Melatonin as adjuvant treatment for coronavirus disease 2019 pneumonia patients requiring hospitalization (MAC-19 PRO): a case series

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ABSTRACT

Treatment for coronavirus disease 2019 (COVID19) pneumonia remains empirical and the search for therapies that can improve outcomes continues. Melatonin has been shown to have anti-inflammatory, antioxidant, and immune-modulating effects that may address key pathophysiologic mechanisms in the development and progression of acute respiratory distress syndrome (ARDS), which has been implicated as the likely cause of death in COVID19. We aimed to describe the observable clinical outcomes and tolerability of high-dose melatonin (hdM) given as adjuvant therapy in patients admitted with COVID19 pneumonia. We conducted a retrospective descriptive case series of patients who: 1) were admitted to the Manila Doctors Hospital in Manila, Philippines, between March 5, 2020 and April 4, 2020; 2) presented with history of typical symptoms (fever, cough, sore throat, loss of smell and/or taste, myalgia, fatigue); 3) had admitting impression of atypical pneumonia; 4) had history and chest imaging findings highly suggestive of COVID19 pneumonia, and, 5) were given hdM as adjuvant therapy, in addition to standard and/or empirical therapy. One patient admitted to another hospital, who one of the authors helped co-manage, was included. He was the lone patient given hdM in that hospital during the treatment period. Main outcomes described were: time to clinical improvement, duration of hospital stay from hdM initiation, need for mechanical ventilation (MV) prior to cardiopulmonary resuscitation, and final outcome (death or recovery/discharge). Of 10 patients given hdM at doses of 36-72mg/day per os (p.o.) in 4 divided doses as adjuvant therapy, 7 were confirmed COVID19 positive (+) by reverse transcription polymerase chain reaction (RT-PCR) and 3 tested negative (-), which was deemed to be false (-) considering the patients’ typical history, symptomatology, chest imaging
findings and elevated bio-inflammatory parameters. In all 10 patients given hdM, clinical stabilization and/or improvement was noted within 4-5 days after initiation of hdM. All hdM patients, including 3 with moderately severe ARDS and 1 with mild ARDS, survived; none required MV. The 7 COVID19(+) patients were discharged at an average of 8.6 days after initiation of hdM. The 3 highly probable COVID19 patients on hdM were discharged at an average of 7.3 days after hdM initiation. Average hospital stay of those not given hdM (non-hdM) COVID19(+) patients who were admitted during the same period and recovered was 13 days. To provide perspective, although the groups are not comparable, 12 of the 34 (35.3%) COVID19(+) non-hdM patients admitted during the same period died, 7/34 (20.6%) required MV; while 6 of 15 (40%) non-hdM (-) by RT-PCR but highly probable COVID19 pneumonia patients also died, 4/15 (26.7%) required MV. No significant side-effects were noted with hdM except for sleepiness, which was deemed favorable by all patients, most of whom had anxiety- and symptom-related sleeping problems previously. HdM may have a beneficial role in patients treated for COVID19 pneumonia, in terms of shorter time to clinical improvement, less need for MV, shorter hospital stay and possibly lower mortality. HdM was well tolerated. This is the first report describing the benefits of hdM in patients being treated for COVID19 pneumonia. Being a commonly available and inexpensive sleep-aid supplement worldwide, melatonin may play a role as adjuvant therapy in the global war against COVID19.

Key words: melatonin, COVID19 pneumonia, adjuvant therapy, cytokine storm, immune system.

1. INTRODUCTION

The main cause of death in severe COVID19 is due to progressive acute lung injury (ALI) and acute respiratory distress syndrome (ARDS) (1, 2), likely triggered by excessive inflammatory and immune responses activating a so-called “cytokine storm” (2, 3). Treatment for COVID19 remains supportive and empirical, and the search for therapies that will improve outcomes continues (4).

Melatonin is an inexpensive sleep-aid supplement commonly available worldwide. It is known for its anti-inflammation (5-8), anti-oxidation (5) and immune-enhancing effects (9-11). It has been shown to reduce the pro-inflammatory cytokines tumor necrosis factor (TNF)-α, interleukin (IL)-1β, IL-6, and IL-8, and increase the level of the anti-inflammatory cytokine IL-10 (5-8). Safety of melatonin in humans up to doses exceeding 100mg/day has been demonstrated (5, 12, 13).

With the known rapidly progressive pathophysiology of severe COVID19 that could be possibly addressed by the anti-inflammatory, antioxidant and immune-modulatory effects of melatonin, this study determined the efficacy and tolerability of high-dose melatonin (hdM) as an adjuvant therapy in patients admitted for COVID19 pneumonia.

2. METHODS

We reviewed the charts of all patients who: 1) were admitted to the Manila Doctors Hospital in Manila, Philippines, between March 5, 2020 and April 4, 2020; 2) presented with history of typical symptoms (fever, cough, sore throat, loss of smell and/or taste, myalgia, fatigue); 3) had admitting impression of atypical pneumonia; 4) had history and chest imaging findings (ground glass opacities in both lung fields) highly suggestive of COVID19 pneumonia, and, 5) were given hdM as adjuvant therapy, in addition to standard and/or empirical therapy. One patient, who one of the authors helped co-manage and was admitted to the University of the
East Ramon Magsaysay Memorial Center, also in Metro Manila, was included in the case series. He was the lone patient given hdM in that hospital during the same period.

Main outcomes noted were: time to clinical improvement (symptomatic improvement or relief, stabilization and/or regression of lung infiltrates by chest x-ray or computed tomography scan, decrease in bio-inflammatory laboratory parameters), need for mechanical ventilation (MV), duration of hospital stay after hdM initiation, and final outcome (death, or recovery and discharge) in all patients given hdM.

To provide perspective, although the groups are not comparable, the mortality rates of hdM patients and those not given hdM (non-hdM) were also determined.

3. RESULTS

Of the 58 patients admitted to our institution between March 5, 2020 to April 4, 2020 for recent history of flu-like symptoms (fever, cough, sore throat, loss of smell and/or taste, myalgia, fatigue) and chest imaging findings of ground glass opacities highly suggestive of COVID19 pneumonia, 9 (15.5%) were given hdM—3 patients on admission and the 6 others, during the course of their hospital stay when no clinical improvement was observed with the standard and empirical treatments they were receiving. HdM was administered at doses between 36 mg/day to 72 mgs/day per os (p.o.) in 4 divided doses. Another COVID19 positive (+) patient identified by reverse transcription polymerase chain reaction (RT-PCR), admitted to another hospital, who one of the authors helped co-manage, was also included in this case series. He was the lone patient given hdM in that hospital during the same period. Of the total of 10 patients given hdM, 7 were confirmed RT-PCR(+), while 3 tested negative(-), but considering their history and clinical presentation including chest imaging findings, treatment for COVID19 pneumonia was continued. Three of the 7 given hdM had moderately severe ARDS (pO2/FiO2 < 200 mm Hg). The 10 patients given hdM had high-risk features: >60 years of age, or younger but with established cardiovascular disease or other comorbidities including diabetes mellitus, hypertension, chronic obstructive pulmonary disease, chronic kidney disease, bronchial asthma and obesity (Table 1).

Table 1. Patient demographics, comorbidities, laboratory findings.

<table>
<thead>
<tr>
<th>P</th>
<th>H</th>
<th>Age/ Sex</th>
<th>Comorbidities</th>
<th>Max O2 supply</th>
<th>Chest Xray</th>
<th>CT scan</th>
<th>V</th>
<th>ABG (worst)</th>
<th>CBC</th>
<th>Procal</th>
<th>Inflam I (LDH, CRP, Ferritin)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>8</td>
<td>64 F</td>
<td>HTN</td>
<td>Face Mask 10 lpm</td>
<td>Pulmonary Edema</td>
<td>Diffusely distributed GGO (crazy paving) pattern and areas of beginning consolidation on both lungs</td>
<td>+</td>
<td>pH 7.45</td>
<td>Hgb 115</td>
<td>LDH 454(451)(525)(540)</td>
<td>hsCRP 30.8</td>
</tr>
<tr>
<td>2</td>
<td>8</td>
<td>64 F</td>
<td>HTN</td>
<td>NC 2 lpm</td>
<td>Bilateral Pneumonia</td>
<td>GGO diffusely scattered in both lungs</td>
<td>+</td>
<td>pH 7.423</td>
<td>Hgb 129</td>
<td>hsCRP 16.87</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>5</td>
<td>68 F</td>
<td>HCVD</td>
<td>NC 2 lpm</td>
<td>Bilateral Infiltrates</td>
<td>Consolidated GGO scattered in both lung fields</td>
<td>+</td>
<td>pH 7.45</td>
<td>Hgb 120</td>
<td>LDH 296</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>6</td>
<td>61 F</td>
<td>(-)</td>
<td>NC 1 lpm</td>
<td>Bilateral infiltrates</td>
<td>Bilateral parenchymal GGO and consolidation</td>
<td>+</td>
<td>pH 7.45</td>
<td>Hgb 129</td>
<td>LDH 183</td>
<td></td>
</tr>
</tbody>
</table>

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| 5 | 9 | 26/ M | HPN spCVD BAIAE | NC 2 lpm | Bilateral infiltrates | Patchy GGO with mixed consolidation in right middle lobe and inferior lingula | pH 7.494 | pCO2: 34.4 | pO2: 104 | O2 sat: 97 | PF ratio: 371 | Hgb 157 | Wbc 3.07 | Neu 67 | Lymp 29 | PC 222 | 0.06 | LDH 294 (348)(268)(208) (215) | hsCRP 0.13 (6.35)(13.29) Ferritin 1154 (757)(663)(389) |
| 7 | 11 | 76/ M | HCVD, CAD, BANIAE | NC 6 lpm | Reticular infiltrates mid to LLLF and RLLF | GGO bilateral upper and Right middle lobe and lingula | pH 7.518 | pCO2: 28.8 | HCO3: 23.2 | pO2: 65.4 | O2 sat: 93.5 | PF ratio: 458 | Hgb 137 | Wbc 7.05 | Neu 73 | Lymp 9 | PC 218 | <0.05 | LDH 212(268)(277)(182) hsCRP 118 (129)(109)(38) Ferritin 869 (852)(627) |
| 8 | 5 | 62/ F | HTN | Room air | Reticular infiltrates on both lower lobes with patchy GGO in RUL | Few reticular and GGO in the superior segment, anterior and medial basal segment of the Right lower lobe and posterior segment of the left lower lobe. | pH 7.438 | pCO2: 37 | HCO3: 24.6 | pO2: 93 | O2 sat: 96.9 | PF ratio: 465 | Hgb 141 | Wbc 6.26 | Neu 47 | Lymp 40 | PC 306 | <0.05 | LDH 212 in/L hsCRP 1.14 Ferritin 108 |
| 9 | 11 | 54/ F | HTN T2DM BANIAE Microvascular ischemia with paroxysmal arrhythmia | NC 3 lpm | Bilateral infiltrates | Multifocal reticular and GGO with signs of early peripheral consolidation scattered in both lung fields. | pH 7.42 | pCO2: 35 | HCO3: 22 | pO2: 53 | O2 sat: 88 | PF ratio: 252 | Hgb 140 | Wbc 8.78 | Neu 63 | Lymp 25 | PC 338 | <0.05 | LDH 240 hsCRP 1.47 Ferritin 393 |
| 10* | 9 | 70/ M | HTN T2DM COPD (current smoker) | Face mask 10 lpm | Interstitial hazed opacities in the bilateral mid lung fields; infiltrates in the right lower lung | GGO, both lungs, bilateral subsegmental atelectatic changes; enlarged mediastinal lymph nodes | pH7.46 | pCO2: 33 | HCO3: 23.5 | pO2: 72 | O2 sat: 95 at 8 lpm by NC | PF ratio: 138 | Hgb 141 | Wbc 3.9 | Neu 60 | Lymp 40 | PC 170 | 0.11 | LDH 277 hsCRP 23.25 Ferritin 3,190 |


Of the 58 patients admitted to our institution for COVID19 pneumonia during the same period, 40 (69%) were confirmed by RT-PCR to be COVID19 (+), while 18 (31%) were (-). All patients were given standard and/or empirical treatment as deemed appropriate by the attending medical teams — composed of an infectious disease specialist, a pulmonologist, and other referral consultants depending on the patients’ comorbidities. The treatment regimen consisted of any of the following as needed: supplemental oxygen, MV, agents to mitigate cytokine storm, oral or parenteral antibiotics, anti-viral agents; vasopressor support, renal-replacement therapy, and hemoperfusion/hemadsorption. (Table 2).
Clinical stabilization and/or improvement was noted within 4-5 days after initiation of hdM in all COVID19(+) patients and highly probable but COVID19 (-) patients given the supplement. None required MV. All 10 patients given hdM survived and the 7 COVID19(+) patients were discharged clinically improved and asymptomatic at an average of 7.3 days after hdM initiation. The average hospital stay of non-hdM COVID19(+) patients who recovered was 13 days.

To provide perspective, although both groups are not comparable, 12 of the 34 (35.3%) COVID19(+) non-hdM patients admitted in our institution during the same period died, 7/34 (20.6%) required MV; while 6 of 15 highly probable but COVID19 (-) non-hdM patients (40%) died, 4/15 (26.7%) required MV.

No significant side-effects were noted in those given hdM except for increased sleepiness.

4. DISCUSSION

Published research suggests the versatility of melatonin as an anti-inflammatory, antioxidant and immuno-modulating agent (5-11). The beneficial effects of hdM as adjuvant therapy in cancer (13-15), severe viral influenza (5, 8) and sepsis (8) have also been demonstrated. Melatonin increases p53/p21Waf1 (wild type p53/21 activated fragment) expression causing anti-proliferative effects in breast cancer cells (16). Moreover, other mechanisms were also identified such as inducing apoptosis in MCF (Michigan Cancer Foundation)-7 human breast cancer cells, and anti-estrogenic effects to Erα signaling pathway (17). In 1997, Lisson et al. (18) published a study wherein breast cancer patients were given melatonin 20 mg/day at bedtime as adjunctive treatment for chemotherapy. The authors concluded that patients given melatonin had less thrombocytopenia, stomatitis and neuropathy, with no significant adverse reactions (19).

In several animal models, melatonin protects against bacterial, viral and parasitic infection through immunomodulation and/or direct or indirect anti-oxidant activity (20). The production

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Table 2. Dose of melatonin and other drugs given.

<table>
<thead>
<tr>
<th>P</th>
<th>Dose of Mel</th>
<th>Mel use at admission</th>
<th>Blood Culture</th>
<th>Antibiotics</th>
<th>Chl/ HChl</th>
<th>Azithro-mycin</th>
<th>Lopinavir/ritonavir</th>
<th>Oselta-mivir</th>
<th>Na ascorbate</th>
<th>Zinc</th>
<th>IL-6 inhibitor</th>
<th>Hemoperfusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>60 mg/day</td>
<td>Day 1</td>
<td>No Growth</td>
<td>Pip-Tazo</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>2</td>
<td>36 mg/day</td>
<td>Day 1</td>
<td>No Growth</td>
<td>Pip-Tazo</td>
<td>HChl</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td></td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>3</td>
<td>48 mg/day</td>
<td>Day 4</td>
<td>No Growth</td>
<td>Meropenem</td>
<td>Chl</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>4</td>
<td>36 mg/day</td>
<td>Day 11</td>
<td>No Growth</td>
<td>Meropenem</td>
<td>Pip-Tazo</td>
<td>Chl</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>5</td>
<td>60 mg/day</td>
<td>Day 3</td>
<td>No Growth</td>
<td>Meropenem</td>
<td>HChl</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>6</td>
<td>36 mg/day</td>
<td>Day 1</td>
<td>No Growth</td>
<td>Pip-Tazo</td>
<td>HChl</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td></td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>7</td>
<td>36 mg/day</td>
<td>Day 7</td>
<td>No Growth</td>
<td>Meropenem</td>
<td>Pip-Tazo</td>
<td>HChl</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>8</td>
<td>48 mg/day</td>
<td>Day 1</td>
<td>No Growth</td>
<td>Ceftriaxone</td>
<td>HChl</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td></td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>9</td>
<td>72 mg/day</td>
<td>Day 1</td>
<td>No Growth</td>
<td>Ceftriaxone</td>
<td>HChl</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td></td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>10</td>
<td>60 mg/day</td>
<td>Day 4</td>
<td>No Growth</td>
<td>Pip-Tazo</td>
<td>HChl</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td></td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

of reactive oxygen species (ROS) is one of the lethal effects of any severe infection. Melatonin has shown beneficial effects in sepsis by decreasing the synthesis of pro-inflammatory cytokines and suppressing inducible nitric oxide (NO) synthase gene expression (21). In a study done by Carrillo-Vico et al. administration of melatonin (10mg) to mice with lipopolysaccharide induced inflammation showed 90% survival. This may be due to the increased levels of anti-inflammatory cytokine interleukin (IL)-10 and decreased concentration of proinflammatory mediators such as TNF, IL-12, and interferon-γ (21). A study in septic newborns performed by Gitto et.al. showed that malondialdehyde (MDA), nitrite production and 4-hydroxynonenals (HDA) were significantly reduced in those patients given melatonin. Moreover, melatonin administration also improved the clinical outcome of those septic newborns (22).

This is the first report on the actual use of hdM in humans afflicted with COVID19 pneumonia. Melatonin is not a direct viricidal agent; but it may help neutralize the deleterious effects of the SARS-CoV-2 that causes COVID19 (5). It may particularly mitigate the maladaptive immune-modulatory response that releases pro-inflammatory cytokines and chemokines triggering a ‘cytokine storm’ that causes extensive multiorgan damage particularly in the lungs (5-8). Zhang et al. postulated that following SARS-CoV-2 infection, ALI/ARDS may result due to a suppressed immune response, excessive inflammation and uncontrolled oxidation resulting in a cytokine storm (5). Melatonin may exert a beneficial role as adjuvant therapy in the regulation of the immune system, inflammation and oxidation stress, to mitigate the complications of ALI/ARDS and related multi-organ complications (5) (Figure 1).

Fig. 1. Pathogenesis of COVID-19 and potential adjuvant use of melatonin.

Zhang R et al. (5) postulated that following SARS-CoV-2 infection, ALI/ARDS may occur due to a suppressed immune response, excessive inflammation and uncontrolled oxidation activating a cytokine storm. Melatonin may exert a beneficial role as adjuvant therapy in the regulation of the immune system, inflammation and oxidation stress, to mitigate the complications of ALI/ARDS and related multi-organ complications. ALI: Acute lung injury; ARDS: Acute respiratory distress syndrome. (Figure is reproduced from (5) with permission).

Secondary hemophagocytic lymphohistiocytosis (sHLH) is a hyperinflammatory syndrome characterized by a fatal hypercytokemia with multiorgan failure. Clinically, it is manifested by unremitting fever, cytopenias, hyper-ferritinemia and in half of patients, ARDS (2). This sHLH
closely resembles the cytokine storm associated with COVID19. In the early stage of coronavirus infection, dendritic and epithelial cells are activated and express a cluster of pro-inflammatory cytokines and chemokines which is manifested as an increase in IL-1β, IL-2, IL-6, IL-8, both IFN-α/β, TNF, CeC motif chemokine 3 (CCL3), CCL5, CCL2, and IP-10 (5) which then contributes to the development of the disease. IL-10 is antiviral; and usually in coronavirus infection, this interleukin is decreased (23, 24). However, in some COVID19 patients, elevated levels of IL-10 have been noted. It is still unknown whether this is a unique feature of COVID19 (24). A marked decrease in IL-10, results in amplification of inflammatory response which then promotes cellular apoptosis or necrosis of the affected cells that lead to increased permeability of blood vessels and aberrant accumulation of inflammatory monocytes, macrophages and neutrophils in the lung alveoli causing ALI. In a study by Smits et.al, they noted that the severity of ALI was accompanied by an elevated expression of inflammatory cytokine, IFN β, rather than the viral load (25). Another study by Channappanavar et.al, noted that the depletion of inflammatory monocytes caused an increased survival rate of coronavirus host without a change in viral load (26). This signifies that if the same mechanism is involved in COVID19, and inhibiting several cytokines may lead to improved outcomes. This is where melatonin may play a big role in COVID19.

Melatonin has anti-inflammatory effects through different pathways. Sirtuin-1 (SIRT1) mediates melatonin’s anti-inflammatory action by inhibiting high mobility group box 1 protein 1 (HMGB1) leading to attenuation of pro-inflammatory macrophages (27). In a study done by Wang et al., regulation of SIRT1 in sepsis-induced ALI reduced lung injury and inflammation. Another mechanism of melatonin is through nuclear factor kappa-B (NF-kB). NF-kB is an inflammatory mediator in ALI and melatonin has been shown to downregulate NF-kB in T cells and lung tissues (28, 29). Moreover, melatonin has an anti-oxidative effect through upregulating anti-oxidative enzymes and downregulating pro-oxidative enzymes (5). In viral infections, virus replication generates oxidized products. In SARS-induced lung injury, there is a production of oxidized low-density lipoprotein which then activates innate immune response causing overproduction of IL-6 in alveolar macrophages via toll-like receptor 4 (TLR4) leading to ALI (29). TLR4 is a therapeutic target for melatonin. In a study by Zhao et al., melatonin was shown to exert an anti-inflammatory and antioxidant effect reducing damage to the white matter after focal brain ischemia in rats by regulating the TLR4/NF-κB pathway (30). Thus, it is likely that administration of melatonin will be beneficial in attenuating the inflammation and oxidation in COVID19 patients.

In Patients 3, 4, 5, 7, 10 who received hdM, the supplement was added to the regimen when clinical improvement was not noted with the standard and empirical drugs the patients were taking (Table 1, 2). In those patients, a temporal relationship between hdM initiation and clinical improvement was observed. Symptomatic improvement was observed after 48-72 hours, soon followed by a decrease in bio-inflammatory parameters.

The regression of ground glass opacities (infiltrates) on chest imaging was observed last. The most frequent chest x-ray and computerized tomography scan findings of interstitial changes and ground glass opacities (GGOs) may occur early in the course of COVID19 pneumonia and have been observed to peak around 10 to 14 days after the onset of symptoms (31-35). However, the radiologic changes may persist even when the patients have already improved clinically and even up to the time of discharge (34). This was noted in most of the patients in this series despite the marked symptomatic relief and decrease in bio-inflammatory parameters. In Patient No. 1 in the series, a 64-year old male diabetic, hypertensive with chronic kidney disease and chronic gout, who had moderately severe ARDS on admission, serial chest x-rays showed bilateral infiltrates/GGOs persisting up to discharge but showing some regression (Figure 2). However, patient was already symptom-free by the fifth hospital day.
with decreasing bio-inflammatory parameters. Thus, regression of chest imaging abnormalities could not be correlated with clinical improvement due to a radiologic lag when compared with other clinical parameters.

It may also be noteworthy that the 10 patients given hdM in this case series all had high risk features: > 60 years of age, or < 60 years of age and with established cardiovascular disease or comorbidities, which have been associated with an increased risk for mortality in COVID19. Three of the 10 hdM patients had moderately severe ARDS, and another one had mild ARDS. None of the 10 given hdM required MV.

In Wuhan, China, age and comorbidities were highly contributory factors among those who died from COVID19 (36, 37). A retrospective, multicenter cohort study done by Zhou et al. which included 191 adult in-patients (≥18 years old) with laboratory confirmed COVID19 from Jinyintan Hospital and Wuhan Pulmonary Hospital (Wuhan, China) showed that the following factors: older age (odds ratio 1·10, 95% CI 1·03–1·17, per year increase; p=0·0043), higher Sequential Organ Failure Assessment (SOFA) score (5·65, 2·61–12·23; p<0·0001), and d-dimer greater than 1 µg/mL (18·42, 2·64–128·55; p=0·0033) on admission were associated with in-hospital mortality (36, 37). This was attributed to the age-dependent defects in T-cell and B-cell function coupled with the excessive production of type 2 cytokines that promoted viral replication and more pronounced pro-inflammatory responses, contributing to an unfavorable outcome. Moreover, the aging process especially when accompanied by comorbidities such as diabetes mellitus and hypertension could contribute to severe infection and even death. The chronic inflammatory and prothrombotic effects seen with aging can also be likely predisposing factors, aggravated by the excessive cytokine responses to a severe systemic infection (38). Major cytokines, particularly IL-6, and levels of d-dimer, which is identified as a marker of thrombin formation and active fibrinolytic activity, could also be implicated as contributing factors that progressively increase the disease process and delay recovery rate in the elderly population.

Figure 2. The representative x-ray graphs of a COVID19 patient before and after melatonin as a adjuvant treatment.

Serial chest x-rays of Patient No.1, a 64-year old male hypertensive with T2DM, CKD, chronic gout, with moderate ARDS and PFR of less than 160 mmHg. Follow-up chest x-ray showed regression of ground glass opacities after melatonin treatment but this finding persisted until discharge although patient was already symptom-free. PFR: paO2/FIO2 ratio (Courtesy of Department of Radiology, Manila Doctors Hospital).
An ambispective cohort study by Li et al., that included 548 hospitalized patients with COVID19, showed that almost half of the cases were severe (39). Compared to non-severe conditions, prevalence of comorbidities was higher among those with severe conditions, namely: diabetes (19.3% vs. 11.1%, p=0.009), hypertension (38.7% vs. 22.2%, p<0.001) and coronary artery disease (10.4% vs. 2.2%, p<0.001). In this case series, among the 10 patients included, 8 had hypertension, 3 had diabetes and 2 had coronary artery disease.

A retrospective cohort study of Wu et al., involving 201 patients with confirmed COVID19 pneumonia admitted to Wuhan Jinyintan Hospital, looked into risk factors in the development of ARDS and death (40). The risk factors associated with the development of ARDS and progression from ARDS to death shown in the study were older age, neutrophilia, higher lactate dehydrogenase and D-dimer. The authors also considered immune response dysfunction leading to a cytokine storm as one possible mechanism of pathogenesis of COVID19 (40).

Three of the 10 hdM patients in these case series had moderately severe ARDS, and another one had mild ARDS. None of the 10 given hdM required MV. Due to the patients’ advanced age and oftentimes presence of comorbid conditions, the pro-inflammatory complications of COVID19 may have devastating effect unless adequately addressed. Due to its anti-inflammatory, anti-apoptotic and anti-oxidant properties, melatonin has a potential role as one of the treatment modalities in COVID19 (5, 41). Melatonin modulates the immune system response, reduces the pro-inflammatory cytokines levels (41) and enhances T helper immune responses (42). Melatonin, in vitro, has also been shown to significantly reduce the population of Th1 CD4 lymphocytes, increase the expression of anti-inflammatory cytokine IL-10 together with the population of IL-10-producing CD4 T cells (9).

HdM, up to a dose of 72 mg/day in 4 divided doses was well-tolerated in this case series. The only side-effect reported was increased sleepiness, which was deemed favorable by all patients, most of whom had anxiety- and symptom-related sleeping problems previously. Melatonin’s encouraging safety profile in this case series is in keeping with literature on exogenous melatonin use. Experimental studies on mice during gestation have shown that very high doses of up to 200 mg/kg/day did not show any significant embryo/fetal toxicity (43). Andersen et al. detailed the possible risks of melatonin in specific patient groups and found little to no side effects even at doses given at 1 gm/day (44). To date, no studies have reported serious adverse effects of melatonin use. Even in extreme doses, short-term (days of administration) use of melatonin in both animal and humans has been consistently shown to be safe, with only mild side effects recorded such as dizziness, headache, nausea, and sleepiness (45, 46). Randomized trials have likewise recorded mild side effects comparable to placebo with long-term (years) of administration of melatonin in adults (47-49). Animal or human toxicity despite high-dose melatonin administration has not been documented (5, 43, 44, 50). There are no reports or studies elucidating neither the positive nor adverse effects of melatonin in pregnancy or lactation. In the elderly, anxiolytic and hypnotic effects of melatonin are regarded as beneficial (51).

There is a paucity of information on the use of melatonin in high-risk groups, and most studies consider the increase in sleepiness a welcome respite from the lack or interference of sleep in critically ill patients (47, 48, 50, 52, 53). A recent randomized trial reported no adverse effects or complications, and a trend towards a shorter ICU stay and total hospital stay after liver resection was noted with a single dose of 50 mg/kg body weight dissolved in 250 mL of milk via gastric tube given before surgery (54).

Melatonin has not caused serious adverse effects in studies of ALI/ARDS in mouse models (55, 56). However as with most research on melatonin for varying clinical scenarios, conclusions for its use in humans may only be extrapolated from translational research in animals, and from the limited yet indirect safety studies in humans that populate the literature (5). More studies are needed to establish concrete safety guidelines in the use of high-dose
melatonin in humans. Specifically, the effects of hdM in COVID19 pneumonia have yet to be elucidated.

In conclusion, high-dose melatonin may have a beneficial role as adjuvant therapy in patients being treated for COVID19 pneumonia, in terms of shorter time to clinical improvement, less need for intubation and MV, shorter hospital stay, and possibly lower mortality. It was well tolerated. Being a commonly available and inexpensive sleep-aid supplement worldwide, it may play a role as adjuvant therapy in the global war vs COVID19. We are preparing to undertake soon a bigger multicenter randomized clinical trial to validate the observations in this report.

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AUTHORSHIP

Rafael Castillo: Conception and design, analysis and interpretation of data, drafting the manuscript and revising it critically for important intellectual content, gave final approval of version to be published; Gino Rei Quizon: Acquisition of data, analysis and interpretation of data, revision of manuscript, gave final approval of version to be published; Felix Eduardo Punzalan: Analysis and interpretation of data, revision of manuscript, gave final approval of version to be published; Dante Morales: Conception and design, or analysis and interpretation of data, gave final approval of version to be published; Mario Joselito Juco: Acquisition of data, analysis and interpretation of data; Arthur Dessi Roman: Acquisition of data, analysis and interpretation of data; Donnah de Leon: Acquisition of data, analysis and interpretation of data; Rafael Bien Guingon: Acquisition of data, analysis and interpretation of data, gave final approval of version to be published. Dun Xian Tan, identification and interpretation of data, suggestions for information to be included in the paper and minor editing of the manuscript.

All authors agreed to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.
CONFLICT OF INTEREST

Rafael R. Castillo: member of speakers bureau and advisory board of Servier, Boehringer-Ingelheim, Menarini, LRI-Therapharma, AstraZeneca, Sanofi, UAP Pharma; officer and owns stocks in Trianon International and FAME Publishing with salary donated to a charitable foundation; Dante Morales: member of Board of Directors of Manila Doctors Hospital, international research grants from AstraZeneca, Bayer Pharma, PHRI Canada; chairman/member of advisory board and speakers’ bureau of Astra Zeneca, LRI Therapharma, Bayer Pharma, Menarini, Pfizer, Sanofi; Mario Juco: medical director of Manila Doctors Hospital; Arthur Dessi Roman: member of speakers bureau of Sanofi, Unilab, Natrapharm, MSD, Terrumo; Felix Eduardo Punzalan: member of speakers bureau of Sanofi, Servier, Aspen, Novartis; member of advisory board of Merck, GX Pharma; Donnah de Leon, Rafael Bien Guingon, Dun Xian Tan and Russel J Reiter have no potential conflicts of interest.

REFERENCES


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