

LATEST GUIDELINES COVID 2020

DEFINITIONS:

- ILI is defined as one with acute respiratory infection
 - with fever $\geq 38^{\circ}\text{C}$ (100.4°F) and
 - cough
 - with onset within last 10 days
- SARI is defined as one with acute respiratory infection
 - with fever $\geq 38^{\circ}\text{C}$ (100.4°F),
 - cough
 - with onset within the last 10 days and
 - requiring hospitalization
- Respiratory failure
 - Represents the failure of the lung to maintain adequate gas exchange
 - Characterized by ABG abnormalities: $\text{PaO}_2 < 60$ mmHg with or without hypercarbia $\text{PaCO}_2 > 46$ mmHg (with drop in $\text{pH} < 7.30$)

COVID 19 RT-PCR POSITIVE PATIENT

1. Management of any COVID 19 patient mandates the Health Care Personnel (HCP) to be in full Personal Protection Equipment(PPE).
2. Patient is Categorized in to three groups:

CATEGORY	Type of patients who are provided treatment and care
Group A	Asymptomatic/Patients with mild symptoms RR<24/m & SpO2>94% in room air
Group B	Symptomatic patient with mild to moderate Pneumonia with no signs of severe disease RR: 24-30/m (or) SPO2: 90%-94% at Room Air
Group C	Symptomatic patient with Severe Pneumonia with RR > 30/min (or) SPO2 < 90% at Room Air (or) less than 94% with oxygen, ARDS, Septic Shock

CLINICAL CATEGORIES

Clinical category	Description	Parameters
Asymptomatic	No Symptoms	SpO2: $\geq 94\%$ in room air RR: $\leq 24/\text{m}$ No evidence of hypoxemia or breathlessness
Mild	Patients with uncomplicated upper respiratory tract infection.	SpO2: $\geq 94\%$ in room air RR: $\leq 24/\text{m}$ No evidence of hypoxemia or breathlessness
Moderate	Pneumonia with no signs of severe disease	SpO2: $94\%-90\%$ in room air RR: $24-30/\text{m}$
Severe	Severe Pneumonia	SpO2: $< 90\%$ room air RR: $> 30/\text{m}$
Critical	Acute Respiratory Distress Syndrome (ARDS)	Onset: new or worsening respiratory symptoms within one week of known clinical insult. Chest imaging (Chest X ray and portable bed side lung ultrasound): bilateral opacities, not fully explained by effusions, lobar or lung collapse, or nodules. Origin of Pulmonary infiltrates: respiratory failure not fully explained by cardiac failure or fluid overload. Need objective assessment (e.g. echocardiography) to exclude hydrostatic cause of infiltrates/ oedema if no risk factor present. Oxygenation impairment in adults: Mild ARDS: $200 \text{ mmHg} < \text{PaO}_2/\text{FiO}_2 \leq 300 \text{ mmHg}$ (with PEEP or CPAP $\geq 5 \text{ cm H}_2\text{O}$) Moderate ARDS: $100 \text{ mmHg} < \text{PaO}_2/\text{FiO}_2 \leq 200 \text{ mmHg}$ with PEEP $\geq 5 \text{ cm H}_2\text{O}$ Severe ARDS: $\text{PaO}_2/\text{FiO}_2 \leq 100 \text{ mmHg}$ with PEEP $\geq 5 \text{ cm H}_2\text{O}$ When PaO_2 is not available, $\text{SpO}_2/\text{FiO}_2 \leq 315$ suggests ARDS (including in non- ventilated patients)

Critical	Septic Shock	Adults: persisting hypotension despite volume resuscitation, requiring vasopressors to maintain MAP ≥ 65 mmHg and serum lactate level > 2 mmol/L
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INVESTIGATIONS

Timing	Mild	Moderate	Severe/Critical
At admission	CBC RBS ECG HbA1C (if Diabetic) D-Dimer (If starting on Tab Favipiravir) RFT S.Electrolyte S. Uric Acid	<ul style="list-style-type: none"> • Complete Blood Count (with N/L RATIO) • LFT, RFT, RBS • S.Electrolytes • 12 lead ECG • CHEST X Ray –PA view • CRP, D-DIMER • S. FERRITIN, S.LDH • PROCALCITONIN • TROP – I & T • PT/INR • ABG • CT Thorax (if Available) • Blood culture (if total count is high) • IL – 6 • S. Cortisol • 2D ECHOCARDIOGRAPHY • COVID Antibody IgM/IgG Tests 	<ul style="list-style-type: none"> • Complete Blood Count (with N/L RATIO) • LFT, RFT, RBS • S. Electrolytes • 12 lead ECG • CHEST X Ray –PA view • CRP, D-DIMER • S. FERRITIN, S.LDH • PROCALCITONIN • TROP – I & T • PT/INR • ABG • CT Thorax (if Available) • Blood culture (if total count is high) • IL – 6 • S. Cortisol • S.Mg²⁺, S.Ca²⁺ • 2D ECHOCARDIOGRAPHY • NTproBNP • HsCRP • S. Lactate • COVID Antibody IgM/IgG Tests
Repeat Daily	–	Complete Blood Count, LFT, RFT ABG	Complete Blood Count, LFT, RFT ABG
Repeat Every 72hrs	If initial D-Dimer is high	CRP, D-DIMER S. FERRITIN, S.LDH Chest X ray	CRP, D-DIMER S. FERRITIN, S.LDH Chest X ray
At the time of discharge	–	CRP, D-DIMER S. FERRITIN, S.LDH Chest X ray	CRP, D-DIMER S. FERRITIN, S.LDH Chest X ray RT-PCR – Nasal & Throat swab

Other Investigations should be done based on patient's Co-morbid status

IDENTIFICATION OF HIGH-RISK PATIENT

CO MORBIDITIES	CLINICALLY	LABORATORY VALUE
Age > 50 yrs	Hypoxia- SPO ₂ < 94%	Lymphopenia (<20) with Neutrophil/Lymphocyte ratio >17
Ischemic Heart Disease	Tachycardia > 100/min	CRP > 100 mg/L
Diabetes	Respiratory Distress RR > 30/min	Serum Ferritin > 300 microg/L
Hypertension	Hypotension Systolic BP < 90 mmHg	LDH > 450
Lung Disease (COPD/Asthma/Post TB Sequele)	Altered Sensorium	D-Dimer > 1000 ng/ml
Chronic Kidney Disease/ Chronic Liver Disease		
Immunosuppression / HIV / Malignancy		
Obesity		

Note: Calculation tool for predicting critically ill COVID-19 at admission can be used as reference tool. (Development of Validation of Clinical risk score to predict the occurrence of critical illness in hospitalised patient with COVID-19. JAMA internal Medicine –published online, May 12/05/2020)

GENERAL MEASURES AND GUIDELINES

- 1) Categorize in to A, B, C based on Symptoms, SpO₂ & Respiratory Rate
- 2) Supportive Care:

- Finger Pulse Oximeter for continuous monitoring of Heart rate and Oxygen saturation
- Start oxygen with Mask at saturation of 94% or lower
- HFNC to be used if there is failed oxygen therapy and Non-invasive ventilation (NIV) to be used appropriately with two limb circuit expiratory filters
- Counselling of COVID19 patients (By Counsellor/psychologist/psychiatrist)
- Normal feeding, no dietary restrictions, good oral hydration
- Maintenance IV fluids (If indicated)
- Maintain blood glucose levels <180 mg/dl.
- If Patient is on ACE inhibitors/ARBs, should be continued
- Avoid using NSAIDs other than Paracetamol Unless Absolutely Necessary
- Avoid using Nebulized drugs to avoid aerosolization of virus. **PREFER MDI with SPACER**
- Antibiotic selection in case of superadded bacterial pneumonia should be according to institution antibiogram.

GROUP A - MILD CASES	
TREATMENT	PRECAUTIONS
<p>ANTIVIRAL THERAPY</p> <ul style="list-style-type: none"> • TAB HYDROXYCHLOROQUININE(HCQ) 400MG BD FOR 1 DAY Followed by 200MG 1-0-1 X 4 DAY for patients in COVID CARE CENTER/HOME ISOLATION <p>(OR)</p> <p>Tab FAVIPIRAVIR 1800mg 1-0-1 on Day 1 f/b 800mg 1-0-1 for 6 days (total 7 days) for PATIENTS IN DCHC</p> <p>(OR)</p> <p>If Tab HCQ/Tab FAVIPIRAVIR is contraindicated, then combination of Cap DOXYCYCLIN 100mg 1-0-1 for 5 days + Tab IVERMECTIN 12mg 1-0-0 for 3 days</p> <ul style="list-style-type: none"> • #Cap Oseltamavir 75mg 1-0-1 for 5 days <p>ANTICOAGULATION</p> <ul style="list-style-type: none"> • Inj ENOXAPARIN 40mg S/C 1-0-0 X 7 DAYS (IF D-DIMER IS MORE THAN 1000NG/ML (OR) X-RAY/CT THORAX SHOWING GROUND GLASS OPACITIES) <p>SUPPORTIVE THERAPY-</p> <ul style="list-style-type: none"> • TAB ZINC 50 MG 0-1-0 X 7 DAYS • TAB VITAMIN C 500 MG 1-1-1 X 7 DAYS • Tab N Acetylcysteine 600mg 1-1-1 If Patients Having Cough 	<ul style="list-style-type: none"> • CATEGORIZATION SHOULD BE REASSESSED REGULARLY • CONTRAINDICATION FOR HCQ- <ul style="list-style-type: none"> • QT INTERVAL > 480ms • Pre-existing cardiomyopathy and cardiac rhythm disorders • History of Unexplained Syncope • Retinopathy, • Hypersensitivity to HCQ or 4-aminoquinoline compounds • G6PD deficiency • Epilepsy • Hypokalemia ($K^+ < 3$ Meq) • Contraindications for Tab FAVIPIRAVIR: Hyperuricaemia, severe hepatic & renal impairment, Pregnant women and lactating mothers • PREGNANCY IS NOT A CONTRAINDICATION FOR HCQ • # Cap OSELTAMAVIR is advised due to possibility of H1N1 co infection along with COVID19 disease with present weather condition. Its usage will be reviewed at a later date.

GROUP B - MODERATE CASES	
TREATMENT	PRECAUTIONS
<p>ANTIVIRAL THERAPY</p> <ul style="list-style-type: none"> • Inj REMDESIVIR 200 mg IV on day 1 followed by 100 mg IV daily for 4 days (total 5 days) <p>IF REMDESIVIR IS NOT AVAILABLE TO START TAB HYDROXYCHLOROQUININE(HCQ) 400MG BD FOR 1 DAY followed by 200MG 1-0-1 X 4 DAY</p> <p>Co-administration of Inj REMDESIVIR with HCQ or chloroquine should be avoided</p> <ul style="list-style-type: none"> • Cap Oseltamavir 75mg 1-0-1 for 5 days <p>STERIODS</p> <ul style="list-style-type: none"> • Inj. Methyl Prednisolone 0.5 -1 mg/kg (or) Inj. Dexamethasone 0.1 – 0.2 mg/kg for 3-5 Days <p>ANTICOAGULATION</p> <ul style="list-style-type: none"> • Inj ENOXAPARIN 40MG S/C 1-0-0 x 7 DAYS 	<p>Contraindications for Inj REMDESIVIR:</p> <ul style="list-style-type: none"> • AST/ALT > 5 times Upper limit of normal (ULN) • Severe renal impairment (i.e., eGFR < 30ml/min/m² or need for hemodialysis) • Pregnancy or lactating females • Children (< 12 years of age) <ul style="list-style-type: none"> • No dose adjustment for Inj REMDESIVIR if eGFR >30ml/min • Formula to calculate eGFR in Adults • eGFR, Male: $(140 - \text{age in years}) \times (\text{weight in kg}) / 72 \times (\text{serum creatinine in mg/dL})$; • eGFR, Female: $(140 - \text{age in years}) \times (\text{weight in kg}) \times 0.85 / 72 \times (\text{serum creatinine in mg/dL})$ <p>STERIODS</p> <ul style="list-style-type: none"> • to be started preferably within 48 hours of admission (or) if oxygen requirement is increasing and if inflammatory markers are increased.

IV ANTIBIOTICS ACCORDING TO LOCAL ANTIBIOGRAM

AWAKE PRONING

- **CONVALASCENT PLASMA THERAPY:** 4 to 13 ml/kg (usually 200 ml single dose given slowly over not less than 2 hours)

SUPPORTIVE THERAPY

- TAB ZINC 50 MG 0-1-0 X 7 DAYS
- TAB VITAMIN C 500 MG 1-1-1 X 7 DAYS
- TAB N-ACETYL CYSTEINE 1-1-1 IN PATIENTS WITH COUGH

- PATIENT SHOULD BE REASSESSED EVERY 12 HRLY AND CONTINOUS MONITORING OF SATURATION.
- START ON OXYGEN-NASAL PRONGS 2-5 L/MIN or FACE MASK 5L/MIN

GROUP C - SEVERE/CRITICAL CASES

TREATMENT	PRECAUTIONS
<p>ANTIVIRAL THERAPY</p> <ul style="list-style-type: none"> • If the patient has not received Inj REMDESIVIR, such patients can be started on Inj REMDESIVIR. Inj REMDESIVIR 200 mg IV on day 1 followed by 100 mg IV daily for 4 days (total 5 days) • Inj. TOCILUZUMAB 8mg/kg (maximum 800 mg at one time) given slowly in 100 ml NS over 1 hour; dose can be repeated once after 12 to 24 hours if needed <p style="text-align: center;">(Or)</p> <p>Inj ITOLIZUMAB: 1st dose – 1.6mg/kg dose iv infusion. Subsequent dose: weekly 0.8mg/kg dose infusion over 4hours if required</p> <ul style="list-style-type: none"> • Cap Oseltamavir 75mg 1-0-1 for 5 days <p>STERIODS</p> <ul style="list-style-type: none"> • Inj. Methyl Prednisolone 1-2 mg/kg (or) Inj. Dexamethasone 0.2 – 0.4 mg /kg for 5-7 Days <p>ANTICOAGULATION</p> <ul style="list-style-type: none"> • Inj ENOXAPARIN 1mg/kg body wt s/c 1-0-1 X 7 DAYS <p>PRONE VENTILLATION</p> <p>Inj CEFTRIAZONE 1gm IV 1-0-1 AND CAN BE ESCALATED ACCORDING TO LOCAL ANTIBIOGRAM OR TREATING PHYSICIAN</p> <p>CONSIDER SEPSIVAC (IF AVAILABLE) 0.3ml INTRADERMAL ONCE A DAY FOR 3 DAYS IN CASE OF SEPTIC SHOCK</p> <p>IV Diuretics in case of evidence of Heart Failure secondary to Myocarditis</p> <p>SUPPORTIVE THERAPY</p> <ul style="list-style-type: none"> • TAB ZINC 50 MG 0-1-0X 7 DAYS • INJ. VITAMIN C 1.5GM IV 6 HOURLY X 5DAYS • TAB N-ACETYL CYSTEINE 1-1-1 	<p>Indication for TOCILUZUMAB/ITOLIZUMAB:-</p> <ol style="list-style-type: none"> 1. IL-6 levels 50-100 fold higher than normal (Normal range 0 - 9.5pg/ml) 2. Worsening trend of the inflammatory markers (Ferritin, LDH, CRP) 3. Deteriorating clinical condition with worsening of PaO2/Fio2 ratio (more than 25% deterioration from the immediate previous value). <p>Contraindications for Inj TOCILUZUMAB/ITOLIZUMAB</p> <p>PLHIV, those with active infections (systemic bacterial/fungal), High Serum. Procalcitonin, Tuberculosis, active hepatitis, Absolute Neutrophil Count < 2000/mm3 and Platelet count < 1,00,000/mm3, hepatic and renal impairment; patients on chronic steroid therapy, Paediatric patients <18 years old; Pregnancy and, Nursing mothers</p> <ul style="list-style-type: none"> • PATIENT SHOULD BE CONTINOUSLY MONITORED • TO START ON OXYGEN WITH FACE MASK WITH NON REBREATHING BAG @ 8-10 lt/m • BASED ON PaO2/Fio2 ratio, HIGH FLOW NASAL OXYGEN (HFNC)/NIV SHOULD BE GIVEN AND IF PATIENT DETERIORATES INTUBATION SHOULD BE CONSIDERED AND LUNG PROTECTIVE VENTILATION TO BE FOLLOWED AS PER ARDSnet PROTOCOL • ABG TO BE DONE REGULARLY FOR MONITORING OF ACIDOSIS AND HYPOXEMIA • INOTROPHIC SUPPORT (NORADRENALINE – TITRATE ACCORDING TO THE MEAN ARTERIAL PRESSURE) • CORRECTION OF ACIDOSIS • MAINTAIN Hb% GREATER THAN 8gm%

SUMMARY OF TREATMENT OF COVID-19 PATIENTS BASED ON CLINICAL CATEGORIES

MILD	MODERATE	SEVERE/CRITICAL
<p>Antiviral Therapy*</p> <p>Tab Hydroxychloroquine(HCQ) 400mg Bd For 1 Day F/B 200mg 1-0-1 X 4 Day for patients in COVID CARE CENTER/HOME ISOLATION</p> <p style="text-align: center;">(OR)</p> <p>Tab FAVIPIRAVIR 1800mg 1-0-1 on Day 1 f/b 800mg 1-0-1 for 6 days for PATIENTS IN DCHC</p> <p style="text-align: center;">(OR)</p>	<p>Antiviral Therapy*</p> <p>Inj REMDESIVIR 200 mg IV on day 1 followed by 100 mg IV daily for 4 days</p> <p style="text-align: center;">(Or)</p> <p>IF REMDESIVIR IS NOT AVAILABLE TO START Tab Hydroxychloroquine(HCQ) 400mg BD For 1 Day F/B 200mg 1-0-1 X 4 Day</p> <p>Co-administration of Inj REMDESIVIR with HCQ or chloroquine should be avoided</p>	<p>Antiviral Therapy*</p> <p>Inj. TOCILUZUMAB 8mg/kg (maximum 800 mg at one time) given slowly in 100 ml NS over 1 hour; dose can be repeated once after 12 to 24 hours if needed</p> <p style="text-align: center;">(Or)</p> <p>Inj ITOLIZUMAB: 1st dose – 1.6mg/kg dose iv infusion. Subsequent dose: weekly 0.8mg/kg dose infusion over 4hours if required</p>

<p>If Tab HCQ/Tab FAVIPIRAVIR is contraindicated, then combination of</p> <p>Cap DOXYCYCLIN 100mg 1-0-1 for 5 days +</p> <p>Tab IVERMECTIN 12mg 1-0-0 for 3 Days</p> <p>Anticoagulation</p> <p>Inj Enoxaparin 40mg S/C 1-0-0 x 7 days (If D-dimer Is More Than 1000ng/ml or X-ray/CT Thorax Showing Ground glass opacity)</p> <p>Supportive Therapy</p> <p>Tab Zinc 50 Mg 0-1-0x 7 Days Tab Vitamin C 500 Mg 1-1-1 X 7 Days Tab N Acetylcysteine 1-1-1 If Patients Having Cough</p>	<p>STERIODS</p> <p>Inj. Methyl Prednisolone 0.5 -1 mg/kg (or) Inj. Dexamethasone – 0.2 mg /kg for 3-5 Days</p> <p>ANTICOAGULATION</p> <p>Inj Enoxaparin 40mg S/C 1-0-0 x 7 days (if Wt >65kg, 60md 1-0-1 for 7days)</p> <p>Iv Antibiotics According to Local Antibiogram</p> <p>Awake Proning</p> <p>Start on oxygen –Nasal Prongs 2- 5l/min or face mask 5l/min</p> <p>CONVALASCENT PLASMA THERAPY:</p> <p>4 to 13 ml/kg (usually 200 ml single dose given slowly over not less than 2 hours</p> <p>Supportive Therapy</p> <p>Tab Zinc 50 Mg 0-1-0x 7 Days Tab Vitamin C 500 Mg 1-1-1 X 7 Days Tab N Acetylcysteine 1-1-1 If Patients Having Cough</p>	<p>STERIODS</p> <p>Inj. Methyl Prednisolone 1-2 mg/kg for 5-7 Days (or) Inj. Dexamethasone 0.2 – 0.4 mg /kg for 5-7 Days</p> <p>ANTICOAGULATION</p> <p>Inj Enoxaparin 1 Mg/Kg Body Weight S/C 1-0-1 X 7days</p> <p>Inj Ceftriaxone 1 Gm Iv 1-0-1 And Can Be Escalated According To Local Antibiogram Or Treating Physician</p> <p>Start on oxygen with face mask+NRM and change over to HFNC/NIV (based on PaO₂/FiO₂)</p> <p>IF PATIENT DETERIORATES with HFNC/NIV trial (repeat ABG after 6hrs suggests worsening of oxygenation) then EARLY INTUBATION SHOULD BE CONSIDERED AND LUNG PROTECTIVE VENTILATION TO BE FOLLOWED AS PER ARDSnet PROTOCOL</p> <p>Prone Ventillation SEPSIVAC 0.3ml</p> <p>INTRADERMAL</p> <p>ONCE A DAY FOR 3 DAYS</p> <p>Supportive Therapy</p> <p>Inj. Vitamin C 1.5gm Iv 6 Hourly X 5 days Tab Zinc 50 Mg 0-1-0x 7 Days Tab N Acetylcysteine 1-1-1 If Patients Having Cough</p>
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1. Continous monitoring of oxygen saturation by pulse oximeter and early diagnosis of hypoxemia is essential in all group of patients
2. Indications and contraindications of the drugs are to be considered before use which is mentioned in detail below
3. Transition of patients between the clinical categories is based on SpO₂, RR & PaO₂/FiO₂ ratio
4. Treatment of all co morbid illness to continue

Special Note:

- *Cap Oseltamavir 75mg 1-0-1 for 5 days to be added to patients of all categories
- All the investigational therapies and drugs approved recently by DGI should be used with caution and after informed consent from the patient

1. Hydroxychloroquine (HCQ)

Dose: Tab HCQ 400MG BD FOR 1 DAY Followed by 200MG 1-0-1 X 4 Days

CONTRAINDICATION FOR HCQ

- QT INTERVAL > 480ms
- Pre-existing cardiomyopathy and cardiac rhythm disorders
- History of Unexplained Syncope
- Retinopathy,
- Hypersensitivity to HCQ or 4-aminoquinoline compounds
- G6PD deficiency
- Epilepsy
- Hypokalemia (K⁺ < 3 Meq)

2. Anticoagulant Agents

Pro Coagulant factors are increased in COVID-19 infection and associated with increased risk of thrombosis

Pneumonia and sepsis are complicated by DIC, but although COVID-19 patients do have abnormalities of coagulation and are not atypical of DIC.

The most marked abnormality is an elevation of D-Dimer (if D-dimer is more than 1000ng/ml) but without a parallel fall in platelet or prolongation of clotting time, this suggests that local rather disseminated thrombin generation and fibrinolysis is taking place

Dose:

Inj ENOXAPARIN 40MG S/C Once daily for mild and moderate. Twice daily in severe cases.

Other options:

- Inj Fondaparinux 2.5mg OD SC
- Unfractionated Heparin 5000 Units BD SC

Contraindications:

ESRD, active bleeding, emergency surgery, platelets < 20,000/mm³, BP >200/120 mmHg)

INVESTIGATIONAL THERAPIES (as per MOHFW)

- 1. Remdesivir** (under Emergency Use Authorization) may be considered in patients with moderate disease (those on oxygen) with none of the following contraindications:
 - AST/ALT > 5 times Upper limit of normal (ULN)
 - Severe renal impairment (i.e., eGFR < 30ml/min/m² or need for hemodialysis)
 - Pregnancy or lactating females
 - Children (< 12 years of age)**Dose:** 200 mg IV on day 1 followed by 100 mg IV daily for 4 days (total 5 days)

- 2. Convalescent plasma (Off Label)** may be considered in patients with moderate disease who are not improving (oxygen requirement is progressively increasing) despite use of steroids. Special prerequisites while considering convalescent plasma include:
 - ABO compatibility and cross matching of the donor plasma
 - Neutralizing titer of donor plasma should be above the specific threshold (if the latter is not available, plasma IgG titer (against S-protein RBD) above 1:640 should be used)
 - Recipient should be closely monitored for several hours post transfusion for any transfusion related adverse events
 - Use should be avoided in patients with IgA deficiency or immunoglobulin allergy

Dose: Dose is variable ranging from 4 to 13 ml/kg (usually 200 ml single dose given slowly over not less than 2 hours)

- 3. Tocilizumab (Off Label)** may be considered in patients with severe disease with progressively increasing oxygen requirements and in mechanically ventilated patients not improving despite use of steroids. Long term safety data in COVID 19 remains largely unknown. Special considerations before its use include:
 - **IL-6 levels 50-100 fold higher than normal (Normal range 0 - 9.5pg/ml)**
 - **Worsening trend of the inflammatory markers (Ferritin, LDH, CRP)**
 - **Deteriorating clinical condition with worsening of PaO₂/Fio₂ ratio (more than 25% deterioration from the immediate previous value)**

The drug is contraindicated in

PLHIV, those with active infections (systemic bacterial/fungal), High Serum. Procalcitonin, Tuberculosis, active hepatitis, Absolute Neutrophil Count < 2000/mm³ and Platelet count < 1,00,000/mm³, hepatic and renal impairment; patients on chronic steroid therapy, Paediatric patients <18 years old; Pregnancy and, Nursing mothers

Dose: 8mg/kg (maximum 800 mg at one time) given slowly in 100 ml NS over 1 hour; dose can be repeated once after 12 to 24 hours if needed

Drugs Recently approved by DGCI

1. ITOLIZUMAB (An anti-CD6 IgG1 monoclonal antibody) Indication:

- 1. IL-6 levels 50-100 fold higher than normal (Normal range 0 - 9.5pg/ml)**
- 2. Worsening trend of the inflammatory markers (Ferritin, LDH, CRP)**
- 3. Deteriorating clinical condition with worsening of PaO₂/Fio₂ ratio (more than 25% deterioration from the immediate previous value).**

Dose: 1st dose – 1.6mg/kg dose iv infusion

- Subsequent dose: weekly 0.8mg/kg dose infusion over 4hours if required based on lung function parameters

Contraindication:

PLHIV, those with active infections (systemic bacterial/fungal), High Serum. Procalcitonin, Tuberculosis, active hepatitis, Absolute Neutrophil Count < 2000/mm³ and Platelet count < 1,00,000/mm³, hepatic and renal impairment; patients on chronic steroid therapy, Paediatric patients <18 years old; Pregnancy and, Nursing mothers

Side effects:

- In trial Infusion reactions have been reported in 15% of the patients
- In clinical practice also infusion reaction ranged from 12% to 15%
- Other adverse events include Diahorea, Pruritus in 7 – 12 % of cases

2. Tab. FAVIPIRAVIR

Mechanism of action: It is considered that favipiravir is metabolized in cells to a ribosyl triphosphate form (favipiravir RTP) and that favipiravir RTP selectively inhibits RNA polymerase involved in influenza viral replication

Indications: mild to moderate cases of COVID19 in adults >18yrs old

Dose: 1800mg bid followed by 800mg bid upto maximum of 14days

Contraindications: Hyperuricaemia, severe hepatic & renal impairment, Pregnant women and lactating mothers

Side Effects: increased Uric Acid levels, diarrhea, decreased neutrophil counts, increase in AST/ALT levels

Drug Interactions: metabolised partly by Aldehyde Oxidase(AO) and partly by Xanthine Oxidase(XO). Precautions for co-administration with Pyrazinamide, Repaglinide, Theophylline, Famciclovir

PRONE VENTILATION

Early self-proning in awake, non-intubated patients – Moderate cases

- Any COVID-19 patient with respiratory embarrassment severe enough to be admitted to the hospital may be considered for rotation and early self-proning.
- Care must be taken to not disrupt the flow of oxygen during patient rotation

Criteria to be fulfilled	Avoid proning
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<ul style="list-style-type: none"> • Patients with oxygen requirement of >4L • Normal mental status • Able to self-prone or change position with minimal assistance 	<ul style="list-style-type: none"> • Hemodynamic instability • Close monitoring not possible
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- Typical protocols include 30–120 minutes in prone position, followed by 30–120 minutes in left lateral decubitus, right lateral decubitus, and upright sitting position

(Caputo ND, Strayer RJ, Levitan R. Academic Emergency Medicine 2020;27:375–378)

Requirements for safe prone positioning in ARDS

- Pre-oxygenate the patient with FiO2 1.0
- Secure the endotracheal tube and arterial and central venous catheters
- Adequate number of staff to assist in the turn and to monitor the turn
- Supplies to turn (pads for bed, sheet, protection for the patient)
- Knowledge of how to perform the turn as well as how to supine the patient in case of an emergency

Contraindications to prone ventilation

- Spinal instability requires special care
- Intra cranial pressure may increase on turning
- Rapidly return to supine in case of CPR or defibrillation

When to start proning in SEVERE CASES?

- P/F ratio <150 while being ventilated with FiO2 >0.6 and PEEP >5 cm H2O When to stop proning?
- When P/F exceeds 150 on FiO2 > 0.6 and > 6 PEEP

What portion of the day should patients be kept prone?

- As much as possible (16-18 hours a day)
- Adult patients with severe ARDS receive prone positioning for more than 12 hours per day (strong recommendation, moderate-high confidence in effect estimates)

(ATS-ERS Guideline. Am J Respir Crit Care Med;2017;195(9):1253-1263)

Oxygen delivery protocol

- SpO2 < 94% ~ Supplement with nasal prongs or simple face mask at 2-5L/min



- Monitor continuous SpO2 with finger pulse oximetry
- If SpO2 < 94% on simple face mask or nasal prongs, change to non-rebreather mask oxygen (NRB) at 10-15L/min



- Oxygen Delivery Devices & approximate FiO2%

100% O ₂ Flow Rate (L/min)	FiO ₂ (%)
Nasal Cannula	
1	24
2	28
3	32
4	36
5	40
6	44
Oxygen Mask	
5–6	40
6–7	50
7–8	60
Mask with Reservoir Bag	
6	60
7	70
8	80
9	90
10	>99
Nonrebreathing Mask	
4–10	60–100
Venturi Mask*	
3 (80)	24
6 (68)	28
9 (50)	35
12 (50)	40
15 (41)	50

*Number in parentheses indicates total flow of entrained room air with Venturi mixture.

HFNO (High Frequency Nasal Oxygen) and NIV (Non-invasive Ventilation)

- When oxygen requirement increases to needing NRB, options of High Frequency Nasal Oxygen (HFNO) or NIV should be considered.
- HFNC flow rates to be set from 30 -60 L/min titrating to maintain SpO₂ ≥ 92%
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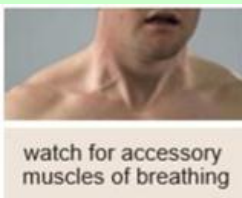
HFNC provides PEEP up to 5-6 cm H₂O and can deliver FiO₂ up to 100%

- If HFNC non-available or patient not maintaining SpO₂ on flow rates up to 60L/min, initiate on non-invasive ventilation (NIV) only with an ICU ventilator with two limbed circuit and expiratory HME filter with a NIV mode available. Caution is to be exercised to not use portable home BiPAP or CPAP machines with single circuit for these patients.



NIV settings:
FiO₂ to be titrated to maintain SpO₂ ≥ 92%.
Pressure Support (IPAP): 12-15cm H₂O (to target tidal volume of 6ml/kg)
PEEP (EPAP): 5-15 cmH₂O as tolerated to achieve SpO₂ ≥ 90-92%
Backup rate: 15 breaths/min
Backup I:E ratio 1:3
Trigger: maximum sensitivity

- Appropriate mask with good seal to be ensured when initiated on NIV. Helmetmasks/hoods if available, to be preferred to minimize aerosol contamination.
- Once initiated on NIV, close monitoring of respiratory variables hourly is important.
- Reassess clinical condition hourly, monitor and observe ABG's 4-6hrly
- Look for signs of clinical improvement in the form of settling tachycardia, improving SpO₂, reduced tachypnea and reduced work of breathing.
- On NIV when there are signs of clinical deterioration in the form of worsening sensorium, increased accessory muscles of breathing, raising Pco₂, worsening pH on ABG ~ failure of NIV has to be considered and patient has to be planned for intubation and mechanical ventilation after consent from the family.



Intubation and Mechanical Ventilation

- Indication for intubation: ARDS with PaO₂/FiO₂ < 200
- Worsening respiratory distress even on NIV
- Patient in Shock

Initial Settings: Controlled Mode ventilation: VCV (volume-controlled ventilation) or PCV (pressure-controlled ventilation)

- Tidal Volume (V_t) 6-8ml/PBW (predicted body weight)
- PEEP 8 – 18 cmH₂O (follow FiO₂-PEEP table) to titrate to target SpO₂ 90-92%
- FiO₂ ~ target SpO₂ 90-92% with lowest FiO₂ possible
- Respiratory rate 14-18/min (maximum up to 35/min)
- Plateau pressure < 30 cmH₂O and driving pressure < 16cmH₂O
- ABG targets: PaO₂ 55-80 mmHg, pH > 7.3
- Measure compliance 6hrly ~ V_t in ml /Pplat – PEEP

Notes:

Predicted body weight (PBW)
 Males = 50 + 2.3 [height (inches) - 60]
 Females = 45.5 + 2.3 [height (inches) - 60]

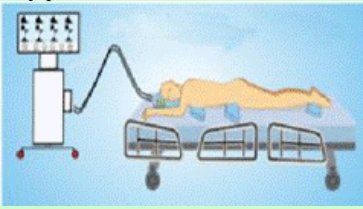
Incremental PEEP FiO₂ table

FiO ₂	0.3	0.4	0.4	0.5	0.5	0.6	0.7	0.7	0.7	0.8	0.9	0.9	0.9	1.0
PEEP	5	5	8	8	10	10	10	12	14	14	14	16	18	18 - 24

Additional steps: If Pplat > 30cmH₂O, reduce V_t upto 4ml/PBW

- If SpO₂ < 88% despite ARDSnet protocol: increase depth of sedation
- Optimize secretions clearance/bronchodilation

- Initiate early muscle relaxant infusion (cis-atracurium or vecuronium)
- Early prone ventilation



Indication for prone ventilation:

- Intubation and mechanical ventilation < 36hrs
- PaO₂/FiO₂ < 150, FiO₂ > 60%, PEEP > 5, Vt 6ml/PBW
- Duration of proning: 12-16 hrs.
- Multiple sessions until favorable trends are achieved.

Adjunctive measures when intubated and mechanically ventilated:

- Antibiotics guided by protocols
- Steps to reduce VAP (ventilator associated pneumonia) by following VAP bundles
- Head-end elevation
- Thrombo-prophylaxis
- Adequate analgesia and sedation

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In absence of ABG facility at the hospitals, use SpO₂/FiO₂ ratio as described in the below table

Derivation of SpO₂/FiO₂ values corresponding to PaO₂/FiO₂ ratios in the combined anesthesia and ARMA database*

SOFA Respiratory score	PaO ₂ /FiO ₂	SpO ₂ /FiO ₂
1	<400	<512
2	<300	<357
3	<200	<214
4	<100	<89

* Data derived from 4728 matched SpO₂/FiO₂ and PaO₂/FiO₂ measurements from the combined anesthesia and ARMA database

Derivation of SpO₂/FiO₂ values corresponding to PaO₂/FiO₂ stratified by PEEP in the ARMA database*

SOFA Respiratory score	PaO ₂ /FiO ₂ ratio		SpO ₂ /FiO ₂ ratio	
		PEEP <8	PEEP 8-12	PEEP >12
1	<400	<502	<515	<425
2	<300	<370	<387	<332
3	<200	<240	<259	<234
4	<100	<115	<130	<129

* Data derived from 2916 matched SpO₂/FiO₂ and PaO₂/FiO₂ measurements from the ARMA database of the ARDS network's NIH study(8)

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