Early Outpatient Treatment of Symptomatic, High-Risk COVID-19 Patients That Should Be Ramped Up Immediately as Key to the Pandemic Crisis

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More than 1.6 million Americans have been infected with severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), and more than 10 times that number carry antibodies to it. High-risk patients with progressing symptomatic disease currently have only hospitalization treatment, with its high mortality, available to them. An outpatient treatment that prevents hospitalization is desperately needed. Two candidate medications have been widely discussed: remdesivir and hydroxychloroquine (HCQ) + azithromycin (AZ). Remdesivir has shown mild effectiveness in hospitalized inpatients, but no trials in outpatients have been registered. HCQ + AZ has been widely misrepresented in both clinical reports and public media, and results of outpatient trials are not expected until September. Early outpatient illness is very different from later florid disease requiring hospitalization, and the treatments differ. Evidence about use of HCQ alone, or of HCQ + AZ in inpatients, is irrelevant with regard to the efficacy of HCQ + AZ in early high-risk outpatient disease. Five studies, including 2 controlled clinical trials, have demonstrated significant major outpatient treatment efficacy. HCQ + AZ has been used as the standard of care in more than 300,000 older adults with multiple comorbid conditions; the estimated proportion of such patients diagnosed with cardiac arrhythmia attributable to the medications is 47 per 100,000 users, among whom estimated mortality is less than 20% (9/100,000 users), as compared with the 10,000 Americans now dying each week. These medications need to be made widely available and promoted immediately for physicians to prescribe.

azithromycin; coronavirus; COVID-19; doxycycline; hydroxychloroquine; remdesivir; SARS-CoV-2; zinc

Abbreviations: AZ, azithromycin; COVID-19, coronavirus disease 2019; DOX, doxycycline; FAERS, FDA Adverse Event Reporting System; FDA, Food and Drug Administration; HCQ, hydroxychloroquine; NIH, National Institutes of Health; RCT, randomized controlled trial; RR, relative risk; SARS-CoV-2, severe acute respiratory syndrome coronavirus 2; TdP, torsades de pointes.

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Aside from the now more than 1.6 million Americans found through testing and public-health reporting to be infected with severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), seropositivity studies in California (1, 2), Colorado (3), and New York City and State (4) suggest that some 10- to 50-fold larger numbers of people carry antibodies to the virus. The workforce and effort required to carry out contact tracing on these tens of millions of Americans is not practical. While these studies have generated some media criticism, recent similar studies of blood donor samples in the Netherlands found 3% with SARS-CoV-2 antibodies (5), and 5% of household volunteers in Spain were found to have antibodies (6). Even allowing for some degree of false-positivity of these antibody tests, they still indicate that appreciably larger fractions of the population have been infected than have been characterized by identified reported cases. “Flattening the curve” by means of social distancing, mask-wearing, and staying at home serves to reduce hospital loads and spread them out over time, but to date it has pushed the infection reproduction number at time $t$ ($R_t$) down only to about 1.0 (7); thus, even
if maintained, over time very large numbers of people in the United States may eventually get the infection. The great majority of infected people are at low risk for progression or will manifest the infection asymptotically. For the rest, outpatient treatment that prevents disease progression and hospitalization is required. Exposures will occur as isolation policies are lifted and people begin to mix, even with various degrees of public isolation such as mask usage and physical separation still in place. Thus, the key to returning society toward normal functioning and to preventing huge loss of life, especially among older individuals, people with comorbidity, and African Americans and Hispanics/Latinos, is a safe, effective, and proactive outpatient treatment that prevents hospitalization in the first place.

All medical treatments have costs and benefits. In an ideal world, randomized double-blinded controlled clinical trials establish evidence for the relative degree of benefit, and if they are large enough, for estimates of the frequencies of adverse events. These trials take time to conduct: to get formal approval, to get funding, to enroll enough eligible patients, to wait for the outcomes to occur, and to analyze the data. In the context of the coronavirus disease 2019 (COVID-19) pandemic, we are presently averaging about 10,000 deaths per week in the United States, under moderately strong isolation policies that have put more than 36 million people out of work. Results of currently ongoing or planned randomized trials for use of a number of outpatient medications are many weeks or months off, and there are no guarantees that the results for these agents, even if statistically significant, will show sufficient magnitudes of effectiveness to be useful clinically. We are rapidly reaching a breaking point in the ability to maintain the status quo; states have begun the process of lifting their restrictions, and we thus need to evaluate what evidence we do have for promising outpatient treatments.

**REVIEW OF EVIDENCE**

Based on laboratory and other preliminary evidence available to date, 2 candidate medication regimens, among many others, have been widely discussed for outpatient treatment: remdesivir (Veklury; Gilead Sciences, Inc., Foster City, California), and hydroxychloroquine (HCQ) plus azithromycin (AZ). Remdesivir has been studied extensively in laboratory work and in animals (8) and for other viral diseases and has good biological properties, suggesting utility for the treatment of SARS-CoV-2 infection. In a study of compassionate use of remdesivir in 53 hospitalized patients with severe disease (9), 13% died, which appears lower than what might have been expected without treatment, though greater than the proportion of deaths in the placebo arm of the Adaptive COVID-19 Treatment Trial (more below). In a randomized, controlled but relatively underpowered trial carried out among nonventilated hospitalized patients with severe disease in China (10), investigators were not able to demonstrate benefit in either improvement or mortality in comparison with placebo. An appreciable fraction of the remdesivir patients left the trial early because of serious adverse events. The Adaptive COVID-19 Treatment Trial, a study of hospitalized patients with advanced lung disease, has released initial results (11) showing that patients on remdesivir had 31% faster recovery than patients on placebo (medians of 11 days and 15 days, respectively)—a difference which was statistically significant—but these results involved patients who did indeed survive. Mortality in the 2 groups (8.0% and 11.6%, respectively) was better for remdesivir but not significantly so (P = 0.059). More specific for consideration here, outpatient use of remdesivir has not been studied. The “secret” report issued by Scientists to Stop COVID-19 recommends widespread use of remdesivir and “as early in infection as possible” (12, p. 7), but no actual evidence as yet shows that it would be helpful for routine outpatient circumstances and disease in humans. The Food and Drug Administration (FDA) recently approved use of remdesivir in the current public-health emergency circumstances, but only for patients with “severe disease defined as SpO2 ≤ 94% on room air, requiring supplemental oxygen, mechanical ventilation, or extracorporeal membrane oxygenation (ECMO)” and “administered in an in-patient hospital setting via intravenous (IV) infusion by a healthcare provider” (13, p. 2). This approval seems specifically not to allow outpatient use. Symptomatic outpatient infection is a pathologically and clinically different disease than the life-threatening inpatient acute respiratory distress syndrome caused by SARS-CoV-2; thus, there is little reason to think that the same treatment would be useful for both (14). In any event, none of 20 currently registered trials is scheduled to provide data on outpatient use of remdesivir, and thus we may not know whether it could be used effectively to prevent hospitalization of symptomatic outpatients unless or until it is actually tried that way.

The other suggestion is the combined regimen of HCQ + AZ (or its variant, HCQ + doxycycline (DOX)). The FDA recently issued guidance to physicians and the general public (15) advising that the combination HCQ + AZ should not generally be used except by critically ill hospital inpatients or in the context of registered clinical trials. The National Institutes of Health (NIH) panel for COVID-19 treatment guidelines says essentially the same thing (16), and a similar statement has been released by major cardiology societies (17). Numerous reviews of HCQ efficacy and adverse events have been and continue to be published. To my knowledge, all of these reviews have omitted the 2 critical aspects of reasoning about these drugs: use of HCQ combined with AZ or with DOX and use in the outpatient setting. For example, a study carried out at Veterans’ Administration medical centers (18) examined treated hospitalized patients and was fatally flawed (19). The same point about outpatient use of the combined medications has been raised by a panel of distinguished French physicians (20) in petitioning their national government to allow outpatient use of HCQ + AZ. It appears that the FDA, NIH, and cardiology society positions have been based upon theoretical calculations about potential adverse events and from measured physiological changes rather than on current real-world mortality experience with these medications and that their positions should be revised. In reviewing all available evidence, I will show that HCQ + AZ and HCQ + DOX are generally safe for short-term use in the
early treatment of most symptomatic high-risk outpatients, where not contraindicated, and that they are effective in preventing hospitalization for the overwhelming majority of such patients. If these combined medications become the standard of care, they are likely to save an enormous number of lives that would otherwise be lost to this endemic disease.

What is the evidence for these assertions? Similar to remdesivir, 16 clinical trials of HCQ + AZ are listed in the ClinicalTrials.gov database (21). Of these, only 5 involve treating outpatients with the combined HCQ + AZ regimen (see Web Table 1, available at https://doi.org/10.1093/aje/kwaa093). For the earliest trial, between now and September, assuming a flat epidemic curve of 10,000 deaths per week, I estimate that approximately 180,000 more deaths will occur in the United States before the trial results are known. The Centers for Disease Control and Prevention has estimated substantially greater numbers of deaths (22).

In this context, we cannot afford the luxury of perfect knowledge and must evaluate, now and on an ongoing basis, the evidence for the benefit and risk of these medications (23). Available evidence of efficacy of HCQ + AZ has been repeatedly described in the media as “anecdotal,” but it most certainly is not. The evidence is not perfect either. Each piece of evidence, contained in each study, must be carefully considered and not dismissed, because in an ideal world such evidence would fall into a lower part of the evidence-quality triangle. Furthermore, and most critical to the correct understanding of what evidence is available, evidence for single agents cannot be extrapolated to apply to combined agents, evidence for one biochemical form of a drug cannot be extrapolated to another form, and even more importantly, evidence for utility or lack thereof or toxicity in hospitalized patients cannot be extrapolated to apply to outpatient use—outpatient use comprising the sole argument for application that I am making in this review.

Thus, for example, studies of chloroquine or HCQ used alone do not bear upon evidence for efficacy of HCQ + AZ or HCQ + DOX. This point has been argued forcefully by the French physicians (20). The first study of HCQ + AZ was controlled but not randomized or blinded and involved 42 patients in Marseilles, France (24). That study showed a 50-fold benefit of HCQ + AZ versus the standard of care (P = 0.0007). In the study, 6 patients progressed, stopped medication use, and left the trial before the day-6 planned outcome measure of swab-sampled nasopharyngeal viral clearance. Reanalysis of the raw study data elsewhere (25) and by myself shows that including these 6 patients does not much change the 50-fold benefit. What does change the magnitude of benefit is presentation with asymptomatic or upper respiratory-tract infection versus lower respiratory-tract infection—the latter cutting the efficacy in half (25-fold vs. standard of care). This shows that the sooner these medications are used, the better their effectiveness, as would be expected for viral early respiratory disease. The average start date of medication use in this study was day 4 of symptoms. The study has been criticized on various grounds that are not germane to the science, but the most salient criticism is the lack of randomization into the control and treatment groups. This is a valid general scientific criticism, but it does not represent epidemiologic experience in this instance. If the study had shown a 2-fold or perhaps 3-fold benefit, that magnitude of result could be postulated to have occurred because of subject-group differences from lack of randomization. However, the 25-fold or 50-fold benefit found in this study is not amenable to lack of randomization as the sole reason for such a huge magnitude of benefit. Further, the study showed a significant, 7-fold benefit of taking HCQ + AZ over HCQ alone (P = 0.035), which cannot be explained by differential characteristics of the controls, since it compares one treatment group with the other, and the treated subjects who received AZ had more progressed pneumonia than the treated subjects receiving HCQ alone, which should otherwise have led to worse outcomes. The study has also been described as “small,” but that criticism only applies to studies not finding statistical significance. Once a result has exceeded a plausible chance finding, greater statistical significance does not contribute to evidence for causation (26). No different conclusion would have resulted had a study with 1,000 patients found the same 50-fold benefit but with a P value of 10−10. Study-size limitation only applies to studies having findings within the play of chance. That is not the case here.

A second study by the Marseilles group involved 1,061 patients who tested positive for SARS-CoV-2 and were treated with HCQ + AZ for at least 3 days and followed for at least 9 days (27). The authors stated that no cardiac toxicity was observed (27). A good clinical outcome and virological cure were seen in 973 patients (92%). Five patients died, and the remainder were in various stages of recovery.

The third piece of evidence involves a cohort of 1,450 patients treated by Dr. Vladimir Zelenko of Monroe, New York. Dr. Zelenko has released a 2-page report (28) describing his clinical reasoning and procedures, dosing conditions and regimen, and patient results through April 28. Symptomatic patients presenting to Dr. Zelenko were treated with 5 days of HCQ + AZ + zinc sulfate if they were considered high-risk, as evidenced by 1 or more of the following: age 60 years or older; high-risk comorbidity; body mass index (weight (kg)/height2 (m2)) ≥30; and mild shortness of breath at presentation. Patients were considered to have COVID-19 on the basis of clinical grounds and started treatment as soon as possible following symptom onset, rather than waiting for test results before starting treatment. Of the 1,450 patients, 1,045 were classified as low-risk and sent home to recuperate without active medications. No deaths or hospitalizations occurred among them. Of the remaining 405 who were treated with the combined regimen, 6 were ultimately hospitalized and 2 died. No cardiac arrhythmias were noted in these 405 patients.

The fourth relevant study was a controlled nonrandomized trial of HCQ + AZ in 636 symptomatic high-risk outpatients in São Paulo, Brazil (29). All consecutive patients were informed about the utility and safety profile of the medications and offered the treatment, and those who declined (n = 224) comprised the control group. Patients were monitored daily by telemedicine. The study outcome was a need for hospitalization, defined as a clinically worsening condition or significant shortness of breath (blood oxygen saturation <90%). Even though the severities of all of the
recorded influenza-like signs and symptoms and of important comorbid conditions (diabetes, hypertension, asthma, stroke) were substantially greater in the treated patients than in the controls, the need for hospitalization was significantly lower: 1.2% in patients starting treatment before day 7 of symptoms, 3.2% for patients starting treatment after day 7, and 5.4% for controls ($P < 0.0001$). No cardiac arrhythmias were reported in the 412 treated patients. The most common side effect of treatment was diarrhea (16.5%), but 12.9% of treated patients presented with diarrhea before treatment began.

Finally, a small study in a long-term-care facility in Long Island, New York, is ongoing (30). This study has been employing HCQ + DOX rather than HCQ + AZ for treatment of high-risk COVID-19 patients. DOX itself has antiviral activity against SARS-CoV-2 at in vitro concentrations of 5.6 μM (median) (31). Among the first 54 residents treated in the Long Island study, 6 were hospitalized and 3 (5.6%) died (30). An unofficial update of these data indicates that of about 200 high-risk patients treated with HCQ + DOX, 9 (4.5%) have died.

The 2 nonrandomized but controlled trials provide important evidence, if not “proof,” of the major efficacy of early use of HCQ + AZ against SARS-CoV-2 infection in symptomatic high-risk outpatients. What can be said about the uncontrolled large case series of treated patients? Standard published case reports provide clinical evidence of the possibility of an exposure-outcome relationship, but not of the regularity, magnitude, or representativeness of such a relationship. The same can be said of case-series reports, meaning that subject entry into the series is not necessarily well-defined and no denominator information is provided from which to gauge what the series represents. However, a large series in the context of known risks of mortality or adverse events can allow for ballpark estimates of the denominator and thus provide a reasonable frame of reference for whether the outcomes are likely to represent beneficial or harmful results. For example, among Connecticut cases aged 60 years or older, mortality at present is 20% (32). Thus, it would be ballpark to estimate that some 20% of the 1,466 treated high-risk patients in the Zelenko and Marseilles cohorts would have died without outpatient HCQ + AZ treatment—293 patients, as compared with the 7 who did die. An alternative is to use the 12%–13% mortality of hospitalized patients in the placebo arms of the remdesivir trials (10, 11). This would give about 180 expected deaths.

ADVERSE EVENTS

Both proposed drug regimens have shown side effects. Remdesivir, in its phase 3 trial of 10-day versus 5-day therapeutic courses in hospitalized patients, produced a range of adverse events in more than 70% of patients in both treatment arms (33). Adverse events requiring medication discontinuation were many fewer: 5% in the 5-day group and 10% in the 10-day group. In the Chinese trial, 12% of remdesivir patients stopped using the medication before the end of the 10-day treatment because of drug-related adverse events (10).

For HCQ + AZ use, the argued issue concerns fatal cardiac arrhythmia: the warnings issued by the FDA, the NIH, and the cardiology societies. Indeed, both HCQ and AZ produce QT prolongation, rare instances of fatal tordes de pointes (TdP) ventricular tachycardia, and long QT-interval syndrome. A number of essays by cardiologists published in the Journal of the American Medical Association and other journals have anxiously warned about these risks but have not examined mortality from them. The sole question is whether these fatal events, or even any fatal cardiac arrhythmia events, would occur with enough frequency that general treatment of noncontraindicated high-risk outpatients with HCQ + AZ would outweigh benefit in preventing hospitalization and mortality. A number of studies have examined hospital inpatient use, but these studies have had major flaws, not least of which is that patients hospitalized with multiple medical problems and more advanced disease do not represent the mortality experience for outpatient use of these medications in patients otherwise well enough not to be hospitalized. One source of data on mortality associated with these medications is the FDA Adverse Event Reporting System (FAERS) database (34). Examination of the database for adverse events reported from the creation of the database in 1968 through 2019 and into the beginning of 2020 shows 1,064 adverse event reports for HCQ, including 200 deaths for all of the cardiac causes that could be both specifically and broadly classified as rhythm-related. Of these, 57 events, including 10 deaths, were attributed to TdP tachycardia and long QT-interval syndrome combined. This concerns the entirety of HCQ use over more than 50 years of data—probably millions of uses and long-term use than the 5 days recommended for COVID-19 treatment. For AZ use, the numbers of reported TdP and long QT-interval syndrome events total 37, of which 2 were deaths. FAERS data are generated by patient, physician, and pharmacist report initiation and probably underrepresent true event occurrences. However, even if the true numbers were 10-fold larger, they would still be minuscule compared with the amounts of medication usage. How much the risk of QT prolongation would be enhanced with HCQ and AZ taken together is unknown, but the FAERS data are generated by patient, physician, and pharmacist report initiation and probably underrepresent true event occurrences. However, even if the true numbers were 10-fold larger, they would still be minuscule compared with the amounts of medication usage. How much the risk of QT prolongation would be enhanced with HCQ and AZ taken together is unknown, but the FAERS database indicates that the major efficacy of early use of HCQ + AZ against SARS-CoV-2 infection in symptomatic high-risk outpatients is 20% (32). Thus, it would be ballpark to estimate that some 20% of the 1,466 treated high-risk patients in the Zelenko and Marseilles cohorts would have died without outpatient HCQ + AZ treatment—293 patients, as compared with the 7 who died. An alternative is to use the 12%–13% mortality of hospitalized patients in the placebo arms of the remdesivir trials (10, 11). This would give about 180 expected deaths.

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these medications and had this shortening, a total of 1 case of TdP tachycardia occurred, and it was not fatal—there were no deaths. Substantial fractions of these hospitalized patients were taking diuretics, which may be contraindicated for HCQ + AZ use in the first place. This arrhythmia issue is a real, physiologically measurable effect of the use of these combined medications, but fatal arrhythmia outcomes are so rare that they are of much lesser clinical significance than the hospitalization and mortality that the drugs prevent. This fact is also clear from the lack of any cardiac arrhythmia events or arrhythmia mortality noted in the 405 Zelenko patients or the 1,061 Marseilles patients or the 412 Brazil patients. Patients were not enrolled in these studies if they had known histories of QTc prolongation. History of cardiac arrhythmia or other possible contraindications for use of HCQ or AZ or DOX is a normal part of workup and clinical judgment in a physician’s choice to use these medications and in how to monitor the patients (see Web Appendix).

Further evidence of the real-world unimportance of arrhythmia and other cardiovascular adverse event endpoints of HCQ + AZ use is given in the large Oxford, United Kingdom-based record-linkage study (41). Fourteen large medical-records databases were examined for all-cause mortality and for 15 specified classes of adverse events among hundreds of thousands of patients with rheumatoid arthritis who had used these drugs. First, 323,122 users of HCQ + AZ were compared with 351,956 users of HCQ + amoxicillin. No significant difference in all-cause mortality was seen: As reported by the authors, the relative risk was 1.36 ($P = 0.10$), and as I calculate from the data provided by the authors in their Web supplement to the paper (41), the relative risk was 1.18 ($P = 0.37$)—either way, a null association within the range of chance. However, the authors selectivity presented from among the 15 analyzed endpoints the 3 most significant associations: those for cardiovascular mortality (relative risk (RR) = 2.19, $P = 0.0088$), chest pain/angina (RR = 1.15, $P = 0.0027$), and heart failure (RR = 1.22, $P = 0.027$). What is misrepresented in the authors’ presentation of these data in this way is that these 3 outcomes were not individually specified to be of more interest than any of the other 12 specific outcomes that they examined, and they did not correct their calculated levels of statistical significance for the 15 classes of outcomes. In lay terms, it was a fishing expedition. When accounting is done by means of the standard Bonferroni correction for multiple comparisons, the respective $P$ values are 0.12, 0.04, and 0.35. The large amount of data in this study thus shows that there is no significant relationship of HCQ + AZ use versus HCQ + amoxicillin use with any of the 15 outcomes specified or with all-cause mortality, except for a just-barely-significant association with chest pain/angina (a 15% higher risk). Even if the chest pain/angina association were a true finding, it would still be of little clinical import for a relatively infrequent outcome in the context of the mortality to be prevented by HCQ + AZ use in widespread symptomatic high-risk outpatient COVID-19 treatment.

Second, the stated concern of the FDA and NIH advisers and the cardiology society opinion restricting use of HCQ + AZ was for fatal TdP tachycardia and long QT-interval syndrome, 2 rare types of cardiac arrhythmia, as well as for cardiac arrhythmia in general. In the Oxford study, Lane et al. (41) examined cardiac arrhythmia outcomes and obtained, for their random-effects meta-analysis result, a relative risk of 1.08 ($P = 0.36$) for HCQ + AZ use versus HCQ + amoxicillin use. The fixed-effects meta-analysis relative risk was 1.04 ($P = 0.41$). This study clearly demonstrates that cardiac arrhythmia adverse events are not appreciably increased by combining HCQ with AZ. In the same study, the investigators compared HCQ use with sulfasalazine use and again found no difference in cardiac arrhythmia risk: For HCQ, there was a slightly lower relative risk of 0.89 ($P = 0.13$). The subjects analyzed in the Oxford study were largely older adults with multiple comorbid conditions in addition to rheumatoid arthritis.

Finally, the Oxford study allows for a direct estimate of the number of arrhythmia events attributable to HCQ + AZ use (41). Among 306,106 people taking sulfasalazine (which is known not to produce QT prolongation), 877 with cardiac arrhythmia were identified (0.287%). In 320,589 people taking HCQ + AZ, 1,068 had arrhythmia (0.333%). The difference, 0.047% or 47 per 100,000 older multimorbidity patients taking HCQ + AZ, is attributable to the HCQ + AZ use. These are events, not fatalities. As I noted above, fatalities according to FAERS comprise less than 20% of HCQ-related arrhythmia events. The maintenance HCQ dose in the Oxford study patients, 200 mg/day, gives as large or larger plasma drug levels as 5 days of HCQ at 400 mg/day, the recommended dose for outpatient COVID-19. These very small numbers of arrhythmias, as well as the null results in this very large empirical study, should therefore put to rest the anxieties about population excess mortality from HCQ + AZ outpatient use, either from cardiac arrhythmia or from all causes.

This discussion shows that the FDA, NIH, and cardiology society warnings about cardiac arrhythmia adverse events, while appropriate for theoretical and physiological considerations about use of these medications, are not borne out in mortality in real-world usage of them. Treatment-failure mortality will be much higher, but even that pales in comparison with the lives saved. It would therefore be incumbent upon all 3 organizations to reevaluate their positions as soon as possible. It is unclear why the FDA, NIH, and cardiology societies made their recommendations about HCQ + AZ use now, when the Oxford study (41) analyzed 323,122 users of HCQ + AZ compared with 351,956 users of HCQ + amoxicillin—that is, that the combination of HCQ + AZ has been in widespread standard-of-care use in the United States and elsewhere for decades, with use being comparable to HCQ + amoxicillin as if it just involved an alternate antibiotic choice, this use being predominantly in older adults with multiple comorbid conditions, and with no such strident warnings about the use given during that time. I note that since DOX is believed to cause even fewer cardiac arrhythmias than AZ, in patients where that is a concern (42), the evidence from long-term-care facilities suggests that HCQ + DOX likely will work about as well.
DISCUSSION

Given that a detailed and dispassionate review of all of the available relevant evidence leads to conclusions about outpatient HCQ + AZ use that are different from those of the FDA and NIH panels (which comprise wider expertise than the cardiology societies), I address how different underlying scientific worldviews might be involved. This is particularly reflected in the Scientists to Stop COVID-19 position about remdesivir use “as early as possible” (12, p. 5)—that is, early outpatient use being implied. All but 1 of the scientists on the Scientists to Stop COVID-19 panel are laboratory or clinical scientists; only 1 is an epidemiologist. Their recommendation for remdesivir use as early as possible was made without either FDA approval or randomized controlled trial (RCT) evidence of efficacy in the outpatient context. This recommendation therefore appears to be an extrapolation from animal and laboratory data and from use in severely ill hospitalized patients. However, a history of epidemiology shows numerous instances of failed extrapolation from animals to humans. As Bracken noted, “Animal research on almost any topic of epidemiologic interest is so heterogeneous and inadequately synthesized that it is possible to selectively assemble a body of evidence from the animal and in-vitro studies that support almost any epidemiologic result” (43, p. 221). For example, some carcinogens have been affirmed in animal studies but carcinogenicity has not been shown in human studies (acrylamide, alar, cyclamate, red dye #2, saccharin) (43). This is partly why the FDA has an approval system of phased RCTs leading to safety and efficacy of use in humans, in the specific contexts in which the drug is intended. It is not a question of off-label use, but of who are the patients for whom to use the medication. For COVID-19, inpatient acute respiratory distress syndrome is typically a florid immune-system overreaction, whereas initial outpatient illness is a viral multiplication problem involving the beginnings of immune response. These are different diseases. Thus, how well remdesivir might perform in outpatients will not be known until it is tried in typical outpatient circumstances, whether in RCTs or in any other unbiased systematic study of such use. Further, to the degree that remdesivir is similar in temporal characteristics to an antiviral agent like oseltamivir, it would be used in general societal contexts where patients must first recognize that they might have symptoms of the disease (and not something else) and go to their physicians or clinics for care and either be rapidly tested as positive with an assay that has negligible false-negative results or be symptomatic enough for the disease to be clinically distinguished and diagnosed, but definably positive in this way not more than 2 days after symptoms start. This is a very narrow temporal window in which to be definitive and to obtain full antiviral effectiveness, and it could be difficult to achieve in general in the mass-treatment circumstances that we are facing. So regardless of the strength of the implied evidence of outpatient efficacy when given shortly after the start of symptoms, remdesivir efficacy might be substantially lower in the context of actual population outpatient usage. This is another reason why empirical studies of medication use in the full context of application are needed.

The extrapolation from laboratory theory to empirical use also seems to underlie resistance to the idea that combined HCQ regimens could work for early outpatient use. HCQ is known to interfere with toll-like receptor signaling, reducing dendritic cell activation and immune response. This would seem to be counterproductive for suppressing SARS-CoV-2 multiplication in early treatment. Again, in extrapolation from physiological theory to human data, the epidemiologic data are definitive. The fact that epidemiologic data to date show strong evidence for efficacy of combined HCQ + AZ in early outpatient treatment, even if there is not “proof” yet at the level of several successful RCTs, is evidence that this medication regimen works in that context. The clash in scientific worldviews is that basic and clinical scientists seem to feel that biological and drug-development evidence for medication use in nonhuman and nonoutpatient contexts can be extrapolated to recommendations for outpatient use without benefit of RCT evidence but do not accept epidemiologic evidence without RCTs, whereas epidemiologists have had career experience with laboratory and animal evidence that did not hold up under epidemiologic study but do reason by including all types of epidemiologic study designs and derive causal conclusions in the standard way following Hill’s aspects (26) on the basis of strong totality of evidence, sometimes even without RCT evidence. There are contexts in which each approach is valid. However, it is not my point to say that remdesivir has little evidence to support its potential outpatient utility, only efficacy considerations that have not been addressed and that could lead to lack of efficacy under general use, but that HCQ + AZ has been directly studied in actual early high-risk outpatient use with all of its temporal considerations and found empirically to have sufficient epidemiologic evidence for its effective and safe employment that way, and that requiring delay of such general use until availability of additional RCT evidence is untenable because of the ongoing and projected continuing mortality. No studies of COVID-19 outpatient HCQ + AZ use have shown higher mortality with such use than without, cardiac arrhythmias included; thus, there is no empirical downside to use of this combined medication.

Some of my medical colleagues still prefer to wait until more studies are done and stronger evidence such as that from RCTs becomes available and government and professional advisory panels reevaluate the evidence. I strongly urge these panels to reconsider the data and arguments discussed above. Substantial fractions of physicians treating COVID-19 patients in Europe and elsewhere report use of HCQ + AZ: 72% in Spain, 49% in Italy, 41% in Brazil, 39% in Mexico, 28% in France, 23% in the United States, 17% in Germany, 16% in Canada, and 13% in the United Kingdom (44)—much of the non-US use in outpatients. HCQ + AZ has been standard-of-care treatment at the 4 New York University hospitals, where a recent study showed that adding zinc sulfate to this regimen significantly cut both intubation and mortality risks by almost half (45). The French physicians are insistent that with careful clinical judgment and supervision, these medications are safe and should be used as early as possible for outpatients, and they provide a detailed clinical guide to their use (20). Until we have quantitative evidence for the utility and safety of other
medications for preventing hospitalization and mortality in high-risk COVID-19 outpatients, the urgency of current mass mortality requires an immediate application of the best treatment that we have available, even if knowledge is imperfect and even if efficacy is yet unproven to the standards of double-blinded RCTs. This problem will get even worse as US states and cities yield to the acute pressure at this moment to begin lifting stay-at-home restrictions and even more people become infected. Some people will have contraindications and will need other agents for treatment or will need to remain in isolation. However, I conclude that for the great majority, HCQ + AZ and HCQ + DOX, preferably with zinc (46), can be this outpatient treatment, at least until we find or add something better, whether that be remdesivir or something else.

It is our ethical and professional obligation not to just stand by “carefully watching” as the old, the infirm, and inner-city residents are killed by this disease and our economy is destroyed by it and offer nothing except high-mortality hospital treatment. We have a solution, albeit imperfect, to attempt to deal with this disease. We have to let physicians employing good clinical judgment use it and informed patients choose it. There is a small chance it may not work, but the urgency of the circumstances demands that we at least start to take that risk and evaluate what happens. If the situation does not improve, we can stop it, but we will know that we did everything we could instead of sitting by and letting hundreds of thousands of people die because we did not have the courage to act according to rational calculations.

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