YNHHS Initial Treatment Algorithm for Hospitalized ADULTS with Non–Severe* COVID-19

Disclaimer: There are no FDA-approved treatments for COVID-19, supportive care is standard of care. Limited treatment data are available & clinical judgment is warranted – Algorithm last updated 5/18/20

Patient with confirmed POSITIVE SARS-CoV-2 by PCR
Assess all patients routinely for clinical trial eligibility (see Appendix 1)
*(If mechanically ventilated or on ECMO, proceed to Severe algorithm)

Oxygen saturation ≤ 94% on room air
(≤ 95% if pregnant)

YES
Continue supportive care
Consider adjunctive treatment

NO
SUPPORTIVE CARE & EVERY 4 HOUR OXYGEN MONITORING

ADJUNCTIVE TREATMENT CONSIDERATIONS
If ≥ 3 Liter O2 requirement
OR ≥ 2 Liter O2 requirement & hs-CRP >70
Tocilizumab x 1 dose
(see Appendix 2 for exclusion criteria)

Remdesivir availability under the EUA is limited. Potential candidates will be identified. Pharmacy will contact primary providers of eligible patients.
Remdesivir has not been FDA approved; remdesivir is authorized by the FDA under and Emergency Use Authorization (EUA)

Consider MICU evaluation if > 4 Liter O2 requirement or hemodynamic instability
(at YNHH see Appendix 4 for suggested triage guidelines)

YNHH: ID consult is not mandatory; consider ID input if immunosuppressed* or clinically decompensating
BH, GH, LMH, or WH: consult ID

COVID-SPECIFIC TESTS
1) Baseline & every 12 hours (for 5 days, then daily thereafter): CRP, D-dimer
2) Baseline & every 12 hours x3: Troponin (continue longer if further testing clinically indicated)
3) Baseline & every 24 hours (for 5 days*): CBC with differential, BMP, LFTs, Ferritin, Procalcitonin, BNP, fibrinogen, PT/PTT, Mg
4) Baseline & ICU transfer: Cytokine panel
5) Baseline and with acute kidney injury (AKI): urinalysis and urine protein/albumin ratio
6) Baseline EKG (see Appendix 3 for QTc recommendations)
7) Repeat Chest X-Ray: if clinical deterioration. (CXR not indicated for discharge or to document clinical improvement)

*May extend longer if clinically indicated
Obtain LFTs daily if on remdesivir

See Page 3 of algorithm for multi-disciplinary management by sub-specialty recommendations
Report suspected adverse events related to therapeutics through RL solutions

Algorithm reviewed by YNHHS SAS and YNHH/YSM Ad-Hoc COVID-19 Treatment Team
YNHHS Initial Treatment Algorithm for **Hospitalized** **ADULTS with Severe** COVID-19

**Disclaimer:** There are no FDA-approved treatments for COVID-19, supportive care is standard of care. Limited treatment data are available & clinical judgment is warranted - Algorithm last updated 5/18/20

Patient with **confirmed POSITIVE** SARS-CoV-2 by PCR

Assess all patients routinely for clinical trial eligibility (see Appendix 1)

Continue supportive care
Consider adjunctive treatment

YNHH: consider ID input as needed
BH, GH, LMH, or WH: consult ID

**ADJUNCTIVE TREATMENT CONSIDERATIONS**

<table>
<thead>
<tr>
<th>If ≥ 3 Liter O2 requirement</th>
</tr>
</thead>
<tbody>
<tr>
<td>OR ≥ 2 Liter O2 requirement &amp; hs-CRP &gt;70</td>
</tr>
<tr>
<td><strong>Tocilizumab</strong> x 1 dose</td>
</tr>
<tr>
<td><em>(see Appendix 2 for exclusion criteria)</em></td>
</tr>
</tbody>
</table>

**Remdesivir** availability under the EUA is limited. Potential candidates will be identified. Pharmacy will contact primary providers of eligible patients.

Remdesivir has not been FDA approved; remdesivir is authorized by the FDA under and Emergency Use Authorization (EUA)

If worsening ARDS after 48 hours:

**Consider methylprednisolone** 40mg Q8H for 72 hours. Reassess for extended course or taper (up to 5-7 days total).
Steroids given at discretion of primary team

If patient on ECMO or planned for ECMO, also see [ECMO algorithm](#)

**COVID-SPECIFIC TESTS**

1) **Baseline & every 12 hours (for 5 days, then daily thereafter):** CRP, D-dimer

2) **Baseline & every 12 hours x3:** Troponin (continue longer if further testing clinically indicated)

3) **Baseline & every 24 hours**: CBC with differential, BMP, LFTs, Ferritin, Procalcitonin, BNP, fibrinogen, PT/PTT, Mg

4) **On ICU admission:** Cytokine panel

5) **Baseline and with acute kidney injury (AKI):** urinalysis and urine protein/albumin ratio

6) **Baseline EKG** (see Appendix 3 for QTc recommendations)

7) **Repeat Chest X-Ray:** if clinical deterioration. *(CXR not indicated for discharge or to document clinical improvement)*

*May extend longer if clinically indicated* 
Obtain LFTs daily if on remdesivir

Report suspected adverse events related to therapeutics through [RL solutions](#)

See Page 3 of algorithm for multi-disciplinary management by sub-specialty recommendations

Algorithm reviewed by YNHHS SAS and YNHH/YSM Ad-Hoc COVID-19 Treatment Team
YNHHS Initial Treatment Algorithm for **Hospitalized** ADULTS with COVID-19

**Disclaimer:** There are no FDA-approved treatments for COVID-19, supportive care is standard of care. Limited treatment data are available & clinical judgment is warranted - **Algorithm last updated 5/18/20**

### Hematologic:
- If D-dimer <5 mg/L: All patients should receive **standard prophylactic anticoagulation and aspirin 81mg daily** unless contraindicated★
- If D-dimer ≥5 mg/L or receiving convalescent plasma: use **weight-based intermediate prophylactic anticoagulation and aspirin 81mg daily** unless contraindicated★
- If confirmed VTE or high clinical suspicion, start **therapeutic dose anticoagulation and aspirin 81mg daily** unless contraindicated★
- If sudden and unexplained change in O2 OR new asymmetrical upper or lower extremity edema, consider venous U/S of affected extremity
- If ferritin >100,000 or D-dimer >10mg/L, consider Hematology consult at discretion of primary team

(*see Appendix 5 for anticoagulation dosing recommendations)

**Aspirin 81mg PO daily**
- Relative contraindications: recent or risk for CNS bleed, use of other anti-platelet therapy, severe thrombocytopenia, allergy, or history of bleeding disorder
- Discontinue at discharge

### Cardiac:
- Monitor electrolytes: **Replete Mg >2, K >4**
- Baseline EKG and monitor telemetry closely for QTc Prolongation (Appendix 2 for recommendations)
- Caution combining QTc prolonging medications
- If significantly elevated troponin or EKG abnormalities and/or hemodynamic instability, consider POCUS for LV function assessment and cardiology consult

### Obstetrics:
- Treatment Protocol is similar.
- Alternative cut-offs for:
  - Treatment administration with oxygen saturation of ≤ 95%
  - D-dimer cutoff for anticoagulation (see Appendix 5b)

Remdesivir is available to pregnant patients under Expanded Access / Compassionate Use requests. Request only if potential benefits outweigh risks.

*Immunosuppressed hosts* include: Cancer treatment within 1 year, the use of immunosuppressive drugs (biologics, chronic prednisone ≥20mg daily), solid organ transplant, bone marrow transplantation, HIV/AIDS (regardless of CD4 count), leukemia, lymphoma, SLE, vasculitis, and pregnancy
YNHHS Algorithm for Hospitalized ADULTS with COVID-19 requiring ECMO

Disclaimer: There are no FDA-approved treatments for COVID-19, supportive care is standard of care. Limited treatment data are available & clinical judgment is warranted – Algorithm last updated 5/18/20

Guidance for Patients with Confirmed COVID-19 and Refractory Respiratory Failure Requiring ECMO

Prior to cannulation
- Goals of care discussion
- Follow YNHHS COVID-19 Severe Algorithm for treatment and testing
- Evaluate for secondary causes of respiratory failure
- Order pre-ECMO cytokine panel

Evaluation / Management of Secondary Causes of Respiratory Failure
- Vigorous pulmonary toilette
- Infection – blood and sputum cultures
- Pulmonary embolism
- Heart failure – limited TTE

ECMO (24-48 hours)
- Order post-ECMO cytokine panel (after ~48 hours)
- Assess eligibility for clinical trials / expanded access protocols

Potential Adjunctive Therapeutic Resources
- Convalescent plasma administration if eligible
- Remdesivir if available via Expanded Access / Compassionate Use
- Consult Allergy / Immunology to help target immune dysregulation
  - Sarilumab trial if eligible (current trial excludes patients who received an IL-6 antagonist in the prior 30 days)
  - Possible repeat tocilizumab dosing
- Cytokine adsorption via ECMO circuit

* Available options are subject to rapid change *

ECMO (48 hours–2 weeks)
- Consider Allergy / Immunology and Infectious Diseases consultation
- Consider adjunctive therapeutic resources

ECMO (2-3 weeks)
- Revisit goals of care discussions if no clinical improvement after addressing potentially reversible processes

Algorithm reviewed by YNHHS SAS and YNHH/YSM Ad-Hoc COVID-19 Treatment Team
### Appendix 1: Active Coronavirus (SARS-CoV)-2 infection Clinical Trials for Hospitalized Patients

<table>
<thead>
<tr>
<th>Drug, study description and rationale for use</th>
<th>Inclusion and Exclusion Criteria</th>
<th>Notable adverse effects</th>
<th>Primary Investigator(s)/ Contact Information</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Drug: Remdesivir</strong></td>
<td></td>
<td></td>
<td>Pl: Onyema Ogbuagu</td>
</tr>
<tr>
<td>Viral RNA dependent RNA polymerase inhibitor</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Rationale:</strong> In-vitro data reveals potent</td>
<td></td>
<td></td>
<td>Contact: <a href="mailto:Onyema.Ogbuagu@yale.edu">Onyema.Ogbuagu@yale.edu</a></td>
</tr>
<tr>
<td>SARS-COV-2 inhibition and early clinical data</td>
<td></td>
<td></td>
<td><a href="mailto:Laurie.Andrews@yale.edu">Laurie.Andrews@yale.edu</a></td>
</tr>
<tr>
<td>shows possible benefit</td>
<td></td>
<td></td>
<td>Contact (GH expanded access trial):</td>
</tr>
<tr>
<td><strong>Description:</strong> A Phase 3</td>
<td></td>
<td></td>
<td><a href="mailto:Gavin.McLeod@greenwichhospital.org">Gavin.McLeod@greenwichhospital.org</a></td>
</tr>
<tr>
<td>Randomized Study to Evaluate the Safety and</td>
<td></td>
<td></td>
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<tr>
<td>Antiviral Activity of Remdesivir (GS-</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>5734™) in Participants with Severe COVID-19</td>
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<td></td>
<td></td>
</tr>
<tr>
<td><strong>Mild / Moderate Disease</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Inclusion</strong></td>
<td>• Aged ≥ 18 years or Adolescents 12 – 18 years weighing &gt; 40 kg</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Lung involvement confirmed with chest imaging</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Coronavirus (SARS-CoV)-2 infection confirmed by polymerase chain reaction (PCR) test ≤ 4 days before randomization (may repeat test if &gt; 4 days)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Willingness of study participant to accept randomization to any assigned treatment arm</td>
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<td></td>
</tr>
<tr>
<td></td>
<td>• Must agree not to enroll in another study of an investigational agent prior to completion of Day 28 of study</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Key Exclusion</strong></td>
<td>• Severe liver disease</td>
<td></td>
<td>Nausea</td>
</tr>
<tr>
<td></td>
<td>• SaO2/SPO2 ≤ 94% in room air condition, or the PaO2/FiO2 ratio &lt; 300 mg Hg</td>
<td></td>
<td>Vomiting</td>
</tr>
<tr>
<td></td>
<td>• Severe renal impairment or receiving renal replacement therapy</td>
<td></td>
<td>Elevated liver enzymes</td>
</tr>
<tr>
<td></td>
<td>• Pregnant or breastfeeding, or positive pregnancy test in a predose examination</td>
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<td></td>
</tr>
<tr>
<td></td>
<td>• Receipt of any experimental treatment for COVID-19 within the 30 days prior to the time of the screening evaluation</td>
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<tr>
<td></td>
<td>• Creatinine clearance &lt; 50 mL/min</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Severe Disease</strong></td>
<td>• Aged ≥ 18 years or Adolescents 12 – 18 years weighing &gt; 40 kg</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Severe Acute Respiratory Syndrome Coronavirus (SARS-CoV)-2 infection confirmed by polymerase chain reaction (PCR) test ≤ 4 days before randomization (may repeat test if &gt; 4 days)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Peripheral capillary oxygen saturation (SpO2) ≤ 94% or requiring supplemental oxygen at screening</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Key Exclusion</strong></td>
<td>• Participation in any other clinical trial of an experimental treatment for COVID-19</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Concurrent treatment with other agents with actual or possible direct acting antiviral activity against SARS-CoV-2 is prohibited &lt; 24 hours prior to study drug dosing</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Evidence of multiorgan failure</td>
<td></td>
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<tr>
<td></td>
<td>• Mechanically ventilated (including V-V ECMO) ≥ 5 days, or any duration of V-A ECMO</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Requiring mechanical ventilation at screening</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Severe liver disease</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Creatinine clearance &lt; 50 mL/min</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Drug: Sarilumab
Monoclonal antibody to IL6 receptor

**Rationale:** IL-6 receptor antagonist may attenuate cytokine release in patients with severe disease

**Description:**
Phase 2/3, Randomized, Double-Blind, Placebo Controlled Study Assessing Efficacy and Safety of Sarilumab for Hospitalized Patients with COVID-19

### Inclusion
- Aged ≥ 18 years
- Evidence of pneumonia and have one of the following disease categories: severe disease, multi-system organ dysfunction or critical disease
- Laboratory-confirmed SARS-CoV-2 infection
- Elevated liver enzymes
- Leukopenia
- Infusion reactions (e.g. flushing, chills)

### Key Exclusion
- Low likelihood of survival after 48 hours from screening
- Presence of neutropenia less than 2000/mm³
- AST or ALT greater than 5 X ULN
- Platelets < 50,000/mm³ prior immunosuppressive therapies
- Use of chronic oral corticosteroids for non-COVID-19 related condition
- Patients who have received IL-6 receptor antagonist within 30 days of study enrollment
- Participation in any other clinical trial of an experimental treatment for COVID-19
- Known or suspected history of tuberculosis
- Suspected or known active systemic bacterial or fungal infection

**PI:** Geoffrey Chupp
**Contact:** Geoffrey.Chupp@yale.edu

### Expanded access program for use of convalescent plasma in COVID-19 patients

**Inclusion**
- Aged ≥ 18 years
- Confirmed positive SARS-CoV-2 infection by PCR
- Severe or Life-threatening disease by the following definitions
- Severe disease
  - Requiring supplemental oxygen with one or more of the following:
    - Non-rebreather
    - High-flow nasal cannula
    - Pulmonary infiltrates with ≥ 3 L via NC with rapid progression
    - Mechanical ventilation
- Life-threatening disease
  - Refractory respiratory failure, or
  - Septic shock, or
  - Multi-organ dysfunction

**Relative Exclusion**
- ≥ 10 days since first positive SARS-CoV-2 PCR
- Confirmed or high suspicion for bacterial or fungal infection
- D-dimer ≥ 5 mg/L or evidence of/suspicion for thrombosis
- Recent bleeding or high risk for bleeding
- Known severe IgA deficiency

**Contacts**
- YNHH: Mahalia.desruisseaux@yale.edu
- BH: Tina.McCurry@bpthosp.org
- GH: James.Sabetta@greenwichhospital.org
- LMH/WH: Christopher.Song@lmhosp.org

**PI:** Geoffrey Chupp
**Contact:** Geoffrey.Chupp@yale.edu
For single patient INDs and emergency use, expanded access may be appropriate when all the following apply:

- Patient has a serious disease or condition, or whose life is immediately threatened by their disease or condition
- There is no comparable or satisfactory alternative therapy to diagnose, monitor, to treat the disease or condition
- Patient enrollment in a clinical trial is not possible
- Potential patient benefit justifies the potential risks of treatment
- Providing the investigational medical product will not interfere with investigational trials that could support a medical product’s development or marketing approval for the treatment indication

There are several steps necessary when undertaking emergency use of a drug including specific investigator, Sponsor, and FDA requirements. If a provider assesses emergency use of a drug is appropriate they should contact the Yale Human Research Protection Program (HRPP) and the Investigational Drug Service (IDS) (203-688-4872) as soon as possible to get assistance in identifying and navigating the applicable requirements.
Appendix 2: Tocilizumab Exclusion Criteria

a. Anticipated immediate death (≤24 hours) regardless of critical care support

b. **Cardiac**: NYHA Class IV heart failure; Severe, inoperable multi-vessel coronary artery disease; Cardiac arrest; Recurrent arrests in the current presentation, or unresponsive to defibrillation or pacing, or unwitnessed out-of-hospital cardiac arrest with poor prognosis

c. **Hepatic**: Cirrhosis with MELD-Na score ≥25 (in patients who are not transplant candidates), alcoholic hepatitis with MELD-Na >30, advanced liver cancer

d. **Neurologic**: Severe dementia leading to dependence in multiple ADLs; Rapidly progressive or end-stage neuromuscular disease

e. **Oncologic**: Advanced malignancy or high-grade primary brain tumors receiving only palliative treatment with estimated 3 or fewer month prognosis.

f. **Pulmonary**: Severe, chronic lung disease with baseline oxygen requirement of >60% FiO2; Primary pulmonary hypertension with NYHA Class III-IV heart failure (and patient refractory to/not a candidate for pulmonary vasodilators)

g. **Trauma**: Severe trauma; Severe burns: age >60 and 50% of total body surface area affected

h. **Functional Status**: Dependent in all ADLs due to a progressive chronic comorbid condition
Appendix 3: Care Pathways for Mitigation of Drug-Induced Malignant Arrhythmias in COVID-19 Patients

Recommendations:
All COVID-19 patients should have the following:

- When ordering an EKG for a COVID 19 patient to monitor their QTc, select the diagnosis “COVID 19” to alert cardiology to expedite the formal reading of the EKG.
- Daily monitoring of electrolytes; maintain K > 4 and Mg > 2
- All unnecessary QT prolonging drugs should be avoided or switched to alternatives whenever possible.

Recommendations:
A flowchart for the monitoring of potential malignant arrhythmias in these patients is shown below.
Appendix 4: YNHH Acute Respiratory Failure with COVID-19 MICU / SDU Triage Guidelines

- RR < 25
  - Obtain ABG
    - pH > 7.32
    - Hypercapnia with pH < 7.32
      - Consider SDU evaluation, reassess in 2-4 hours
      - Consult MICU
  - Hypercapnia with pH < 7.32
    - Consult MICU

- >4L NC with O2 sat < 93%
  - Consult MICU

- RR > 25 +/- AMS +/- inability to manage secretions
  - Obtain ABG and consult MICU
Appendix 5a: Anticoagulation Dosing Guidelines (Non-Pregnant Patients)

Administer aspirin 81mg PO daily to all patients unless contraindicated. Discontinue aspirin at discharge.

<table>
<thead>
<tr>
<th>DOAC Dosing</th>
<th>D-dimer</th>
<th>BMI &lt; 40kg/m2</th>
<th>BMI ≥ 40kg/m2</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>DOAC</strong></td>
<td><strong>D-dimer ≥ 5 mg/L Intermediate Dose Prophylaxis</strong></td>
<td><strong>Confirmed VTE treatment, high clinical suspicion or clotting of dialysis lines/tubing</strong></td>
<td></td>
</tr>
<tr>
<td>Apixaban</td>
<td>5mg PO Q12H regardless of renal function</td>
<td>10mg PO Q12H x 7 days followed by 5mg PO Q12H (limited data for 10mg in CrCl &lt; 25 or Cr &gt; 2.5)</td>
<td></td>
</tr>
<tr>
<td>Rivaroxaban (may favor in BMI ≥ 40kg/m2)</td>
<td>20mg Q24H Avoid use with CrCl &lt; 30mL/min</td>
<td>15mg PO Q12H x 21 days followed by 20mg PO Q24H Avoid use with CrCl &lt; 30mL/min</td>
<td></td>
</tr>
</tbody>
</table>

Enoxaparin is the preferred form of anticoagulation

Relative contraindications for aspirin: recent or risk for CNS bleed, use of other anti-platelet therapy, severe thrombocytopenia, allergy, or history of bleeding disorder

Target anti-Xa levels between 0.3 – 0.7 units/mL

Patients receiving treatment should continue full dose anticoagulation for 3 months

Consult pharmacy for assistance with dosing recommendations, if needed
Seek hematology input for further recommendations on treatment as needed, including duration and extended prophylaxis for discharge
Appendix 5b: Anticoagulation Dosing Guidelines (Pregnant Patients)

Administer aspirin 81mg PO daily to all patients unless contraindicated. Discontinue aspirin at discharge.

<table>
<thead>
<tr>
<th>D-dimer</th>
<th>BMI &lt; 40 kg/m²</th>
<th>BMI ≥ 40 kg/m²</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 3.5 mg/L</td>
<td>CrCl ≥ 30 mL/min</td>
<td>CrCl ≥ 30 mL/min</td>
</tr>
<tr>
<td>Prophylaxis</td>
<td>• Enoxaparin 40mg sq daily</td>
<td>• Enoxaparin 40mg sq Q12H*</td>
</tr>
<tr>
<td></td>
<td>CrCl &lt; 30mL/min</td>
<td>CrCl &lt; 30mL/min</td>
</tr>
<tr>
<td></td>
<td>• Enoxaparin 30mg sq daily</td>
<td>• Enoxaparin 0.5mg/kg sq Q12H*</td>
</tr>
<tr>
<td>≥ 3.5 mg/L</td>
<td>CrCl ≥ 30 mL/min</td>
<td>CrCl ≥ 30 mL/min</td>
</tr>
<tr>
<td>or receiving convalescent plasma</td>
<td>• Enoxaparin 0.5mg/kg sq Q12H*</td>
<td>• Enoxaparin 0.5mg/kg sq Q12H*</td>
</tr>
<tr>
<td>Intermediate Dose Prophylaxis</td>
<td>CrCl &lt; 30mL/min</td>
<td>CrCl &lt; 30mL/min</td>
</tr>
<tr>
<td></td>
<td>• Enoxaparin 0.5mg/kg sq Q12H*</td>
<td>• Enoxaparin 0.5mg/kg sq Q12H*</td>
</tr>
<tr>
<td>≥ 7 mg/L</td>
<td>CrCl ≥ 30 mL/min</td>
<td>CrCl ≥ 30 mL/min</td>
</tr>
<tr>
<td>Confirmed VTE or high clinical suspicion</td>
<td>• Enoxaparin 1mg/kg sq Q12H</td>
<td>• Enoxaparin 1mg/kg sq Q12H</td>
</tr>
<tr>
<td>TREATMENT</td>
<td>CrCl &lt; 30mL/min</td>
<td>CrCl &lt; 30mL/min</td>
</tr>
<tr>
<td></td>
<td>• Enoxaparin 1mg/kg sq Q24H</td>
<td>• Enoxaparin 1mg/kg sq Q24H</td>
</tr>
</tbody>
</table>

Dosing weight for PREGNANT patients should be actual body weight and POST-PARTUM dosing should be PRE-PREGNANCY weight

Relative contraindications for aspirin: recent or risk for CNS bleed, use of other anti-platelet therapy, severe thrombocytopenia, allergy, or history of bleeding disorder

*Target anti-Xa levels between 0.3 – 0.7 units/mL

Consult pharmacy for assistance with dosing recommendations, if needed
Seek hematology input for further recommendations on treatment as needed, including duration
## Appendix 6

### Possible medications for COVID-19

(Subject to change as more data becomes available and based on medication availability)

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dose</th>
<th>Mechanism</th>
<th>Rationale for use</th>
<th>Notable Adverse Reactions</th>
<th>Other considerations</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>IMMUNOMODULATING AGENTS</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td><strong>Tocilizumab</strong> (1-7)</td>
<td>8mg/kg IV x 1 dose (actual body weight; dose max 800 mg)</td>
<td>Monoclonal antibody to IL6 receptor</td>
<td>IL-6 receptor antagonist may attenuate cytokine release in patients with severe disease</td>
<td>Headache, Elevated liver enzymes, Infusion reactions (e.g. flushing, chills)</td>
<td>The use of IL-6 levels should NOT guide decision to administer tocilizumab at this time</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Retrospective data suggest possible benefit (clinical trials ongoing)</td>
<td></td>
<td>Additional doses not indicated at this time</td>
</tr>
<tr>
<td><strong>Remdesivir</strong> (8-13)</td>
<td>200mg IV once followed by 100mg IV daily for 5 days</td>
<td>Viral RNA dependent RNA polymerase inhibitor</td>
<td>In-vitro data reveals potent SARS-CoV-2 inhibition and early clinical data shows possible benefit</td>
<td>Nausea, vomiting, Elevated liver enzymes, Rectal bleeding</td>
<td>Remdesivir was authorized (not approved) by the FDA through an Emergency Use Authorization (EUA). Availability under the EUA is limited. Potential candidates will be identified. Pharmacy will contact primary providers of eligible patients.</td>
</tr>
</tbody>
</table>

### Medications which may be available through Clinical Trials or Expanded Access

(Subject to change as more data becomes available and based on medication availability)

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dose</th>
<th>Mechanism</th>
<th>Rationale for use</th>
<th>Notable Adverse Reactions</th>
<th>Other considerations</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Remdesivir</strong> (8-13)</td>
<td>200mg IV once followed by 100mg IV daily for 5 or 10 days</td>
<td>Viral RNA dependent RNA polymerase inhibitor</td>
<td>In-vitro data reveals potent SARS-CoV-2 inhibition and early clinical data shows possible benefit</td>
<td>Nausea, vomiting, Elevated liver enzymes, Rectal bleeding</td>
<td>Remdesivir remains available through clinical trials until 5/29/20, with the ability to enroll 5 patients per arm (10 patients total)</td>
</tr>
<tr>
<td></td>
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<td></td>
<td></td>
<td></td>
<td>Available for pregnant patients and patients on ECMO under Expanded Access; request only if benefits outweigh risks</td>
</tr>
<tr>
<td><strong>Convalescent Plasma</strong> (14-18)</td>
<td>One ABO compatible unit</td>
<td>Individual (not pooled) plasma from a recovered COVID19 patient</td>
<td>Transfer of potentially neutralizing antibodies which could diminish viral pathogenesis</td>
<td>Transfusion reactions, Potential to increase hypercoagulability</td>
<td>Available through expanded access, not a trial</td>
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<td>Each unit may contain variable titers of anti-SARS-CoV-2 antibodies with differing avidity</td>
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<td>Cannot be used in patients with IgA deficiency due to risk of anaphylaxis</td>
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<td>Use with intermediate dosing anticoagulation (see Appendix 5 above)</td>
</tr>
</tbody>
</table>
### IMMUNOMODULATING AGENTS

| Sarilumab (19-21) | Clinical Trial dosing | Monoclonal antibody to IL6 receptor | IL-6 receptor antagonist may attenuate cytokine release in patients with severe disease | Elevated liver enzymes | Leukopenia | Infusion reactions (e.g. flushing, chills) | Available through clinical trial only at this time |

#### Medications NOT currently recommended as first line for COVID-19

(Can be considered in certain cases after discussion with Infectious Diseases and Pharmacy)

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dose</th>
<th>Mechanism</th>
<th>Rationale for possible efficacy</th>
<th>Rationale for NOT including as first line agent</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Hydroxychloroquine (HQC)</strong> (22-36)</td>
<td>400mg PO q12h x 24h, then 200mg q12h x 4 days for a 5 day total duration</td>
<td>• Prevents acidification of endosomes interrupting cellular functions and replication</td>
<td>• In-vitro data shows potent SARS-COV-2 inhibition and early clinical data shows possible benefit</td>
<td>• Available data from clinical trials does not demonstrate benefit. Risks outweigh benefits given theoretic risk for cardiac arrhythmia.</td>
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<td>• Prevents viral entry via ACE2 binding</td>
<td>• HCQ was found more potent than chloroquine in inhibiting SARS-CoV-2 in vitro</td>
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<td></td>
<td></td>
<td>• Reduction of viral infectivity</td>
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<tr>
<td></td>
<td></td>
<td>• Immunomodulator</td>
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<tr>
<td><strong>Lopinavir/Ritonavir (37-40)</strong></td>
<td>N/A</td>
<td>Viral protease inhibitor</td>
<td>• In-vitro data reveals potent SARS-COV-2 inhibition</td>
<td>• Limited availability, poor tolerability (such as GI side effects) and recent data demonstrated questionable clinical efficacy</td>
</tr>
<tr>
<td><strong>Atazanavir (41)</strong></td>
<td>N/A</td>
<td>Viral protease inhibitor</td>
<td>• More potent binding to the virus compared to other protease inhibitors in vitro (lower than lopinavir)</td>
<td>• Mild indirect hyperbilirubinemia is common and not indicative of hepatic dysfunction</td>
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<td>• Drug more widely available than other PI’s including lopinavir/ritonavir and better tolerated</td>
<td>• CYP enzyme inhibitor (3A4, 2C8) monitor/discuss with pharmacy potential for drug-drug interactions</td>
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<td>• For patients with NG/OG/NJ open capsules for enteral administration</td>
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<td>• Atazanavir needs an acidic environment for absorption and therefore antacids, H2 blockers, proton pump inhibitors (PPIs) should be avoided. If these agents must be given the administration should be separated as below:</td>
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<td>○ Atazanavir should be given 2 hours before or 1 hour after antacids</td>
</tr>
</tbody>
</table>
Atazanavir should be given at the same time as the H2 blocker or the atazanavir should be given 10 hours after or 2 hours before the H2 blocker

• For PPIs avoid concomitant use

<table>
<thead>
<tr>
<th>Drug &amp; Combination</th>
<th>Dose &amp; Duration</th>
<th>Relevant Details</th>
<th>Additional Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Azithromycin (42)</td>
<td>500 mg x 1, followed by 250 mg q24h x 4 days</td>
<td>• Not well defined; possible immunomodulator</td>
<td>• In a small study, combination of HCQ and azithromycin was associated with significant a reduction in SARS-CoV-2 viral load</td>
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<td>• Very limited data on use of azithromycin alone or in combination with other agents</td>
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<td>• Gautret, et al. study is limited by small sample size (only 6 patients received HCQ &amp; azithromycin combination) and those patients had lower viral loads than other included patients</td>
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<td>• Combination of HCQ and azithromycin and atazanavir can increase the risk for QTc prolongation</td>
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<tr>
<td>Darunavir/Cobicistat (43)</td>
<td>N/A</td>
<td>• Viral protease inhibitor</td>
<td>• In-vitro data shows SARS-COV-2 inhibition</td>
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<td>• Decreased binding to viral protease compared to atazanavir. No clinical data at this time</td>
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<tr>
<td>Ribavirin (44, 45)</td>
<td>N/A</td>
<td>• Viral RNA polymerase inhibitor and inhibition of elongation of RNA fragments</td>
<td>• In vitro data for use in SARS-CoV and MERS-CoV indicates possible activity</td>
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<td>• Limited evidence for SARS-CoV-2 and toxicity risk outweighs benefit of use</td>
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<td>• Typically used with interferon</td>
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<td>• Studied in patients with other coronaviruses with mixed results</td>
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<tr>
<td>Oseltamivir (46)</td>
<td>N/A</td>
<td>• Inhibits influenza virus neuraminidase blocking viral release</td>
<td>• Activity against influenza virus</td>
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<td>• No current data to support use of this drug.</td>
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<td>• Additionally, <strong>SARS-CoV-2 does not use neuraminidase in the replication cycle</strong> so mechanistically there would be no benefit</td>
</tr>
<tr>
<td>Nitazoxanide (47)</td>
<td>N/A</td>
<td>• Augments host antiviral response</td>
<td>• In-vitro data reveals SARS-COV-2 inhibition</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• No clinical data available</td>
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<tr>
<td>IMMUNOMODULATING AGENTS</td>
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<tr>
<td><strong>Interferon-beta</strong> (38-40, 48)</td>
<td>N/A</td>
<td>• Immunomodulator</td>
<td>• Possible activity against SARS-CoV and MERS-CoV&lt;br&gt;• Typically used in combination with ribavirin</td>
</tr>
<tr>
<td><strong>Corticosteroids</strong> (49-53)</td>
<td>If indicated per protocol: Methylprednisolone 40mg q8hr IV for three days, then re-assess</td>
<td>• Inhibit production of inflammatory cytokines that regulate neutrophil and T-cell responses leading to immune suppression</td>
<td>• May be helpful in attenuating cytokine release in patients with severe disease</td>
</tr>
<tr>
<td><strong>Intravenous immunoglobulin (IVIG)</strong> (54, 55)</td>
<td>N/A</td>
<td>• Neutralizing antibodies against the virus</td>
<td>• May have both antiviral and immunomodulatory effects&lt;br&gt;• A recent observational study reported clinical and radiographic improvement in 3 patients who received high dose IVIG at time of respiratory distress</td>
</tr>
<tr>
<td><strong>Baricitinib</strong> (56, 57)</td>
<td>N/A</td>
<td>• Janus Kinase (JAK) inhibitor binding cyclin G - associated kinase, may inhibit viral entry via endocytosis</td>
<td>• May have targeted antiviral and immunomodulatory effect with less side-effects at an effective dose than other JAK inhibitors</td>
</tr>
<tr>
<td><strong>Zinc</strong> (58, 59)</td>
<td>N/A</td>
<td>• Directly impairs RNA synthesis in SARS-CoV by inhibiting the replication and transcription complex, as well as RNA-dependent RNA polymerase.</td>
<td>• Increasing intracellular zinc concentrations may inhibit RNA synthesis</td>
</tr>
</tbody>
</table>
Chloroquine has been demonstrated to be a zinc ionophore. All data is based on in vitro studies only.

Ascorbic acid & Thiamine (60-63) | N/A | • Unclear; ?role in septic shock/ARDS | • ? benefit in septic shock/ARDS

- No published peer reviewed studies in the medical literature were found to support the usage of these vitamins for COVID-19. There are ongoing clinical trials assessing possible benefit.
- Two recently published open-label studies evaluating the use of vitamin C alone and in combination in other types of infections, associated with septic shock and acute respiratory distress syndrome (ARDS) showed no clear evidence of benefit. It cannot be concluded that intravenous vitamin C or thiamine is an effective treatment of ARDS (resulting from COVID-19, or otherwise).

References:

10. Sciences G. Study to Evaluate the Safety and Antiviral Activity of Remdesivir (GS-5734™) in Participants With Severe Coronavirus Disease (COVID-19). NCT042928992020.
11. Sciences G. Study to Evaluate the Safety and Antiviral Activity of Remdesivir (GS-5734™) in Participants With Moderate Coronavirus Disease (COVID-19) Compared to Standard of Care Treatment. NCT042927302020.