 30 mL/minute: no adjustme			
	Cl < 30 mL/minute: 30 mg subci		d he frequently mentared as accumulation may accur with repeated doors	
			d be frequently monitored, as accumulation may occur with repeated doses. rotocol for alternative anticoagulation.	
Pediatrics:	in neparin-induced thrombocyte	ppenna (nii), please follow mon nii p		
	prophylaxis doses:			
		subcutaneously every 12 hours		
		cents: 0.5 mg/kg/dose subcutaneous	ly every 12 hours	
		recommendations (use with caution ar		
			els frequently monitored, as accumulation may occur with repeated doses.	
	ysis: Not dialyzable and supple			
Enoxaparin monit				
 Routine ant 	i-Xa levels are not recommende	ed.		
 If an anti-Xa 	a level is deemed necessary, it :	should be drawn 4-6 hours after enoxa	aparin administration with an anti-Xa goal of 0.2- 0.4 units/mL for prophylaxis and 0.5-1 Units/ml	for therapeutic dose.
 Consider re 	-checking anti-Xa if the patient	experiences active bleeding or has e	vidence of renal dysfunction while on enoxaparin therapy	
	to Anticoagulation (Bleeding R			
			bleeding /hematoma, congenital bleeding disorder	
		>44 seconds, fibrinogen <100 g/dL, o	or platelet <50,000/microliter	
Consider Avoiding				
			nt is likely to require an invasive procedure within 24 hours of starting enoxaparin, Neurosurgical p	procedure, Pelvic fracture within past 48
		< 5-7 days ago), Uncontrolled hyperte	nsion	
Criteria for Manag	mmatory Syndrome in Childre	<u>en (MIS-C)</u>		
		or (>28.0°C for >24 hours or report of	f subjective fever lasting ≥24 hours), laboratory evidence of inflammation (Including, but not limite	ad to one or more of the following: on
			vated neutrophils; reduced lymphocytes; and low albumin), and evidence of clinically severe illness	
		spiratory, hematologic, gastrointesti		
	ive plausible diagnoses			
		infection by RT-PCR, serology, or ant	tigen test; or COVID-19 exposure within the 4 weeks prior to the onset of symptoms	
		· · · · · · · · · · · · · · · · · · ·	· · · · · · · · · · · · · · · · · · ·	
i				

وزارة الصحة Supportive care and antiviral treatment of suspected or confirmed COVID-19 infection Ministry of Health

(Version 2.9) May 19th, 2021

COVID-19 Testing*	Category	Supportive Care	Supportive Care Pharmacotherapy Precautions						
immunomodulatory – Supportive (arrhythmia) – Thrombopro – Antiviral the	y therapy should also be considered f	or antiviral therapy if they are re signs and symptoms should be or other potentially life-threat	not already rece e admitted to the	hospital. Admission to a pediatric intensive care unit is appropriate for children					
I	Immunomodulator	Dosing		Safety monitoring					
"Medication Relat MIS-C with o disease or s OR	rednisolone see below table ted Information" or without features of Kawasaki igns of myocardial dysfunction ritical COVID-19 with evidence of	 IVIG 2 g/kg + methylpredn 1 mg/kg every 12 hours (n mg for 12 hours) for 5 days IVIG 2 g/kg + methylpredn of 15 to 30 mg/kg/d for 3 	maximum of 30 /s nisolone bolus	 Potential adverse reactions: anaphylaxis, Infusion reaction, hemolysis, transaminitis, aseptic meningitis 					
dilation/and OR	features of shock or coronary artery eurysm ritical COVID-19 with evidence of	 1-2 mg/kg/day divided BID prednisolone, methylprednisolone, methylprednisolone, methylprednisolone, mg/m2 daily (dexametha 	nisolone)	(see precautions above)					

Abbreviations:

ARDS: Acute respiratory distress syndrome, COVID-19: Coronavirus Disease 2019, CBC: Complete Blood Count, CRP: C-Reactive Protein, ECMO: Extracorporeal Membrane Oxygenation, IL6: Interleukin 6, LFT: Liver Function Test, PCR: Polymerase Chain Reaction, ECG: Electrocardiogram, G6PD: Glucose-6-Phosphate Dehydrogenase, ACEI: Angiotensin-converting enzyme inhibitors, ARBs: Angiotensin II receptor blockers, MI: Myocardial infarction, MIS-C: Multisystem Inflammatory Syndrome in Children, CSS: Cytokine Storm Syndrome, mechanical ventilation (MV), noninvasive mechanical ventilation (NIV), high-flow nasal canula (HFNC)

Footnotes:

*Testing for SARS-COV2 virus shall be performed in accordance with published case definition by Saudi CDC guidelines. ³High risk patients have one or more: 1. Elderly (age > 65 years), 2. With underlying end organ dysfunction, 3. Diabetes, 4. History of cardiovascular disease, 5. History of pulmonary disease, 6. Immunocompromised, and/or 7. Pregnancy

وزارة الصحة Ministry of Health Supportive care and antiviral treatment of suspected or confirmed COVID-19 infection

Medication	Contraindication	Major Drug Interactions	Required dose adjustment	Pregnancy and Lactation
Paracetamol (acetaminophen)	 Hypersensitivity to acetaminophen or any component of the formulation Severe hepatic impairment or active liver disease 	 Acetaminophen may increase the levels/effects of: Busulfan; Dasatinib; Imatinib; Local Anesthetics; Mipomersen; Phenylephrine (Systemic); Prilocaine; Sodium Nitrite; SORAfenib; Vitamin K Antagonists The levels/effects of Acetaminophen may be increased by: Alcohol (Ethyl); Dapsone (Topical); Dasatinib; Flucloxacillin; Isoniazid; MetyraPONE; Nitric Oxide; Probenecid; SORAfenib 	 Requires dose adjustment with patient with hepatic impairment <u>See MoH online formulary</u> 	 Oral paracetamol is considered safe in normal therapeutic doses for short-term use as a minor analgesic/antipyretic in pregnancy. Consider Administering IV paracetamol to a pregnant woman only if clearly needed. Carefully assess maternal benefit and fetal risk before administering IV paracetamol during labor and delivery.
Remdesivir	 Safety and efficacy not established 	 Avoid Concomitant Use: There are no known interactions where it is recommended to avoid concomitant use. Increased Effect/Toxicity: There are no known significant interactions involving an increase in effect. Decreased Effect: There are no known significant interactions involving a decrease in effect. 	 No dose adjustment studied 	 Not studied
Favipiravir	 Hematopoietic tissues such as decreased RBC production, and increases in liver function parameters Testis toxicity was also noted Teratogenic 	Acyclovir, Adefovir dipivoxil, Afatinib, Allopurinol, Almotriptan, Alprostadil, Ambrisentan, Aminohippuric acid, Aminophenazone, Amiodarone, Amitriptyline, Amodiaquine, Anastrozole, Antipyrine, Apalutamide, Apixaban, Atorvastatin, Avatrombopag, Avibactam, Azelastine, Baricitinib, Belinostat, Benzyl alcohol, Benzylpenicillin, Betrixaban, Bisoprolol, Bosutinib, Brentuximab vedotin, Brigatinib, Bumetanide, Buprenorphine, Cabazitaxel, Canagliflozin, Captopril, Cefaclor, Cefazolin, Cefdinir, Cefotiam, Ceftibuten, Ceftizoxime, Celecoxib, Cephalexin, Ceritinib, Cerivastatin, Chloroquine, Cholic Acid, Cidofovir, Cimetidine, Cisapride, Citrulline, Clobazam, Clomifene, Cobimetinib, Colchicine, Conjugated estrogens, Copanlisib, Crizotinib, Cyclophosphamide, Cyclosporine, Dabigatran etexilate, Zafirlukast, Zalcitabine, Zidovudine, Zopiclone	 No dose adjustment studied 	- Contraindicated
Tocilizumab	 Known hypersensitivity to tocilizumab or any component of the formulation Active infections 	 Avoid Concomitant Use: Anti-TNF Agents; BCG (Intravesical); Belimumab; Biologic Disease-Modifying Antirheumatic Drugs (DMARDs); Cladribine; Natalizumab; Pimecrolimus; Tacrolimus (Topical); Vaccines (Live) Increased Effect/Toxicity: Anti-TNF Agents; Biologic Disease-Modifying Antirheumatic Drugs (DMARDs); Fingolimod; Leflunomide; Natalizumab; Siponimod; Vaccines (Live) The levels/effects of Tocilizumab may be increased by: Belimumab; Cladribine; Denosumab; Ocrelizumab; Pimecrolimus; Roflumilast; Tacrolimus (Topical); Trastuzumab Tocilizumab may decrease the levels/effects of: BCG (Intravesical); Coccidioides immitis Skin Test; CYP3A4 Substrates (High risk with Inducers); Nivolumab; Pidotimod; Sipuleucel-T; Smallpox and Monkeypox Vaccine (Live); Tertomotide; Vaccines (Inactivated); Vaccines (Live) The levels/effects of Tocilizumab may be decreased by: Echinacea 	 Requires dose adjustment with patient with hepatotoxicity <u>See MoH online formulary</u> 	 Fetal risk cannot be ruled out
Baricitinib	 Hypersensitivity to Baricitinib or any component of formulation 	 Need therapy modification and monitoring:5-Aminosalicylic Acid Derivatives, Chloramphenicol (Ophthalmic), CloZAPine Deferiprone, Denosumab, Echinacea, Fingolimod, Leflunomide, Nitisinone, Nivolumab, Pidotimod, Pretomanid, Probenecid, Promazine, Roflumilast, Sipuleucel-T, and Tertomotide Avoid combination: Vaccines (Live), Talimogene Laherparepvec, Tacrolimus (Topical), Belimumab, Biologic Disease-Modifying Antirheumatic Drugs, Cladribine, Cladribine, Dipyrone, Natalizumab, Pimecrolimus, 	 Requires dose adjustment with patient with renal and liver impairment 	 Not recommended in breastfeeding Information related to pregnancy is limited
Systemic Dexamethasone	 Concomitant use of more than a single dose of dexamethasone with rilpivirine Hypersensitivity to dexamethasone or any component of the product Systemic fungal infection 	 Avoid concomitant use of DexAMETHasone (Systemic) with any of the following: Aldesleukin; BCG (Intravesical); Cladribine; Conivaptan; Desmopressin; Fusidic Acid (Systemic); Idelalisib; Indium 111 Capromab Pendetide; Lapatinib; Lasmiditan; Macimorelin; Mifamurtide; MIFEPRIStone; Natalizumab; Pimecrolimus; Rilpivirine; Simeprevir; Tacrolimus (Topical); Upadacitinib 	 Use cautiously in the elderly at the lowest possible dose <u>See MoH online formulary</u> 	 Pregnant or breastfeeding women, use prednisolone (Oral) or intravenous hydrocortisone instead of dexamethasone.

وزارة الصحة Ministry of Health Supportive care and antiviral treatment of suspected or confirmed COVID-19 infection

Medication Related	Information			
Medication	Contraindication	Major Drug Interactions	Required dose adjustment	Pregnancy and Lactation
Inhaled budesonide (Pulmicort®)	 Hypersensitivity to budesonide Allergenic cross-reactivity for corticosteroids is limited Patients with cirrhosis 	 Diminish the effect of: Aldesleukin and Cosyntropin Enhance the effect/toxicity of: Desmopressin and Loxapine Increase the serum concentration of Budesonide: CYP3A4 Inhibitors Diminish the effect of Budesonide: Tobacco 	 Use cautiously in hepatic impairment See MoH online formulary 	 Present in breast milk.
IVIG	 Hypersensitivity to IVIG or any component of the formula Documentation of allergic cross-reactivity 	- MMR, varicella vaccines	 Use cautiously with Renal impairment due to risk of immune globulin-induced renal dysfunction; the rate of infusion and concentration of solution should be minimized. Discontinue if renal function deteriorates. See MoH online formulary 	
Enoxaparin	 Active major bleeding History of immune-mediated heparin-induced thrombocytopenia within the past 100 days or in presence of circulating antibodies Hypersensitivity to benzyl alcohol (present in multi-dose formulation) - Hypersensitivity to enoxaparin. 	 Avoid combination: Vorapaxar: May enhance the adverse/toxic effect of Anticoagulants. More specifically, this combination is expected to increase the risk of bleeding. Urokinase: May enhance the anticoagulant effect of Anticoagulants. Rivaroxaban: Anticoagulants may enhance the anticoagulant effect of Rivaroxaban Omacetaxine: Anticoagulants may enhance the adverse/toxic effect of Omacetaxine MiFEPRIStone: May enhance the adverse/toxic effect of Anticoagulants. Specifically, the risk of bleeding may be increased Hemin: May enhance the anticoagulant effect of Anticoagulants. Edoxaban: May enhance the anticoagulant effect of Anticoagulants. Dabigatran Etexilate: May enhance the anticoagulant effect of Anticoagulants. Apixaban: May enhance the anticoagulant effect of Anticoagulants. 	 Renal impairment (CrCl 30 to 80 mL/min): No adjustment necessary Renal impairment (CrCl less than 30 mL/min): reduce usual recommended dose by 50%. <u>See MoH online formulary</u> 	 Low molecular weight heparin (LMWH) does not cross the placenta; increased risks of fetal bleeding or teratogenic effects have not been reported (Bates 2012).

Drug Adminis	tration in patients wit	h Swallowing Difficulties				
Drug	Formulation	Remarks				
Favipiravir	Tablets	 Tablets can be crushed and mixed with liquid. 				
Baricitinib	Tablet	 Tablets can be mixed with room temperature water. 				
			Administration via	Dispersion Volume	Container Rinse Volume	
			 Oral dispersion 	10 mL	10 mL	
			- Gastrostomy tube	15 mL	15 mL	
			 Nasogastric tube 	30 mL	15 mL	

وزارة الصحة Ministry of Health Supportive care and antiviral treatment of suspected or confirmed COVID-19 infection

(Version 2.9) May 19th, 2021

References:

- 1. Ministry of Health. Coronavirus Diseases 19 (COVID-19) guidelines. March 2020, version 1.2
- Jin, Y.H., et al., A rapid advice guideline for the diagnosis and treatment of 2019 novel coronavirus (2019-nCoV) infected pneumonia (standard version). Mil Med Res, 2020. 7(1): p. 4. 2.
- Li, J.Y., et al. The epidemic of 2019-novel-coronavirus (2019-nCoV) pneumonia and insights for emerging infectious diseases in the future. Microbes Infect, 2020. 3.
- 4. Living Dong, Shasha Hu, JianiunGao, Discovering drugs to treat coronavirus disease 2019 (COVID-19), Drug Discov & Ther 2020;14(1):58-60
- Zhou F, Yu T, Du R, Fan G, Liu Y, Liu Z, et al. Clinical course and risk factors for mortality of adult inpatients with COVID-19 in Wuhan, China: a retrospective cohort study. Lancet Published Online First: 11 March 5. 2020. doi:10.1016/S0140-6736(20)30566-3
- Interim clinical guidance for patients suspected of/ confirmed with COVID-19 in Belgium. 19 March 2020, version 4. 6.
- American Society of Health-System Pharmacists. Assessment of Evidence for COVID-19-Related Treatments. 03-21-2020. 7.
- NHS Thromboprophylaxis and anticoagulation in COVID-19 infection. Imperial College Healthcare V 0.1 08.04.2020 Covid Treatment Group 8.
- 9. J. Grein, N. Ohmagari, D. Shin, G. Diaz, et al. Compassionate Use of Remdesivir for Patients with Severe Covid-19. The new england journal of medicine. April 10, 2020
- University of Liverpool COVID-19 resources, www.covid19-druginteractions.org updated 3 April 2020 10.
- 11. Thachil, Jecko, et al. "ISTH interim guidance on recognition and management of coagulopathy in COVID-19." Journal of Thrombosis and Haemostasis 18.5 (2020): 1023-1026.
- 12. Tang, Ning, et al. "Abnormal coagulation parameters are associated with poor prognosis in patients with novel coronavirus pneumonia." Journal of Thrombosis and Haemostasis (2020).
- 13. Oiu, Haivan, et al. "Clinical and epidemiological features of 36 children with coronavirus disease 2019 (COVID-19) in Zheijang, China: an observational cohort study." The Lancet Infectious Diseases (2020).
- Bauman, Mary E., et al. "Evaluation of enoxaparin dosing requirements in infants and children." Thrombosis and haemostasis 101.01 (2009): 86-92. 14.
- 15. Lee, Young R., et al. "Stratifying therapeutic enoxaparin dose in morbidly obese patients by BMI class: a retrospective cohort study." Clinical Drug Investigation 40.1 (2020): 33-40.
- 16. Li, Wei, et al. "Chest computed tomography in children with COVID-19 respiratory infection." Pediatric radiology (2020): 1-4.
- 17. Schloemer. Nathan J., et al. "Higher doses of low-molecular-weight heparin (enoxaparin) are needed to achieve target anti-Xa concentrations in critically ill children." Pediatric Critical Care Medicine 15.7 (2014): e294-e299.Uptodate last access May 20, 2020
- 18. Micromedex last access May 20, 2020
- Geoffrey D. Barnes, et al. Thromboembolism and anticoagulant therapy during the COVID-19 pandemic: interim clinical guidance from the anticoagulation forum. Journal of Thrombosis and Thrombolysis. 21 May 19. 2020
- 20. World Health Organization Clinical management of COVID-19: interim guidance, 18 May 2020.
- A Trial of Favipiravir and Hydroxychloroquine combination in Adults Hospitalized with moderate and severe Covid-19 CLINICAL TRIAL PROTOCOL, King Abdullah International Medical research Center, Protocol V1 21. date April 5th, 2020.
- 22. Randomised Evaluation of COVid-19 thERapY (RECOVERY) Trial on dexamethasone, 16 June 2020
- 23. Llitios JF, Leclerc M, Chochois C, et al, High incidence of venous thromboembolic events in anticoagulated severe COVID-19 patients, J Thromb Haemost, 2020.
- 24. Tang N. Bai H. Chen X. Gong J. Li D. Sun Z. Anticoagulant treatment is associated with decreased mortality in severe coronavirus disease 2019 patients with coagulopathy. J Thromb Haemost. 2020;18(5):1094-1099.
- 25. Ludvigsson JF. Systematic review of COVID-19 in children shows milder cases and a better prognosis than adults. Acta Paediatr. 2020
- 26. Loi M, Branchford B, Kim J, Self C, Nuss R. COVID-19 anticoagulation recommendations in children. Pediatr Blood Cancer. 2020.
- 27. American Society of Hematology. https://www.hematology.org/covid-19/covid-19-and-coagulopathy. http://www.hematology.orgcovid-covid-and-coagulopathy.
- 28. Thachil J, Tang N, Gando S, et al. ISTH interim guidance on recognition and management of coagulopathy in COVID-19. J Thromb Haemost. 2020: 18(5): 1023-1026
- Panel on COVID-19 Treatment. COVID-19 Treatment Guidelines. 29.
- Available at https://www.covid19treatmentguidelines.nih.gov/overview/ 30.
- 31. Kache, S., Chisti, M.J., Gumbo, F. et al. COVID-19 PICU guidelines: for high- and limited-resource settings. Pediatr Res (2020)
- World Health Organization, Pocket Book for Hospital Care of Children; Guidelines for the Management of Common Illness with Limited Resources (World Health Organization, Geneva, 2013) 32.
- Food and Drug Administration. FACT SHEET FOR HEALTHCARE PROVIDERS EMERGENCY USE AUTHORIZATION (EUA) OF BARICITINIB. https://www.fda.gov/media. Accessed December 7, 2020 33.
- National Institute of Allergy and Infectious Diseases. Adaptive COVID-19 Treatment Trial 2 (ACTT-2). Clinical Trials.gov Identifier: NCT04401579 34.
- Kalil AC, Patterson TF, Mehta AK, et al. Baricitinib plus Remdesivir for Hospitalized Adults with Covid-19. N Engl J Med. 2020 Dec 11. doi: 10.1056/NEJMoa2031994. Epub ahead of print. PMID: 33306283. 35.
- 36. Alhazzani, Waleed et al. "Surviving Sepsis Campaign: guidelines on the management of critically ill adults with Coronavirus Disease 2019 (COVID-19)," Intensive care medicine vol. 46.5 (2020): 854-887. doi:10.1007/s00134-020-06022-5
- 37. McCullough PA. Favipiravir and the Need for Early Ambulatory Treatment of SARS-CoV-2 Infection (COVID-19), Antimicrob Agents Chemother, 2020 Nov 17:64(12):e02017-20, doi: 10.1128/AAC.02017-20, PMID: 32967849; PMCID: PMC7674042.
- Joshi S, Parkar J, Ansari A, Vora A, Talwar D, Tiwaskar M, Patil S, Barkate H. Role of favipiravir in the treatment of COVID-19. Int J Infect Dis. 2021 Jan; 102:501-508. doi: 10.1016/j.ijid.2020.10.069. Epub 2020 Oct 38. 30. PMID: 33130203; PMCID: PMC7831863



(Version 2.9) May 19th, 2021

39. Ramakrishnan S, Nicolau DV Jr, Langford B, Mahdi M, Jeffers H, Mwasuku C, Krassowska K, Fox R, Binnian I, Glover V, Bright S, Butler C, Cane JL, Halner A, Matthews PC, Donnelly LE, Simpson JL, Baker JR, Fadai NT, Peterson S, Bengtsson T, Barnes PJ, Russell REK, Bafadhel M. Inhaled budesonide in the treatment of early COVID-19 (STOIC): a phase 2, open-label, randomised controlled trial. Lancet Respir Med. 2021 Apr 9:S2213-2600(21)00160-0. doi: 10.1016/S2213-2600(21)00160-0. Epub ahead of print. Erratum in: Lancet Respir Med. 2021 Apr 14;: PMID: 33844996; PMCID: PMC8040526.

Summary of Protocol changes

- Adjustment under Thromboprophylaxis
 - Referring patients with Heparin-induced thrombocytopenia to MoH protocol
 - Continuation of anticoagulation is subject to assessment of VTE risk by the treating medical team.