

COMMENT

Use of hydroxychloroquine in multidrug protocols for SARS-CoV-2

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Abstract

We review the available evidence supporting the use of hydroxychloroquine-based multidrug protocols in the treatment of COVID-19, in response to a recently published editorial by TMJ. *Tasman Medical Journal* 2024; 6: 27-32

Introduction

We read with interest an editorial by Millar¹ concerning the role of hydroxychloroquine in the treatment of COVID-19 patients. Community standard of care multidrug therapies for COVID-19 were based on signals of benefit and acceptable safety.²⁻⁹ At the onset of the pandemic, there was insufficient time for large prospective randomized controlled trials (RCT) to validate community standard of care protocols. In such studies, randomization should handle the validity threats of selection bias and both known and unknown confounders, however successful randomization requires a large number of patients with outcome events (e.g. hospitalizations, deaths), to ensure that the patients experiencing these events are also randomized.¹⁰ As an example, the number n of events needed to randomize a dichotomous equiprobable confounder variable (i.e. similar to male/female) within $x = 10\%$ margin (with 95% confidence) can be obtained by bounding the ratio σ/μ of standard deviation to mean with $2(\sigma/\mu) = 2(1/n)^{1/2} \leq x = 0.1$, thus requiring $n \geq 400$ expected events. Therefore, for RCTs with a mortality endpoint, if one assumes $p = 2\%$ case fatality rate (CFR), then the estimated sample size required to achieve sufficient

randomization is $N = n/p \geq 20,000$, which is impractical in the midst of an emergency.

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Clinicians understood quickly that no single drug was going to be necessary nor sufficient to treat acute COVID-19 with its three phases of viral replication, cytokine storm, and thrombosis. Hydroxychloroquine was part of the initial multidrug protocol used by Zelenko from March 2020.¹¹ On April 28, 2020, Zelenko published a letter,^{12,13} also reproduced in his posthumous autobiography,¹⁴ reporting the details of his hydroxychloroquine-based multidrug protocol and his patient outcomes. Zelenko's protocol consisted of risk stratifying patients as high or low, and treating the high-risk patients with hydroxychloroquine (200 mg twice daily for 5 days), azithromycin (500 mg once daily for 5 days), and zinc sulfate (50 mg elemental zinc for 5 days).² He defined three categories of patients at high risk: (a) all older than 60 years of age; (b) those that were immuno-compromised or had comorbidities or whose BMI was $\geq 30 \text{ kg/m}^2$; (c) all patients not satisfying the previous two conditions who developed

shortness of breath. By April 28, 2020, Zelenko had treated 405 high-risk patients resulting in 6 hospitalizations and 2 deaths. No hospitalizations or

deaths were observed amongst the other 1,045 low-risk patients who received only supportive care. He

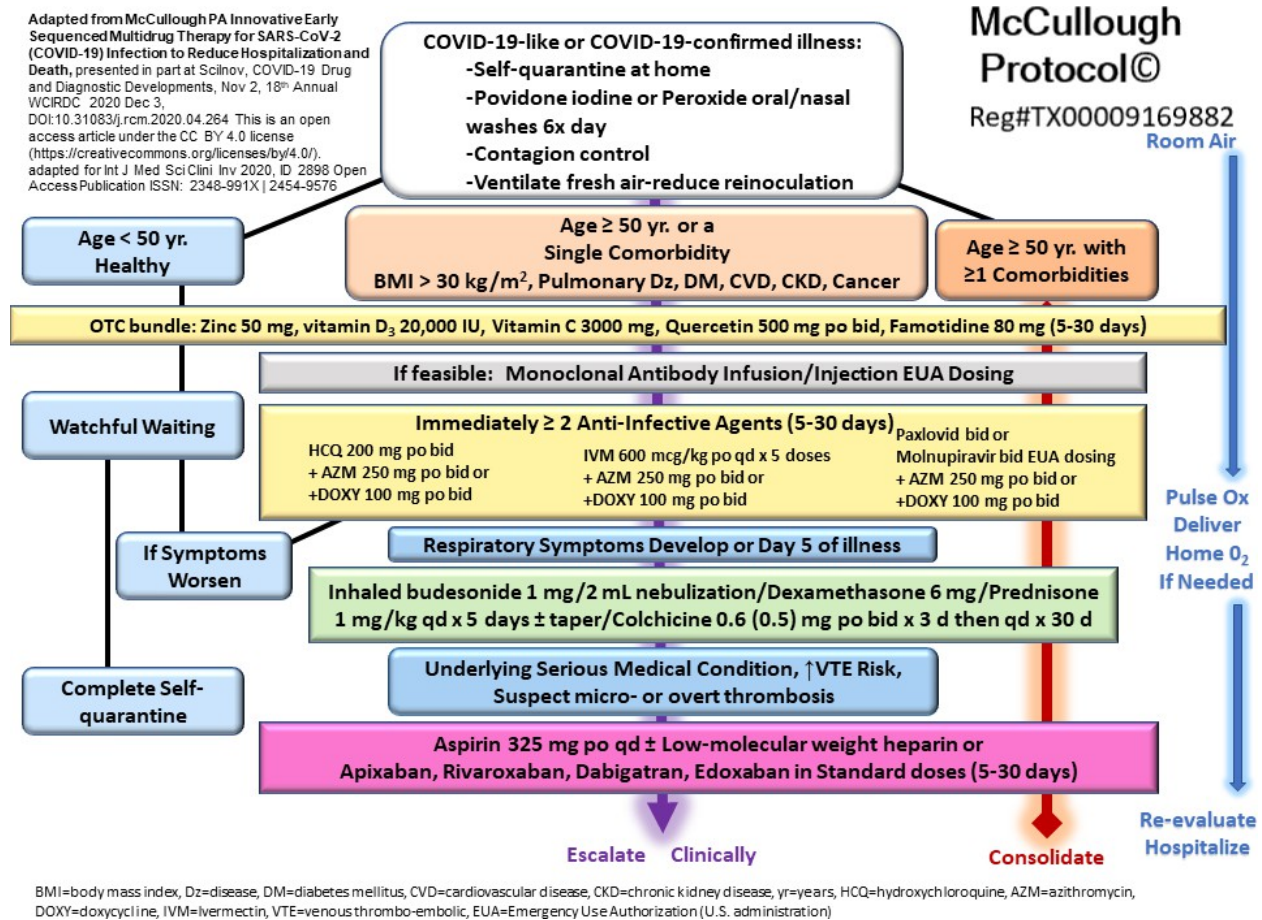


Figure 1: McCullough's protocol^{5-7,19} of sequenced multidrug pre-hospital treatment of the three stages of COVID-19: viral proliferation, cytokine injury, and thrombosis. No single drug is necessary nor sufficient.

improved on his triple drug protocol by introducing budesonide nebulization and oral dexamethasone at the beginning of May 2020, and selective use of apixaban near the end of May 2020¹⁵. By June of that year he had treated 800 high-risk patients, resulting cumulatively in 12 hospitalizations and 2 deaths.^{14,16,17} While public health policy in the United States opposed the adoption of Zelenko's protocol,¹⁸ the community standard of care developed from that point forward to the widely adopted McCullough protocol^{3-5,19} (Fig. 1).

In 2022, we proposed a statistical technique for comparing a case series (N, a) of N patients that received treatment with a negative events (e.g. hospitalizations, deaths, etc.) against historical controls that lower-bound

the probability x of a negative event without treatment by an inequality $p_1 < x$.¹⁵ Our technique calculates from (N, a) an efficacy threshold x_0 and a random selection bias threshold x_1 , both dependent on the desired level of confidence $1 - p_0$ (Fig. 2). Then, $p_1 > x_0$ implies the existence of treatment efficacy by the preponderance of evidence, meaning that it is more likely than not that the observed effect cannot be entirely accounted for by random selection bias, thus justifying an emergency adoption. Likewise, $p_1 > x_1$ implies that the existence of some treatment efficacy is clear and convincing, meaning that we can have $1 - p_0$ confidence that the observed benefit cannot be entirely accounted for by random selection bias, at which point there is no longer sufficient equipoise to ethically justify a randomized

controlled trial against placebo. Here, *random selection bias* refers to any possible selection bias that can result by randomly choosing N patients out of the entire population. This analysis can be used only with regimens that have known acceptable safety, limiting its applicability to treatments using safe repurposed medications.

Because Zelenko treated only high-risk patients, with increased likelihood of death relative to the general population, we can compare his case series against observed outcomes over the entire United States population. This comparison is biased towards the null hypothesis, but a positive result that overcomes this bias

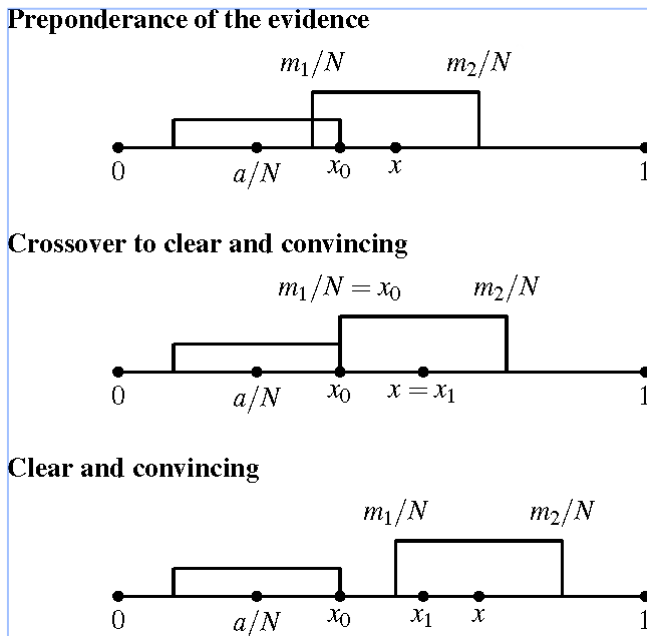


Figure 2: Statistical comparison of a case series (N, a) of N treated patients with a negative events against the population-level probability x of a negative event without treatment. A positive finding depends on the relative position between the confidence interval for the probability of negative event with treatment (on the left) and the confidence interval $[m_1/N, m_2/N]$ for the probability of a negative event without treatment for N randomly selected patients. This figure is adapted from the graphical abstract of Gkioulekas et al.¹⁵ under the terms of the CC-BY-4.0 license.

is sufficient. For $(N, a) = (405, 2)$ we obtained $x_0 =$

1.8% and $x_1 = 4.0\%$ and for $(N, a) = (800, 2)$ we obtained $x_0 = 1.0\%$ and $x_1 = 2.0\%$, using 95% confidence.¹⁵ During 2020, the case fatality rate (CFR) in the United States ranged between 2% and 6%.²⁰ Using $p_1 = 2\%$, it follows that by the end of April 2020 the mortality rate reduction benefit was supported by the preponderance of evidence, with crossover to clear and convincing by June 2020. Similar analysis shows clear and convincing hospitalization rate reduction by the end of April 2020.¹⁵

Furthermore, from a case series of 10,429 outpatients, treated in Marseilles, France by Raoult's group in the IHU Méditerranée Infection Institute with hydroxychloroquine and azithromycin, in addition to standard of care, through the end of December 2020,²¹ we identified a case series of 1495 high-risk patients (age ≥ 60 years) with 5 reported deaths, whereas no deaths were reported for the other 8,934 patients. Using $(N, a) = (1495, 5)$ gives a random selection bias threshold $x_1 = 1.4\%$ for 95% confidence which compares favorably with the CFR in France which ranged from 2% to above 14% during 2020, indicating a clear and convincing finding of mortality rate reduction.^{15,20} The standard of care used by Raoult's group included zinc supplementation, enoxaparin for patients older than 70 or with comorbidities, and dexamethasone, for patients with high viral loads, inflammatory pneumonopathy, or based on clinical judgment.²¹ The mortality rate for patients receiving only this standard of care was 2.1% for high-risk patients with age ≥ 60 years (11 deaths out of 520 high-risk patients) and no deaths reported for the other 1594 patients,²¹ which was lower than one would have expected for untreated high-risk patients.¹⁵ It was also 7-fold larger than the 0.3% mortality rate observed for high-risk patients in the $(N, a) = (1495, 5)$ case series who were treated with hydroxychloroquine and azithromycin in addition to the standard of care.

Recently, Raoult released his dataset of 30,423 COVID-19 patients²²⁻²⁴ treated through the end of 2021. An independent analysis of his data, using propensity score matching and logistic regression, has shown that hydroxychloroquine and azithromycin were associated with 58% reduction of the composite endpoint of ICU

admissions and deaths, whereas azithromycin alone was associated with 27% reduction over the same endpoint.²⁵ This result also implies that the positive results observed, when using hydroxychloroquine and azithromycin in combination, cannot be exclusively attributed to azithromycin alone. Further evidence has been reviewed by Luzariaga and Iglesias.²⁶

The premise underlying Zelenko's protocol was to reduce the viral multiplication rate and enable the immune system to clear the virus before the infection invades the lungs.^{2,27} Subsequently, McCullough's protocol (Fig. 1) recognized that COVID-19 is a triphasic illness, with viral proliferation followed by cytokine injury and thrombosis, requiring a carefully timed sequenced treatment of each phase.^{5-7,19} Consequently, RCTs^{28,29} of hydroxy-chloroquine on hospitalized patients, at the last two stages of the illness, do not extrapolate to outpatients treated during the first stage. To prevent hospitalization, any treatment intervention should be administered early, preferably within 3 days from the onset of symptoms,³⁰ unlike the 8-day window used by the TOGETHER trial.³¹ Spivak *et al*³² is underpowered and tested hydroxychloroquine monotherapy, thus not necessarily generalizable to Zelenko's triple-drug therapy. Furthermore, although the inclusion criteria only allowed patients up to 72 hours after a positive COVID-19 test, this does not account for the unknown additional delay between onset of symptoms and testing.

We concur with Millar's skepticism¹ concerning meta-analyses based on studies that use a multiplicity of treatments. At minimum, outpatient studies need to be separated from inpatient studies and considered separately. The effect size obtained from a meta-analysis is quantitatively meaningful when the underlying studies investigate very similar treatment protocols. Furthermore, his comments¹ suggesting more evidence is needed are well taken. There remain opportunities for large clinical trials for the treatment of high-risk recurrent infections.

Conclusion

It is our interpretation that hydroxychloroquine played an important role in preventing hospitalizations and deaths due to COVID-19, particularly in 2020 with the

more virulent strains. Widespread use of nasal sprays and gargles, aspirin, vitamin D, ivermectin, nirmatrelvir/ritonavir, molnupiravir, favipiravir, colchicine, corticosteroids, and anticoagulants (Fig. 1) in protocols all contributed to the benefits of early treatment which were widely favored over therapeutic nihilism in the pre-hospital phase. In case of a future pandemic, involving a novel disease, doctors should be encouraged to attempt treatments with repurposed medications based on biological plausibility, signals of benefit, and acceptable safety. Article 37 of the 2013 Helsinki declaration allows the use of unproven treatments if "*proven interventions do not exist or other known interventions have been ineffective*" and the unproven treatment "*offers hope of saving life, reestablishing health or alleviating suffering*".³³ When these efforts result in case series of treated patients that show a large magnitude of benefit, then statistical comparison with historical controls can be used to support the strength of association between treatment and improved outcomes.^{15,34} As evidence accumulates, the Bradford Hill criteria framework can be used to assess the support for a causality claim,^{35,36} as an inference to the best explanation.^{37,38} This evidence can be gathered rapidly and form the basis for an agile emergency response to future pandemics, if public health is willing to leverage the clinical experience of medical doctors at the front lines.

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Editor's note: This paper is one of three manuscripts received in response to our editorial review. As the topic of hydroxychloroquine in SARS-CoV-2 infection and repurposed drugs in general is of some importance the Journal has made its

columns available to the authors of these papers without comment. Further comment is welcome.

References

1. Millar JA. The myth of hydroxychloroquine in the treatment of SARS-CoV-2 infection. *Tasman Med J*. 2024; 6: 13-16.
2. Derwand R, Scholz M, Zelenko V. COVID-19 outpatients - Early risk-stratified treatment with zinc plus low dose hydroxychloroquine and azithromycin: A retrospective case series study. *Int J Antimicrob Agents*. 2020; 56: 106214.
3. McCullough PA, Kelly RJ, Ruocco G, Lerma E *et al*. Pathophysiological basis and rationale for early outpatient treatment of SARS-CoV-2 (COVID-19) infection. *Am J Med*. 2020; 134: 16-22.
4. McCullough PA. Innovative early sequenced multidrug therapy for SARS-CoV-2 (COVID-19) infection to reduce hospitalization and death. *Int J Med Sci Clin Invent*. 2020; 7: 5139-50.
5. McCullough PA, Alexander PE, Armstrong R, Arvinte C *et al*. Multifaceted highly targeted sequential multidrug treatment of early ambulatory high-risk SARS-CoV-2 infection (COVID-19). *Rev Cardiovasc Med*. 2020; 21: 517-30.
6. Procter BC, Ross C, Pickard V, Smith E *et al*. Clinical outcomes after early ambulatory multidrug therapy for high-risk SARS-CoV-2 (COVID-19) infection. *Rev Cardiovasc Med*. 2021; 21: 611-4.
7. Procter BC, Ross C, Pickard V, Smith E *et al*. Early ambulatory multidrug therapy reduces hospitalization and death in high-risk patients with SARS-CoV-2 (COVID-19). *Int J Innov Res Med Sci*. 2021; 6: 219-21.
8. Chetty S. Elucidating the pathogenesis and Rx of COVID reveals a missing element. *Mod Med*. 2020; 45: 28-31.
9. Stone JC, Ndarukwa P, Scheim DE, Dancis BM *et al*. Changes in SpO₂ on room air for 34 severe COVID-19 patients after ivermectin-based combination treatment: 62% normalization within 24 hours. *Biologics*. 2022; 2(3): 196-210.
10. Risch H. Pausibility, not science has dominated public discussions of the COVID pandemic. *Am J Econ Sociol*. 2023;82: 411-424.
11. Zelenko V. Correspondence from Dr Vladimir Zelenko on Treatment of COVID-19 in New York; March 23, 2020. Available at: <https://drelef.org/zelenko/Vladimir-Zelenko-treatment.pdf> (accessed July 20, 2022).
12. Zelenko V. To all medical professionals around the world [letter]; April 28, 2020. Available at: <https://drelef.org/zelenko/Zelenko-memo-April.pdf> (accessed July 20, 2022).
13. Risch HA. Early outpatient treatment of symptomatic, high-risk COVID-19 patients that should be ramped-up immediately as key to the pandemic crisis. *Am J Epidemiol*. 2020; 189: 1218-26.
14. Zelenko V, Hamachek B. Zelenko: How To Decapitate The Serpent. Carbondale CO: Pierruci Publishing; 2022.
15. Gkioulekas E, McCullough PA, Zelenko V. Statistical analysis methods applied to early outpatient COVID-19 treatment case series data. *COVID*. 2022; 2(8): 1139-82.
16. Zelenko V. To Dr. Moshe Bar Siman Tov [letter]; June 14, 2020. Available at: <http://drelef.org/zelenko/Zelenko-memo-June-Dr-Shemtov-letter.pdf> (accessed July 20, 2022).
17. Risch HA. The author replies. *Am J Epidemiol*. 2020; 189: 1444-9.
18. Hatfill S. The intentional destruction of the national pandemic plan. *J Am Phys Surg*. 2021; 26: 74-6.
19. Palazzuoli A, Beltrami M, McCullough PA. Acute COVID-19 management in heart failure patients: A specific setting requiring detailed inpatient and outpatient hospital care. *Biomedicines*. 2023; 11: 790.
20. Mathieu E, Ritchie H, Rodes-Guirao L, Appel C *et al*. Coronavirus pandemic (COVID-19). *Our World in Data*. 2023. <https://ourworldindata.org/coronavirus>.
21. Millon M, Lagier JC, Tissot-DuPont H, Ravaux I *et al*. Early treatment with hydroxychloroquine and azithromycin in 10,429 COVID-19 outpatients: A monocentric retrospective cohort study. *Rev Cardiovasc Med*. 2021; 22: 1063-72.
22. Brouqui P, Raoult D. Construction, quality control and regulatory aspect of a database of 30,423 COVID-19 patients cared for at the IHU Méditerranée Infection France. *Biomed J Sci Tech Res*. 2023; 52(3): 43799-804.
23. Millon M, Cortaredona S, Delorme L, Colson P *et al*. Monocentric retrospective cohort of 30,423 COVID-19 patients; 2023. <https://doi.org/10.57760/sciencedb.07803> - Science Data Bank.
24. Brouqui P, Million M, Parola P, McCullough PA *et al*. Outcomes after early treatment with hydroxychloroquine and azithromycin: An analysis of a database of 30,423 COVID-19 patients. *New Microbes New Infect*. 2023; 55: 101188.
25. Lounnas V, Gkioulekas E, Rendell M, Lacout A *et al*. An independent analysis of a retrospective cohort of 30,423 Covid-19 patients treated at IHU-Mediterranean in Marseille, France: Part 1, Efficacy of early treatment with hydroxychloroquine and azithromycin. *Arch Microbiol Immunol*. 2024; 8: 51-66.
26. Luzariaga A, Iglesias J. The need to revisit hydroxychloroquine in early treatment strategies of COVID-19. In: Varon J, Marik PE, Rendell M, Iglesias J *et al*, editors. *Controversies in the Pandemic*. New Delhi, India: Jaypee Brothers Medical Publishers; 2024. .
27. Derwand R, Scholz M. Does zinc supplementation enhance the clinical efficacy of chloroquine/hydroxychloroquine to win today's battle against COVID-19? *Med Hypotheses*. 2020; 142: 109815.
28. The RECOVERY Collaborative Group. Effect of

- hydroxychloroquine in hospitalized patients with Covid-19. *NEJM*. 2020; 383: 2030-40.
29. WHO Solidarity Trial Consortium. Repurposed antiviral drugs for Covid-19 - Interim WHO Solidarity trial results. *NEJM*. 2021; 384: 497-511.
 30. Fazio S, Bellavite P, Zanolin E, McCullough PA *et al*. Retrospective study of outcomes and hospitalization rates of patients in Italy with a confirmed diagnosis of early COVID-19 and treated at home within 3 days or after 3 days of symptom onset with prescribed and non-prescribed treatments between November 2020 and August 2021. *Med Sci Monit*. 2021; 27: e935379.
 31. Reis G, Silva EADSM, Silva DCM, Thabane L *et al*. Effect of early treatment with hydroxychloroquine or lopinavir and ritonavir on risk of hospitalization among patients with COVID-19: The TOGETHER randomized clinical trial. *JAMA Netw Open*. 2021; 4(4): e216468.
 32. Spivak AM, Barney BJ, Greene T, Holubkov R *et al*. A randomized clinical trial testing hydroxychloroquine for reduction of SARS-CoV-2 viral shedding and hospitalization in early outpatient COVID-19 infection. *Microbiol Spectr*. 2023; 11(2): e0467422.
 33. World Medical Association. World Medical Association Declaration of Helsinki: Ethical principles for medical research involving human subjects. *JAMA*. 2013; 310(20): 2191-4.
 34. Rendell M. Commentary on 'Statistical analysis methods applied to early outpatient COVID-19 treatment case series data' by Gkioulekas, McCullough and Zelenko: A return back to the future. *J Health Care Commun*. 2022; 7 (10): 130-5.
 35. Hill AB. The environment and disease: Association or causation? *Proc R Soc Med*. 1965; 58(5): 295-300.
 36. Howick J, Glasziou P, Aronson JK. The evolution of evidence hierarchies: what can Bradford Hill's 'guidelines for causation' contribute? *JRSM*. 2009; 102(5): 186-94.
 37. Harman GH. The inference to the best explanation. *Philos Rev*. 1965; 74(1): 88-95.
 38. Ward AC. The role of causal criteria in causal inferences: Bradford Hill's 'aspects of association'. *Epidemiol Perspect & Innov*. 2009; 6: 2.