ORIGINAL RESEARCH

Implications for Systemic Approaches to COVID-19: Effect Sizes of Remdesivir, Tocilizumab, Melatonin, Vitamin D3, and Meditation

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Introduction: COVID-19 poses a chronic threat to inflammatory systems, reinforcing the need for efficient anti-inflammatory strategies. The purpose of this review and analysis was to determine the efficacy of various interventions upon the inflammatory markers most affected by COVID-19. The focus was on the markers associated with COVID-19, not the etiology of the virus itself.

Methods: Based on 27 reviewed papers, information was extracted on the effects of COVID-19 upon inflammatory markers, then the effects of standard treatments (Remdesivir, Tocilizumab) and adjunctive interventions (vitamin D_3 , melatonin, and meditation) were extracted for those markers. These data were used to approximate effect sizes for the disease or interventions via standardized mean differences (SMD).

Results: The data that were available indicated that adjunctive interventions affected 68.4% of the inflammatory markers impacted by COVID-19, while standard pharmaceutical medication affected 26.3%.

Discussion: Nonstandard adjunctive care appeared to have comparable or superior effects in comparison to Remdesivir and Tocilizumab on the inflammatory markers most impacted by COVID-19. Alongside standards of care, melatonin, vitamin D₃, and meditation should be considered for treatment of SARS-COV-2 infection and COVID-19 disease.

Keywords: COVID-19, inflammation, cytokine storm, effect size, antivirals, corticosteroids, melatonin, vitamin D, meditation, whole health

Introduction

The COVID-19 pandemic has had pervasive adverse consequences, especially in the United States.¹ The combination of a highly contagious virus with significant hospitalization and mortality rates has resulted in devastating loss of human life and social stability. These costs continue to increase as research uncovers wider and more serious consequences for those who survive the initial infection, ranging from autoimmune disorders to fatal cardiopulmonary events.² The diverse and unpredictable nature of these complications point to a disease that is more complex than a respiratory virus.³ It is increasingly apparent that SARS-CoV-2 infection can lead to an inflammatory disorder affecting numerous bodily systems, with lasting effects on systemic inflammation that will linger long after the virus is eliminated.

Evidence suggests that approaching this systemic disease with systemic adjunctive interventions such as vitamin D_3 , melatonin, and meditation could prove more effective than targeting acute symptoms solely with pharmacological means such as

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Remdesivir and Tocilizumab.^{4–6} The purpose of this work is to track the effects of various catalysts, including Remdesivir, Tocilizumab, melatonin, meditation, vitamin D_3 , and COVID-19 itself, on a variety of inflammatory markers. The initial hypothesis explores whether adjunct interventions show any significant effects on inflammatory markers that Remdesivir and Tocilizumab do not affect. If so, that would imply those interventions may be viable complement in the treatment of COVID-19.

Inflammatory Model of Disease

A significant body of evidence suggests that acute and chronic inflammation accompanies and underlies many noncommunicable diseases.⁷ Inflammation is critical to a healthy host's defense against infection and injury, however, when continually activated it contributes to the ongoing pathophysiology of many chronic diseases. A persistent, low-level inflammatory response affects the innate immune system, adaptive immune system, and inflammatory mediators.⁸ The dynamic mechanisms of chronic inflammation can lead to tissue damage, oxidative stress, angiogenesis, and fibrosis in a variety of tissues, including the nervous system.⁹

Many of the most damaging effects of COVID-19 are related to cascading systemic inflammation and resulting cytokine storms, affecting respiratory, cardiac, nervous, gastrointestinal, and immune systems concurrently.^{10,11} Patterns of comorbidity demographics indicate that COVID-19 both initiates inflammatory dysregulation and exacerbates existing systemic inflammation.¹² The prevalence and magnitude of stressors contributing to preexist-ing inflammation make a catalyst like COVID-19 especially dangerous.¹³

COVID Reveals Lingering Vulnerability

Though vaccines have been developed to eventually curb the spread of the virus itself, there are already more than 135 million infected so far, a number that is expected to rise for months.¹⁴ It is important to note that the elimination of the virus, whether in an individual host or in a population, does not eliminate the increased health risks. Some individuals who have fully cleared the virus still exhibit significantly impaired cardiopulmonary, central nervous, and peripheral nervous systems.¹⁵ This suggests an eventual potential spike in pulmonary fibrosis, cardiopulmonary events, neuronal damage, Alzheimer's disease, Parkinson's disease, and multiple sclerosis.^{16–18} In addition to direct health impacts, the COVID-19 pandemic has highlighted the vulnerabilities of healthcare systems. The heavy load on hospitals was likely exacerbated by high rates of metabolic syndrome, a low-grade inflammatory condition that is suspected to afflict 10–40% of the developed world's population and correlates to COVID-19 severity and comorbidities.¹⁹ Lengthy recovery times and high prevalence of comorbidity resulted in overcrowded hospitals with rates of readmission as high as 1/6.²⁰ If COVID-19 were to mutate enough to escape the vaccines or if a new inflammatory pathogen emerges, the population is currently even more vulnerable than at the beginning of the current pandemic, with few effective mitigating interventions.

Ending the spread of COVID-19 will not address the health needs of the survivors, nor erase the conditions that exacerbated the pandemic. In fact, authorities have predicted that the long-term public health impact of COVID-19 will be profound; and in December 2020, Congress provided \$1.15 billion in funding for the National Institutes of Health to support research into the prolonged health consequences of SARS-CoV-2 infections.²¹

Mechanism of Action: COVID-19

While the inflammatory reaction and release of cytokines in healthy individuals is a necessary and normal reaction to injury and infection, in the case of inflammatory diseases cascading overproduction of cytokines can occur. COVID-19 often dysregulates pro-inflammatory chemokines that continue to stimulate this reaction until multiorgan damage occurs, often with fatal consequences for the infected individual.²² The mechanism suspected in severe occurrences of the SARS-CoV2 virus is the cytokine storm, an event involving hyperactivity of immune cells that produce pro-inflammatory cytokines, referred to as hypercytokinemia, which can lead to severe or even life-threatening cases of multiple organ failures.²³

Even though inflammatory symptoms may be more visible in particular parts of the body, the nature of cytokine storms is systemic, not acute or localized. Once chemokine and cytokine interactions have begun to spiral out of homeostasis, a system-wide feedback loop can be established which triggers effects in an extremely diverse set of bodily functions and organs.²⁴ Pulmonary complications, blood clots, and neurological issues are closely interwoven with systemic inflammation, making treatment of individual symptoms difficult.²⁵

Elevated TGF-beta plasma levels have been observed in respiratory diseases such as COPD and COVID-19 and maintain a central role in the pathophysiology of lung fibrosis during inflammation.^{26,27} In COVID-19 patients, TGF-beta upregulation contributes to cytokine release syndrome, pulmonary edema and acute respiratory distress syndrome via increased capillary permeability.^{28–31} The complement system further drives TGF-beta release from platelets to promote systemic inflammation and coagulopathy.³² SARS-CoV-2 infections also promote increased MAPK pathway mediated inflammation.^{33,34}

Figure 1 was created by this research team to outline the cellular and systemic progression of COVID-19 and the inflammatory response:

Mechanism of Action: Systemic Adjunct Interventions

The type of systemic dysregulation that both fuels the worst cases of COVID-19 and poses a lingering risk has been the subject of significant research relating to the

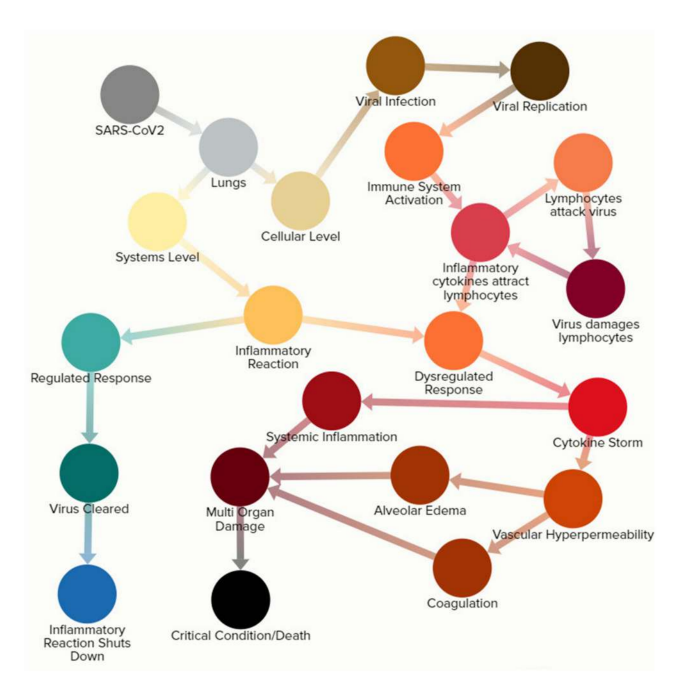


Figure 1 Progression and feedback loops associated with cytokine storms. Note: Courtesy of the Whole Health Institute.

efficacy of non-pharmacological adjunct interventions.³⁵ A particular area of focus has been referred to as "integrative medicine," which incorporates systems-level approaches to health that include behavioral changes, mental practices, and nutritional adjustments, interventions which have been associated with significant improvements in stress levels, pain, cardiovascular health, and metabolic health.³⁶ Examples of these adjunct interventions that are relevant to COVID-19 treatment include meditation, vitamin D₃ (cholecalciferol), and melatonin.^{37–39}

Targeted modulation of key cytokines such as TGF-beta with natural products such as artemisinin, an antimalarial lactone derived from *Artemisia annua*, may potentially help prevent pulmonary fibrosis in patients with respiratory diseases such as COVID-19 and could act as a potent cytokine-modulating adjunctive therapy as well, but insufficient data was present to include artemisinin or TGF-beta in this analysis.^{40–43}

The mechanisms of action for these interventions are still being studied, but existing evidence suggests regulating influences that have significant effects on dysregulated inflammatory processes. Vitamin D_3 , for example, has been shown to improve immune function while also curbing cascading inflammation, with numerous studies linking it to reduced severity and deaths in COVID-19 cases.^{44–46} The mechanism(s) behind melatonin is slightly less understood but potentially has the intriguing benefit of upregulating antiviral function while downregulating runaway inflammation.⁴⁷

Meditation has been linked to diverse psychological and physiological changes that improve homeostasis and diminish inflammatory feedback loops, though its mechanisms are still widely debated.⁴⁸ Theoretical modeling suggests behavioral health practices such as meditation and yoga may provide some degree of adjunctive efficacy against general infectious conditions through enhancements of endogenous levels of melatonin and other antiinflammatory or immuno-modulating processes.⁴⁹ This model was expanded to directly relate to COVID-19, with multidisciplinary evidence supporting the premise.

None of these adjunct interventions have been linked to any dangerous side effects during normal administration. It is important to note that the efficacy of an adjunct treatment does not replace or disprove standards of care. Such interventions are intended to supplement medical best practices, not replace them.

Central Role of Systems Approach

The systemic nature of cascading inflammation makes it difficult to treat with interventions that are more targeted in nature. Many of the pharmacological interventions that are capable of significant system-wide effects, such as the leading COVID-19 medications Remdesivir and Tocilizumab, are often accompanied by negative side-effects.^{50,51} Therefore, it is imperative to establish adjunctive interventions that can address the broader, holistic factors behind COVID-19 and do not result in dangerous side effects or numerous contraindications.

While a reductionist model is ideal for acute conditions, COVID-19 and its complications require a systems model that incorporates multiple levels of connection, causality, and interaction. There is insufficient data to calculate a full systems analysis of COVID-19 and potential interventions, but a mapping of the effects of the disease and its treatments is an important step toward understanding the most effective way to address this pandemic and any others like it. At the time of the writing of this research no such systems model for COVID-19 is known.

Methodology

The purpose of this article is to conduct a review and comparative analysis of the effect sizes of COVID-19, Remdesivir, Tocilizumab, meditation, vitamin D_3 , and melatonin on a variety of inflammatory markers.

Data

Figure 2 reflects the compilation process for this research, through which a collection of search terms was designed to capture a wide sampling of sources related to quantitative interactions between inflammatory processes/markers and COVID-19, Remdesivir, Tocilizumab, melatonin, meditation, and vitamin D₃. Remdesivir and Tocilizumab were selected for analysis as a result of being the predominant pharmacological intervention for COVID at the time of data-gathering. Vitamin D₃ and melatonin were included as adjunctive interventions due to their repeated use in COVID infections, while meditation was also included as an adjunct intervention to analyze a potentially anti-inflammatory intervention that did not introduce external properties to the subject.⁵² Though there are multiple meditative practices, the meditation studies included in this review focused primarily on mindfulness meditation.

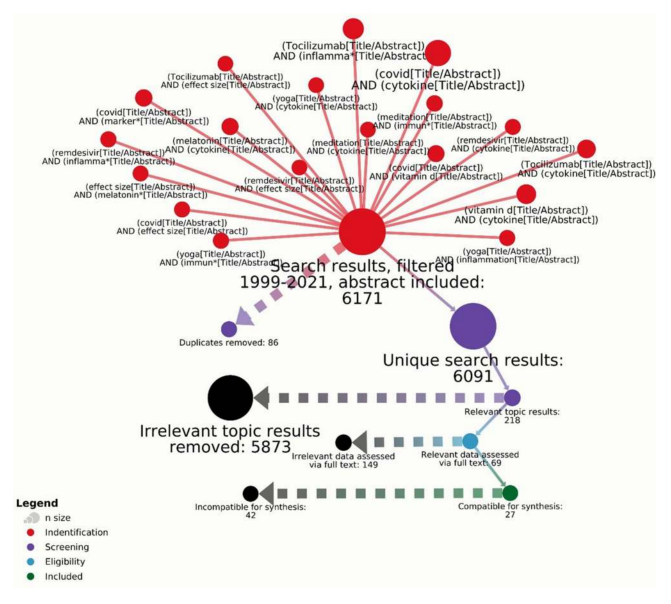


Figure 2 Outline of the source selection process for this review. The process was relatively heterogeneous but primarily focused on adult populations across the world.

Due to the extremely recent nature of COVID-19 and the lack of significant research into many adjunctive approaches, selection also focused on known effects on biomarkers associated with SARS-CoV2 infection, not only on effects specific to COVID-19 infection. The markers tracked in interventions was moderately heterogeneous across studies and availability was a limiting factor in which interventions could be effectively tracked. The search terms were filtered to include studies providing an abstract and published 1999–2021, the date range being intended to ensure current terminology and comprehension of inflammatory disorders. Table 1 lists the search terms developed internally for this study: Searches were conducted primarily through PubMed and subjected to topical identification, screening for relevance, full-text reading for practically relevant data, and included if all of the above allowed coherent synthesis and analysis. Metrics that were compiled in systematic reviews were compared to the original publication to ensure fidelity.

The final results for review and analysis include: Aiswarya et al,⁵³ 2021; Akbari et al, 2020;⁵⁴ Alamdari et al, 2015;⁵⁵ Bartels et al, 2013;⁵⁶ Bower et al, 2015;⁵⁷ Cahn et al, 2017;⁴⁸ Carlson et al, 2003,⁵⁸ 2004,⁵⁹ 2007;⁶⁰ Celinski et al, 2014;⁶¹ Chen et al, 2020;⁶² L. Chen et al, 2020;⁶³ Cichoz-Lach et al, 2010;⁶⁴ Epel et al, 2016;⁶⁵

Table I	Search	Terms	Included	in	Selection	of	Sources	for
Review a	nd Analy	rsis						

Search Terms
(CovidTitle/Abstract) AND (effect sizeTitle/Abstract)
(Effect sizeTitle/Abstract) AND (melatonin*Title/Abstract)
(CovidTitle/Abstract) AND (vitamin dTitle/Abstract)
(RemdesivirTitle/Abstract) AND (cytokineTitle/Abstract)
(TocilizumabTitle/Abstract) AND (cytokineTitle/Abstract)
(RemdesivirTitle/Abstract) AND (effect sizeTitle/Abstract)
(TocilizumabTitle/Abstract) AND (effect sizeTitle/Abstract)
(RemdesivirTitle/Abstract) AND (inflamma*Title/Abstract)
(TocilizumabTitle/Abstract) AND (inflamma*Title/Abstract)
(CovidTitle/Abstract) AND (cytokineTitle/Abstract)
(MeditationTitle/Abstract) AND (cytokineTitle/Abstract)
(MeditationTitle/Abstract) AND (immun*Title/Abstract)
(CovidTitle/Abstract) AND (marker*Title/Abstract)
(YogaTitle/Abstract) AND (inflamma*Title/Abstract)
(YogaTitle/Abstract) AND (cytokineTitle/Abstract)
(YogaTitle/Abstract) AND (immun*Title/Abstract)
(MelatoninTitle/Abstract) AND (cytokineTitle/Abstract)
(Vitamin dTitle/Abstract) AND (cytokineTitle/Abstract)

Gonzalez-Garcia et al, 2014;⁶⁶ Javanmard et al, 2016;⁶⁷ Lengacher et al, 2013;⁶⁸ Qin et al, 2020a,⁶⁹ 2020b;⁶⁹ Rausch-Fan et al, 2002;⁷⁰ Raygan et al, 2019;⁷¹ Rosenkranz et al, 2016;⁷² Shete et al, 2017;⁷³ Smolen et al, 2008;⁷⁴ Wang et al, 2020;⁷⁵ Witek-Janusek et al, 2008;⁷⁶ Zhang et al, 2020.³⁹

Biomarker Selection

Disease markers were selected for inclusion if there were sufficient empirical evidence from research to calculate COVID-19 effects (Table 2).^{39,48,53–76} Intervention effects were included if they affected at least one inflammatory marker.

Exceptions to This Rule Include CD3

Though there were insufficient data to estimate effect sizes with COVID-19 status, there was evidence of close associations between CD3 and COVID-19's inflammatory effects.⁷⁷

Table 2 Markers,	Largely	Inflammatory,	Selected	from	Those
Most Impacted by	COVID-	19			

Markers	
CD19	IL-6
CD3	IL-6 classic
CD4	IL-6 trans
CD8	IL-8
Cortisol	INF-λ
CRP	Lymphocytes
Eosinophil	Monocyte
ESR (erythrocyte sedimentation rate)	NK cell (natural killer cell)
Ferritin	Th Memory cells
IL-10	Th Naïve cells
IL-2	TNF a
IL-4	

Cortisol

Cortisol was widely recorded as a predictive marker of severe COVID-19 cases and was tracked with sufficient data across several interventions, so this marker was included despite insufficient data to calculate COVID-19 effect sizes.⁷⁸

IL-6

There were ample data among both COVID-19 research and the various interventions to calculate effects on interleukin 6, but a specific distinction deserves note. IL-6 is often reported as a pro-inflammatory marker and is one of the most dysregulated cytokines affected by COVID-19, but there have also been interactions with IL-6 that were wildly inconsistent across studies, sometimes demonstrating behavior directly opposing what is expected of an inflammatory marker.^{28,56} A possible explanation involves research indicating that there are two forms of IL-6, a proinflammatory IL-6 trans and a regulating antiinflammatory IL-6 classic.⁷⁹ All effects on IL-6 were calculated and regulating effects categorized as IL-6 classic, while dysregulating effects were categorized as IL-6 trans. Note that this is an internal categorization and does not reflect differentiation in the data. Should the premise of two forms of IL-6 be discounted the effect sizes toward a combined IL-6 will not be affected.

Effect Size vs Significance

Effect size by standardized mean difference (SMD or d) differs from statistical significance (p) and is the primary metric explored in this paper. This is driven by the fact that significance reports how likely a difference is due to chance, while effect size reports how large the difference is. When attempting to compare the potential effects of diverse interventions, p alone is insufficient to determine efficacy. The need for effect sizes when examining tangible effects was explained by the American Statistical Association:

Statistical significance is not equivalent to scientific, human, or economic significance. Smaller p-values do not necessarily imply the presence of larger or more important effects, and larger p-values do not imply a lack of importance or even lack of effect.⁸⁰

The majority of sources used in this analysis did not provide effect sizes. When sources did not include a reported standardized mean difference, an effect size (d) was calculated using test mean (M), control mean, and pooled standard deviations (SD). When sources did not include standard deviations, SD was calculated using sample size (n), and a minimum/maximum range of reported values. When sources only included weighted mean difference, these were converted to standardized mean difference using the WMD value, n, the minimum/ maximum range of reported values, and the SD. If the above data were not available the study was designated as incompatible for synthesis and excluded.

 $d = (M_2 - M_1)/SD_{pooled}$ $SD_{pooled} = \sqrt{((SD_1^2 + SD_2^2)/2)}$

Where SMD were categorized the size guidelines of Cohen's d were followed with some adjustments:⁸¹

- Below 0.1 is negligible effect size.
- 0.1–0.2 is very small effect size.
- 0.2–0.5 is small effect size.
- 0.5–0.8 is medium effect size.
- 0.8–1.0 is large effect size.
- Above 1.0 is very large effect size.

Though extensive effort was made to ensure accuracy, due to the diversity of metrics and conversion to different measurement systems, all SMD effect sizes reported in this analysis should be considered approximate.

System Analysis

This analysis is a preliminary step toward a systems-level examination of COVID-19 effects and interventions. A systemic disease requires a systemic research approach, which itself requires a comprehension of a wide array of interconnections. Though the scope of this initial paper does not include a complex systems model, multiple network analyses were performed. These metrics will inform not only the effects of individual interventions, but the density of overlap between the effects of the disease and interventions in connection with different inflammatory markers.

Results

Given the complex nature of the immune system and cytokines in particular, simple categorization is problematic, but broad generalities have been applied. Markers whose increase is usually indicative of dysregulated inflammation have been labelled as inflammatory markers, while markers whose increase is usually indicative of a healthy, regulated response are labelled as antiinflammatory markers. Catalyst effects which increase anti-inflammatory markers are labelled as regulating while catalyst effects which increase inflammatory markers are labelled as dysregulating. These are only broad tendencies and shall not be interpreted as universal qualities nor precise descriptions of the nuanced behavior of these markers.

The studies incorporated into this analysis included numerous severity levels of COVID-19, different dosages of Remdesivir and Tocilizumab, and various durations of adjunctive interventions. A limiting factor is the heterogeneity between states of illness and relative effect sizes on inflammatory markers, an example being the fact most COVID-19 patients were tracked only after developing acute symptoms while adjunctive interventions sometimes tracked otherwise healthy patients. In an attempt to detect broader patterns the individual results for each catalyst were combined and averaged. Figure 3 represents averaged SMD for each catalyst, a more detailed set of metrics is available in Appendix A.^{39,48,53–76}

COVID-19 Effects

As expected for a systemic disease, Table 3 indicates COVID-19 exerted significant dysregulating effects across a wide variety of markers.^{39,48,53–76}

COVID-19's effects were mostly very large, including major dysregulating effects on CD19, CD4, CD8, C-reactive



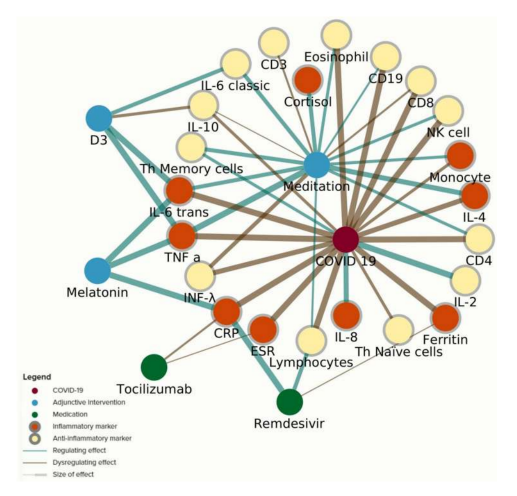


Figure 3 Data point map of catalysts, inflammatory markers, and effect size of connection. Color of the circles identifies whether an inflammatory marker, anti-inflammatory marker, adjunct intervention, medication, or COVID-19 is being referenced. Blue connecting lines indicate a regulating effect from the catalyst to the marker. Brown connecting lines indicate a regulating effect from the catalyst to the marker. The thickness of the connecting line represents the relative effect size of the catalyst on the marker.

protein, eosinophils, erythrocyte sedimentation rate, ferritin, IL-6 trans, overall lymphocytes, natural killer cells, and TNF-a. There was a very large regulating effect on IL-2, which may be a correlation with the body's heightened attempts to curb the runaway inflammation. There were large dysregulating effects on IL-4 and INF- λ , with a large regulating effect on IL-8. COVID-19 had small dysregulating effects on IL-10, monocyte levels, and Th naïve cells, with a small improvement in Th memory cells.

Pharmacological Effects

Despite a large body of studies examining Remdesivir and Tocilizumab, few incorporated sufficient data to calculate effect size on specific markers. Where greater detail was available methodology and differentiation of interventions are listed in <u>Appendix A</u>.

Remdesivir

The data that were available in Table 4 indicated Remdesivir had a very large effect at regulating levels of C-reactive protein, moderate effects at increasing lymphocyte levels, and a negligible dysregulating effect on ferritin levels.^{39,48,53–76}

Tocilizumab

No significant improvements were detected in the averaged SMD for Tocilizumab, as shown in Table 5.^{39,48,53–76} Tocilizumab showed a very small dysregulating effect on C-reactive protein and a negligible dysregulating effect on erythrocyte sedimentation rates.

Adjunctive Intervention Effects

Where greater detail was available methodology and differentiation of interventions are listed in <u>Appendix A</u>.

Catalyst	Marker	Impact	Effect Size	SMD		
COVID 19	CD19	Dysregulating	Very large	-2.04303		
COVID 19	CD4	Dysregulating	Very large	-1.56414		
COVID 19	CD8	Dysregulating	Very large	-2.13114		
COVID 19	CRP	Dysregulating	Very large	3.989791		
COVID 19	Eosinophil	Dysregulating	Very large	-4.12104		
COVID 19	ESR	Dysregulating	Very large	3.153453		
COVID 19	Ferritin	Dysregulating	Very large	2.691077		
COVID 19	IL-2	Regulating	Very large	2.199888		
COVID 19	IL-6 trans	Dysregulating	Very large	2.820931		
COVID 19	Lymphocytes	Dysregulating	Very large	-2.88716		
COVID 19	NK cell	Dysregulating	Very large	-2.87907		
COVID 19	TNF a	Dysregulating	Very large	2.424624		
COVID 19	IL-4	Dysregulating	Large	0.964237		
COVID 19	IL-8	Regulating	Large	-0.92577		
COVID 19	INF-λ	Dysregulating	Large	-0.98216		
COVID 19	IL-10	Dysregulating	Small	-0.45513		
COVID 19	Monocyte	Dysregulating	Small	0.440168		
COVID 19	Th Memory cells	Regulating	Small	0.32566		
COVID 19	Th Naïve cells	Dysregulating	Small	-0.2364		

 Table 3 Effects of COVID-19 on Markers, Whether the Effect Was Presumed Regulating or Dysregulating, and the Size of the Effect

 Using Modified Cohen's d. Differentiation by Severity and/or Time Period of Observation Detailed in Appendix A

Table 4 Effects of Remdesivir on Markers, Whether the EffectWas Presumed Regulating or Dysregulating, and the Size of theEffect Using Modified Cohen's d. Differentiation by Severity and/or Time Period of Observation Detailed in Appendix A

Catalyst	Marker	Impact	Effect Size	SMD
Remdesivir	CRP	Regulating	Very large	-3.34148
Remdesivir	Lymphocytes	Regulating	Medium	0.727273
Remdesivir	Ferritin	Dysregulating	Negligible	0.010854

Table 5 Effects of Tocilizumab on Markers, Whether the EffectWas Presumed Regulating or Dysregulating, and the Size of theEffect Using Modified Cohen's d. Differentiation by Severity and/or Time Period of Observation Detailed in Appendix A

Catalyst	Marker	Impact	Effect Size	SMD
Tocilizumab	CRP	Dysregulating	Very small	0.101553
Tocilizumab	ESR	Dysregulating	Negligible	0.018939

Catalyst	Marker	Impact	Effect Size	SMD
D3	IL-6 trans	Regulating	Very large	-1.65169
D3	TNF a	Regulating	Very large	-2.1139
D3	IL-6 classic	Regulating	Medium	0.772109
D3	IL-10	Dysregulating	Small	-0.39462

Vitamin D₃

In Table 6 Vitamin D_3 demonstrated very large reductions in TNF-a and IL-6 trans levels, moderate effects on increasing IL-6 classic levels, and small dysregulation of IL-10.^{39,48,53–76}

Meditation

Table 7 indicates that meditation affected a very wide variety of markers, more than any other catalyst besides COVID-19.^{39,48,53–76} Meditation demonstrated very large regulating effects in TNF-a and large regulating effects in IL-4. Moderate regulating effects were seen in cortisol, IL-6 trans (decreased), IL-6 classic (increased), and Th memory cells, with moderate dysregulation to INF- λ . There were small regulating effects in CD4, eosinophils, monocytes, and natural killer cells. Meditation showed very small regulating effects in CD19 and overall lymphocytes, with a very small dysregulating effect on CD3 and CD8. There was a negligible dysregulating effect on IL-10.

It is notable that the research indicating there is both a pro-inflammatory and anti-inflammatory form of IL-6 suggests they would be inversely correlated.²⁵ Mean meditation effect sizes show an almost exact inverse correlation between positive and negative effects on IL-6 (0.58 v. -0.57), lending evidence to this premise. **Table 7** Effects of Meditation on Markers, Whether the Effect Was Presumed Regulating or Dysregulating, and the Size of the Effect Using Modified Cohen's d. Included Studies Focused Primarily on Mindfulness Meditation. Differentiation by Severity and/or Time Period of Observation Detailed in <u>Appendix A</u>

Catalyst	Marker	Impact	Effect Size	SMD
Meditation	TNF a	Regulating	Very large	-1.01499
Meditation	IL-4	Regulating	Large	-0.8638
Meditation	Cortisol	Regulating	Medium	-0.74358
Meditation	IL-6 classic	Regulating	Medium	0.58209
Meditation	IL-6 trans	Regulating	Medium	-0.57073
Meditation	INF-λ	Dysregulating	Medium	-0.715
Meditation	Th Memory cells	Regulating	Medium	0.561265
Meditation	CD4	Regulating	Small	0.3826
Meditation	Eosinophil	Regulating	Small	0.31
Meditation	Monocyte	Regulating	Small	-0.293
Meditation	NK cell	Regulating	Small	0.472
Meditation	CD19	Regulating	Very small	0.149333
Meditation	CD3	Dysregulating	Very small	-0.11597
Meditation	CD8	Dysregulating	Very small	-0.17333
Meditation	Lymphocytes	Regulating	Very small	0.151667
Meditation	IL-10	Dysregulating	Negligible	-0.095

Table 8 Effects of Melatonin on Markers, Whether the EffectWas Presumed Regulating or Dysregulating, and the Size of theEffect Using Modified Cohen's d. Differentiation by Severity and/or Time Period of Observation Detailed in Appendix A

Catalyst	Marker	Impact	Effect Size	SMD
Melatonin	CRP	Regulating	Very large	-1.815
Melatonin	IL-6 trans	Regulating	Very large	-2.05
Melatonin	TNF a	Regulating	Very large	-1.89

Melatonin

As seen in Table 8, melatonin did not affect as many markers as meditation, but it had the greatest number of very large effects of any intervention or medication, with major improvements in C-reactive protein, IL-6 trans, and TNF-a.^{39,48,53–76}

Analysis Effect Size Comparison

Comparisons were examined where two or more catalysts affected a given marker. The markers tracked are grouped into the broad categories of leukocytes and cytokines, with a third category including only one marker, erythrocyte sedimentation rate.

Leukocytes

As expected, COVID-19 had significantly larger dysregulating effects than any intervention on leukocytes, illustrated in Figure $4^{.39,48,53-76}$ A surprising finding was the wide variety of markers improved by meditation.

- Meditation had opposite effects to COVID-19 in five out of eight leukocyte comparisons tracked, including CD19, CD4, eosinophils, overall lymphocytes, monocytes, and natural killer cells.
- CD8: There was a very small increase in CD8 with meditation, approximately 8.1% the size of the COVID-19 effect on CD8.
- Th memory cells: A notable effect was a significantly greater effect by meditation on the production of Th memory cells than was stimulated by introduction of the virus alone. This finding has implications for antiviral strategies.
- Lymphocytes: Remdesivir could only be tracked to overall lymphocytes, but in this instance outperformed meditation in effects contrary to COVID-19. This suggests efficacy for improving sustainable lymphocyte production, though the systemic effects are difficult to track.

Cytokines/Chemokines

Figure 5 demonstrates that though COVID-19 had significantly greater effects on leukocytes than any of the interventions, influences on cytokines were more

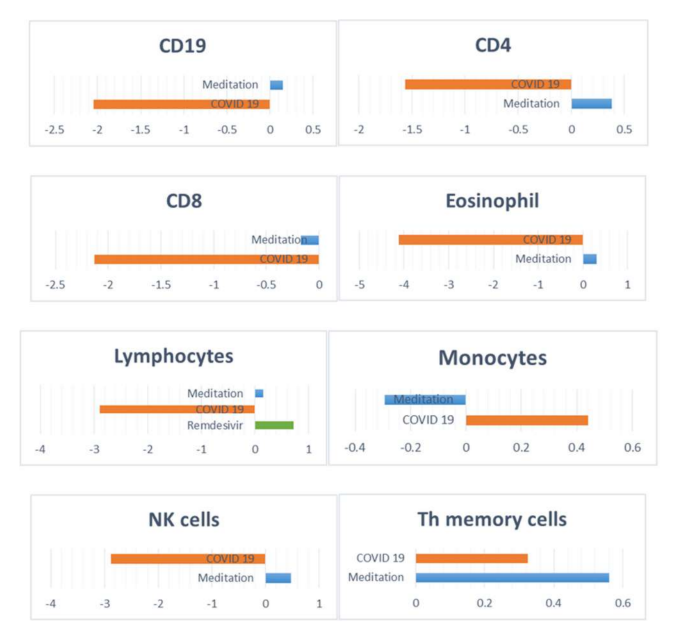


Figure 4 Comparative effects on leukocytes by COVID-19, medication, and adjunct interventions. Horizontal bars track the effects of interventions or COVID-19, the X-axis indicates the effect size via standard mean difference (SMD). Tracked markers include CD19, CD4 (Th helper cells), CD8, eosinophils, lymphocytes, monocytes, NK cells (natural killer cells), and TH memory cells.

competitive.^{39,48,53–76} There were also more data for cytokine effects, allowing more comprehensive comparisons.

- C-reactive protein: This marker was tracked across four catalysts, Tocilizumab, Remdesivir, melatonin, and COVID-19. Tocilizumab had a very small dysregulating effect, but both Remdesivir and melatonin showed very large effects in opposition to COVID-19's influence, Remdesivir showing the largest single effect of any intervention on CRP.
- Ferritin: Remdesivir had a negligible effect on ferritin.
- IL-4: Meditation had an opposing effect to COVID-19 on IL-4 of an equivalent amount, both having large effect sizes.
- IL-6 classic: Both meditation and vitamin D₃ had a medium effect on IL-6 classic, the potentially regulating version of the IL-6 cytokine. It is notable that meditation and vitamin D₃ both had effects that increased and decreased IL-6 at approximately



Figure 5 Comparative effects on cytokines by COVID-19, medication, and adjunct interventions. Horizontal bars track the effects of interventions or COVID-19, the X-axis indicates the effect size via standard mean difference (SMD). Tracked markers include CRP (C-reactive protein), ferritin, IL-4 (interleukin 4), IL-6 classic (regulatory interleukin 6), IL-6 trans (dysregulated interleukin 6), IL-20 (interleukin 10), INF λ (interferons λ), and TNF-a (tumor necrosis factor alpha).

inverse amounts depending on the study, lending further evidence to the concept of a trans and classic form of IL-6.

• IL-6 trans: As the most prominent indicator of cytokine storms, IL-6 trans was tracked by multiple catalysts, though insufficient data were found for Remdesivir and Tocilizumab. Melatonin and vitamin D_3 both showed a very large effect on IL-6 trans, though still somewhat smaller than the opposing effect of COVID-19. Meditation showed a medium sized effect on IL-6 trans. It is notable that meditation and vitamin D_3 both had effects that increased and decreased IL-6 at approximately inverse amounts depending on the study, lending further evidence to the concept of a trans and classic form of IL-6.

- IL-10: Provided unexpected results, with COVID-19, vitamin D₃, and meditation all increasing the marker to some degree, meditation to a significantly smaller degree.
- INF-λ: Meditation had a medium sized effect on INFλ that corresponded with a large effect by COVID-19. The complex relationship of interferons with various immune functions makes this correlation difficult to analyze.
- Tumor necrosis alpha: Along with IL-6, TNF-a is a cytokine frequently linked to dysregulated inflammation. Melatonin, vitamin D₃, and meditation all showed a very large effect in opposition to COVID-19's very large effect.

ESR

Figure 6 tracks erythrocyte sedimentation rate, a common hematological test to measure levels of inflammation.^{39,48,53–76}

• ESR: Tocilizumab showed a negligible effect on ESR in comparison to COVID-19's very large effect.

System Connections

Meaningful system analysis requires an understanding of the connections between parts of the system. This work is not intended to conduct such an analysis but can provide information on the connections to be used in future research. These simple network metrics reflect the number and direction of connections compiled in this research.

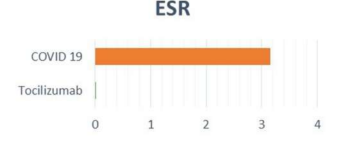


Figure 6 Comparative effects on ESR (erythrocyte sedimentation rate) by COVID-19, medication, and adjunct interventions.

Degree Centrality

The simplest of network metrics, these numbers represent the total number of connections of any kind that belong to a data point. Degree centrality is useful for determining hubs of interaction but cannot determine direction of influence.

Indegree

Unlike degree centrality, indegree measures directional connections, specifically the data points receiving incoming connections. Given the focus of this research, indegree is primarily made up of inflammatory markers. Higher values reflect a greater degree of influence from the rest of the system.

Outdegree

Whereas indegree measures incoming connections, outdegree tracks the number of outgoing connections from a given data point. Given the focus of this research, indegree is made up of catalysts. Higher values reflect a greater degree of influence upon the rest of the system. These connections were calculated in Table 9 based on the relationships discovered in the analyzed body of research.^{39,48,53–76}

An examination of the degrees of connection revealed COVID-19 and meditation were by far the most widely influential catalysts. The markers most affected by any catalyst included IL-6 trans, TNF-a, and CRP, followed by IL-10 and overall lymphocytes.

Discussion

The initial hypothesis asks whether adjunct interventions show any significant effects on inflammatory markers that are dysregulated by COVID-19. These effects were compared to Remdesivir and Tocilizumab, some of the leading SARS-CoV2 medications, to determine whether there are any unique benefits. The findings demonstrate that adjunct interventions not only affect a greater number of markers than Remdesivir or Tocilizumab, the effect sizes of adjunct treatments are often as large or larger. Melatonin, vitamin D₃, and meditation affected 23 inflammatory markers while Remdesivir and Tocilizumab affected 5. Adjunctive interventions affected 68.4% of the inflammatory markers impacted by COVID-19, standard pharmaceutical medication affected 26.3% of the inflammatory markers impacted by COVID-19. Melatonin, vitamin D₃, and meditation had very large or large effects on 7 inflammatory markers, Remdesivir and Tocilizumab had very large or large effects on 1 inflammatory marker, though that single effect was extremely large.

Degree Centrality	Value	Indegree	Value	Outdegree	Value
COVID 19	19	IL-6 trans	4	COVID 19	19
Meditation	16	TNF a	4	Meditation	16
D3	4	CRP	4	D3	4
IL-6 trans	4	IL-10	3	Melatonin	3
TNF a	4	Lymphocytes	3	Remdesivir	3
CRP	4	CD8	2	Tocilizumab	2
Melatonin	3	IL-4	2	CD8	0
IL-10	3	CD19	2	Cortisol	0
Remdesivir	3	ESR	2	IL-2	0
Lymphocytes	3	IL-6 classic	2	IL-6 trans	0
CD8	2	Eosinophil	2	IL-4	0
IL-4	2	INF-λ	2	CD19	0
CD19	2	CD4	2	IL-10	0
ESR	2	NK cell	2	IL-8	0
IL-6 classic	2	Th Memory cells	2	ESR	0
Eosinophil	2	Monocyte	2	TNF a	0
INF-λ	2	Ferritin	2	IL-6 classic	0
CD4	2	Cortisol	1	Eosinophil	0
NK cell	2	IL-2	1	Lymphocytes	0
Th memory cells	2	IL-8	1	CRP	0

Table 9 Top Twenty Data Points and V	alues of the Network Metrics, Tracking D	Degree Centrality, Indegree, and Outdegree
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Though the impacts of Tocilizumab are clearly smaller in both effect size and number of markers affected, comparisons between Remdesivir, melatonin, vitamin $D_{3,}$ and meditation are significantly more nuanced. In comparison to Remdesivir the adjunctive treatments affected a greater quantity of markers, with a far greater number of large effects, yet Remdesivir exhibited a uniquely large effect size on C-reactive proteins. Uneven distribution of both effect sizes and markers affected suggests different mechanisms operating behind the adjunctive treatments and Remdesivir. In this scenario the adjunct treatments and Remdesivir might operate synergistically, influencing different inflammatory pathways via different mechanisms.

Melatonin, vitamin D₃, and meditation should be considered for treatment of COVID-19 alongside standards of care. Though the precise interactions between markers and disease outcomes are unclear, evidence suggests these interventions have significant, quantifiable effects on the inflammatory markers associated with COVID-19's dysregulation.

Implications

There are short and long-term implications behind the evidence supporting the efficacy of adjunct interventions to improve inflammatory dysregulation. In the immediate future these adjunct interventions could be considered for use alongside standard medication. The cost-efficiency and lack of serious side effects or contraindications associated with reasonable applications of meditation, vitamin D_3 , and melatonin makes for a simple, affordable, and easy combination with most treatment plans. The increased scope of regulating systemic inflammation could result in significantly lower rates of cytokine storms, potentially saving many lives.

The long-term benefit to this approach lies in the focus on systemic dysregulation rather than acute events. The current healthcare system is highly vulnerable to the ripple effects of inflammatory viruses on chronically ill patients, posing high individual and institutional risk. COVID-19 is particularly systemic in its pathology, yet there are many communicable and non-communicable diseases that are heavily influenced by interconnected, dysregulated inflammation. A greater focus on systemic diseases and interventions is highlighted in this pandemic, but advances in this field could have lasting benefits across numerous fields of medicine. It is likely there is no single solution to either the pandemic or these institutional weaknesses, but rather a combination of interventions and policies are necessary to stimulate the entire organism to adapt.

Further research, both into the adjunct interventions listed here and into system-wide interventions overall, should be considered warranted and pressing.

Limitations

This research has several conditions and limits that must be considered. The act of aggregating, calculating, approximating, and averaging effect sizes from very disparate sources creates the possibility of imprecision at multiple levels. As referenced above, differences in the initial health conditions and observation intervals of the participants in these studies are complicating factors that warrant examination. The consistent findings of data from distinct sources lends credibility to the overall findings, but review of the differentiated data is available and encouraged in Appendix A. The selection, characterization, and contextualization of highly complex inflammatory markers is likewise inherently approximate. Limited data on markers associated with many interventions also prevented a wider comparative analysis among standard medical approaches. In particular, it is important to note that although insufficient data were found to calculate effect sizes on IL-6 by Tocilizumab, this intervention is primarily known as an IL-6 inhibitor, so it can be assumed it has a significant effect on that marker.⁸²

While Tocilizumab and Remdesivir were being studied as leading interventions when this study began, by the time of publication they have been largely supplanted by more effective treatments. Comparison with contemporary standards of care is warranted. Most COVID-19 effects were tracked after admission to a hospital or care center, creating statistical sampling issues in relation to asymptomatic or misdiagnosed infections. Long-term examination of changes in inflammatory loads among asymptomatic infections is necessary to expand marker effect size accuracy.

Several emergent Phase II/III studies on artemisinin and derivatives have released preliminary data suggesting a reduced time of conversion to negativity and reduction in severe symptoms.⁸³ Artemisia and its bioactive constituents represent promising examples of adjunct therapeutics for broad spectrum antiviral effects, reduction in tissue fibrosis, and inhibition of key cytokines such as TGF-beta, benefits that have been well established in malaria treatments. However, the artemisinin studies did not evaluate serum cytokines and could not be included in the current meta-analysis; thus future clinical trials administering these compounds and others should extensively evaluate serum biomarkers.

Finally, a critical limitation is a lack of understanding between the mechanisms of inflammatory markers and the epidemiological behavior of COVID-19. This paper explores effects upon inflammatory markers associated with COVID-19, but the complex dynamics of those markers cannot yet be used to predict differences in treatment outcomes. Further exploration of the threshold for cytokine storms is necessary before any precise benefits in reducing individual markers can be determined.

Disclosure

The authors report no conflicts of interest for this work.

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