Management Protocol for Long Haul COVID-19 Syndrome (LHCS)

The approach outlined below is a consensus protocol based on a collaboration led by Dr. Mobeen Syed ("Dr. Been"), Dr. Tina Peers, and the FLCCC Alliance. Given the lack of clinical treatment trials of Long Haul COVID-19 Syndrome, these recommendations are based on the pathophysiologic mechanisms of COVID-19 and post-viral illnesses along with our collective experience observing profound and sustained clinical responses achieved with the treatment approaches below.

This protocol has also been used to treat post-vaccine inflammatory syndromes with similar success. As with all FLCCC Alliance protocols, the components, doses, and durations will evolve as more clinical data accumulates.

If the patient presents with shortness of breath or low oxygen levels: Refer to lung specialist if available, otherwise perform chest imaging (CT preferred) to assess for secondary organizing Pneumonia (OP). If findings consistent with secondary OP found, initiate Corticosteroid Therapy as below. May need to repeat or prolong course of treatment if symptoms or oxygen needs persist.

1. **FIRST LINE THERAPIES**
   - **IVERMECTIN**: 0.2 mg/kg body weight. Once daily for 1 week.¹
   - **PREDNISONE**: 10–15 mg daily for 3 weeks. Taper to 10 mg for three days, then 5 mg for three days and then stop.¹
   - Low dose **NALTREXONE** (LDN): Begin with 1 mg daily and increase to 4.5 mg as required. May take 2–3 months for full effect.
   - **OMEGA-3 FATTY ACIDS**: Vascepa, Lovaza or DHA/EPA 4 g per day. Omega-3 fatty acids play an important role in the resolution of inflammation by inducing resolvin production
   - **VITAMIN D**: The majority of those with post-COVID-19 syndrome continue to have hypovitaminosis D. See tables 1 or 2 for vitamin D supplementation.

   If symptoms do not improve after 1–2 weeks continue steroids, omega-3 fatty acids and Naltrexone and add second line medications.

2. **SECOND LINE THERAPIES**
   - **FLUVOXAMINE** (low dose): 25 mg once daily. Stop if the symptoms increase. Caution with the use of other anti-depressants and psychiatric drugs. Taper and discontinue once symptoms improve.
   - **ATORVASTATIN**: 20–40 mg once daily. Caution in patients with Postural Orthostatic Tachycardia Syndrome (POTS); may exacerbate symptoms.

3. **THIRD LINE THERAPY**
   - **MARAVIROC**: 300 mg PO twice a day

   If 6–8 weeks have elapsed and significant symptoms persist this drug can be considered. Note that maraviroc can be expensive and carries the risk of significant side effects and interactions with other medications.

4. **OPTIONAL ADJUNCTIVE THERAPIES** (in order of priority)
   - **Curcumin**: 500 mg BID (has anti-inflammatory and immunomodulating properties and has been demonstrated to repolarize macrophages).
   - **Nigella Sativa**: 40 mg/kg/day (1 tsp ≈ 3.3 grams) – like curcumin it has anti-inflammatory and immunomodulating properties.
   - **Vitamin C**: 500 mg BID (vitamin C inhibits histamine and repolarizes monocytes).
   - **Melatonin**: 2–8 mg at night (slow release/extended release) with attention to sleep hygiene. Increase dose from 1 mg as tolerated (may cause severe nightmares at high dosages).
   - **Kefir, probiotic yogurt and/or Bifidobacterium Probiotics** (e.g., Daily Body Restore) together with **Prebiotics** (e.g. XOS Prebiotic, Bio Nutrition Pre-Biotic) to normalize the microbiome. Prolonged dysbiosis has been reported following COVID-19 infection.
   - **Behavioral modification, mindfulness therapy and psychological support** may help improve survivors’ overall well-being and mental health.
   - **Luteolin 100–200 mg day or Quercetin** 250 mg day (or mixed flavonoids). Luteolin and quercetin have broad spectrum anti-inflammatory properties. These natural flavonoids inhibit mast cells, and have been demonstrated to reduce neuroinflammation.
   - **H1 receptor blockers** (for mast cell activation syndrome): **Loratadine** 10 mg daily, or **Cetirizine** 5–10 mg daily, or **Fexofenadine** 180 mg — daily.
   - **H2 receptor blockers** (for mast cell activation syndrome): **Famotidine** 20–40 mg, or **Nizatidine** 150 mg — twice daily as tolerated.
   - **Montelukast**: 10 mg/day (for mast cell activation syndrome). Caution as may cause depression is some patients.
   - **Anti-androgen therapy**: **Spironolactone** 50–100 mg twice a day, and **Dutasteride** 1 mg daily.

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1. Relative contraindications: 1) Patients on Warfarin require close monitoring and dose adjustment.
2. Pregnant or lactating women require a more in-depth risk/benefit assessment.
2. Side effects may include: Increased appetite, mood changes, insomnia, raised blood glucose, dyspepsia.
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Tables

Table 1. Guidance on upfront loading dose regimens to replenish Vitamin D stores in the body

<table>
<thead>
<tr>
<th>Serum vitamin D (ng/mL)**</th>
<th>Vitamin D dose, 50,000 IU capsules: Initial and weekly ***</th>
<th>Duration (weeks)</th>
<th>Total amount for deficit correction (IU, in millions) ****</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 10</td>
<td>300,000</td>
<td>8 – 10</td>
<td>1.5 – 1.8</td>
</tr>
<tr>
<td>11–15</td>
<td>200,000</td>
<td>8 – 10</td>
<td>1.0 – 1.2</td>
</tr>
<tr>
<td>16–20</td>
<td>200,000</td>
<td>6 – 8</td>
<td>0.8 – 1.0</td>
</tr>
<tr>
<td>21–30</td>
<td>100,000</td>
<td>4 – 6</td>
<td>0.5 – 0.7</td>
</tr>
<tr>
<td>31–40</td>
<td>100,000</td>
<td>2 – 4</td>
<td>0.3 – 0.5</td>
</tr>
<tr>
<td>41–50</td>
<td>100,000</td>
<td>2 – 4</td>
<td>0.2 – 0.3</td>
</tr>
</tbody>
</table>

* A suitable daily or weekly maintenance dose should start after completing the schedule.
** For conversion of ng/mL to nmol/L, multiply by 2.5.
*** Mentioned replacement doses can be taken as single cumulative doses or spread out through the week.
**** Estimated deficit of vitamin D needed to replenish body stores.

(Table adapted with permission from S.J. Wimalawansa)

Table 2. Vitamin D dosing in the absence of a baseline Vitamin D level

<table>
<thead>
<tr>
<th>Body-weight category</th>
<th>Dose (IU) kg/day</th>
<th>Dose (IU)/day</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Daily dose (IU)</td>
<td>Once a week</td>
</tr>
<tr>
<td>BMI ≤ 19 (under-weight)</td>
<td>40 – 70</td>
<td>≈ 2,000 – 4,000</td>
</tr>
<tr>
<td>BMI 20–29 (non-obese person)</td>
<td>70 – 100</td>
<td>≈ 5,000 – 7,000</td>
</tr>
<tr>
<td>BMI 30–39 (obese persons)</td>
<td>100 – 150</td>
<td>≈ 9,000 – 15,000</td>
</tr>
<tr>
<td>BMI ≥ 40 (morbidly obese persons)</td>
<td>150 – 200</td>
<td>≈ 16,000 – 30,000</td>
</tr>
</tbody>
</table>

(Table adapted with permission from S.J. Wimalawansa)

Clinical signs and symptoms can be grouped in the following clusters. The reason for this grouping is to allow organ specific targeted therapy/individualized therapy.

1. Respiratory: shortness of breath, congestion, persistent cough, etc.
2. Neurological/psychiatric: brain fog, malaise, tiredness, headaches, migraines, depression, inability to focus/concentrate, altered cognition, insomnia, vertigo, panic attacks, tinnitus, anosmia, phantom smells, etc.
3. Musculoskeletal: myalgias, fatigue, weakness, joint pains, inability to exercise, post-exertional malaise, inability to perform normal activities of daily life (ADL’s).
5. Autonomic: Postural tachycardia syndrome (POTS), abnormal sweating.
6. Gastrointestinal disturbance: Anorexia, diarrhea, bloating, vomiting, nausea, etc.
7. Dermatologic: Itching, rashes, dermatographia
8. Mucus membranes: Running nose, sneezing, Burning and itchy eyes.

Approach to Treatment

The treatment approach should be individualized according to the grouping of clinical signs and symptoms. However, in general, it is likely that patients who did not receive adequate antiviral treatment (e.g. ivermectin, etc.) during the acute symptomatic phase and adequate anti-inflammatory/macrophage repolarization therapy (e.g. corticosteroids, statins, omega-3 fatty acids, fluvoxamine, ivermectin, etc.) during the acute phase of COVID-19 are much more likely to develop the Post-COVID-19 Syndrome.

In patients with ongoing respiratory symptoms, chest imaging is suggested (preferably a chest CT scan). Those with unresolved pulmonary inflammation (organizing pneumonia with ground glass opacification) should be treated with a course of corticosteroids. Low-dose prednisolone/ methylprednisolone (10 mg/day) for six weeks is suggested. [521] However, the patients’ symptoms and CRP should be followed closely as a dose escalation may be required in those who respond poorly. An unknown number of patients who have recovered from COVID-19 organizing pneumonia will develop pulmonary fibrosis with associated limitation of activity. Pulmonary function testing demonstrates a restrictive type pattern with decreased residual volume and DLCO. [506] These patients should be referred to a pulmonologist with expertise in pulmonary fibrosis. Anti-fibrotic therapy may have a role in these patients, [473–476] however additional data is required before this therapy can be more generally recommended. As discussed above, the serotonin receptor blocker cyproheptadine may reduce the risk of pulmonary fibrosis. [364]

Similar to patients who have recovered from septic shock, [522] a prolonged (many months) immune disturbance with elevated pro- and anti-inflammatory cytokines may contribute to the LHCS. This is likely the consequence of monocyte activation syndrome and monocyte repolarization therapy is therefore indicated. Activated microglia may contribute to the neurological symptom’s characteristic of LHCS. A cytokine panel may allow targeted anti-inflammatory therapy (Maraviroc in patients with high CCR5 levels). It should be noted that much like omega-3 fatty acids, corticosteroids have been demonstrated to increase expression of pro-resolving lipids including Protectin D1 and Resolvin D4. [523]
Naltrexone is a well-known opioid antagonist used in chronic opiate abuse. Naltrexone is classically prescribed in daily doses of at least 50 mg taken orally. Paradoxically, low dose naltrexone (LDN) in a dose between 1 to 5 mg has been demonstrated to have anti-inflammatory, analgesic and neuromodulating properties. Specifically, LDN has been shown to reduce glial inflammatory response by modulating Toll-like receptor 4 signaling in addition to systemically upregulating endogenous opioid signaling by transient opioid-receptor blockade. [315,524] LDN typically in a dose of 4.5 mg has been used successfully to treat fibromyalgia, Crohn's disease, multiple sclerosis, and complex chronic pain syndromes as well as many chronic pain syndromes. [315,524] LDN may be particularly useful in the treatment of LHCS as is inhibits activated macrophages/monocytes and microglia. [524,525] Once activated, microglia produce inflammatory and excitatory factors that can cause sickness behaviors such as pain sensitivity, fatigue, cognitive disruption, sleep disorders, mood disorders, and general malaise; clinical features typical of those found with LHCS.

References

Disclaimer
The I-RECOVER protocol is borne of clinical experience only and thus is meant solely for educational purposes to health care providers regarding potentially beneficial empiric treatment approaches for Long Haul COVID-19 Syndrome. Never disregard professional medical advice because of something you have read on our website and releases. This is not intended to be a substitute for professional medical advice, diagnosis, or treatment in regards to any patient. Treatment for an individual patient is determined by many factors and thus should rely on the judgement of your physician or qualified health care provider. Always seek their advice with any questions you may have regarding your medical condition or health.

Please check our homepage www.flccc.net regularly for updates of our COVID-19 Protocols! – New medications may be added and/or dose changes to existing medications may be made as further scientific studies emerge.